



UNIFORMED SERVICES UNIVERSITY OF THE HEALTH SCIENCES
F. EDWARD HEBERT SCHOOL OF MEDICINE
4301 JONES BRIDGE ROAD
BETHESDA, MARYLAND 20814-4799



GRADUATE EDUCATION
(301) 295-3913
FAX: (301) 295-6772

APPROVAL SHEET

Title of Dissertation: "*Helicobacter Pylori* Transmission and Risk Factors for Infection in Rural China"

Name of Candidate: CAPT Linda Brown
Doctor of Public Health Degree
8 December 1999

Dissertation and Abstract Approved:

David Cruess, Ph.D.
Department of Preventive Medicine & Biometrics
Committee Chairperson

2/2/00

Date

Terry Thomas, Ph.D.
Department of Preventive Medicine & Biometrics
Committee Member

01/10/2000

Date

Gary Gackstetter, COL, USAF, MC
Department of Preventive Medicine & Biometrics
Committee Member

03/07/99

Date

Andre Dubois, M.D., Ph.D.
Department of Medicine
Committee Member

01/10/2000

Date

Mitchell Gail, M.D., Ph.D.
National Cancer Institute, NIH
Committee Member

01/13/2000

Date



Report Documentation Page				Form Approved OMB No. 0704-0188	
Public reporting burden for the collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Washington Headquarters Services, Directorate for Information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington VA 22202-4302. Respondents should be aware that notwithstanding any other provision of law, no person shall be subject to a penalty for failing to comply with a collection of information if it does not display a currently valid OMB control number.					
1. REPORT DATE 1999		2. REPORT TYPE		3. DATES COVERED 00-00-1999 to 00-00-1999	
4. TITLE AND SUBTITLE Helicobacter Pylori Transmission and Risk Factors for Infection in Rural China				5a. CONTRACT NUMBER	
				5b. GRANT NUMBER	
				5c. PROGRAM ELEMENT NUMBER	
6. AUTHOR(S)				5d. PROJECT NUMBER	
				5e. TASK NUMBER	
				5f. WORK UNIT NUMBER	
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) Uniformed Services University of the Health Sciences,F. Edward Hebert School of Medicine,4301 Jones Bridge Road,Bethesda,MD,20814-4799				8. PERFORMING ORGANIZATION REPORT NUMBER	
9. SPONSORING/MONITORING AGENCY NAME(S) AND ADDRESS(ES)				10. SPONSOR/MONITOR'S ACRONYM(S)	
				11. SPONSOR/MONITOR'S REPORT NUMBER(S)	
12. DISTRIBUTION/AVAILABILITY STATEMENT Approved for public release; distribution unlimited					
13. SUPPLEMENTARY NOTES					
14. ABSTRACT see report					
15. SUBJECT TERMS					
16. SECURITY CLASSIFICATION OF:			17. LIMITATION OF ABSTRACT	18. NUMBER OF PAGES 223	19a. NAME OF RESPONSIBLE PERSON
a. REPORT unclassified	b. ABSTRACT unclassified	c. THIS PAGE unclassified			

COPYRIGHT STATEMENT

The author hereby certifies that the use of any copyrighted material in the dissertation entitled:

***"HELICOBACTER PYLORI* TRANSMISSION AND RISK FACTORS FOR INFECTION IN RURAL CHINA"**

beyond brief excerpts is with the permission of the copyright owner, and will save and hold harmless the Uniformed Services University of the Health Sciences from any damage which may arise from such copyright violations.

ABSTRACT

Title of Dissertation: *Helicobacter pylori* Transmission and Risk Factors for Infection in Rural China

Name, degree, year: Linda Morris Brown, Dr.P.H., 1999

Thesis directed by: Terry L. Thomas, Ph.D., Associate Professor, Department of Preventive Medicine & Biometrics

Helicobacter pylori (*H. pylori*) is one of the most common bacterial infections among humans, but little is known about its mode of transmission. A cross-sectional study of 3288 adults aged 35-69 from Shandong Province, China was conducted to assess some possible risk factors that may be associated with *H. pylori* infection in this high prevalence area of China. In-person home interviews (lasting approximately 15 minutes) were conducted in Chinese from October 1997-May 1998. The response rate was 96.4%. Maximum likelihood estimates and bootstrap confidence intervals (CI) of the association between the prevalence of *H. pylori* infection and demographic, lifestyle and some common environmental exposures were computed using polychotomous logistic regression.

The *H. pylori* serostatus of the study participants was positive 60.6%, negative 31.0%, and indeterminate 8.4%. Source of drinking water, especially water from a shallow village well (OR=1.8, 95%CI=1.4-2.3), was associated with increased prevalence of *H. pylori* infection. ORs were also elevated for infrequent hand washing before meals (OR=1.6, 95%CI=1.0-2.5) and bathing in a pond or ditch (OR=1.6, 95%CI=1.0-2.4). ORs were also associated with median village education level, ranging from 1.0 for villages classified as high, to

1.7 (95%CI=1.4-2.1) for villages classified as medium, to 2.4 (95%CI=2.0-3.0) for villages classified as low. ORs decreased slightly with increased consumption of all allium vegetables combined. The ORs were reduced for having a cat as a pet during childhood (OR=0.7, 95%CI=0.5-1.0) and/or adulthood (OR=0.6, 95%CI). No significant associations were seen with any measure of cigarette smoking or alcoholic beverage consumption. Additionally, crowding or density factors as a child were not related to a higher prevalence of *H. pylori* infection as an adult.

The results of the multi-variate modeling revealed prevalence rates that ranged from 75% to 46%, prevalence rate ratios that ranged from 1.0-1.5, and relative odds ratios that ranged from 1.0-8.6. Although a lack of heterogeneity in the population may have hampered the ability to detect actual associations, the finding that water may contribute to *H. pylori* infection deserves further evaluation. It is also noteworthy that cat ownership was not associated with an increased prevalence of *H. pylori*. The results of this study suggest that person-to-person and waterborne transmission, but not zoonotic transmission, are likely routes of *H. pylori* infection in this rural Chinese population.

***HELICOBACTER PYLORI* TRANSMISSION
AND RISK FACTORS FOR INFECTION
IN RURAL CHINA**

by

Linda Morris Brown

**Dissertation submitted to the Faculty of the
Department of Preventive Medicine and Biometrics of the
Uniformed Services University of the Health Sciences
in partial fulfillment of the requirements for the degree
of Doctor of Public Health 1999**

ACKNOWLEDGMENTS

I thank my advisor Dr. Terry Thomas and the other members of my dissertation committee - Drs. David Cruess, Andre Dubois, Gary Gackstetter, and Mitchell Gail for their advice and helpful comments; my colleagues at the National Cancer Institute - Drs. Joseph F. Fraumeni, Jr., Mitchell Gail, Frank Groves, and Wei-cheng You for their technical and administrative support; my colleagues at the Beijing Institute for Cancer Research - Drs. Yunsheng Chang, Wei-dong Liu, Jun-Ling Ma, Kai-feng Pan, and Lian Zhang for their technical assistance and management of field activities; John Heinrich (Westat, Inc.) for his assistance in data file management; Stella Semiti and Jerome Mabi (Information Management Systems) for computer support; and the residents of the 13 Linqu villages who participated in this study.

DEDICATION

To Richard, Leslie, Jeffrey, and Rocky for their support and understanding.

TABLE OF CONTENTS

Approval Sheet	i
Copyright Statement	ii
Abstract	iii
Title Page	v
Acknowledgments	vi
Dedication	vii
List of Tables	xiii
List of Maps	xv
Chapter I. Background and Rationale	1
Background	1
Modes of Transmission	1
Epidemiology	2
Rationale	3
Research Questions	4
Chapter II. Literature Review	5
Introduction	5
Microbiologic Characteristics	6
Diagnostic Tests Used In Epidemiologic Studies	6
Biopsy-based Tests	7
Serologic Tests	7
Breath Test	9
Disease Associations	9

Epidemiology	12
Descriptive Studies/Prevalence	12
Risk Factors	15
Smoking	16
Alcohol Use	17
Diet	18
Occupational Exposures	19
Waterborne Exposures	21
Exposure To Animals/Pets	22
Hygiene Practices	23
Density/Crowding	24
Social Factors	24
Family History Of Gastric Disease/Ulcer	26
Possible Routes Of Transmission	26
Person-To-Person Transmission	26
Institutionalized Populations	26
Familial Exposures	28
Oral-Oral Route	30
Fecal-Oral Route	33
Waterborne Transmission	34
Zoonotic/Vectorborne Transmission	35
Iatrogenic Transmission	37
Summary Of Published Literature	39

Chapter III. Methods	43
Study Population	43
Description Of Area	46
Study Design	47
Interview And Questionnaire	48
Reclassification Of <i>H. Pylori</i> Status	50
Data Analysis	50
Human Subjects And Confidentiality	53
Investigators	53
Chapter IV. Results	55
Demographic Characteristics	55
Risk Factors	56
Cigarette Smoking	56
Alcoholic Beverage Consumption	57
Dietary Factors	57
Occupation	58
Water-Related Exposures	58
Exposure To Animals And Pets	60
Sanitation/Hygiene Factors	60
Density/Crowding Factors	61
Social Factors	62
Gastric Factors	62
Multi-Variate Analysis	63

Chapter V. Discussion	67
Research Question 1.	67
Research Question 2.	68
Research Question 3.	70
Research Question 4.	71
Research Question 5.	71
Research Question 6.	72
Research Question 7.	73
Research Question 8.	74
Strengths Of The Study	76
Limitations Of The Study	78
Conclusions	80
Future Directions	81
Global Issues	81
Studies Within The Intervention Trial Cohort	82
Public Health Significance of <i>H. pylori</i> Research in Linq County	83
Tables	85
Appendix 1	155
Maps	157
Appendix 2	159
English Version of Questionnaire	160
Consent Form	173
Chinese Version of Questionnaire	176

Bibliography	184
--------------------	-----

LIST OF TABLES

Table II-1.	Prevalence of <i>Helicobacter pylori</i> in various adult populations ..	86
Table II-2.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and smoking	89
Table II-3.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and alcohol use	92
Table II-4.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and diet	95
Table II-5.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and occupational exposures	98
Table II-6.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and waterborne exposures	102
Table II-7.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and exposure to pets and other animals	104
Table II-8.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and hygiene practices	107
Table II-9.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and density/crowding	110
Table II-10.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and social factors	115
Table II-11.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and family history of gastric disease/ulcer	121
Table II-12.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and institutionalized populations	123
Table II-13.	Studies assessing the relationship between <i>Helicobacter pylori</i> infection and familial exposures	125
Table IV-1.	Distribution of cases and controls for selected variables by <i>Helicobacter pylori</i> status	129

Table IV-2.	Association between prevalence of <i>Helicobacter pylori</i> infection and cigarette smoking	132
Table IV-3.	Association between prevalence of <i>Helicobacter pylori</i> infection and alcoholic beverage consumption	134
Table IV-4.	Association between prevalence of <i>Helicobacter pylori</i> infection and dietary variables	135
Table IV-5.	Association between prevalence of <i>Helicobacter pylori</i> infection and occupation variables	138
Table IV-6.	Association between prevalence of <i>Helicobacter pylori</i> infection and water-related exposures	139
Table IV-7.	Association between prevalence of <i>Helicobacter pylori</i> infection and animals and pets	141
Table IV-8.	Association between prevalence of <i>Helicobacter pylori</i> infection and hygiene factors	143
Table IV-9.	Association between prevalence of <i>Helicobacter pylori</i> infection and density/crowding factors	146
Table IV-10	Association between prevalence of <i>Helicobacter pylori</i> infection and social factors	149
Table IV-11	Association between prevalence of <i>Helicobacter pylori</i> infection and gastric factors	151
Table IV-12.	Prevalence rate, prevalence rate ratio, and relative odds ratio of <i>Helicobacter pylori</i> infection for combinations of models according to age, percent of time washed hands before eating, number of children in the household, and village education level	152
Table IV-13.	Prevalence rate, prevalence rate ratio, and relative odds ratio of <i>Helicobacter pylori</i> infection for combinations of models according to age, percent of time washed hands before eating, number of children in the household, and source of drinking water	154

LIST OF MAPS

Map Of China Showing Location Of Shandong Province	157
Map Of Shandong Province Showing Location Of Linqu County	158

CHAPTER I. BACKGROUND AND RATIONALE

BACKGROUND

Helicobacter pylori (*H. pylori*) is one of the most common bacterial infections in humans worldwide,¹ and remains a public health problem in both developed and developing countries.² This spiral-shaped bacterium present on human gastric mucosa was first isolated in 1982 by Warren and Marshall.³ Initially it was classified as *Campylobacter pylori*, but in 1989 was included in a new genus, *Helicobacter* and renamed *Helicobacter pylori*.⁴ Since its discovery, *H. pylori* has been recognized as a major cause of gastritis, and a risk factor for duodenal ulcer disease, gastric ulcer disease, and gastric lymphoma in humans.^{5,6} Additionally, the International Agency for Research on Cancer (IARC) Working Group concluded in June 1994 that there was sufficient evidence to consider *H. pylori* a human carcinogen for gastric cancer.⁷ However, some researchers now believe that the epidemiologic evidence is contradictory and that IARC was premature in its Group 1 designation because *H. pylori* appears to play a role only in the initial steps that result in chronic inflammation (a common occurrence in much of the world), but not in the later steps that lead to carcinogenesis.^{8,9}

Modes of transmission

Despite the worldwide interest in *H. pylori* infection and its sequelae, little is known about its mode of transmission. Most epidemiologic data support a

person-to-person mode of transmission (oral-oral, fecal-oral, or gastro-oral). This mode of transmission is further supported by the discovery of *H. pylori* in saliva, dental plaque, feces, and vomitus, but the details of transmissibility of *H. pylori* infection from these sources is not known. There has also been some support for water and sewage-born transmission. Although the principle reservoir for *H. pylori* infection appears to be the human stomach, *H. pylori* has been isolated from monkeys, baboons, and domestic cats. Knowledge of the manner of *H. pylori* transmission is fundamental to the development of strategies for controlling and eventual eradication of this infectious disease.

Epidemiology

Because acute infection with *H. pylori* is generally asymptomatic, it is not possible to ascertain when infection occurs using clinical criteria.¹⁰

Epidemiologic studies of the prevalence of *H. pylori* are usually conducted by using serologic tests that detect IgG or IgA antibodies to *H. pylori* infection or by ¹³C urea breath tests (UBT) that detect urease activity produced by the bacteria.¹¹

Although a few studies have attempted to estimate incidence rates and seroconversion rates by looking at serial serologic surveys within a population, most of the information on the epidemiology and rates of *H. pylori* infection in geographically and demographically diverse populations comes from sero-prevalence studies. Studies of *H. pylori* prevalence have generally been cross-sectional in design and have included random surveys of the general population,

groups of healthy volunteers, military recruits, groups of hospitalized or institutionalized children and adults, or adults attending outpatient clinics. In addition, a number of studies have investigated intra-familial clustering of *H. pylori* infection and infection patterns in infants.

The prevalence of *H. pylori* in developing countries may reach more than 70% compared with less than 40% in developed countries. *H. pylori* infection varies widely by geographic area, age, race, and socioeconomic status (SES). Residence in a developing country, increasing age, black or Hispanic ethnicity, and lower education and income have been associated with higher *H. pylori* prevalence. Low SES during childhood, child feeding practices such as premastication of food by the mother, and poor sanitation/hygiene have been implicated.

RATIONALE

Since 1983 the National Cancer Institute (NCI) has been collaborating with the Beijing Institute for Cancer Research (BICR) and School of Oncology, Beijing Medical University to assess reasons for the exceptionally high rates of stomach cancer in Linqu, a rural county in Shandong Province in northeast China with high prevalences of precancerous gastric lesions.¹² Because Linqu county also has high rates of *H. pylori* infection,¹³ it was an ideal area to conduct research into the etiology of *H. pylori* infection and its possible mode(s) of transmission.

A population-based sample with known *H. pylori* status was assembled from 13 randomly selected Linqu villages enrolled in a randomized intervention trial to inhibit progression of precancerous gastric lesions. This stable population was used to evaluate etiologic factors that influence the prevalence of *H. pylori* infection in Shandong Province, China. The study was established to measure factors characteristics of both the environment and the host. Factors of particular interest include measures of adult and childhood socioeconomic status, source of water, personal hygiene, sharing of utensils, potential food contamination, and exposure to animals.

Research questions

The specific aims of the study were to determine whether there is an association between *H. pylori* sero-positivity and

- (1) drinking water source (i.e., shallow well, deep well, pond, river, ditch, or running water), storage, or treatment;
- (2) adult sanitation and hygiene (e.g., hand washing, use of soap, bathing, teeth brushing, and sharing of cups);
- (3) social class factors;
- (4) use of tobacco or alcohol products;
- (5) presence of gastro-esophageal reflux;
- (6) raw fruit and vegetable consumption or preparation (i.e. washing or peeling);
- (7) adult exposure to pets and other domestic animals;
- (8) childhood exposures (e.g. domestic animals, crowding, and eating premasticated food).

CHAPTER II. LITERATURE REVIEW

INTRODUCTION

Helicobacter pylori (*H. pylori*), a spiral-shaped pathogenic bacterium found on the human gastric mucosa, was first isolated by Warren and Marshall in 1982³ and soon after linked with chronic antral gastritis and peptic ulceration.¹⁴ Initially it was classified as *Campylobacter pylori*, but in 1989 was included in a new genus, *Helicobacter* and renamed *Helicobacter pylori*.⁴ Although it has been less than twenty years since its “discovery”, thousands of articles have been written about *H. pylori*, one of the most common human bacterial infections in the world.¹ Since a complete review of all the *H. pylori* literature is beyond the scope of this dissertation, this paper will include brief reviews of the microbiologic characteristics of *H. pylori*, the diagnostic tests utilized in epidemiologic studies, and the association of *H. pylori* with gastric cancer and other diseases. The primary focus will be on the epidemiology and transmission of *H. pylori* infection in adults including reviews of *H. pylori* prevalence in various countries, risk factors for *H. pylori* infection, and hypothesized modes of transmission. Studies of children will be added where appropriate to supplement the adult literature. The major emphasis will be on scientific articles from the recent literature from 1995 through mid 1999, but selected scientific papers published in peer-reviewed journals prior to 1995 will also be included.

MICROBIOLOGIC CHARACTERISTICS

H. pylori is an S-shaped or curved gram-negative rod. It has from two to six flagella that give it the mobility to withstand rhythmic gastric contractions and penetrate the gastric mucosa. It measures 2.4-4.0 μm in length and 0.5-1.0 μm in width. The principle reservoir for *H. pylori* infection appears to be the human stomach, especially the antrum. However, it does not colonize areas of the stomach where intestinal metaplasia or dysplasia are present.¹⁵ *H. pylori* contains a large urease enzyme protein that produces urease which allows the organism to survive in the acidic stomach by creating an alkaline environment. *H. pylori* produces a number of virulence factors including cytotoxin associate gene A (*cagA*) and vacuolating cytotoxin (*vacA*) that may be associated with pathologic mechanisms.^{16,17} Approximately 10-20% of the population appear resistant to chronic infection with *H. pylori*.¹⁸ Chronic infection appears to be influenced by host genetic factors or markers such as ABO blood group and Lewis blood-group antigen,¹⁹ and to differences in susceptibility to particular strains of *H. pylori*.²⁰ *H. pylori* also has a dormant coccoid form that it reverts to when faced with unfavorable environmental conditions.¹⁵

DIAGNOSTIC TESTS USED IN EPIDEMIOLOGIC STUDIES

Because acute infection with *H. pylori* is generally asymptomatic, it is not possible to ascertain when infection occurs on symptomatic or clinical grounds.¹⁰ Most epidemiologic studies of the prevalence of *H. pylori* infection usually

employ serologic tests or ^{13}C urea breath tests (UBT). Biopsy-based methods are often utilized in hospital or clinical settings.

Biopsy-based diagnostic tests

In the early 1990's, the diagnosis of *H. pylori* infection was based on either the isolation of bacteria from gastric biopsy specimens obtained from endoscopy or by identification of the bacteria on stained biopsy sections.^{14,21} Today, a positive culture obtained from endoscopy is usually augmented with a biopsy urease test (CLO test) and/or histology is often used as a “gold standard” to diagnose patients with active *H. pylori* infection.^{22,23} While biopsy/culture is 100% specific, it is not 100% sensitive; this method can evaluate only a relatively small portion of the stomach; it is not able to differentiate among the various strains of *H. pylori*; and it is highly invasive and expensive.

Serologic tests

In 1988 Perez-Perez et al, developed a non-quantitative, enzyme-linked immunosorbent assay (ELISA) to detect *H. pylori* antibodies in serum.²¹ A major advantage of this serologic test is that it allows large numbers of subjects to be screened quickly and relatively inexpensively, thus it is a good test to use in epidemiologic studies.²⁴ The prevalence of *H. pylori* (in either fresh or stored serum) is usually found by using serologic tests that detect IgG antibodies to *H. pylori* infection, although IgA and IgM antibodies have also been utilized.²⁵

There are, however, several limitations to the use of serologic tests. 1) Since no single antigen is recognized by sera from all subjects, antigen reagent preparations should contain multiple strains of *H. pylori*.^{26,27} It has also been suggested that assays based on indigenous strains may perform better.²⁸ In fact, a study in Henan Province, China found a much higher prevalence of *H. pylori* using a biopsy-based technique (85.6%) than using serology (56.2%).²⁹ The authors noted that one explanation for the large discrepancy may be that the antigen used for serology was derived from a single *H. pylori* strain isolated in Germany. However, other studies have found little difference in outcome between local and referent strains.^{30,31} 2) It is difficult to define the cut-off value that divides positive from negative subjects. One approach to improve both sensitivity and specificity, although this biases both values upward, is to include a gray or indeterminate zone for subjects whose values can not be considered truly positive or truly negative. In an epidemiologic study comparing the characteristics of positive and negative subjects, such an approach would help to minimize misclassification by not including those with equivocal results. An ROC (receiver operating characteristic) is often used to determine the effect on the test of varying the cut-off value or to compare the performance among different tests.²³ 3) The test is sensitive to changes in reagents and laboratory conditions; thus sera collected at different points in time (e.g., before and after treatment or longitudinally at yearly intervals) should be run together in the same ELISA plate. 4) Serology is not an appropriate test to use immediately following treatment for *H. pylori*. It takes a variable amount of time (anywhere from 6 months to two

years or in some cases never) for an elevated serum IgG value to return to baseline values. Some researchers suggest considering IgG values that have fallen by 50% or more six months after treatment as an indication of successful *H. pylori* eradication.^{26,32}

Breath Test

The UBT using either ^{13}C or ^{14}C is a noninvasive, non-quantitative test that determines current *H. pylori* status by detecting urease activity produced by the bacteria. This test has recently been approved by the FDA for routine use. Although not universal, several researchers now consider the breath test a non-invasive “gold standard”.^{23,26} Generally, this is a useful test to use following antibiotic treatment since gastric urease is only present in the stomach when the bacteria that make the urease are present. However, false negatives can result in treated subjects if bacteria are present in too small a number to produce detectable urease. For the small percentage of people infected by other bacteria that also produce urease (generally less than 5% in most populations), false positives can result. UBT is generally more expensive than serology, but its use in epidemiologic studies, especially studies utilizing children as study subjects, is increasing.

DISEASE ASSOCIATIONS

H. pylori is thought to be indigenous to the human population and is well-adapted to exist in the human stomach for the lifetime of its host.¹⁶ Spontaneous

eradication of *H. pylori* from the gastric mucosa, as measured by sero-reversion, is a relatively rare event; 0.1-1.1% annually.³³ Infection with *H. pylori* can result in chronic gastritis, a cellular infiltrate of immunocompetent lymphocytes, and IgA, IgG, and IgM-secreting plasma cells in the gastric mucosa.³⁴ Infection is generally asymptomatic with only a small percentage of those infected developing clinical disease.³⁵ *H. pylori* has been recognized as a major cause of gastritis and is associated with duodenal ulcer disease, gastric ulcer disease, and gastric lymphoma in humans.^{5,6}

Environmental and host genetic factors appear to be important in the progression of *H. pylori* initiated gastritis into more serious outcomes. Additionally, variation in age at acquisition of *H. pylori* has been proposed as a possible factor to explain the observation that the same organism, *H. pylori*, apparently produces different effects on the gastric mucosa that result in different clinical outcomes.⁶ According to a hypothesis proposed by Blaser et al., early age at acquisition of *H. pylori* infection may result in more intense inflammation and the early development of atrophic gastritis and subsequent risk of gastric ulcer and/or gastric cancer.³⁶ Later acquisition of infection would induce a different series of gastric changes that would favor the development of duodenal ulcer.³⁶ High rates of gastric cancer in areas where infection is common in early childhood support this hypothesis. Other host and environmental factors such as hygiene practices and diet may also play a role in the acquisition of infection and the expression of clinical disease.¹⁸

The model of gastric cancer postulated by Correa suggests a progression from chronic gastritis to gastric atrophy, to intestinal metaplasia, to dysplasia, to cancer.³⁷ In 1991, nested case-control studies of gastric carcinoma in the United States and Britain suggested that *H. pylori* may be a cofactor in the pathogenesis of gastric cancer.³⁸⁻⁴⁰ Recent investigators have continued to find an increased risk of non-cardia, gastric cancer in *H. pylori* positive compared with *H. pylori* negative subjects.^{41,42} An International Agency for Research on Cancer (IARC) Working Group felt that there was sufficient evidence in 1994 to classify *H. pylori* as a human carcinogen for gastric cancer.⁷ However, some researchers now believe that the epidemiologic evidence is contradictory and that IARC was premature in its Group 1 designation because *H. pylori* appears to play a role only in the initial steps that result in chronic inflammation (a common occurrence in much of the world), but not in the later steps that lead to carcinogenesis.^{8,9,43} Intervention studies that include treatment for *H. pylori* for subjects at different stages in the progression toward cancer will be informative in clarifying the "*H. pylori* = gastric cancer" controversy.

On the other hand, some *H. pylori* strains, particularly *cag+*, appear to be protective against adenocarcinomas of the esophagus and gastric cardia.^{41,44,45} Also, the prevalence of *H. pylori* infection appears to be lower in persons with gastroesophageal reflux disease (the major risk factor for Barrett's esophagus which is strongly associated with adenocarcinoma of the esophagus) than in controls.^{46,47}

Recently, there has been some concern that *H. pylori* infection may be associated with increased risk of coronary heart disease possibly due to a low-grade systemic inflammatory response or an increase in concentrations of circulating coagulation factors.⁴⁸⁻⁵⁰ However, the results from several prospective studies,^{48,51,52} and a meta analysis of more than 20 epidemiological studies⁵³ suggest that *H. pylori* is probably not an important contributor to coronary heart disease.

EPIDEMIOLOGY

Descriptive studies/Prevalence

H. pylori infection is ubiquitous and a public health problem in both developed and developing countries. The prevalence of *H. pylori* infection varies widely by geographic area, age, race, and socioeconomic status (SES). Because it is not possible to ascertain when infection occurs on clinical grounds, most of the information on the rates of *H. pylori* in geographically and demographically diverse populations comes from sero-prevalence studies. This has major disadvantages for epidemiologists since it is generally not possible to distinguish between factors associated with acquiring versus maintaining *H. pylori* infection.⁵⁴ Studies of *H. pylori* prevalence among adults have generally been cross-sectional in design and have included random surveys of the general population,^{13,25,48,55-70} groups of healthy volunteers,⁷¹⁻⁸¹ military personnel,⁸²⁻⁸⁴ students,^{85,86} employed workers,^{54,87,88} groups of institutionalized patients,⁸⁹⁻⁹² or patients attending hospitals or out-patient clinics.^{29,49,93-101} Included in Table II-1

is the prevalence of *H. pylori* in adult populations from various geographic areas of the world.

H. pylori prevalence in developing countries may reach 70% or more compared with 40% or less in developed countries. The acquisition rate of *H. pylori* appears to be more rapid in developing than developed countries.^{69,102} In a rural village of Linqu County, Shandong Province, China, a study of 98 children noted that nearly 70% of children aged 5-6 years were infected with *H. pylori*,¹⁰³ a rate similar to that reported for adults in that area,¹³ suggesting that most infection takes place early in childhood. Since the annual rate of seroconversion in adult populations in developed countries appears to be small (on the order of 0.2-1.0%³³) the increasing prevalence by age, seen in most developed countries, is most likely due to a birth cohort effect where older populations had higher rates of infection in the past probably due to poorer sanitation.^{69,104-107} Two recent studies of young adults, one of Israeli backpackers to Southeast Asia, South America, and Africa¹⁰⁸ and the other of military personnel deployed to the Persian Gulf for Desert Storm⁸² found much higher annual rates of seroconversion, 6.4% and 7.3%, respectively, suggesting that adults can seroconvert at higher rates than expected under unusual circumstances.

In the United States, differences by race are evident, with whites having a substantially lower seroprevalence of *H. pylori* than either blacks or Hispanics.^{49,77,79,80,83,84,109} In a study by Replogle et al in California,⁴⁹ the ORs for being *H. pylori* seropositive, given African-American and Hispanic ethnicity, were 4.1 (2.2-7.4) and 3.1 (1.6-6.2), respectively. Similar risk estimates for blacks

(OR=4.4, CI=3.0-6.3) and Hispanics (OR= 4.2, CI=2.1-8.6) were seen in a study of US Army recruits⁸⁴ and a study of US Navy and Marine Corps personnel⁸³ (OR=4.2, CI=2.9-6.0 for blacks; OR=3.9, CI=2.4-6.3, for Hispanics) Ethnic differences were also evident in New Zealand, where *H. pylori* was most prevalent in Pacific Islanders, intermediate in Maori, and least prevalent in Europeans⁸⁸ and in Belgium where Caucasians of Belgian decent (22.6%) had much lower rates than subjects who originated from the Mediterranean countries of Tunisia and Morocco (62.5%).⁹⁹ After adjusting for age and SES, the RR for Maori and Pacific Island subjects compared to Europeans were 1.4 (1.1-1.8) and 1.8 (1.4-2.2), respectively.⁸⁸ These differences in *H. pylori* prevalence by race/ethnicity may reflect differences in social and/or hygiene factors, or the wide-spread use of anti-microbials for treatment of other common infections, especially during childhood.¹¹⁰ This variability may also be explained by differences in ethnic or genetic predisposition to infections.¹⁸

Although some studies reported an excess for *H. pylori* of one gender over the other,^{49,84} overall, there are no striking gender differences in *H. pylori* prevalence. Differences in *H. pylori* prevalence by SES factors can be striking and will be presented in detail under the risk factor section. The high rates evident in institutionalized populations⁹⁰⁻⁹² will be discussed in relation to *H. pylori* transmission. Several factors require consideration when comparing rates of *H. pylori* positivity among studies. Antigenic responses to *H. pylori* may vary. If the antigens used in a serologic test were not specific to the population tested, then seroprevalence could be greatly underestimated.²⁶⁻²⁸ In addition, differences in

seroprevalence could be due to differing test characteristics (i.e., specificity), or laboratory conditions.¹¹¹

Risk Factors

In addition to determining the prevalence of *H. pylori* in various geographic areas, a number of studies included a questionnaire component designed to investigate risk factors for *H. pylori* positivity. The quality of these studies was variable and in many cases the statistical procedures utilized were not well described. Also, it was not always clear whether prevalence rates for the various groups compared were standardized either directly or indirectly to adjust for differences in age structures. The most common tests used to determine statistical significance were the chi-square test (for independence and homogeneity), Fisher's exact test, and the Student's t test. Crude and adjusted relative risks (RRs) and odds ratios (ORs) were calculated using the Mantel-Haenszel procedure and unconditional logistic regression, respectively. ORs were the most common measure of association presented.

While the OR is a legitimate measure of association in its own right, it can be used as an estimate of the RR only when the incidence of disease in the population studied is rare. In addition, the cases and noncases included in the study should be representative of all cases and noncases in the population from which the study subjects came. Because this "rare disease assumption" does not hold for *H. pylori* infection in any of the populations studied, the OR should not be considered an approximation of the RR for studies of *H. pylori*. In

addition, since these are cross-sectional studies, the outcome measure is prevalence of *H. pylori* infection at the time blood was drawn for serology, biopsy and culture were performed, or the breath test samples obtained. Thus, the measure of association in a logistic regression model is the prevalence odds ratio which compares the odds of being infected with *H. pylori* in the exposed to the odds of being infected with *H. pylori* in the unexposed/referent group.

Described in subsequent tables and text are the major factors investigated for their possible association with *H. pylori* positivity. The following topics are included: smoking, alcohol use, diet, occupational exposures, waterborne exposures, exposure to pets and other animals, hygiene practices, density/crowding, social factors, and family history of gastric disease/ulcer.

Smoking

Table II-2 presents a summary of results from several recent studies that evaluated the possible association between *H. pylori* infection and smoking. Whereas studies by Lin et al,⁵⁷ Zober et al,¹¹² Hamajima et al,¹¹³ Murray et al,⁶¹ Fontham et al,¹⁰⁹ and Bateson¹¹⁴ found *H. pylori* sero-positive subjects overall more likely to be current smokers than sero-negative subjects, their results were often not consistent by race or gender. For example, the study by Hamajima et al found an OR of 7.8 for *H. pylori* infection among current male smokers, but only an OR of 1.2 among female current smokers.¹¹³ Conversely, the study by Lin et al, found a significant association with current smoking among females (OR=2.8), but not among males.⁵⁷ The positive finding (OR=1.7) reported by

Fontham et al held for blacks (OR=3.1), but not whites (OR=0.6)¹⁰⁹ and the study by Lin et al found no association with intensity or age started smoking.⁵⁷ At least 12 recent studies found no significant association with current smoking or any other measure of tobacco use.^{13,54,63-65,68,75,88,92,115,116} and one recent study from Japan found a significant negative association with current smoking.⁸⁷ Some authors have suggested that these contradictory results may be due to uncontrolled confounding by social class^{61,87} or to differential antibiotic use since smoking appears to effect treatment success.¹¹³ While one can not rule out that an association between smoking and *H. pylori* infection may exist, such a hypothesis is not strongly supported by the current literature.

Alcohol use

The results from recent epidemiologic studies looking at the relationship between alcohol consumption and *H. pylori* infection are presented in Table II-3. None of the studies found a positive association between alcohol consumption and *H. pylori* infection, but many noted a non-statistically significant reduction in risk. The studies by Brenner et al, that 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.^{115,117,118} In two of these studies, the association was stronger for wine than beer.^{117,118} Several studies did not adequately control for potential confounding variables or did not present the actual risk estimates or prevalences in their papers, 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 due to its production of urease. Therefore, it is not surprising that the reduction in gastric pH which may accompany alcohol intake, would have little effect on the prevalence of *H. pylori*.¹¹⁹ However, alcohol is known to possess direct antimicrobial properties that appear to be greater for wine than for other types of alcoholic beverages.¹²⁰ The differing results may be due to the different methodologies employed or to real differences in either the type/amount of alcohol consumed and its effect on *H. pylori* in different populations.

Diet

Dietary associations with *H. pylori* are presented in Table II-4. Although the studies cover many different types of populations and include studies of both adults and children, there are some consistent associations that suggest that nutritional status may be related to *H. pylori* infection. Goodman et al^{121,122} and Fontham et al¹⁰⁹ found significantly reduced ORs and negative gradients in risk of *H. pylori* infection with increased consumption of fruits and/or vegetables. An intervention study by Jarosz et al found that 30% of *H. pylori* infected patients with chronic gastritis treated with vitamin C for four weeks had apparent *H. pylori* eradication compared with 0% in the control group.¹²³ Trends of decreasing risk with increasing consumption of vitamin C was observed in studies by Goodman et al¹²¹ and Fontham et al;¹⁰⁹ however, Malaty et al found high levels of vitamin C to be associated with *H. pylori* infection in twins reared apart.¹²⁴ Goodman et al

also found high levels of beta-carotene to be protective.¹²¹ In contrast, consumption of raw/uncooked vegetables was related to risk of having *H. pylori* infection in studies by Goodman et al (OR=2.0 for 3 or more servings per day)¹²² and Hopkins et al (OR=3.2).¹²⁵ The cause of this increased risk has not been determined, but may have been due to contaminated water or soil,¹²⁵ or contamination by a vector such as the housefly.¹²⁶ The role of food prepared under less than ideal sanitary conditions as a possible mechanism of *H. pylori* transmission, was suggested by Begue et al who found elevated risks for consumption of food obtained from street vendors in Peru.¹²⁷

Occupational exposures

Occupational exposures were considered by several researchers (Table II-5) investigating whether people working in certain occupations with potentially greater exposure to *H. pylori* had increased prevalence of infection. Bohmer et al in a study of inhabitants of institutions for the intellectually disabled found that most of the patients were seropositive (82.8%).⁹⁰ He also reported a higher rate of seropositivity (31.6%) among employees such as the nursing staff with intensive contact with institutionalized inhabitants than among employees such as medical staff, speech-trainers, secretarial staff, and drivers with little or no direct contact (14.1%).⁹⁰ Risk of infection from potential exposure to these bacteria in waste water was investigated by Friis et al in a study of Swedish sewage workers.¹²⁸ They found that seroprevalence did not differ between sewage workers and other age-SES-location-matched municipal workers.¹²⁸

Shelly and Haddadin reported that *H. pylori* infection did not appear to be an occupational hazard for anesthesiologists in the U.S. even though they are potentially exposed to secretions from the stomachs of their patients while performing endotracheal intubations.¹²⁹

Since *H. pylori* has been cultured from both saliva and dental plaque, studies were conducted by Lin et al,¹³⁰ Banatvala et al,¹³¹ and Malaty et al¹³² to determine whether dentists, dental nurses, and dental workers were at increased risk for *H. pylori* infection. Dentists and dental workers were not found to have higher rates of seropositivity than controls in any of these studies. When Lin et al compared the prevalence of *H. pylori* in dentists and dental nurses to that in endoscopists and endoscopy nurses they found endoscopists (80%) to have significantly higher rates than dentists (21%), but the rates in the two types of nurses were not significantly different.¹³⁰

There has been conflicting data regarding the prevalence of *H. pylori* in endoscopy staff. In a group of eleven studies that focused on this occupation, four found that medical staff who performed endoscopies had a higher prevalence of *H. pylori* than medical staff who did not¹³³⁻¹³⁶ and three reported the prevalence of *H. pylori* to be significantly higher in endoscopists compared to population controls.¹³⁷⁻¹³⁹ The results of a recent study by Mones et al in Spain that used the UBT indicated that medical professionals, more specifically gastroenterologists performing endoscopies, do not have a higher prevalence of *H. pylori* infection than do healthy controls.¹⁴⁰ Braden et al found no increased risk of *H. pylori* infection in endoscopy staff (physicians and nurses) compared to

the general medical staff (physicians and nurses), but did find a risk for all medical staff compared to controls.¹⁴¹ On the other hand, Rudi et al found that neither exposure to patients in an acute care hospital (Group 2) nor exposure to endoscopic procedures (Group 3) increased the rate of *H. pylori* infection.¹¹⁶ In a study published by Nishikawa et al, *H. pylori* overall seroprevalence among endoscopists and endoscopy nurses did not differ significantly from that in healthy controls, although the prevalence among younger (<40 years old) endoscopy staff was significantly higher than among younger controls (24.4% vs 12.0%, $p < 0.05$).¹⁴² It is possible that differences in medical practices, including the use of gloves and other protective equipment, may be responsible for these discrepancies. To obtain a more definitive answer, a large prospective study of endoscopists would be required.¹⁴³ The probable risk for endoscopists, but not dentists suggests that gastric mucous may be a better medium for transmission of *H. pylori* than saliva.¹³³

Waterborne Exposures

Water has been suggested as a possible source of *H. pylori* infection. Recent studies that accessed the relationship between *H. pylori* and waterborne exposures are listed in Table II-6. Studies in Columbia, rural China, and Lima, Peru found that water source may be related to risk of *H. pylori* infection.^{13,122,144} Three waterborne factors were linked to higher risks of *H. pylori* infection in Colombian children, drinking water from a stream, swimming in a stream, and swimming in a swimming pool.¹²² Klein et al found that the water supply in Lima,

Peru may be vulnerable to bacterial contamination, especially if it is stored in a cistern or utilized through central community water taps.¹⁴⁴ Although not significantly different due to the small percentage of subjects reporting use of a pond or ditch as a source of drinking water, Zhang et al¹³ found a substantially higher seroprevalence of *H. pylori* (88%) among those obtaining their water from a surface-water source compared to subjects who obtained their water from a well (73%). Other studies found no differences by water source.^{67,74,76,125,145} These studies included work by Teh et al who found no excess risk of *H. pylori* for subjects who obtained their water from the river⁶⁷ and Hopkins et al who found no significant excess risk for either swimming near contaminated beaches or bathing in local rivers, irrigation ditches, or lakes.¹²⁵

Exposure to Animals/Pets

Table II-7 identifies studies that assessed whether exposure to pets or other domestic animals, especially cats, was related to infection with *H. pylori*. A study in Germany by Rothenbacher et al found that adults who owned a cat as a child had a significantly higher prevalence of *H. pylori*.⁸⁵ Most studies, however, found no association with having a pet either during childhood or as an adult.^{78,85,93,97,127,146} and two studies in the US found pet owners to have significantly lower rates of *H. pylori* infection.^{80,147} Although it is possible that lower rates in pet owners may be due to confounding by social class, one large population-based study in Canada that adjusted for social class found no association between pet ownership and a history of peptic ulcer disease.¹⁴⁸

Exposure to sheep was implicated in two recent studies.^{122,149} In a study of children from rural Columbia, Goodman et al reported that children who “played with sheep” had a higher risk of *H. pylori* infection (OR=4.5).¹²² A study by Dore et al revealed that the prevalence of *H. pylori* was significantly higher among Sardinian shepherds occupationally exposed to sheep than among their non-exposed family members or among Sardinian blood donors.¹⁴⁹ Overall, these studies only weakly support a role for domestic pets or other animals in the transmission of *H. pylori*. In light of the interesting findings with sheep, further investigation of exposure to other domestic animals is warranted.

