

بسم الله الرحمن الرحيم

## **Republic of Sudan**



## Ministry of Higher Education and Scientific Research University of Shendi

## Faculty of Graduate Studies and Scientific Research

## Association of Anti *Helicobacter pylori* IgG Antibodies in Typhoid Patients in Atbara Teaching Hospital

A thesis submitted in partial fulfillment of Master degree in Medical Laboratory Sciences

(Medical Microbiology)

By

Dalal Mohamed Osman Fadol

B.Sc. Medical Laboratory Science, Shendi University

(2006)

Supervisor:

Dr. Hadia Abass Eltaib (PhD)

Assistant professor of Molecular Biology -Faculty of Medical Laboratory Sciences - Shendi University

August-2018

الآية

هال تعالى:

حدق الله العظيم

البغرة: آيـــة 269



## Dedication

Happily, I would like to dedicate this simple attempt to:

My parents

Who learn me the first alphabetic

To my mother

Spring of love

To my sisters

To my teachers whom educate me letters

To my friends

## Acknowledgement

My great full thanks to ALLAH who gave me health and power to finish this work.

I would like to thanks my supervisor

#### Dr. Hadia Abass Eltaib

for this stimulating suggestion, help, knowledge, experience and encouragement helped me in all time of study.

My teachers in Shendi University.

My friends **Ahmed Wedaa**, **Abd Elbasit Mohammed**, and all friends help me to complete my research.

I would to express my deeply thanks to all persons contributed with me and help me to complete this piece of work.

## Abstract

#### **Back ground:**

Infections with H.pylori remain the most common bacterial infection that will increase the susceptibility to other gastrointestinal infections and typhoid fever.

#### **Objectives:**

The aim of study was to evaluate the prevalence of *H.pylori* among typhoid patients at Atbara teaching hospital, and also to identify the risk factors related to infection.

#### Method:

A cross sectional study was conducted in Atbara city, during the period from May to July 2018. Using ELISA technique for serodetection of anti-*Helicobacter pylori* IgG.

#### **Result:**

Anti- *Helicobacter pylori* IgG was determined in sera of 72 cases, 59 (81.9%) out of them were positive.

There was significant association between present of *H.pylori* and typhoid fever in related to age, resident, tribe, smokers, and coffee drinkers as risk factors with (P.value=0.000, 0.006,0.000,001,0.02) respectively.

#### **Concolusion:**

Strong association between H.pylori and typhoid fever was detected among study population which increased with age.

#### Key wards:

H.pylori, Enteric fever, IgG, Risk factors, Atbara.

### ملخص البحث

#### خلفيه:

تعد الإصابة بالجرثومة الحلزونية البوابية الأكثر شيوعاً والتي تزيد مـــن احتماليـــة الإصـــابة بالأمراض الأخرى المتعلقة بالجهاز الهضمي وكذلك مرض الحمى التايفية. **الأهداف:** 

كان الهدف من هذه الدراسة هو تقييم انتشار الجرثومة الحلزونية البوابية في مرضي الحمـــى التايفية بمستشفى عطبرة التعليمي, وكذلك تحديد عوامل الخطر.

#### الطريقة:

دراسة مقطعيه أجريت في مدينة عطبرة في الفترة من مايو ٢٠١٨ حتى يونيو ٢٠١٨م. بواسطة تقنية الاليزا تم الكشف عن وجود الجسم المضاد من نوع IgG للجرثومة الحلزونية البوابية.

#### النتائج:

هنالك ترافق ما بين الجرئومة الحلزونية البوابية والحمى التايفية في مصل ٧٢ من الأشخاص تحت الدراسة (٥٩ من ٧٢) ٩٨٩ كانت نتائجهم ايجابية للفحص. كما أن الانتشار المصلي يتأثر بعامل العمر والسكن والقبيلة كما أن هناك فروقات ذات دلاله إحصائية بين الانتشار المصلي وعوامل الخطر كالتدخين وشرب القهوة حيث كانت القيم الاحتمالية لهم ( ٠٠,٠٠، و ٢٠,٠٠، ، ٠٠,٠٠، , ٠,٠٠٠) على التوالي.

#### الخلاصة:

هناك علاقة قوية بين الإصابة بين الجرثومة الحلزونية البوابية والحمى التايفية تزيد مع العمر.

#### الكلمات المفتاحية:

الجرثومة الحلزونية البوابية، الحمى التايفية، الجسم المضاد IgG، عوامل الخطر,عطبرة.

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## List of abbreviations

Term	Meaning
AIDS	Acquired Immune Deficient Syndrome
A <sub>w</sub>	Water activity
CO <sub>2</sub>	carbon dioxide
DNA	Deoxyribo-nucleic acid
ELISA	Enzyme Linked Immune Assay
g/dl	Gram per deciliter
H.pylori	Helicobacter pylori
HIV	Human Immune deficiency Virus
HLA	Human leukocyte Antigen
IgG	Immuno globulin G
mμ	Micrometer
Mg/dl	Milligram per deciliter
Мр	Macrophage
Nacl	Sodium chloride
NANO <sub>2</sub>	Sodium nitrite
Ph	Positive hydrogen
SPSS	Statistical Package for Social Science
Vi	Virulence
WHO	World Health Organization

## **Chapter One**

Introduction

Rationale

**Objectives** 

#### **1-1 Introduction**

Helicobacter pylori is spiral- shaped bacteriam that inftects well over 30% of the worlds populations. in some countries it infects more than 50% of population. This is, there fore, one of the most common bacterial infections known to mankind between 1979-1982. Australian pathologist, robin warren and Astralian gastroenterologist barrymarshal, identified H.pylori and suggested a link to the development of stomach ulcers since this discovery, the world health organization has declared the bacteria to be a class I carcinogen. It invades the mucosal lining of the stomach and cause of up to 95% of duodenal and up to 75% gastric ulcers and has been. Associated with gastric cancer and lymphoma(Blaster., 1999). Dispite intense investigation to the spread of H.pylori, the precise mode of transmission remain unclear. oral to oral or faecal to oral are the most likely routes at this stage most infection occur in childhood crowded living condition poor sanitation poor personal hygine and poor water supply correlate with higher rates of infection (which can approach 80% of the population in developing world (Harford et al.2000). H.pylori infects both genders eqally. The presence of H.pylori in the stomach induces a chronic, active, inflammation in almost everyone infected. majority of people with H. pylori ,however ,are asymptomatic fewer than10% of individuals colonies with H. pylori develop peptic ulcer disease gastric cancer or mucosa associated lymph tissue lymphoma (Harford et al., 2000).