Hygiene Practices

Presented in Table II-8 are studies that assessed the relationship 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 higher seroprevalence of *H. pylori*.¹⁵⁰ Some of these practices included having no WC/bathroom,^{73,93} refrigerator,⁹³ or hot water supply^{73,93,100} in the house when the subject was a child; sharing cups as children and having mothers who didn’t use soap when they washed their hands;¹²² having mothers prechew the food for their young children;¹⁵¹ use of chopsticks;⁶³ not usually washing ones hands after going to the toilet;⁶² and living in a relatively small area with extremely limited sanitary facilities.¹⁵² Other hygiene practices during adulthood such as sharing towels⁹⁷ and type of toilet/bathroom facilities^{67,73,78} were not strongly related to *H. pylori* infection.

Density/Crowding

A variety of density measures both during childhood and adulthood are presented in Table II-9. Some measure of overcrowding or high density during either childhood or as an adult such as living in a crowded environment,¹²⁴ sibship size,^{67,86,87,93,97} number of persons or children in the home,^{49,55,62,73,74,122} number of persons per room,^{60,73,78,93,100,153} crowding index,^{77,98,147} having to share a room or bed with a parent,^{78,85,153,154} or living in a crowded space in a submarine¹⁵² was consistently related to *H. pylori* positivity in all recent studies that evaluated this factor. In Japanese self-defense officials, those of lower rank stay in group barracks until they marry, thus the finding of a higher prevalence of *H. pylori* in this group may reflect crowding during young adulthood.⁵⁴ The positive association of *H. pylori* with high density environments, especially during childhood, suggest that crowded household quarters may facilitate the transmission of infection among siblings and other family members. This finding is consistent with the data on intrafamilial clustering of *H. pylori* presented in Table II-13.

Social Factors

Studies assessing the relationship between *H. pylori* infection and social factors are presented in Table II-10. Social factors were independently associated with *H. pylori* status in a variety of studies throughout the world. The most commonly used measures were SES-based occupation (usually based on

the Registrar General's categories that separated jobs into professional occupations, managerial occupations, skilled occupations, semiskilled occupations, and unskilled occupations), education, and income. Occupation-based SES was associated with *H. pylori* seroprevalence in studies in Ireland,^{61,66,72,153} Italy,⁹³ South Wales,⁴⁸ New Zealand,⁸⁸ and the UK, in a study by Webb et al,⁷⁸ but not one by Mendall et al.¹⁰⁰ Income was related to *H. pylori* infection in Australia,⁵⁷ Brazil,⁵⁹ Russia in children, but not adults,⁹⁸ and the US,^{84,147} but not in Taiwan.⁶⁷ Low education was significantly related to higher risk of *H. pylori* in many,^{49,60,62,63,68,79,80,85} but not all studies.^{65,86,88} Low SES as defined differently by various investigators was also associated with a higher seroprevalence of *H. pylori* in most studies in which it was evaluated.^{54,74,77,124,125} In Japanese self-defense workers, low rank was associated with less education and lower incomes than the middle and high ranks.⁵⁴

Although social class has consistently been linked to *H. pylori* in a number of studies, it is probably not a risk factor in itself, but rather represents differences in a number of lifestyle factors that are related to risk of *H. pylori* infection. In other words, low social class whether measured by occupation, education, or income is a surrogate marker for a set of negative environmental characteristics such as poor housing, poor hygiene, unemployment, access to medical care, stressful home or work environments, poor nutrition, or exposure to other infectious agents that may increase the likelihood of an individual being exposed to *H. pylori* and may render an individual more susceptible to *H. pylori* infection if exposed. In addition, social class measures reflect differences in

economic, political, or cultural indicators that may differ between racial and ethnic groups.

Family History of Gastric Disease/Ulcer

The seven studies that evaluated the relationship between *H. pylori* infection and family history of gastric disease/ulcer are presented in Table II-11. The results are generally inconclusive. Whereas the study by Brenner et al found a significantly elevated risk in children whose mothers, but not fathers had ulcer disease,¹⁵⁵ the study by Kikuchi et al found a significantly elevated risk in public service workers whose father, but not mother had a history of ulcer disease.⁸⁷ The study by Gasbarrini et al found significantly elevated risks of *H. pylori* infection for subjects whose siblings, but not parents had a history of ulcer disease⁶⁴ and the study by Martin-de-Argila et al reported a significantly greater percentage of *H. pylori* positive than *H. pylori* negative subjects with a first degree relative with peptic ulcer disease.⁷⁵ The other three studies listed in Table II-11 found no significant association between *H. pylori* infection and family history of ulcer.^{65,73,145}

POSSIBLE ROUTES OF TRANSMISSION

Person-to-Person Transmission

Institutionalized Populations

Studies assessing the relationship between *H. pylori* infection and institutionalized populations are listed in Table II-12. The study of

institutionalized young mentally and physically handicapped adults in Austria by Lambert et al was one of the first to investigate the prevalence of *H. pylori* infection in institutionalized patients as a way to evaluate the possibility of person-to-person transmission of *H. pylori*.⁹² He found a much higher prevalence of *H. pylori* in residents in 1989 (75%) compared to age-sex-matched controls (23%) and to stored serum taken from the same subjects in 1977 (34%). The prevalence of infection was related to length of stay in 1977, but not in 1989.⁹² Fifty-one *H. pylori* negative subjects in 1977 were positive in 1989, for an annual seroconversion rate of 7.4%,⁹² much higher than the rate of 0.2-1% typical of most developed countries.

Significantly higher rates of *H. pylori* infection were also seen in other institutionalized populations. In a study in England by Harris et al, *H. pylori* seroprevalence was higher in adult residents of a hospital for people with severe learning difficulties (87%) than for controls from the local community (41%).⁹¹ However, there were no differences in *H. pylori* seroprevalence by age or duration of stay among the residents.⁹¹ Bohmer et al, found a strong association between *H. pylori* positivity and length of institutionalization among adult inhabitants of two large institutes for the intellectually disabled in the Netherlands.⁹⁰ Risks were elevated for subjects with IQs<50 and those who regurgitated their food suggesting that these factors may increase the spread of *H. pylori* due to less than adequate sanitary practices among these residents.⁹⁰ In a study among Japanese patients institutionalized with neurologic impairments, Kimura et al found *H. pylori* to be significantly more prevalent

among institutionalized patients (81.1%) than among patients living at home (20%) or healthy Japanese.⁸⁹ Percent positivity was found to increase with both age and duration of institutionalization.⁸⁹ In addition, 18 of 38 seronegative patients (47.4%) seroconverted after one year.⁸⁹ For this population of patients, possible routes of transmission include salivary secretions or fecal-oral contamination. Among institutionalized schizophrenic patients in Belgium, the risk of *H. pylori* infection was elevated (OR=3.0, CI=1.4-7.3) compared with volunteer blood donor controls.¹⁵⁶ Again, the prevalence in patients increased with length of stay in the institution.¹⁵⁶

Institutionalized children in Hong Kong with profound neurodevelopmental disabilities were found by Lewindon et al to have significantly higher rates of *H. pylori* infection (55.4%) than age-matched controls (8%).¹⁵⁷ In a group of normal children in Russia, Malaty et al found the prevalence of *H. pylori* infection to be greater in children from orphanages and communal apartments (64%) than in children with families (40%).⁹⁸ Although a common source of exposure can't be ruled out in several of these studies, they do lend support to the hypothesis that most *H. pylori* infection is transmitted person-to-person.

Familial Exposures

A number of studies listed in Table II-13 looked at the relationship between *H. pylori* infection and intrafamilial clustering of *H. pylori*. Studies in Italy,¹⁵⁸ Belgium,¹⁵⁹ Austria,¹⁶⁰ and Canada¹⁶¹ noted an increased prevalence of *H. pylori* infection in family members of *H. pylori* positive children compared to

family members of *H. pylori* negative children and/or age-matched control groups. Excess risks were found in both parents and siblings of infected children with no consistent differences apparent by gender of parent. Similarly, a higher prevalence of *H. pylori* infection in children who lived in households with *H. pylori* positive relatives was reported in studies conducted in Germany,¹⁵⁵ China,^{162,163} Poland,¹⁵⁰ and Russia.⁹⁸

Several studies evaluated intraspousal transmission of *H. pylori* infection. Singh et al in India¹⁶⁴ and Brenner et al in Germany¹⁶⁵ found a strong relationship between partner's infection status in studies conducted among couples from the general population and among employees of a health insurance company and their partners, respectively. In the Indian study, 60% of *H. pylori* positive spouses had seroconverted at the one-year follow-up.¹⁶⁴ The German study reported an increase in risk with the time lived with an infected partner.¹⁶⁵ Georgopoulous et al in Greece¹⁶⁶ and Parente et al in Italy¹⁶⁷ found higher rates among spouses in *H. pylori* positive than *H. pylori* negative patients and/or matched controls. However, in the Italian study, rates of positivity differed substantially (73% and 33%, respectively) for spouses and controls aged 20-34, but were the same (75% and 74%, respectively) for spouses and controls older than 50.¹⁶⁷ This is similar to a study by Ma et al conducted in an area of China where the overall prevalence rate of *H. pylori* in adults was high (68%) and differed little by age.¹⁰³ Ma et al reported no significant difference in the seroprevalence of *H. pylori* in spouses of *H. pylori* positive and *H. pylori* negative subjects.¹⁰³ Several studies performed DNA fingerprinting to determine the

specific strains of *H. pylori* harbored by family members.^{166,168-170,247} They found the same strain of *H. pylori* to be present in spouses, siblings, and parents.^{166,168-170,247}

These studies taken as a whole strongly support the concept of intrafamilial clustering of *H. pylori* infection. They suggest that either person-to-person transmission occurred in these families possibly due to close interpersonal contact, or that family members shared a genetic predisposition to *H. pylori* infection, or that family members were exposed to a common source of infection.

Oral-oral Route

Many scientists have hypothesized that the oral-oral route is the most likely method of *H. pylori* transmission, especially in developed countries. Elevated prevalence of *H. pylori* within families and institutionalized populations provide support for this route of transmission. However, it is not clear whether *H. pylori* is a constant or intermittent inhabitant of the oral cavity.^{171,172} Because there are numerous bacteria in the oral cavity that show urease activity, the specificity of urease-based tests may be too low to be useful in detecting oral *H. pylori* infection.^{173,174} Polymerase chain reaction (PCR) is a very sensitive assay technique, but positive results can not confirm the viability of the bacteria and whether it is able to transmit disease.¹⁷⁴ Microbiological culture of *H. pylori* is still the recognized “gold standard” for the diagnosis of infection.¹⁷⁵ The most likely

routes of oral infection include saliva, dental plaque, and refluxed gastric contents or vomit.

Recent studies by Leung et al,¹⁷⁶ Li et al,^{177,178} Namavar et al,¹⁷⁹ Shimada et al,¹⁸⁰ and Mapstone et al¹⁸¹ have detected *H. pylori* DNA in saliva in *H. pylori* positive subjects using PCR assays. *H. pylori* has been successfully isolated from saliva by Ferguson et al¹⁸² and Pytko-Polonczk et al.¹⁷⁴ However, several other researchers were not successful.^{179,183-185} Evidence that *H. pylori* might be transmitted by saliva comes from a study by Megraud who found a higher risk of *H. pylori* associated with premastication of food by mothers in Western Africa who feed their infants (OR=2.9, CI=0.9-9.0).¹⁵¹ In addition, Chow et al reported that Chinese immigrants in Australia who used chopsticks to eat from communal dishes had a significantly higher prevalence of *H. pylori* infection (64.8%) than those who did not (42.3%).⁶³ However, Leung et al were able to detect *H. pylori* DNA in both saliva and chopstick washings in only 1/15 subjects with *H. pylori* DNA in saliva and in only 1/45 subjects with *H. pylori* documented by UBT, suggesting that this mode of transmission is probably not common.¹⁷⁶ A study by Luzzza et al in rural Italy found no evidence to suggest that *H. pylori* and Epstein Barr Virus (the etiologic agent of infectious mononucleosis, a common infection that is transmitted by the oral-oral route) share a common mode of transmission.⁹³

Several investigators have been successful in isolating *H. pylori* from dental plaque of infected subjects, but the percent positive has varied from 1% to 88%.^{173,174,185-187} Other investigators have not been able to culture *H. pylori* from

dental plaque,¹⁸⁸⁻¹⁹⁰ or to detect its presence using an indirect immunoperoxidase technique.¹⁹¹ The inability of some researchers to culture *H. pylori* could be due to the presence of the viable but unculturable coccoid form of *H. pylori*, the presence of too small a number of organisms to detect, or the presence of too many other types of bacteria in the mouth that inhibit the growth of *H. pylori*.¹⁷² Peach et al in a study of Australian adults selected from electoral rolls found that positive *H. pylori* status was associated with high plaque score, OR=1.7 (1.1-2.7) and visiting the dentist less than once a year, OR=4.4 (0.8-23.0).⁶² Hardo et al in a study of dyspeptic patients in the UK found no association between *H. pylori* infection and time between visits to the dentist, number of times per week brushed teeth, oral hygiene index, and periodontal status scale.¹⁸⁸ In a study of patients with duodenal ulcer and hospital employees with and without occupational exposure to *H. pylori*, Dore-Davin et al found no correlation between *H. pylori* status as determined by PCR on saliva and dental plaque and status as determined by UBT.¹⁹²

Because *H. pylori* acquisition rates are higher in children, especially in countries with less than adequate sanitation, Axon has suggested that *H. pylori* is an epidemic disease of childhood transmitted through a gastro-oral route by mucousy vomit.¹⁹³ Others feel that recovery of *H. pylori* in the oral cavity is most likely associated with gastroesophageal reflux or regurgitation of stomach contents.^{172,185,194} Additional support for transmission of *H. pylori* via stomach contents comes from a case report of possible transmission of *H. pylori* infection between adults by ingestion of vomit during mouth-to-mouth resuscitation¹⁹⁵ and

from a study by Bohmer et al who found a higher prevalence of *H. pylori* among institutionalized intellectually disabled adults who regurgitated their food (OR=2.0, CI=1.1-3.6).⁹⁰

Fecal-oral Route

Another possible route of *H. pylori* transmission is the fecal-oral route. *H. pylori* DNA has been detected from feces of infected subjects by some researchers,^{179,180,196} but not others.¹⁹⁷ Recently, Gramley et al found detectable *H. pylori* DNA in the feces of 73% of infected subjects.¹⁹⁶ Isolation of *H. pylori* by fecal culture, has been performed by a number of investigators from around the world.^{179,198,199} However, isolation of *H. pylori* from feces has been problematic for some researchers, especially for those unable to obtain fresh fecal samples. Delay in processing the samples could have resulted in the small number of *H. pylori* organisms present being overgrown by other fecal bacteria or possibly the transformation of *H. pylori* into its viable but not culturable coccoid form.²⁰⁰ Studies by Hazell et al in China²⁰¹ and Webb et al in the UK²⁰² noted serum antibodies to *H. pylori* and hepatitis A, a sensitive marker of fecal-oral exposure, but did not find strong evidence supporting community-wide fecal-oral spread of *H. pylori* via food or water.^{201,202} Other studies that found elevated risks of *H. pylori* infection associated with lack of hand washing suggest that the fecal-oral route of transmission may be possible in some populations.^{62,122}

Waterborne Transmission

Experimental tests have shown that under physical or chemical stress *H. pylori* is able to convert into its viable, but non-culturable coccoid form.²⁰³ It has also been demonstrated that *H. pylori* can live for several days in milk and tap water in its infectious bacillary form^{204,205} and in river water for several months in its coccoid form.²⁰⁶ Although researchers have failed to convert coccoid to bacillary form in culture,²⁰⁷ it has not yet been determined whether *H. pylori* can revert from its coccoid to its infectious form in humans.²⁰⁸ A study by Cellini et al found evidence of reversion in mice,²⁰⁹ however, Eaton et al was not able to replicate this in gnotobiotic pigs.²¹⁰ Support for waterborne transmission comes from epidemiologic studies conducted in Columbia,¹²² rural China,¹³ and Lima, Peru¹⁴⁴ that found that water source may be related to risk of *H. pylori* infection. The finding of *H. pylori* positive drinking and sewage water samples by PCR assays in Peru provide additional evidence that waterborne transmission may be important in areas of the world with less than adequate water quality.^{204,211} The possible role of waterborne transmission of *H. pylori* is further supported by a study by Baker and Hegarty in Pennsylvania.²¹² They reported a strong association ($p < 0.02$) between consumption of well water, contaminated by *H. pylori*, and *H. pylori* infection in those consuming the water.²¹² In addition, an analysis of surface and ground water samples in Pennsylvania and Ohio found 61% of the samples to be contaminated with *H. pylori*.²¹³

Zoonotic/Vectorborne Transmission

Although the principle reservoir for *H. pylori* infection appears to be people, *H. pylori* has been isolated from non-human primates and domestic cats.^{20,214,215} Human *H. pylori* has been shown to infect monkeys under controlled laboratory conditions.²¹⁴ However, even if *H. pylori* occurs naturally in monkeys, they are unlikely to represent a major route of transmission of *H. pylori* infection to humans since close contact between non-human primates and humans in most of the world is limited.²¹⁴

Handt et al were the first to report isolation of *H. pylori* from domestic cats.^{215,216} Their laboratory was able to experimentally infect naïve cats with *H. pylori*, to culture *H. pylori* from feline salivary and gastric sections, and to find *H. pylori* DNA in feline feces and dental plaque.^{217,218} Although PCR cannot determine the viability of the *H. pylori* organism, these studies raised the possibility that *H. pylori* could be transmitted from cats to humans via saliva, vomit, or feces. Recently, El-Zaatari et al found no evidence of *H. pylori* infection when they examined 25 stray cats.²¹⁹ Instead, they reported that *H. helmanni* was the organism responsible for the chronic gastritis seen in these cats.²¹⁹ The epidemiologic evidence is also inconsistent. A significant association between *H. pylori* infection and cat ownership in childhood was reported in a study of adults in Germany,⁸⁵ but not in the U.K.²²⁰ These results suggest that *H. pylori* infection is probably uncommon in cats and is probably not a major public health problem for cat owners.

Gnotobiotic pigs have been successfully infected with *H. pylori*, but there is currently no convincing evidence that swine are a reservoir for *H. pylori*, even though the monogastric pig stomach has anatomical and physiologic similarities with the human and non-human primate stomachs.²¹⁴ There has also been a suggestion in the epidemiologic literature that handling sheep may be a risk factor for *H. pylori* infection.^{122,149} In addition, in Sardinia, 32 of 32 sheep were seropositive for *H. pylori*, *H. pylori* DNA was detected in the mucosal strips from the stomach of 3 of 10 sheep or lambs, and *H. pylori* DNA was present in raw sheep milk.^{149,221}

The most recent reservoir suggested for *H. pylori* transmission is the housefly. In controlled experimental studies, Grubel et al found that houseflies infected with *H. pylori* in the laboratory could harbor viable *H. pylori* bacteria in their intestines as well as on their body hairs.²²² In further investigations, these researchers captured wild flies from rural, agricultural, and metropolitan areas in the U.S., Japan, Poland, and Egypt.¹²⁶ They found high levels of *H. pylori*-contaminated flies in Egypt (33%) and Poland (57%), areas where there is a high seroprevalence of *H. pylori* in children.¹²⁶ However, the investigators also uncovered high levels in California (38%), an area where the seroprevalence of *H. pylori* is not very high.¹²⁶ The possibility that flies could transmit *H. pylori* from contaminated feces to food or mucosal surfaces is indirectly supported by a number of epidemiologic studies in which subjects without indoor bathroom facilities, especially during childhood, had a higher seroprevalence of *H. pylori* than subjects with indoor facilities.^{73,78,93,122} However, evidence is lacking that *H.*

pylori can be isolated from flies that have been in contact with *H. pylori* infected feces and that *H. pylori* can be transmitted from contaminated flies to food in a quantity sufficient to cause active infection in humans.²²³ In a recent experimental study by Osato et al²²⁴ researchers were unable to recover *H. pylori* from houseflies that were fed human feces infected with *H. pylori*. Therefore, it seems less likely that the domestic housefly serves as a major vector for *H. pylori* transmission.

Iatrogenic Transmission

Endoscopy is a common medical procedure used to diagnose and manage gastrointestinal disease. Because of the complex structure of the endoscope and the difficulty in disinfecting them, the possibility of iatrogenic infection in patients following endoscopy is a potential risk factor not only for *H. pylori*, but also for other infectious diseases such as hepatitis B, hepatitis C, tuberculosis, and possibly HIV.^{225,226} In fact, nosocomial transmission of *H. pylori* is the only proven mode of transmission.²²⁶ According to Tytgat, the rate of iatrogenic infection may approximate 4 per 1000 endoscopies (0.4%) when the prevalence of *H. pylori* in the endoscoped population is around 60%.²²⁷ He suggests that the rate of iatrogenic infection may reach 1% in areas of the world utilizing improper disinfection techniques.²²⁷ The retrospective study by Langenberg et al in the Netherlands had a rate of 1.1% for *H. pylori* negative patients to develop iatrogenic infection from endoscopy when alcohol was used as the disinfectant instead of gluteraldehyde.²²⁸

Even before the “discovery” of *H. pylori*, Ramsey et al reported that 17 of 37 (45.9%) subjects participating in studies of gastric acid secretion became hypochlorhydric²²⁹ (a condition that occurs during the acute phase of *H. pylori* infection following gastric biopsy²³⁰). Iatrogenic infection is also responsible for postendoscopic acute gastric mucosal lesions (sub-acute *H. pylori* infection) found in Japan that often follows endoscopy as a result of the ineffective disinfection methods utilized.^{194,228,230,231} Proper cleaning requires use of a detergent and brushing (and often use of an enzymatic cleaner) to remove blood, mucous, and tissue from the endoscope channels prior to disinfection.^{225,227} In 1990, the working party to the World Congresses of Gastroenterology recommended that the endoscope be soaked in 2% activated glutaraldehyde for at least 5-10 minutes, 10 minutes being sufficient to prepare the instrument for use in any patient about to undergo endoscopy.²³²

Rohr et al investigated the prevalence of *H. pylori* in patients attending hospitals in San Paulo, Brazil (most of which did not follow CDC endoscopy cleaning guidelines).²³³ He found *H. pylori* positive patients to have had a greater number of prior endoscopies than *H. pylori* negative patients, although the differences were not statistically significant.²³³ A study by Kaneko et al in Japan found that careful cleaning of the endoscope with glutaraldehyde, but not alcohol, may be sufficient to avoid iatrogenic transmission of *H. pylori* infection.²³⁴ Biopsy forceps typically penetrate the gastric mucosa and are difficult to clean.²²⁷ Therefore, sterilization of the forceps or preferably use of disposable forceps is essential.²²⁷ Studies in Japan¹⁹⁴ and Taiwan²³⁵ suggest that mechanical washing

of the endoscope is superior to manual washing in preventing iatrogenic spread of *H. pylori*, especially in situations where recommended disinfection procedures may not be routinely followed. Fantry et al, in a study in Baltimore, Maryland found that endoscopes were frequently contaminated (61% of the time) with *H. pylori* following procedures on *H. pylori* infected patients, but that risk of infection was minimal if proper disinfection methods were utilized.²²⁶ A study of disinfection procedures for endoscopes in 20 Japanese hospitals found that *H. pylori* infection following endoscopy was due to inadequate disinfection procedures rather than to any resistance of *H. pylori* to disinfectants.²³⁶

SUMMARY OF PUBLISHED LITERATURE

H. pylori is a ubiquitous bacterium with approximately 50% of the world's population estimated to be infected.²³⁷ The prevalence of *H. pylori* infection varies widely by geographic area, age, race, ethnicity, and socioeconomic status. Rates appear to be higher in developing than developed countries with most of the infections occurring during childhood. *H. pylori* causes chronic gastritis and has been associated with several serious diseases of the gastrointestinal tract, including duodenal ulcer and gastric cancer. Since its "discovery" in 1982 by Warren and Marshall, *H. pylori* has been the topic of extensive research.

A number of studies have included questionnaire components that have been used to investigate factors possibly related to the etiology of *H. pylori* infection. The majority of recent studies have found no significant association with current smoking or any other measure of tobacco use. Depending upon the

population studied, alcohol consumption appears to either have no association with *H. pylori* infection or to have a slight protective effect. Adequate nutritional status, especially frequent consumption of fruits and vegetables and vitamin C, appear to protect against infection with *H. pylori* although the mechanism of action is not known. In contrast, food prepared under less than ideal conditions, or exposed to contaminated water or soil may increase risk of *H. pylori* infection. Although the evidence is still contradictory, the occupational group that appears to be at greatest risk of becoming infected with *H. pylori* is endoscopists. Since risks are not elevated for dentists, this suggests that gastric mucous may be a better medium for transmission of *H. pylori* than saliva. Drinking water source has been related to risk of *H. pylori* infection in some but not all studies. Since *H. pylori* has recently been detected in surface and shallow ground water, this is an area that deserves further investigation. Most studies have found no association between adult pet ownership and risk of *H. pylori* infection. Although several studies have suggested sheep as a possible source of *H. pylori* transmission, this hypothesis deserves additional investigation. Since only a few studies have investigated childhood exposure to animals, this also warrants further research. Overall, inadequate sanitation practices such as having no bathroom, refrigerator, or hot water supply during childhood and improper handwashing practices after using the toilet appear to be related to higher prevalence of *H. pylori* infection. All recent studies that evaluated crowding/high density living conditions using measures such as sibship size, number of persons in the home or per room, crowding index, or having to share a room or

bed with a parent, noted a positive association with prevalence of *H. pylori*. This suggests that crowded conditions may facilitate the transmission of infection among family members and is consistent with data on intrafamilial and institutional clustering of *H. pylori* infection. Low social class, whether measured by education, occupation, or income, has been consistently linked to *H. pylori* infection. Although probably not a risk factor in itself, social class represents differences in lifestyle factors that may be related to the risk of *H. pylori* infection. Only a small number of studies have evaluated the relationship between family history of gastric disease and *H. pylori* infection and the results have been inconclusive.

Understanding the route of transmission of *H. pylori* is important if public health measures to prevent the spread of *H. pylori* are to be implemented. Nosocomial transmission of *H. pylori* is the only proven mode of transmission. Difficulty in adequately disinfecting the endoscope is the major cause of iatrogenic infection in patients following endoscopy. For the general population, the most likely mode of transmission is person to person, either by the oral-oral route (through saliva or vomitus) or possibly by the fecal-oral route. The person-to-person mode of transmission is supported by the higher incidence of infection among institutionalized children and adults and the clustering of *H. pylori* infection within families. Also lending support to this concept is the detection of *H. pylori* DNA in saliva, dental plaque, gastric juice, and feces. The gold standard, isolation of *H. pylori* by culture from these sources has been performed by some investigators, but has proven to be problematic for others. Waterborne

transmission, probably due to fecal contamination, may be an important source of infection, especially in parts of the world where untreated water is common. Recent studies in the United States have linked clinical *H. pylori* infection with the consumption of *H. pylori* contaminated well water. This is an area of research worthy of further investigation. Although the principle reservoir for *H. pylori* is people, *H. pylori* has been isolated from non-human primates and domestic cats. However, even if *H. pylori* occurs naturally in non-human primates, it is unlikely to be a major route of transmission since in most of the world direct contact between humans and monkeys is limited. Although *H. pylori* has been isolated in domestic cats, additional research in this area has suggested that *H. pylori* is probably uncommon in domestic cats and thus is probably not a major concern for cat owners. The most recent reservoir suggested for *H. pylori* transmission is the housefly. However, evidence is lacking that *H. pylori* can be transmitted to humans from flies that have been in contact with *H. pylori* infected feces. Nevertheless, the hypothesis is appealing since flies are known to carry many other infectious agents. Knowledge of the epidemiology and mode of transmission of *H. pylori* are important to prevent its spread and may be useful in identifying high risk populations, especially in areas with high rates of gastric lymphoma, gastric cancer, and gastric ulcer.

CHAPTER III. METHODS

STUDY POPULATION

In 1989, as part of a previous epidemiologic study conducted by the National Cancer Institute (NCI) and the Beijing Institute for Cancer Research (BICR) in Linqu county, Shandong Province, China, the names of all residents aged 35-64 were transcribed from village lists in 14 villages selected at random from five townships.¹² The names of all individuals aged 35-39, from the same 14 villages, were added to the study roster in 1994, in preparation for a joint NCI/BICR randomized intervention trial to inhibit progression of precancerous gastric lesions. Chinese health officials visited the 4,326 individuals on the roster, explained the study, and invited their participation in a screening program that included gastroscopy, gastric biopsy, and phlebotomy.

Of the 4,326 possible participants, 210 subjects were excluded from the study because they were not medically eligible for endoscopy. Specific reasons for exclusion included: a bleeding disorder, any type of cancer, a history of liver disease, renal disease, chronic obstructive pulmonary disease, or other life-threatening illness. Also excluded from the trial were 226 subjects who refused endoscopy (afraid, felt there was no need, or couldn't tolerate the procedures) and all the eligible subjects from one village (291) because they participated in an earlier pilot study of the intervention agents. As a result, the study population for the intervention trial consisted of 3,599 subjects from 13 villages who had undergone gastroscopy, gastric biopsy, and phlebotomy.

A 5-ml blood sample was drawn from each potential study subject prior to the beginning of the trial. Serum was separated and aliquoted in the field, stored immediately at -20 degrees C, and then transferred to a -70 degrees C freezer at BICR. One aliquot of serum (0.5 ml) was tested in Beijing by Dr. Lian Zhang, BICR, for IgG antibodies to *H. pylori*. *H. pylori* strains cultured from gastric biopsies of two patients in Linq County were used to provide the antigenic preparation for serology. The two strains were grown on blood agar plates for 48 hours and then harvested in distilled water. The cell suspensions were sonicated six times for 30 seconds each. The protein concentrations were measured by the Markwell et al. modification of the Lowry method (1978) and diluted to 10 µg/ml, and the soluble material from the two strains was pooled for an ELISA procedure.²¹ Microtiter plates were prepared using 1 µg total protein per well. All assays were performed on coded samples in duplicate and then repeated. Each microtiter plate was read at 414 nm. Each subject was considered positive for *H. pylori* infection if the ELISA absorbance reading for IgG or IgA was 1.0 or higher, a cutoff based on examination of the distribution of readings in relation to a group of uninfected persons and reference sera. Although the sensitivity and specificity of this *H. pylori* antigen preparation derived from two Chinese strains was not determined, a similar preparation from five Chinese strains using a cut-point of 0.514 yielded a specificity of 94.9% based on 39 *H. pylori*-negative (by histology, culture, and direct urease assay) U.S. children.³⁰ The sensitivity for the five-strain antigen in 132 Chinese subjects with biopsy-confirmed *H. pylori* infection was 100%.³⁰ A somewhat lower

sensitivity and a somewhat higher specificity might be expected for the two-strain antigen and cut-point of 1.0 utilized in this study. The intra-assay and interassay variations based on positive and negative control sera were <10%.

In order to participate in the 42-month intervention trial, all study subjects had to sign an informed consent indicating their willingness to participate.

Thirty-nine subjects elected not to participate. An additional 149 subjects were excluded because they were deceased (14), were "out-of-scope" for age or village (42), had a history of allergy to antibiotics (67), or they were not tested for antibodies to *H. pylori* (26). The remaining 3411 male and female subjects from 13 villages were enrolled in the intervention trial in September, 1995. They were distributed as follows:

Village	Male	Female	Total
Xi Quan	131	116	247
Li Hu Zhuang	129	131	260
Wang Jia Zhuang	125	117	242
Xi Si Hou	63	49	112
Li Jia Gou	179	154	333
Suo Zhuang	198	200	398
Huang Ai Quad	78	90	168
Yang Jia He	127	140	267
Xin Zhuang	148	136	284
Guo Jia Zhuang	196	165	361
Hou Jia He	212	194	406
Hou He Ye	92	90	182
Nan Yang He	75	76	151
Total	1,753 (51.4%)	1,658 (48.6%)	3,411

DESCRIPTION OF AREA

Shandong Province is one of the least economically developed provinces in China and Linqu County is one of the poorest counties in Shandong, with 3/4 of the land area located in mountainous regions resulting in infertile soil and periodic severe droughts. Maps showing the location of Shandong Province and Linqu County are found in Appendix 1. Linqu county has a population of about 900,000 and over 600 people die from stomach cancer each year (42% of all deaths from cancer). The annual stomach cancer mortality rates for Linqu County are 70 per 100,000 for men and 26 per 100,000 for women.

The 13 rural villages range in size from 642 to 1845 inhabitants with an average population per village of about 800. Farming is the major occupation and source of income for residents. The average annual per capita income is about \$200. Cash crops include wheat, sweet potatoes, peanuts, and apples. There is variation in the socio-economic status among the villages due to differences in the ability of the land to support farming. There is also variation in the source of water, with one village using a public well, other villages using private wells, and still others using running water piped in from a village water tower.

Most of the villages are well planned and consist of brick houses built in the last 10-15 years. Houses range from three to five rooms and have court yards which contain the families's animals (e.g., ducks, chickens, pigs). Floors are either dirt, cement, or brick. All homes have electricity, but the lighting is limited with many homes having single light bulbs hanging from the ceiling.

Although most homes have a TV set, many have ceiling fans, and a few have telephones, they lack other modern conveniences such as refrigerators, washing machines, dish washers, and air conditioners. Kitchens and pit privies/outhouses are located outside the main house. Food is stored in one room of the house. A typical meal consists of pancakes or steamed bread, some vegetables, and sometimes salty vegetables with a little meat. Villagers buy food at the village store or at the free market held every five days in the town or larger villages. All the elementary age children attend school (the larger villages have their own elementary school), but 42% of the adult residents have had no schooling. The larger villages have their own village doctor; the smaller villages share doctors. The village leader and a village committee oversee the administration of each village.

STUDY DESIGN

This study is a double-blinded cross-sectional investigation of 3411 *H. pylori* infected and uninfected adults enrolled in a joint NCI/BICR randomized intervention trial to inhibit progression of precancerous gastric lesions in Linq County, Shandong Province, China. Responses to a structured questionnaire (Appendix 2) were used to evaluate factors that influence the transmission of *H. pylori* infection in this rural area of China. Sample size calculations indicated that the study should have adequate power (at least 90%) to detect a prevalence rate ratio of 1.2 or greater.

INTERVIEW AND QUESTIONNAIRE

In-person interviews lasting approximately 15 minutes were conducted in Chinese in the homes of *H. pylori* infected and uninfected subjects by trained BICR field staff from October 1997 - May 1998. Both the interviewers and the interviewees were blinded to the *H. pylori* sero-status of the study participants. The questionnaire (Appendix 2) was designed to study risk factors for infectious disease in general and *H. pylori* in particular and to investigate potential modes of transmission. Information was collected on childhood socioeconomic factors, personal hygiene, and exposures related to food, drinking water, waste water, and animals. Topics ascertained in the questionnaire included background and demographic information (marital status, education, occupation, income), household composition (number of people, number of young children, bed sharing, animals), water and personal habits (source of drinking water; water storage; water treatment; washing of chopsticks, bowls and cups; sharing of cups; hand washing practices; bathing practices; teeth brushing), eating habits (consumption of sweet and sour pancakes and raw fruit and vegetables, whether raw fruit and vegetables are washed or peeled before eating, and presence of gastro-esophageal reflux), smoking and drinking habits (ever, age start, current status, frequency, and duration), and childhood household composition (type of floor, number of people, number of children, animals, bed sharing, premastication of food).

The questionnaire was designed by the principal investigator, CAPT Linda Morris Brown, with the assistance of scientists from BICR, NCI, and the

Uniformed Services University of the Health Sciences (USUHS). Drs. Jun-Ling Ma and Wei-dong Liu supervised the training of six Chinese interviewers and the administration of the Chinese-version questionnaire to study subjects. The 10-15 minute interview used a pre-coded structured questionnaire. The coding manual was developed by NCI and BICR. All data were keyed by a BICR coder and 100% verified by a second coder. A 10% sample of the questionnaires were mailed to NCI for verification by the principal investigator. Dr. Wei-cheng You reviewed a sample of tape-recorded interviews to check agreement with completed questionnaires.

After the data were keyed and verified they were computer edited. Range checks were developed to ensure that specific data items had acceptable codes and logic checks were developed to ensure that there was consistency between related data items. A computer disk containing the edited data was mailed to the principal investigator. Frequency distributions for all data items were produced and reviewed for inconsistencies. An analytic file was prepared by Information Management Systems, a NCI computer contractor, under the direction of the principal investigator.

Subject's current occupation was recorded verbatim. These responses were grouped into the following categories: administrative, technical/clerical/sales, precision production, operator/fabricator/laborer, service occupations, farming, and animal worker based on the Standard Occupational Classification Manual 1980 (U.S. Department of Commerce). Data on median village education level was supplied by the Chinese collaborators. These data

were used to group villages into three education level categories: high (27% had an elementary school education), medium (22% had an elementary school education), and low (15% had an elementary school education). Questionnaire data were merged with a file containing information on *H. pylori* and pathology test results and a file containing information on garlic and other allium vegetable consumption.

RECLASSIFICATION OF *H. PYLORI* STATUS

For the Intervention Trial, a subject was considered seropositive if at least one of two optical density readings for IgG or one of two optical density readings for IgA was 1.0 or greater. All other subjects were considered seronegative. For the purposes of the cross-sectional study, more conservative definitions of seropositive and seronegative were employed that included the establishment of a gray or indeterminate zone of values ($\pm 10\%$ of the cutoff value of 1.0). A subject was considered seropositive if both IgG optical density readings were 1.1 or higher and seronegative if both IgG values were 0.9 or lower and neither IgA value was 1.0 or greater. Subjects with missing IgG values were included in the gray zone.