Typhoid fever a serious systemic illness that each year affects over 20 million people, predominantly in developing countries (Crump et al., 2004). Infection with *Salmonella typhi* is transmitted by the faecal–oral route and in several epidemiological studies risk factors were identified that suggested either waterborne transmission or foodborne transmission (Luby *et al.*, 1998). The determination of the relative contribution of distinct environmental risk factors for transmission of disease is essential to focus local control strategies. Also host-related risk factors for infection have been examined, identifying both genetic factors (Dunstan *et al.*, 2001) as well as concurrent *Helicobacter pylori* 

infection, that was interpreted as a cause of a reduced gastric acid barrier (Bhan et al., 2002). A high incidence of salmonellosis has been observed in individuals with surgically induced or other types of achlorhydria (pernicious anaemia and chronic atrophic gastritis) according to Kunz and Waddell (1956). Also H. pylori infection may exert an effect on the secretion of gastric acid. Approximately 50% of the world's population is infected with *H. pylori* (Torres et al., 2000), and even higher prevalences have been reported in developing countries (Bardhan., 1997), where acquisition occurs at a younger age than in the developed world (Blaser., 1999). Active infection with *H.pylori* is associated with a transient hypochlorhydria that may be present for several months (Harford et al., 2000). Furthermore, H. pylori-induced chronic gastritis of the body of the stomach reduces acid secretion and persistent hypochlorhydria constitutes a risk for the development of gastric cancer. In the absence of the acid-mediated inhibition of gastric gastrin release, the serum gastrin concentration increases. In contrast, antral-predominant, body-sparing gastritis due to H.pylori increases gastric acid secretion, resulting in duodenal ulcer disease (El Omar et al., 2000). Consequently, the association between H. pylori infection as an indicator of hypochlorhydria and the susceptibility to other gastrointestinal infections is ambiguous. An increased susceptibility to enteric infections in *H. pylori*-infected individuals, as measured by anti-*H. pylori* IgG response, was documented for cholera (Shahinian et al., 2000) and typhoid fever (Bhan et al., 2002). However, the evidence for the association of H. pylori infection and diarrhoea is conflicting (Sullivan et al., 1990) and even a protective effect of *H. pylori* infection was demonstrated (Rothenbacher *et al.*, 2000).

### 1-2 Rationale

Noticed association of *H.pylori* and typhoid fever during duty work in Atbara hospital.and to detect the presence of correlation between them as the result of previous study.

A little information about seroprevalence of *H.pylori* in Atbara city and its surrounding villages.

The usages of tankers for saving water for a long time in surrounding villages which may be a source of infection with both *H.pylori* and typhoid fever.

### **1-3 Objectives**

#### 1-3-1General objective:

To detect association between anti *H.pylori* antibodies and typhoid fever patients in Atbara teaching hospital.

#### 1-3-2 Specific objectives:

- To determine occurrence of IgG antibodies against *H.pylori* in patient with typhoid fever.
- To identify major risk factors associated with *Helicobacter pylori* (gender, age, smoking, coffee consumption).
- To correlate between seroprevalence of *H.pylori* and typhoid fever .

## **Chapter Tow**

Literature review

#### 2- Literature review

#### 2-1 Helicobacter pylori:

This organism was earlier known as *Campylobacter pylori* and believed to be the casual agent of chronic gastritis. Strong evidence has accumulated regarding its close association with gastric and duodenal ulcers. It is also being incriminated as responsible for initiating the process of metaplasia in gastric epithelium which ultimately may lead to carcinoma of stomach (Rajesh & Rattan, 2004).

#### 2-1-1 Morphology of Helicobacter pylori:

In stomach this organism takes form of short spirals or S shaped gram negative bacterium which is about  $3\mu$ m long, 0.5-1.0 $\mu$ m wide with a wavelength of about 2.5 $\mu$ m. after growth on laboratory media *Helicobacter pylori* is visible under the microscope as curved gram negative (Rajesh & Rattan, 2004).

#### 2-1-2 Physiology and growth conditions:

The physiological characteristics of *H.pylori* have received relatively little attention. It is a Gram negative spiral-shaped bacteria, although its morphology is not constant. Under adverse conditions it becomes coccoid, but there is controversy about the nature of the coccoid form. Some researchers have stated that this form is either a contaminant or a dead bacterium (Kusters *et al.*,1996), but others consider it to be a metabolically active form that cannot be cultured in vitro (Bode *et al.*,1993; Nilius *et al.*,1993). It has also been suggested that some cocci can revert to their original spiral shape (Andersen *et al.*,1997). *H.pylori* is microaerophilic; optimal growth occurs in the presence of 5–15% oxygen (Goodwin,1989). Incubation in air results in reduced survival(West *et al.*,1992) and it grows poorly under anaerobic conditions (Goodwin and Armstrong,1990). The presence of 5% CO2 seems to provide optimal conditions, while 10% CO2 led to a loss in cultivability in one study (Donelli *et al.*,1998).

#### 2-1-2-1Carbon source:

Glucose is not necessary for growth (Albertson *et al.*,1998; Reynolds & Penn,1994). Cell yield is not influenced by the presence of glucose, pyruvate, succinate, or citrate, but survival is enhanced by their presence. Prolonged incubation with carbon sources improves the viability of the organism (Albertson *et al.*,1998). *H. pylori* depends on the presence of various amino acids for growth, including arginine, histidine, isoleucine, leucine, methionine, phenylalanine, and valine. Some strains also need alanine,serine, proline, and tryptophan (Reynolds & Penn,1994).

#### 2-1-2-2 PH and water activity:

*H.pylori* can be cultured in environments within a pH range of 4.5–9. And at low pH values (e.g. 3.5), the addition of urea increases survival. NaNO<sub>2</sub> has no effect if concentrations range from 0 mg/ml to 400 mg/ml; growth is not possible at NaCl concentrations of 52.5 g/l. The pathogen is sensitive to environments with a low water activity ( $A_w$ ): growth is inhibited at values <0.98. In one study, *Helicobacter pylori* concentrations became undetectable in nutrient-rich laboratory medium within three days when the  $A_W$  was 0.96 (Jiang & Doyle,1998).

#### 2-1-2-3 Temprature:

*H. pylori* only grows at temperatures of 30–37°C. All the required growth conditions are met in the gastrointestinal tract of all warm-blooded animals. At temperatures below 30°C, *H. pylori* could survive in some foods, such as fresh fruit and vegetables, fresh poultry or fish, fresh meats, and some dairy products (Banwart,1979). *H. pylori* survived at 30°C in laboratory media (Jiang & Doyle,1998), water(West *et al.*, 1992), and milk (Fan *et al.*,1998), and survived longer at lower temperatures (Jiang & Doyle, 1998).

#### 2-1-3 Reservoir:

The human stomach appears to be the environment most suitable for the organism's growth; there are no significant animal or environmental reservoirs for strains infecting humans. *H. pylori* has been isolated from domestic,

commercially reared cats (Handt et al., 1994; Fox, 1995) and it has been suggested that it might be a zoonotic pathogen with transmission occurring from cats to humans. However, there have been no data to support this hypothesis. For example, after adjusting for potential confounders in a study of 447 factory workers in the United Kingdom, there was no association between H. pylori seropositivity and cat ownership during childhood (Webb et al., 1996). In Ulm, Germany, in 1996–97 among schoolchildren in first grade, neither contact with pets in general nor contact with specific kinds of animals was positively associated with infection (Bode et al., 1998). The possibility that H. pylori might be a zoonotic pathogen transmitted from animals other than cats has also been considered (Dunn et al., 1997; Fox, 1995), but the organism has never been isolated from animals slaughtered for consumption, such as pigs (Fox, 1995). It has been isolated from some non-human primates, such as macaque monkeys. However, because contact between humans and other primates is rare, it is unlikely that these other animals play an important role in the transmission to humans. It is possible that the inability to isolate the organism from other animals may be due to the difficulty of detecting the bacterium in materials other than gastric tissue (Fox, 1995).

#### 2-1-4 Route of transmission:

1- Person to person transmission:

Several studies have assessed the relation between *H.pylori* infection and institutionalized populations. Significantly higher rates of *Helicobacter pylori* infection also were found in other institutionalized populations (Lambert *et al.*, 1995).

Familial exposures. Most of the selected studies that looked at the relation between *H.pylori* infection and intrafamilial clustering of *Helicobacter pylori* did not use DNA fingerprinting to confirm that relatives had the same strain of *H.pylori* (Mitchell *et al.*, 1993).

2- Fecal-oral route:

Also another possible method of *H.pylori* transmission is the fecal-oral route. *H. pylori* DNA has been detected in feces of infected subjects by some researchers (Namavar *et al.*, 1995; Shimada *et al.*, 1994; Gramley *et al.*, 1999) but not others (Van Zwet *et al.*, 1994). Recently, Gramley *et al.* found detectable *H. pylori* DNA in the feces of 73 percent of infected subjects. Isolation of *H.pylori* by fecal culture has been performed by a number of investigators from around the world. However, isolation of *H. pylori* from feces has been problematic for some researchers, especially for those unable to obtain fresh feces. Delay in processing could have resulted in the small number of *H. pylori* organisms present being overgrown by other fecal bacteria. Recently, Parsonnet et al. were able to culture *H.pylori* from cathartic-induced diarrheal stools in 7 of 14 *H. pylori* infected subjects but not from normal stools (Parsonnet *et al.*, 1999).

3-Vector borne or zoonotic transmission:

*H. pylori* has been isolated from nonhuman primates and domestic cat. (Handt *et al.*, 1994)

4- Iatrogenic transmission:

Risk factor for latrogenic transmission of *H.pylori* is endoscopy Because of the complex structure of the endoscope and difficulty in disinfecting it (Fantry *et al.*, 1995).

#### 2-1-5 diseases caused by Helicobacter pylori:

Most infected persons are never suffering from any symptoms related to infection, however, *H. pylori* causes chronic active, chronic persistent, and atrophic gastritis in adults and children. Infection with *H. pylori* also causes duodenal and gastric ulcers. Infected persons have a 2- to 6-fold increased risk of developing gastric cancer and mucosal associated- lymphoid-type lymphoma compared with their uninfected counterparts. The role of *H. pylori* in non-ulcer dyspepsia remains unclear.

Active infection with *H. pylori* is associated with a transient hypochlorhydria that may be present for several months (Harford *et al.*, 2000). Furthermore, *H.* 

*pylori*-induced chronic gastritis of the body of the stomach reduces acid secretion and persistent hypochlorhydria constitutes a risk for the development of gastric cancer (Suerbaum & Michetti, 2002; El Omar *et al.*, 2000).

#### 2-1-5-1 Symptoms of ulcer:

The most common ulcer symptom is gnawing or burning pain in the epigastrium. This pain typically occurs when the stomach is empty, between meals and in the early morning hours, but it can also occur at other times. It may last from minutes to hours and may be relieved by eating or by taking antacids. Less common ulcer symptoms include nausea, vomiting, and loss of appetite. Bleeding can also occur; prolonged bleeding may cause anemia leading to weakness and fatigue. If bleeding is heavy, hematemesis, hematochezia, or melena may occur (Center for disease Control and Prevention., 1998).

#### 2-1-5-2 Risk factor of *H.pylori* infection:

The major factors investigated for their possible association with *H. pylori* positivity are The following topics included: smoking, alcohol consumption, diet, occupational exposures, waterborne exposures, hygiene practices, density/crowding, social factors, and family history of gastric disease.

**Smoking:** Studies have assessed the possible association between *H. pylori* infection and smoking. Whereas some found that *H.* pylori seropositive subjects were overall more likely than seronegative subjects to be current smokers, results were often not consistent by race or gender (Lin *et al.*, 1998; Fontham *et al.*, 1995).

**Alcohol consumption:** None of several recent epidemiologic studies of the relation between alcohol consumption and *H. pylori* infection found a positive association, but many noted a nonstatistically significant reduction in risk, (Brenner *et al.*, 1997; Brenner *et al.*, 1999 (a,b)) who incorporated a quantitative measure of alcohol consumption while controlling for potential confounding factors, found a significant negative association with alcohol consumption, especially at moderate to high levels. In two of these studies, the association was stronger for wine than for beer.

Several studies did not adequately control for potential confounding variables or did not present the actual risk estimate or prevalence; thus, it is difficult to evaluate whether alcohol consumption has a "protective" effect on the prevalence of *H. pylori. H.pylori* is better able to survive in the acid environment of the stomach than other bacteria are because of its production of urease. Therefore, it is not surprising that the reduction in pH that may accompany alcohol consumption would have little effect on the prevalence of *H. pylori* (Jenkins, 1997). However, alcohol is known to have direct antimicrobial effects that appear to be more pronounced for wine than for other types of alcoholic beverages (Klontz, 1999). The differing results may be due to the different methodologies used or to real differences in either the type or amount of alcohol consumed and its effect on *H. pylori* in different populations.

**Diet:** Studies have also looked at dietary associations with *H. pylori*. Although the studies cover many different types of populations and include both adults and children, some consistent associations suggest that nutritional status may be related to *H. pylori* infection (Goodman *et al.*, 1997).

**Occupational exposures:** Occupational exposures have been studied by several researchers to determine whether people working in certain occupations with potentially greater exposure to *H. pylori* had an increased prevalence of infection.( Bohmer *et al.*, 1997; Friis *et al.*, 1996)

Waterborne exposures. Water has been suggested as a possible source of *H*. *pylori* infection (Goodman *et al.*, 1996).

**Hygiene practices:** Studies also have assessed the relation between H. pylori infection and various hygiene practice indicators in a number of countries. Overall, poor hygiene practices, especially during childhood, appear to be related to a higher seroprevalence of *H. pylori* (Goodman *et al.*, 1996).

Family history *of gastric disease*. Studies have also evaluated the relation between *H.pylori* infection and family history of gastric disease (Brenner *et al.*, 2000).

#### 2-1-6 Diagnosis of H.pylori:

Several methods may be used to diagnose *H. pylori* infection. Serological tests that measure specific *H. pylori* IgG antibodies can determine if a person has been infected. The sensitivity and specificity of these assays range from 80% to 95% depending upon the assay used.

Another diagnostic method is the breath test. In this test, the patient is given either 13C- or 1 4C-labeled urea to drink. *H. pylori* metabolizes the urea rapidly, and the labeled carbon is absorbed. This labeled carbon can be measured as CO2 in the patient's expired breath to determine whether *H. pylori* is present. The sensitivity and specificity of the breath test ranges from 94% to 98%. Upper esophagogastroduodenal endoscopy is considered the reference method of diagnosis. During endoscopy, biopsy specimens of the stomach and duodenum are obtained and the diagnosis of *H. pylori* can be made by several methods:

• The biopsy urease test - a colorimetric test based on the ability of *H. pylori* to produce urease; it provides rapid testing at the time of biopsy.

• Histologic identification of organisms - considered the gold standard of diagnostic tests.

• Culture of biopsy specimens for *H. pylori*, which requires an experienced laboratory and is necessary when antimicrobial susceptibility testing is desired (Center for Disease Control and Prevention., 1998).

#### 2-1-7Treatment of *H.pylori*:

*H.pylori* sensitive to penicillin, cephalosporins, tetracycline, erythromycin, rifampicin, aminoglycosides and nitrofurans. (Rajesh & Rattan, 2004).