DATA ANALYSIS

Since this is a cross-sectional study, the outcome of interest is the prevalence of *H. pylori* infection as indicated by serology. The measure of association used in the standard analysis is the prevalence odds ratio. Because

the data were multinomial, (i.e., data in a discrete distribution associated with events which might have more than two outcomes - negative, positive, and indeterminate) maximum likelihood estimates were computed using polychotomous logistic regression (BMDP PR, 1990). Age, as a continuous variable, was included in all logistic models to control for potential confounding. Because subjects may share unmeasured risk factors for *H. pylori* infection with members of their families, the infection status may be correlated within families, conditional on measured covariates. Because standard calculations of variances based on the polychotomous logistic model assume that responses are independent, conditional on covariates, they can be misleading in this application. Therefore, the bootstrap based on resampling families with replacement was used to estimate variances and covariances of estimated log relative odds parameters, $\hat{\beta}_1$ and $\hat{\beta}_2$ which are defined precisely in equation [1] below and of functions of $\hat{\beta}_1$ and $\hat{\beta}_2$. To estimate the needed variances and covariances, 100 bootstrap replicates were used.

The prevalence odds ratio may not provide a good estimate of the relative risk, because *H. pylori* positivity is common in this population (more than 60%) and thus does not meet the rare disease assumption. However, prevalence rate ratios can be calculated directly from these data because they come from a cross-sectional sample of a known population. Using predictive models, the relative odds ratio (ROR), the prevalence rate (PR), and the prevalence rate ratio

(PRR) were calculated for subjects aged 40, 50, and 60 using polychotomous logistic regression as described next.

Let X be a $P+1$ dimensional vector of covariates and let β_1 and β_2 be corresponding $(P+1) \times 1$ vectors of polychotomous logistic regression coefficients corresponding to the outcomes $D=1$ (*H. pylori* positive) and $D=2$ (*H. pylori* indeterminate). The vector X has 1 as its first component, corresponding to an intercept, and $D=0$ corresponds to *H. pylori* negative. The basic polychotomous logistic model is then, for $i=1$ or 2 ,

$$P(D=i|X) = \exp(\beta_i^T X) / \{1 + \exp(\beta_1^T X) + \exp(\beta_2^T X)\}, \quad [1]$$

where T indicates transposition. Suppose X_0 is a reference covariate pattern, then the relative odds ratio for $i=1$ is:

$$ROR = \{[P(D=1|X)/P(D=0|X)]/[P(D=1|X_0)/P(D=0|X_0)]\} = \exp\{\beta_1^T (X - X_0)\}. \quad [2]$$

Note that ROR does not depend on the intercept β_{10} , but only on the p other components of β_1 , namely $(\beta_{11}, \dots, \beta_{1p})^T$. Prevalence rates of *H. pylori* infection,

$$PR = P(D=1|X)$$

are calculated directly from [1]. Prevalence rate ratios (PRRs) comparing covariate level X with the reference level are computed from [1] using

$$PRR = P(D=1|X) / P(D=1|X_0).$$

The variables used in the two main models were: Model 1 - frequency of hand washing before meals (>50% of the time, <50% of the time), number of children in the household (0-1, 2+), and median village education level (high, medium, low); Model 2 - frequency of hand washing before meals (>50% of the time, <50% of the time), number of children in the household (0-1, 2+), and

source of drinking water (deep private well, shallow private well/deep village well, shallow village well). Median village education level and source of drinking water were included in separate models because they both represented differences in *H. pylori* serostatus by village and thus were correlated. 95 % confidence intervals (CIs) were calculated from the 2.5th and 97.5th quantiles of the bootstrap distribution, based on resampling and using 1000 bootstrap replicates.

HUMAN SUBJECTS AND CONFIDENTIALITY

The principal investigator received clinical exemption from the Office of Management and Budget (OMB) Clearance for the questionnaire and Institutional Review Board (IRB) approval from the BICR IRB, the Westat, Inc. IRB, the NCI IRB, and the USUHS IRB. All information obtained from the study was and will be treated as confidential. A study ID number replaced the subject's name on all documents received by NCI. The informed consent for the intervention trial included permission for the administration of a questionnaire at 12 and 24 months to collect basic information on health and lifestyle factors. The questionnaire for this study was submitted to the IRBs as an amendment to the original protocol for the Intervention Trial.

INVESTIGATORS

CAPT Linda Morris Brown was responsible for the overall design, management, and data analysis for the study. Assistance in the design of the questionnaire was supplied by Drs. Jun-Ling Ma, Wei-dong Liu, and Wei-cheng

You. Primary support and management of field activities was provided by the BICR under the direction of Drs. Jun-Ling Ma and Wei-dong Liu. They were responsible for translating the questionnaire into Chinese and monitoring field support activities including the training of interviewers, interviewing subjects, and monitoring quality control measures. The questionnaire was back-translated by a Chinese translator under contract to Westat, Inc. and by Ms. Katherine Chen, a NCI employee and was found to agree with the original English version.

CHAPTER IV. RESULTS

Interviews were completed with 3,288 (96.4%) of the 3,411 study subjects enrolled in the Shandong Intervention Trial. Reasons for non-response included death (54 subjects; 1.6%), dropped out of trial (33 subjects; 1.0%), lost to follow-up (10 subjects; 0.3%), and refusal (26 subjects; 0.8%). The *H. pylori* serostatus of the participants was positive for 1994 subjects (60.6%), negative for 1019 subjects (31.0%), and indeterminate for 275 (8.4%).

DEMOGRAPHIC CHARACTERISTICS

Table IV-1 presents the number of subjects by *H. pylori* serostatus according to agegroup, gender, gastric pathology category, and village. The percent positive decreased with age ranging from 63% in those aged less than 40 years to 57% in those aged 55 and older. The percent positive was higher in females (63%) than males (59%). The percent positive varied greatly by gastric pathology category. Subjects with normal gastric mucosa or superficial gastritis have the lowest prevalence of *H. pylori* infection (28%); intermediate levels (46%) were seen for subjects with mild chronic atrophic gastritis (CAG); higher levels (67%-77%) were seen for subjects with other CAG, intestinal metaplasia, and dysplasia. The highest prevalence of infection was seen for subjects with cancer (80%), although this figure was based on only five subjects. There was also significant variation by village ($\chi^2=120.76$, d.f.=12, $p<0.001$). The highest

rate (76%) was seen among subjects from Wang Jia Zhuang, whereas the lowest rate (47%) was seen among subjects from Guo Jia Zhuang.

All analyses were run using polychotomous logistic regression. Since results from the indeterminate group have no clear meaning, the only results reported are for *H. pylori* positive compared to *H. pylori* negative subjects. All results were adjusted for age as a continuous variable. Other variables such as village and village education status were not found to be significant confounders or effect modifiers, and thus were not included in the logistic regression models. Presented for each variable is the prevalence odds ratio (OR), the 95% confidence interval (CI) computed directly from the polychotomous logistic regression, and the bootstrap 95% CI that allows for additive familial correlation.

RISK FACTORS

Cigarette Smoking

Table IV-2 presents prevalence ORs for *H. pylori* infection and smoking-related factors. The odds of ever smoking cigarettes was slightly lower for *H. pylori* positive subjects (OR=0.9, 95%CI=0.7-1.0), however, there were no consistent trends according to the age first started smoking, the number of cigarettes smoked per day, the number of years smoked, or the lifetime number of packs smoked. In addition, the OR for past smoking was slightly lower than the ORs for current smokers (0.8 vs. 0.9), but the difference was not significant.

Alcoholic beverage consumption

Prevalence ORs for the association between *H. pylori* infection and alcoholic beverage consumption are presented in Table IV-3. In these data, there was no association between infection status and alcohol use overall or with any measure of alcohol consumption including age first started drinking, number of times drank per week, or with number of years drank. Among drinkers, the odds of being *H. pylori* positive was slightly reduced for current smokers (OR=0.9) and slightly elevated for past smokers (OR=1.2), although both sets of confidence intervals included 1.0.

Dietary factors

Associations between the prevalence of *H. pylori* infection and dietary factors are presented in Table IV-4. There was a pattern of decreasing ORs with increasing consumption of allium vegetables (ranging from 1.0 for less than 11 jin per year to 0.8 for greater than 21 jin per year; 1 jin ~ 500 grams), although the trend was not significant (p for trend=0.50). The gradients in the ORs for the individual types of allium vegetables (garlic, garlic stalk, scallions, and chives) were less consistent. There was no apparent association between *H. pylori* infection and the number of times per month raw fruits and vegetables were eaten. ORs were reduced the less frequent raw fruits and vegetables were washed, reaching OR=0.7 (95%CI=0.5-0.9) for subjects who never washed their raw fruit and vegetables. No association was seen with the frequency that subjects peeled their raw fruits and vegetables before eating. The number of

sour or fermented pancakes eaten per year was associated with a slightly elevated OR, reaching 1.2 (95%CI=0.8-1.7) in the highest consumption category; whereas the number of sweet or nonfermented pancakes eaten per year was associated with a reduced OR (OR=0.8, 95%CI=0.7-0.9). The OR was only slightly elevated (OR=1.1) for subjects who preferred their food moderately or very salty.

Occupation

Table IV-5 presents ORs for the association between the prevalence of *H. pylori* infection and various occupations. The majority of subjects (75.5% of those *H. pylori* positive and 75% of those *H. pylori* negative) reported having an occupation as a farmer. Although none of the ORs were statistically different from 1.0, elevated ORs were associated with employment in precision production and service occupations; reduced ORs were associated with employment in technical/clerical/sales, operator/fabricator/laborer, and as animal workers.

Water-related exposures

Presented in Table IV-6 are ORs for the prevalence of *H. pylori* infection and its relationship with water-related exposures. Source of drinking water was found to vary between *H. pylori* positive and negative subjects. Compared to subjects who obtained their drinking water from deep private wells, ORs were significantly elevated for subjects who obtained their water from a shallow village well (OR=1.8, 95%CI=1.4-2.3). ORs were also elevated for subjects who used a

shallow private well (OR=1.3), a deep village well (OR=1.4), and running/spring water (OR=1.4). A reduced OR was seen for subjects who obtained their drinking water from a pond/river/ditch (OR=0.8), but the majority of those subjects had obtained their drinking water from a river. Three *H. pylori* positive subjects and one *H. pylori* negative subject reported obtaining their drinking water from a pond or ditch. Similarly, a greater percentage of *H. pylori* positive subjects reported washing diapers in a pond or ditch (OR=2.2, 95%CI=0.6-7.8). There was a slight decrease in the ORs associated with increased number of days that drinking water was stored in a jar, ranging from 1.0 for 0, 0.9 for 1 or 2, and 0.8 for 3 or more; however, all CIs overlapped. There appeared to be no protective effect of boiling drinking water.

An elevated OR was associated with washing or bathing when the weather was warm (OR=1.3, 95%CI=1.0-1.6). The highest ORs were seen for subjects who bathed in a pond or ditch (OR=1.6, 95%CI=1.0-2.4) or who bathed at a public bathhouse (OR=2.8, 95%CI=0.9-8.4). However, there did not appear to be a consistent trend with the number of times bathed when warm. The OR for washing or bathing when the weather was cold was only slightly elevated (OR=1.1). Overall there was no association between *H. pylori* infection and swimming in a river or reservoir, however, the OR was slightly elevated (OR=1.2) for subjects who reported swimming in a river or reservoir more than four times per year.

Exposure to animals and pets

Presented in Table IV-7 are ORs for the association between the prevalence of *H. pylori* infection and exposure to pets and other domestic animals. ORs were reduced for keeping any animal in the house as an adult (OR=0.7, 95%CI=0.5-1.0), and significantly reduced for keeping only one animal in the house as an adult (OR=0.5, 95%CI=0.3-0.9). ORs were reduced for keeping cats (OR=0.6, 95%CI=0.3-1.1) or rabbits (OR=0.7, 95%CI=0.4-1.1), but not dogs (OR=1.2, CI=0.3-4.5). There was also a modest reduction in the OR associated with keeping any animal in their courtyard as an adult (OR=0.8, 95%CI=0.5-1.2). ORs were reduced for all 11 types of animals studied that were kept in the courtyard, including cats (OR=0.7), dogs (OR=0.9), and sheep (OR=0.3).

Overall, there was no association between *H. pylori* infection and keeping animals in the house when the subject was ten years old. However, the ORs associated with keeping cats (OR=0.7, 95%CI=0.5-1.0) or dogs (OR=0.5, 95%CI=0.3-1.0) in the house when the subject was ten years old were reduced. The OR also was not elevated for subjects who had a job working with animals (OR=0.5, 95%CI=0.2-1.2).

Sanitation/hygiene factors

ORs for the association between the prevalence of *H. pylori* infection and certain sanitation or hygiene factors are presented in Table IV-8. The OR associated with spitting onto the ground was slightly reduced (OR=0.9, 95%CI=0.8-1.0), but no gradient was evident for the number of times per month that subjects spit onto the ground. There was also no association with the number of times per month that subjects brushed their teeth. The OR was elevated for subjects who reported washing their hands before eating less than half the time (OR=1.6, 95%CI=1.0-2.5) or never (OR=3.8, 95%CI=0.5-31.0). On the other hand, reduced ORs were seen for subjects who did not always wash their hands after bowel movements and for subjects who never washed their hands before preparing meals. No consistent patterns were seen with frequency of washing hands or body with soap, frequency of cup washing after use, the temperature of the water used to wash chopsticks and bowls, or frequency of pre-chewing their child's food. Most subjects always washed their chopsticks and bowls after eating, but few always washed them with detergent. The ORs were reduced for subjects who shared cups with their family less than half the time, for subjects who never kissed young children on the lips or who were never kissed as children by their parents on the lips, and for subjects whose house where they lived when they were ten years old had a floor of some material other than dirt.

Density/crowding factors

Presented in Table IV-9 are the prevalence ORs for the association between *H. pylori* infection and density/crowding factors. As an adult, there was no consistent association with total number of people in the household and the OR was only slightly elevated for sharing a bed with a spouse (OR=1.1, 95%CI=0.9-1.4). However, the ORs were elevated for more than one child in the household (OR=2.0, 95%CI=1.0-3.7), sharing a bed with more than two people (OR=2.1, 95%CI=1.1-3.8), and sharing a bed with more than one child (OR=1.7, 95%CI=1.0-2.8). On the other hand, the ORs were less than 1.0 for density factors when the subject was 5 or 10 years old such as number of people in the household and number of people shared a bed with.

Social factors

The association between social factors and prevalence of *H. pylori* infection is presented in Table IV-10. The most striking finding was with village education level; the OR increased with decreasing village education level, from 1.0 for high, to 1.7 (95%CI=1.4-2.1) for medium, to 2.4 (95%CI=2.0-3.0) for low (p for trend<0.001). No consistent patterns were seen for socioeconomic variables at the individual level (e.g., subject's education level, spouse's education level, or subject's annual income). The ORs were reduced for subjects who were divorced or never married (OR=0.4) compared with those who were married, but the number of subjects in those categories was small.

Gastric factors

The association between the prevalence of *H. pylori* infection and gastric factors is presented in Table IV-11. Overall, there was no association between ever having gastric reflux and prevalence of *H. pylori* infection. However, the ORs were slightly reduced for subjects who had reflux 4-8.9 (OR=0.9) and nine or more (OR=0.8) times per month.

MULTI-VARIATE ANALYSIS

Presented in Table IV-12 are the prevalence rates (PRs), prevalence rate ratios (PRRs) and relative odds ratios (RORs) for combinations of models according to age (40, 50, or 60), percent of time washed hands before eating (<50% or >50%), number of children in the household (0-1 or 2+), and village education level (low, medium, or high). Within each age category, the referent group is subjects who washed their hands >50% of the time, who had 0-1 children in their household, and who came from a village with high education level. The RORs are the same for each age category because the parameter estimate for age when two groups being compared have the same age, drops out.

There were only small variations in the PR by age, with the highest PR seen for subjects aged 40 and the lowest for subjects aged 60. For example, the PR for a subject aged 40 who washed their hands <50% of the time, had 2+ children in their household, and came from a low education village was 74%; the PR for a 60 year old person with the same factors was 71%. Within each age category, the PR was highest for those subjects from a low education village with

at least one other risk factor (i.e., washed hands <50% of the time or had 2+ children in the household). The lowest PR for each age category (51% for age 40, 48% for age 50, and 46% for age 60) was for subjects with none of the risk factors (i.e., they washed their hands >50% of the time and had fewer than 2 children in their household) who lived in a high education village. Within age categories, PR was strongly related to village education level, with the highest PRs seen for subjects who lived in low education villages, intermediate values for subjects who lived in medium education villages, and the lowest PRs seen for subjects who lived in villages with the highest education level.

The PRRs varied little by age. Most of the PRRs were statistically different from 1.0, ranging from 1.0 to 1.5. Because the background rate of *H. pylori* is high in this population, even a small PPR has a big impact. An inverse relationship between prevalence of *H. pylori* infection and village education level was apparent, with the subjects who lived in the lowest education villages having the largest PRRs and subjects who lived in the highest education villages the smallest. In general, PRRs were lowest for subjects who washed their hands <50% of the time and who had 2 or more children in their household.

Unlike the PRRs, the RORs showed great variation both within and among village education categories. Within a village education category the greatest RORs was seen for subjects with both risk factors (washed hands <50%, had 2+ children); the lowest was seen for subjects with neither risk factor (washed hands >50%, had 0-1 children. For example, among subjects who lived in villages with low education the RORs ranged from 8.6 for subjects who had

both risk factors, 5.0 or 4.1 for subjects who had one risk factor, and 2.4 for subjects who had neither risk factor. The RORs also varied greatly by village education level. The RORs ranged from 8.6 (95%CI=3.9-22.2) for subjects with both risk factors who lived in a village with the lowest education level, to 6.0 (95%CI=2.7-16.2) for subjects who lived in a village with an intermediate education level, to 3.6 (95%CI=1.6-9.3) for subjects who lived in a village with the highest education level.

Presented in Table IV-13 are the PRs, PRRs and RORs for combinations of models according to age (40, 50, or 60), percent of time washed hands before eating (<50% or >50%), number of children in the household (0-1 or 2+), and source of drinking water (shallow village well, shallow private well or deep private well, and deep private well. Within each age category, the referent group for the PRRs and RORs is subjects who washed their hands >50% of the time, who had 0-1 children in their household, and who had a deep private well.

The PRs, PRRs, and RORs followed similar patterns to those seen in Table IV-12, but the values were lower for each combination of variables. Overall, the PRs, PRRs, and RORs were highest for subjects who obtained their drinking water from a shallow village well, intermediate for subjects whose drinking water came from a shallow private well or a deep village well, and lowest for those who obtained their drinking water from a deep private well. For example, the PRs ranged from 73% (95%CI=63-81%) for subjects aged 40 who washed <50% of the time, had less than two children, and obtained their drinking water from a shallow village well to 49% (95%CI=43-55%) for subjects aged 60

who washed >50% of the time, had less than two children, and obtained their drinking water from a deep private well. The PRRs ranged from 1.0-1.4. The RORs also showed variation by source of drinking water. The RORs ranged from 7.0 (95%CI=3.4-17.6) for subjects with both risk factors whose drinking water came from a shallow village well, to 4.9 (95%CI=2.3-12.2) for subjects whose drinking water came from a shallow private well or a deep village well, to 3.9 (95%CI=1.9-9.4) for subjects whose drinking water came from a shallow village well.

CHAPTER V. DISCUSSION

This large cross-sectional study conducted among adults living in 13 villages in Shandong Province, China provided the opportunity to assess risk factors related to the prevalence of *H. pylori* infection. Factors of particular interest included measures of adult and childhood socioeconomic status, source of drinking water, personal hygiene and sanitation factors, sharing of utensils, potential food contamination, and exposure to animals.

RESEARCH QUESTION 1:

Is there an association between H. pylori positivity and drinking water source (i.e., shallow well, deep well, pond, river, ditch, or running water), storage, or treatment?

There is some suggestion in the data, that source of drinking water is potentially related to *H. pylori* infection. Compared with subjects who obtained their drinking water from deep private wells, ORs were elevated for subjects who obtained their water from wells that were more apt to be contaminated with fecal or other material containing *H. pylori* (shallow private wells (OR=1.3), deep village wells (OR=1.4), or shallow village wells (OR=1.8)). The possible role of waterborne transmission of *H. pylori* is supported by a study in the U.S. that reported a strong association between consumption of well water contaminated by *H. pylori* and *H. pylori* positive status in persons consuming the water.²¹² The finding of *H. pylori* DNA in the drinking water in Peru suggests that waterborne

transmission may be important in areas of the world, such as Peru and China with untreated water. Klein et al found that the water supply in Lima, Peru may be vulnerable to bacterial contamination, especially when it is stored in a cistern;¹⁴⁴ however, this study detected a slight protective effect for subjects who stored their water in a jar for more than three days. Although it has been shown that *H. pylori* can live for several days in tap water in its infectious bacillary form,²⁰⁵ longer storage may have resulted in the conversion of *H. pylori* into its coccoid form.²⁰³ Surprisingly, in this study there was no evidence of a protective effect of always boiling water before drinking it. Although the data from this study are not totally consistent, the elevated ORs for some sources of drinking water suggest that the high prevalence of *H. pylori* infection in this study area might be related to the presence of *H. pylori* in the drinking water.

RESEARCH QUESTION 2:

Is there an association between H. pylori positivity and adult sanitation and hygiene (e.g., hand washing, use of soap, bathing, teeth brushing, and sharing of cups)?

Several studies have reported an association between poor hygiene practices and *H. pylori* infection; however, most were either conducted among children^{122,150} or asked about childhood exposures.^{73,93} The questionnaire for this study sought information about the frequency as an adult of hand washing before eating meals, after bowel movements, and before preparing meals. The most consistent finding was for subjects who washed their hands before eating

meals less than half the time (OR=1.6) or never (OR=3.8). There was no protective effect of washing hands, body, or chopsticks and bowls with soap or detergent. Although the majority of subjects reported always sharing cups with their family, there was a protective effect for subjects who shared cups less than half the time. These findings were in contrast to those by Peach et al in Australia⁶² who found an elevated risk for adult subjects who did not usually wash their hands after going to the toilet (OR=4.1), but no association with frequency of sharing cups or washing hands before eating. In this study, ORs were slightly reduced for subjects who never kissed young children on the lips, whereas no association with either frequency of teeth brushing or spitting onto the ground was apparent. Several studies found the prevalence of *H. pylori* infection to be higher in subjects who did not have a bathroom, indoor toilet, or running water;^{73,78,93} modern necessities not found in any of the homes of the study subjects. Since all of the rural Chinese study subjects ate with chopsticks, it was not possible to confirm the elevated OR of 2.5 reported among Chinese immigrants in Australia who used chopsticks. In this study, there was a borderline association for subjects who bathed when the weather was warm (OR=1.3) that reached 1.6 for subjects who washed in a pond or ditch. However, there was no gradient in risk with the number of times per month that the subject bathed. There was a slight (OR=1.2) non-significant risk for subjects who swam in a river or reservoir four or more times per year. Higher prevalence of *H. pylori* infection has been linked to swimming in a stream in Colombia,¹²² but not to swimming near contaminated beaches or bathing in local rivers, irrigation

ditches, or lakes in Chile.¹²⁵ These data suggest that hand washing before meals and elimination of familial cup sharing may be important health measures to reduce the prevalence of *H. pylori* infection in China. Also, source of water used for bathing, especially pond or ditch water, may be related to increased prevalence of *H. pylori* infection.

RESEARCH QUESTION 3:

Is there an association between H. pylori positivity and social class factors?

Social class factors (i.e., SES-based occupation, education, income, SES) have been associated with *H. pylori* infection status in a number of studies throughout the world.^{48,49,54,57,59-63,66,68,72,74,77-80,84,85,88,93,124,125,147,153} This study noted a strong inverse relationship between village education level and prevalence of *H. pylori* infection, that reached 2.4 for those in the lowest compared to the highest village education category. However, there were no consistent associations in this study with individual measures of social class including number of years of education (subject or spouse), and subject's annual family income. These data suggest that density factors and group SES factors may be associated with the prevalence of *H. pylori* infection in rural China. The absence of an association between *H. pylori* and individual social class factors may result from a lack of heterogeneity in the study population.

RESEARCH QUESTION 4:

*Is there an association between *H. pylori* positivity and use of tobacco or alcohol products?*

Similar to some,^{13,54,63-65,68,75,87,88,92,116} but not all^{57,61,109,112-114} recent studies, this study found no significant association with current smoking or any other measure of cigarette use including age first started smoking, number smoked per day, number of years smoked, and lifetime packs smoked. All of the recent epidemiologic studies that looked at the relation between alcohol use and *H. pylori* infection reported either no association^{54,61-63,68,92} or a reduction in risk,^{57,109,115,117,118} that was stronger for wine than beer and more apparent at moderate to high levels of alcoholic beverage consumption. Similarly, this study found no association between *H. pylori* infection and ever/never use of alcohol, and only a slight (OR=0.9) nonsignificant reduction in the OR for subjects who drank alcohol seven or more times per week. The relative homogeneity among drinkers in this study including the lack of heavy drinkers may have limited the ability to detect a protective effect from alcohol even if it has antimicrobial properties.¹²⁰ These findings suggest that neither smoking nor drinking are related to the prevalence of *H. pylori* infection in this population.

RESEARCH QUESTION 5:

*Is there an association between *H. pylori* positivity and presence of gastro-esophageal reflux?*

Some studies have suggested that *H. pylori* infection, which may diminish acid secretion, appears to be lower in persons with gastroesophageal reflux disease (GERD).^{46,47} GERD, the major risk factor for Barrett's esophagus, is associated with adenocarcinoma of the esophagus.^{46,47} In this study, there were slight nonsignificant protective effects for subjects who reported the occurrence of gastric reflux 4-8.9 (OR=0.9) and nine or more (OR=0.8) times per month. These results suggest that *H. pylori* infection in Linqu county may be associated with a decreased prevalence or severity of GERD.

RESEARCH QUESTION 6:

Is there an association between H. pylori positivity and raw fruit and vegetable consumption or preparation (i.e. washing or peeling)?

Increased consumption of allium vegetables has been suggested as a protective factor for stomach cancer^{238,239} and garlic has been shown to inhibit *H. pylori* in vitro,^{240,241} but not in vivo.²⁴² In this study, there was a pattern of decreasing ORs with increased consumption of all allium vegetables combined, but the pattern was less consistent for each individual type of allium vegetable (garlic, garlic stalk, scallions, chives). Several studies have reported significantly reduced ORs and negative gradients in risk of *H. pylori* infection with increased consumption of fruits and/or vegetables, vitamin C, and beta-carotene.^{109,121-123} In contrast, consumption of raw/uncooked vegetables was associated with elevated risk of *H. pylori* infection.^{122,125} In these data, there was no apparent association between *H. pylori* infection and the number of times per month raw fruits and

vegetables were eaten or with the frequency that subjects peeled their raw fruits and vegetables before eating. ORs were reduced the less frequently raw fruits and vegetables were washed.

These data do not support a role for raw fruit and vegetable consumption in the prevalence of *H. pylori* infection in this study population, except for a possible protective effect of heavy intake of garlic and other allium vegetables. The reduced ORs for subjects who washed their fruit or vegetables infrequently suggests the possibility of transmission of *H. pylori* to raw fruits and vegetables via water, but not soil.

RESEARCH QUESTION 7:

Is there an association between H. pylori positivity and adult exposure to pets and other domestic animals?

Although *H. pylori* has been isolated from laboratory domestic cats,^{215,216} no evidence of *H. pylori* infection was found in an investigation of stray cats.²¹⁹ Similarly, the epidemiologic literature suggests that adult ownership of a cat or other pet is not associated with a higher prevalence of *H. pylori* infection.^{62,78,80,97,148} In fact, one study in the U.S. found pet owners to have significantly lower rates of *H. pylori* infection.⁸⁰ In this study, ORs were reduced for adults who kept cats (OR=0.6) or any animal (OR=0.7) in the house or any animal in the courtyard (OR=0.8). ORs were reduced for all types of animals that were kept in the study subject's courtyard, including cats (OR=0.7) and sheep (OR=0.3, although based on very small numbers). Exposure to sheep was

implicated in a recent study that found a high prevalence of *H. pylori* infection among Sardinian shepherds occupationally exposed to sheep.¹⁴⁹ In agreement with studies in Australia and Germany that reported no association with occupational handling of animals,^{62,85} the OR in this study was not elevated for subjects who had a job working with animals. Thus, these data for rural China support epidemiologic studies in other parts of the world that found ownership of a cat or other domestic animal not to be associated with increased prevalence of *H. pylori* infection.

RESEARCH QUESTION 8:

*Is there an association between *H. pylori* positivity and childhood exposures (e.g. domestic animals, crowding, and eating premasticated food)?*

Although most studies found no association between *H. pylori* infection and having a pet during childhood,^{93,127,147} several studies did report a positive association with cat ownership as a child^{85,146,220} A study from rural Columbia found higher *H. pylori* prevalence among children who played with sheep.¹⁴⁹ In this study, there was no association between *H. pylori* infection and presence of cats (OR=0.7) or any animal (OR=1.0) in the household when the subject was ten years old. No information was sought on childhood exposure to sheep or other farm animals.

In these data, there was no indication that crowding or density factors as a child (e.g., number of people in the household when the subject was ten year old, the number of young siblings (< five years old, < ten years old) when the

subject was ten, the number of people the subject shared a bed with at ages five and ten) was related to a higher prevalence of *H. pylori* infection as an adult. This is in contrast to many recent studies that found that some measure of high density during childhood such as sibship size,^{67,87,93,243} number of persons or children in the home,^{49,62,122} number of persons per room,^{73,78,100,150} or having to share a room or bed with a parent^{85,154} was associated with prevalence of *H. pylori* infection.

According to the literature, poor hygiene or sanitation practices, especially during childhood such as having no bathroom, refrigerator, or hot water supply in the house,^{73,93,100,150} sharing cups and having a mother who didn't use soap when she washed her hands or a mother who prechewed their food when they were a child^{122,151} was associated with a higher prevalence of *H. pylori* infection. This study was unable to evaluate some of these factors because no subject had a bathroom, refrigerator, or hot water supply in the house as a child. The ORs were slightly reduced for subjects who were never kissed as children by their parents on the lips and for subjects whose house where they lived when they were ten years old had a floor of some material (brick, wood, cement) other than dirt. The OR was also reduced for subjects who reported that their parents never pre-chewed their food as a child, but the association was not consistent across exposure categories. These findings suggest that childhood exposures, as recalled during adulthood, are not important predictors of adult prevalence of *H. pylori* infection in this study population.

STRENGTHS OF THE STUDY

This cross-sectional investigation allowed the opportunity to evaluate some possible etiologic factors that influence the prevalence of *H. pylori* infection in Shandong Province, China. The benefit of a cross-sectional study over a case-control study is that it is based on a sample of the general population, thus both ORs and PRRs can be calculated. An additional benefit of a cross-sectional study over a cohort study is that it can be carried out over a relatively short time period.

Bias, or a systematic error in the risk estimates due to a flaw in the design or conduct of a study, is always of concern in epidemiologic studies. Non-response bias can be a problem if the response rates are low since people who don't respond are generally not a representative group of study subjects and thus may have a different risk profile than responders. This type of bias was not of great concern in this study because of the high (over 96%) participation rates. Selection bias may occur if the way in which infected and noninfected subjects were selected induced an apparent, but not true, association with an exposure. *H. pylori* status was determined using a serologic test. For the Intervention Trial, a subject was considered seropositive if at least one of two optical density readings for IgG or one of two optical density readings for IgA was 1.0 or greater. All other subjects were considered seronegative. To improve sensitivity and specificity for the cross-sectional study, a more conservative definition based primarily on IgG values that included the establishment of a gray or indeterminate zone ($\pm 10\%$ of the cutoff value of 1.0) was utilized. The

classification of *H. pylori* serostatus was determined scientifically based on the Elisa optical density test results. Because study investigators were blinded to the *H. pylori* serostatus of study subjects, selection bias is unlikely to have played a role in the results.

The quality of the information collected was comparable between *H. pylori* positive and *H. pylori* negative subjects (information bias). The same structured questionnaire was administered to all study subjects in person by trained interviewers. *H. pylori* is generally asymptomatic so subjects and interviewers were unaware of infection status. In addition, a survey research company, Westat, Inc. was responsible for assigning ID numbers and determining the randomization for the intervention trial; none of the NCI or BICR investigators were aware of the treatment status of participants. Thus, neither recall bias (differential memory on the part of diseased subjects in an attempt to explain their disease) nor interviewer bias (occurs when interviewers are aware of the disease status of subjects and are not objective in their interviewing techniques) could have played a role in this study. Bootstrap confidence intervals were calculated to avoid underestimation of the standard errors due to intrafamilial correlations. This did not appear to be a problem, however, as both the bootstrap and regular confidence intervals were similar. Although there may be some degree of misclassification or inaccuracies in the data acquisition, because of the reasons stated above, inaccuracies are not likely to differ systematically (differential misclassification) according to *H. pylori* serostatus.

LIMITATIONS OF THE STUDY

There are also several limitations of this study. Similar to all cross-sectional studies, the outcome and the exposure are assessed simultaneously so it is often difficult to separate cause and effect. Also, the outcome is measured as prevalence rather than incidence of disease and will reflect *H. pylori* infections that could have occurred from early childhood to several weeks before the blood sample was drawn.

Because this study assessed the effects of many factors related to prevalence of *H. pylori* infection in other populations, it is surprising that more significant associations were not found. Non-differential misclassification (an error in the data that is not related to disease or exposure status), may have hampered the ability to detect actual associations. A lack of heterogeneity in this rural Chinese population may have limited the ability to detect differences between *H. pylori* positive and negative subjects since it is not possible to detect any causes of disease that are ubiquitous or relatively invariant within the population under study.²⁴⁴ Not only is this a potential problem for adult exposures, but it is even more likely that subjects would have shared common environmental factors before the cultural revolution; factors that may have influenced their acquisition of *H. pylori* infection during childhood. It is also possible that almost everyone was exposed to the *H. pylori* organism during childhood and it was differences in host factors and host susceptibility that determined who became clinically infected.

Another reason for the lack of significant findings may be a result of non-differential misclassification. In this study, many questions did not seek quantitative responses so it may have been difficult for study subjects to fit their answers into the qualitative categories of “Always, Sometimes, Never”.

Answering quantitative questions also may have been difficult for the study subjects with little, if any, formal education. In addition, recall of information from childhood may have been difficult and fraught with error. For binary exposures and for continuous variables when the error was independent of the true value, any bias introduced by non-differential misclassification would have shifted the risk estimate toward the null.²⁴⁵ Thus, the observed ORs are probably closer to 1.0 than they would have been if there were no errors in the data. Because errors are unlikely to be independent of true values in self-reported questionnaire data, when the exposure variable was polytomous (i.e., more than two categories) the bias from non-differential misclassification could be strong enough to either exaggerate or reverse the estimate of the effect of a trend.²⁴⁵

Because of the problem of multiple comparisons, the possibility that some of the positive findings in this study were due to chance can't be ruled out. Based on an alpha level of 0.05 five significant findings for every 100 ORs or PRRs calculated would be expected. Thus, it is important to consider the strength and consistency of the association and biologic plausibility when deciding which exposure factors may be causally related to prevalence of infection.

In the multivariate analyses there were notable discrepancies in the magnitude of the associations between the RORs and the PRRs. In these data the ROR may not have been a good estimator of the PRR because the outcome of interest was not rare in the study population. Also, because the model was based on polychotomous logistic regression, the estimates for the RORs and PRRs were based on different denominator data. The RORs compared the prevalence of exposure in persons who were *H. pylori* positive to the prevalence of exposure in persons who were *H. pylori* negative. As a result, persons with indeterminate serostatus were excluded from this analysis. The PRR compared the prevalence of being *H. pylori* positive (relative to the whole population which included persons with indeterminate serostatus) in persons with the exposure to the prevalence of being *H. pylori* positive in persons without the exposure.

CONCLUSIONS

In summary, this study found elevated ORs for group, but not for individual social class factors. Determinants of density or crowding during adulthood, but not during childhood appeared to be related to the prevalence of *H. pylori* infection. Source of drinking water or bathing water may be associated with *H. pylori* serostatus. Use of alcohol, tobacco, raw fruits and vegetables (with the possible exception of allium vegetables) and exposure to animals during adulthood or childhood were not associated with increased prevalence of *H. pylori* infection in this rural Chinese population.

Based on the findings for the density and hygiene factors, as well as data from another study in the area that found a higher prevalence of *H. pylori* infection in the children of infected parents,¹⁰³ person-to-person transmission appears to be a likely route of infection in these rural Chinese villages. However, the data are not sufficient to determine whether the route was oral-oral, fecal-oral, or gastro-oral. A plausible role of waterborne transmission in this population was suggested by the possible association between *H. pylori* infection and source of drinking water and bathing water. However, a role for zoonotic transmission in the spread of *H. pylori* infection in the study area appears unlikely.

FUTURE DIRECTIONS

Global issues

Infection with *H. pylori* is generally asymptomatic, thus it may be well-adapted to exist in the human stomach for the lifetime of its host. Because only a small percentage of those infected develop severe disease (duodenal ulcer, gastric ulcer, gastric lymphoma, or gastric cancer)^{5,7,57} and *H. pylori* may be protective against other conditions such as adenocarcinoma of the esophagus,^{46,47} the public health response is not clear.

Possible questions that may require addressing include the following:

- (1) Because *H. pylori* is an organism thought to be indigenous to the human population, should it be eliminated? If yes, what is the role of public health

in discovering the route(s) of transmission, educating the public, and reducing or eliminating spread?

- (2) Should populations be screened for *H. pylori* infection? If yes, should all populations be screened or only those at high risk of gastric lymphoma or gastric cancer?
- (3) Should *H. pylori* be eliminated in all persons in whom it is detected or only in those with severe symptoms or at high risk of gastric lymphoma or gastric cancer?
- (4) Does elimination of *H. pylori* in middle-aged adults protect against the development of gastric cancer?
- (5) Should a vaccine against *H. pylori* be used, and if so, who should be vaccinated? Should its use be universal or restricted to children living in populations at high risk of stomach cancer like parts of rural China?

Studies within the intervention trial cohort

To further investigate waterborne factors as a possible transmission mechanism for *H. pylori* infection in Shandong Province, I plan to conduct a small nested case-control study in collaboration with investigators from Pennsylvania State University. To evaluate whether an association exists between *H. pylori*-infected drinking water and *H. pylori* serostatus, the drinking water sources of approximately 50 *H. pylori* positive and 50 *H. pylori* negative households would be tested for the presence of *H. pylori* and compared with the serostatus of household members.

As part of the Intervention Trial, blood was drawn and *H. pylori* serostatus determined on all trial participants annually from 1996-1999. To further investigate risk factors related to the prevalence of *H. pylori* infection, questionnaire responses among subjects randomized to the *H. pylori* treatment arm whose disease has remained eradicated, who seroconverted after their infection was considered to be eradicated, and whose infection was never successfully eradicated could be compared.

Public health significance of *H. pylori* research in Linqu County

This study confirmed the high prevalence of *H. pylori* infection among residents of Linqu County. In addition, it identified factors that may reduce *H. pylori* prevalence in rural China (i.e., hand washing, and higher village SES-associated factors) and others that may increase *H. pylori* prevalence (i.e., “contaminated” water used for drinking, bathing, and washing of fruits and vegetables eaten raw). Knowledge of *H. pylori* etiology and transmission are fundamental to devising approaches for controlling this infectious disease. Improvements such as increased personal and family hygiene; overall economic development, including refrigeration and better sanitation; and treatment of the water supply may have a large impact on public health in rural China.