#### 2-2 Typhoid fever:

Typhoid fever is caused by *Salmonella typhi*, a Gram-negative bacterium. A very similar but often less severe disease is caused by *Salmonella* serotype *paratyphi* A. The nomenclature for these bacteria is confused because the criteria for designating bacteria as individual species are not clear. Two main views on the nomenclature of the genus *Salmonella* have been discussed. Le Minor and Popoff suggested that two species should be recognized: *Salmonella* 

*bongori* and *Salmonella enterica*. *S. enterica* included six subspecies, of which subspecies I (one) contained all the pathogens of warm-blooded animals. *S. typhi* was a serotype within subspecies I: *Salmonella enterica* subspecies I serotype *typhi*. This proposal was rejected by the International Judicial Commission because the name was not well known to clinicians and its use might cause accidents endangering health or life. The original rules therefore remain in force. Ezaki and colleagues have noted in the International Journal of Systematic and Evolutionary Microbiology that the correct nomenclature for the causal agent of typhoid fever is *Salmonella typhi* and have requested that the current subspecific status of serotype *paratyphi* A should be raised to specific status, i.e. *Salmonella paratyphi* A (WHO, 2003).

*S. typhi* has several unique features, the genetic basis of many of which is known as a result of early genetic studies and the recent sequencing of the whole genome. Although many genes are shared with *E. coli* and at least 90% with *S. typhimurium*, there are several unique clusters of genes known as pathogenicity islands and many more single genes that seem to have been acquired by *S. typhi* during evolution. *S. typhi* can be identified in the laboratory by several biochemical and serological tests. One of the most specific is that of polysaccharide capsule Vi, which is present in about 90% of all freshly isolated *S. typhi* and has a protective effect against the bactericidal action of the serum of infected patients. This capsule provides the basis for one of the commercially available vaccines. Vi antigen is present in some other bacteria (*Citrobacter freundii*, *Salmonella paratyphi C* and *Salmonella dublin*) but not in exactly the same genetic context. The ratio of disease caused by *S. typhi* to that caused by *S. paratyphi* is about 10 to 1 in most of the countries where this matter has been studied (WHO., 2003).

#### 2-2-1 Pathogenisity:

During an acute infection, *S. typhi* multiplies in mononuclear phagocytic cells before being released into the bloodstream. After ingestion in food or water, typhoid organisms pass through the pylorus and reach the small intestine. They

rapidly penetrate the mucosal epithelium via either microfold cells or enterocytes and arrive in the lamina propria, where they rapidly elicit an influx of macrophages (Mp) that ingest the bacilli but do not generally kill them. Some bacilli remain within Mp of the small intestinal lymphoid tissue. Other typhoid bacilli are drained into mesenteric lymph nodes where there is further multiplication and ingestion by Mp. It is believed that typhoid bacilli reach the bloodstream principally by lymph drainage from mesenteric nodes, after which they enter the thoracic duct and then the general circulation. As a result of this silent primary bacteraemia the pathogen reaches an intracellular haven within 24 hours after ingestion throughout the organs of the reticuloendothelial system (spleen, liver, bone marrow, etc.), where it resides during the incubation period, usually of 8 to 14 days. The incubation period in a particular individual depends on the quantity of inoculum, i.e. it decreases as the quantity of inoculum increases, and on host factors. Incubation periods ranging from 3 days to more than 60 days have been reported. Clinical illness is accompanied by a fairly sustained but low level of secondary bacteraemia (~1-10 bacteria per ml of blood) (WHO.,2003).

#### 2-2-2 Symptoms:

The clinical presentation of typhoid fever varies from a mild illness with lowgrade fever, malaise, and slight dry cough to a severe clinical picture with abdominal discomfort and multiple complications. Many factors influence the severity and overall clinical outcome of the infection. They include the duration of illness before the initiation of appropriate therapy, the choice of antimicrobial treatment, age, the previous exposure or vaccination history, the virulence of the bacterial strain, the quantity of inoculums ingested, host factors (e.g. HLA type, AIDS or other immunosuppression) and whether the individual was taking other medications such as H2 blockers or antacids to diminish gastric acid. Patients who are infected with HIV are at significantly increased risk of clinical infection with *S. typhi* and *S. paratyphi* (Gotuzzo *et al.*, 1991). Evidence of *Helicobacter*  *pylori* infection also represents an increased risk of acquiring typhoid fever (WHO, 2003).

Acute non-complicated disease: Acute typhoid fever is characterized by prolonged fever, disturbances of bowel function (constipation in adults, diarrhoea in children), headache, malaise and anorexia. Bronchitic cough is common in the early stage of the illness. During the period of fever, up to 25% of patients show exanthem (rose spots), on the chest, abdomen and back (WHO, 2003).

**Complicated disease:** Acute typhoid fever may be severe. Depending on the clinical setting and the quality of available medical care, up to 10% of typhoid patients may develop serious complications. Since the gut-associated lymphoid tissue exhibits prominent pathology, the presence of occult blood is a common finding in the stool of 10-20% of patients, and up to 3% may have melena. Intestinal perforation has also been reported in up to 3% of hospitalized cases. Abdominal discomfort develops and increases. It is often restricted to the right lower quadrant but may be diffuse. The symptoms and signs of intestinal perforation and peritonitis sometimes follow, accompanied by a sudden rise in pulse rate, hypotension, marked abdominal tenderness, rebound tenderness and guarding, and subsequent abdominal rigidity. A rising white blood cell count with a left shift and free air on abdominal radiographs are usually seen (WHO, 2003).

Altered mental status in typhoid patients has been associated with a high casefatality rate. Such patients generally have delirium or obtundation, rarely with coma. Typhoid meningitis, encephalomyelitis, Guillain-Barré syndrome, cranial or peripheral neuritis, and psychotic symptoms, although rare, have been reported. Other serious complications documented with typhoid fever include haemorrhages (causing rapid death in some patients), hepatitis, myocarditis, pneumonia, disseminated intravascular coagulation, thrombocytopenia and haemolytic uraemic syndrome. In the pre-antibiotic era, which had a different clinical picture, if patients did not die with peritonitis or intestinal haemorrhage, 15% of typhoid fever cases died with prolonged persistent fever and diseases for no clear reason. Patients may also experience genitourinary tract manifestations or relapse, and/or a chronic carrier state may develop (WHO, 2003).

**Carrier state:** 1-5% of patients, depending on age, becomes chronic carriers harboring *S.typhi* in the gallbladder (Edelman & Levine Myron ,1986).

#### 2-2-3 Contamination and transmission:

Humans are the only natural host and reservoir. The infection is transmitted by ingestion of food or water contaminated with faeces. Ice cream is recognized as a significant risk factor for the transmission of typhoid fever. Shellfish taken from contaminated water, and raw fruit and vegetables fertilized with sewage, have been sources of past outbreaks.

The highest incidence occurs where water supplies serving large populations are contaminated with faeces. Epidemiological data suggest that waterborne transmission of *S. typhi* usually involves small inocula, whereas foodborne transmission is associated with large inocula and high attack rates over short periods. The inoculum size and the type of vehicle in which the organisms are ingested greatly influence both the attack rate and the incubation period. In volunteers who ingested 109 and 108 pathogenic *S. typhi* in 45 ml of skimmed milk, clinical illness appeared in 98% and 89% respectively. Doses of 105 caused typhoid fever in 28% to 55% of volunteers, whereas none of 14 persons who ingested 103 organisms developed clinical illness. Although it is widely believed that *Salmonella* is transmitted via the oral route, the transmission of *S. typhimurium* via the respiratory route has been demonstrated in a mouse model (Ivanoff *et al.*, 1980).