Its parent study, a randomized intervention trial involving treatment of *H. pylori* in persons at various pathologic stages in the progression toward gastric cancer, should have a global impact by helping determine the efficacy of *H. pylori* eradication in adults at different points in this continuum. If treatment

proves successful, then routine screening for *H. pylori* and subsequent treatment may be a viable option in affluent countries with access to preventive medical services. However screening and treatment may not be efficacious in developing countries with large numbers of affected individuals. The high cost and technical difficulties involved in administering effective treatment programs may be prohibitive. In these countries, vaccination (once developed and tested) may be a better alternative. Because chronic gastritis and other gastric cancer precursors are common; *H. pylori* rates are high in children (50% at ages 3-4) as well as adults; use of antibiotics is low; and the population is well characterized, cooperative, and homogeneous, Linqu County may be an ideal location to conduct a placebo-controlled *H. pylori* vaccine trial among infants or very young children.

TABLES

Table II-1. Prevalence of *Helicobacter pylori* in various adult populations

Author, Year, Ref No.	Country	No. in study	Population	Age range	Diagnostic Test	Rate
Alagathanam, 1999 ³⁵	India	354	General upper class population	12-70+	Serology	49.4%
Kimura, 1999 ⁸⁹	Japan	196	Institutionalized	3-64	Serology	81.1%
Wong, 1999 ⁷¹	China Hong Kong Changle	397 1456	Healthy volunteers	36-65	Serology	80.4% 58.4%
Buckley, 1998 ⁷²	Ireland	1000	Blood donors	18-60	Serology	43.0%
Kawasaki, 1998 ⁸⁶	Nepal	1142	General population	4-93	Serology	56.8%
Kikuchi, 1998 ⁴⁷	Japan	4361	Public service workers	19-69	Serology	30.5%
Lin, 1998 ³⁷	Australia	273	General population	20-80	Serology	38.0%
Luzza, 1998 ⁸³	Rural Italy	705	Out-patients	1-87	Serology	63.0%
Rothenbacher, 1998 ⁸⁵	Germany	337	Adult students	50-85	UBT	34.8%
Senra-Varela, 1998 ⁸⁸	Spain	332	General population	Adult	Serology	43.0%
Souto, 1998 ⁸⁹	Brazil	164	General population	20-90	Serology	84.7%
Strachan, 1998 ⁴¹	South Wales	1796	General population	45-59	Serology	70.0%
Stroffolini, 1998 ¹⁶	Italy	1659	Military students	17-24	Serology	17.5%
Torres, 1998 ⁶⁰	Mexico	5997	General population	20-90	Serology	81.3%
Us, 1998 ⁸⁴	Turkey	364	Out-patients	20-80	Serology	69.0%
Zober, 1998 ¹¹³	Germany	6143	Chemical workers	17-64	Serology	38.2%
Ahmad, 1997 ⁸⁵	Bangladesh	181	Out-patients	20-44	Serology	92.0%
Bohmer, 1997 ⁸⁰	Netherlands	338	Institutionalized	11-89	Serology	82.8%
		254	Healthy employees	18-50		27.2%
Furuta, 1997 ⁸⁶	Japan	1043	Out-patients	16-65	Serology	40.0%
Murray, 1997 ⁸¹	Ireland	4742	General population	12-64	Serology	50.5%
Peach, 1997 ⁸²	Australia	217	General population	Adult	Serology	30.6%
Rothenbacher, 1997 ⁸⁷	Germany	501	Out-patients	15-79	UBT	23.4%
Shinichi, 1997 ⁸⁴	Japan	566	Workers	50-55	Serology	79.3%
Taylor, 1997 ⁴²	US	204	Army personnel	19-37	Serology	37.0%
Andersen, 1996 ³⁵	Denmark	3589	General population	30-60	Serology	25.9%

Author, Year, Ref No.	Country	No. in study	Population	Age range	Diagnostic Test	Rate
Breuer, 1996 ³³	Germany	260	Blood donors	18-61	Serology	39.2%
Fraser, 1996 ³⁴	New Zealand	579	Workers	40-64	Serology	56.0%
		190	European			36.0%
		195	Maori			57.0%
		194	Pacific Islanders			73.0%
Malaty, 1996 ³⁴	Korea	161	Healthy volunteers	20-75	Serology	75.0%
Malaty, 1996 ³⁴	Russia	213	Out-patients	20-75	Serology	88.0%
Martin-de-Argila, 1996 ³⁵	Spain	381	Out-patients	5-77	Serology	53.0%
Matysiak-Budnik, 1996 ³⁶	Poland	656	Blood donors	0-85	Serology	73.0%
Zhang, 1996 ³⁷	Rural China	2646	General population	35-64	Serology	72.0%
Blecker, 1995 ³⁸	Belgium	618	Pregnant women out-patients	18-40	Serology	22.6%
	Caucasian	562				62.5%
	Mediterranean	56				85.6%
Chang-Claude, 1995 ³⁸	China	194	Asymptomatic subjects	15-26	Biopsy	59.5%
Chow, 1995 ³⁹	Australia	328	General population	25-64	Serology	
			Chinese immigrants			
Gasbarrini, 1995 ⁴⁴	San Marino	2237	General population	18-70+	Serology	51.0%
Gilboa, 1995 ⁴⁵	Rural Israel	377	General population	30-90	Serology	72.0%
Harris, 1995 ⁴¹	UK	424	Institutionalized	18-106	Serology	87.0%
		267	Healthy controls	18-80+		40.9%
Hyams, 1995[61]	US	1000	Navy and Marine Corps personnel	17-50	Serology	25.0%
	White	730				17.5%
	Black	168				46.4%
	Hispanic	76				44.7%
	Other	24				29.2%
Lambert, 1995 ⁴²	Australia	122	Institutionalized	19-47	Serology	75.0%
		273	Population controls			23.0%
Murray, 1995 ⁴⁶	Ireland	2115	General Practitioner lists	25-64	Serology	57.6%
Replogle, 1995 ⁴⁸	California	556	Out-patients	20-39	Serology	27.2%

Author, Year, Ref No.	Country	No. in study	Population	Age range	Diagnostic Test	Rate
Malaty, 1994 ⁷⁷	Texas Hispanics Blacks	90 60	Healthy volunteers	19-49	Serology	47.0% 65.0%
Smoak, 1994 ⁸⁴	US White Black Hispanic	938 536 324 47	Army recruits	17-26	Serology	26.3% 14.0% 44.0% 38.0%
Teh, 1994 ⁶⁷	Taiwan	823	General population	1-40+	Serology	54.4%
Webb, 1994 ⁷⁸	UK	471	Male volunteers	18-64	Serology	37.0%
EUROGAST, 1993 ⁸⁸	Japan Poland Denmark Minnesota	386 171 157 198	General population	25-34 55-64 25-34 55-64 25-64 55-64 25-64 55-64	Serology	61.0% 89.0% 69.0% 89.0% 15.0% 30.0% 15.0% 34.0%
Malaty, 1992 ⁷⁹	Texas Hispanics Blacks Whites	89 89 89	Healthy volunteers	20-75	Serology	65.0% 66.0% 26.0%
Mendall, 1992 ¹⁰⁰	UK	215	In- and out-patients	18-82	Serology	33.0%
Mitchell, 1992 ⁸⁹	China	1727	General population	1-50	Serology	44.2%
Graham, 1991 ⁸⁰	Texas Blacks Whites	246 239	Healthy volunteers	15-80	Serology	70.0% 34.0%
Polish, 1991 ¹⁰¹	Colorado	370	Out-patients	15-70	Serology	23.0%
Sitas, 1991 ⁷⁰	Wales	749	General population	30-75	Serology	56.9%
Loffeld, 1990 ⁸¹	Netherlands	401	Blood donors	19-65	Serology	35.0%

Table II-2. Studies assessing the relationship between *Helicobacter pylori* infection and smoking

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Kikuchi, 1998 ⁸⁷	Japan	4361	Public service workers aged 19-65	Ex-smoker, OR=1.0 (0.8-1.2) Current smoker, OR=0.8 (0.7-0.9)	Strength: Control for confounding by logistic regression
Lin, 1998 ⁸⁷	Australia	273	General population	Ex-smoker, OR=1.8 (0.9-3.3) Current smoker, OR=1.9 (1.2-2.8)	Strength: Population-based sample; control for confounding by logistic regression Weakness: No association with intensity or age started smoking
Strachan, 1998 ⁴⁸	South Wales	1796	Prospective Heart Disease Study volunteers aged 45-59	HP prevalence: never smoke (68.8%), ex-smoker (70.4%), cigar/pipe (68.9%), 1-14 cigs/day (74.4%), 15-24 cigs/day (71.7%), ≥25 cigs/day (69.2%) [p for heterogeneity=0.70]	Weakness: No control for confounding; statistical test used not specified
Zober, 1998 ¹¹²	Germany	6143	Chemical production workers aged 17-64	HP prevalence: never smoke (33.7%), ex-smoker (42.4%), 1-15 cigs/day (37.4%), >15 cigs/day (42.6%)	Weakness: Sample not representative of general population; no control for confounding; statistical tests not utilized
Brenner, 1997 ¹¹⁵	Germany	501	Out-patients aged 15-79	Former smoker, OR=1.5 (0.8-2.7) Current smoker, OR=1.6 (0.8-3.0)	Strength: control for confounding by logistic regression Weakness: Sample not representative of general population Note – UBT used
Hamajima, 1997 ¹¹⁵	Japan	192	Out-patients aged 20-70+	Male ex-smoker, OR=2.0 (0.6-7.0) Male current smoker, OR=7.8 (2.0-30.4) Female ex-smoker, OR=0.1 (0.0-2.1) Female current smoker, OR=1.2 (0.2-6.2)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population; no clear biologic mechanism; <10% of females smoked; inconsistent findings by gender

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Murray, 1997 ⁶¹	Ireland	4742	General population sample aged 12-64	Ex-smoker, OR=1.2 (1.0-1.5) Current smoker<20/day, OR=1.0 (0.8-1.2) Current smoker≥20/day, OR=1.3 (1.1-1.7)	Strength: Sample representative of general population; control for confounding by logistic regression; Weakness: Possible bias from low response rate (58%)
Peach, 1997 ⁶²	Australia	217	General population sample from electoral roll	No association with tobacco use	Strength: Sample representative of general population; control for confounding by logistic regression Weakness: Specific ORs not presented
Rudi, 1997 ¹¹⁶	Germany	457	Hospital staff aged 16-73	No significant association with tobacco use	Weakness: Sample not representative of general population; specific results not presented
Shinchi, 1997 ⁵⁴	Japan	566	Male self-defense officials aged 50-55	HP prevalence: never smoke (81.5%), ex-smoker (75.7%), <25 cigs/day (81.2%), ≥25 cigs/day (77.5%) [p for trend=0.48] (excluding ex-smokers)	Weakness: Sample not representative of general population; low serology specificity (76%); no control for confounding
Fraser, 1996 ⁸¹	New Zealand	579	Workforce survey of subjects aged 40-64	Ex-smoker, RR=1.1 (0.9-1.3) Current smoker, RR=0.9 (0.8-1.1)	Strength: Presented RR estimates adjusted for age and ethnicity Weakness: Not representative of general population; no control for SES factors
Martin-de-Argila, 1996 ⁷⁵	Spain	381	Healthy volunteers aged 5-77	Of those who smoked ≥5 cigarettes/day, 54%=HP+ and 46%=HP- (p=0.69)	Weakness: Sample not representative of general population; no control for confounding
Zhang, 1996 ¹³	China	2646	General population of rural Chinese aged 35-64	No significant association with smoking	Strength: Subjects representative of general population; control for confounding by logistic regression Weakness: Specific ORs not presented
Chow, 1995 ⁶³	Australia	328	General population of Chinese immigrants aged >25	Ex-smoker, OR=1.7 Current smoker, OR=1.3 [p>0.05]	Strength: Subjects representative of general population Weakness: Adjusted ORs from logistic regression not presented

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Fontham, 1995 ¹⁰⁹	New Orleans	321	Endoscopy clinic patients aged 18-75	Current smoker, OR=1.7 (1.0-2.9) White smoker, OR=0.6 (0.2-1.8) Black smoker, OR=3.1 (1.5-6.5)	Strength: control for confounding by logistic regression; population of uniform social class Weakness: Sample not representative of general population; inconsistent findings by race Note – biopsy and slide staining used
Gasbarrini, 1995 ⁶⁴	San Marino	2237	General population sample aged 20- 70+	No significant association with smoking	Strength: Subjects representative of general population Weakness: ORs not presented; no control for confounding
Gilboa, 1995 ⁸⁵	Rural Israel	377	General population	No significant association with smoking	Strength: Subjects representative of general population Weakness: ORs not presented; no control for confounding
Lambert, 1995 ⁹²	Australia	122	Institutionalized medically and physically handicapped young adults	No significant association with smoking	Strength: Controls representative of general population; ORs adjusted for confounding by logistic regression Weakness: Specific ORs for tobacco not presented
EUROGAST, 1993 ⁶⁸	17 Populations	3194	General population Samples aged 25- 34 and 55-64	Ex-smoker, OR=1.2 (0.9-1.5) Current smoker, OR=1.0 (0.8-1.2)	Strength: Subjects representative of general population; control for confounding by logistic regression
Bateson, 1993 ¹¹⁴	Australia	464	Out-patients with normal upper digestive tracts	HP prevalence: never smoke & ex- smoker (35.5%), current smoker (49.6%) [p<0.01]	Weakness: Sample not representative of general population; no control for confounding Note – biopsy with CLO test used

Table II-3. Studies assessing the relationship between *Helicobacter pylori* infection and alcohol use

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Brenner, 1999 ¹¹⁶	Germany	1785	Participants in German National Health and Nutrition Survey aged 18-88	>0-10 g ethanol/day, OR=0.9 (0.8-1.1) 10-<20 g ethanol/day, OR=0.8 (0.6-1.0) ≥20 g ethanol/day, OR=0.7 (0.6-0.9) Association stronger for wine than beer	Strength: Population-based sample; control for confounding by Cox regression Weakness: Population included only light and moderate drinkers
Brenner, 1999 ¹¹⁷	Germany	425	Health insurance company and their household members aged 15-69	≤75 g ethanol/wk, OR=0.6 (0.3-1.2) >75 g ethanol/wk, OR=0.3 (0.1-0.8) [p for trend=0.017] Association stronger for wine than beer	Strength: Control for confounding by logistic regression Weakness: Sample not representative of general population; included members from the same household Note – UBT used
Kikuchi, 1998 ⁸⁷	Japan	4361	Public service workers	Occasional drinker, OR=0.9 (0.7-1.1) Ex-drinker, OR=0.7 (0.3-1.6) Current drinker, OR=0.8 (0.7-1.1)	Strength: 87% participation rate Weakness: ORs only adjusted for age and sex
Lin, 1998 ⁸⁷	Australia	273	General population	Alcohol consumption, OR=0.7 (0.5-1.1)	Strength: Population-based sample; control for confounding by logistic regression Weakness: Adjusted risks not presented for categorical data
Brenner, 1997 ¹¹⁵	Germany	501	Out-patients aged 15-79	≤75 g ethanol/wk, OR=0.9 (0.5-1.6) >75 g ethanol/wk, OR=0.3 (0.2-0.7)	Strength: Control for confounding by logistic regression Weakness: Sample not representative of general population Note – UBT used
Murray, 1997 ⁶¹	Ireland	4742	General population sample aged 12-64	< Recommended limit (RL), OR=0.9 (0.7-1.1) ≤2 times RL, OR=1.0 (0.8-1.3) >2 times RL, OR=1.2 (0.9-1.8)	Strength: Sample representative of general population; control for confounding by logistic regression; Weakness: Possible bias from low response rate (58%)

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Peach, 1997 ⁶²	Australia	217	General population sample from electoral roll	No association with alcohol use	Strength: Sample representative of general population; control for confounding by logistic regression Weakness: Specific ORs not presented
Shinichi, 1997 ⁶⁴	Japan	566	Male self-defense officials aged 50-55	HP prevalence: never drink (80.0%), ex-drinker (61.9%), <50 ml/day (81.5%), ≥50 ml/day (78.0%) [p for trend=0.54] (excluding ex-drinkers)	Weakness: Sample not representative of general population; low serology specificity (76%); no control for confounding
Fraser, 1996 ⁶⁵	New Zealand	579	Workforce survey of subjects aged 40-64	Drinkers, RR=0.9 (0.8-1.1)	Strength: Presented RR estimates adjusted for age and ethnicity Weakness: Not representative of general population; no control for SES factors
Martin-de-Argila, 1996 ⁶⁵	Spain	381	Healthy volunteers aged 5-77	Of those who drank >60 g ethanol/day, 55%=HP+ and 45%=HP- [p=0.78]	Weakness: Sample not representative of general population; no control for confounding
Chow, 1995 ⁶³	Australia	328	General population of Chinese immigrants aged ≥25	Drink < weekly, OR=0.8 Drink weekly, OR=0.9 [p>0.05]	Strength: Subjects representative of general population Weakness: Adjusted ORs from logistic regression not presented
Fontham, 1995 ¹⁰⁹	New Orleans	321	Endoscopy clinic patients aged 18-75	1-2 drinks/wk, OR=0.4 (0.2-0.8) ≥3 drinks/wk, OR=0.6 (0.3-1.4) [p for trend=0.25]	Strength: control for confounding by logistic regression; population of uniform social class Weakness: Sample not representative of general population Note – biopsy and slide staining used
Lambert, 1995 ⁹²	Australia	122	Institutionalized medically and physically handicapped young adults	No significant association with alcohol use	Strength: Controls representative of general population; ORs adjusted for confounding by logistic regression Weakness: Specific ORs for alcohol not presented

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
EUROGAST, 1993 ⁶⁸	17 Populations	3194	General population sample aged 25-34 and 55-64	Some alcohol, OR=1.0 (0.8-1.2)	Strength: Subjects representative of general population; control for confounding by logistic regression

Table II-4. Studies assessing the relationship between *Helicobacter pylori* infection and diet

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Jarosz, 1999 ¹²³	Poland	51	<i>H. pylori</i> infected patients with chronic gastritis aged 21-60	8/27 (30%) had apparently successful <i>H. pylori</i> eradication after 4 weeks of vitamin C treatment (5 g/day) compared with 0/24 untreated controls [p=0.01]	Strength: Patients randomized to treatment; levels of vitamin C measured in gastric juice and serum Weakness: Follow-up limited to 4 weeks post treatment
Begue, 1998 ¹²⁷	Peru	104	Gastroenterology clinic patients aged 0-17	Food from street vendors: never [referent]; once/month, OR=2.7 (1.0-7.6); once/week, OR=2.5 (0.2-99)	Strength: ORs adjusted for age by stratification Weakness: Subjects not representative of general population, no control for other confounding factors Note – biopsy and stain used
Malaty, 1998 ¹²⁴	Sweden	270 twin pairs	Swedish Twin Registry adults, mean age=65.5	Consumption of high levels of ascorbic acid associated with <i>H. pylori</i> infection in twin reared apart. (p=0.04)	Strength: Study of twin pairs controlled for genetic effects Weakness: used paired t test – no adjustment for confounding
Goodman, 1997 ¹²¹	Columbia	684	Population-based sample of rural children aged 2-9 years	Servings of fruits & vegetables/day: ½-1 ½, OR=19.1 (4.0-91.9); 2-2 ½, OR=1.8 (1.0-3.4); 3-5 ½, OR=1.0 [referent]; 6-12, OR=0.6 (0.3-1.4) Vitamin C in mg/day: 10-39, OR=7.2 (1.5-34.1); 40-79, OR=1.3 (0.7-2.4); 80-119, OR=1.0 [referent]; 120-641, OR=0.6 (0.3-1.1) Beta-carotene in IU/day: 96-299, OR=3.1 (1.2-7.9); 300-599, OR=2.3 (1.1-5.1); 600-899, OR=1.9 (0.9-3.8); ≥900, 1.0 [referent]	Strength: Population-based sample; control for confounding by logistic regression Weakness: Possibility for differential misclassification if mothers of HP-children more socially aware of proper dietary responses; diet difficult to classify; variables highly correlated Note – UBT used

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Hamajima, 1997 ¹¹³	Japan	192	Out-patients aged 20-70+	Males: rice for breakfast, OR=3.7 (1.3-10.8); soybean paste soup every day, OR=5.2 (1.8-15.2) Females: pickled Chinese cabbage weekly, OR=2.8 (1.1-7.5); lettuce, OR=2.9 (1.1-7.8)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population; no clear biologic mechanism
Peach, 1997 ⁶²	Australia	217	General population sample from electoral roll	No association with vegetarian diet	Strength: Sample representative of general population; control for confounding by logistic regression Weakness: Specific ORs not presented
Shinchi, 1997 ⁵⁴	Japan	566	Male self-defense officials aged 50-55	HP prevalence for Tofu: low (88.6%), middle (76.7%), high (76.1%) [p for trend=0.01, but no longer significant after Bonferroni correction] No other food items related to HP+	Strength: Adjusted for rank (SES measure) by the direct method Weakness: Sample not representative of general population; low serology specificity (76%); multiple comparisons
Goodman, 1996 ¹²²	Columbia	684	Population-based sample of rural children aged 2-9 years	Raw vegetables in servings/day: 2, OR=1.5 (0.8-2.9); ≥3, OR=2.0 (0.9-4.6) Fruits & vegetables in servings/day: ≥2, OR=0.1 (0.0-0.3) Milk : ≥1/2 cups/day, OR=0.5 (0.3-1.1)	Strength: Population-based sample; control for confounding by logistic regression Weakness: Possibility for differential misclassification if mothers of HP- children more socially aware of proper dietary responses; diet difficult to classify; variables highly correlated Note – UBT used
Zhang, 1996 ¹¹	China	2646	General population aged 35-64	HP prevalence for eat sour pancakes: yes (76%), no (70%) [p<0.05]	Strength: Subjects representative of general population; control for confounding by logistic regression

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Fonham, 1995 ¹⁰⁹	New Orleans	321	Endoscopy clinic patients aged 18-75	Fruits/wk: 7-14, OR=0.8 (0.4-1.6); ≥ 14 , OR=0.5 (0.2-1.0) [p for trend=0.04] Vitamin C in mg/day: Middle 2 quartiles, OR=0.8 (0.4-1.5); Highest quartile, OR=0.4 (0.2-0.8) [p for trend=0.02]	Strength: Control for confounding by logistic regression; population of uniform social class; used validated reduced dietary questionnaire Weakness: Sample not representative of general population Note – biopsy and slide staining used
Hopkins, 1993 ¹²⁵	Chile	1815	General population <35 years of age	Uncooked vegetables , OR=3.2 (1.9-5.7) Uncooked shellfish , OR=1.3 (1.1-1.7)	Strength: Subjects representative of general population; control for confounding by logistic regression
Webberley, 1992 ²⁴⁶	UK	241	Out-patients aged 18-70	HP prevalence similar for Asian vegans (36%) and Asian meat eaters (32%) HP prevalence higher in Asians than Caucasians (18%)	Weakness: Subjects not representative of general population; no control for confounding; statistical methods not described

Table II-5. Studies assessing the relationship between *Helicobacter pylori* infection and occupational exposures

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Monés, 1999 ⁴⁰	Spain	180 44 189	Gastroenterologists Non-gastroenterologists Healthy controls	HP prevalence: gastroenterologists (53.3%) [endoscopists (53.3%); non-endoscopists (53.5%)], non-gastroenterologists (50.0%), controls (51.9%)	Strength: Comparison of endoscopists with non-endoscopists and gastroenterologists with other physicians as well as controls; Weakness: No control for confounding Note – UBT used
Lin, 1998 ³⁰	Australia	92 40 295 39 107	Dentists Dental nurses Controls Gastroenterologists Endoscopy nurses	Dentists vs controls, OR=0.6 (0.3-1.1) Dental nurses vs controls, OR=1.0 (0.4-2.9) HP prevalence: gastroenterologists (80%) vs dentists (21%) [p<0.01], especially those employed ≥11 years HP prevalence for endoscopy (38%) vs dental (23%) nurses [p>0.05]	Strength: Representative sample of dentists and dental nurses; population-based sample of controls; control for confounding by age and sex by logistic regression Weakness: Assumes exposure differences due to occupational rather than some other exposure
Nishikawa, 1998 ⁴²	Japan	92 29 101	Gastroenterologists Endoscopy nurses Healthy controls	HP prevalence: endoscopy personnel (29.8%), controls (24.8%) [p=0.406]	Strength: Controls matched to endoscopy personnel by age Weakness: No control for confounding factors other than age
Shelley, 1998 ³⁸	US	54	Anesthesiologists	HP prevalence 4% in anesthesiologists compared to 19% in 2 published US control groups [p<0.01]	Weakness: no control for confounding; controls groups from different population than anesthesiologists
Bohmer, 1997 ³⁸	Netherlands	254	Employees of institutes for Intellectually Disabled aged 18-50	HP prevalence: staff with close physical contact to patients (31.6%) vs other staff (14.1%) [p=0.002] OR=2.8 (1.1-7.0) for those employed >5 years	Weakness: Statistical tests used not specified, no apparent control for confounding

Author, Year, Ref No,	Country	No. in Study	Population	Outcome	Comments
Braden, 1997 ¹⁴¹	Germany	922 538 169 66 413	Gastroenterologist Other physicians Endoscopy nurses Other nurses Controls	Case/control comparisons: endoscopy physicians vs controls, OR=1.6 (1.3-2.1); other physicians vs controls, OR=1.5 (1.2-2.0); endoscopy nurses vs controls, OR=1.8 (1.2-2.6); other nurses vs controls, OR=1.6 (1.0-2.6); endoscopy staff vs general medical staff, OR=1.1 (0.9-1.3)	Strength: Comparison of endoscopists with other physicians as well as controls; control for confounding by age by logistic regression Weakness: Study can't differentiate risk between transmission by endoscopy and colonoscopy since these physicians do both in Germany Note – UBT used
Potts, 1997 ¹³⁵	UK	30 30	Gastroenterologists Respiratory doctors	15/30 gastroenterologists & 3/30 respiratory physicians had positive breath tests for HP [p<0.001] Risk may be lower for gastroenterologists who wear gloves	Weakness: Specifics of population selection not clear; statistical tests used not specified
Rudi, 1997 ¹¹⁶	Germany	110 272 75	1) Nonmedical staff 2) Medical staff 3) Endoscopy staff	HP prevalence: Group 1 (35.5%), Group 2 (34.6%), Group 3 (24.0%). No significant differences in the 3 groups	Strength: Adjusted for age, duration of experience, and number of years worked in unit Weakness: Statistical tests used not specified
Friis, 1996 ¹³⁸	Sweden	151 138	Sewage workers Other municipal workers	No increased risk for HP infection among sewage workers, RR=1.0 (0.7-1.5) and OR=0.90 (CI=0.5-1.5)	Strength: RRs calculated, ORs controlled for confounding by age and region by logistic regression; referent group of workers frequency matched by age, SES, and region Weakness: No control for confounding effects of other potential risk factors
Goh, 1996 ¹³⁴	Malaysia	82 53	Endoscopy staff Non-endoscopy staff	Endoscopy staff had a higher prevalence of HP than non-endoscopy staff (32.9% vs 11.3%, p=0.004)	Weakness: No control for confounding factors; mean age 2.9 years older for endoscopy staff Note – UBT used

Author, Year, Ref No,	Country	No. in Study	Population	Outcome	Comments
Liu, 1996 ¹³⁵	China	1050 702	Medical staff Healthy out-patient controls	Medical staff vs controls, OR=2.9 (2.4-3.4); endoscopists vs internists, OR=2.2 (1.4-3.7); endoscopist vs general nurses, OR=2.4 (1.4-4.1) Worked as an endoscopist for ≥ 5 years, OR=3.7 (1.4-10.1)	Strength: Comparison of endoscopists with other medical staff and "healthy" controls; includes a large variety of medical staff Weakness: No control for confounding other than age
Su, 1996 ¹³⁶	Taiwan	70 64	Endoscopists Other physicians	HP prevalence: endoscopists (80.0%) vs other physicians (51.6%) [$p<0.05$] HP prevalence: < 30 endoscopies/week (70.3%) vs ≥ 30 endoscopies/week (90.9%) [$p=0.01$]	Strength: Excluded subjects with serology values in the "gray zone"; control doctors has similar exposures except for endoscopy Weakness: No comparison with general population controls
Banatvala, 1995 ¹³⁷	Wales	70 47 62 179	Dentists Clinical students Pre-clinical students Matched controls	HP prevalence: dentists (16%), clinical dental students (6%), pre-clinical dental students (10%). Case/control comparisons: Dentists (OR=0.6, CI=0.2-1.5), clinical students (OR=1.0, CI=0.3-3.7), pre-clinical students (OR=1.0, CI=0.2-5.4)	Strength: Controls matched on age, sex, SES; matched analysis controlled for these confounding factors Weakness: Controls not representative of general population; response rate in dentists=69%
Chong, 1994 ¹³⁸	US	111 11 510	Gastroenterologists Endoscopy nurses Blood donors	HP prevalence: gastroenterologists/endoscopy nurses (53%) vs controls (14%) [$p<0.001$] HP prevalence: gastroenterologists (52%), endoscopy nurses (55%)	Strength: Subgroup analyses were done on "comparable" endoscopy professionals and controls Weakness: Volunteer control group obtained from one geographic area (Charlottesville, VA) may not be comparable to endoscopy group attending advanced course in Miami, FL - lacked controls except for US born Caucasians

Author, Year, Ref No,	Country	No. in Study	Population	Outcome	Comments
Lin, 1994 ¹³⁸	Australia	39 107 25 42 273	Gastroenterologists Endoscopy nurses Internists General nurses Population controls	Case/control comparisons: gastroenterologists, OR=5.0 (CI=2-12); Internists, OR=1.0 (0.3-3.1); endoscopy nurses, OR=0.6 (0.3-1.2); general nurses, OR=0.9 (0.3-2.5) Gastroenterologists vs internists, OR=1.8 (1.1-4.6)	Strength: Controls representative of general population; subjects age- and sex matched Caucasians; logistic regression analysis controlled for these confounding factors
Malaty, 1992 ¹³⁹	US	239	Dental workers volunteers from Texas	Dentists + dental hygienist vs. assistants + dental students, OR=2.3 (1.0-5.3) HP prevalence: white dental workers (16%), nonwhite dental workers (46%); white community (27%), black community (58%)	Strength: Compared dentists with other dental workers as well as with asymptotic controls; control for confounding by logistic regression Weakness: Control group may not be comparable on age and other factors
Mitchell, 1989 ¹³⁹	Australia	33 68 35 715	Gastroenterologists Endoscopy nurses General practitioners Normal blood donors	HP prevalence: gastroenterologists (51%) vs. blood donors (21%) [$p<0.01$]; endoscopy nurses (19%) and general practitioners (28%) not different from blood donors	Strength: Comparison of endoscopists with endoscopy nurses, and general practitioners with volunteer controls Weakness: Control group not representative of general population, may be unusual since highest HP+ found in the youngest age group (21-30)

Table II-6. Studies assessing the relationship between *Helicobacter pylori* infection and waterborne exposures

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Elitsur, 1998 ¹⁴⁵	US	1164	West Virginia children	Source of water supply, city or well, was not significantly related to HP status	Strength: large study of children; controlled for confounding by multiple regression Weakness: Subjects not representative of "healthy" children; bias may have occurred in selection of participants; results hard to interpret
Goodman, 1996 ¹²²	Columbia	684	Population-based sample of rural children aged 2-9 years	Lifetime drinking water source: Well/pump only [referent] Tap, OR=1.0 (0.5-2.0) Stream, OR=2.8 (1.2-6.8) Swims in rivers/streams: < once a year [referent] several times /year, OR=3.3 (1.2-9.4) Swims in swimming pools: < once a year [referent] several times /year, OR=3.6 (1.5-8.5)	Strength: Population-based sample; control for confounding by logistic regression Weakness: Possibility for differential misclassification if mothers of HP- children more socially aware of proper hygiene responses; variables highly correlated Note – UBT used
Malaty, 1996 ⁷⁴	Korea	161	Adult volunteers aged 20-75	Water source: well [referent] City, OR=1.4 (1.2-2.6) Bottled, OR=1.7 (1.4-6.8)	Strength: Control for confounding by logistic regression Weakness: Sample not representative of general population; over representation of middle class
Matysiak-Budnik, 1996 ⁷⁶	Poland	181	Healthy volunteers aged 0-34	Used well water, OR=1.1 (0.2-7.3)	Weakness: Sample not representative of general population; no control for confounding; high rate of HP+
Zhang, 1996 ¹³	China	2646	General population of rural adults aged 35-64	HP prevalence - drinking water source: Ponds or ditches (88%) Deep wells (73%) Surface wells (72%) No statistically significant differences	Strength: Subjects representative of general population; control for confounding by logistic regression Weakness: Only a small percentage obtained their water from ditches

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Teh, 1994 ⁶⁷	Taiwan	555 268	Adults Children General population	Drinking water source: Tap water [referent] River water, OR=0.9 (0.6-1.5) Well water, OR=1.2 (0.8-1.7)	Strength: Subjects representative of general population; age-adjusted ORs Weakness: No control for other confounding factors
Hopkins, 1993 ¹²⁵	Chile	1815	General population <35 years of age	Swimming near contaminated beaches, OR=1.1 (0.8-1.5) Bathing in fresh water rivers, irrigation ditches, or lakes, OR=1.4 (0.8-2.4)	Strength: Subjects representative of general population; control for confounding by logistic regression
Klein, 1991 ¹²⁴	Peru	407	Children aged ≤ 12 of low and high SES	External vs. internal water sources, OR=3.4 [p=0.001] Municipal vs. community wells (high income only), OR=12.6 [p=0.02]	Strength: Control for age by logistic regression Weakness: Subjects not representative of general population; confounded by SES as no subject of high SES had external water sources Note – UBT used

Table II-7. Studies assessing the relationship between *Helicobacter pylori* infection and exposure to pets and other animals

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Dore, 1999 ¹⁸	Sardinia	123 30 509	Shepherds Family members Blood donors aged 17-75	HP prevalence among shepherds (98%) greater than among family members (73%), OR=21.9 (4.5-106) or blood donors (43%), OR=78.8 (20-311)	Strength: Serology confirmed by endoscopy Weakness: Controls not representative of general population, no control for confounding
McIsaac, 1999 ¹⁸	Canada	15779	National population health survey	No association between dog/cat ownership and history of peptic ulcer disease (OR=1.1, CI=0.9-1.4)	Strength: Sample representative of general population; large study population; control for confounding by logistic regression Weakness: Dogs and cats grouped together; only asked about current pet ownership; diagnosis of peptic ulcer not independently validated
Begue, 1998 ¹⁷	Peru	104	Gastroenterology patients aged 0-17	No association with exposure to domestic animals (dogs, cats, birds)	Weakness: Subjects not representative of general population, no control for confounding factors Note – biopsy and stain used
Bode, 1998 ¹⁸	Germany	685	General population sample of preschool children	Has pets (OR=1.1, CI=0.6-2.4) Has cat (OR=1.9, CI=0.7-5.1) Has rabbit (OR=1.7, CI=0.6-4.7)	Strength: Population-based sample; control for confounding by logistic regression Weakness: Prevalence of infection only 6.3%; little power to detect significant association Note – UBT used
Luzza, 1998 ⁹¹	Rural Italy	705	Adult and Children Out-patients aged 1-87	No difference in the percentage of HP+ and HP- subjects who had household pets during childhood	Weakness: Subjects not representative of general population; no control for confounding; no statistical tests

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Rothenbacher, 1998 ⁸⁵	Germany	337	Adult students aged 50-80	No association between HP infection and occupational handling of animals or having a pet or a dog in the household during childhood HP prevalence - cat in the house during childhood: No (30.7%) Yes (44.7%) [p=0.017]	Strength: Assess current HP status with UBT Weakness: Not representative of general population; no control for confounding; no detail of statistical test used
Peach, 1997 ⁶³	Australia	217	General population sample from electoral roll	No association with occupational or recreational contact with animals	Strength: Sample representative of general population; control for confounding by logistic regression Weakness: Specific ORs not presented
Rothenbacher, 1997 ⁹⁷	Germany	501	Out-patients aged 15-79	No association between HP infection and having a pet, dog, or cat in the household (compared % positivity)	Weakness: Sample not representative of general population; no control for confounding except by age Note – UBT used
Goodman, 1996 ¹²²	Columbia	684	Population-based sample of rural children aged 2-9 years	Household has rabbits (OR=0.3, CI=0.1-0.7) Child plays with sheep (OR=4.5, CI=0.7-30.5)	Strength: Population-based sample; control for confounding by logistic regression Weakness: Estimate for contact with sheep lacks precision Note – UBT used
Staat, 1996 ¹²⁷	US	2581	6-19 years old participants in National survey	HP prevalence – any pets in household: Any (19.0%) None (31.5%) [p<0.05]	Strength: Population-based sample; prevalences weighted to account for oversampling and lack of response
Webb, 1996 ²⁰⁰	UK	447	Male volunteers aged 18-65	Pets during childhood: None [referent] Cat (OR=1.4, CI=0.7-2.9) Other (OR=0.6, CI=0.6-2.5)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population
Webb, 1994 ¹⁸	UK	471	Male volunteers aged 18-65	HP prevalence - household pets: No (34.7%) Yes (38.2%) [p=0.5]	Weakness: Subjects not representative of general population; no control for confounding

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Graham, 1991 ⁴⁰	Texas Blacks Whites	246 239	Healthy volunteers aged 15-80	Have pets, OR=0.5 [p=0.009]	Strength: control for confounding by logistic regression Weakness: Subjects not representative of general population; owning pets thought to be surrogate for high social class Note -- UBT used

Table II-8. Studies assessing the relationship between *Helicobacter pylori* infection and hygiene practices

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Luzza, 1998 ⁹³	Rural Italy	705	Out-patients	Possession of a bathroom (HP+=85%, HP-=92%), a refrigerator (HP+=43%, HP-= 62%), and a hot water supply (HP+=26%, HP-= 48%) during childhood was reported by fewer HP+ than HP- subjects (not significant after adjustment for age)	Weakness: Subjects not representative of general population; no control for confounding; statistical results not presented
Peach, 1997 ⁸²	Australia	217	General population sample from electoral roll	Did not usually wash hands after toilet, OR=4.1 (0.5-35.0) No association with frequency of sharing toothbrush or cup or washing hands before eating	Strength: Sample representative of general population; control for confounding by logistic regression Weakness: only a small number reported infrequent handwashing; ORs not presented for sharing toothbrush or cup or washing hands before eating
Rothenbacher, 1997 ⁹⁷	Germany	501	Out-patients aged 15-79	HP prevalence - shared towel with another household member: Yes (15.9%), No (25.9%) [p=0.199]	Weakness: Sample not representative of general population; no control for confounding except by age Note - UBT used
Breuer, 1996 ⁹⁵	Germany	260	Blood donors aged 18-61	No WC in house now, OR=1.0 (0.4-3.0) No WC in house age 8, OR=2.6 (1.1-6.4) No hot water now, OR=3.2 (0.2-35.6) No hot water age 8, OR=1.8 (0.9-3.7)	Strength: All subjects approached participated Weakness: Subjects not representative of general population; no control for confounding;
Czkwianianc, 1996 ¹³⁰	Poland	240	Children aged 6 months-17 years who were patients at hospitals and clinics	Prevalence of HP+ higher in children who lived in households where the level of hygiene and sanitation were the poorest [p<0.005]	Weakness: Sample not representative of general population; no control for confounding; statistical tests not specified

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Goodman, 1996 ¹²²	Columbia	684	Population-based sample of rural children aged 2-9 years	Shares cups, OR=1.8 (1.1-3.1) Mother rarely washes hands with soap, OR=2.7 (1.1-6.6) Location of latrine relative to hand washing facility: < 25 meters [referent] ≥ 25 meters, OR=1.6 (0.8-2.9) No latrine, OR=2.2 (0.9-5.3)	Strength: Population-based sample; control for confounding by logistic regression Weakness: Possibility for differential misclassification if mothers of HP-children more socially aware of proper hygiene responses; exposures difficult to classify; variables highly correlated Note – UBT used
Chow, 1995 ⁶¹	Australia	328	General population of Chinese immigrants aged ≥25	Used chopsticks, OR=2.5 [p<0.001]	Strength: Subjects representative of general population; Weakness: Adjusted ORs from logistic regression not presented
Mcgraud, 1995 ¹⁵¹	Western Africa	38 97	HP+: mother and child HP-: mother and child	Premastication of infant's food by mother, OR=2.9 (0.9-9.0)	Weakness: Pilot study embedded in review paper; no details for design or analysis provided; no apparent control for confounding factors Note – serology and/or breath test was used
Teh, 1994 ⁶⁷	Taiwan	555 268	Adults Children General population	Have other than flushing type toilet, OR=1.0 (0.6-1.6)	Strength: Subjects representative of general population; age-adjusted ORs Weakness: No control for other confounding factors
Webb, 1994 ¹⁴	UK	471	Male volunteers aged 18-65	HP prevalence – bathroom: No (42.0%) vs. Yes (35.1%) [p=0.2] HP prevalence – inside toilet: No (40.3%) vs. Yes (35.5%) [p=0.3]	Weakness: Subjects not representative of general population; no control for confounding

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Hammermeister ¹³² 1992	Germany	64	Submarine crews	HP prevalence – German submarine crews (31.7%) vs. German air force recruits (16.2%), OR=2.4 [p<0.05]	Strength: Prospective study, with 3-4 blood samples in a 1 year period - no seroconversion in any group Weakness: Authors used logistic regression to control for age, but not SES
	France	74 51 135	New Air Force staff Infantry ≥3 years New Infantry	HP prevalence – French infantry regular officers (13.7%) vs. French infantry recruits (18.5%) NS	
Mendall, 1992 ¹⁰⁰	UK	215	Subjects aged 18-82 attending a health-screening clinic	No fixed hot water supply in house at age 8, OR=4.3 (1.9-10.0)	Strength: Control for confounding by logistic regression; age 8 used as mid-point of childhood and to enhance recall Weakness: Subjects not representative of general population

Table II-9. Studies assessing the relationship between *Helicobacter pylori* infection and density/crowding

Author, Year, Ref No.	Country	No. In Study	Population	Outcome	Comments
Alagathanam, 1999 ⁵⁵	India	354	General upper class population aged 12-70+	HP prevalence – Family size <5 (45.4%) vs. ≥5 (56.7%) [p<0.05]	Strength: Representative of an urban class population Weakness: No control for confounding; no adjustment for multiple subjects from same household
Kikuchi, 1998 ⁸⁷	Japan	4361	Public service workers	Sibship size: 1 [referent] 2-3, OR=1.2 (0.8-1.6) 4+, OR=1.5 (1.0-2.1) [p=0.013]	Strength: 87% participation rate Weakness: ORs only adjusted for age and sex
Luzza, 1998 ⁹¹	Rural Italy	705	Out-patients	Number of people/room during childhood: HP+=3.2±2.4, HP-=2.1±1.9 [p<0.5] Number of siblings: HP+=4.6±2, HP-=3.7±2 [p<0.05]	Weakness: Subjects not representative of general population; control for confounding in logistic regression, but ORs based on continuous variables not that informative compared to means
Malaty, 1998 ¹²⁴	Sweden	270 twin pairs	Swedish Twin Registry adults, mean age=65.5	Crowded living conditions and poor economic conditions associated with HP+ in twins reared apart (p=0.02)	Strength: Study of twin pairs controlled for genetic effects Weakness: used paired t test – no adjustment for confounding
Rothenbacher, 1998 ⁸⁵	Germany	337	Adults students aged 50-80	Shared a bed with siblings or parents during childhood, OR=2.6 (1.1-6.1)	Strength: Control for confounding by logistic regression Weakness: Not representative of general population Note – UBT used
Stroffolini, 1998 ⁸⁶	Italy	1659	Military students	≥1 sibling, OR=1.3 (1.0-1.7)	Strength: control for confounding by logistic regression Weakness: sample not representative of general population; sample may be too heterogeneous

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Torres, 1998 ⁶⁰	Mexico	9116	General population aged 1-39	Crowding (persons/room): 1.5 [referent] 1.6-3.5, OR=1.3 (1.1-1.5) ≥3.6, OR=1.4 (1.2-1.6)	Strength: Sample representative of general population; age-adjusted ORs Weakness: no control for other confounding factors
Peach, 1997 ⁶³	Australia	217	General population sample from electoral roll	≥6 people in household at age 0-16, OR=2.5 (1.1-5.5)	Strength: Sample representative of general population; control for confounding by logistic regression
Rothenbacher, 1997 ⁶⁷	Germany	501	Out-patients aged 15-79	Sibship size, 0 [referent] 1, OR=1.2 (0.5-2.7) 2-3, OR=1.5 (0.7-3.4) ≥4, OR=2.7 (1.2-6.1)	Strength: Control for confounding by logistic regression Weakness: Sample not representative of general population Note – UBT used
Shinchi, 1997 ⁶⁴	Japan	566	Male self-defense officials aged 50-55	HP prevalence – rank Low (82.5%) Middle (75.2%) High (75.0%) [p for trend=0.48]	Weakness: Sample not representative of general population; low serology specificity (76%); no control for confounding
Breuer, 1996 ⁶⁵	Germany	260	Blood donors aged 18-61	A) ≥1 child in household at present, OR=1.8 (1.0-3.2) B) >1 person/room now, OR=2.4 (1.2-4.9) C) >1 person/room at age 8, OR=2.1 (1.1-4.3)	Strength: All subjects approached participated; control for confounding by logistic regression for A and C Weakness: Subjects not representative of general population; only control for age for B
Clemens, 1996 ¹⁵⁴	Bangladesh	257	Population-based sample of children aged 2-5	HP+ children had a higher median number of persons per sleeping room (3.8 vs. 3.2) [p<0.05]	Strength: Sample representative of general population; control for confounding Weakness: Population may have been too homogeneous to see striking SES differences
Czkwianiance, 1996 ¹⁵⁰	Poland	240	Children aged 6 months-17 years who were patients at hospitals and clinics	Prevalence of HP+ higher in children who lived in households where number of persons per room was higher [p<0.005]	Weakness: Sample not representative of general population; no control for confounding; statistical tests not specified

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Goodman, 1996 ¹²²	Columbia	684	Population-based sample of rural children aged 2-9 years	Number of persons in family: 3 [referent] 4-5, OR=3.7 (1.2-11.4) 6-8, OR=6.3 (1.9-20.5) 9-15, OR=8.0 (2.2-28.9)	Strength: Population-based sample; control for confounding by logistic regression Note – UBT used
Malaty, 1996 ¹⁴	Korea	161	Adult volunteers aged 20-75	≥5 family members in house, OR=1.2 (1.1-4.5)	Strength: Control for confounding by logistic regression Weakness: Sample not representative of general population; over representation of middle class
Malaty, 1996 ¹⁴	Russia	213	Asymptomatic adults aged 20-75	Crowding index, Low [referent] Moderate, OR=0.7 (0.6-2.2) High, OR=1.4 (1.1-4.5)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population; Russian society is very homogeneous
McCallion, 1996 ¹⁵³	Ireland	367	3-15 year old outpatients	Household density (persons/room), <0.7 [referent] 0.7-1.0, OR=2.0 (1.0-3.9) >1.0, OR=3.4 (1.7-6.6) Sharing a bed with a parent, Never [referent] 1-2 nights/week, OR=2.3 (1.2-4.3) >2 nights/week, OR=3.0 (1.4-6.4)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population
Staat, 1996 ¹⁴⁷	US	2581	6-19 year old participants in National survey	Crowding index (persons/room) <0.5 [referent] 0.5-0.99, OR=1.8 (1.0-3.4) 1.0-1.99, OR=2.1 (1.3-3.4) ≥2, OR=5.6 (2.9-10.9)	Strength: Population-based sample; control for confounding by logistic regression

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Replogle, 1995 ⁴⁵	California	556	Health plan members aged 20- 39	Number of children Lived with as a child, Didn't live with children [referent] 1-2, OR=1.1 (1.0-1.2) 3-4, OR=1.3 (1.1-1.5) ≥5, OR=1.4 (1.2-1.5) [p for trend≤0.01] Lived with children as an adult, OR=1.9 (1.4-2.6)	Strength: Calculated RRs Weakness: Subjects not representative of general population; no control for confounding
Malaty, 1994 ⁷⁷	Texas	150	Black & Hispanic Healthy volunteers aged 19-49	Crowding index, Low [referent] Middle, OR=3.1 (2.1-4.1) High, OR=4.5 (3.3-5.7)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population
Teh, 1994 ⁶⁷	Taiwan	268	Children General population	Sibship size, 0-1 [referent] 2, OR=1.6 (0.7-3.8) ≥3, OR=3.9 (1.6-9.5)	Strength: Subjects representative of general population; age-adjusted ORs Weakness: No control for other confounding factors
Webb, 1994 ⁷⁸	UK	471	Male volunteers aged 18-65	Crowding - >1 per room, OR=1.8 (1.1-4.0) Shared a bed, OR=1.6 (1.0-2.6)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population; collinearity of density measures

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Hammermeister 1992 ¹³²	Germany	64	Submarine crews	HP prevalence – German submarine crews (31.7%) vs. German air force recruits (16.2%), OR=2.4 [p<0.05]	Strength: Prospective study, with 3-4 blood samples in a 1 year period - no seroconversion in any group Weakness: Authors used logistic regression to control for age, but not SES
	France	74 51 135	New Air Force staff Infantry ≥3 years New Infantry	HP prevalence – French infantry regular officers (13.7%) vs. French infantry recruits (18.5) NS	
Mendall, 1992 ¹⁰⁰	UK	215	Subjects aged 18-82 attending a health-screening clinic	Persons per room in childhood <0.7 [referent] 0.7-0.99, OR=1.4 (0.4-4.4) 1.0-1.29, OR=4.0 (1.4-11.8) ≥1.3, OR=6.2 (1.8-18.6)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population

Table II-10. Studies assessing the relationship between *Helicobacter pylori* infection and social factors

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Buckley, 1998 ⁷²	Ireland	1000	Blood donors aged 18-60	HP prevalence – social class*: I/II (36.9%) III (44.0%) IV/V (50.0%) [p<0.01]	Weakness: No control for confounding; statistical test used unclear; non-representative population
Lin, 1998 ⁷³	Australia	273	General population Aged 20-80	Household income (increase in risk with decrease in income), OR=1.5 (1.1-2.4)	Strength: Population-based sample; control for confounding by logistic regression Weakness: Adjusted risks not presented for categorical data
Luzza, 1998 ⁷⁴	Rural Italy	705	Out-patients aged 1-87	Nonmanual occupation: OR=0.7 (0.5-0.9) Childhood social class low: HP+=54%, HP-=32%	Strength: Occupation controlled for confounding by logistic regression Weakness: Subjects not representative of general population; no control for confounding or statistical tests for childhood social class
Malaty, 1998 ⁷⁴	Sweden	270 twin pairs	Swedish Twin Registry adults, mean age=65.6	Crowded living conditions and poor economic conditions associated with HP+ in twins reared apart (p=0.02)	Strength: Study of twin pairs controlled for genetic effects Weakness: used paired t test – no adjustment for confounding
Rothenbacher, 1998 ⁷⁵	Germany	337	Adult students aged 50-80	Education of father: ≤9 years [referent] 10-11 years, OR=0.8 (0.5-1.5) ≥12 years, OR=0.4 (0.2-0.7)	Strength: Control for confounding by logistic regression Weakness: Not representative of general population Note – UBT used
Souto, 1998 ⁷⁶	Brazil	184	General population aged 10-90	Annual family income – decrease in risk with increase in income [p for trend=0.013]	Strength: Population-based sample Weakness: No control for confounding

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Strachan, 1998 ⁴⁴	South Wales	1796	Population-based Prospective Heart Disease Study participants aged 45-59	HP prevalence - Social class at entry*: I/II (60.9%) IIINM (65.8%) IIIM (72.7%) IV/V (76.7%) [p<0.0001] Years of father's schooling ≤8, OR=1.2 (0.9-1.6)	Weakness: No control for confounding; statistical test used not specified
Stroffolini, 1998 ⁴⁶	Italy	1659	Military students		Strength: control for confounding by logistic regression Weakness: sample not representative of general population; sample may be too heterogeneous
Torres, 1998 ⁴⁸	Mexico	9116	General population aged 1-39	Education level College [referent] High school, OR=1.4 (1.2-1.8) Basic, OR=2.0 (1.6-2.4) Illiterate, OR=2.4 (1.7-3.4)	Strength: Sample representative of general population; age-adjusted ORs Weakness: no control for other confounding factors
Murray, 1997 ⁴¹	Ireland	4742	General population sample aged 12-64 from electoral roll	HP prevalence - social class*: Increasing trend from I (43.0%) to V (68.1%) [p for trend>0.001] OR=2.8 (1.8-4.4) for V vs. I	Strength: Sample representative of general population; control for age and sex by logistic regression; Weakness: Possible bias from low response rate (58%)
Peach, 1997 ⁴³	Australia	217	General population sample from electoral roll	Negative HP status significantly associated with increasing education, OR=0.6 (0.4-1.0) [p<0.05]	Strength: Sample representative of general population; control for confounding by logistic regression
Rothenbacher, 1997 ⁴⁷	Germany	501	Out-patients	HP prevalence - school education: ≤9 years (30.5%) 10-11 years (20.5%) >12 years (14.4%) [p=0.051]	Weakness: Sample not representative of general population; no control for confounding except by age Note - UBT used

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Shinchi, 1997 ³⁴	Japan	566	Male self-defense officials aged 50-55	HP prevalence -- rank Low (82.5%) Middle (75.2%) High (75.0%) [p for trend=0.48]	Weakness: Sample not representative of general population; low serology specificity (76%); no control for confounding
Fraser, 1996 ³⁵	New Zealand	579	Workforce survey of subjects aged 40-64	The RR of HP infection for SES and income were not statistically significant after adjusting for age and ethnicity (European, Maori, Pacific Islander). However, HP status varied with SES in Europeans: 1 & 2 [high] (21%) 3 (42%) 4 (45%) 5 & 6 [low] (54%) [p=0.011]	Strength: Presented RR estimates for SES in total group adjusted for age and ethnicity Weakness: Not representative of general population; SES in Europeans does not appear to be adjusted for age
Malaty, 1996 ³⁶	Korea	161 252	Volunteer Adults Children	Adult social class, High [referent] Middle, OR=1.0 (1.0-2.2) Low, OR=1.7 (1.7-18.0) Child social class, High [referent] Middle, OR=2.5 (1.1-5.5) Low, OR=5.2 (1.5-17.4)	Strength: Control for confounding by logistic regression Weakness: Sample not representative of general population; over representation of middle class
Malaty, 1996 ³⁸	Russia	213 307	Adults Children Out-patients	Income below the poverty level was associated with HP seroprevalence in children (OR=1.4, CI=1.2-2.5), but not in adults (OR=0.3, CI=0.2-0.8)	Strength: Control for confounding by logistic regression for poverty level in adults Weakness: Subjects not representative of general population; control only for age in analysis of poverty level for children; Russian society is very homogeneous

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
McCallion, 1996 ¹³³	Ireland	367	3-15 year old outpatients	Social class – manual vs. nonmanual, OR=1.6 (1.0-2.6)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population
Staat, 1996 ¹⁴⁷	US	2581	6-19 year old participants in National survey	Income below poverty line, OR=1.5 (1.0-2.2) Head of household education level < 12 years, OR=1.8 (1.2-2.7)	Strength: Population-based sample; control for confounding by logistic regression
Chow, 1995 ⁶³	Australia	328	General population of Chinese immigrants aged >25	Years of education, ≥ 13 [referent] 10-12, OR=2.1 7-9, OR=1.5 <7, OR=3.9 [p<0.001]	Strength: Subjects representative of general population Weakness: Adjusted ORs from logistic regression not presented
Gilboa, 1995 ⁶⁵	Rural Israel	377	General population sample aged 30-90	There was no difference in HP seroprevalence with educational level	Strength: Subjects representative of general population Weakness: ORs not presented; no control for confounding
Murray, 1995 ⁶⁶	Ireland	2115	Subjects aged 25-64 selected from General Practitioner lists	HP prevalence – social class*: I (38.0%) II (45.3%) IIINM (52.4%) IIIM (66.5%) IV (70.4%) V (66.0%)	Strength: Subjects representative of general population Weakness: ORs not presented; no control for confounding
Replogle, 1995 ⁴⁹	California	556	Health plan members aged 20-39	Years of education, < 12 [referent] 12, OR=0.6 (0.5-0.9) 12-16, OR=0.4 (0.3-0.5) >16, OR=0.3 (0.2-0.5)	Strength: Calculated RRs Weakness: Subjects not representative of general population; no control for confounding

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Malaty, 1994 ⁷⁷	Texas	150	Black & Hispanic healthy volunteers aged 19-49	Present social class not High, OR=2.2 (1.5-2.9) Childhood social class, High [referent] Middle, OR=8.9 (7.3-10.5) Low, OR=54.6 (52.7-56.5)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population
Teh, 1994 ⁶⁷	Taiwan	823	Children and Adults General population	Family income/month, ≤\$55 [referent] \$556-\$926, OR=1.0 (0.6-1.8) \$927-\$1481, OR=1.0 (0.6-1.8) >\$1482, OR=0.9 (0.5-1.7)	Strength: Subjects representative of general population; age-adjusted ORs Weakness: No control for other confounding factors
Webb, 1994 ⁷⁸	UK	471	Male volunteers	Manual occupation, OR=2.2 (1.1-4.6) Father manual occupation, OR=2.0 (0.9-4.9)	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population
EUROGAST, 1993 ⁶⁸	17 Populations	3194	General population sample aged 25-34 and 55-64	Education level, Primary [referent] Secondary, OR=0.8 (0.6-1.0) Higher, OR=0.6 (0.4-0.8)	Strength: Subjects representative of general population; control for confounding by logistic regression
Hopkins, 1993 ¹²⁵	Chile	1815	General population <35 years of age	Low SES, OR=1.7 (1.3-2.1)	Strength: Subjects representative of general population; control for confounding by logistic regression
Malaty, 1992 ⁷⁹	Texas	108	Asymptomatic healthy Hispanic volunteers	Education level, College graduate [referent] High school graduate, OR=2.7 (1.6-12) Didn't graduate high school, OR=3.6 (1.8-9.7)	Strength: Control for confounding by logistic regression; study of risk factors in Hispanics Weakness: Subjects not representative of general population

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Mendall, 1992 ¹⁰⁰	UK	215	Subjects aged 18-82 attending a health-screening clinic	Current social class, I/II [referent] IIIM, OR=0.7 (0.3-2.1) IIIM, OR=2.2 (0.8-5.5) IV/V, OR=0.6 (0.2-1.7) [p for trend=0.9]	Strength: Control for confounding by logistic regression Weakness: Subjects not representative of general population
Graham, 1991 ⁸⁰	Texas Blacks Whites	246 239	Healthy volunteers aged 15-80	There was a significant inverse gradient between age-adjusted frequency of educational level and HP infection [p=0.001]	Strength: control for confounding by logistic regression Weakness: Subjects not representative of general population Note – UBT used

Social class – I=professional occupations, II=managerial occupations, III=skilled occupations (subdivided as nonmanual (NM) and manual (M)), IV=semiskilled occupations, and V=unskilled occupations

Table II-11. Studies assessing the relationship between *Helicobacter pylori* infection and family history of gastric disease/ulcer

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Brenner, 1998 ¹⁵⁵	Germany	863	Pre-school children out-patients	History of ulcer in mother, OR=11.7 (3.8-36.2) History of ulcer in father, OR=1.6 (0.4-5.5)	Strength: 79% participation rate; both variables controlled for confounding by logistic regression Weakness: mother's history only adjusted for age and sex
Elitsur, 1998 ¹²⁵	West Virginia	303	1-20 year old children attending hospitals/clinics	No difference in the percentage of HP+ (9.8%) and HP- (9.4%) children with a family history of peptic ulcer disease [p=0.86]	Strength: large study of children; controlled for confounding by multiple regression Weakness: Subjects not representative of "healthy" children; bias may have occurred in selection of participants; results hard to interpret
Kikuchi, 1998 ¹⁷	Japan	4361	Public service workers	History of gastric disease in mother, OR=1.2 (0.9-1.7) History of gastric disease in father, OR=1.5 (1.2-1.8)	Strength: 87% participation rate; father's history controlled for confounding by logistic regression Weakness: mother's history only adjusted for age and sex
Breuer, 1996 ⁷³	Germany	260	Blood donors aged 18-61	Family history of ulcer, OR=1.3 (0.6-2.5)	Strength: All subjects approached participated Weakness: Subjects not representative of general population; no control for confounding
Martin-de-Argila, 1996 ⁷⁵	Spain	381	Healthy volunteers aged 5-77	Of those whose first degree relatives had peptic ulcer disease, 63%=HP+ and 37%=HP- [p=0.028]	Weakness: Sample not representative of general population; no control for confounding
Gasbarrini, 1995 ⁶⁴	San Marino	2237	General population sample aged 20-70+	History of ulcer in mother, OR=0.7 (0.5-1.1) History of ulcer in father, OR=1.2 (0.9-1.5) History of ulcer in sibling, OR=1.5 (1.1-2.1)	Strength: Subjects representative of general population; control for confounding by logistic regression

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Gilboa, 1995 ⁶⁵	Rural Israel	377	General population sample aged 30-90	No significant association between HP infection and family history of peptic ulcer	Strength: Subjects representative of general population Weakness: ORs not presented; no control for confounding

Table II-12. Studies assessing the relationship between *Helicobacter pylori* infection and institutionalized populations

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Kimura, 1999 ⁹⁸	Japan	196	Institutionalized neurologically impaired patients aged 3-64	Prevalence rate among institutionalized (81.1%) higher than healthy Japanese population or in home care patients with severe neurologic impairment (20%) [p,0.001]. Prevalence in inpatients increased with duration of institutionalization [p<0.01]. One year seroconversion rate=47.4%	Weakness: Prevalence rates not age-adjusted; no control for confounding
Bohmer, 1997 ⁹⁰	Netherlands	338	Institutionalized adults from 2 institutes for the intellectually disabled aged 11-89	Prevalence rate among institutionalized (82.8%) higher than Dutch population (approximately 50%) IQ<50, OR=1.9 (1.1-3.6) Rumination, OR=2.0, (1.1-3.6) Institutionalized longer than 15 years, OR=22 (3.9-128)	Weakness: Statistical tests used not specified; no apparent control for confounding
DeHert, 1997 ¹⁵⁶	Belgium	89 89	Institutionalized schizophrenic patients Volunteer blood donor controls	Prevalence of HP+ greater in patients (34.8%) than controls (14.6%), OR=3.0 (1.4-7.3) Prevalence in patients increased with length of stay in institution	Strength: Cases and controls matched for age and gender Weakness: Statistical tests used not specified; no apparent control for confounding; not clear if matched analysis was done Note – Letter to the Editor
Lewindon, 1997 ¹⁵⁷	Hong Kong	157 50	Institutionalized children aged 1-15 "Age-matched" out-patient controls	Prevalence of HP+ greater in patients (55.4%) than controls (8%) [p<0.0002]	Weakness: Mean age of in-patients 2.2 years older than mean age of out-patients; controls not representative of general population; statistical tests used not specified

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Breuer, 1996 ³³	Germany	260	Blood donors aged 18-61	Boarding school/orphanage during childhood, OR=2.6 (0.8-8.2)	Strength: All subjects approached participated Weakness: Subjects not representative of general population; no control for confounding; only 5% had not lived at home during childhood so power reduced
Malaty, 1996 ³⁴	Russia	307	1-19 year old children	Children from orphanages and communal apartments vs. those who have families, OR=2.1 (1.2-2.5)	Weakness: Subjects not representative of general population; control only for age;
Harris, 1995 ³¹	UK	424 267	Adults residents of a hospital for people with severe learning difficulties Age-sex matched controls	Prevalence of HP+ greater in residents of hospital (87%) than in controls (41%) No differences in prevalence of HP+ for residents by age or duration of stay	Weakness: Controls not representative of general population; no control for confounding; not clear how age and sex controlled for in analysis since numbers of patients and controls varied by agegroup
Lambert, 1995 ³²	Australia	122 273	Institutionalized medically and physically handicapped young adults Population-based age-sex matched controls	Prevalence of HP+ greater in residents of hospital (75%) than in controls (23%), OR=13.4 (6.5-29) [p<0.001] Prevalence of HP+ greater in these same residents in 1989 (75%) than in 1977 (34%) after adjustment for age, OR=2.4 (1.1-5.3) [p<0.03] Duration of stay was significantly related to HP status in 1977, but not 1989 The annual seroconversion rate was 7.4%	Strength: Controls representative of general population; ORs adjusted for confounding by logistic regression Weakness: Not clear how age and sex controlled for in comparison of HP prevalence

Table II-13. Studies assessing the relationship between *Helicobacter pylori* infection and familial exposures

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Brenner, 1999 ¹⁶⁵	Germany	110	Employees of a health insurance company Their partners	Prevalence of HP+ greater in employees with HP+ partner (42%) than HP- partner (7%), OR=7.0 (CI=1.8-26.7) Risk increased with number of years lived with HP+ partner	Strength: Studied healthy employees not patients; controlled for confounding by logistic regression Weakness: No DNA fingerprinting; can't rule out common source of infection Note -- UBT used
Rothenbacher, 1999 ¹⁶³	Germany	1143	General population sample of first grade children	Risks for child being HP+ according to parental HP status: Father HP+ , OR=3.8 (0.8-19.1) Mother HP+ , OR=7.9 (4.0-15.7)	Strength: Representative sample of children; controlled for confounding by logistic regression Weakness: Data only available on one parent Note -- UBT used
Singh, 1999 ¹⁶⁴	India	25 25	General population subjects Married spouses	Prevalence of HP+ greater in spouses of HP+ subjects (83.3%) than HP- subjects (28.5%) [$p<0.01$] 3 of 5 (60%) initially HP- spouses with HP+ spouses were HP+ at one year follow-up	Strength: Subjects selected from general population Weakness: No DNA fingerprinting; can't rule out common source of infection; no control for confounding Note -- biopsy, rapid urease test, histologic exam, and serology used
Brenner, 1998 ¹⁵⁵	Germany	863	Pre-school children out-patients	History of ulcer in mother, OR=11.7 (3.8-36.2) History of ulcer in father, OR=1.6 (0.4-5.5)	Strength: 79% participation rate; both variables controlled for confounding by logistic regression Weakness: mother's history only adjusted for age and sex; presence of ulcer used as surrogate for presence of HP infection
Chalkauskas, 1998 ¹⁷⁰	Lithuania	13 from 6 families	Dyspeptic patients and their relatives	Members of 2/6 families harbored same HP strain	Strength: Performed DNA fingerprinting Weakness: Can't rule out common source of infection

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Ma, 1998 ¹⁰³	China	49	General population sample Children 3-12	Risks for children being HP+ according to parental HP status - At least one parent HP+, OR=30.4 (4.0-232.2) Father HP+ , OR=10.1 (1.6-63.2) Mother HP+ , OR=30.1 (2.8-325.2) Risk for one spouse being HP+ when other spouse was HP+, OR=1.5 (0.6-3.7)	Strength: Representative sample of children; control for confounding by age and gender by logistic regression; bootstrap technique used to test differences between parents Weakness: small number of children; not total independence of children (7/41 had other sib participants); data not available on both mother and father for all 49 children; Note – UBT used for children
Bonamico, 1996 ¹⁵⁸	Italy	121 41	Relatives of HP+ children attending a pediatric clinic	Prevalence of HP+ in relatives of HP+ children (68%) was higher than in various control populations. The relative-specific prevalences were: mother (70%), father (67%), siblings (67%), other relative cohabitants (56%)	Weakness: No control group specific to study
Czkwianianc, 1996 ¹⁵⁰	Poland	240	Children aged 6 months-17 years who were patients at hospitals and clinics	Prevalence of HP+ higher in children who lived in households with HP seropositive relatives [p<0.005]	Weakness: Sample not representative of general population; no control for confounding; statistical tests not specified
Georgopoulos, 1996 ¹⁶⁶	Greece	64 64	Ulcer patients Spouses	Prevalence of HP+ greater in spouses of HP+ patients (78%) than HP- patients (20%), OR=14.0 (2.3-144.6) 8/18 couples colonized with identical HP strains	Strength: Chromosomal DNA from 18 couples analyzed Weakness: No control for confounding Note – biopsy, CLO, and culture used
Malaty, 1996 ¹⁶¹	Russia	307	1-19 year old children	Child has HP+ siblings, OR=2.4 (2.1-2.7)	Weakness: Subjects not representative of general population; control only for age
Mitchell, 1996 ¹⁶⁹	Australia	3	Children	Twin boys and younger sister found to be infected with identical strains of HP	Strength: Analyzed antigen recognition pattern

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Parente, 1996 ¹⁶⁷	Italy	124 124 248	HP+ patients with ulcer Spouses Volunteer blood donors	Prevalence of HP+ greater in spouses of HP+ patients (71%) than in matched controls (58%) [p<0.05]	Strength: Control for confounding by logistic regression; spouses matched to controls by sex, age, area, SES Weakness: Controls not representative of general population
Schutze, 1995 ¹⁶⁸	Austria	18	Patients with chronic ulcer in whom HP had been eradicated	HP reinfection occurred in 2/18 patients by a HP strain identical to that carried by the spouse	Strength: Performed DNA fingerprinting Weakness: Can't rule out common source of infection
Blecker, 1994 ¹⁶⁹	Belgium	35	Family members of HP+ children	Prevalence of HP+ greater in family members of HP+ children (48.6%) compared to general population group (17.8%)	Weakness: Poor participation rate of families; lack of independence among members of same family; comparability regarding age of control population not known
Malaty, 1994 ²⁴⁷	Sweden	269 twin pairs	Swedish Twin Registry adults, mean age=65.5	Variation in the acquisition of HP infection estimated to be 57% for genetic factors, 20% for shared environmental factors, and 23% for nonshared environmental factors	Strength: Intraclass correlations used to estimate influence of genetic and environmental effect in monozygotic and dizygotic twin pairs reared together and apart Weakness: Didn't assess specific risk factors
Barnford, 1993 ¹⁴⁶	Ireland	4 families	Parents of HP+ children	Members of 2 families harbored subtypes or strains of HP with the same DNA fingerprint	Strength: Performed DNA fingerprinting Weakness: Can't rule out common source of infection Note – biopsy and culture used
Mitchell, 1993 ¹⁶⁰	Austria	27 11 69	HP+ children HP- children Age-sex matched blood donors & children	Prevalence of HP+ greater in family members of HP+ children (76.7%) than HP- children (15%) and age-matched blood donor controls (21.7%) [p<0.001] Prevalence of HP+ greater in parents of HP+ children (81.1%) greater than in siblings (69.4%)	Strength: Compared HP+ prevalence in relatives of HP+ and HP- children as well as in age-sex matched controls Weakness: No control for other factors such as SES; lack of independence among members of same family

Author, Year, Ref No.	Country	No. in Study	Population	Outcome	Comments
Drum, 1990 ⁶¹	Canada	67 22 37	Parents Siblings Child controls	Prevalence of HP+ greater in parents of HP+ children (73.5%) than HP- children (24.2%) Prevalence of HP+ greater in mothers of HP+ children (83.3%) than fathers (62.5%) Prevalence of HP+ greater in siblings of HP+ children (81.8%) than HP- children (2%) or similarly aged controls (14%)	Strength: Control for children from same hospital that served the patient and their siblings Weakness: No control for factors such as SES related to HP status

Table IV-1. Distribution of cases and controls for selected variables by *Helicobacter pylori* serostatus

Factors	<u><i>H. pylori</i> +</u>		<u><i>H. pylori</i> -</u>		<u>Indeterminate</u>	
	N 1994	(col%) (row%)	N 1019	(col%) (row%)	N 275	(col%) (row%)
<u>Age in Years</u>						
<40	484	(24) (63)	219	(21.5) (28)	67	(24) (9)
40-44	530	(27) (62.5)	250	(24.5) (29)	68	(25) (8)
45-54	537	(27) (60)	290	(28) (32)	66	(24) (7)
55-59	170	(8.5) (57)	102	(10) (34)	27	(10) (9)
60+	273	(14) (57)	158	(15.5) (33)	47	(17) (10)
<u>Gender</u>						
Male	978	(49) (59)	536	(53) (32)	155	(56) (9)
Female	1016	(51) (63)	483	(47) (30)	120	(44) (7)
<u>Grouped Gastric Pathology Categories</u>						
Normal, SG	21	(1) (28)	47	(5) (64)	6	(2) (8)
CAG (III) mild	546	(27) (46)	523	(51) (45)	104	(38) (9)
Other CAG	188	(9) (77)	37	(4) (15)	19	(7) (8)
IM supperficial	183	(9) (67)	71	(7) (26)	21	(8) (8)
IM deep	656	(33) (71)	190	(19) (21)	78	(28) (8)

Factors	<u><i>H. pylori</i> +</u>		<u><i>H. pylori</i> -</u>		<u>Indeterminate</u>	
	N 1994	(col%) (row%)	N 1019	(col%) (row%)	N 275	(col%) (row%)
Dysplasia	298	(15) (68)	105	(10) (24)	34	(12) (8)
Cancer	4	(0.2) (80)	1	(0.1) (20)	0	---
Missing	98	(5) (63)	45	(4) (29)	13	(5) (8)
<u>Village name</u>						
Xi Quan	155	(8) (67)	52	(5) (22)	26	(9) (9)
Li Hu Zhuang	133	(7) (56)	81	(5) (34)	25	(9) (11)
Wang Jia Zhuang	183	(9) (76)	44	(4) (18)	13	(5) (5)
Xi Si Hou	61	(3) (56)	39	(4) (36)	9	(3) (8)
Li Jia Gou	215	(11) (67)	85	(8) (26)	22	(8) (7)
Suo Zhuang	246	(12) (64)	101	(10) (26)	40	(14.5) (10)
Huang Ai Quad	125	(6) (75)	32	(3) (19)	9	(3) (5)
Yang Jia He	182	(9) (71)	60	(6) (23)	14	(5) (5)
Xin Zhuang	140	(7) (50)	114	(11) (41)	25	(9) (9)
Guo Jia Zhuang	160	(8) (47)	152	(15) (44)	32	(12) (9)
Hou Jia He	214	(11) (55)	146	(14) (37)	31	(11) (8)

Factors	<i>H. pylori</i> +		<i>H. pylori</i> -		Indeterminate	
	N 1994	(col%) (row%)	N 1019	(col%) (row%)	N 275	(col%) (row%)
Hou He Ye	87	(4) (49)	71	(7) (40)	19	(7) (11)
Nan Yang He	93	(5) (64)	42	(4) (29)	10	(4) (7)

Table IV-2. Association between prevalence of *Helicobacter pylori* infection and cigarette smoking

Smoking Category	Positive	Negative	OR^{a,b}	95% CI	95% CI^c
<u>Smoked cigarettes</u>					
Never	1166	553	1.0	---	---
Ever	828	466	0.9	0.7-1.0	0.7-1.0
<u>Age first started smoking</u>					
<18	142	96	0.7	0.5-0.9	0.5-0.9
18-20	329	164	1.0	0.8-1.2	0.8-1.2
21-25	170	102	0.8	0.6-1.0	0.6-1.0
>25	187	104	0.9	0.7-1.2	0.7-1.2
<u>Number smoked per day</u>					
<10	169	93	0.9	0.7-1.2	0.7-1.1
10-19	174	92	0.9	0.7-1.2	0.7-1.2
20	338	205	0.8	0.6-1.0	0.6-1.0
>20	146	76	0.9	0.7-1.2	0.7-1.2
<u>Number of years smoked</u>					
<20	180	105	0.8	0.6-1.0	0.6-1.0
20-24	186	85	1.0	0.8-1.3	0.8-1.3
25-29	147	80	0.9	0.6-1.2	0.6-1.2
30-39	182	114	0.8	0.6-1.0	0.6-1.0
>39	133	82	0.9	0.6-1.2	0.6-1.2
<u>Lifetime packs of cigarettes smoked</u>					
<3595	164	95	0.8	0.6-1.1	0.7-1.0
3595-6862	176	79	1.1	0.8-1.4	0.8-1.4
6863-9490	181	98	0.9	0.7-1.1	0.7-1.1
9491-14235	152	100	0.7	0.6-1.0	0.6-1.0
>14235	154	94	0.8	0.6-1.1	0.6-1.1

Smoking Category	Positivee	Negative	OR^{a,b}	95% CI	95% CI^c
<u>Smoking status</u>					
Current	749	420	0.9	0.7-1.0	0.7-1.0
Past	79	46	0.8	0.6-1.2	0.6-1.3

^aThose who never smoked cigarettes were referent category for all OR calculated.

^bAdjusted for age in a polychotomous logistic model.

^cCalculated using bootstrap technique.

Table IV-3. Association between prevalence of *Helicobacter pylori* infection and alcoholic beverage consumption

Drinking Category	Positive	Negative	OR^{a,b}	95% CI	95% CI^c
<u>Drank alcohol</u>					
Never	1092	548	1.0	---	---
Ever	902	471	1.0	0.8-1.1	0.8-1.1
<u>Age first started drinking</u>					
<18	151	73	1.0	0.8-1.4	0.8-1.4
18-20	345	175	1.0	0.8-1.2	0.8-1.2
21-25	166	102	0.8	0.6-1.0	0.6-1.0
>25	240	121	1.0	0.8-1.3	0.8-1.3
<u>Number of times drank per week</u>					
<3	217	102	1.0	0.8-1.3	0.8-1.4
3-6	229	119	1.0	0.7-1.2	0.7-1.2
7-9	271	146	0.9	0.8-1.2	0.7-1.2
>9	183	104	0.9	0.7-1.2	0.7-1.2
<u>Number of years drank</u>					
<20	262	96	1.3	1.0-1.7	1.0-1.6
20-24	198	112	0.8	0.7-1.1	0.6-1.1
25-34	271	171	0.8	0.6-1.0	0.6-1.0
>34	171	92	1.0	0.8-1.4	0.8-1.4
<u>Drinking status</u>					
Current	843	446	0.9	0.8-1.1	0.8-1.1
Past	59	25	1.2	0.8-2.0	0.8-1.9

^aThose who never drank were referent category for all OR calculated.

^bAdjusted for age in a polychotomous logistic model.

^cCalculated using bootstrap technique.