#### 2-2-4 Diagnosis of typhoid fever:

Laboratory diagnosis of typhoid fever depends upon following parameter:

- Isolation of causative agent.
- Detection of microbial antigen.
- Titration of antibody against causative agent.

#### 2-2-4-1Cultural method:

The method of choice for isolation of causative agents of typhoid fever is blood culture.

- Blood culture: with all aseptic precautions, about 10 ml of blood should withdrawn. This large quantity of blood is required because in many cases number of bacteria in blood is too little (just one bacteria per ml). as far as possible sample should be collected prior the administration of any antibacterial therapeutic agent to patient.
- Bone marrow culture: it may give positive result when blood culture fail, particularly in patients admitted to hospital after prolonged antibiotic therapy.
- Stool culture: a spoonful of feaces should be collected in clean container. Numerous selective media are available.
- Urine culture: it can cultured either as such or the deposit obtained after centrifugation, duodenal fluid, and bile can be processed in the same way as fecal specimen is processed (Rajesh &Rattan, 2004).

#### 2-2-4-2 Slide agglutination test:

If on bases of biochemical reaction the organism has been identified as *Salmonella*, its identify can be confirmed with slide agglutination test and serotype ascertained. The identify of genus can be confirmed by observing for agglutination with polyvalent O antisera and polyvalent H antisera against *salmonellae*. For identification of serotype the isolate is reacted with group specific antisera followed by monovalent O specific antiserum. Similarly H antigen in phase 1 or phase 2 can be determined to reach at final antigenic structure of organism (Rajesh& Rattan, 2004).

#### 2-2-4-3 Detection of microbial antigen:

The circulating *Salmonella* antigen may be detected in blood of patients with typhoid fever by coagulation method. ELISA has also been attempted by some workers to detect Vi antigen in the urine of patients.

#### 2-2-4-4 Titration of antibody:

Large numbers of serological tests have been devised to detect and titrate the antibody against common agents of typhoid fever:

- Widal tube agglutination test.
- Widal slide agglutination test.
- Indirect haemagglutiation test.
- Counterimmunoelectrophoresis.
- ELISA.

(Rajesh & Rattan, 2004).

#### 2-2-5 Treatment of Typhoid Fever:s

*H.pylori* sensitive to Ciprofloxacin, Azithromycin, Tetracycline, Ceftriaxone, Trimethoprim and Sulfamethazole, and Chloromphenicol. (Rajesh & Rattan, 2004).

#### 2-2-6 Previous study:

A case control study conducted in urban slum community in south Delhi from November 1995 to October 1996 to determine the association between typhoid fever and *Helicobacter pylori*, 83 case subjects of culture-proven typhoid fever were identified through one year surveillance of subject age 0-40 years, two agesex matched neighborhood. control subjects selected for each case subject. Serum anti H.pylori immunoglobulin G antibodies were measured in case and neighborhood control subjects, the result was a significant association between typhoid fever and *H.pylori* that anti *H.pylori* IgG detected in 64% of case subjects (53 of 83) and 50% (83 of 166) neighborhood control subjects (Maharaj, 2002).

## **Chapter Three**

## **Materials and Methods**

#### **3-** Materials and Methods

#### 3-1 Study design:

This was a cross-sectional hospital based study conducted in Atbara Teaching Hospital.

#### 3-2 Study area:

Atbara Town is a town located in River Nile State away from Khartoum about 310 kilometers to the north-east direction, between latitude 14. and 17 degrees north and longitude 33 and 59 degrees east.

#### **3-3 Study duration:**

The study was performed during period from 20 May to 20 July at Royal Laboratory in Atbara City.

#### **3-4 Study population:**

The target people in this study were those with typhoid fever with exclusion of patients that have been infected with typhoid fever during the last six month.

#### 3-5 Sampling and sample size:

Seventy two serum samples were collected from diagnosed typhoid patients with symptoms of illness using sterile syringe. The subject's veins were sterilized with 70% alcohol using impregnated cotton then puncture was made with needle smoothly and whole blood was collected from the vein in sterile plain container. Then after blood was clotted it was centrifuged at 3000 for 5 minutes. Patients were grouped into males and females, and according to age grouped from 15-30 , and more than 30 years.

#### 3-6 Data collection:

#### **3-6-1 Data collection tool:**

The data collected from the patient using closed answer questionnaire.

#### 3-6-2 Data analysis:

The collected data were analyzed with SPSS program version 18.

#### **3-7 Ethical consideration:**

After approval of the research from the ethical committee of medical laboratory sciences ,All patient under study were informed about the objective of research , the verbal consent was taken from them before enrolled under study.

#### 3-8 anti *H.pylori* ELISA:

ELISA kit purchased from EUROIMMUN company containing microplate wells  $12 \times 8$ , three calibrator 1,2 and 3, positive and negative control, enzyme conjugate, sample buffer, wash buffer, chromogen/substrate solution and stop solution.

#### **3-8-1 Principle of test:**

The ELISA test kit provides a semi-quantitative in vitro assay for human antibodies of the IgG class against *H.pylori* in serum or plasma. The kit contains microtiter strips each with 8 break-off reagent wells coated with *H.pylori* antigens. In the first reaction step, diluted patients samples are incubated in the well. In case of positive samples, specific IgG antibodies will bind to antigens. To detect the bound antibodies, a second incubation is carried out using an enzyme-labelled anti-human IgG (enzyme conjugate) catalyzing a colour reaction.

#### 3-8-2 Preparation of sample and washing buffer:

Patients samples were diluted 1:101 in sample buffer 10  $\mu$ l serum in 1.0 ml sample buffer and mixed by vortexing.

Concentrated wash buffer diluted from 10 x to 1.0x by adding 100ml of of washing buffer to 900 ml of distilled water.

#### 3-8-3 Procedure of semi-quantitative analysis:

For semi-quantitative method calibrator 2 along with positive, negative controls and patients samples.

First step, sample incubation: 100  $\mu$ l was pipetted per well and incubated for 30 minutes at room temperature, then the well were manually washed by empting the well and subsequently washed three times using 300  $\mu$ l of working wash buffer.

Second step, conjugate incubation:  $100\mu$ l of enzyme conjugate (peroxidaselabled anti human IgG) was pipetted into each microplate wells and incubated for 30 minutes at room temperature, then the well were manually washed by emptying the well and subsequently washed three times using 300 µl of working wash buffer.

Third step, substrate incubation:  $100\mu$ l of chromogen/substrate (TMB/H<sub>2</sub>O) solution was pipetted into each microplate wells and incubated for 15 minutes at room temperature.

Stop the reaction  $100\mu$ l of stop solution(0.5 M Sulphuric acid) was pipetted into each microplate wells in the same order and at the same speed as chromogen/substrate solution was introduced.

Last step:Photometric measurement was done by using ELISA reader which performed as absorbance in filter 492 nm.

#### **3-8-4 interpretation of result:**

Results had been interpretated by calculation a ratio of the extinction value of control or patient sample over the extinction value of the calibrator 2.