Table IV-4. Association between prevalence of *Helicobacter pylori* infection and dietary variables

Dietary Variables	Positive	Negative	OR^a	95%CI	95%CI^b
<u>Jin^c of garlic eaten per year</u>					
<2	420	212	1.0	---	---
2	351	162	1.1	0.8-1.4	0.8-1.4
3-4	349	156	1.1	0.8-1.4	0.8-1.4
5	274	163	0.8	0.6-1.1	0.6-1.1
>5	321	175	0.9	0.7-1.2	0.7-1.2
<u>Jin^c of garlic stalk eaten per year</u>					
0	495	216	1.0	---	---
1	206	89	1.0	0.7-1.3	0.7-1.3
2	394	227	0.7	0.6-0.9	0.6-0.9
3-4	284	148	0.8	0.6-1.1	0.6-1.0
>4	336	188	0.8	0.6-1.0	0.6-1.0
<u>Jin^c of scallions eaten per year</u>					
<3	352	130	1.0	---	---
3-4	267	135	0.7	0.5-1.0	0.6-0.9
5-7	433	256	0.6	0.5-0.8	0.5-0.8
8-10	411	227	0.7	0.5-0.9	0.5-0.9
>10	252	120	0.8	0.6-1.0	0.6-1.1
<u>Jin^c of chives eaten per year</u>					
<3	353	178	1.0	---	---
3-4	199	101	1.0	0.7-1.3	0.7-1.3
5-7	442	249	0.9	0.7-1.1	0.7-1.1
8-10	460	225	1.0	0.8-1.3	0.8-1.2
>10	261	115	1.1	0.8-1.5	0.8-1.5

Dietary Variables	Positive	Negative	OR^a	95%CI	95%CI^b
<u>Jin^c of allium vegetables eaten per year</u>					
<11	354	161	1.0	---	---
11-15	342	172	0.9	0.7-1.1	0.7-1.1
16-21	339	173	0.9	0.7-1.1	0.6-1.2
22-30	361	195	0.8	0.6-1.1	0.6-1.0
>30	319	167	0.8	0.6-1.1	0.7-1.1
<u>Number of times per month ate raw fruits and vegetables</u>					
Never ate	103	54	1.0	---	---
Ever ate	1891	964	1.0	0.7-1.4	0.7-1.4
<3	296	150	1.0	0.7-1.5	0.7-1.5
3-5.9	391	222	0.9	0.6-1.3	0.6-1.3
6-9.9	408	219	0.9	0.6-1.3	0.6-1.4
10-15.9	438	201	1.1	0.7-1.6	0.7-1.6
≥16	358	172	1.0	0.7-1.5	0.7-1.5
<u>Frequency washed their raw fruits and vegetables</u>					
Always	1179	553	1.0	---	---
> half the time	386	215	0.8	0.7-1.0	0.7-1.0
< half the time	249	141	0.8	0.6-1.0	0.6-1.0
Never	77	56	0.7	0.5-1.0	0.5-0.9
<u>Frequency peeled their raw fruits and vegetables</u>					
Always	314	174	1.0	---	---
> half the time	252	105	1.3	1.0-1.7	0.9-1.8
< half the time	309	162	1.0	0.8-1.3	0.7-1.4
Never	1016	524	1.0	0.8-1.3	0.8-1.3
<u>Lian^d of sour pancakes eaten per year</u>					
Never ate	1493	791	1.0	---	---
Ever ate	501	228	1.1	1.0-1.4	0.9-1.4
<271	166	79	1.1	0.8-1.5	0.8-1.5
271-900	166	75	1.1	0.9-1.5	0.8-1.6
>900	168	73	1.2	0.9-1.6	0.8-1.7

Dietary Variables	Positive	Negative	OR^a	95%CI	95%CI^b
<u>Lian^d of sweet pancakes eaten per year</u>					
Never ate	1005	464	1.0	---	---
Ever ate	989	555	0.8	0.7-0.9	0.7-0.9
<225	337	174	0.9	0.7-1.1	0.7-1.1
225-525	322	194	0.7	0.6-0.9	0.6-0.9
>525	330	186	0.8	0.6-1.0	0.6-1.0
<u>Preferred saltiness of food</u>					
Lightly salty	419	239	1.0	---	---
Moderately salty	415	205	1.1	0.9-1.4	0.9-1.5
Very salty	1160	575	1.1	0.9-1.3	0.9-1.4

^aAdjusted for age in a polychotomous logistic model.

^bCalculated using bootstrap technique.

^c1 jin ~ 500 grams.

^d1 lian ~ 50 grams.

Table IV-5. Association between prevalence of *Helicobacter pylori* infection and occupation variables

Occupation^a	Positive	Negative	OR^{b,c}	95% CI	95% CI^d
Administrative	112	56	1.0	0.7-1.4	0.7-1.3
Technical/clerical/sales	16	10	0.8	0.4-1.7	0.3-2.0
Precision production	62	22	1.4	0.8-2.2	0.8-2.2
Operator/fabricator/laborer	28	21	0.7	0.4-1.2	0.3-1.2
Service occupations	427	201	1.2	1.0-1.4	0.9-1.4
Farming	1505	764	1.0	0.8-1.2	0.8-1.1
Animal worker	9	11	0.5	0.2-1.1	0.2-1.2

^aThe number of subjects obtained by summing all occupations exceeds the total number of subjects in the study since some subjects reported having more than one occupation.

^bThose not employed in each occupation are the referent group for each occupation.

^cAdjusted for age in a polychotomous logistic model.

^dCalculated using bootstrap technique.

Table IV-6. Association between prevalence of *Helicobacter pylori* infection and water-related exposures

Water Variables	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Source of drinking water</u>					
Deep private well	235	163	1.0	---	---
Shallow private well	1004	543	1.3	1.0-1.6	1.0-1.7
Deep village well	70	35	1.4	0.9-2.2	0.8-2.3
Shallow village well	648	250	1.8	1.4-2.3	1.4-2.3
Pond/river/ditch	25	22	0.8	0.4-1.5	0.4-1.5
Running/spring water	12	6	1.4	0.5-3.8	0.6-3.4
<u>Number of days store drinking water in a jar</u>					
None	496	225	1.0	---	---
Some	1498	794	0.9	0.7-1.0	0.7-1.0
<3	538	271	0.9	0.7-1.1	0.7-1.1
3	338	186	0.8	0.7-1.1	0.7-1.1
>3	621	337	0.8	0.7-1.0	0.7-1.0
<u>Frequency that water is boiled</u>					
Always	1372	704	1.0	---	---
Often	511	255	1.0	0.8-1.2	0.8-1.2
Sometimes	77	33	1.1	0.8-1.7	0.7-1.8
Never	33	26	0.7	0.4-1.1	0.4-1.1
<u>Location washed diapers</u>					
Never washed diapers	634	360	1.0	---	---
Washed in house	574	294	1.1	0.9-1.3	0.9-1.3
Washed in river/stream	941	430	1.2	1.0-1.5	1.0-1.4
Washed in pond/ditch	12	3	2.2	0.6-7.8	NA ^c

Water Variables	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Location washed/bathed when weather was warm</u>					
Never washed	218	146	1.0	---	---
Ever washed	1776	873	1.3	1.0-1.6	1.0-1.6
Washed at home	616	298	1.3	1.0-1.7	1.0-1.7
Washed in court yard	572	292	1.2	1.0-1.6	1.0-1.5
Washed in stream	521	270	1.2	1.0-1.6	1.0-1.6
Washed in pond/ditch	108	43	1.6	1.0-2.4	1.0-2.4
Washed in public bath	18	4	2.8	0.9-8.4	NA ^c
<u>Number of times per month washed/bathed when weather was warm</u>					
Never washed	218	146	1.0	---	---
1-4	511	261	1.3	1.0-1.6	1.0-1.7
4.1-8.0	391	180	1.4	1.0-1.8	1.0-1.8
8.1-25.9	479	226	1.3	1.0-1.8	1.0-1.8
≥26	395	206	1.2	0.9-1.6	0.9-1.7
<u>Washed/ bathed when weather was cold</u>					
No	1767	919	1.0	---	---
Yes	227	100	1.1	0.9-1.5	0.9-1.5
<u>Number of times per year swim in river/reservoir</u>					
None	130	352	1.0	---	---
Some	135	267	1.0	0.7-1.4	0.7-1.4
1-4	61	34	0.9	0.6-1.3	0.6-1.3
>4	62	25	1.2	0.8-1.9	0.7-2.0

^aAdjusted for age in a polychotomous logistic model.

^bCalculated using bootstrap technique.

^cNot available - does not meet asymptotic assumption for bootstrap technique.

Table IV-7. Association between prevalence of *Helicobacter pylori* infection and animals and pets

Animal Variables	Positive	Negative	OR^a	95%CI	95%CI^b
<u>Number of animals kept in the house as an adult</u>					
No animals	1897	947	1.0	---	---
Some animals	96	72	0.7	0.5-0.9	0.5-1.0
1	38	37	0.5	0.3-0.8	0.3-0.9
2-4	22	13	0.9	0.4-1.7	0.4-1.7
>4	33	19	0.9	0.5-1.6	0.5-1.7
<u>Types of animals kept in the house as an adult</u>					
No animals	1897	947	1.0	---	---
Cats	33	27	0.6	0.4-1.0	0.3-1.1
Dogs	7	3	1.2	0.3-4.5	NA ^c
Rabbits	47	34	0.7	0.4-1.1	0.4-1.1
<u>Types of animals kept in the courtyard as an adult</u>					
No animals	62	27	1.0	---	---
Some animals	1932	992	0.8	0.5-1.3	0.5-1.2
Cats	26	15	0.7	0.3-1.5	0.3-1.6
Chickens	1617	856	0.8	0.5-1.2	0.4-1.4
Cows	307	189	0.6	0.4-1.1	0.4-1.1
Dogs	365	164	0.9	0.6-1.5	0.6-1.5
Donkeys	68	40	0.7	0.4-1.3	0.4-1.3
Ducks	956	502	0.8	0.5-1.2	0.5-1.3
Geese	253	145	0.7	0.4-1.2	0.4-1.2
Goats	488	242	0.8	0.5-1.4	0.5-1.4
Pigs	1469	759	0.8	0.5-1.2	0.5-1.2
Rabbits	346	180	0.8	0.5-1.3	0.5-1.4
Sheep	2	3	0.3	0.04-1.7	NA ^c

Animal Variables	Positive	Negative	OR^a	95%CI	95%CI^b
<u>Number of animals kept in the house at age 10</u>					
No animals	1677	853	1.0	---	---
Some animals	312	159	1.0	0.8-1.2	0.8-1.2
1	71	49	0.7	0.5-1.0	0.5-1.1
2-4	97	43	1.1	0.8-1.6	0.7-1.6
>4	131	56	1.1	0.8-1.6	0.8-1.5
<u>Types of animals kept in the house at age 10</u>					
No animals	1677	853	1.0	---	---
Cats	70	52	0.7	0.5-1.0	0.5-1.0
Dogs	18	17	0.5	0.3-1.0	0.3-1.0
Rabbits	76	59	1.1	0.9-1.5	0.9-1.4
<u>Had job working with animals</u>					
No	1981	1006	1.0	---	---
Yes	9	11	0.5	0.2-1.1	0.2-1.2

^aAdjusted for age in a polychotomous logistic model.

^bCalculated using bootstrap technique.

^cNot available - does not meet asymptotic assumption for bootstrap technique.

Table IV-8. Association between prevalence of *Helicobacter pylori* infection and hygiene factors

Hygiene factors	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Number of times per month spit onto the ground</u>					
Never spit	695	331	1.0	---	---
Ever spit	1298	687	0.9	0.8-1.1	0.8-1.0
<30	278	152	0.9	0.7-1.1	0.7-1.1
30-89	289	158	0.9	0.7-1.1	0.7-1.1
90-180	463	233	0.9	0.8-1.2	0.8-1.2
>180	268	144	0.9	0.7-1.1	0.7-1.1
<u>Number of times per month brushed teeth</u>					
Never brushed	1197	628	1.0	---	---
Ever brushed	797	390	1.0	0.9-1.2	0.8-1.3
<6	229	110	1.0	0.8-1.3	0.8-1.3
6-15	302	163	0.9	0.7-1.1	0.7-1.1
>15	266	117	1.1	0.9-1.4	0.9-1.5
<u>Frequency of hand washing before eating meals</u>					
Always	1689	867	1.0	---	---
> half the time	237	130	1.0	0.8-1.2	0.7-1.2
< half the time	61	21	1.6	1.0-2.6	1.0-2.5
Never	7	1	3.8	0.5-31.0	NA ^c
<u>Frequency of hand washing after bowel movements</u>					
Always	961	427	1.0	---	---
> half the time	493	294	0.7	0.6-0.9	0.6-0.9
< half the time	333	187	0.8	0.7-1.0	0.7-1.0
Never	207	111	0.9	0.7-1.1	0.7-1.1
<u>Frequency of hand washing before preparing meals</u>					
Always	1299	645	1.0	---	---
> half the time	88	48	0.9	0.7-1.3	0.6-1.4
< half the time	30	13	1.2	0.6-2.3	0.6-2.2
Never	6	5	0.6	0.2-2.0	0.2-2.2

Hvgiene factors	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Frequency of hand washing with soap</u>					
Always	706	338	1.0	---	---
> half the time	518	270	0.9	0.8-1.1	0.8-1.1
< half the time	591	315	0.9	0.8-1.1	0.8-1.1
Never	179	96	0.9	0.7-1.2	0.7-1.2
<u>Frequency of body washing with soap</u>					
Always	814	387	1.0	---	---
> half the time	265	142	0.9	0.7-1.1	0.7-1.1
< half the time	270	122	1.1	0.8-1.4	0.8-1.4
Never	426	221	1.0	0.8-1.2	0.8-1.2
<u>Frequency of washing chopsticks and bowls</u>					
Always	1928	982	1.0	---	---
> half the time	59	32	0.9	0.6-1.5	0.6-1.4
< than half the time	5	5	0.5	0.2-1.9	NA ^c
Never	2	0	---	---	---
<u>Frequency of washing chopsticks and bowls with detergent</u>					
Always	29	12	1.0	---	---
> half the time	25	10	1.0	0.4-2.8	0.4-3.0
< half the time	253	127	0.8	0.4-1.7	0.4-1.7
Never	1686	868	0.8	0.4-1.6	0.4-1.6
<u>Temperature of water used to wash chopsticks and bowls</u>					
Hot	13	7	1.0	---	---
Cold	1861	963	1.3	0.5-3.4	0.4-4.2
Both	120	49	1.0	0.4-2.5	0.3-3.2
<u>Frequency of cup washing after use</u>					
Always	1800	901	1.0	---	---
> half the time	109	78	0.7	0.5-0.9	0.5-0.9
< half the time	66	29	1.1	0.7-1.8	0.7-1.9
Never	19	11	0.9	0.4-1.9	0.4-2.0

Hvgiene factors	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Frequency of sharing cups with family</u>					
Always	1847	915	1.0	---	---
> half the time	31	19	0.9	0.5-1.5	0.4-1.8
< half the time	24	23	0.6	0.3-1.0	0.3-1.1
Never	92	62	0.8	0.6-1.1	0.6-1.1
<u>Frequency of pre-chewing child's food</u>					
Always	853	413	1.0	---	---
Sometimes	251	110	1.1	0.8-1.4	0.8-1.4
Never	890	496	0.9	0.7-1.0	0.7-1.0
<u>Frequency of kissing young (< age 10) children on the lips</u>					
Always	381	172	1.0	---	---
Sometimes	625	327	0.9	0.7-1.1	0.7-1.1
Never	988	520	0.9	0.7-1.1	0.7-1.1
<u>Frequency of parents pre-chewing your food as a child</u>					
Always	1588	803	1.0	---	---
Sometimes	61	29	1.1	0.7-1.7	0.7-1.7
Never	89	56	0.8	0.6-1.1	0.6-1.1
<u>Frequency of parents kissing you on the lips as a child</u>					
Always	388	187	1.0	---	---
Sometimes	407	198	1.0	0.8-1.3	0.8-1.2
Never	458	251	0.9	0.7-1.1	0.7-1.1
<u>Type of flooring in house at age 10</u>					
Ground	1979	1009	1.0	---	---
Brick	11	8	0.7	0.3-1.9	0.3-2.2
Cement	2	2	0.5	0.1-3.8	NA ^c
Wood	1	0	~	---	---

^aAdjusted for age in a polychotomous logistic model.

^bCalculated using bootstrap technique.

^cNot available - does not meet asymptotic assumption for bootstrap technique.

Table IV-9. Association between prevalence of *Helicobacter pylori* infection and density/crowding factors

Hvgiene factors	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Number of people in household as an adult</u>					
<3	407	228	1.0	---	---
3	269	132	1.0	0.8-1.4	0.7-1.4
4	662	342	0.9	0.7-1.2	0.6-1.2
5	454	223	1.0	0.7-1.3	0.7-1.3
>5	201	94	1.0	0.7-1.4	0.7-1.5
<u>Number of children in household as an adult</u>					
0	1458	767	1.0	---	---
1	402	196	1.0	0.9-1.3	0.8-1.3
>1	55	14	2.0	1.1-3.6	1.0-3.7
<u>Number of people shared a bed with as an adult</u>					
0	283	166	1.0	---	---
1	1309	640	1.1	0.9-1.4	0.9-1.4
2	254	150	0.9	0.7-1.2	0.6-1.2
>2	67	17	2.1	1.2-3.7	1.1-3.8
<u>Number of children shared a bed with as an adult</u>					
Never shared a bed	283	166	1.0	---	---
Shared with child	429	232	1.0	0.8-1.3	0.7-1.3
1	337	199	0.9	0.7-1.2	0.7-1.2
>1	82	25	1.7	1.0-2.8	1.0-2.8
Shared bed with adult	1203	579	1.2	0.9-1.4	0.9-1.4
<u>Shared a bed with spouse as an adult</u>					
Never shared a bed	283	166	1.0	---	---
Yes	1513	741	1.1	0.9-1.4	0.9-1.4
No	119	70	0.9	0.7-1.3	0.7-1.3

Hygiene factors	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Number of people in household when 10 years old</u>					
<5	339	153	1.0	---	---
5	377	197	0.8	0.7-1.1	0.6-1.1
6	460	237	0.8	0.6-1.1	0.6-1.1
7	363	207	0.7	0.6-1.0	0.6-1.0
>7	452	225	0.8	0.7-1.1	0.7-1.1
<u>Number of siblings less than 5 years old in household when subject was 10</u>					
No siblings <5	739	366	1.0	---	---
Yes siblings <5	1150	599	0.9	0.8-1.1	0.8-1.1
1	672	372	0.9	0.7-1.1	0.7-1.1
2	426	208	1.0	0.8-1.2	0.8-1.2
>2	44	15	1.4	0.8-2.5	0.8-2.6
No siblings	88	41	1.1	0.8-1.7	0.7-1.7
<u>Number of siblings less than 10 years old in household when subject was 10</u>					
No siblings <10	372	187	1.0	---	---
Yes siblings <10	1522	783	1.0	0.8-1.2	0.8-1.2
1	514	267	1.0	0.8-1.2	0.7-1.1
2	546	309	0.9	0.7-1.1	0.8-1.2
>2	449	201	1.1	0.8-1.4	0.8-1.4
No siblings	88	41	1.1	0.8-1.7	0.8-1.7
<u>Number of people shared a bed with at age 10</u>					
None	106	46	1.0	---	---
1	516	228	1.0	0.7-1.4	0.6-1.5
2-3	663	358	0.8	0.6-1.2	0.5-1.2
4	296	164	0.8	0.5-1.2	0.5-1.2
>4	406	218	0.8	0.6-1.2	0.5-1.2

Hygiene factors	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Number of people shared a bed with at age 5</u>					
<2	314	128	1.0	---	---
2	437	227	0.8	0.6-1.0	0.6-1.0
3	506	281	0.7	0.6-0.9	0.5-1.0
4	341	186	0.7	0.6-1.0	0.6-1.0
>4	249	127	0.8	0.6-1.1	0.6-1.1

^aAdjusted for age in a polychotomous logistic model.

^bCalculated using bootstrap technique.

Table IV-10. Association between prevalence of *Helicobacter pylori* infection and social factors

Social factors	Positive	Negative	ORa	95% CI	95% CI^b
<u>Marital status</u>					
Married	1831	921	1.0	---	---
Widowed	146	78	1.1	0.8-1.4	0.8-1.5
Divorced or	2	3	0.4	0.06-2.2	NA ^c
Never married	15	17	0.4	0.2-0.9	0.2-0.9
<u>Number of years of subject's education</u>					
<1	693	349	1.0	---	---
1-3	358	193	0.9	0.7-1.1	0.7-1.2
4-6	522	254	1.0	0.8-1.2	0.8-1.2
7-9	356	196	0.8	0.6-1.0	0.6-1.1
>9	65	27	1.1	0.7-1.8	0.7-1.8
<u>Number of years of spouse's education</u>					
<1	584	324	1.0	---	---
1-3	323	168	1.0	0.8-1.3	0.8-1.3
4-6	484	219	1.2	0.9-1.4	0.9-1.4
7-9	389	175	1.1	0.9-1.4	0.9-1.4
>9	45	32	0.7	0.4-1.1	0.4-1.1
<u>Village education level</u>					
High	448	376	1.0	---	---
Medium	768	374	1.7	1.4-2.1	1.4-2.1
Low	778	269	2.4	2.0-2.9	2.0-3.0

Social factors	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Subject's annual income in Yuan^d</u>					
<2,000	330	183	1.0	---	---
2,000-3,999	555	264	1.1	0.8-1.3	0.8-1.3
4,000-5,999	421	225	0.9	0.7-1.2	0.7-1.2
6,000-7,999	269	156	0.8	0.6-1.1	0.6-1.1
>8,000	412	189	1.0	0.8-1.4	0.8-1.4

^aAdjusted for age in a polychotomous logistic model.

^bCalculated using bootstrap technique.

^cNot available using bootstrap technique.

^d10 yuan ~ \$1.25 U.S.

Table IV-11. Association between prevalence of *Helicobacter pylori* infection and gastric factors

Gastric factors	Positive	Negative	OR^a	95% CI	95% CI^b
<u>Number of times per month have gastric reflux</u>					
Never	1089	555	1.0	---	---
Ever	901	464	1.0	0.8-1.1	0.8-1.1
<0.43	300	156	1.0	0.8-1.2	0.8-1.2
0.43-3.9	357	165	1.1	0.9-1.3	0.9-1.3
4.0-8.9	117	63	0.9	0.7-1.3	0.7-1.4
>9	127	80	0.8	0.6-1.1	0.6-1.1

^aAdjusted for age in a polychotomous logistic model.

^bCalculated using bootstrap technique.

Table IV-12. Prevalence rate, prevalence rate ratio, and relative odds ratio of *Helicobacter pylori* infection for combinations of models according to age, percent of time washed hands before eating, number of children in the household, and village education level

Age	% Wash Hands	# Children in House	Village Education	Prevalence Rate	Prevalence Rate Ratio ^a	Relative Odds Ratio ^b
40	<50%	2+	low	74 (59-86)	1.4 (1.1-1.7)	8.6 (3.9-22.2)
40	>50%	2+	low	75 (64-84)	1.5 (1.2-1.7)	5.0 (2.7-10.7)
40	<50%	0-1	low	74 (66-83)	1.5 (1.3-1.7)	4.1 (2.5-7.8)
40	>50%	0-1	low	70 (67-72)	1.4 (1.3-1.5)	2.4 (2.0-3.0)
40	<50%	2+	med	70 (54-83)	1.4 (1.1-1.7)	6.0 (2.7-16.2)
40	>50%	2+	med	69 (58-79)	1.4 (1.1-1.6)	3.5 (1.9-7.4)
40	<50%	0-1	med	69 (59-78)	1.4 (1.2-1.6)	2.8 (1.7-5.2)
40	>50%	0-1	med	63 (60-66)	1.2 (1.1-1.3)	1.7 (1.4-2.0)
40	<50%	2+	high	61 (43-77)	1.2 (0.9-1.5)	3.6 (1.6-9.3)
40	>50%	2+	high	60 (47-72)	1.2 (0.9-1.4)	2.1 (1.2-4.4)
40	<50%	0-1	high	59 (48-70)	1.2 (0.9-1.4)	1.7 (1.1-3.1)
40	>50%	0-1	high	51 (47-55)	1.0	1.0
50	<50%	2+	low	72 (57-85)	1.5 (1.1-1.8)	8.6 (3.9-22.2)
50	>50%	2+	low	73 (62-82)	1.5 (1.3-1.7)	5.0 (2.7-10.7)
50	<50%	0-1	low	73 (64-81)	1.5 (1.3-1.7)	4.1 (2.5-7.8)
50	>50%	0-1	low	68 (65-70)	1.4 (1.3-1.5)	2.4 (2.0-3.0)
50	<50%	2+	med	68 (52-82)	1.4 (1.1-1.7)	6.0 (2.7-16.2)
50	>50%	2+	med	68 (56-78)	1.4 (1.1-1.6)	3.5 (1.9-7.4)
50	<50%	0-1	med	67 (57-76)	1.4 (1.2-1.6)	2.8 (1.7-5.2)
50	>50%	0-1	med	60 (57-63)	1.2 (1.1-1.4)	1.7 (1.4-2.0)
50	<50%	2+	high	59 (41-75)	1.2 (0.9-1.5)	3.6 (1.6-9.3)
50	>50%	2+	high	58 (45-70)	1.2 (0.9-1.4)	2.1 (1.2-4.4)
50	<50%	0-1	high	56 (46-68)	1.2 (1.0-1.4)	1.7 (1.1-3.1)
50	>50%	0-1	high	48 (45-52)	1.0	1.0

Age	% Wash Hands	#Children in House	Village Education	Prevalence Rate	Prevalence Rate Ratio^a	Relative Odds Ratio^b
60	<50%	2+	low	71 (54-84)	1.5 (1.2-1.9)	8.6 (3.9-22.2)
60	>50%	2+	low	71 (59-81)	1.5 (1.3-1.8)	5.0 (2.7-10.7)
60	<50%	0-1	low	71 (62-80)	1.5 (1.3-1.8)	4.1 (2.5-7.8)
60	>50%	0-1	low	66 (61-69)	1.4 (1.3-1.6)	2.4 (2.0-3.0)
60	<50%	2+	med	66 (50-81)	1.4 (1.1-1.8)	6.0 (2.7-16.2)
60	>50%	2+	med	66 (53-76)	1.4 (1.1-1.7)	3.5 (1.9-7.4)
60	<50%	0-1	med	65 (55-74)	1.4 (1.2-1.7)	2.8 (1.7-5.2)
60	>50%	0-1	med	58 (54-62)	1.3 (1.2-1.4)	1.7 (1.4-2.0)
60	<50%	2+	high	57 (39-74)	1.2 (0.9-1.6)	3.6 (1.6-9.3)
60	>50%	2+	high	55 (42-68)	1.2 (0.9-1.5)	2.1 (1.2-4.4)
60	<50%	0-1	high	54 (43-66)	1.2 (1.0-1.4)	1.7 (1.1-3.1)
60	>50%	0-1	high	46 (42-50)	1.0	1.0

^a Subjects of the same age who washed hands before eating >50% of the time, who had <2 children in their household, and who lived in a village with a high education level were the referent category for all PRR calculated.

^b Subjects of the same age who washed hands before eating >50% of the time, who had <2 children in their household, and who lived in a village with a high education level were the referent category for all ROR calculated.

Table IV-13. Prevalence rate, prevalence rate ratio, and relative odds ratio of *Helicobacter pylori* infection for combinations of models according to age, percent of time washed hands before eating, number of children in the household, and source of drinking water

Age	% Wash Hands	# Children in House	Type of Well	Prevalence Rate	Prevalence Rate Ratio ^a	Relative Odds Ratio ^b
40	<50%	2+	shallow village	70 (52-84)	1.3 (0.9-1.6)	7.0 (3.4-17.6)
40	>50%	2+	shallow village	72 (61-82)	1.3 (1.1-1.6)	3.9 (2.2-8.3)
40	<50%	0-1	shallow village	73 (63-81)	1.3 (1.1-1.6)	3.3 (1.9-6.2)
40	>50%	0-1	shallow village	68 (64-71)	1.2 (1.1-1.4)	1.8 (1.4-2.4)
40	<50%	2+	shallow private	69 (52-83)	1.2 (0.9-1.6)	4.9 (2.3-12.3)
40	>50%	2+	shallow private	68 (58-78)	1.2 (1.0-1.5)	2.7 (1.6-5.7)
40	<50%	0-1	shallow private	69 (59-78)	1.2 (1.1-1.5)	2.3 (1.3-4.3)
40	>50%	0-1	shallow private	61 (58-64)	1.1 (1.0-1.2)	1.3 (1.0-1.6)
40	<50%	2+	deep private	61 (42-78)	1.1 (0.8-1.4)	3.9 (1.9-9.4)
40	>50%	2+	deep private	61 (49-74)	1.1 (0.9-1.3)	2.2 (1.2-4.5)
40	<50%	0-1	deep private	62 (50-74)	1.1 (0.9-1.3)	1.8 (1.1-3.2)
40	>50%	0-1	deep private	55 (49-60)	1.0	1.0
50	<50%	2+	shallow village	68 (50-83)	1.3 (0.9-1.7)	7.0 (3.4-17.6)
50	>50%	2+	shallow village	70 (58-80)	1.3 (1.1-1.6)	3.9 (2.2-8.3)
50	<50%	0-1	shallow village	71 (61-80)	1.4 (1.1-1.6)	3.3 (1.9-6.2)
50	>50%	0-1	shallow village	65 (62-68)	1.3 (1.1-1.4)	1.8 (1.4-2.4)
50	<50%	2+	shallow private	66 (50-81)	1.3 (0.9-1.6)	4.9 (2.3-12.3)
50	>50%	2+	shallow private	66 (55-76)	1.3 (1.0-1.5)	2.7 (1.6-5.7)
50	<50%	0-1	shallow private	66 (57-76)	1.3 (1.1-1.5)	2.3 (1.3-4.3)
50	>50%	0-1	shallow private	58 (56-61)	1.1 (1.0-1.3)	1.3 (1.0-1.6)
50	<50%	2+	deep private	58 (39-76)	1.1 (0.7-1.5)	3.9 (1.9-9.4)
50	>50%	2+	deep private	59 (46-7)	1.1 (0.9-1.4)	2.2 (1.2-4.5)
50	<50%	0-1	deep private	59 (48-71)	1.1 (0.9-1.4)	1.8 (1.1-3.2)
50	>50%	0-1	deep private	52 (47-57)	1.0	1.0

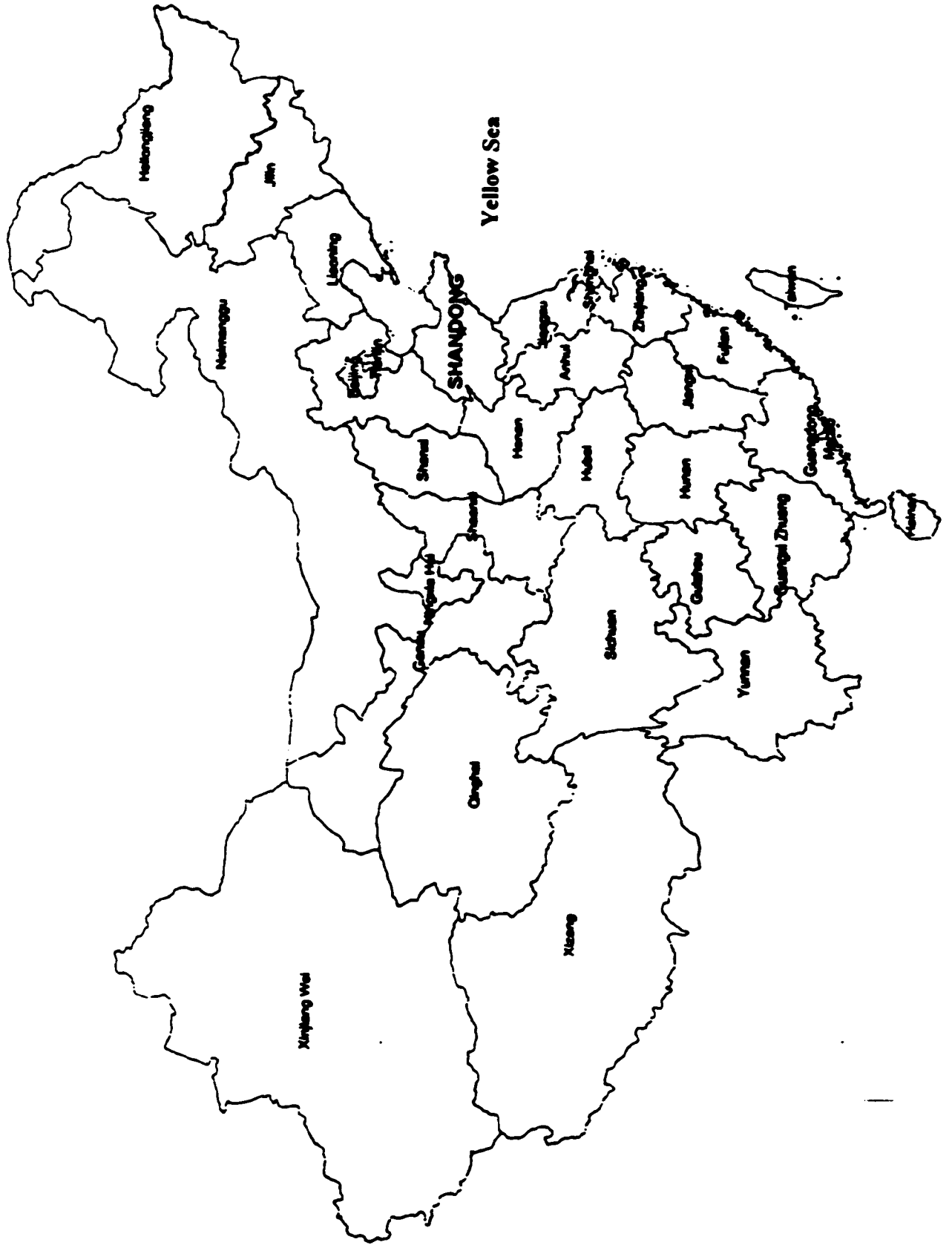
Age	%Wash Hands	#Children in House	Type of Well	Prevalence Rate	Prevalence Rate Ratio^a	Relative Odds Ratio^b
60	<50%	2+	shallow village	66 (47-82)	1.3 (0.9-1.7)	7.0 (3.4-17.6)
60	>50%	2+	shallow village	68 (54-78)	1.4 (1.1-1.7)	3.9 (2.2-8.3)
60	<50%	0-1	shallow village	68 (58-79)	1.4 (1.2-1.7)	3.3 (1.9-6.2)
60	>50%	0-1	shallow village	63 (59-67)	1.3 (1.1-1.4)	1.8 (1.4-2.4)
60	<50%	2+	shallow private	64 (46-79)	1.3 (0.9-1.7)	4.9 (2.3-12.3)
60	>50%	2+	shallow private	64 (52-75)	1.3 (1.0-1.6)	2.7 (1.6-5.7)
60	<50%	0-1	shallow private	64 (53-74)	1.3 (1.1-1.6)	2.3 (1.3-4.3)
60	>50%	0-1	shallow private	56 (52-59)	1.1 (1.0-1.3)	1.3 (1.0-1.6)
60	<50%	2+	deep private	56 (36-74)	1.1 (0.8-1.5)	3.9 (1.9-9.4)
60	>50%	2+	deep private	56 (42-70)	1.1 (0.9-1.4)	2.2 (1.2-4.5)
60	<50%	0-1	deep private	57 (45-68)	1.2 (0.9-1.4)	1.8 (1.1-3.2)
60	>50%	0-1	deep private	49 (43-55)	1.0	1.0

^aSubjects of the same age who washed hands before eating >50% of the time, who had <2 children in their household, and whose drinking water came from a deep private well are the referent category for all PRRs.

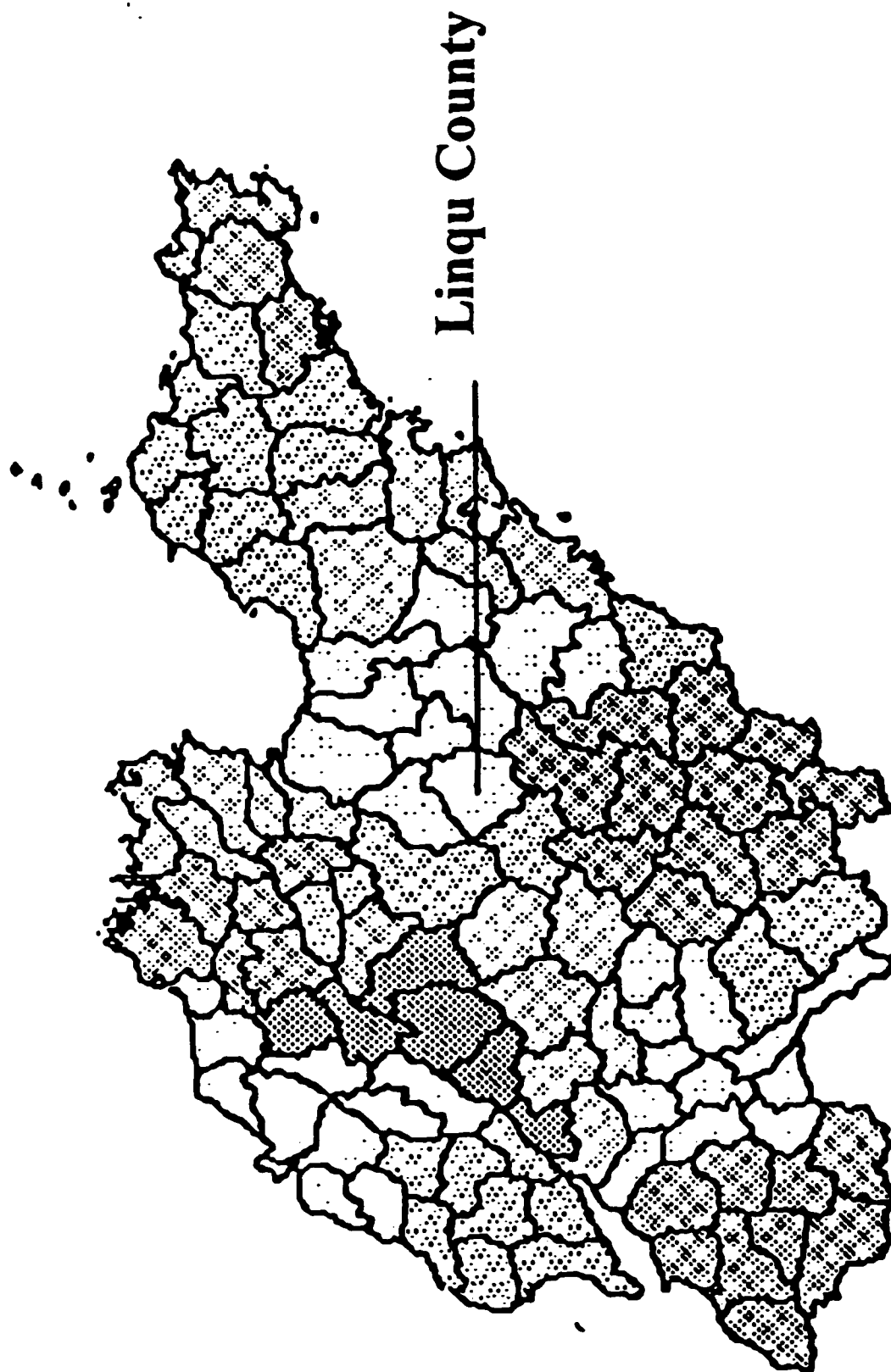
^bSubjects of the same age who washed hands before eating >50% of the time, who had <2 children in their household, and whose drinking water came from a deep private well are the referent category for all RORs.

APPENDIX 1

CHINA



SHANDONG PROVINCE



APPENDIX 2

SHANDONG INTERVENTION STUDY QUESTIONNAIRE

Village_____

Study Subject ID_____

Sex: 1 male 2 female

Date of Birth _____
YR MO DADate of Interview _____
YR MO DA

Time Start Interview __:__

Time End Interview __:__

Total Time __ minutes

Interviewer ID _____

INTRODUCTION:

Thank you for agreeing to participate in this study. During this interview, I will be asking questions about your household composition now and during childhood, your washing, smoking, drinking, and eating habits, and general background information about you and your family. Your cooperation is very important to this research study and will help us learn more about the health of adults in Linqu. I would like to remind you that your participation is voluntary and that the information you provide will be kept confidential and will be used only for research purposes.

Section 1. Background

First, I'd like to ask a few background questions.

1.1 Are you currently (READ):

Married.....1
 Widowed.....2 (skip to 1.4)
 Divorced or separated, or..3 (skip to 1.4)
 Never married.....4 (skip to 1.4)

1.2 How many years of education does your spouse have?

_____ Years
 None.... 0

1.3 What is your spouse's current occupation?

Grain Farmer.....1
 Fruit Farmer.....2
 Vegetable Farmer.....3
 Village Leader.....4
 Not working.....5
 Other, Specify _____

1.4 How many years of education do you have?

_____ Years
 None.... 0

1.5 What is your current occupation?

Grain Farmer.....1
 Fruit Farmer.....2
 Vegetable Farmer.....3
 Village Leader.....4
 Not working.....5
 Other, Specify _____

1.6 What is the total amount of income earned last year by people who live in your house?

_____ Yuan

Section 2. - Household Composition

Now I would like to ask some questions about your current household.

- 2.1 How many people including yourself live in your house?

_____ People

- 2.2 How many children 10 years old and younger live in your house?

_____ Children

- 2.3 Do you share a bed with someone else in your household?

No.....0 (Skip to 2.5)

Yes.....1

- 2.4 Whom do you share a bed with? (Circle all that apply)

Spouse.....0

Children.....1 How many? _____

Other.....2 specify _____

- 2.5 What kind of animals live inside your house with you? (Circle all that apply)

None.....0

Cats.....1 How many? _____

Dogs.....2 How many? _____

Rabbits.....3 How many? _____

Other, specify _____

- 2.6 What kind of animals live outside your house in the court yard? (Circle all that apply)

None.....0

Dogs.....1

Pigs.....2

Sheep.....3

Ducks.....4

Cows.....5

Donkeys.....6

Cats.....7

Chickens.....8

Rabbits.....9

Goats.....10

Other, specify _____

Section 3. Water and Personal Habits (Personal Hygiene)

Now I would like to ask some questions about your current source of water and washing habits.

3.1 What is your usual source of drinking water?

Deep private well..... 1
 Shallow private well.....2
 Deep village well.....3
 Shallow village well.....4
 Pond.....5
 River.....6
 Ditch.....7
 Running water.....8

3.2 Do you store your drinking water in a jar?

No.....0 (Skip to 3.4)
 Yes.....1

3.3 If yes, How long do you store your drinking water?

_____ Number of days

Or

_____ Number of weeks

3.4 In the summer, when you drink water, how often is it boiled?

Always.....0
 More than half the time...1
 Less than half the time ..2
 Never.....3

3.5 How often are your chopsticks and bowls washed after meals?

Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3 (skip to 3.8)

3.6 When your chopsticks and bowls are washed, what kind of water do you usually use?

Hot water.....0
 Hot & cold water equally..1
 Cold water.....2

- 3.7 When your chopsticks and bowls are washed, how often is detergent used?
- Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3
- 3.8 How often is you cup washed after you use it?
- Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3
- 3.9 How often do members of your family share cups?
- Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3
- 3.10 How often do you wash your hands before eating?
- Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3
- 3.11 How often do you wash your hands after bowel movements?
- Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3
- 3.12 How often do you wash your hands before preparing food?
- Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3
 Don't prepare food.....9

3.13 How often do you use soap when you wash your hands?

Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3

3.14 When it is cold outside, how many times per day, week, month, or year do you wash yourself?

___ Times per day
 ___ Times per week
 ___ Times per month
 ___ Times per year
 ___ Never wash...997 (skip to 3.16)

3.15 When it is cold, where do you usually wash yourself?

At Home0
 River.....1
 Stream.....2
 Pond.....3
 Ditch.....4
 Public Bath House.....5

3.16 When it is warm outside, how many times per day, week, month, or year do you wash yourself?

___ Times per day
 ___ Times per week
 ___ Times per month
 ___ Times per year
 ___ Never wash...997 (skip to 3.19)

3.17 When it is warm, where do you usually wash yourself?

In the house.....0
 In the yard.....1
 River.....2
 Stream.....3
 Pond.....4
 Ditch.....5
 Public Bath House.....6

3.18 How often do you use soap when you wash yourself?

Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3

3.19 How many times per day, week, month, or year do you spit onto the ground?

☐ Times per day
☐ Times per week
☐ Times per month
☐ Times per year
☐ Never spit...997

3.20 How many times per day, week, or month do you brush your teeth?

☐ Times per day
☐ Times per week
☐ Times per month
☐ Never997

3.21 How often do you swim in the river or reservoir?

☐ Times per year
☐ Never997

3.22 Have you ever washed diapers?

No.....0 (Skip to 3.24)
 Yes.....1

3.23 Where did you wash the diapers?

At Home.....0
 River.....1
 Stream.....2
 Pond.....3
 Ditch.....4

3.24 How often did you feed your child (children) by prechewing their food?

Often.....0
 Sometimes.....1
 Never.....2

3.25 How often did you kiss your child (children) on the mouth when they were less than ten years old?

Often.....0
 Sometimes.....1
 Never.....2

Section 4. Eating Habits

Now I would like to ask some questions about your eating habits and health.

- 4.1 How do you prefer the saltiness of your food? Do you prefer lightly salty, moderately salty, or very salty?

Lightly salty.....1
 Moderately salty.....2
 Very salty.....3

- 4.2 Do you eat sour pancake?

Yes.....1
 No.....0 (Skip to 4.3)

For how many months did you eat sour pancake last year? _____ Months

On average how many days per month did you eat it? _____ Days

On average how many Lian did you eat per day? _____ Lian

- 4.2 Do you eat sweet pancake?

Yes.....1
 No.....0 (Skip to 4.4)

For how many months did you eat sweet pancake last year? _____ Months

On average how many days per month did you eat it? _____ Days

On average how many Lian did you eat per day? _____ Lian

- 4.4 How many times per day, week, month, or year do you eat raw fruits or vegetables in season?

_____ Times per day
 _____ Times per week
 _____ Times per month
 _____ Times per year
 Never eat ..997 (Skip to 4.7)

- 4.5 How often are your raw fruits and vegetables washed before you eat them?

Always.....0
 More than half the time...1
 Less than half the time...2
 Never.....3

4.6 How often are your raw fruits and vegetables peeled before you eat them?

Always.....0
More than half the time...1
Less than half the time...2
Never.....3

4.7 How many times per day, week, month, or year do you have gastric reflux?

___ Times per day
___ Times per week
___ Times per month
___ Times per year
Never have reflux ..997

Section 5. Smoking and Drinking Habits

Now I would like to ask some questions about your smoking and drinking habits.

5.1 Did you ever smoke cigarettes?

No.....0 (Skip to 5.6)

Yes.....1

5.2 How old were you when you started smoking?

_____ years old

5.3 Do you currently smoke cigarettes?

No.....0

Yes.....1

5.4 How many cigarettes do (did) you usually smoke each day?

_____ cigarettes per day

5.5 How many years have you smoked (did you smoke) cigarettes?

_____ number of years

5.6 Did you ever drink alcoholic beverages?

No.....0 (Skip to Section 6)

Yes.....1

5.7 How old were you when you started drinking alcoholic beverages?

_____ years old

5.8 Do you currently drink alcoholic beverages?

No.....0

Yes.....1

5.9 How many times do (did) you usually drink each week?

_____ times per week

5.10 How many years have you drunk (did you drink) alcoholic beverages?

_____ number of years

Section 6. Childhood Household Composition

Now I would like to ask some questions about your household when you were a child.

6.1 When you were 10 years old, what kind of floor did your house have?

Cement.....0
 Brick.....1
 Ground.....2
 Other, specify _____

6.2 When you were 10 years old, how many people including yourself, lived in your house?

_____ People

6.3 When you were 10 years old, how many brothers and sisters did you have?

_____ siblings
 (If none skip to 6.5)

6.4 What were the ages of your brothers and sisters when you were 10 years old?

_____ years old
 _____ years old
 _____ years old
 _____ years old
 _____ years old

5 When you were 10 years old, what kind of animals lived inside your house with you?
 (Circle all that apply)

None.....0
 Cats.....1 How many? _____
 Dogs.....2 How many? _____
 Rabbits...3 How many? _____
 Other, specify _____

6.6 When you were 10 years old, how many people did you share a bed with?

_____ People

6.7 When you were 5 years old, how many people did you share a bed with?

_____ People

6.8 When you were a small child, how often did your parents prechew your food for you?

Often.....0

Sometimes.....1

Never.....2

Don't know.....9

6.9 When you were a small child, how often did your parents kiss you on the mouth?

Often.....0

Sometimes.....1

Never.....2

Don't know.....9

I have nothing else to ask today. Thank you for answering my questions. The information you have given us is very important.

Interviewer Comments

1. The subject's cooperation was:

Very Good.....1
 Good.....2
 Fair.....3
 Poor.....4

2. The overall quality of this interview is:

High Quality.....1
 Generally Reliable....2
 Questionable.....3
 Unsatisfactory.....4

3. The main reason(s) for the unsatisfactory or questionable quality is (are) that the subject
 (Code all that apply)

Was too01
 Had poor hearing, or speech.....02
 Was evasive or suspicious.....03
 Was upset or depressed.....04
 Was bored or disinterested.....05
 Did not understand the questions..06
 Could not remember events well....07
 Was confused or disoriented.....08
 Other, Specify_____

4. How many individuals besides the study subject were present during the interview?

_____ Number present

5. Who was present during the interview? (Circle all that apply)

Spouse.....1
 Mother.....2
 Father.....3
 Daughter.....4 How many? _____
 Son.....5 How many? _____
 Other, specify_____

INFORMED CONSENT

Subject I.D. number:

Purpose of the Research

In order to improve the people's health, the Beijing Institute for Cancer Research, the Weifang Medical Institute, and the Linqu Bureau of Health in collaboration with the United States National Cancer Institute are initiating a medical research trial among individuals like yourself who participated in earlier medical examinations. The study will help us to test the effectiveness of several treatments in fighting the high rate of stomach cancer in Linqu. You are invited to join this 42-month research study.

What is Involved

The study involves taking three types of active treatment or placebo, answering some questions, having samples of your blood and breath collected, and having a gastroscopic examination.

1. Treatments:

One treatment involves an antibiotic medication to help cure a bacterial infection that occurs in the stomach and that may increase risk of stomach cancer. Another is a garlic supplement, and a third is a vitamin/mineral supplement. The treatments involve taking 3 capsules before the morning meal and 3 before the evening meal. Bottles containing each type of capsule (either the active ingredient or a placebo) will be distributed monthly. The study will be "blinded" in that we will not know which combinations of the active treatments you will be taking until the trial is over (unless there is a medical need to know the treatment). This helps ensure the scientific validity of the trial. One in eight participants will receive only placebo.

2. Information Collection:

You will also be given a short questionnaire at the time of enrollment into the trial and at months 12, 24, and 42 to collect basic information on health and lifestyle factors.

3. Blood Collections:

At months 12, 24, and 42 you will be asked to have 5 ml of blood taken from your arm. This blood will be used to determine the effect of treatment on bacteria, and some will be stored for later studies related to this project. Every three months we will ask 60 different randomly selected participants to have 10 ml of blood taken from their arm to measure nutrients and garlic compounds, which can be used to assess pill-taking compliance.

4. Breath Collections:

At about 3 and 6 months into the trial, you will be asked to take a teaspoon of a liquid containing natural protein byproducts and to breathe into a tube or bag. Your breath will be used to assess the effectiveness of treatment for bacteria. A randomly selected group of 200 participants will be asked to have their breath collected at months 12, 24, and 42.

5. Gastroscopic Examination:

After 42 months, you will be asked to have another gastroscopic examination like you have had before. This exam, done by a skilled gastroenterologist, will tell us whether the pills you have been taking have helped improve stomach conditions.

Risks

There is only minimal risk to you in participating in this research study. Vitamin and mineral supplements and the ingredients in the garlic preparation are widely available around the world. There is very little evidence of toxicity associated with these agents. A pilot study in one of our 14 villages revealed no serious side-effects. There are also no known side-effects associated with the breath test.

The treatment for bacteria (amoxicillin/omeprazole), however, has been associated with side effects in some cases. The most common symptoms observed in the pilot study in Linqu were diarrhea (13%), rashes (8%), and abdominal bloating (5%). Also reported were a few complaints of constipation and nausea. In rare instances a sore mouth can result, and in extremely rare instances more severe reactions can occur. Please tell us if you have had allergic reactions, such as skin rashes to drugs in the past. Also, if you enroll in the trial, please tell us if any of these symptoms (especially rashes) develop during the trial.

Blood drawing may cause a small amount of pain when the needle is passed into the vein, but should not cause long term pain. After blood is taken sometimes there can be a small bruise or soreness at the site.

While there have been no serious complications in over 8000 gastroscopic procedures of this type, there is a very small chance that bleeding may occur. You will be instructed how to detect and treat such problems.

Benefits

If you agree to participate you will become part of a research study which may help us understand more about this bacterial infection and about garlic and vitamin supplements and whether the treatments may lower the risk of changes in the stomach that could lead to cancer. You also may benefit by having your health monitored closely and possibly having stomach cancer diagnosed at an early, potentially curable stage.

Confidentiality of Records

The results of the study we are conducting will be reported in medical journals or at scientific meetings, but we will very carefully keep and handle the records of the study to ensure that your identification will not be made public.

Participation Is Voluntary

You must volunteer to be in the study. You will not be paid for being in the study, and if you want to leave the study at any time, you can and you will still be able to receive medical care.

If you refuse to be in the study, there will be no penalty or loss of your usual benefits to which you are otherwise entitled and you may discontinue your participation in the study at any time without penalty or loss of your otherwise entitled benefits. Nonparticipation will not effect your relationship with the Linqu Health Bureau and you will also be able to take advantage of future activities which the Beijing Institute for Cancer Research may conduct in Linqu.

Who to Contact with Questions

If you have any questions you can contact _____ in your village or any of the persons doing the study.

I understand the information about the study that is contained in the consent form and it has been discussed with me. I have had all my questions answered. I hereby consent to take part in the study.

Signature or Fingerprint of Adult Participant

Date Signed

I administered the informed consent and believe that the study subject understands the purpose of the study and the possible risks and benefits.

Signature of BICR Field Staff

Date Signed

山东干预试验研究问卷调查表

村庄:_____ 组号_____ 研究对象编号(ID):_____

姓名:_____ 性别:1.男 2.女 出生日期____年____月

调查日期:____年____月____日 调查开始时间:_____ 调查结束时间:_____ 总计时间:_____分

调查员编号:_____ 调查员签字:_____

介绍:

谢谢您同意参加这项研究.这次调查(访问)我将问您以下问题:您现在及儿童时期的家庭成员,您的刷牙、吸烟、饮酒及饮食习惯,您及您家庭的基本情况。您的合作对这次调查研究非常重要,将有助于我们更多地了解临朐县成年人的健康问题。我想提醒您,您的参与是自愿的,您提供的信息(内容)我将予以保密,并仅用于研究。

第一部分 基本情况

我先问您几个基本问题:

1.1 您现在的婚姻状况如何?

在婚……………1

丧妻或丧夫…2(跳到 1.4)

离婚或分居…3(跳到 1.4)

从未…4(跳到 1.4)

1.2 您配偶上过几年学? _____年

没上过学…0

1.3 您配偶目前从事什么工作?

粮农……………1

果农……………2

菜农……………3

村干部…………4

没工作…………5

其它,具体说明_____

1.4 您上过几年学?

_____年

没上过学…0

1.5 您现在从事什么工作?

粮农……………1

果农……………2

菜农……………3

村干部…………4

其它,具体说明_____

1.6 去年您全家总共收入多少钱? _____元

第二部分 家庭情况

现在我想问您目前的家庭情况

2.1 您家有几口人(包括您在内)? _____人(如只有1人,跳到2.5)

2.2 您家10岁及以下的孩子有几个? _____个

2.3 在家里您与别人合用一张床吗?

不……0(跳到2.5)

是……1

2.4 您与谁合用一张床?(在相应符号上划圈)

配偶……0

孩子……1 几个_____?

其它……2 具体说明_____

2.5 您家屋内养有什么动物?(在相应的符号上划圈)

无……0

猫……1 几只_____

狗……2 几只_____

兔子……3 几只_____

其它,具体说明_____

2.6 您家院内养有什么动物?(在相应符号上划圈)

无……0

狗……1

猪……2

绵羊……3

鸭……4

牛……5

驴……6

猫……7

鸡……8

兔子……9

山羊……10

其它,具体说明_____

第三部分 饮水和个人习惯(个人卫生)

3.1 您平时饮用什么水?

私人深井……1

私人浅井……2

村里深井.....3
 村里浅井.....4
 池塘.....5
 河水.....6
 沟渠.....7
 自来水.....8

3.2 您将饮用水存在缸里吗?

不.....0(跳到 3.4)

是.....1

3.3 如是,您把水存放多长时间? _____天,或____星期

3.4 在夏天,饮用开水的情况如何?

总是.....0

多半时间是.....1

多半时间不是.....2

从不.....3

3.5 饭后碗筷刷洗情况如何?

总是洗.....0

多半洗.....1

多半不洗.....2

从不洗.....3(跳到 3.8)

3.6 您家用什么水洗碗筷?

热水.....0

热水凉水各半.....1

凉水.....2

3.7 您家刷洗碗筷时使用洗洁剂吗?

总是.....0

多半是.....1

多半不是.....2

从来不.....3

3.8 您家的茶杯用后洗刷吗?

总是.....0

多半是.....1

多半不是.....2

从来不.....3

3.9 您家的茶杯合用吗?

总是.....0

多半是.....1

多半不是.....2

从来不.....3

3.10 您饭前洗手吗?

总是.....0

多半是.....1

多半不是.....2

- 3.11 您便后洗手吗? 从不.....3
- 总是.....0
多半是.....1
多半不是.....2
从不.....3
- 3.12 您做饭之前洗手吗?
- 总是.....0
多半是.....1
多半不是.....2
从不.....3
不做饭.....9
- 3.13 您洗手时用肥皂吗?
- 总是.....0
多半是.....1
多半不是.....2
从不.....3
- 3.14 当气温变冷时,您多长时间洗一次澡?
- _____次/天
_____次/周
_____次/月
从不.....997(跳到 3.16)
- 3.15 天冷时,您一般在哪儿洗澡? 屋里.....0
江河.....1
小河.....2
池塘.....3
沟渠.....4
浴池.....5
- 3.16 当气温暖和时,您多长时间洗一次澡?
- _____次/天
_____次/周
_____次/月
从不.....997(跳到 3.19)
- 3.17 天暖时,您一般在哪儿洗澡? 屋里.....0
院子里.....1
江河.....2
小河.....3
池塘.....4
沟渠.....5
浴池.....6
- 3.18 您洗澡时用肥皂吗?
- 总是.....0
多半是.....1
多半不是.....2

3.19 您随地吐痰吗?

_____次/天
 _____次/周
 _____次/月
 从不.....997

3.20 您刷牙吗?

_____次/天
 _____次/周
 _____次/月
 从不.....997

3.21 您常在河里或水库里游泳吗?

_____次/年
 从不.....997

3.22 您洗过婴儿的尿布吗?

不.....0(跳到 3.24)
 是.....1

3.23 您在什么地方洗婴儿尿布?

家里.....0
 江河.....1
 小河.....2
 池塘.....3
 沟渠.....4

3.24 您经常将食物嚼烂喂婴儿吗?

经常.....0
 有时.....1
 从不.....2

3.25 您经常口对口亲吻您的孩子(10岁以下)吗?

经常.....0
 有时.....1
 从不.....2

第四部分 饮食习惯

现在我想问您一些有关饮食和健康的问题

4.1 您喜欢吃咸还是淡? 淡.....1
 中等.....2
 咸.....3

4.2 您吃酸煎饼吗? 是.....1

不.....0(跳到 4.3)

去年一共吃了几个月的酸煎饼? _____月
 平均一个月吃几天? _____天
 平均一天吃几两? _____两

4.3 您吃甜煎饼吗? 是.....1

不……0(跳到 4.4)

去年一共吃了几个月的甜煎饼? _____ 月

平均一个月吃几天? _____ 天

平均一天吃几两? _____ 两

4.4 您生吃水果或蔬菜吗?

_____ 次/天

_____ 次/周

_____ 次/月

_____ 次/年

从不吃……997(跳到 4.7)

4.5 生吃水果或蔬菜前常洗吗?

总是……0

多半是……1

多半不是……2

从来不……3

4.6 生吃水果或蔬菜前常削皮吗?

总是……0

多半是……1

多半不是……2

从来不……3

4.7 您有没有过胃内返酸?

_____ 次/天

_____ 次/周

_____ 次/月

_____ 次/年

从没有过……997

第五部分 吸烟和饮酒习惯

现在我想问一下您的吸烟与饮酒情况

5.1 您吸过卷烟吗?

否……0(跳到 5.6)

是……1

5.2 您几岁开始吸烟? _____ 岁

5.3 目前您还吸卷烟吗?

否……0

是……1

5.4 通常您一天吸几支烟? _____ 支/天

5.5 您的吸烟史有多长? _____ 年

5.6 您有饮酒史吗?

无……0(跳到第 6 部分)

有……1

5.7 您几岁开始饮酒? _____ 岁

5.8 现在您还饮酒吗?

不.....0

是.....1

5.9 通常您一周饮几次酒? _____次/周

5.10 您的饮酒史有多长? _____年

第六部分 童年时的家庭情况

现在我想问您小时候家庭的一些情况

6.1 您 10 岁时,家里的地面是用什么铺成的?

水泥.....0

砖.....1

土.....2

其它,具体说明 _____

6.2 您 10 岁时家里有几口人?(包括您在内)_____人

6.3 您 10 岁时兄弟姐妹几个(不包括您本人)? _____人

(如没有,跳到 6.5)

6.4 您 10 岁时他(她)们分别几岁?(已婚的要注明)

_____	_____	岁
_____	_____	岁
_____	_____	岁
_____	_____	岁
_____	_____	岁

6.5 您 10 岁时,您屋内养有什么动物?(在相应符号上划圈)

无.....0

猫.....1 _____只

狗.....2 _____只

兔子.....3 _____只

其它,具体说明 _____

6.6 您 10 岁时,几人与您同用一张床? _____人

6.7 您 5 岁时,几人与您同用一张床? _____人

6.8 您小时候父母经常将食物嚼烂后喂您吗?

经常.....0

有时.....1

从不.....2

不知道...997

6.9 您小时候您父母口对口亲吻过您吗?

经常.....0

有时.....1

从不.....2

不知道...997

我问完了,谢谢您回答我的问题,您给我们提供的情况很重要。

调查对象的合作情况：
 很好.....1
 好.....2
 可以.....3
 不好.....4

这次调查的质量：
 质量高.....1
 一般可靠.....2
 可能有问题.....3
 不能令人满意.....4

可能有的问题或令人不满意的原因(在所有的原因符号上划圈)

病重.....01
 听力不行或口齿不清...02
 推诿或多疑.....03
 心烦或压抑.....04
 无聊或不感兴趣.....05
 听不懂问话.....06
 记忆不好.....07
 拒绝或不配合.....08
 其它,具体说明_____

除调查对象外,调查时还有几人在场? _____人

有谁在场?(在相应的符号上划圈)

配偶.....1
 母亲.....2
 父亲.....3
 女儿.....4 _____人
 儿子.....5 _____人
 其它,具体说明_____

BIBLIOGRAPHY

1. Cave DR. How is *Helicobacter pylori* transmitted? *Gastroenterology* 1997; 113:S9-14.
2. Graham DY. Evolution of concepts regarding *Helicobacter pylori*: from a cause of gastritis to a public health problem. *Am.J.Gastroenterol.* 1994; 89:469-472.
3. Warren JR, Marshall BJ. *Lancet* 1983; 1:1273-1275.
4. Goodwin CS, Armstrong JA, Chilvers T, Peters M, Colins MD, Sly L, et al. Transfer of *Campylobacter pylori* and *Campylobacter mustelae* to *Helicobacter* gen. nov. as *Helicobacter pylori* comb. nov. and *Helicobacter mustelae* comb. nov., respectively. *Int.J.Sys.Bacteriol.* 1989; 39:397-405.
5. Graham DY. Benefits from elimination of *Helicobacter pylori* infection include major reduction in the incidence of peptic ulcer disease, gastric cancer, and primary gastric lymphoma. *Prev.Med.* 1994; 23:712-716.
6. McColl KE. What remaining questions regarding *Helicobacter pylori* and associated diseases should be addressed by future research? View from Europe. *Gastroenterology* 1997; 113:S158-S162
7. IARC Working Group. Schistosomes, Liver Flukes, and *Helicobacter pylori*. 1994; 61, 177-241. Lyon, France: World Health Organization, International Agency for Research on Cancer.
8. Cheli R, Crespi M, Testino G, Citarda F. Gastric cancer and *Helicobacter pylori*: biologic and epidemiologic inconsistencies. *J.Clin.Gastroenterol.* 1998; 26:3-6.
9. Crespi M, Citarda F. *Helicobacter pylori* and gastric cancer: what is the real risk? *Gastroenterologist.* 1998; 6:16-20.
10. Parsonnet J. The incidence of *Helicobacter pylori* infection. *Aliment.Pharmacol.Ther.* 1995; 9 Suppl 2:45-51:45-51.

11. Dunn BE, Cohen H, Blaser MJ. *Helicobacter pylori*. Clin.Microbiol.Rev. 1997; 10:720-741.
12. You WC, Blot WJ, Li JY, Chang YS, Jin ML, Kneller R, et al. Precancerous gastric lesions in a population at high risk of stomach cancer. Cancer Res. 1993; 53:1317-1321.
13. Zhang L, Blot WJ, You WC, Chang YS, Kneller RW, Jin ML, et al. *Helicobacter pylori* antibodies in relation to precancerous gastric lesions in a high-risk Chinese population. Cancer Epidemiol.Biomarkers.Prev. 1996; 5:627-630.
14. Marshall BJ, Warren JR. Unidentified curved bacilli in the stomach of patients with gastritis and peptic ulceration. Lancet 1984; 1:1311-1315.
15. Lambert JR, Lin SK, Aranda-Michel J. *Helicobacter pylori*. Scand.J.Gastroenterol.Suppl. 1995; 208:33-46:33-46.
16. Blaser MJ. Ecology of *Helicobacter pylori* in the human stomach. J.Clin.Invest. 1997; 100:759-762.
17. Atherton JC. The clinical relevance of strain types of *Helicobacter pylori*. Gut 1997; 40:701-703.
18. Fallone CA. Determinants of ethnic or geographical differences in infectivity and transmissibility of *Helicobacter pylori*. Can.J.Gastroenterol. 1999; 13:251-255.
19. Moriai T, Hirahara N. Clinical course of acute gastric mucosal lesions caused by acute infection with *Helicobacter pylori* [letter]. N.Engl.J.Med. 1999; 341:456-457.
20. Dubois A, Berg DE, Incecik ET, Fiala N, Heman-Ackah LM, Perez-Perez GI, et al. Transient and persistent experimental infection of nonhuman primates with *Helicobacter pylori*: implications for human disease. Infect.Immun. 1996; 64:2885-2891.
21. Perez-Perez GI, Dworkin BM, Chodos JE, Blaser MJ. *Campylobacter pylori*

- antibodies in humans. *Ann.Intern.Med.* 1988; 109:11-17.
22. Faigel DO, Childs M, Furth EE, Alavi A, Metz DC. New noninvasive tests for *Helicobacter pylori* gastritis. Comparison with tissue-based gold standard. *Dig.Dis.Sci.* 1996; 41:740-748.
 23. Feldman RA, Evans SJ. Accuracy of diagnostic methods used for epidemiological studies of *Helicobacter pylori*. *Aliment.Pharmacol.Ther.* 1995; 9 Suppl 2:21-31:21-31.
 24. Taylor DN, Blaser MJ. The epidemiology of *Helicobacter pylori* infection. *Epidemiol.Rev.* 1991; 13:42-59:42-59.
 25. Andersen LP, Rosenstock SJ, Bonnevie O, Jorgensen T. Seroprevalence of immunoglobulin G, M, and A antibodies to *Helicobacter pylori* in an unselected Danish population. *Am.J.Epidemiol.* 1996; 143:1157-1164.
 26. Atherton JC. Non-endoscopic tests in the diagnosis of *Helicobacter pylori* infection. *Aliment.Pharmacol.Ther.* 1997; 11 Suppl 1:11-20:11-20.
 27. Hook-Nikanne J, Perez-Perez GI, Blaser MJ. Antigenic characterization of *Helicobacter pylori* strains from different parts of the world. *Clin.Diagn.Lab.Immunol.* 1997; 4:592-597.
 28. Bodhidatta L, Hoge CW, Churnratanakul S, Nirdnoy W, Sampathanukul P, Tungtaem C, et al. Diagnosis of *Helicobacter pylori* infection in a developing country: comparison of two ELISAs and a seroprevalence study. *J.Infect.Dis.* 1993; 168:1549-1553.
 29. Chang-Claude J, Raedsch R, Waldherr R, von Wulffen H, Crespi M, Yang GR, et al. Prevalence of *Helicobacter pylori* infection and gastritis among young adults in China. *Eur.J.Cancer Prev.* 1995; 4:73-79.
 30. Groves FD, Zhang L, Li JY, You WC, Chang YS, Zhao L, et al. Comparison of two enzyme-linked immunosorbent assay tests for diagnosis of *Helicobacter pylori* infection in China. *Cancer Epidemiol.Biomarkers.Prev.* 1997; 6:551-552.

31. Lindkvist P, Asrat D, Nilsson I, Tsega E, Olsson GL, Wretling B, et al. Age at acquisition of *Helicobacter pylori* infection: comparison of a high and a low prevalence country. *Scand.J.Infect.Dis.* 1996; 28:181-184.
32. Parsonnet J. *Helicobacter pylori*. *Infect.Dis.Clin.North Am.* 1998; 12:185-197.
33. Xia HH, Talley NJ. Natural acquisition and spontaneous elimination of *Helicobacter pylori* infection: clinical implications. *Am.J.Gastroenterol.* 1997; 92:1780-1787.
34. Sipponen P. *Helicobacter pylori* gastritis--epidemiology. *J.Gastroenterol.* 1997; 32:273-277.
35. Blaser MJ. The role of *Helicobacter pylori* in gastritis and its progression to peptic ulcer disease. *Aliment.Pharmacol.Ther.* 1995; 9 Suppl 1:27-30:27-30.
36. Blaser MJ, Chyou PH, Nomura A. Age at establishment of *Helicobacter pylori* infection and gastric carcinoma, gastric ulcer, and duodenal ulcer risk. *Cancer Res.* 1995; 55:562-565.
37. Correa P. *Helicobacter pylori* and gastric cancer: state of the art. *Cancer Epidemiol.Biomarkers.Prev.* 1996; 5:477-481.
38. Parsonnet J, Friedman GD, Vandersteen DP, Chang Y, Vogelmann JH, Orentreich N, et al. *Helicobacter pylori* infection and the risk of gastric carcinoma. *N.Engl.J.Med.* 1991; 325:1127-1131.
39. Nomura A, Stemmermann GN, Chyou PH, Kato I, Perez-Perez GI, Blaser MJ. *Helicobacter pylori* infection and gastric carcinoma among Japanese Americans in Hawaii. *N.Engl.J.Med.* 1991; 325:1132-1136.
40. Forman D, Newell DG, Fullerton F, Yarnell JW, Stacey AR, Wald N, et al. Association between infection with *Helicobacter pylori* and risk of gastric cancer: evidence from a prospective investigation. *BMJ.* 1991; 302:1302-1305.

41. Hansen S, Melby KK, Aase S, Jellum E, Vollset SE. *Helicobacter pylori* infection and risk of cardia cancer and non-cardia gastric cancer. A nested case-control study. *Scand.J.Gastroenterol.* 1999; 34:353-360.
42. Queiroz DM, Mendes EN, Rocha GA, Oliveira AM, Oliveira CA, Cabral MM, et al. Serological and direct diagnosis of *Helicobacter pylori* in gastric carcinoma: a case-control study. *J.Med.Microbiol.* 1999; 48:501-506.
43. Nyren O. Is *Helicobacter pylori* really the cause of gastric cancer? *Semin.Cancer Biol.* 1998; 8:275-283.
44. Chow WH, Blaser MJ, Blot WJ, Gammon MD, Vaughan TL, Risch HA, et al. An inverse relation between *cagA*⁺ strains of *Helicobacter pylori* infection and risk of esophageal and gastric cardia adenocarcinoma. *Cancer Res.* 1998; 58:588-590.
45. Oberg S, Peters JH, Nigro JJ, Theisen J, Hagen JA, DeMeester SR, et al. *Helicobacter pylori* is not associated with the manifestations of gastroesophageal reflux disease. *Arch.Surg.* 1999; 134:722-726.
46. Blaser MJ. Hypothesis: the changing relationships of *Helicobacter pylori* and humans: implications for health and disease. *J.Infect.Dis.* 1999; 179:1523-1530.
47. Wu JC, Sung JJ, Ng EK, Go MY, Chan WB, Chan FK, et al. Prevalence and distribution of *Helicobacter pylori* in gastroesophageal reflux disease: a study from the East. *Am.J.Gastroenterol.* 1999; 94:1790-1794.
48. Strachan DP, Mendall MA, Carrington D, Butland BK, Yarnell JW, Sweetnam PM, et al. Relation of *Helicobacter pylori* infection to 13-year mortality and incident ischemic heart disease in the caerphilly prospective heart disease study. *Circulation* 1998; 98:1286-1290.
49. Replogle ML, Glaser SL, Hiatt RA, Parsonnet J. Biologic sex as a risk factor for *Helicobacter pylori* infection in healthy young adults. *Am.J.Epidemiol.* 1995; 142:856-863.
50. Parente F, Bianchi PG. *Helicobacter pylori* infection and ischaemic heart disease: is there a link? *Ital.J.Gastroenterol.Hepatol.* 1998; 30:119-123.

51. Folsom AR, Nieto FJ, Sorlie P, Chambless LE, Graham DY. *Helicobacter pylori* seropositivity and coronary heart disease incidence. Atherosclerosis Risk In Communities (ARIC) Study Investigators. *Circulation* 1998; 98:845-850.
52. Ossewaarde JM, Feskens EJ, De Vries A, Vallinga CE, Kromhout D. *Chlamydia pneumoniae* is a risk factor for coronary heart disease in symptom-free elderly men, but *Helicobacter pylori* and cytomegalovirus are not. *Epidemiol.Infect.* 1998; 120:93-99.
53. Danesh J. Is there a link between chronic *Helicobacter pylori* infection and coronary heart disease? *Eur.J.Surg.Suppl.* 1998; 27-31.
54. Shinchi K, Ishii H, Imanishi K, Kono S. Relationship of cigarette smoking, alcohol use, and dietary habits with *Helicobacter pylori* infection in Japanese men. *Scand.J.Gastroenterol.* 1997; 32:651-655.
55. Alaganantham TP, Pai M, Vaidehi T, Thomas J. Seroepidemiology of *Helicobacter pylori* infection in an urban, upper class population in Chennai. *Indian J.Gastroenterol.* 1999; 18:66-68.
56. Kawasaki M, Kawasaki T, Ogaki T, Itoh K, Kobayashi S, Yoshimizu Y, et al. Seroprevalence of *Helicobacter pylori* infection in Nepal: low prevalence in an isolated rural village. *Eur.J.Gastroenterol.Hepatol.* 1998; 10:47-50.
57. Lin SK, Lambert JR, Nicholson L, Lukito W, Wahlqvist M. Prevalence of *Helicobacter pylori* in a representative Anglo-Celtic population of urban Melbourne. *J.Gastroenterol.Hepatol.* 1998; 13:505-510.
58. Senra-Varela A, Lopez-Saez JB, Gomez-Biondi V. Prevalence of *Helicobacter pylori* infection in two Spanish regions with different incidence of gastric cancer. *Eur.J.Epidemiol.* 1998; 14:491-494.
59. Souto FJ, Fontes CJ, Rocha GA, de Oliveira AM, Mendes EN, Queiroz DM. Prevalence of *Helicobacter pylori* infection in a rural area of the state of Mato Grosso, Brazil. *Mem.Inst.Oswaldo.Cruz.* 1998; 93:171-174.
60. Torres J, Leal-Herrera Y, Perez-Perez G, Gomez A, Camorlinga-Ponce M, Cedillo-Rivera R, et al. A community-based seroepidemiologic study of

- Helicobacter pylori* infection in Mexico. *J.Infect.Dis.* 1998; 178:1089-1094.
61. Murray LJ, McCrum EE, Evans AE, Bamford KB. Epidemiology of *Helicobacter pylori* infection among 4742 randomly selected subjects from Northern Ireland. *Int.J.Epidemiol.* 1997; 26:880-887.
 62. Peach HG, Pearce DC, Farish SJ. *Helicobacter pylori* infection in an Australian regional city: prevalence and risk factors. *Med.J.Aust.* 1997; 167:310-313.
 63. Chow TK, Lambert JR, Wahlqvist ML, Hsu-Hage BH. *Helicobacter pylori* in Melbourne Chinese immigrants: evidence for oral- oral transmission via chopsticks. *J.Gastroenterol.Hepatol.* 1995; 10:562-569.
 64. Gasbarrini G, Pretolani S, Bonvicini F, Gatto MR, Tonelli E, Megraud F, et al. A population based study of *Helicobacter pylori* infection in a European country: the San Marino Study. Relations with gastrointestinal diseases. *Gut* 1995; 36:838-844.
 65. Gilboa S, Gabay G, Zamir D, Zeev A, Novis B. *Helicobacter pylori* infection in rural settlements (Kibbutzim) in Israel. *Int.J.Epidemiol.* 1995; 24:232-237.
 66. Murray LJ, Bamford KB, O'Reilly DP, McCrum EE, Evans AE. *Helicobacter pylori* infection: relation with cardiovascular risk factors, ischaemic heart disease, and social class. *Br.Heart J.* 1995; 74:497-501.
 67. Teh BH, Lin JT, Pan WH, Lin SH, Wang LY, Lee TK, et al. Seroprevalence and associated risk factors of *Helicobacter pylori* infection in Taiwan. *Anticancer Res.* 1994; 14:1389-1392.
 68. Anonymous. Epidemiology of, and risk factors for, *Helicobacter pylori* infection among 3194 asymptomatic subjects in 17 populations. The EUROGAST Study Group. *Gut* 1993; 34:1672-1676.
 69. Mitchell HM, Li YY, Hu PJ, Liu Q, Chen M, Du GG, et al. Epidemiology of *Helicobacter pylori* in southern China: identification of early childhood as the critical period for acquisition. *J.Infect.Dis.* 1992; 166:149-153.