#### Formula of the ratio:

extinction of control or patient sampl	e
extinction of calibrator 2	

- ratio < 0.8	negative
- ratio ≥0.8 to <1.1	borderline
- ratio ≥1.1	positive

# **Chapter Four**

Results

#### **4-Results**

-A total of (72) serum sample collected from typhoid fever patients, attending A, 40 (56%) were male and 32(44%) were female. As shown in **Table (4-1)**.

-Distribution of study population according to Resident there was 23(32%) from Urban and 49(68%) from the Rural, as mentioned in **Table (4-2)**.

-In regard to tribes, 21(29%) from Gaalia tribe, 11(15%) Shaygia, 17(24%) Rashayda and 23 (32%) from other tribes as revealed in **Table (4-3)**.

-According to present of typhoid fever among study population in different ages, 29(40%) in age group (15-30), 43(60%) within age more than 30 years as mentioned in **Table (4-4)**.

-A total of (92) serum sample collected from Atbara hospital, found that, (72) typhoid patient 59 of them were positive (81.9%) and out of (20) non typhoid subjects ,10 of them were positive (10%)As shown in **Table (4-5)**.

-Frequency of *H.pylori* according to present of typhoid fever among study population there were 59(81.9%) infected with H.pylori and 13(18.1%) were not infected as presented in **Table(4-6)**.

- In regard to frequency of infected patients with H. pylori, out of 72, 59 (81.9%) were infected, from them 33(55.9%) were males, and 26(44.1%) were females, while the rest were not as shown in **Table (4-7)**.

- In relationship between residents and Helicobacter pylori, found that; 13(22%) from urban and 46(78%) from rural as presented in **Table (4-8)**.

- According to relationship between tribes and infection with H.pylori, 18 (30.5%) Gaalia, 8(13.6%) Shaygia, 16 (27.1%) Rashayda, and others 17 (28.8%) as revealed in **Table ( 4-9 ).** 

- Regarding relationship between age groups and infection with H. pylori, found that; 21 (35.6%) within (15-30) age, and 38(64.4%) in age more than 30 years as presented in **Table (4-10)**.

- In related to risk factors of infected with H .pylori, 34 (57%) were smoking while 25(42.4%) were not as shown in **Table (4-11).** Also found 37(62.8%)

were drinking coffee, and the remaining 22(37.2%) were not as tabulated in **Table (4-12).** 

Gender	Total number	Percentage
Male	40	56%
Female	32	44%
Total	72	100

Table (4-1): Show the distribution of the study population according to gender:

Table (4-2): Show the distribution of the study samples according to Resident:

Resident	Total number	Percentage
Urban	23	32%
Rural	49	68%
Total	72	100

Table (4-3): Show the distribution of the study samples according to Tribe:

Tribe	Total number	Percentage
Gaalia	21	29%
Shaygia	11	15%
Rashayda	17	24%
Other	23	32%
Total	72	100

Age group	Total number	Percentage%
15 - 30	29	40
More than 30	43	60
Total	72	100

Table (4-4): Show the distribution of the study population according toAge:

 Table (4-5) Frequency of H.pylori in typhoid patients and non typhoid

 subjects:

Sample	Positive	Negative	Total	Frequency
Typhoid patients	59	13	72	81.9%
Non typhoid Subjects	10	10	20	50%

### p.value=0.000

 Table (4-6): Frequency of *H.pylori* among study population:

Infection with H. Pylori	Frequency	Percentage
Infected	59	81.9%
Non infected	13	18.1%
Total	72	100

P.value=0.000

Gender	Positive	Negative
Male	33 (55.9%)	7 (53.8%)
Female	26 (44.1%)	6 (46.2%)
Total	59 (100%)	13 (100%)

 Table (4-7): Relationship between infection with H.pylori &gender:

p.value=0.003

 Table (4-8): Relationship between infection with H.pylori & Resident:

Resident	Positive	Negative
Urban	13 (22.0%)	10 (76.9%)
Rural	46 (78.0%)	3 (23.1%)
Total	59 (100%)	13 (100%)

p.value=0.006

Tribe	Positive	Negative	P.value
Gaalia	18 <b>(30.5%)</b>	3 (23.1%)	0.001
Shaygia	8 (13.6%)	3 (23.1%)	0.132
Rashayda	16 <b>(27.1%)</b>	1 (7.7%)	0.000
Other	17 <b>(28.8%)</b>	6 (46.2%)	0.022
Total	59 (100%)	13 (100%)	

Age group	Positive	Negative
15 - 30	21 (35.6%)	8 (61.5%)
More than 30	38 (64.4%)	5 (38.5%)
Total	59 (100%)	13 (100%)

Table (4-10) Relationship between infection with *H.pylori and age*:

p.value=0.000

### Table (4-11): Relationship between infection with H.pylori & smoking behavior:

Smoking behavior	Positive	Negative
Smoker	34 (58.0%)	8 (61.5 %)
Non smoker	25 (42.0%)	5 (38.5%)
Total	59 (100%)	13 (100%)

p.value=0.001

Table (4-12): Relationship between infection with of *H.pylori* & coffee drinking:

Coffee drinking	Positive	Negative
Yes	37 (62.8%)	3(23.1%)
No	22 (37.2%)	10(76.9%)
Total	59 (100%)	13 (100%)

p.value = 0.02

# **Chapter Five**

Discussion

Conclusion

Recommendations

### 5-1 Discussion

The prevalence of *H. pylori* infection varies widely by geographic area, age, and race. Because it is not possible to ascertain when infection occurs clinically (Parsonnet, 1995), most of the information on the rates of *H. pylori* in geographically and demographically diverse populations comes from seroprevalence studies. The main finding in this cross-sectional study that; the prevalence of *H.pylori* among typhoid patients were 59 (81.9%) out of 72 patients. And10(50%) among 20 controls samples of non typhoid subjects. The prevalence was higher in patients with typhoid more than other non typhoid subjects this consistent with study done in India (Maharaj, 2002), and this more likely to be that *H.pylori* change in acid barrier of stomach by causing transient hypochlorhydria (Harford *et al.*, 2000) which facilitate infection with *Salmonella*.

Regarding to infection of H. pylori among study population, 33(55.9%) were male, while 26(44.1%) were female, the prevalence of *H.pylori* was higher in male than female and there is statistically significant relationship (*P*.value = 0.03) this agree with (Yvonne &*Rob.*, 2001). and the reason for increased infection in males because males were more exposed to the risk factors that increased infection such as smoking and alcohol consumption.

The study revealed that the infection of *H.pylori* present more in rural 49 (68%) compared to Urban 23(32%), this is statistically significant (P.value=0.006), and agree with (Parsonnet, 1995) may be due to A scarcity of running and non purified water and preserved of water in a tank for long time before used in rural area.

In regarding to H.pylori infection and tribes, 21(29%) were Gaalia, 11 (15%) Shaygia, 17(24%) Rashayda and 23 (32%) from other tribes. Gaalia tribe more likely to be infected with (p.value=0.001) because they represent indigenous population of the region.

The highest frequency of infection with H.pylori occurred in age group more than 30 years 38 (64.4%) flowed by 21(35.6%) from the age group within (15-

30). Noticed that the disease was increased with ages. However, high infection may be attributed to long time exposure to the causative agent during their early childhood and decreased the immunity. (Mitchell *et al.*, 1992), and there was statistically significant relationship (P. value=0.000).