70. Sitas F, Forman D, Yarnell JW, Burr ML, Elwood PC, Pedley S, et al. *Helicobacter pylori* infection rates in relation to age and social class in a population of Welsh men. *Gut* 1991; 32:25-28.
71. Wong BC, Lam SK, Ching CK, Hu WH, Kwok E, Ho J, et al. Differential *Helicobacter pylori* infection rates in two contrasting gastric cancer risk regions of South China. China Gastric Cancer Study Group. *J.Gastroenterol.Hepatol.* 1999; 14:120-125.
72. Buckley MJ, O'Shea J, Grace A, English L, Keane C, Hourihan D, et al. A community-based study of the epidemiology of *Helicobacter pylori* infection and associated asymptomatic gastroduodenal pathology. *Eur.J.Gastroenterol.Hepatol.* 1998; 10:375-379.
73. Breuer T, Sudhop T, Hoch J, Sauerbruch T, Malfertheiner P. Prevalence of and risk factors for *Helicobacter pylori* infection in the western part of Germany. *Eur.J.Gastroenterol.Hepatol.* 1996; 8:47-52.
74. Malaty HM, Kim JG, Kim SD, Graham DY. Prevalence of *Helicobacter pylori* infection in Korean children: inverse relation to socioeconomic status despite a uniformly high prevalence in adults. *Am.J.Epidemiol.* 1996; 143:257-262.
75. Martin-de-Argila C, Boixeda D, Canton R, Mir N, de Rafael L, Gisbert J, et al. *Helicobacter pylori* infection in a healthy population in Spain. *Eur.J.Gastroenterol.Hepatol.* 1996; 8:1165-1168.
76. Matysiak-Budnik T, Knapik Z, Megraud F, Lubczynska-Kowalska W, Gosciniak G, Bouchard S, et al. *Helicobacter pylori* infection in Eastern Europe: seroprevalence in the Polish population of Lower Silesia. *Am.J.Gastroenterol.* 1996; 91:2513-2515.
77. Malaty HM, Graham DY. Importance of childhood socioeconomic status on the current prevalence of *Helicobacter pylori* infection. *Gut* 1994; 35:742-745.
78. Webb PM, Knight T, Greaves S, Wilson A, Newell DG, Elder J, et al. Relation between infection with *Helicobacter pylori* and living conditions in childhood: evidence for person to person transmission in early life. *BMJ.* 1994; 308:750-753.

79. Malaty HM, Evans DG, Evans DJJ, Graham DY. *Helicobacter pylori* in Hispanics: comparison with blacks and whites of similar age and socioeconomic class. *Gastroenterology* 1992; 103:813-816.
80. Graham DY, Malaty HM, Evans DG, Evans DJJ, Klein PD, Adam E. Epidemiology of *Helicobacter pylori* in an asymptomatic population in the United States. Effect of age, race, and socioeconomic status. *Gastroenterology* 1991; 100:1495-1501.
81. Loffeld RJ, Stobberingh E, van Spreeuwel JP, Flendrig JA, Arends JW. The prevalence of anti-*Helicobacter* (*Campylobacter*) *pylori* antibodies in patients and healthy blood donors. *J.Med.Microbiol.* 1990; 32:105-109.
82. Taylor DN, Sanchez JL, Smoak BL, DeFraites R. *Helicobacter pylori* infection in Desert Storm troops. *Clin.Infect.Dis.* 1997; 25:979-982.
83. Hyams KC, Taylor DN, Gray GC, Knowles JB, Hawkins R, Malone JD. The risk of *Helicobacter pylori* infection among U.S. military personnel deployed outside the United States. *Am.J.Trop.Med.Hyg.* 1995; 52:109-112.
84. Smoak BL, Kelley PW, Taylor DN. Seroprevalence of *Helicobacter pylori* infections in a cohort of US Army recruits. *Am.J.Epidemiol.* 1994; 139:513-519.
85. Rothenbacher D, Bode G, Peschke F, Berg G, Adler G, Brenner H. Active infection with *Helicobacter pylori* in an asymptomatic population of middle aged to elderly people. *Epidemiol.Infect.* 1998; 120:297-303.
86. Stroffolini T, Rosmini F, Ferrigno L, Fortini M, D'Amelio R, Matricardi PM. Prevalence of *Helicobacter pylori* infection in a cohort of Italian military students. *Epidemiol.Infect.* 1998; 120:151-155.
87. Kikuchi S., Kurosawa M., Sakiyama T. *Helicobacter pylori* risk associated with sibship size and family history of gastric diseases in Japanese adults. *Jpn.J.Cancer Res.* 1999; 89:1109-1112.
88. Fraser AG, Scragg R, Metcalf P, McCullough S, Yeates NJ. Prevalence of *Helicobacter pylori* infection in different ethnic groups in New Zealand children and adults. *Aust.N.Z.J.Med.* 1996; 26:646-651.

89. Kimura A, Matsubasa T, Kinoshita H, Kuriya N, Yamashita Y, Fujisawa T, et al. *Helicobacter pylori* seropositivity in patients with severe neurologic impairment. *Brain Dev.* 1999; 21:113-117.
90. Bohmer CJ, Klinkenberg-Knol EC, Kuipers EJ, Niezen-de Boer MC, Schreuder H, Schuckink-Kool F, et al. The prevalence of *Helicobacter pylori* infection among inhabitants and healthy employees of institutes for the intellectually disabled. *Am.J.Gastroenterol.* 1997; 92:1000-1004.
91. Harris AW, Douds A, Meurisse EV, Dennis M, Chambers S, Gould SR. Seroprevalence of *Helicobacter pylori* in residents of a hospital for people with severe learning difficulties. *Eur.J.Gastroenterol.Hepatol.* 1995; 7:21-23.
92. Lambert JR, Lin SK, Sievert W, Nicholson L, Schembri M, Guest C. High prevalence of *Helicobacter pylori* antibodies in an institutionalized population: evidence for person-to-person transmission. *Am.J.Gastroenterol.* 1995; 90:2167-2171.
93. Luzzza F, Imeneo M, Maletta M, Paluccio G, Nistico S, Perticone F, et al. Suggestion against an oral-oral route of transmission for *Helicobacter pylori* infection: a seroepidemiological study in a rural area. *Dig.Dis.Sci.* 1998; 43:1488-1492.
94. Us D, Hascelik G. Seroprevalence of *Helicobacter pylori* infection in an Asymptomatic Turkish population. *J.Infect.* 1998; 37:148-150.
95. Ahmad MM, Rahman M, Rumi AK, Islam S, Huq F, Chowdhury MF, et al. Prevalence of *Helicobacter pylori* in asymptomatic population--a pilot serological study in Bangladesh. *J.Epidemiol.* 1997; 7:251-254.
96. Furuta T, Kamata T, Takashima M, Futami H, Arai H, Hanai H, et al. Study of transmission routes of *Helicobacter pylori* in relation to seroprevalence of hepatitis A virus. *J.Clin.Microbiol.* 1997; 35:1891-1893.
97. Rothenbacher D, Bode G, Winz T, Berg G, Adler G, Brenner H. *Helicobacter pylori* in out-patients of a general practitioner: prevalence and determinants of current infection. *Epidemiol.Infect.* 1997; 119:151-157.

98. Malaty HM, Paykov V, Bykova O, Ross A, Graham DP, Anneger JF, et al. *Helicobacter pylori* and socioeconomic factors in Russia. *Helicobacter*. 1996; 1:82-87.
99. Blecker U, Lanciers S, Hauser B, Vandenplas Y. *Helicobacter pylori* positivity in Belgium. *Acta Gastroenterol.Belg*. 1995; 58:31-34.
100. Mendall MA, Goggin PM, Molineaux N, Levy J, Toosy T, Strachan D, et al. Childhood living conditions and *Helicobacter pylori* seropositivity in adult life. *Lancet* 1992; 339:896-897.
101. Polish LB, Douglas JMJ, Davidson AJ, Perez-Perez GI, Blaser MJ. Characterization of risk factors for *Helicobacter pylori* infection among men attending a sexually transmitted disease clinic: lack of evidence for sexual transmission. *J.Clin.Microbiol*. 1991; 29:2139-2143.
102. Pounder RE, Ng D. The prevalence of *Helicobacter pylori* infection in different countries. *Aliment.Pharmacol.Ther*. 1995; 9 Suppl 2:33-9:33-39.
103. Ma JL, You WC, Gail MH, Zhang L, Blot WJ, Chang YS, et al. *Helicobacter pylori* infection and mode of transmission in a population at high risk of stomach cancer. *Int.J.Epidemiol*. 1998; 27:570-573.
104. Roosendaal R, Kuipers EJ, Buitenwerf J, van Uffelen C, Meuwissen SG, van Kamp GJ, et al. *Helicobacter pylori* and the birth cohort effect: evidence of a continuous decrease of infection rates in childhood. *Am.J.Gastroenterol*. 1997; 92:1480-1482.
105. Sipponen P, Kosunen TU, Samloff IM, Heinonen OP, Siurala M. Rate of *Helicobacter pylori* acquisition among Finnish adults: a fifteen year follow-up. *Scand.J.Gastroenterol*. 1996; 31:229-232.
106. Sipponen P. *Helicobacter pylori*: a cohort phenomenon. *Am.J.Surg.Pathol*. 1995; 19 Suppl 1:S30-6:S30-S36
107. Ribeiro VL, Filho JS, Barbosa AJ. Lymphocytic gastritis and *Helicobacter pylori*: a Brazilian survey [letter]. *J.Clin.Pathol*. 1998; 51:83-84.

108. Potasman I, Yitzhak A. *Helicobacter pylori* serostatus in backpackers following travel to tropical countries. *Am.J.Trop.Med.Hyg.* 1998; 58:305-308.
109. Fontham ET, Ruiz B, Perez A, Hunter F, Correa P. Determinants of *Helicobacter pylori* infection and chronic gastritis. *Am.J.Gastroenterol.* 1995; 90:1094-1101.
110. Malaty HM, Graham DY, Wattigney WA, Srinivasan SR, Osato M, Berenson GS. Natural history of *Helicobacter pylori* infection in childhood: 12-year follow-up cohort study in a biracial community. *Clin.Infect.Dis.* 1999; 28:279-282.
111. Cockburn M, Cox B. The effect of measurement error on the determination of *Helicobacter pylori* prevalence. *Epidemiology.* 1997; 8:205-209.
112. Zober A, Schilling D, Ott MG, Schauwecker P, Riemann JF, Messerer P. *Helicobacter pylori* infection: prevalence and clinical relevance in a large company. *J.Occup.Environ.Med.* 1998; 40:586-594.
113. Hamajima N, Inoue M, Tajima K, Tominaga S, Matsuura A, Kobayashi S, et al. Lifestyle and anti-*Helicobacter pylori* immunoglobulin G antibody among outpatients. *Jpn.J.Cancer Res.* 1997; 88:1038-1043.
114. Bateson MC. Cigarette smoking and *Helicobacter pylori* infection. *Postgrad.Med.J.* 1993; 69:41-44.
115. Brenner H, Rothenbacher D, Bode G, Adler G. Relation of smoking and alcohol and coffee consumption to active *Helicobacter pylori* infection: cross sectional study. *BMJ.* 1997; 315:1489-1492.
116. Rudi J, Toppe H, Marx N, Zuna I, Theilmann L, Stremmel W, et al. Risk of infection with *Helicobacter pylori* and hepatitis A virus in different groups of hospital workers. *Am.J.Gastroenterol.* 1997; 92:258-262.
117. Brenner H, Rothenbacher D, Bode G, Adler G. Inverse graded relation between alcohol consumption and active infection with *Helicobacter pylori*. *Am.J.Epidemiol.* 1999; 149:571-576.

118. Brenner H, Berg G, Lappus N, Kliebsch U, Bode G, Boeing H. Alcohol consumption and *Helicobacter pylori* infection: results from the German National Health and Nutrition Survey. *Epidemiology*. 1999; 10:214-218.
119. Jenkins DJ. *Helicobacter pylori* and its interaction with risk factors for chronic disease. *BMJ*. 1997; 315:1481-1482.
120. Klontz KC. Does imbibing alcohol protect against enteric pathogens? *Epidemiology*. 1999; 10:207-209.
121. Goodman KJ, Correa P, Tengana AH, DeLany JP, Collazos T. Nutritional factors and *Helicobacter pylori* infection in Colombian children. *J.Pediatr.Gastroenterol.Nutr*. 1997; 25:507-515.
122. Goodman KJ, Correa P, Tengana AH, Ramirez H, DeLany JP, Guerrero PO, et al. *Helicobacter pylori* infection in the Colombian Andes: a population- based study of transmission pathways. *Am.J.Epidemiol*. 1996; 144:290-299.
123. Jarosz M, Dzieniszewski J, Dabrowska-Ufniaz E, Wartanowicz M, Ziemiński S, Reed PI. Effects of high dose vitamin C treatment on *Helicobacter pylori* infection and total vitamin C concentration in gastric juice. *Eur.J.Cancer Prev*. 1998; 7:449-454.
124. Malaty HM, Graham DY, Isaksson I, Engstrand L, Pedersen NL. Co-twin study of the effect of environment and dietary elements on acquisition of *Helicobacter pylori* infection. *Am.J.Epidemiol*. 1998; 148:793-797.
125. Hopkins RJ, Vial PA, Ferreccio C, Ovalle J, Prado P, Sotomayor V, et al. Seroprevalence of *Helicobacter pylori* in Chile: vegetables may serve as one route of transmission. *J.Infect.Dis*. 1993; 168:222-226.
126. Grubel P, Huang L, Masubuchi N, Stutzenberger FJ, Cave DR. Detection of *Helicobacter pylori* DNA in houseflies (*Musca domestica*) on three continents [letter]. *Lancet* 1998; 352:788-789.
127. Begue RE, Gonzales JL, Correa-Gracian H, Tang SC. Dietary risk factors associated with the transmission of *Helicobacter pylori* in Lima, Peru. *Am.J.Trop.Med.Hyg*. 1998; 59:637-640.

128. Friis L, Engstrand L, Edling C. Prevalence of *Helicobacter pylori* infection among sewage workers. *Scand.J.Work.Environ.Health* 1996; 22:364-368.
129. Shelley KH, Haddadin AS. Is *Helicobacter pylori* infection an occupational hazard for anesthesiologists? *Anesth.Analg.* 1998; 87:973-974.
130. Lin SK, Lambert JR, Schembri MA, Nicholson L, Johnson IH. The prevalence of *Helicobacter pylori* in practising dental staff and dental students. *Aust.Dent.J.* 1998; 43:35-39.
131. Banatvala N, Abdi Y, Clements L, Herbert AM, Davies J, Bagg J, et al. *Helicobacter pylori* infection in dentists--a case-control study. *Scand.J.Infect.Dis.* 1995; 27:149-151.
132. Malaty HM, Evans DJJ, Abramovitch K, Evans DG, Graham DY. *Helicobacter pylori* infection in dental workers: a seroepidemiology study. *Am.J.Gastroenterol.* 1992; 87:1728-1731.
133. Potts LF, Lewis SJ, Mountford RA. Prevalence of *Helicobacter pylori* in respiratory physicians performing bronchoscopy: a comparison with gastroenterologists using the carbon 13 urea breath test. *Helicobacter.* 1997; 2:152-154.
134. Goh KL, Parasakthi N, Ong KK. Prevalence of *Helicobacter pylori* infection in endoscopy and non- endoscopy personnel: results of field survey with serology and 14C-urea breath test. *Am.J.Gastroenterol.* 1996; 91:268-270.
135. Liu WZ, Xiao SD, Jiang SJ, Li RR, Pang ZJ. Seroprevalence of *Helicobacter pylori* infection in medical staff in Shanghai. *Scand.J.Gastroenterol.* 1996; 31:749-752.
136. Su YC, Wang WM, Chen LT, Chiang W, Chen CY, Lu SN, et al. High seroprevalence of IgG against *Helicobacter pylori* among endoscopists in Taiwan. *Dig.Dis.Sci.* 1996; 41:1571-1576.
137. Chong J, Marshall BJ, Barkin JS, McCallum RW, Reiner DK, Hoffman SR, et al. Occupational exposure to *Helicobacter pylori* for the endoscopy professional: a sera epidemiological study. *Am.J.Gastroenterol.* 1994; 89:1987-1992.

138. Lin SK, Lambert JR, Schembri MA, Nicholson L, Korman MG. *Helicobacter pylori* prevalence in endoscopy and medical staff. *J.Gastroenterol.Hepatol.* 1994; 9:319-324.
139. Mitchell HM, Lee A, Carrick J. Increased incidence of *Campylobacter pylori* infection in gastroenterologists: further evidence to support person-to-person transmission of *C. pylori*. *Scand.J.Gastroenterol.* 1989; 24:396-400.
140. Mones J, Martin-de-Argila C, Samitier RS, Gisbert JP, Sainz S, Boixeda D. Prevalence of *Helicobacter pylori* infection in medical professionals in Spain. *Eur.J.Gastroenterol.Hepatol.* 1999; 11:239-242.
141. Braden B, Duan LP, Caspary WF, Lembcke B. Endoscopy is not a risk factor for *Helicobacter pylori* infection--but medical practice is. *Gastrointest.Endosc.* 1997; 46:305-310.
142. Nishikawa J, Kawai H, Takahashi A, Seki T, Yoshikawa N, Akita Y, et al. Seroprevalence of immunoglobulin G antibodies against *Helicobacter pylori* among endoscopy personnel in Japan. *Gastrointest.Endosc.* 1998; 48:237-243.
143. Williams CL. *Helicobacter pylori* and endoscopy. *J.Hosp.Infect.* 1999; 41:263-268.
144. Klein PD, Graham DY, Gaillour A, Opekun AR, Smith EO. Water source as risk factor for *Helicobacter pylori* infection in Peruvian children. Gastrointestinal Physiology Working Group. *Lancet* 1991; 337:1503-1506.
145. Elitsur Y, Short JP, Neace C. Prevalence of *Helicobacter pylori* infection in children from urban and rural West Virginia. *Dig.Dis.Sci.* 1998; 43:773-778.
146. Bode G, Rothenbacher D, Brenner H, Adler G. Pets are not a risk factor for *Helicobacter pylori* infection in young children: results of a population-based study in Southern Germany. *Pediatr.Infect.Dis.J.* 1998; 17:909-912.
147. Staat MA, Kruszon-Moran D, McQuillan GM, Kaslow RA. A population-based serologic survey of *Helicobacter pylori* infection in children and

- adolescents in the United States. *J.Infect.Dis.* 1996; 174:1120-1123.
148. McIsaac WJ, Leung GM. Peptic ulcer disease and exposure to domestic pets. *Am.J.Public Health* 1999; 89:81-84.
 149. Dore MP, Bilotta M, Vaira D, Manca A, Massarelli G, Leandro G, et al. High prevalence of *Helicobacter pylori* infection in shepherds. *Dig.Dis.Sci.* 1999; 44:1161-1164.
 150. Czekwianianc E, Bak-Romaniszyn L, Malecka-Panas E, Suski S, Woch G. Prevalence of *Helicobacter pylori* in children dependently on age and living conditions. *J.Physiol.Pharmacol.* 1996; 47:203-207.
 151. Megraud F. Transmission of *Helicobacter pylori*: faecal-oral versus oral-oral route. *Aliment.Pharmacol.Ther.* 1995; 9 Suppl 2:85-91:85-91.
 152. Hammermeister I, Janus G, Schamarowski F, Rudolf M, Jacobs E, Kist M. Elevated risk of *Helicobacter pylori* infection in submarine crews. *Eur.J.Clin.Microbiol.Infect.Dis.* 1992; 11:9-14.
 153. McCallion WA, Murray LJ, Bailie AG, Dalzell AM, O'Reilly DP, Bamford KB. *Helicobacter pylori* infection in children: relation with current household living conditions. *Gut* 1996; 39:18-21.
 154. Clemens J, Albert MJ, Rao M, Huda S, Qadri F, Van Loon FP, et al. Sociodemographic, hygienic and nutritional correlates of *Helicobacter pylori* infection of young Bangladeshi children. *Pediatr.Infect.Dis.J.* 1996; 15:1113-1118.
 155. Brenner H, Rothenbacher D, Bode G, Adler G. Parental history of gastric or duodenal ulcer and prevalence of *Helicobacter pylori* infection in preschool children: population based study. *BMJ.* 1998; 316:665
 156. De Hert M, Hautekeete M, De Wilde D, Peuskens J. High prevalence of *Helicobacter pylori* in institutionalized schizophrenic patients [letter]. *Schizophr.Res.* 1997; 26:243-244.
 157. Lewindon PJ, Lau D, Chan A, Tse P, Sullivan PB. *Helicobacter pylori* in an

- institution for disabled children in Hong Kong. *Dev.Med.Child Neurol.* 1997; 39:682-685.
158. Bonamico M, Monti S, Luzzi I, Magliocca FM, Cipolletta E, Calvani L, et al. *Helicobacter pylori* infection in families of *Helicobacter pylori*- positive children. *Ital.J.Gastroenterol.* 1996; 28:512-517.
 159. Blecker U, Lanciers S, Mehta DI, Vandenplas Y. Familial clustering of *Helicobacter pylori* infection. *Clin.Pediatr.(Phila.)* 1994; 33:307-308.
 160. Mitchell HM, Bohane T, Hawkes RA, Lee A. *Helicobacter pylori* infection within families. *Int.J.Med.Microbiol.Virol.Parasitol.Infect.Dis.* 1993; 280:128-136.
 161. Drumm B, Perez-Perez GI, Blaser MJ, Sherman PM. Intrafamilial clustering of *Helicobacter pylori* infection. *N.Engl.J.Med.* 1990; 322:359-363.
 162. Eckert MW, McKnight CA, Lee JA, Araya J, Correa P, Cohn IJ, et al. Early gastric cancer and *Helicobacter pylori*: 34 years of experience at Charity Hospital in New Orleans. *Am.Surg.* 1998; 64:545-550.
 163. Rothenbacher D, Bode G, Berg G, Knayer U, Gonser T, Adler G, et al. *Helicobacter pylori* among preschool children and their parents: evidence of parent-child transmission. *J.Infect.Dis.* 1999; 179:398-402.
 164. Singh V, Trikha B, Vaiphei K, Nain CK, Thennarasu K, Singh K. *Helicobacter pylori*: evidence for spouse-to-spouse transmission. *J.Gastroenterol.Hepatol.* 1999; 14:519-522.
 165. Brenner H, Rothenbacher D, Bode G, Dieudonne P, Adler G. Active infection with *Helicobacter pylori* in healthy couples. *Epidemiol.Infect.* 1999; 122:91-95.
 166. Georgopoulos SD, Mentis AF, Spiliadis CA, Tzouveleakis LS, Tzelepi E, Moshopoulos A, et al. *Helicobacter pylori* infection in spouses of patients with duodenal ulcers and comparison of ribosomal RNA gene patterns. *Gut* 1996; 39:634-638.

167. Parente F, Maconi G, Sangaletti O, Minguzzi M, Vago L, Rossi E, et al. Prevalence of *Helicobacter pylori* infection and related gastroduodenal lesions in spouses of *Helicobacter pylori* positive patients with duodenal ulcer. *Gut* 1996; 39:629-633.
168. Schutze K, Hentschel E, Dragosics B, Hirschl AM. *Helicobacter pylori* reinfection with identical organisms: transmission by the patients' spouses. *Gut* 1995; 36:831-833.
169. Mitchell HM, Hazell SL, Kolesnikow T, Mitchell J, Frommer D. Antigen recognition during progression from acute to chronic infection with a *cagA*-positive strain of *Helicobacter pylori*. *Infect.Immun.* 1996; 64:1166-1172.
170. Chalkauskas H, Kersulyte D, Cepulienė I, Urbonas V, Ruzeviciene D, Barauskiene A, et al. Genotypes of *Helicobacter pylori* in Lithuanian families. *Helicobacter.* 1998; 3:296-302.
171. Nguyen AM, el-Zaatari FA, Graham DY. *Helicobacter pylori* in the oral cavity. A critical review of the literature. *Oral Surg.Oral Med.Oral Pathol.Oral Radiol.Endod.* 1995; 79:705-709.
172. Madinier IM, Fosse TM, Monteil RA. Oral carriage of *Helicobacter pylori*: a review. *J.Periodontol.* 1997; 68:2-6.
173. Pustorino R, Nicosia R, D'Ambra G, Di Paola M, Brugnoletti O, Grippaudo G, et al. The mouth-stomach crossing of *Helicobacter pylori*. *Riv.Eur.Sci.Med.Farmacol.* 1996; 18:183-186.
174. Pytko-Polonczyk J, Konturek SJ, Karczewska E, Bielanski W, Kaczmarczyk-Stachowska A. Oral cavity as permanent reservoir of *Helicobacter pylori* and potential source of reinfection. *J.Physiol.Pharmacol.* 1996; 47:121-129.
175. Thomas E, Jiang C, Chi DS, Li C, Ferguson DAJ. The role of the oral cavity in *Helicobacter pylori* infection. *Am.J.Gastroenterol.* 1997; 92:2148-2154.
176. Leung WK, Sung JJ, Ling TK, Siu KL, Cheng AF. Use of chopsticks for eating and *Helicobacter pylori* infection. *Dig.Dis.Sci.* 1999; 44:1173-1176.

177. Li C, Musich PR, Ha T, Ferguson DAJ, Patel NR, Chi DS, et al. High prevalence of *Helicobacter pylori* in saliva demonstrated by a novel PCR assay. *J.Clin.Pathol.* 1995; 48:662-666.
178. Li C, Ha T, Ferguson DAJ, Chi DS, Zhao R, Patel NR, et al. A newly developed PCR assay of *H. pylori* in gastric biopsy, saliva, and feces. Evidence of high prevalence of *H. pylori* in saliva supports oral transmission. *Dig.Dis.Sci.* 1996; 41:2142-2149.
179. Namavar F, Roosendaal R, Kuipers EJ, de Groot P, van der Bijl MW, Pena AS, et al. Presence of *Helicobacter pylori* in the oral cavity, oesophagus, stomach and faeces of patients with gastritis. *Eur.J.Clin.Microbiol.Infect.Dis.* 1995; 14:234-237.
180. Shimada T, Ogura K, Ota S, Terano A, Takahashi M, Hamada E, et al. Identification of *Helicobacter pylori* in gastric specimens, gastric juice, saliva, and faeces of Japanese patients [letter]. *Lancet* 1994; 343:1636-1637.
181. Mapstone NP, Lynch DA, Lewis FA, Axon AT, Tompkins DS, Dixon MF, et al. Identification of *Helicobacter pylori* DNA in the mouths and stomachs of patients with gastritis using PCR. *J.Clin.Pathol.* 1993; 46:540-543.
182. Ferguson DAJ, Li C, Patel NR, Mayberry WR, Chi DS, Thomas E. Isolation of *Helicobacter pylori* from saliva. *J.Clin.Microbiol.* 1993; 31:2802-2804.
183. Bhatia SJ, Abraham P. *Helicobacter pylori* in the Indian environment. *Indian J.Gastroenterol.* 1995; 14:139-144.
184. Leung WK, Sung JJ, Ling TK, Siu KL, Cheng AF. Does the use of chopsticks for eating transmit *Helicobacter pylori*? [letter]. *Lancet* 1997; 350:31
185. Oshowo A, Gillam D, Botha A, Tunio M, Holton J, Boulos P, et al. *Helicobacter pylori*: the mouth, stomach, and gut axis. *Ann.Periodontol.* 1998; 3:276-280.
186. Cheng LH, Webberley M, Evans M, Hanson N, Brown R. *Helicobacter pylori* in dental plaque and gastric mucosa. *Oral Surg.Oral Med.Oral*

Pathol.Oral Radiol.Endod. 1996; 81:421-423.

187. Cellini L, Allocati N, Piattelli A, Petrelli I, Fanci P, Dainelli B. Microbiological evidence of *Helicobacter pylori* from dental plaque in dyspeptic patients. *New Microbiol.* 1995; 18:187-192.
188. Hardo PG, Tugnait A, Hassan F, Lynch DA, West AP, Mapstone NP, et al. *Helicobacter pylori* infection and dental care. *Gut* 1995; 37:44-46.
189. Luman W, Alkout AM, Blackwell CC, Weir DM, Plamer KR. *Helicobacter pylori* in the mouth--negative isolation from dental plaque and saliva. *Eur.J.Gastroenterol.Hepatol.* 1996; 8:11-14.
190. Kamat AH, Mehta PR, Natu AA, Phadke AY, Vora IM, Desai PD, et al. Dental plaque: an unlikely reservoir of *Helicobacter pylori*. *Indian J.Gastroenterol.* 1998; 17:138-140.
191. Savoldi E, Marinone MG, Negrini R, Facchinetti D, Lanzini A, Sapelli PL. Absence of *Helicobacter pylori* in dental plaque determined by immunoperoxidase. *Helicobacter.* 1998; 3:283-287.
192. Dore-Davin C, Heitz M, Yang H, Herranz M, Blum AL, Cortes-Theulaz I. *Helicobacter pylori* in the oral cavity reflects handling of contaminants but not gastric infection. *Digestion* 1999; 60:196-202.
193. Axon AT. Review article: is *Helicobacter pylori* transmitted by the gastro-oral route? *Aliment.Pharmacol.Ther.* 1995; 9:585-588.
194. Katoh M, Saito D, Noda T, Yoshida S, Oguro Y, Yazaki Y, et al. *Helicobacter pylori* may be transmitted through gastrofiberscope even after manual Hyamine washing. *Jpn.J.Cancer Res.* 1993; 84:117-119.
195. Figura N. Mouth-to-mouth resuscitation and *Helicobacter pylori* infection [letter]. *Lancet* 1996; 347:1342
196. Gramley WA, Asghar A, Frierson HFJ, Powell SM. Detection of *Helicobacter pylori* DNA in fecal samples from infected individuals. *J.Clin.Microbiol.* 1999; 37:2236-2240.

197. van Zwet AA, Thijs JC, Kooistra-Smid AM, Schirm J, Snijder JA. Use of PCR with feces for detection of *Helicobacter pylori* infections in patients. *J.Clin.Microbiol.* 1994; 32:1346-1348.
198. Thomas JE, Gibson GR, Darboe MK, Dale A, Weaver LT. Isolation of *Helicobacter pylori* from human faeces. *Lancet* 1992; 340:1194-1195.
199. Kelly SM, Pitcher MC, Farmery SM, Gibson GR. Isolation of *Helicobacter pylori* from feces of patients with dyspepsia in the United Kingdom. *Gastroenterology* 1994; 107:1671-1674.
200. Sahay P, Axon AT. Reservoirs of *Helicobacter pylori* and modes of transmission. *Helicobacter.* 1996; 1:175-182.
201. Hazell SL, Mitchell HM, Hedges M, Shi X, Hu PJ, Li YY, et al. Hepatitis A and evidence against the community dissemination of *Helicobacter pylori* via feces. *J.Infect.Dis.* 1994; 170:686-689.
202. Webb PM, Knight T, Newell DG, Elder JB, Forman D. *Helicobacter pylori* transmission: evidence from a comparison with hepatitis A virus. *Eur.J.Gastroenterol.Hepatol.* 1996; 8:439-441.
203. Bode G, Mauch F, Malfertheiner P. The coccoid forms of *Helicobacter pylori*. Criteria for their viability. *Epidemiol.Infect.* 1993; 111:483-490.
204. Hulten K, Han SW, Enroth H, Klein PD, Opekun AR, Gilman RH, et al. *Helicobacter pylori* in the drinking water in Peru. *Gastroenterology* 1996; 110:1031-1035.
205. Fan XG, Chua A, Li TG, Zeng QS. Survival of *Helicobacter pylori* in milk and tap water. *J.Gastroenterol.Hepatol.* 1998; 13:1096-1098.
206. Vincent P. Transmission and acquisition of *Helicobacter pylori* infection: evidences and hypothesis. *Biomed.Pharmacother.* 1995; 49:11-18.
207. Sorberg M, Nilsson M, Hanberger H, Nilsson LE. Morphologic conversion of *Helicobacter pylori* from bacillary to coccoid form. *Eur.J.Clin.Microbiol.Infect.Dis.* 1996; 15:216-219.

208. Owen RJ. Microbiological aspects of *Helicobacter pylori* infection. *Commun.Dis.Rep.CDR.Rev.* 1993; 3:R51-R56
209. Cellini L, Allocati N, Angelucci D, Iezzi T, Di Campi E, Marzio L, et al. Coccoid *Helicobacter pylori* not culturable in vitro reverts in mice. *Microbiol.Immunol.* 1994; 38:843-850.
210. Eaton KA, Catrenich CE, Makin KM, Krakowka S. Virulence of coccoid and bacillary forms of *Helicobacter pylori* in gnotobiotic piglets. *J.Infect.Dis.* 1995; 171:459-462.
211. Westblom TU. Molecular diagnosis of *Helicobacter pylori*. *Immunol.Invest.* 1997; 26:163-174.
212. Baker KH, Hegarty JP. Presence of *Helicobacter pylori* in Drinking Water is Associated with Clinical Infection. *Gastroenterology* 1999;
213. Hegarty JP, Dowd MT, Baker KH. Occurrence of *Helicobacter pylori* in Surface Water in the United States. *Applied Micro.* 1999;
214. Fox JG. Non-human reservoirs of *Helicobacter pylori*. *Aliment.Pharmacol.Ther.* 1995; 9 Suppl 2:93-103:93-103.
215. Handt LK, Fox JG, Dewhirst FE, Fraser GJ, Paster BJ, Yan LL, et al. *Helicobacter pylori* isolated from the domestic cat: public health implications. *Infect.Immun.* 1994; 62:2367-2374.
216. Handt LK, Fox JG, Stalis IH, Rufo R, Lee G, Linn J, et al. Characterization of feline *Helicobacter pylori* strains and associated gastritis in a colony of domestic cats. *J.Clin.Microbiol.* 1995; 33:2280-2289.
217. Fox JG, Batchelder M, Marini R, Yan L, Handt L, Li X, et al. *Helicobacter pylori*-induced gastritis in the domestic cat. *Infect.Immun.* 1995; 63:2674-2681.
218. Fox JG, Perkins S, Yan L, Shen Z, Attardo L, Pappo J. Local immune response in *Helicobacter pylori*-infected cats and identification of *H. pylori* in saliva, gastric fluid and faeces. *Immunology* 1996; 88:400-406.

219. el-Zaatari FA, Woo JS, Badr A, Osato MS, Serna H, Lichtenberger LM, et al. Failure to isolate *Helicobacter pylori* from stray cats indicates that *H. pylori* in cats may be an anthroponosis—an animal infection with a human pathogen. *J.Med.Microbiol.* 1997; 46:372-376.
220. Webb PM, Knight T, Elder JB, Newell DG, Forman D. Is *Helicobacter pylori* transmitted from cats to humans? *Helicobacter.* 1996; 1:79-81.
221. Dore MP, Sepulveda AR, Osato MS, Realdi G, Graham DY. *Helicobacter pylori* in sheep milk [letter]. *Lancet* 1999; 354:132
222. Grubel P, Hoffman JS, Chong FK, Burstein NA, Mepani C, Cave DR. Vector potential of houseflies (*Musca domestica*) for *Helicobacter pylori*. *J.Clin.Microbiol.* 1997; 35:1300-1303.
223. Vaira D, Holton J. Vector potential of houseflies (*Musca domestica*) for *Helicobacter pylori*. *Helicobacter.* 1998; 3:65-66.
224. Osato MS, Ayub K, Le HH, Reddy R, Graham DY. Houseflies are an unlikely reservoir or vector for *Helicobacter pylori*. *J.Clin.Microbiol.* 1998; 36:2786-2788.
225. Axon AT. Disinfection of endoscopic equipment. *Baillieres.Clin.Gastroenterol.* 1991; 5:61-77.
226. Fantry GT, Zheng QX, James SP. Conventional cleaning and disinfection techniques eliminate the risk of endoscopic transmission of *Helicobacter pylori*. *Am.J.Gastroenterol.* 1995; 90:227-232.
227. Tytgat GN. Endoscopic transmission of *Helicobacter pylori*. *Aliment.Pharmacol.Ther.* 1995; 9 Suppl 2:105-10:105-110.
228. Langenberg W, Rauws EA, Oudbier JH, Tytgat GN. Patient-to-patient transmission of *Campylobacter pylori* infection by fiberoptic gastroduodenoscopy and biopsy. *J.Infect.Dis.* 1990; 161:507-511.
229. Ramsey EJ, Carey KV, Peterson WL, Jackson JJ, Murphy FK, Read NW, et al. Epidemic gastritis with hypochlorhydria. *Gastroenterology* 1979;

76:1449-1457.

230. Graham DY, Alpert LC, Smith JL, Yoshimura HH. Iatrogenic *Campylobacter pylori* infection is a cause of epidemic achlorhydria. *Am.J.Gastroenterol.* 1988; 83:974-980.
231. Shimada T, Terano A, Ota S, Takikawa H, Sumino S. Risk of iatrogenic transmission of *Helicobacter pylori* by gastroscopes [letter]. *Lancet* 1996; 347:1342-1343.
232. Axon AT. Disinfection and endoscopy: summary and recommendations. Working party report to the World Congresses of Gastroenterology, Sydney 1990. *J.Gastroenterol.Hepatol.* 1991; 6:23-24.
233. Rohr MR, Castro R, Morais M, Brant CQ, Castelo FA, Ferrari JA. Risk of *Helicobacter pylori* transmission by upper gastrointestinal endoscopy. *Am.J.Infect.Control.* 1998; 26:12-15.
234. Kaneko H, Mitsuma T, Kotera H, Uchida K, Furusawa A, Morise K. Are routine cleaning methods sufficient to remove *Helicobacter pylori* from endoscopic equipment? [letter]. *Endoscopy.* 1993; 25:435
235. Wu MS, Wang JT, Yang JC, Wang HH, Sheu JC, Chen DS, et al. Effective reduction of *Helicobacter pylori* infection after upper gastrointestinal endoscopy by mechanical washing of the endoscope. *Hepatogastroenterology.* 1996; 43:1660-1664.
236. Akamatsu T, Tabata K, Hironga M, Kawakami H, Uyeda M. Transmission of *Helicobacter pylori* infection via flexible fiberoptic endoscopy. *Am.J.Infect.Control.* 1996; 24:396-401.
237. Mitchell HM. The epidemiology of *Helicobacter pylori*. *Curr.Top.Microbiol.Immunol.* 1999; 241:11-30:11-30.
238. Gao CM, Takezaki T, Ding JH, Li MS, Tajima K. Protective effect of allium vegetables against both esophageal and stomach cancer: a simultaneous case-referent study of a high-epidemic area in Jiangsu Province, China. *Jpn.J.Cancer Res.* 1999; 90:614-621.

239. You WC, Blot WJ, Chang YS, Ershow A, Yang ZT, An Q, et al. *Allium* vegetables and reduced risk of stomach cancer. *J.Natl.Cancer Inst.* 1989; 81:162-164.
240. Sivam GP, Lampe JW, Ulness B, Swanzy SR, Potter JD. *Helicobacter pylori*—in vitro susceptibility to garlic (*Allium sativum*) extract. *Nutr.Cancer* 1997; 27:118-121.
241. Cellini L, Di Campli E, Masulli M, Di Bartolomeo S, Allocati N. Inhibition of *Helicobacter pylori* by garlic extract (*Allium sativum*). *FEMS Immunol.Med.Microbiol.* 1996; 13:273-277.
242. Graham DY, Anderson SY, Lang T. Garlic or jalapeno peppers for treatment of *Helicobacter pylori* infection. *Am.J.Gastroenterol.* 1999; 94:1200-1202.
243. Edwards CN, Douglin CP, Prussia PR, Garriques SA, Levett PN. Epidemiology of *Helicobacter pylori* infection in Barbados. *West.Indian Med.J.* 1997; 46:3-7.
244. Schwartz S, Carpenter KM. The right answer for the wrong question: consequences of type III error for public health research. *Am. J. Public Health* 1999; 89:1175-1180.
245. Wacholder S. When measurement errors correlate with truth: surprising effects of nondifferential misclassification. *Epidemiology.* 1995; 6:157-161.
246. Webberley MJ, Webberley JM, Newell DG, Lowe P, Melikian V. Seroepidemiology of *Helicobacter pylori* infection in vegans and meat-eaters. *Epidemiol.Infect.* 1992; 108:457-462.
247. Malaty HM, Engstrand L, Pedersen NL, Graham DY. *Helicobacter pylori* infection: genetic and environmental influences. A study of twins. *Ann.Intern.Med.* 1994; 120:982-986.
248. Bamford KB, Bickley J, Collins JS, Johnston BT, Potts S, Boston V, et al. *Helicobacter pylori*: comparison of DNA fingerprints provides evidence for intrafamilial infection. *Gut* 1993; 34:1348-1350.