In regarding to risk factors, found that; highly infection with H.pylori occurred among smokers 42 (58%), statistically significant (P.value= 0.001) and this agree with most recent researches (Zhang *et al.*, 1996; Fraser *et al.*, 1996; Brenner *et al.*, 1997).

Also the study revealed association between H.pylori infection and coffee drinkers 40(56%), which shown statistically significant (P.value = 0.02), this agree with (Brenner *et al.*, 1997; Brenner *et al.*, 1999a,b) may be due to decreased in stomach pH which have little effect on the H.pylori infection.

### **5-2** Conclusion

The current study revealed relationship between typhoid fever and infection with H. Pylori. The infection with H. Pylori increased with age, tribes, and other risk factors.

### **5-3 Recommendations**

- An effort by the government for eradication of *Helicobacter pylori* which increase risk of infection with typhoid fever.
- Health education program should be taken place especially on personal hygiene.
- Food must be prepared properly and drinking water taken from safe and clean source.
- Research with large sample size will powered by estimation gastric acid for better result.

## **Chapter Six**

References

Appendices

#### References

- Albertson N., Wenngren I., Sjo stro m J-E (1998). Growth and survival of Helicobacter pylori in defined medium and susceptibility to Brij 78. *Journal of Clinical Microbiology*, 36: 1232–1235.
- Andersen AP., Elliott DA., Lawson M., Barland P., (1997). Growth and morphological transformations of Helicobacter pylori in broth media. *Journal of Clinical Microbiology*, 35: 2918–2922.
- 3. Banwart GJ, (1979). Basic food microbiology. Westport, CT, AVI Publishing.
- 4. Bardhan PK.,(1997). Epidemiological features of *Helicobacter pylori* infection in developing countries. *Clin Infect Dis*; 25:973–978.
- Bhan MK., Bahl R., Sazawal S., (2002). Association between *Helicobacter* pylori infection and increased risk of typhoid fever. J Infect Dis; 186:1857– 1860.
- Blaser MJ., (1999). Hypothesis: the changing relationships of *Helicobacter* pylori and humans: implications for health and disease. J Infect Dis; 179:1523–1530.
- Bode G.,Rothenbacher D., Brenner H., Adler G., (1998). Pets are not a risk factor for Helicobacter pylori infection in young children: results of a population-based study in Southern Germany. *Pediatric Infectious Disease Journal*; 17: 909–912.
- Bode G., Mauch F., Malfertheiner P., (1993). The coccoid forms of Helicobacter pylori. Criteria for their viability. *Epidemiologyand Infection*; 111: 483–490.
- Bohmer CJ., Klinkenberg-Knol EC., Kuipers EJ., (1997). The prevalence of *Helicobacter pylori* infection among inhabitants and healthy employees of institutes for the intellectually disabled. *Am J Gastroenterol*; 92:1000-4.
- 10. Brenner H., Bode G., Boeing H., (2000). *Helicobacter pylori* infection among offspring of patients with stomach cancer. *Gastroenterology*; 118:31-5.

- (a) Brenner H., Rothenbacher D., Bode G., (1999). Inverse graded relation between alcohol consumption and active infection with *Helicobacter pylori*. *Am J Epidemiol*; 149:571-6.
- (b) Brenner H., Berg G., Lappus N., (1999). Alcohol consumption and *Helicobacter pylori* infection: results from the German National Health and Nutrition Survey. *Epidemiology*; 10:214-18.
- 13.Brenner H., Rothenbacher D., Bode G., (1997). Relation of smoking and alcohol and coffee consumption to active *Helicobacter pylori* infection: cross sectional study. *BMJ*; 315:1489-92.
- 14.Crump JA., Luby SP., Mintz ED., (2004). The global burden of typhoid fever. Bull World Health Organ; 82:346–353.
- 15.Center of Disease Control and Prevention.,(1998), *H.pylori*, fact sheet for health care provider; 1-4.
- 16.Donelli G., Matarrese P., Dainlli B., Taraborelli T., (1998). The effect of oxygen on the growth and cell morphology of *Helicobacter pylori*. *FEMS Microbiology Letters*; 168: 9–15.
- 17. Dunn BE., Cohen H., Blaser MJ., (1997). Helicobacter pylori. Clinical Microbiology Reviews; 10: 720–741.
- 18.Dunstan SJ., Stephens HA., Blackwell JM., (2001). Genes of the class II and class III major histocompatibility complex are associated with typhoid fever in Vietnam. J Infect Dis; 183:261–268.
- 19.Edelman R., and Levine Myron M., (1986). Summary of an international workshop on typhoid fever. *Reviews of Infectious Diseases*; 8(3): 329-47.
- 20.El Omar EM., Carrington M., Chow WH., (2000). Interleukin-1 polymorphisms associated with increased risk of gastric cancer. *Nature*; 404:398–402.
- 21.Fantry GT., Zheng QX., James SP., (1995). Conventional cleaning and disinfection techniques eliminate the risk of endoscopic transmission of *Helicobacter pylori. Am J Gastroenterol*; 90:227-32.

- 22.Fraser AG, Scragg R, Metcalf P, 1996). Prevalence of *Helicobacter pylori* infection in different ethnic groups in New Zealand children and adults. *Aust N Z J Med* ;26: 646-51.
- 23.Fan X-G., Chua A., Li TG., Zeng QS., (1998). Survival of Helicobacter pylori in milk and tap water. *Journal of Gastroenterology and Hepatology*; 13: 1096– 1098.
- 24.Fontham ET., Ruiz B., Perez A., (1995). Determinants of *Helicobacter pylori* infection and chronic gastritis. *Am J Gastroenterol;* 90:1094-101.
- 25.Fox JG., (1995). Non-human reservoirs of *Helicobacter pylori*. Alimentary *Pharmacology and Therapeutics*; 9 (Suppl. 2): 93–103.
- 26. Friis L., Engstrand L., Edling C., (1996). Prevalence of *Helicobacter pylori* infection among sewage workers. *Scand J Work Environ Health* ; 22:364-8.
- 27. Goodman KJ., Correa P., Tengana Aux HJ., (1996). *Helicobacter pylori* infection in the Colombian Andes: a population-based study of transmission pathways. Am J Epidemiol; 144: 290-9.
- 28.Goodman KJ., Correa P., Tengana AH., (1997). Nutritional factors and Helicobacter pylori infection in Colombian children. J Pediatr Gastroenterol Nutr; 25:507-15.
- 29.Goodwin CS., (1989). Campylobacter pylori: detection and culture. In: Rathbone BJ, Heatly RV, eds. Campylobacter pylori and gastroduodenal disease. Oxford, Blackwell Scientific Publications; 60–62.
- 30.Goodwin CS.,and Armstrong JA., (1990). Microbiological aspects of Helicobacter pylori (Campylobacter pylori). European Journal of Clinical Microbiology; 9: 1–13.
- 31.Gotuzzo E., Frisancho O., Sanchez J., Liendo G., Carillo C., Black RE., Morris JG., (1991). Association between the acquired immunodeficiency syndrome and infection with *Salmonella typhi* or *Salmonella paratyphi* in an endemic typhoid area. *Archives of Internal Medicine*; 151: 381-2.

- 32.Gramley WA., Asghar A., Frierson HFJ., (1999). Detection of *Helicobacter pylori* DNA in fecal samples from infected individuals. *J Clin Microbiol*; 37:22-36.
- 33. Handt LK., Fox JG., Dewhirst FE., (1994). *Helicobacter pylori* isolated from the domestic cat: public health implications. *Infect Immun*; 62:2367-74.
- 34.Harford WV., Barnett C., Lee E., Perez-Perez G., Blaser MJ., Peterson WL., (2000). Acute gastritis with hypochlorhydria: report of 35 cases with long term follow up. *Gut*; 47:467–472.
- 35. Ivanoff B., Cordel J., Robert D., Fontanges R., (1980). Importance de la voie respiratoire dans la salmonellose expérimentale de la souris Balb/c. *Comptes Rendus de l'Académie des Sciences* (Paris); 1271-4.
- 36.Jenkins DJ., (1997). *Helicobacter pylori* and its interaction with risk factors for chronic disease. *BMJ*; 315:1481-2.
- 37.Jiang X.,and Doyle MP., (1998). Effect of environmental and substrate factors on survival and growth of *Helicobacter pylori*. *Journal of Food Protection*; 61: 929–933.
- 38.Klontz KC., (1999). Does imbibing alcohol protect against enteric pathogens? *Epidemiology*; 10:207-9.
- 39.Kunz LJ., Waddell WR., (1956). Association of *Salmonella enteritis* with operations on the stomach. *N Engl J Med*; 255:555–559.
- 40.Kusters JG., Gerrits MM., Van den Brouke-Grauls CMJE., (1996). The morphological conversion of H. pylori from bacillary to coccoid forms is not an active process. In: Abstracts of the 36<sup>th</sup> Interscience Conference on Antimicrobial Agents and Chemotherapy. Washington, DC, *American Society for Microbiology*; 25.
- 41.Lambert JR., Lin SK., Sievert W., (1995). High prevalence of *Helicobacter pylori* antibodies in an institutionalized population: evidence for person-to-person transmission. *Am J Gastroenterol;* 90:2167-71.

- 42.Lin SK., Lambert JR., Nicholson L., (1998). Prevalence of *Helicobacter pylori* in a representative Anglo-Celtic population of urban Melbourne. J Gastroenterol Hepatol; 13: 505-10.
- 43.Luby SP., Faizan MK., Fisher-Hoch SP., (1998). Risk factors for typhoid fever in an endemic setting, Karachi, Pakistan. *Epidemiol Infect*; 120:129–138.
- 44.**Maharaj K, (2002).** Association between *Helicobacter pylori* infection and increase risk of typhoid fever.*J Infect Dis;* 186(12):1857-1860
- 45. Mitchell HM., Bohane T., Hawkes RA., (1993). *Helicobacter pylori* infection within families. *Int J Med Microbiol Virol Parasitol Infect Dis*; 280:128-36.
- 46.Mitchell HM., LiYY., Hu PJ., (1992). Epidemiology of *Helicobacter pylori* in southern China: identification of early childhood as the critical period for acquisition. *J Infect Dis*; 166:149-53.
- 47.Namavar F., Roosendaal R., Kuipers EJ., (1995). Presence of *Helicobacter pylori* in the oral cavity, oesophagus, stomach and faeces of patients with gastritis. *Eur J Clin Microbiol Infect Dis*; 14:234-7.
- 48.Nilius M., Strohle A., Bode G., Malfertheiner P.,(1993). Coccoid like forms (clf) of Helicobacter pylori. Enzyme activity and antigenicity. International Journal of Medical Microbiology, Virology, Parasitology and Infectious Diseases; 280: 259–272.
- 49. Parsonnet J,(1995). The incidence of *Helicobacter pylori* infection.
- 50. Aliment Pharmacol Ther;9(suppl 2):45-51
- 51. Parsonnet J., Shmuely H., Haggerty T., (1999). Fecal and oral shedding of *Helicobacter pylori* from healthy infected adults. *JAMA*; 282:2240-5.
- 52. Rajesh B., and Rattan LI., (2004). Essentials of medical microbiology. New Delhi: *Jitender P Vij*, 3<sup>d</sup> ed: 248-314.
- 53.Reynolds DJ., and Penn CW., (1994). Characteristics of Helicobacter pylori growth in a defined medium and determination of its amino acid requirements. *Microbiology*; 140: 2649–2656.
- 54. Rothenbacher D., Blaser MJ., Bode G., Brenner H., (2000). Inverse relationship between gastric colonization of *Helicobacter pylori* and diarrheal

illnesses in children: results of a population-based cross-sectional study. *J Infect Dis*; 182:1446–1449.

- 55.Shahinian ML., Passaro DJ., Swerdlow DL., Mintz ED., Rodriguez M., Parsonnel J., (2000). *Helicobacter pylori* and epidemic *Vibrio cholerae* O1 infection in Peru. *Lancet*; 355:377–378.
- 56.Shimada T., Ogura K., Ota S., (1994). Identification of *Helicobacter pylori* in gastric specimens, gastric juice, saliva, and faeces of Japanese patients. (Letter). *Lancet*; 343:1636-7.
- 57.Suerbaum S., and Michetti P., (2002). *Helicobacter pylori* infection. *N Engl J Med*; 347:1175–1186.
- 58.Sullivan PB., Thomas JE., Wight DG., (1990). Helicobacter pylori in Gambian children with chronic diarrhoea and malnutrition. Arch Dis Child; 65:189–191.
- 59. Torres J., Perez-Perez G., Goodman KJ., (2000). A comprehensive review of the natural history of *Helicobacter pylori* infection in children. *Arch Med Res*; 31:431–469.
- 60.van Zwet AA., Thijs JC., Kooistra-Smid AM., (1994). Use of PCR with feces for detection of *Helicobacter pylori* infections in patients. *J Clin Microbiol*; 32:1346-8.
- 61. Webb PM., Knight T., Elder JB., Newell DG., Forman D., (1996). Is *Helicobacter pylori* transmitted from cats to humans? *Helicobacter*; 1: 79–81.
- 62. West AP., Miller MR., Tompkins DS., (1992). Effect of physical environment on survival of *Helicobacter pylori*. *Journal of Clinical Pathology*; 45: 228–231.
- 63.**WHO., (2003).** Background document: The diagnosis, treatment and prevention of typhoid fever; 1-7.
- 64. Yvonne T.H.P. van Duynhoven and Rob de Jonge, (2001). Transmission of *Helicobacter pylori*: a role for food?, *Bulletin of the World Health Organization*; 79 (5).

65. Zhang L, Blot WJ, You WC, (1996). *Helicobacter pylori* antibodies in relation to precancerous gastric lesions in a highrisk Chinese population. *Cancer Epidemiol Biomarkers Prev*; 5:627-30.

### **Appendix I**

#### University of Shendi

College of Graduate Studies and Scientific Research.

Sero-detection of anti Helicobacter pylori IgG antibodies in typhoid patients

### in Atbara Teaching Hospital

Name: Serial No: Tribe:..... Resident: Occupation: a- Gender: 1/ Male ( 2/ Female ) ( ). b- Age: 2/More than 30 ( 1/15-30 ( ) ). c- Smoking Behavior: 1/Yes ( 2/No( ) ). d- Alcohol consumption: 2/No ( ) 1/Yes(). e- Coffee drinking: 1/Yes ( 2/No ( ) ). f- Patient anti H.pylori ELISA result: 2/Negative ( 1/ Positive ( ) ).

### Appendix II:





