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Genetic Aspects of Helicobacter pylori and Campylobacter jejuni

by

Koji Hiratsuka

A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE OF DOCTOR OF PHILOSOPHY

DEPARTMENT OF MEDICAL MICROBIOLOGY AND IMMUNOLOGY

EDMONTON, ALBERTA

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FACULTY OF GRADUATE STUDIES AND RESEARCH

The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research for acceptance, a thesis entitled **Genetic Aspects of Helicobacter pylori** and **Campylobacter jejuni** submitted by Koji Hiratsuka in partial fulfillment of the requirements for the degree of Doctor of Philosophy

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This thesis is dedicated to Ursula, my partner in life.

Your dedication to our family, the sacrifices you have made

and your love have made this thesis possible.

Abstract

Genomic libraries of Helicobacter pylori UA802 were constructed, using the lambda cloning vector λ -DASH II. From these libraries, ribcsomal RNA (rRNA) genes, and genes encoding proteins involved in copper transport (hpCopA and hpCopP) were isolated. The DNA sequences of the 23S and 5S rRNA genes were determined, and used for phylogenetic classification of H. pylori UA802. The phylogenetic studies showed that H. pylori UA802 is closely related to Campylobacter spp., but belongs in its own distinct genus.

Pulsed-field gel electrophoresis (PFGE) analyses were performed on *H. pylori* UA802. From these data a genomic restriction map was constructed with the enzymes *NruI*, *NotI*, and *SfiI*. A number of genes were localized on the map by hybridization to Southern blots of PFGE gels.

In 1990 two Campylobacter-specific DNA probes, pDT1719 and pDT1720, were developed. For this research, the *C. jejuni* UA580 DNA inserts of these two probes were sequenced. Two partial open reading frames (ORF), ORF1 and ORF2, were found in pDT1719 and one partial ORF, ORF3, was found in pDT1720. Computer homology searches were conducted for both the nucleotide and predicted amino acid sequences of the three ORFs. No sequences homologous to ORF3 could be found. The predicted amino acid sequence of ORF1 was found to have homology with the HtrA proteins of *S. typhimurium* and *E. coli*. The predicted amino acid sequence of ORF2 had

homology with several proteins which have been associated with the environmental response elements of two-component regulatory systems. ORF1 and ORF2 appeared to be linked together in an operon and may represent a two-component regulatory system of *C. jejuni* which responds to heat shock.

The N-terminal sequence of the first thirty amino acids of the major outer membrane protein (MOMP) of *C. jejuni* UA580, which has been shown to be a porin, was obtained. This amino acid sequence was found to have 59% identity with the amino acid sequence of two membrane proteins of *Campylobacter rectus*, one of which has also been shown to be a porin.

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List of Abbreviations

aa amino acid

BHI Brain Heart Infusion

BMC Boerhinger Mannheim Canada

bp base pair

Brinkman Brinkman Instruments (Canada) Ltd.

BRL Gibco/BRL Life Technologies Inc.

BSA bovine serum albumin

3-[cyclohexylamino]-1-propanesulfonic

CAPS acid

CFU colony forming units

Da Daltons

DEPC diethyl pyrocarbonate

dH₂O Distilled water

Difco Difco Laboratories

DIG digoxygenin

DNase deoxyribonuclease

EDTA ethylenediamine tetra acetic acid

ETOH ethanol

GCG Genetics Computer Group

GM gastric metaplasia

hr hour

ICN ICN Biochemicals Canada Ltd.

kb kilobase

kDa kilodalton

LB Luria broth

LBM Luria broth + 0.5% maltose + 10mM MgSO₄

LMP low melting point

MeOH methanol

min minute

MOMP major outer membrane protein

MOPS morpholinopropanesulfonic acid

NaOAc sodium acetate

NT nick translation

nt nucleotide

OD optical density

ORF open reading frame

PAGE polyacrylamide gel electrophoresis

PCR Polymerase chain reaction

PEG polyethylene glycol

price pulsed-field gel electrophoresis

PFU plaque forming units

Pharmacia Pharmacia Biotech Inc.

PMSF phenylmethylsulfonyl fluoride

RNase ribonuclease

rRNA ribosomal RNA

r·DNA ribosomal DNA

s second

SDS sodium dodecyl sulfate

Sigma Chemical Company

ssc salt/sodium citrate

TAFE Transverse Alternating Field

Electrophoresis

TB Tris borate

10 mM Tris·HCl pH 7.5 1 mM EDTA

TE

 ${\tt N,N,N',N'-Tetra-methylethylenediamine}$ TEMED

upstream activation region UAR

United States Biochemica USB

Chapter 1

Introduction

A. Helicobacter pylori

Studies of the bacterium Helicobacter pylori have changed our view of human gastric disease. The human stomach has not normally been considered a hospitable environment for bacterial growth, so suggestions that H. pylori colonized the stomach and caused gastroduodenal disease were initially met with a great deal of skepticism (Marshall, 1983; Warren, 1983). Research over the past several years has gradually removed this doubt, and H. pylori has now become widely accepted as an etiological agent for chronic type B gastritis, characterized by gastric inflammation (Marshall, et al., 1985; Goodwin et al., 1986; Cover & Blaser, 1992a). H. pylori infection is also believed to play a role in both peptic and duodenal ulceration (Goodwin et al., 1986, Dixon, 1991). In addition, there is evidence that chronic infection with H. pylori increases the risk of gastric adenocarcinoma (Hebbel, 1940; Correa et al., 1990; Forman et al., 1990; Talley et al., 1991; Parsonnet et al., 1991) and gastric lymphoma (Parsonnet et al., 1994).

A.1. The history of H. pylori

Although spiral Gram-negative organisms had been observed in human gastric biopsies as early as 1874, the involvement of these organisms in the pathogenesis of gastroduodenal

disease was unclear. Culturing these organisms in vitro proved difficult. Thus, aside from microscopic observations, In 1983 the organism that very little was known about them. we now call Helicobacter pylori was first cultured from human gastric biopsy specimens in Perth, Australia (Marshall, 1983; Initially the name Campylobacter pyloridis Warren, 1983). was chosen because of physiological and morphological similarities of C. pyloridis with organisms from the genus Campylobacter (Marshall et al., 1987). The name was later changed to C. pylori (Marshall and Goodwin, 1987). However, even early observations of flagellar morphology suggested that there were obvious differences which set this organism apart from Campylobacter spp. (Marshall and Warren, 1984). Later studies of fatty acid profiles, ultrastructure (Goodwin et al., 1987) and ribosomal RNA (Romaniuk et al., 1987) suggested that the organism designated C. pylori was not a true Campylobacter and that the creation of a new genus was In 1989 the genus Helicobacter was created and in order. H. pylori was included in this genus along with the animal pathogen H. mustelae (Goodwin et al., 1989). Currently there are twelve species in the genus Helicobacter (Table 1-1), but with improved methods for the isolation and cultivation of organisms, this number could increase.

A.2. Microbiology and pathogenesis of H. pylori

A number of reviews have been published describing the morphological and physiological characteristics of *H. pylori*

Table 1-1. Members of the genus Helicobactera

		,
Species	Host	Source
H. pylori	Human, monkey, cat	Stomach
H. nemestrinae	Macaque monkey	Stomach
H. acinonyx	Cheetah	Stomach
H. heilmannii	Mammals	Stomach
H. felis	Cat, dog	Stomach
H. rappini	Mammals	Stomach, intestine
H. mustelae	Ferret	Stomach
H. muridarum	Rat, mouse	Stomach, intestine
H. hepaticus	Mouse	Liver
H. canis	Dog	Intestine
H. fenneliae	Human	Intestine
H. cinaedi	Human	Intestine

a- From Cover and Blaser, 1995

(Dick, 1990; Good and Armstrong, 1990; Goodwin and Worsley, 1993). *H. pylori* is a Gram-negative organism which exists as either a curved rod when grown on solid media or as a spiral rod in liquid media or in vivo. It is highly motile, with four to six polar, sheathed flagella. It is a microaerophilic organism and, upon prolonged exposure to oxygen, will assume a viable but non-culturable coccoid form. *H. pylori* has a low G + C content calculated to be 35.2 mol% (Béji et al., 1988). The genome size of *H. pylori* has been estimated to be approximately 1700 kb (Taylor et al., 1992a; Bukanov and Berg, 1994; Jiang, 1994).

Enzymology

A variety of enzymes are produced by H. pylori, including large amounts of urease (Owen et al., 1986; Mobley et al., 1988), superoxide dismutase and catalase (Lior & Johnson, 1985), as well as proteases (Slomiany et al., 1987; Baxter et al., 1989), lipase (Sarosiek et al., 1988), DNase, and oxidase. Urease is of particular interest, as the production of this extracellular enzyme distinguishes H. pylori from other spiral organisms. Urease activity has been used as the basis of diagnostic tests to detect the presence of H. pylori (McNulty & Wise, 1985; Marshall & Surveyor, 1988). In addition, numerous studies have indicated that urease is one of the important factors in colonization of the stomach by H. pylori (Hazell and Lee, 1986; Marshall. et al., 1990; Eaton et al., 1991; Triebling et al., 1991; Segal et al., 1992; Mégraud et al., 1992). It has been hypothesized that urease

activity produces a cloud of ammonia protecting the bacterium from the acidic environment of the stomach long enough for the bacterium to penetrate the mucosal lining. The two structural genes ureA and ureB encode the urease subunits and were among the first genes from H. pylori to be characterized (Clayton et al., 1989a, 1989b; Labigne et al., 1991; Cussac et al., 1992). To date nine genes are thought to be involved in the expression of urease activity in H. pylori (Cover and Blaser, 1995).

Flagella

The flagella have also been implicated in the pathogenesis of *H. pylori*. Four to six uni-polar flagella are normally present, although bipolar flagella have been observed under scanning electron microscopy. The sheath sometimes extends to form a terminal bulb at the end of each flagellum. The flagella confer motility to the organism and, combined with the spiral shape of the organism, are thought to aid in the penetration of the mucous layer and the colonization of the stomach lining (Eaton et al., 1989). *H. pylori* has two flagellin genes, designated flam (Leying et al., 1991) and flam (Suerbaum et al., 1993). Studies have shown that flam, but not flam, is required to assemble functional flagella (Haas et al., 1993; Suerbaum et al., 1993).

Adhesin

H. pylori specifically colonizes the mucosal epithelium of the stomach with a marked preference for the antium (Blaser, 1987). The bacteria are found closely associated with

mucosal cells and can be observed in the intercellular junctions (Hazell et al., 1986, Chen et al., 1986). As well, H. pylori cells have been observed colonizing the duodenum at sites of gastric metaplasia (GM) (Wyatt et al., 1987; Wyatt et This observed tissue-specificity suggests the al., 1990). existence of H. pylori-specific receptors in the antrum of the stomach and in GM tissue in the duodenum (Lingwood, An N-acetylneuraminyl-lactose-binding hemagglutinin (NLBH) associated with the cell-surface has been identified, which can specifically bind to the N-acetylneuraminyl- $\alpha(2-3)$ component galactopyranosyl (NeuAc(2-3)Gal) N-acetylneuraminyl-lactose (NeuAc-lactose) on the surface of erythrocytes (Evans et al., 1988). Evans et al. postulated that this protein mediated the interaction between H. pylori and the gastric epithelium. The gene encoding this adhesin, designated hpaA, has been cloned and sequenced (Evans et al., 1993).

The adhesion of attaching and effacing enteropathogenic Escherichia coli to the surface of intestinal epithelial cells elicits the formation of adhesion pedestals (Moon, et al, 1988). These pedestals are the result of the polymerization of F-actin. The adhesion of H. pylori to epithelial cells was originally characterized as a similar "attaching and effacing" interaction (Hemalatha et al., 1991). However, polymerized F-actin adhesion pedestals could not be observed either in vitro, in cell cultures infected with H. pylori, or in vivo, in human gastric mucosal biopsies

of individuals infected with *H. pylori* (Dytoc et al.,1993). It has been demonstrated that phosphatidylethanolamine (Dytoc et al.,1993) and the human blood group antigen Lewis^b (Borén et al., 1993) are important for the adhesion of *H. pylori* to human epithelial cells.

Cytotoxin

Studies of cytotoxin production by *H. pylori* revealed that about half of the strains tested produced a protein, found in the supernatants of broth cultures, which caused a non-lethal vacuolization of eucaryotic cells (Leunk et al., 1988; Cover et al., 1990). This cytotoxic activity was both heat-labile and protease sensitive. Two proteins of 87 kDa and 128 kDa were observed in the supernatants with cytotoxic activity.

The 87 kDa protein was shown to have the vacuolating activity (Cover & Blaser, 1992b). The gene encoding the toxin, vacA, was sequenced from a number of toxin-producing H. pylori strains (Phadnis et al., 1994; Schmitt and Haas, 1994; Telford et al., 1994; Cover et al., 1994). sequence of the vacA gene from a toxin-negative strain was also obtained. This sequence was significantly different when compared to the sequence of the toxin-positive vacA gene (Cover et al., 1994). The exact role of cytotoxin in the pathogenesis of H. pylori is undetermined. Purified cytotoxin caused gastric lesions in mice similar to those observed in human subjects infected with H. pylori (Telford et al., 1994). This implied that expression of the cytotoxin may play a central role in the pathogenesis

H. pylori. Other studies indicated that patients infected with a strain of H. pylori expressing the cytotoxin had a higher risk of developing duodenal ulceration (Figura et al., 1989; Tee et al., 1993). All strains of H. pylori appeared to possess a vacA gene. However, there were marked differences in both the nucleotide and amino acid sequences between the vacA genes from the strain which did express cytotoxic activity and the strain which did not (Cover et al., 1994).

The role of the 128 kDa protein, designated CagA (Cytotoxin Associated Gene), in expression of cytotoxic activity is not clear. Expression of the CagA protein appears to be closely associated with expression of cytotoxin (Xiang et al., 1995), although the expression of cytotoxin is not dependent on CagA expression (Tummuru et al., 1994). It was also found, in hybridization studies, that not all H. pylori strains possess a cagA gene express the CagA protein (Xiang et al., 1995). The region upstream of the cagA gene may also be associated with cytotoxin expression, as some strains with the CagA-VacA-phenotype which possessed a cagA gene had deletions of several kilobases (kb) in this region. The cagA genes from two strains of H. pylori have been cloned and sequenced (Tummuru et al., 1993; Covacci et al., 1993).

Infection with *H. pylori* has been shown to induce the production of the interleukin IL-8 as part of the inflammatory response (Crabtree et al., 1994; Sharma et al.,

1995). Higher levels of IL-8 were induced by the CagA+VacA+ phenotype. However, isogenic mutants with CagA-VacA+, CagA+VacA-, or CagA-VacA- phenotypes showed no decrease in IL-8 induction (Sharma et al., 1995). This indicated that these two proteins were not directly involved in the IL-8 response.

A.3. Genetics of H. pylori

Although interest in genetic studies of *H. pylori* has increased steadily, relatively few *H. pylori* genes have been characterized. This is due, in part, to difficulties in cloning and characterizing genes from *H. pylori* (Clayton et al., 1989a; Covacci et al., 1993). The susceptibility of cloned *H. pylori* DNA to the host restriction modification systems, due to the methylation patterns, is thought to be one of the contributing factors to these difficulties (Phadnis et al., 1993).

Aside from the genes mentioned above (the urease genes, flaA, flaB, hpaA, vacA, and cagA), a number of other genes have been cloned and characterized. These include a species-specific 26 kDa protein gene (O'Toole et al., 1991), a bacterial ferritin gene (pfr) (Frazier et al., 1993), a lipoprotein gene (lpp20) (Kostrzynska et al., 1994), a superoxide dismutase gene (SOD) (Spiegelhalder et al., 1993), two heat-shock protein genes (hspA, hspB) (Suerbaum et al., 1994), a mucinase gene (hap) (Smith et al., 1994), a gyrase gene (gyrA) (Moore et al., 1995), a P-type ATPase gene (hpcopA) (Ge et al., 1995) and a cation-binding protein

(hpcopP) (Ge et al., 1995). In addition, the complete nucleotide sequences for the 16S ribosomal RNA genes of five different strains of *H. pylori* have been determined (Eckloff et al., 1994).

B. P-type ATPase of H. pylori

Chronic infection with *H. pylori* has been associated with the development of peptic and duodenal ulceration. Treatment of ulcers has involved the use of substituted benzimidazoles such as omeprazole and lansoprazole. These compounds act by blocking sulfhydral-groups of gastric P-type ATPases thus reducing the acidity of the gastric environment and reducing the severity of ulceration. It has been found that these compounds also have anti-bacterial activity against *H. pylori* (Iwahi et al., 1991; Loo et al., 1992). In fact, treatment with omeprazole alone (Stolte and Bethke, 1990) or in combination with antibiotics (Unge et al., 1989: Gellini et al., 1991) has been shown to eradicate *H. pylori* from some infected patients.

The target site of the omeprazole in *H. pylori* has not been elucidated. Omeprazole was shown to inhibit activity of purified urease (Bugnoli et al., 1993). Since urease has been shown to be an important colonization factor of *H. pylori*, this suggested that omeprazole could reduce the ability of *H. pylori* to colonize an individual by inhibiting urease activity. However, minimum inhibitory concentration

(MIC) studies of both urease positive and negative strains of *H. pylori* showed that both strains were equally inhibited by omeprazole (Bugnoli et al., 1993) and that inhibition of growth by omeprazole was directed through a target site other than the urease. Although a P-type ATPase from *H. pylori* has been cloned and characterized (Ge et al., 1995; this thesis) this particular ATPase did not appear to act as a target for omeprazole in vitro. This does not rule out the possibility of another P-type ATPase or the existence of another protein in *H. pylori* that is targeted by omeprazole.

C. Ribosomal RNA

The assembly of amino acids into proteins is a complex process involving many different components. Ribosomal RNA (rRNA) plays a central role in this process. The structure and function of rRNA is highly conserved throughout all life-The similarities and differences among these molecules are used in cataloging and sorting living organisms into their appropriate phylogenetic categories (Woese, 1987). All living organisms can be placed into one of three primary kingdoms based on 16S rRNA sequence. Eucaryotes form one kingdom, while procaryotes are divided into two primary kingdoms, Eubacteria and Archaebacteria (Woese, 1987). kingdom has many divisions. The kingdom Eubacteria is The genus Helicobacter is divided into 10 major classes. found in the class Proteobacteria, the purple bacteria. class was formerly divided into 4 subdivisions, alpha, beta, gamma and delta. A fifth subdivision, epsilon, has been created based on partial 16S and 23S hybridization studies (Vandamme et al., 1991). Within this subdivision there are three distinct rRNA clusters. Helicobacter spp. are grouped with Wolinella succinogenes, Flexispira rappini, and some yet unnamed Campylobacter spp. in rRNA cluster III. The majority of Campylobacter spp. are grouped together in rRNA cluster I, while members of the genus Arcobacter, along with some Campylobacter-like organisms, make up rRNA cluster II.

Procaryotic ribosomes contain three species of rRNA, 23S, 16S and 5S, which interact with ribosomal proteins to form the mature ribosome. Normally, single copies of the 16S and 23S genes and one or two copies of the 5S gene are clustered together in an operon. Several copies of these operons are usually found scattered throughout the genome. The genes for the rRNA species are not always linked together. This is the case in some strains of Borrelia spp. (Schwartz et al., 1992), Campylobacter jejuni and Campylobacter coli (Taylor et al., 1992b), Mycoplasma hyopneumoniae (Taschke et al., 1986), Mycoplasma gallisepticum (Chen and Finch, 1989), Leptospira interrogans (Fukunaga and Mifuchi, 1989), Pirellula marina (Liesack and Stackebrandt, 1989), Thermoplasma acidophilum (Ree and Zimmermann, 1990), Thermus thermophilus (Hartmann and Erdmann, 1989) and H. pylori (Taylor et al., 1992a; Bukanov and Berg, 1994). The fact that rRNA genes can be found separate from each other raises questions about regulation of rRNA expression. It was assumed that the

relative amounts of the various rRNA species were equal because they were transcribed together in operons (Nomura et al., 1984). In E. coli, for example, the 16S, 23S and 5S genes are co-transcribed as a 30S rRNA precursor, which is later processed to give equimolar amounts of the three rRNA species. In organisms where the rRNA genes are not linked, some form of regulation to ensure that the rRNA species are represented in the proper proportions may be required.

D. Genomic studies of H. pylori

A genomic restriction map of H. pylori strain UA802 has been constructed using pulsed-field gel electrophoresis (PFGE) (Taylor et al., 1992a). This map was made using the restriction endonucleases NotI and NruI. A variety of genetic elements have been localized on this map. In 1994, Bukanov and Berg, through the use of a contiguous cosmid library, constructed a physical map of H. pylori strain Comparisons of the maps produced by these two NCTC11638. groups, as well as comparisons of PFGE patterns of different strains of H. pylori (Taylor et al., 1992a), show a large degree of variability in both the restriction patterns and the arrangement of genetic elements from strain to strain. This genomic diversity has also been observed in restriction patterns using conventional electrophoresis (Majewski et al., 1988; Clayton et al., 1989), as well as arbitrary primer PCR (Akopyantz et al., 1992). The reasons for this variability are unclear. A number of hypotheses have been put forth

including differences in methylation patterns, a high frequency of silent mutations (i.e. mutations which do not cause phenotypic change), genetic uptake or exchange and recombination with other strains or bacteria, as well as both global and local rearrangements of the chromosome. Environmental pressures during the course of infection of a new host may also trigger chromosomal rearrangements (Taylor et al., 1992a).

E. The major outer membrane protein of Campylobacter jejuni

campylobacter spp. are recognized as one of the leading causes of bacterial gastroenteritis world-wide. *C. jejuni* is the species most commonly isolated from man. In the United States alone, *C. jejuni* has been estimated to be responsible for up to two million cases of acute diarrhea each year (Blaser and Reller, 1981). The pathogenic mechanisms of *C. jejuni* in the etiology of disease are still not clear. Adhesion to the intestinal epithelium is one of the first steps in the pathogenesis of this organism. A number of outer membrane proteins have been implicated which may be involved in adhesion. One of these proteins is the 45 kDa major outer membrane protein (MOMP) which has been shown to be a porin (Huyer et al., 1986).

In 1990, our laboratory developed non-radioactive DNA probes for the identification of *C. jejuni* and *C. coli* (Taylor and Hiratsuka, 1990). Initially, these probes were

isolated from a λ-gtll expression library of *C. jejuni* UA580 using polyclonal antisera directed against the 45 kDa MOMP. The antiserum used was also reactive with a number of other minor components. Cloned DNA inserts from two positive λ-gtll clones were further subcloned into pUCl3 and these plasmid clones were used as probes. Plasmid pDT1719 contained an 1845 bp insert and could be used to identify both *C. jejuni* and *C. coli* by DNA hybridization. Under reduced stringency pDT1719 was also able to weakly hybridize *C. laridis*. The plasmid pDT172C contained a 1475 bp insert and specifically hybridized to *C. jejuni* chromosomal DNA. Further work to characterize the cloned DNA has yet to be completed.

F. Objectives

F.1. H. pylori

An objective of this research was to construct genomic libraries of *H. pylori* strains and use these libraries to clone various genetic elements. These genomic libraries of *H. pylori* would provide a source of DNA for future research in isolation and characterization of other *H. pylori* genes. Construction of the genomic libraries is Jescribed in Chapter 3.

The genomic libraries were used to clone genes involved in P-type ATPase activity. The cloning of these genes is described in Chapter 4. Characterization of P-type ATPase

genes cloned in this work has been published (Ge et al., 1995).

Another objective was to clone and sequence the 23S and 5S genes of H. pylori (Chapter 4), and to use these sequences to investigate phylogenetic relationships of H. pylori to other organisms. As mentioned previously, the sequences of the 16S genes from five different strains of H. pylori have already been deduced (Eckloff et al., 1994). Studies utilizing the 23S and 5S genes of H. pylori have been limited to DNA-TRNA hybridizations. Complete sequences for these genes have not been reported. The nucleotide sequences of these genes will give us a clearer picture of where H. pylori fits phylogenetically in relation to other closely related organisms, such as Campylobacter species. H. pylori has already been associated phylogenetically with these organisms based on 16S and 23S hybridization studies.

The revision of a genomic restriction map, constructed previously by Taylor et al., 1992a, was another objective of this research. These revisions included the addition of sites for a third restriction enzyme, SfiI, and confirmation that H. pylori UA802 has two copies of rRNA genes, as observed in other strains of H. pylori (Bukanov and Berg, 1994; Jiang, 1994).

F.2. Campylobacter jejuni major outer membrane protein The objectives for the study of the *C. jejuni* major outer membrane protein (MOMP) were two-fold. First, the characterization, including sequence analysis, of two

Campylobacter DNA probes initially isolated using MOMP specific antiserum was carried out. Sequence analysis was useful in identifying genetic elements which had been cloned. The second objective was to obtain amino-acid sequence of purified MOMP protein. No sequence data for either the nucleotide sequence or amino-acid sequence has yet been published for the MOMP of *C. jejuni*.

Chapter 2

Materials and Methods

A. Bacterial strains, growth conditions and vectors A.1. Helicobacter pylori strains

The Helicobacter pylori strains used for this research are listed in Table 2-1. A number of different growth media support the growth of H. pylori (Goodwin & Armstrong, 1990, Goodwin & Worsley, 1993). For this research H. pylori cultures were grown on brain heart infusion (BHI) agar (Oxoid, Oxoid Ltd., Basingstoke, UK) with yeast extract (5%; Difco, Difco Laboratories, Detroit, MI), vancomycin (6 mg/l; Sigma, Sigma Chemical Company, St. Louis, MO, USA), amphotericin B (10 mg/l: Sigma) and ..5% animal serum (equine or bovine; Morse's Biological Supplies, Edmonton, AB, Canada) as previously described (Taylor et al., 1992). Cultures were incubated for 2-4 days in anaerobic jars with 5% H₂, 5% CO₂, and 85% N₂ (Linde Canada Inc., Edmonton, AB, Canada) at 37°C.

BHI broth (Oxoid) with 20% glycerol (BDH, BDH Inc., Toronto, ON, Canada) was used as cryogenic storage media for long-term storage of H. pylori cultures. For cryogenic storage, 1.5 ml of the BHI/glycerol medium were inoculated with a heavy loopful of H. pylori from a plate culture. Aliquots of 100 μ l were dispensed into 1.0 ml cryogenic storage vials (Nalgene Co., Rochester, NY, USA) and the

Table 2-1. Helicobacter pylori strains used in this research

Strain	Source	Origin of cultures
UA802	Clinical isolate	University of Alberta Hospital, Edmonton, AB, Canada
UA763	National Collection of Type Cultures, NCTC11639	Laboratory Center for Disease Control, Ottawa, ON, Canada (LCDC)
UA764	Clinical isolate	LCDC
UA765	Clinical isolate	LCDC

cultures were flash frozen in liquid nitrogen. The frozen samples were stored at -82°C and have been maintained for at least 2 years without significant loss of viability.

A.2. Campylobacter jejuni

Campylobacter jejuni UAS80 (NCTC11168) was used for study of the *C. jejuni* major outer membrane protein (MOMP) and the development of a *C. jejuni* specific DNA probe. These cultures were grown on Mueller-Hinton agar (Oxoid) plates at 37°C in 7% CO₂, 85% humidity for 1-2 days in a Napco E-series model 5100 CO₂ incubator (National Appliance Co., Tualatin, OR, USA).

A.3. Escherichia coli strains

The Escherichia coli strains used in this study are listed in Table 2-2. These strains were grown at 37°C either in Miller Luria broth (LB)(Difco) or on Miller LB-plates containing 1.5% agar (Difco). Growth media were appropriately supplemented with antibiotics or other growth factors dependent on the requirements of the procedures used.

E. coli JM83, DH5 α and DH5 α MCR were used as host strains for pUC20 (Chapter 3). E. coli LE392 was the host strain used for cosmids pSa747 and pHC79. E. coli KW251 was the host strain for λ -GEM11, while SRB and SRB P2 were used as hosts for λ -DASH.

Solid media used for screening pUC20 plasmids were supplemented with carbenicillin (500 mg/l; Ayerst) or ampicillin (100 mg/l; Ayerst), X-gal (100 mg/l; Gibco/BRL),

Table 2-2. Escherichia coli strains used .n this research

Strain	Genotype	Source
JM83	ara, Δ(lac-proAB) rpsL φ80 lacZM15	Bethesda Research Laboratories (BRL)
DH10B	F-mcrA Δ(mrr-hsdRMS-mcrBC) φ80d lacZΔM15 ΔlacX74 deoR recAl endAl Δ(ara,leu)7697 araD139 galU galK nupG rpsL	BRL
ER1793	fhuA2 Δ(lacZ)rl supE44 e14- trp-31 his-1 rpsL104 xyl-7 mtl-2 metBl Δ(mcrC-mrr) 114::IS10	New England Biolabs
DH5a	$F^-\phi 80dlacZ\Delta M15$ $\Delta (lacZYA-argF)U169$ deoR recAl endAl hsdR17(r_K^- , m_K^+) supE44 λ^- thi-1 gyrA96 relA1	Gibco/BRL
DH5αMCR	F-mcrA Δ(mrr-hsdRMS-mcr3C) φ80dlacZΔM15 Δ(lacZYA-argF)U169 deoRrecAl endAl supE44 λ-thi-1 gyrA96 relAl	Gibco/BRL
KW251	F' supE44 galK2 galT22 metB1 hsdR2 mcrB1 mcrA- agrA81::Tn10 recD1014 tetracyclineR	Promega
SRB	e14-(mcrA) Δ(mcrCB-hsdSMR-mrr)171 sbcC recJ uvrC umuC::Tn5 supE44 lac gyrA96 relA1 thi-1 endA1[F' proAB lacIqZΔM15] kanamycinR	Stratagene
SRB P2	el4-(mcrA) Δ(mcrCB-hsdSMR-mrr)171 sbcC recJ uvrC umuC::Tn5 supE44 lac gyrA96 relA1 thi-1 endA1[F' proAB lacIqZΔM15] spi- gam- red- kanamycin ^R	Stratagene
LE392	F- hsd R514 (r_k - m_k -) sup E44 sup F58 lac YI or $\Delta(lac$ IZY)6 gal K2 gal T22 met B1 trp R55 λ -	Boehringer Mannheim Company (BMC)

and IPTG (0.5 mM; BMC). For growth and propagation of pUC20 strains and pHC79 strains, solid media was supplemented with carbenicillin (500 mg/l) or ampicillin (100 mg/l). For plasmid and pHC79 DNA isolation, ampicillin (100 mg/l) was added to broth media. Solid and liquid media used for work with cosmid pSa747 were supplemented with 30 mg/l kanamycin (Sigma).

For work with lambda phage, MgSO₄ was added to all media, to stabilize lambda phage particles. The MgSO₄ was added as a 1.0 M so, tion through Millex GV $0.22\mu m$ micro-filtration filter units (Millipore) using a 10 ml syringe (B-D) to a final concentration of 10 mM. Maltose was also added to top agar and liquid media to a final concentration of 0.5% (w/v), to induce production of the maltose binding protein which acts as the receptor for the phage.

A.4. Vectors

For the construction of genomic libraries of H. pylori described in Chapter 3, puc20 (plasmid, BMC), pSa747 (cosmid, Tait et al., 1983), pHC79 (cosmid, BMC), λ -GEM11 (Promega), and λ -DASH (Stratagene) were used. For subcloning and double stranded sequencing the plasmid pBluescript (Stratagene) was used.

B. Cloning techniques

B.1.Isolation of chromosomal DNA

Genomic DNA was isolated from Helicobacter pylori strains UA763, UA764, UA765, and UA802, and Campylobacter jejuni

strain UA580 using a modified version of a previously described procedure (Ezaki et al., 1988). Briefly, organisms were grown on appropriate solid media. Cells were collected with a sterile loop and suspended in 1.0 ml of Mueller-Hinton broth in a 1.5 ml microfuge tube. The cells were harvested immediately, without incubation, in an Eppendorf model 5415C microfuge (Brinkman, Brinkman Instruments (Canada) Ltd., Rexdale, ON, Canada) at 16,000 x q for 30 s. The supernatant was removed and the cell pellet was resuspended in 300 μl of isolation buffer (Appendix A). RNase A (1.0 mg/ml;Pharmacia, Pharmacia Biotech Inc., Baie d'Urfe, PQ, Canada) was added to a final concentration of $10\mu q/ml$. Sodium dodecyl sulfate (SDS, 20% w/v solution; ICN, ICN Biochemicals Canada Ltd., Mississauga, ON, Canada) was added to a final concentration of 0.5%. The suspension was incubated at 65°C The suspension was then extracted twice with 500 μ l of phenol:chloroform:iso-amyl alcohol (25:24:1) and the aqueous phase was transferred to a new tube. was filled with chilled (-20°C) 95% ethanol (ETOH) and the DNA was spooled out of solution using a drawn-out glass micropipette. The DNA pellet was rinsed in chilled 95% ETOH and dissolved in 300 μ l of TE (Appendix A). residual ETOH, 30 μ l of 3.0 M sodium acetate (NaOAc) (pH 7.0) and 1.0 ml of chilled 95% ETOH were added. recovered by centrifugation in a microfuge for 5 min at 16,000 x g, washed once with chilled 70% ETOH and desiccated. The dry pellet was dissolved in an appropriate amount of TE to give a final concentration of DNA between 0.1-1.0 $\mu g/\mu l$. The purity of the DNA samples was checked and the concentration of the DNA samples was estimated by electrophoresis on 0.5% agarose gels (Section C.1). The gels were also useful in determining the amount of shearing which may have occurred during the isolation process. If the DNA appeared pure and was not significantly sheared it was used for further experimentation.

B.2. Restriction endonuclease digestion of DNA

Restriction endonuclease digestions were carried out in conditions suggested by the manufacturers of the enzymes. For genomic library construction, chromosomal DNA was either completely or partially digested with the appropriate restriction endonuclease to give ends compatible with the insertion site of the chosen vector.

Complete restriction endonuclease digestion of DNA

For complete restriction endonuclease digestions DNA was digested with 1-5 units of enzyme/ μ g DNA for 2-16 hr in conditions recommended by the manufacturer of the enzyme. Digestions were stopped with the addition of 1/10th volume 0.5 M EDTA (pH 8.0). In some instances, digestions were immediately used for electrophoretic analysis. For these samples 1/10th volume of loading dye (Appendix A) was added. For the construction of plasmid libraries of H. pylori UA763 (Chapter 3), complete digestions of pUC20 DNA or H. pylori UA763 DNA were carried out with the enzymes BgIII, EcoRI or

SstI. The lambda vectors and pHC79 were digested to completion with BamHI, while the pSa747 was cut with BglII.

For chromosomal DNA to be used for genomic library construction it was necessary to remove EDTA and any residual enzyme, which would interfere with the ligation reactions. To do this the DNA was precipitated by the addition of $1/10^{\rm th}$ volume of 3.0 M NaOAc and 3 volumes of chilled 95% ETOH. The DNA was recovered in a microfuge and the pellet was washed twice with 200 μ l of chilled 70% ETOH. The DNA was desiccated and the pellet was dissolved in 10 μ l of either TE or double-distilled water.

Genomic restriction digestions of the *H. pylori* strains UA763 and UA802 were carried out as described in Chapter 4, using the restriction endonuclease *BglII*.

Partial digestion of chromosomal DNA for cloning

Partial digestion of chromosomal DNA was carried out for the construction of genomic cosmid and lambda libraries of H. pylori strains UA763 and UA802 (Chapter 3). The genomic DNA was partially digested with either Sau3AI or BgIII. Approximately 50 μ g of chromosomal DNA were digested in a volume of 125 μ l with 0.1-0.5 units of enzyme/ μ g of DNA under conditions recommended by the manufacturer of the enzyme. At 5, 10, 15, and 30 min 25 μ l aliquots were removed and added to tubes containing 5μ l of 0.5M EDTA (pH 8.0). After 2 hr 5 μ l of 0.5 M EDTA (pH 8.0) were added to the final 25 μ l of the digestion. The digestions were pooled and 10 μ l of 3M NaOAc (pH 7.0) and 300 μ l of chilled 95% ETOH were added. The DNA

was precipitated overnight at -20°C. The DNA was recovered by centrifugation in a microfuge and the pellet was washed once with chilled 70% ETOH. The pellet was desiccated and dissolved in 100 μ l of TE.

B.3. Sodium chloride gradient size fractionation of DNA

Partially digested DNA was passed through a continuous 13.0 ml 1.0-5.0 M sodium chloride gradient in order to fractionate the DNA fragments by size. The gradients were made from the bottom up using an Easy Cast™ gradient maker (Owl Scientific Plastics Inc.) connected to a model P-1 peristaltic pump (Pharmacia) and mixed with a vibrating mixer (Haakebuchler Instruments Inc.). To form the gradients, 6.9 ml of 1.0 M NaCl in TE were added to the front chamber of the gradient maker and 6.3 ml of 5.0 M NaCl in TE were added to The mixer and the peristaltic pump were the back chamber. switched on and the flow from the back chamber was opened. The centrifuge tube was filled from the bottom up through the peristaltic pump via a plastic tube connected to siliconized glass FISHERbrand disposable 100 μ l micro-pipette (Fisher, Fisher Scientific, Edmonton, AB, Canada). The partial digestions were carefully layered on top of the gradients and the gradients were centrifuged at 38,000 rpm for 4 hr at 22°C in a Beckman L-80 ultracentrifuge (Beckman, Beckman Instruments (Canada) Inc., Mississauga, ON, Canada) using a Beckman model SW40 swinging bucket rotor. fractionate the gradients, the peristaltic pump was connected to a fraction collector and 300 μ l fractions were collected from the bottom of the tubes into 1.5 ml microfuge tubes. The fractions were de-salted by first precipitating the DNA with 1.0 ml of chilled 95% ETOH. The pellets were then washed several times in 70% ETOH at room temperature until they lost their grainy texture (precipitating salt). pellets were air dried (5-10 min) and were dissolved in 100 μ l of TE. To check the range of fragment sizes of the DNA in each fraction, 10 μ l of each fraction were subjected to electrophoresis, with appropriate standards, on a 0.4% agarose gel. For cosmid libraries, DNA fragments of greater Lambda libraries required DNA than 23 kb were needed. fragments in the range of 10-22 kb. The fractions containing DNA fragments in the correct size ranges were re-precipitated by adding 10 μ l of 3M NaOAc (pH 7.0) and 300 μ l of chilled The pellets were dissolved in TE and pooled to 95% ETOH. give a final volume of 50 μ l. This pooled DNA was then used for ligation to either cosmid or phage vectors.

B.4. Preparation of vector DNA for cloning Preparation of plasmid and cosmid vectors

Plasmid and cosmid vectors were digested to completion with restriction enzymes to give ends compatible with the passenger DNA fragments. In some cases double digestions of vectors were used to give directional clones. If a digestion buffer compatible with both enzymes was available, the digestions were carried out with both enzymes added simultaneously. If no compatible buffer was available, the

DNA was digested with the first enzyme, precipitated, using $1/10^{\rm th}$ volume NaOAc and 3-4 volumes chilled 95% ETOH, and redissolved in buffer suitable for the second digestion. Salt and residual enzymes which could interfere with ligation were removed either by precipitation of the DNA using NaOAc ($1/10^{\rm th}$ volume) and chilled 95% ETOH (3-4 volumes) or by centrifuging the digested vector through an $0.45\mu \rm m$ Ultrafree®-probind filter unit (Millipore Corp., Bedford, MA, USA).

Preparation of lambda vectors

The lambda replacement vectors, λ -GEM11 and λ -DASH (see Chapter 3 for description of the vectors), were first digested to completion with restriction enzymes that gave ends compatible with the passenger DNA fragments (BamHI). Right and left arms of the vector were then isolated by electrophoresis through a 0.5% low melting point (LMP) agarose gel, followed by excision and purification of the arms from the LMP gel (Section C.5). The arms were dissolved in TE and pooled in a final volume of 15 μ l and used for ligation to passenger DNA. In some cases vector arms which had already been restricted and purified were commercially available.

B.5. Ligation of vector to passenger DNA

For ligation of DNA fragments to either plasmid or cosmid vectors, a passenger:vector molar ratio of approximately 2:1 was used. For ligation of DNA fragments to lambda vectors arms, a passenger:vector arms molar ratio of approximately 1:1 was used. The ligations were carried out at 4°C

overnight, 15°C overnight or room temperature for 2-4 hr in buffer conditions specified by the manufacturer of the enzyme (Cibco/BRL Life Technologies Inc. (BRL), Burlington, ON, Canada). In general, about 1-4 μ g of total DNA were used for ligation reactions with 1 unit of enzyme in a final volume of 10 μ l.

B.6. Transformation of competent cells

Competent E. coli cells obtained commercially or made in our laboratory were used for transformation of plasmid or cosmid vectors. Competent cells were prepared as described (Ausubel et al., 1987). Briefly, the appropriate strain of bacterial culture was inoculated into 100 ml of suitable The culture was grown overnight (16 hr) and liquid medium. harvested in a 250 ml centrifuge bottle at 4°C for 5 min at 5,000 rpm using a Beckman model JA-14 rotor in a Beckman The cells were then gently model J 2-21 centrifuge. resuspended in 10 ml of chilled Ca-buffer (Appendix A) and transferred to a sterile 17 x 100 mm capped polypropylene culture tube. The cells were incubated on ice for 30 min and collected at 4°C for 5 min at 5,000 rpm using a Beckman model JA-20.1 rotor. The cells were gently resuspended in 3.0 ml of a chilled solution of 60 mM $CaCl_2$, 15% (v/v) glycerol, and 10 mM PIPES, pH 7.0. The competent cells were divided into 300 μ l aliquots in 1.5 ml microfuge tubes. If the cells were not used immediately, they were flash-frozen in liquid nitrogen and stored at -80°C.

To transform cells, four serial 10-fold dilutions of the ligated DNA were made in $100\mu l$ volumes of chilled Ca-buffer to give aliquots containing 1.0 μl ,0.1 μl , 0.01 μl , and 0.001 μl of ligated DNA solution. Twenty-five μl of competent cells were added to each tube and the tubes were incubated on ice for 90-120 min. The cells were then heat-shocked at 42°C for 2 min and plated onto suitable solid media, supplemented with antibiotics as required. The plates were incubated at 37°C with the agar layer up (lid down) overnight. The following day, colonies were counted to determine the efficiency of ligation and transformation. Isolated colonies were selected for screening (see Section B.8 below).

B.7. In vitro packaging and transfection of lambda and cosmid DNA

Lambda or cosmid DNA was packaged into viable bacteriophage heads before transfection. Several commercially prepared in vitro lambda packaging extract kits were available. For this research, extracts from Boehringer Mannheim Canada Ltd. (BMC, Laval, PQ, Canada), United States Biochemical (USB, Cleveland, OH, USA), and Stratagene (La Jolla, CA, USA) were all used following the protocols accompanying the packaging kits. For the packaging reactions 4-6 μ g of ligated DNA in a volume of 4 μ l were used.

For transfection, 5 ml overnight cultures of appropriate bacterial cells were grown in sterile 17×100 mm polypropylene culture tubes in LBM (Appendix A). The cells were harvested at 4°C for 5 min at 5,000 rpm using a Beckman

model JA-20.1. The cells were resuspended in 1 ml of 10 mM MqSO4 and placed on ice. Serial 10-fold dilutions of the packaged DNA suspensions were made in 10 mM MqSO4 and divided into 100 μ l aliquots. To each aliquot, 50 μ l of cells were The cells were incubated for 20 min at 37°C. The added. cells were then added to 2.5 ml of top agar (culture medium, 10 mM MgSO₄, 0.5% maltose, 0.7% agarose held at 48°C) in 13 x 100 mm test tubes, mixed rapidly, and plated on solid The top agar was allowed to solidify at room temperature for 10 min and the plates were incubated lids up at 38-40°C overnight. The following day the lambda plaques or cosmid-containing colonies were enumerated to determine the titer of the packaged lambda/cosmid suspensions. Five ml of SM buffer (Appendix A) were overlaid on plates with greater than 200 lambda plaques. These plates were then incubated overnight at 4°C and the following day the suspensions were transferred to sterile 17 x 100 mm capped polypropylene tubes (Fisher), and 1 ml of chloroform was These 5 ml suspensions served as lambda library added. stocks and were re-plated for screening (see Section B.8). Isolated cosmid colonies were screened as described in Section B.8.

B.8. Replica filters for screening Screening of plasmid and cosmid clones

For the screening of plasmid or cosmid clones, isolated colonies from the primary plates (from Section B.6 or Section B.7 above) were spotted, using a sterile tooth pick, in

arrays onto two plates of solid growth medium. On one plate sterile 82 mm nitrocellulose disks (MSI, Micron Separation Inc., Westboro MA, USA), marked for orientation and in some cases marked with grid lines, were carefully layered onto solid growth media and the colonies were spotted directly onto the nitrocellulose. The second plate did not have a nitrocellulose disk and was used as a master plate. The plates were incubated at the appropriate temperature (usually 37°C) overnight or until the colonies reached approximately The filter plate was processed by first 5 mm in diameter. carefully removing the filter with sterile forceps and placing the filter colony-side up for 5 min on Whatman 3MM chromatography paper which had been soaked in 10% SDS. This step allowed lysis of the cells. The filter was then transferred, colony side up, to Whatman 3MM chromatography paper pre-soaked in a denaturing solution (Appendix A) and then allowed to soak in this solution for 10 min. The filter was then neutralized by floating it on neutralizing solution (Appendix A) for 10 min. The filter was then rinsed in 2 x SSC, air dried and the DNA was fixed to the filter by baking (no vacuum) overnight at 65°C. The filters were screened by hybridization as described below (Section C.4).

Screening of lambda clones

Lambda plates, made as described in Section B.7, were screened by transferring the lambda DNA to nitrocellulose disks. Nitrocellulose disks were marked for identification and orientation, then sterilized by autoclaving. Sterile

forceps were used to layer these disks, with the markings down, onto the plates (ie. in contact with the plate surface). The filters were allowed to soak on the plate for During this period the bottoms of the plates were marked to orient the filters. The filters were then carefully removed and floated plaque-side up on denaturing solution for 30-45 s. The filters were then floated on neutralizing solution for 5 min and rinsed in 2 x SSC The filters were air dried and the DNA was (Appendix A). fixed onto the filter by baking (no vacuum) at 65°C The filters were screened by hybridization as overnight. described below (Section C.4). Positive plaques were isolated using sterile toothpicks, and placed in 500 μ l of SM buffer with 50 μ l of chloroform and stored in this solution indefinitely at 4°C. The positive plaque stock was amplified by re-plating 50 μ l of the phage suspension (Section B.7). This amount of phage suspension was normally enough to give confluent plaque formation (ie. the plate was clear). plate was overlaid with 5 ml of SM (Appendix A) and incubated The overlay was then transferred to overnight at 4°C. sterile 17 x 100 mm capped polypropylene tubes (Fisher) and 1 ml of chloroform was added to each tube. This stock solution could be kept at 4°C indefinitely.

Four plates with greater than 200 plaques of each of the BglII and Sau3AI λ -DASH genomic libraries of H. pylori UA802, were screened with PCR probes labelled by nick translation with [32P]-dATP (Chapter 4). Approximately 2000

plaques/library were screened. Positive clones were isolated using sterile toothpicks and stock suspensions were made, as $d\varepsilon$ scribed above. Some of the positive clones were further characterized by restriction digestion analyses and sequence analyses.

B.9. Isolation of plasmid/cosmid DNA Mini-preparations of plasmid/cosmid DNA

Mini-scale plasmid/cosmid DNA isolations were carried out using methods modified from those of Birnboim and Doly broth cultures, supplemented with Five m (1979).appropriate antibiotics, were inoculated with single colonies harboring the plasmid/cosmid to be isolated. The cultures were grown in sterile 17 x 100 mm capped polypropylene tubes (Fisher) overnight at 37°C in a tube roller (Model TC-7; NBSC, New Brunswick Scientific Co., Edison, NJ, USA). cells were harvested in a benchtop centrifuge (Sorvall GLC 2B General Laboratory Centrifuge, DuPont Instruments) for 3 min at 4000 rpm, resuspended in 100 μ l of Birnboim & Doly solution 1 (Appendix A) and transferred to 1.5 ml microfuge Two hundred μ l of Birnboim and Doly solution 2 tubes. (Appendix A) were added and mixed by inverting the tube. After 1 min, 150 μ l of Birnboim and Doly solution 3 (Appendix The tubes were then A) were added and mixed by inversion. centrifuged in a microfuge at 16,000 x g for 15 min. Pelleted material was removed from the tube using sterile toothpicks and the tubes were filled with chilled 95% ETOH. These tubes were then incubated at -20°C for at least 2 hr.

The precipitated DNA was then collected in a microfuge for 2 min and washed twice with chilled 70% ETOH. The final ETOH wash was removed and the DNA pellets were dessicated in a vacuum dessicator (Fisher) for 5 min. The DNA was then dissolved in TE + 1μ g/ml RNase A (BMC). In some instances the DNA was further cleaned by phenol/chloroform extraction and precipitation or by passing the DNA solution through a 0.45 μ m Ultrafree®-probind filter unit (Millipore).

Large scale isolation of plasmid/cosmid DNA

Large scale DNA isolations were done either by use of a DNA isolation kit (Quiagen) or by using methods modified from those of Birnboim and Doly (1979). The modified Birnboim and Doly isolations were carried out as follows. Two hundred ml culture media, appropriately supplemented with o f antibiotics, were inoculated with a single colony of bacterial culture harboring the plasmid/cosmid of interest. The cultures were incubated overnight at 37°C in a controlled Cells were then environment incubator shaker (NBSC). harvested in 250 ml centrifuge bottles at 4°C for 5 min at The cells were 5,000 rpm using a Beckman model JA-14. resuspended in 2.5 ml of Birnboim and Doly solution 1 and transferred to a 50 ml polypropylene centrifuge tube. cell suspension was chilled on ice for 10 min. Five ml of Birnboim and Doly solution 2 well added and mixed by inversion using Parafilm® (American National Can, Greenwich, CT, USA) to cover the tube mouth. After 2 min of incubation on ice, 3.0 ml of Birnboim and Doly solution 3 were added and the suspension was again mixed by inversion. Incubation was continued on ice for 30 min and the tube was then centrifuged at 12,500 rpm at 4°C for 30 min using a Beckman model JA-20 The supernatant was transferred to a new 50 ml rotor. centrifuge tube and 30 ml of chilled 95% ETOH were added. The tube was mixed by inversion and incubated for at least 2 hr at -20°C. The tube was then centrifuged at 10,000 rpm at 4°C for 10 min using a Beckman model JA-20 rotor. supernatant was discarded and the pellet was dried in a dessicator for 10 min. The pellet was dissolved in 9.0 ml of TE, and 9.0 g of CsCl were added. The CsCl was dissolved and the samples were then transferred into quick seal centrifuge tubes (Ultra-Clear®, Beckman). Ethidium bromide was added to give a final concentration of approximately 150 $\mu \mathrm{g/ml}$, the tube was filled to the top with a solution of CsCl (1.0 g/mlThe tubes were then in TE), and the tube was sealed. centrifuged for 22 hr at 22°C, at 55,000 rpm using a Beckman Ti70 rotor in a Beckman L-8 ultracentrifuge. The plasmid band was then removed from the centrifuge tube using a 16 gauge needle (B-D, Becton Dickinson and Company, Rutherford, NJ, USA) and a 3.0 ml syringe (B-D), and transferred to a 17 x 100 mm capped polypropylene tube. ethidium bromide was removed by extraction with iso-amyl alcohol (Sigma) and the DNA was precipitated with 2.5 volumes The pellet was washed several times of chilled 95% ETOH. with chilled 70% ETOH, until the pellet was no longer "grainy". The DNA was then air-dried for 15 min and redissolved in TE to give a final concentration of approximately 0.25-1.0 $\mu g/\mu l$.

B.10. Isolation of lambda DNA

Preparation of host cells for transfection

Single isolated colonies of the host cells were inoculated into 5 μ l of LBM-broth (Appendix A) in sterile 17 x 100 mm capped polypropylene tubes. One tube produced enough cells for 20 mini preparations or one large scale preparation. The cells were grown overnight at 37°C in a tube roller (NBSC). The following morning the cells were harvested in a benchtop centrifuge (Sorvall) for 3 min at 4000 rpm. The cells were then resuspended in 1.0 ml of 10 mM MgSO₄ and used for transfection.

Mini-preparations of lambda DNA

In a 1.5 ml microfuge tube, approximately 2.5 x 10^7 plaque forming units (PFU) of phage were combined with 50 μ l of host cells in a final volume of 200 μ l. The phage were allowed to transfect the host cells at 37° C for 15-20 min and the suspension was then added to 4 ml of LBM-broth in sterile 17-100 mm capped polypropylene tubes. The broth culture was grown for 9-12 hr. on a tube roller at 37° C. Three hundred μ l of chloroform and 1 ml of 5.0 M NaCl were added and the suspension was incubated for an additional 10 min at 37° C. Cellular debris was removed from the phage suspension by centrifugation at 11,000 rpm at 4° C for 15 min using a Beckman model JA-20.1 rotor. The supernatant was transferred

to a fresh tube and the phage particles were precipitated by the addition of 1.25 ml of 40% w/v polyethylene glycol (PEG) The phage particles 8000 and incubated on ice for 20 min. were collected by centrifugation at 11,000 rpm at 4°C for 15 Phage were resuspended in 400 μ l of TM (Appendix A) and transferred to 1.5 ml microfuge tubes. Pancreatic DNase and RNase A were added to a final concentration of $1\mu g/ml$ and the tubes were incubated for at least 30 min at 37°C. The suspension was extracted twice against one volume of The bacteriophage particles were then collected chloroform. by centrifugation at 60,000 rpm at 22°C for 1 hr in a Beckman TLA-100.2 rotor in a Beckman TL-100 tabletop ultracentrifuge. The supernatant was carefully decanted leaving a "glassy" precipitate at the bottom of the tubes. The pellet was suspended in 300 $\mu 1$ of TM and transferred to a microfuge tube. The centrifuge tube was washed with an additional 100 μ l of TM, and wash was pooled with the original phage suspension. Proteinase K (at a final concentration of 50 $\mu q/ml$), SDS (at a final concentration of 0.1%) and EDTA (at a final concentration of 20 mM) were added and the suspension was incubated at 65°C for 20 min. The solution was then extracted once with one volume of phenol, once with one volume of phenol:chloroform and once with one volume of chloroform. The DNA was then precipitated by the addition of 1/10th volume of 3 M NaOAc and 3 volumes of chilled 95% ETOH. The tube was incubated for at least one hr at -20°C. was collected in a microfuge and washed twice with 300 μl of chilled 70% ETOH. The pellet was allowed to air-dry for about 15 min and was dissolved in an appropriate volume of TE to give a final concentration of approximately 0.2-1.0 $\mu g/\mu l$. The oncentration and purity of the DNA were assessed on a 0.6% agarose gel.

Large scale isolation of lambda DNA

The procedure for large scale isolation of lambda DNA was similar to the mini-preparation of lambda DNA but with the following changes. In a 13 x 100 mm test-tube, approximately 6×10^8 PFU of phage were used to transfect 1.0 ml of host The transfected cells were then added to 100 ml of cells. LBM in sterile 250 Erlenmeyer flask. The flasks were incubated overnight at 39°C in a controlled environment incubator shaker (NBSC). Two hundred and fifty ml centrifuge bottles were used for removal of cellular debris and centrifugation was carried out at 6,000 rpm at 4°C for 15 min with a Beckman JA-14 rotor. PEG precipitation of the phage particles was carried out as above with scaled-up volumes. The phage were resuspended in 750 μ l of TM and transferred to 1.5 ml microfuge tubes. The suspension was treated with pancreatic DNase ($1\mu g/ml$) and RNase A ($1\mu g/ml$) and extracted twice against one volume of chloroform. The bacteriophage particles were then collected by centrifugation at 60,000 rpm at 22°C for 1 hr in a Beckman TLA-100.2 rotor in a Beckman TL-100 tabletop ultracentrifuge. The phage particles were treated with proteinase K, SDS and EDTA as above and incubated at 65°C for 20 min. Phenol, phenol:chloroform and chloroform extractions were carried out as described and $1/10^{\rm th}$ volume of 3 M NaOAc and 3 volumes of chilled 95% ETOH were added. The DNA was spooled out of the tube with a drawn-out glass micropipette and dissolved in 300 μ l of TE. The DNA was re-precipitated with $1/10^{\rm th}$ volume of 3 M NaOAc and 3 volumes of chilled 95% ETOH. The DNA was washed twice with 300 μ l of chilled 70% ETOH, air-dried, and dissolved in TE to give a final concentration of approximately 0.2-1.0 μ g/ μ l. The concentration and purity of the DNA were assessed on a 0.6% agarose gel.

C. DNA manipulation techniques

C.1. Agarose gel electrophoresis

Agarose gel electrophoresis was used for a variety of purposes such as restriction digestion analysis of DNA, sizing of intact or fragmented DNA, purity checks of DNA, Southern transfer and hybridization analysis, and analysis of PCR products. The concentration and size of the agarose gels as well as the duration of electrophoresis were determined by the size range of the DNA to be separated. Denaturing agarose gels were used in the analysis of RNA samples (Section F.2). PFGE was used for separation of large DNA fragments for genomic restriction map construction (Section D.3). Low melting point agarose gels were used for extraction of DNA fragments (Section C.5).

Gels were run in 0.5 x Tris-borate (TB) buffer (45 mM Tris base, 45 mM boric acid, 1 mM EDTA, pH 8.3) using horizontal

gel electrophoresis chambers (mini sub-cell, wide mini sub-cell, Bio-Rad, Bio-Rad Laboratories Ltd., Mississauga, ON, Canada; Aquabogue Model 850, Aquabogue Machine Shop, Aquabogue, NY, USA). Bio-Rad model 1450 (Bio-Rad) or Fisher Biotech model FB 105 (Fisher) power supplies were used for electrophoresis. Either a 1 kb ladder (BRL) or a 123 bp ladder (BRL) were used as size standards for linear DNA.

C.2. Southern transfer

DNA was transferred from agarose gels to either supported nitrocellulose (NitroPlus; MSI) or nylon transfer membrane (Hybond; Amersham, Amersham Canada Ltd., Oakville, ON, Canada) by the methods of Southern (Southern, 1975). The DNA was fixed to the nitrocellulose membranes by baking (no vacuum) at 65°C for a minimum of 2 hr. For nylon membranes, the DNA was cross-linked to the membranes in a Stratalinker® UV crosslinker oven (Stratagene).

C.3. Preparation of probes for hybridization

Nick translation was used for labelling probes with radioactive nucleotides. Random priming was used for labelling probes with either non-radioactive digoxygenin (DIG) or with radioactive nucleotides.

Nick translation

The DNA to be labelled (0.3-1.0 μ g) was incubated in a 500 μ l microfuge tube in NT buffer (Appendix A) with 1.0 μ l [α^{32} P]-dATP (3000 mCi/mmol, 10 mCi/ml; Dupont Canada Inc., Mississauga, ON, Canada), 2.0 μ l of DNAse (0.2 μ g/ml; BMC) and 1.0 μ l of DNA polymerase I (5 units; BMC) in a final

volume of 20 μ l for a minimum of 90 min at 15°C in a Lauda K-2/RD circulating water bath (Brinkman). For some labelling experiments $[\alpha^{32}P]$ -dCTP (3000 mCi/mmol, 10 mCi/ml; Dupont) was used instead of $[\alpha^{32}P]$ -dATP. For these experiments dATP replaced dCTP in the 1 X NT buffer. Some probes were labelled using a commercial Nick Translation Kit (BRL) following the protocols specified by the manufacturer of the kit, using the same incubation times and temperatures as The labelled DNA was then precipitated by the addition of 1/10th volume of 3.0 M NaOAc (pH 7) and 3 volumes of chilled 95% ETOH (-20°C). The DNA was collected by centrifugation in microfuge for 2 min. The supernatant was carefully removed, placed in an microfuge tube, and discarded to an appropriately shielded radioactive discard receptacle. The pellet was washed once with 80 μ l of chilled 70% ETOH and collected in the microfuge. Again the supernatant was discarded to the radioactive discard receptacle. The pellet was allowed to dry for 15 min and redissolved in 100-200 μ l of TE. One μ l of the labelled DNA was analyzed in a Beckman model LS6800 scintillation counter to determine specific activity.

Random primer labelling

Probe DNA $(0.3-1.0~\mu g)$ was denatured by heating at 95°C for 10 min in a 500 μ l microfuge tube. The denatured DNA was chilled rapidly on wet ice and 2.0 μ l of random hexanucleotide mix $(0.8~\mu g/\mu l)$; University CORE DNA Services, Calgary, AB, Canada), 2.0 μ l 10 X NT buffer (Appendix A),

1.0 μ l [α^{32} P]-dATP (3000 mCi/mmol, 10 mCi/ml; Dupont) and 1 μ l of Klenow polymerase (1 unit/ μ l; BRL) were added to the tube and the volume was adjusted to 20 μ l with sterile distilled water. For DIG labelling the 10 X NT buffer and [α^{32} P]-dATP were replaced by 2.0 μ l dNTP mix (Appendix A). The labelling mixture was then incubated at 37°C for at least 60 min. In most cases the labelling reaction was allowed to incubate overnight. The probe DNA was then precipitated and redissolved in 100 μ l of TE. For the DIG-labelled probe, 4.0 M LiCl was used in place of the 3.0 M NaOAc. The [α^{32} P]-dATP labelled probes were counted in a scintillation counter.

C.4. Hybridization

Membranes prepared as outlined above (Sections B.8 and C.2) were hybridized with either radioactive or DIG-labelled Membranes were pre-hybridized by probes as follows. immersing the filters in hybridization solution (Appendix A). It was necessary to wet nitrocellulose membranes in 2 X SSC prior to pre-hybridization. Pre-hybridization was carried out either in sealable plastic freezer bags, or shallow sealable plastic containers at 37°C for at least 4 hr for radioactive probes and at least 1 hr for DIG-labelled probes. For hybridization of an average size filter (10 X 15 cm), a volume of 5.0 ml for the sealable bags or 50 ml for the plastic containers was used. Labelled probes were boiled in a water bath for 2-10 min and were then added to the pre- 10^{6} hybridization solution. Generally, 1 x CPM of radioactively-labelled probe/filter or ≥10 ng/ml of DIG-

labelled probe were added. The hybridization was then allowed to proceed, depending on the desired stringency, at $37-42^{\circ}\text{C}$ for a minimum of 4 hr.

Processing membranes screened with radioactive probe

Radioactively-probed filters were washed twice after hybridization in 2 X SSC, 0.1% SDS wash solution at 65°C for 30 min. Excessive background signals were removed by washing in 0.2 X SSC, 0.1% SDS at 65°C for an additional 30 min. This process was repeated if the background was still high. The washed membranes were dried and exposed at -82°C to Kodak X-OMAT AR X-ray film (Kodak, Eastman Kodak Co., Rochester, NY, USA) in a shielded X-ray cassette with an intensifying screen (Cronex Lightning plus, Dupont). The exposure times were varied to give the best results with the minimum amount of background. In most cases overnight exposure was adequate.

Processing membranes screened with DIG-labelled probe

Following hybridization, membranes probed with DIG-label were first washed at room temperature in 2 X SSC, 0.1% SDS twice for 5 min. This was followed by two 15 min washes in 0.1 X SSC, 0.1% SDS at 65°C. The membranes were briefly rinsed in wash buffer (Appendix A), then equilibrated for 30 min in blocking buffer (Appendix A). The membranes were then immersed in a solution of anti-DIG-alkaline phosphatase conjugate diluted 1:5000 in blocking buffer for 30 min at room temperature. Approximately 20 ml of diluted conjugate/100 cm² surface area of membrane were used. To

remove excess conjugate, the membranes were then washed twice for 15 min at room temperature with wash buffer. membranes were equilibrated for 5 min at room temperature in PPD-buffer (Appendix A) and immersed in a solution of Lumigen PPD (BMC) diluted 1:100 in PPD-buffer. Following this the membranes were removed and the Lumigen solution was allowed to drain from the membrane. Excess solution was removed by briefly blotting the membranes (DNA side up) on Whatmann 3MM The membranes were then wrapped in chromatography paper. Saran Wrap® or sealed in sealable freezer bags without drying and incubated for 15 min at 37°C. The wrapped membranes were then exposed to Kodak X-OMAT AR® X-ray film in an X-ray Exposure times were varied cassette at room temperature. depending on the strength of the signal and amount of Usually exposure times of 15-30 min were background. sufficient. If membranes were not allowed to dry, they could To re-probe a membrane it was first washed in be re-probed. sterile distilled water, then washed twice at 37°C in a solution of 0.2 M NaOH and 0.1 %SDS. The membrane was rinsed briefly in 2 x SSC and hybridized again as described above.

C.5. Low melting point agarose extraction of DNA fragments

Electrophoresis of DNA was carried out using standard procedures through low melting point agarose gels of appropriate concentrations. The gels were stained with ethidium bromide and the appropriate DNA fragment(s) were excised and excess agarose was trimmed off. The agarose

blocks containing DNA were placed into microfuge tubes with 100-200 μ l of TE buffer. The LMP agarose was melted by In order to separate the incubation at 65°C for 10 min. agarose from the DNA sample, the suspensions were extracted twice with one volume of phenol. The aqueous phase was carefully removed and the volume of the DNA solutions was reduced by extracting the aqueous layer with n-butanol (Anachemia Science, Montreal, PQ, Canada). This step also removed any traces of ethidium bromide which might have still been bound to the DNA. The DNA solutions were extracted once against one volume of chloroform and the DNA was precipitated by adding 1/10 volume of 3 M NaOAc and 2-3 volumes of 95% ETOH and incubating at -70°C for 30 min or -20°C overnight. The DNA was collected in a microfuge and the supernatant was discarded. The DNA pellets were washed once with 100 μl of The ETOH wash was chilled 70% ETOH and centrifuged again. removed and the DNA pellets were dessicated. The pellets were redissolved in an appropriate volume of TE.

C.6. Polymerase chain reaction (PCR) methods

PCR was used to generate specific DNA fragments for use as probes (Chapter 4). Conditions were varied depending on the Taq polymerase used, the degree of homology of the primer to the template DNA, and the size of the expected PCR product. In most cases, basic conditions, described below, were adequate. Using Promega Taq polymerase, the reactions were run in 1 X reaction buffer (provided as a 10 X solution by the manufacturer), with 1.25 mM MgCl₂, 0.125 mM dNTP (0.125

mM of dATP,dCTP, dGTP, and dTTP), 20 pmol of each primer, and up to 100 ng of template DNA with 2.5 units of enzyme. Forty cycles of 60s at 94°C, 60s at 45°C and 120s at 72°C were used.

With BRL Tag polymerase, the basic conditions were 1 X reaction buffer (provided as a 10 X solution by the manufacturer), with 1.5 mM MgCl₂, 0.125 mM dNTP, 20 pmol of each primer, and up to 100 ng of template DNA with 2.5 units of enzyme. Forty cycles of 40s at 94°C, 40s at 45°C and 90s at 72°C were used.

PCR was used to generate probes for the P-type ATPase as well as the 16S and 23S genes of *H. pylori* UA802 (chapter 4). Promega and BRL *Taq* polymerase were used for the amplification in conditions described above. The primers used are described in Chapter 4.

C.7. Partial digestion mapping of lambda clones

Rapid construction of crude restriction maps of lambda clones was accomplished using partial digestion mapping. Briefly, lambda clones were first digested to completion with the enzyme to be mapped, to determine the sizes of the digested fragments of the cloned DNA. The clone was then partially digested with the enzyme, fractionated on 0.5% agarose gels (Section C.1), and the DNA was transferred to nylon or nitrocellulose membranes by Southern transfer (Section C.2). These blots were hybridized with purified right arms of the lambda vector (Section C.5) labelled with either radioactive or non-radioactive label (Section C.3).

The resulting ladder of partially digested fragments which hybridized with the right arm of the vector was used to determine the order of the fragments in the lambda clones, and crude restriction maps were constructed.

D. Pulsed Field Gel Electrophoresis

D.1. Preparation of blocks for PFGE

Pulsed field gel electrophoresis is a method by which large DNA fragments, up to several hundred kb, can be resolved on To prevent shearing of the large DNA an agarose gel. fragments it was necessary to first immobilize bacterial cells in agarose blocks. Bacterial cells from a single plate of pure culture were suspended in 1.0 ml of suspension buffer (Appendix A) in a 1.5 ml microfuge tube. The cells were harvested for 20 s in a microfuge and resuspended in 0.75 ml The cells were then added to a 1.5% of suspension buffer. agarose solution (1.5% agarose in 0.25 M EDTA, 10 mM Tris HCl pH 7.5, kept at 58°C) and dispensed into a PFGE block mold. The blocks were allowed to solidify at room temperature for 10 min. The blocks were then transferred from the mold into 5 ml of lysozyme buffer (Appendix A) in a capped 17 x 100 mm polypropylene tube and incubated at 37°C for 2 hr. The blocks were then transferred to 5 ml of deproteination solution (Appendix A) in a capped 17 x 100 mm polypropylene tube and incubated at 58°C for 1-2 days. Residual proteinase activity was removed by three 15 min washes in PMSF-buffer (Appendix A) at room temperature. The blocks were then washed 10-15 times in 5 ml of PFGE block-washing buffer for 20 min at room temperature.

D.2. Restriction digestion of embedded DNA

The washed blocks containing the embedded DNA were sliced into 2 x 2 mm slices using a clean razor blade. Slices were pre-soaked in 400 μ l of the appropriate restriction enzyme buffer at room temperature for 15 min. The enzyme buffer was removed, 400 μ l of fresh buffer was added, and the slices were incubated for an additional 15 min. This process was repeated 5 times. After the final so μ the enzyme buffer was removed and 400 μ l of enzyme buffer with 2 unit/slice of restriction enzyme were added. The digestions were then incubated at the appropriate digestion temperature for 6-16 hr

D.3. PFGE gels

Gels were run using either the LKB 2015 Pulsaphor plus PFGE system or the Beckman Gene Line II Pransverse Alternating Field Electrophoresis (TAFE) system. The concentration of agarose used for PFGE was either 0.75% or 1.0% in PFGE running buffer (Appendix A). For the TAFE 1.0% agarose gels were used in TAFE running buffer (Appendix A). The gels were cast in molds provided by the manufacturers. The agarose blocks containing the embedded DNA were placed into wells formed in the gels and sealed into place using 1.5% low melting point agarose. PFGE gels were run in PFGE running buffer with constant voltage set at 190v. The gels were run at 8°C and pulse-times were varied from 5 s to 60 s. Pulse

times were varied directly with the size of the fragments to be separated. The duration of the pulse times varied depending on the range of fragment sizes in a given preparation. Once electrophoresis was complete the gels were processed as described in earlier sections either for Southern blots (Section C.2) or for LMP-agarose fragment extraction (Section C.5). For Southern blots of PFGE gels, the gels were first immersed in 0.25 M HCl for 15 min prior to the normal denaturation/neutralization steps.

E. DNA sequencing

E.1. Double-stranded dideoxy-sequencing

Double-stranded dideoxy-sequencing reactions were carried out using $[\alpha^{-35}S]$ -dATP as described (Wang, 1988) or following the protocols included in the Sequenase® dideoxy-sequencing kit (USB). Approximately 2-6 μg of DNA were used for each sequencing reaction. Using these methods the DNA sequences of the cloned DNA from the two Campylobacter probes, pDT1719 and pDT1720, were determined (hapter 6).

E.2. Cycle sequencing with ^{33}P

Cycle sequencing was carried out with $[\gamma^{-33}P]$ -ATP according to protocols included with the dsDNA Cycle Sequencing System (BRL). The reactions were carried out in a Perkin Elmer model 480 DNA thermocycler (Perkin Elmer) for forty cycles of 30s at 94°C, 30s at 50°C, and 40s at 72°C.

E.3. Sequence analysis

Sequencing gels were run in TBE-buffer (Appendix A) using either the BRL model S1 sequencing gel electrophoresis apparatus or the Bio-Rad Sequi-gen II sequencing cell system. The gels were composed of 5% acrylamide (19:1 acrylamide:bisacrylamide; Bio-Rad) and 8 M urea in TBE-buffer. To catalyze the solidification of the gels, ammonium persulfate to a final concentration of 0.5 mg/ml and N,N,N',N'-Tetramethylethylenediamine (TEMED) to a final concentration of 0.5 μ l/ml were added immediately prior to casting. Gels were run with constant power using either an EC703 power supply (E-C Apparatus Corp., St. Petersburg, FL, USA) or an LKB 2297 macrodrive S power supply (LKB). Gels were pre-rur before samples were loaded to allow the gels to reach an optimal temperature for separation, usually between 45-55°C. Samples were denatured at 95°C for 2-5 min immediately prior to loading.

Once the samples were run for the desired time, the gels were immobilized on sheets of Whatman 3MM chromatography filter paper and dried at 80°C for 1 hr on a model SE1160 slab gel dryer (Hoefer Scientific, San Francisco, CA, USA). The dried gels were placed in X-ray film cassettes and exposed to Kodak XAR5® (Kodak) or BioMax® (Kodak) X-ray film, usually overnight. In some instances longer exposures were necessary.

The sequence data was read from the developed film and entered into either the Microgenie computer analysis system (Beckman) or the Wisconsin Sequence Analysis Package

(Genetics Computer Group (GCG), Madison, WI, USA) for computer analysis. Computer analysis of the rRNA gene sequences in Chapter 4, was carried out using the Wisconsin Sequence Analysis Package. Phylogentic comparisons were carried out using the pileup program of the Wisconsin Sequence Analysis Package. This program will align multiple sequences by the progressive alignment method of Feng and Doolittle (1987). The pileup program is also able to plot the comparisons in the form of dendograms. In Chapter 6, the DNA sequence of the two Campylobacter probes, pDT1719 and pDT1720, was entered and assembled using Microgenie. Analyses of the sequences and homology searches were carried out with the Basic local alignment search tool (BLAST; Altshul et al., 1990).

E.4. Construction of deletion mutants for sequencing

Deletion mutations of some clones of the 23S/5S genes were carried out to obtain sequential deletions of the 23S/5S genes for sequencing. The deletions were generated using the Promega Erase-a-base system following protocols recommended by the manufacturer with the following changes. Instead of 25 30-second time points, 6 30-second time points were used for deletions. Reagent volumes were adjusted accordingly. Deletion reactions were carried out at 34°C with 400 units of enzyme/5 μ g of DNA. At this temperature 250-350 bases/min were deleted. A reaction volume of 30 μ l was used and 5 μ l aliquots were taken for each time point and added to 15 μ l of

S1 nuclease mix. Eighty μ l of ligase mix were then added to each sample.

F. RNA techniques

F.1. Isolation of total RNA

Total RNA was isolated using a modification of the methods of Konkel et al. (1994). All solutions used in this section were made up in deionized water treated with diethyl pyrocarbonate (DEPC; Aldrich Chemical Co., Milwaukee, WI, USA; refer to Appendix A under DEPC-water). H. pylori cultures were spread on solid media and grown as described above. A sterile loop was used to resuspend the cells in 500 μ l of protoplast buffer (Appendix A). suspended cells were divided into two microfuge tubes and 250 μ l of 5% SDS were added. The cell suspensions were incubated for 15 min at 70°C. The cell suspensions were then cooled to room temperature and extracted twice with 300 μl phenol. The aqueous layer was then extracted once with 300 μ l phenol:chloroform (1:1) and twice with 300 μ l chloroform. Following this, 250 μ l of 5.0 M NaCl were added and the suspensions were incubated on ice for 10 min. The precipitate was removed by centrifugation for 2 min in a microfuge and the supernatant was divided into two new microfuge tubes. One ml of chilled 95% ETOH was added and the DNA was spooled out, rinsed in 95% ETOH and dissolved in 300 μ l of TE. The remaining supernatant was incubated at -20°C for at least 2 hr. The RNA was collected in a microfuge and allowed to air dry. The RNA pellets were then dissolved in a final volume of 200 μ l of DEPC-water. This solution was then treated with RNase-free DNase I (BMC) and the enzyme was removed by centrifugation of the samples through a Millipore ultrafree®-probind 0.45 μ m filter unit. The samples were de-salted by centrifugation through Chromaspin 100 columns (Clontech, Palo Alto, CA, USA) equilibrated with DEPC-water.

F.2. Denaturing RNA agarose gels

The purity of the RNA samples was checked by electrophoresis in 1.2% denaturing agarose gels (Appendix A) in 1 x MoPs buffer (Appendix A). RNA samples were prepared for electrophoresis by adding 10 μ l of denaturing mix (Appendix A) and 2 μ l of loading dye to 5 μ l of RNA sample and incubating at 65°C for 10 min. After electrophoresis the gel was stained in ethidium bromide for 20 min and destained in dH₂O for at least 1 hr. The RNA could then be transferred to nylon membranes, following normal Southern transfer procedures, without denaturation or neutralization. The RNA was cross-linked to the nylon membrane in a Stratalinker® UV crosslinker oven (Stratagene).

F.3. Primer extension

Primer extension was carried out as described by Newnham and Taylor (1994), using superscript II reverse transcriptase (BRL) with the following changes. Instead of incorporating $[\alpha^{-35}S]$ -dATP, $[\gamma^{-33}P]$ end-labelled primers were used. End-

labelling was carried out as described in the Cycle Sequencing System (BRL) protocol.

G. SDS-PAGE and Western blots

Crude preparations of the MOMP of C. jejuni UA580 were subjected to SDS-PAGE analysis and proteins were blotted to SDS-PAGE was carried out as previously PVDF membranes. Samples were electrophoresed described (Laemmli, 1970). through a 5% stacking gel (Appendix A) and a 10% separating gel (Appendix A) in a Bio-Rad Mini-PROTEAN II cell in SDS-PAGE running buffer (Appendix A). Prior to loading the protein samples, 1/10th volume of SDS-PAGE running dye (Appendix A) was added to the samples and the samples were boiled for 1-2 min. Proteins were run with constant current of no greater than 40 mA. When sufficient separation was obtained, the proteins were then electro-blotted to PVDF membranes (Matsudaira, 1987). Electro-blotting was carried out in a Bio-Rad apparatus in CAPS buffer (Appendix A) with constant current (300 mA) for 30 min. The membrane was then immersed in fresh staining solution (Appendix A) for 5 min. The membrane was then destained with several changes of destaining solution (Appendix A), rinsed with dH2O several times and allowed to dry in air. The stained membrane could then be stored in a sealable plastic freezer bag indefinitely The stained PVDF membranes were sent to the at -20°C. Alberta Peptide Institute (API, Department of Biochemistry, University of Alberta, Edmonton, Canada) and the 45 kDa MOMP protein was excised from the membrane for N-terminal amino acid sequence analysis.

Chapter 3

Construction of genomic libraries of H. pylori.

A. Introduction

An objective of this research was to construct genomic libraries of H. pylori DNA which would provide a source of cloned H. pylori DNA fragments from which specific genes or regions of DNA could be isolated. Such libraries will become increasingly useful as the mechanisms of pathogenesis of H. pylori and the factors involved are delineated. discussed in Chapter 1, a high degree of genomic variability has been observed between different strains of H. pylori; this variability may be related to the different clinical symptoms of H. pylori infection. Isolation and comparisons of virulence factors from various strains may provide us with information about the mechanisms of the different clinical presentations of H. pylori infection (ie. gastritis, ulceration). The availability of genomic libraries for different strains of H. pylori would facilitate these studies. In addition the genomic libraries can be used to clone other genes or regions of interest, such as the rRNA genes, to further our understanding of this organism.

A major consideration for the construction of genomic libraries was the selection of a vector system. A wide variety of plasmid, cosmid and lambda vectors are available, each having its wan advantages and disadvantages. The selected vector system should produce genomic libraries which

would provide a convenient source of DNA for isolation and characterization of genetic elements.

A.1. Plasmid vectors

There are many convenient plasmid vectors, with a variety of selection and screening markers, that are commercially available. These vectors are useful for efficient cloning of fragments from 0-10 kb in size (Maniatis et al., 1982). The number of clones needed to statistically represent 99% of a genome (N) can be calculated using the formula

(Maniatis et al., 1982). One assumption made in this formula is that all parts of the genome are equally represented in the array of fragments to be cloned. In practice this is not always true. Toxicity of cloned elements and the restriction modification system of the host strain may make some regions difficult to clone. In addition, larger fragments may not be cloned as efficiently and could be poorly represented. To circumvent these problems, construction of plasmid libraries using a number of different restriction enzymes would be necessary.

The size of the *H. pylori* strain UA802 genome has been estimated to be approximately 1.7 Mb, as determined by Taylor et al. (1992a) using PFGE. By using the above formula, with the average insert size as 5 kb and the genome as 1.7 Mb, a minimum of 1563 clones would be needed to represent 99% of

the genome. This figure (1563) represents the optimal number of clones needed assuming all regions of the chromosome were represented equally. As mentioned, this is not always the case, and more than one library would likely need to be constructed, using a variety of restriction enzymes. An increase in the number of clones screened, beyond what is needed to statistically represent the genome, would decrease the chances of missing some of the regions that are difficult to clone.

Because a minimum of 1563 clones is needed to represent the genome, and because of the logistical problems of long-term storage of such a large number of clones, plasmid libraries are generally used for cloning specific genes. Since the objectives of this research included the isolation of specific genes such as the P-type ATPase and the rRNA genes, the construction of plasmid libraries was attempted.

A.2. Cosmid vectors

Cosmid vectors are very convenient for the rapid construction of genomic libraries, as large fragments (up to 50 kb) of DNA can be cloned. Cosmid vectors combine convenience of plasmid vectors with the cloning capacity of lambda vectors by incorporating the lambda cos (cohesive end) site into plasmid vectors. The cos site is the signal recognized by the lambda packaging machinery which packages DNA of appropriate length into phage heads. The maximum and minimum sizes of DNA which can be packaged to produce viable phage particles are approximately 53 kb and 37 kb

respectively (Chauthaiwale et al., 1992). The large size of the fragments which can be cloned in cosmid vectors means that far fewer clones are needed to statistically represent a given genome. However, the larger size of fragments also increases the chance of cloning toxic regions. Thus, as with the plasmid vectors, the chromosome may not be equally represented.

Since far fewer clones are needed to represent the genome using cosmid libraries, these libraries can easily be cryogenically stored and re-screened at later dates with probes as they become available. In fact, in a recent review, Fonstein and Haselkorn (1995) recommended the use of cosmid vectors for construction of genomic "encyclopedias" of organisms which could be used as convenient sources of DNA for isolation and characterization of genes.

A.3. Lambda vectors

Many lambda cloning systems are now available commercially. These vectors are classified as either insertion or replacement vectors. With insertion vectors, the passenger DNA is inserted into a non-essential region of the vector. Some vectors include the *lacZ/blue-white* selection cassette for added convenience in screening. Since the maximum size of DNA which can be packaged to produce viable phage particles is approximately 53 kb (Chauthaiwale et al., 1992), insertion vectors are limited to cloning DNA fragments of 0-10 kb (Maniatis et al., 1982). The minimum number of clones needed to represent 99% of a genome is the same as

with plasmid vectors (1563). However, the screening, storage and handling of lambda libraries is easier than plasmid or cosmid libraries.

The size of inserts which can be cloned in replacement vectors is significantly greater than can be cloned in insertion vectors. This is because the passenger fragments replace a non-essential region of the lambda DNA which is approximately 14 kb in size. Thus, inserts of 9-23 kb can be cloned into most replacement vectors. This makes them quite convenient for the construction of genomic libraries, as the number of plaques to be screened is reduced when compared with the insertion vectors (487 compared to 1563). In addition, lambda libraries can be stored as phage suspensions for several years, and can be screened repeatedly with probes as they become available.

B. Materials and Methods

Please refer to Chapter 2 for detailed methodology used in library construction.

C. Results

C.l Construction of plasmid libraries of H. pylori UA763

Initial work on the production of genomic libraries of H. pylori DNA was conducted using the plasmid vector pUC20. Chromosomal DNA from H. pylori UA763 was digested with the restriction enzymes $Bg^{7}II$ or SstI. This DNA was cloned into

puc20 and transformed into a variety of $E.\ coli$ host strains. Table 3-1 shows the results of transformation of $E.\ coli$ host strains DH5 α or DH5 α MCR with puc20 DNA ligated with $H.\ pylori$ UA763 DNA. Transformations were repeated several times and the best results obtained are presented. The $E.\ coli$ DH5 α and DH5 α MCR competent cells were obtained commercially from BRL. The expected transformation efficiencies of these competent cells were 10^6-10^8 colonies/ μg of plasmid DNA (BRL), while the competent cells made in the laboratory were expected to give efficiencies of 10^6-10^7 colonies/ μg of plasmid DNA (Maniatis et al., 1982). Construction of plasmid libraries of $H.\ pylori$ UA763 was also attempted using EcoRI, however, for reasons which could not be determined, no recombinants were observed.

With the BglII and SstI puc libraries transformed into the DH5 α host, the number of transformants obtained was extremely low (120-500 and 160-400 colonies/ μ g transformed DNA respectively; Table 3-1). These white colonies were screened with radioactively-labelied H. pylori UA763 DNA, and only a few of the clones hybridized (5/42 BglII colonies using the E. coli DH5 α strain; 7/65 SstI colonies using the E. coli DH5 α strain; 57/552 DglII colonies using the E. coli DH5 α MCR strain; 52/723 SstI colonies using the E. coli DH5 α MCR strain). Mini-preparations of plasmid DNA showed that the non-hybridizing white colonies harbored plasmids equal in size to or slightly smaller than wild-type pUC20

Table 3-1. Transformation efficiency of plasmid DNA in E. coli DH5 α and DH5 α MCR strains

		rick scrains		
Insert DNAª	Host strain	No. of coloniesb	Amount of DNA used for transformation	Transformants per µg DNA
None- EcoRI + control	DH5α	342 ^c	0.1 ng	3.42 x 10 ⁶
None- EcoRI - control	DH5α	8.0°	100.0 ng	80
None- EcoRI + control	DH5α MCR	271.0°	0.1 ng	2.71 x 10 ⁶
None- EcoRI - control	DH5α MCR	13.0°	100.0 ng	130
<i>H. pylori</i> UA763 <i>Eco</i> RI	DH5a	0	0.1 ng	0
<i>H. pylori</i> UA763 <i>Eco</i> RI	DH5α	0	1.0 ng	0
<i>H. pylori</i> UA763 <i>Eco</i> RI	DH5α	0	10.0 ng	0
H. pylori UA763 BglII	DΗ5α	0.3 (0)	1.0 ng	3.0×10^2
H. pylori UA763 BglII	DH5α	5.0 (1)	10.0 ng	5.0 x 10 ²
E. pylori UA763 BglII	DH5α	12.0 (4)	100.0 ng	1.2 x 10 ²
H. pylori UA763 SstI	DH5α	1.6 (0)	1.0 ng	1.δ x 10 ³
H. pylori UA763 SstI	DH5α	4.C (1)	10.0 ng	4.0 x 10 ²
H. pylori UA763 SstI	DH5α	18.0 (6)	100.0 ng	1.8 x 10 ²
H. pylori UA763 BglII	DH5α MCR	56.0(19)	1.0 ng	5.6 x 10 ⁴
H. pylori UA763 BglII	DH5a MCR	128.0(38)	10.0 ng	1.28 x 10 ⁴
H. pylori UA763 BglII	DH5α MCR	too many to count	300.0 ng	
H. pylori UA763 SstI	DH5a MCR	58.0(19)	1.0 ng	6.8 x 10 ⁴
H. pylori UA763 Ss I	DH5α MCR	173.0(33)	0.0 ng	1.73 x 10 ⁴
H. pylori UA763 SstI	DH5α LICF	too many to count	? 0 ng	

a- puc20 DNA was cut with appropriate endonuclease and ligated to insert DNA. No ligase was added to -controls

b- average of three plates. Numbers in brackets represent the number of colonies which hybridized with labelled H. pylori DNA

c- number of blue colonies on a single plate were counted for the negative controls

with no insert. This indicated that either small deletions were occurring in the multiple cloning site of pUC20 or that the H. pylori DNA was not stable and was Deing deleted.

Use of the commercially prepared competent DH5lphaMCR strain allowed for a greater transformation efficiency than the other E. coli host strains. This strain has been highly altered genetically to reduce the effects of the restriction modification barrier. Even so, the number of clones which carried H. pylori DNA was far below the expected number of transformants/ μg of DNA. The manufacturer of these competent cells states that the optimal transformation of wild-type, uncut pUC20 would yield 1 x 10^8 colonies/ μ g of transformed DNA (BRL catalog 1993-94). The values obtained for the re-ligation of digested vectors (positive control) was 2.7 X 10^6 colonies/ μ g. This was within the acceptable range of 10^6 - 10^8 colories/ μ g of plasmid DNA (BRL catalog 1993-94). The number of recombinants observed for the genomic libraries were, in contrast, 4-6 orders of magnitude lower than optimal.

Since no efficient plasmid libraries could be obtained, construction of cosmid libraries was attempted. For this research the cosmids pSa747 (Tait et al., 1983) and pHC79 (BMC) were used to construct genomic cosmid libraries of H. pylori. Table 3-2 shows the highest transfection efficiency obtained using these cosmids. Variation of the vector:passenger ratios, variation of the amounts of DNA

Table 3-2. Efficiency of cosmid libraries in E. coli LE392

				
Insert DNAª	Vector	No. of colonies hybrid-izing to H. priori	Amount of packaged DNA used/plate for transfection	CFU ^c /μg D NA
H. pylori UA763 BglII	pSa747	4 (37)	3 μl (30 ng)	4.1î x 10 ²
H. pylori UA763 Sau3AI	pSa747	8 (72)	3 µl (30 ng)	4.66 x 10 ²
H. pylori UA802 EglII	pSa747	7 (56)	3 μl (30 ng)	6.22 x 10 ²
H. pylori UA802 Sau3AI	pSa747	11 (93)	3 μ1 (30 ng)	1.03 x 10 ³
H. pylori UA763 BglII	рнс79	18 (132)	3 μl (30 ng)	1.47 x 10 ³
H. pylori UA763 Sau3AI	рнс79	2 (147)	3 μl (30 ng)	1.63 x 10 ³
H. pylori UA802 BglII	рнс79	11 (133)	$3 \mu 1 (30 \text{ ng})$	1.48 x 10 ³
H. pylori UA802 Sau3AI	рнс79	12 (184)	3 μ1 (30 ng)	2.04 % 103

a- partial digests of genomic H. pylori DNA were ligated to

vector cut with .amHI
b- total of 3 plates. Numbers in brackets represent total number of colonies observed

c- colony forming units

packaged, and purification of ligated DNA did not improve these results. With wild type lambda packaged in vitro, the optimal expected plaque forming units (PFU) would be 1 x 10^9 plaques/ μ g of packaged DNA (Stratagene catalog 1992, p.42). The transfection frequencies were far below what were expected for optimal transfection (133-267 CFU/ μ g packaged DNA for pSa747 clones; 67-600 CFU/ μ g packaged DNA for pHC79 clones; Table 3-2).

C.3. Construction of lambda genomic libraries of H.

The construction of lambda libraries of H. Pylori DNA was started when problems in cosmid library construction became apparent. Initially the lambda vector λ -GEM11 (Promega) was used to construct libraries of H. Pylori UA763 and UA802 genomic DNA, nowever, the efficiency of these libraries seemed low. The vector λ -DASH II (Stratagene) was suggested as an alternative lambda vector which had been used successfully in other labs (Dr. J. Elliot, personal communication). This vector was used to construct genomic libraries of H. Pylori UA802. H. Pylori UA802 was selected because a genomic restriction map of this strain had been constructed in our laboratory (Taylor et al., 1992a).

Transfections of the λ -GEM11 libraries into the *E. coli* strain KW251 gave unsatisfactory results (258-410 PFU/ μ g packaged DNA for λ -GEM11; Table 3-3). In addition to the low number of plaques observed, the size of the plaques was smaller than those observed on control plates (0.5 mm

Table 3.3. Transfection efficiency of lambda libraries

Insert DNAª	E. coli host strain	Vector	No.of plaquesb	PFU ^c /µg packaged DNA
H. pylori UA763 BglII	KW251	λ-GEM11	138	276
H. pylori UA763 Sau3AI	KW251	λ-GEM11	205	410
H. pylori UA802 BglII	KW251	λ-GEM11	156	312
H pylori UA802 Sau3AI	KW251	λ-GEM11	129	258
H. pylori UA802 BglII	SRB/ SRB-P2	λ-DASH II	378 ^d	1.23 x 10 ⁴
H. pylori UA802 Sau3AI	SRB/ SRB-P2	λ-DASH II	366 ^e	1.22 x 10 ⁴

- a- partial digests of *H. pylori* genomic DNA were ligated with <code>PamHI</code> lambda vector arms
- b- average of 3 plates. For each plate 50 μl (~500 ng packaged DNA) of undiluted phage suspension were used for transfection
- c- plaque forming units
- d- average of 10 plates. For each plate 3 $\mu 1$ (~30 ng packaged DNA) of undiluted phage suspension were used for transfection
- e- average of 4 plates. For each plate 3 μl (~30 ng packaged DNA) of undiluted phage suspension were used for transfection

compared to 1-2 mm). Plaque-lifts were hybridized with radioactively-labelled H. pylori DNA. Because of the small size and low number of the plaques, the results were difficult to interpret. It did appear that all of the plaques hybridized with the radioactively-labelled H. pylori DNA. The figures presented in Table 3-3 for the λ -GEM11 libraries represent the best values obtained with this lambda system. A number of parameters were varied during these attempts to make a library including variation of vector:passenger ratios, adjustment of total amounts of DNA to be packaged, and purification of ligated DNA through DNA purification columns (Chroma-spin TE 1000), all with similar results.

Results for the λ -DASH II libraries of H. pylori UA802 transfected into E. coli SRB were considerably better than those observed for the $\lambda\text{-GEM11}$ libraries (1.22-1.23 x 10^4 PFU/ μ g packaged DNA; Table 3-3). The size of the plaques observed (1-3 mm) was also increased. In addition, hybridization of plaque-lifts of these plates clearly showed that all of the plaques contained H. pylori DNA. The numbers obtained were still far below those expected for construction As mentioned above, an optimal of lambda libraries. efficiency of 1 x 10 9 plaques/ μ g of packaged DNA was expected. Even though the values obtained were low, there were sufficient PFU to statistically represent several genomes. A minimum of 487 clones was calculated to represent 99% of the genome, and approximately 2000 plaques were obtained for each library.

D. Discussion

D.1. Plasmid libraries

Since downstream characterization of genes cloned in plasmid vectors is much more convenient than in cosmid or lambda vectors, construction of plasmid libraries of H. pylori UA763 was attempted. However, no efficient libraries using pUC20 as a vector were produced from this The results showed that less than 10% of the colonies contained H. pylori DNA. The highest transformation rates for the plasmid libraries of H. pylori DNA were obtained using the E. coli host strain DH5aMCR. This strain carried mutations of the restriction modification system (Table 3-4). This is the system by which foreign DNA is recognized by the host and destroyed. In E. coli several genes are involved in this complex system (Bickle and Krüger, Thus, it appeared that the restriction modification barrier was one of the factors responsible for difficulties observed in construction of plasmid libraries. Even with the DH5aMCR E. com strain, however, the efficiency of library construction was still low, suggesting that other factors could also be involved.

These results correlated with those obtained in a recent study of the choice of host strains for the construction of H. pylori genomic lambda libraries (Phadnis et al., 1993). The mcrC-hsdRMS-mrr loci seemed to be of particular

Tab' ~ 3-4. Restriction phenotypes of some E. coli strains

Strain	Restriction Phenotype	Source
JM83	mcrA+ mcrBC+ hsdR+ hsdM+ hsdS+ mrr+	BRL
DH5a	mcrA+ mcrBC+ hsdR- hsdM+ hsdS+ mrr+	BRL.
DH5@MCR	mcrA- mcrBC- hsdR- hsdM- hsdS- mrr-	BRL
LE392	mcrA- mcrBC+ hsaR- hsdM+ hsdS+ mrr+	вмс
KW251	mcrA- mcrBC- hsdR- hsdM+ hsdS+ mrr+	Promega
SRB	mcrA- mcrBC- hs@R- hsdM- hsdS- mrr-	Stratagene
SRB P2	mcrA- mcrBC- hsdR- hsdM- hsdS- mrr-	Stratagene
нв101	mcrA+ mcrBC- hsdR- hsdM- hsdS- mrr-	BRL
ER1793	mcrA+ mcrBC- hsdR- hsdM- hsdS- mrr	NEB

- mcrA encodes restriction of DNA with cytosine methylated at the sequence 5'-CmeCGG-3'(Raleigh and Wilson, 1986)
- mcrBC- encodes restriction of DNA with cytosine methylated at the sequence 5'-GmeC-3' (Raleigh and Wilson, 1986)

- mrr encodes restriction of DNA with adenosine methylated at the sequences 5'-CmeAG-3' or 5'-GmeAC-3' (Waite-Rees et al., 1991)

importance, as lambda libraries constructed in strains carrying mutations in these alleles had the highest efficiency. These libraries produced approximately 2000 plaques from 5 μ l of undiluted phage suspension. Although it was not stated in the study, standard packaging reactions require up to 5 μ g of DNA in total and yield approximately 500 μ l of suspension. Most commercial in vitro packaging systems will give 1 x 10^7 -1 x 10^9 plaques/ μ g of packaged DNA. Thus, the observed efficiency of the libraries constructed by Phadnis et al., using the E. coli host strain ER1793, could be estimated to be 8 x 10^4 - 4 x 10^5 plaques/ μ q packaged DNA, which was still far below the expected efficiency for such a This suggested, as did the results we this research, that other factors must also have contributed to decreased library efficiency.

The results of the cloning of *H. pylori* DNA presented here were similar to those obtained with *Streptococcus pneumoniae* DNA. In the construction of genomic libraries of *S. pneumoniae*, it was found that cloned *S. pneumoniae* DNA was highly unstable in *E. coli* when using high copy number vectors such as pBR325 (Chen and Morrison, 1987). Morrison and Jaurin (1990) first suggested that the low G + C content of the *S. pneumoniae* DNA (39 mol.%; Marmur and Doty, 1969) resulted in a high incidence of *L. coli* promoter-like sequences, which had strong enough comoter activity to interfere with the stability of the cloned DNA. This hypothesis was a ser contradicted in a study showing that

promoters of sufficient strength to cause plasmid instability were probably rare and that even sequences resembling relatively strong E. coli promoters did not seem to affect clone stability (Dillard and Yothers, 1991). Although sequence-related effects could not be ruled out in the later study, Dillard and Yothers concluded that problems associated with plasmid genomic library construction of S. pneumoniae were attributable to multiple effects and not specifically to promoter-like sequences. In addition to the sequence-related effects, cloning of lethal elements and vector-specific problems (copy number) were also cited as possible contributors to the problem. Since H. pylori has a lower G + C content (35.2 mol%; Béji et al., 1988) than that of S. pneumoniae, sequence-related factors, such as E. coli promoter-like sequences, could interfere with genomic library construction. Cloning of lethal elements and plasmid-related effects are also factors which could inverfere with H. pylori library construction, analogous to the situation with S. pneumoniae.

Of all the researchers who constructed genomic libraries of H. pylori, only two groups used plasmid vectors. Plasmid libraries were used to clone the cagA (Covacci et al., 1993) and vacA (Telford et al., 1994: genes. These groups used pBluescript (Stratagene), a phagemid derivative of pUC19, and screened 70,000-85,000 recombinants to obtain their clones. Even with this number of clones neither the cagA nor the vacA were cloned as intact genes but were cloned as overlapping

fragments. These genes probably represented elements which were toxic to the host strains and thus could not be cloned as entire genes.

Some of the genes clored for this research (rRNA genes) have homologs in the host strains. In the case where the host chromosome might hybridize to the probe used for screening, plasmid isolations of each individual clone would be necessary. Thus, use of screening procedures such as those used by Covacci et al. (1993) and Telford et al. (1994) would have been impractical for this work, as it was not feasible to isolate and screen plasmid DNA from 70,000-85,000 clones. Other studies in which plasmid vectors were used in the cloning of H. pylori genes involved cloning of specific restriction fragments from genomic digests (Leying et al., 1992; O'Toole et al., 1991; Smith et al., 1994).

Efficient genomic DNA libraries of *H. pylori* could not be produced for this research using the plasmid vector pUC20. Restriction modification was one of the barriers which had to be overcome in the construction of plasmid libraries of *H. pylori* DNA, as indicated by the increased number of recombinants obtained using the DH5aMCR host strain. Other factors, such as those associated with sequence-specific interference (promoter-like sequences), cloning of lethal elements, and vector related problems (copy number), may also have contributed to difficulties observed in plasmid library construction, similar to problems observed by other researchers (Chen and Morrison, 1987; Covacci et al., 1993;

Telford et al., 1994). It appears, therefore, that plasmid vectors may not be well suited for genomic library construction of *H. pylori* DNA.

D.2. Cosmid library construction

When it became apparent that plasmid libraries were not adequate for the purposes of this research, the construction of cosmid libraries was begun in 1990. Since Clayton et al. ssfully used pHC79 to clone urease genes (1989a) had the cosmid pSa747 was successfully used from H. pyl Azotobacter vinelandii nitrogen fixation for the clc genes (Hiratsuka, 1987), these two vectors were chosen for library construction. Unfortunately, an efficient cosmid library of H. pylori DNA was not constructed because, as with the plasmid libraries, only a small percentage of the clones carried H. pylori DNA. This was probably due to the E. coli host strain used for growth and propagation of the cosmid While the The E. coli strain used was LE392. libraries. LE392 strain has been mutated for some of the restriction sodification functions (mcrA hsdR), it still retains some restriction modification properties (mcrBC+ hsdMS+ mrr+)(Table 3-4). As mentioned, mutations at the mcrC-hsdRMS-mrr loci are critical for efficient library construction of H. pylori In fact, Phadnis et al. (1993) found LE392 to be the DNA. least efficient strain for construction of lambda libraries. The transfection efficiencies of the cosmid libraries in an (mcrC-hsdRMS-mrr) background were not tested for this

research. This was because, by this time, I had constructed H. pylori genomic libraries using the lambda vector $\lambda DASH$ II, and therefore, further work on cloning was concentrated on the lambda libra: The cosmid pHC79 has been used successfully by a number of groups (Clayton et al., 1989a; Spiegelhalder et al., 1993; Moore et al., 1995). All of these groups used the E. coli strain HB101 for growth and propagation of their libraries. The strain HB101 carries a (mcrC-hsdRMS-mrr) mutation while retaining the mcrA+phenotype.

While it was possible that the efficiencies of the cosmid libraries might have been greater if they had been transfected into a (mcrC-hsdRMS-mrr) host, other factors may also have contributed to the low number of recombinants observed for the cosmid ibraries. Factors mentioned previously which could affect the construction of plasmid libraries (low G + C effects, lethal alleles) would be magnified in cosmid clones due to the size of the inserted DNA. Evidence to support this hypothesis was obtained during the construction of a physical genetic map of H. pylori using an ordered cosmid library (Bukanov and Berg, 1994). library was successfully constructed using the cosmid vector Lorist6 in the strain DH5 α . This host strain carried a mutation in only the hsdM loci and was not mutated for any other restriction modification function (Table 3-4). However, the vector was modified by the addition of rhodependent transcription terminators bracketing the cloning site. Such modifications have been used to counteract the instability of cloned DNA associated with the presence of promoter or promoter-like sequences (Chen and Morrison, 1987). In addition, the replication of this vector was regulated by the phage λ replication system. Under this system the copy number of the recombinant cosmids was far more stable that of the standard ColEl replication system. Using this vector Bukanov and Berg were able to construct a genomic library of H. pylori NCTC11638.

The results described here supported observations that LE392 was not an efficient host for *H. pylori* DNA library construction (Phadnis et al., 1993). Results of Bukanov and Berg (1994) indicated that genomic libraries of *H. pylori* DNA could be constructed independent of restriction modification using the vector Lorist6. Problems associated with lethal alleles would still be present. However, construction of more than one library using a variety of enzymes should overcome these problems. In addition, use of a (mcrc-hsdRMS-mrr)- host may improve library efficiency.

D.3. Lambda library construction

The construction of lambda genomic libraries of *H. pylori* DNA was begun when difficulties with the cosmid libraries were encountered. At this time a restriction map of *H. pylori* UA802 had been constructed in this lab (Taylor et al., 1992a), so efforts to construct genomic libraries of *H. pylori* were concentrated on this strain. The efficient construction of lambda libraries of *H. pylori* genomic DNA

proved to be difficult. No useful H. pylori DNA libraries were constructed using λ -GEM11 in the host KW251. libraries of H. pylori UA802 DNA were successfully constructed using the vector $\lambda\text{-DASH}$ II in the host strain SRB, but with low efficiencies. These results were consistent with the later findings of Phadnis et al. (1993) that the E. coli strains KW251 and SRB were relatively poor hosts for genomic library construction of H. pylori DNA. Phadnis et al. obtained 2 plaques/ μ l of phage suspension using E. coli KW251 as the host. Similarly, using E. coli KW251 as the host for the H. pylori UA802 libraries, three plaques/ μ l of phage suspension (an average of 155.5 plaques/50 μ l) were obtained. With E. coli SRB, al. obtained 4 plaques/ μ l of phage suspension. research, using the host E. coli SRB, 124 plaques/ μ l of phage suspension were obtained (an average of 372 plaques/3 μ l). The differences observed in the efficiency of the SRB host may be a reflection of the different lambda vectors used. Phadnis et al. used the vector λ -GEM11 (Promega), while I used λ -DASH II (Stratagene).

Other researchers have also reported difficulties in construction of lambda libraries of *H. pylori* DNA. Clayton et al. (1989a) reported difficulty using another lambda expression vector λ EMBL3. Although sufficient numbers of recombinants were screened, the libraries they constructed were not representative of the entire *H. pylori* genome. Similar findings were made by Covacci et al. in 1993, in

their attempt to clone the cagA gene using the vectors λ - σ t11, λ -EMBL3 and λ -DASH. This gene could not be cloned using λ -EMBL3 and λ -DASH and only portions of the cagA were cloned using λ -qtll. Another example of a library which was not representative of the entire genome was that used in the construction of the ordered cosmid map of H. pylori NCTC11638 (Bukanov and Berg, 1994). This map contained three gaps for which cosmid recombinants could not be isolated. Ιn addition, there were three sequences which were poorly represented in the cosmid library, one of which was cagA, which had been shown by earlier studies to be difficult to clone (Covacci et al., 1993; Tummuru et al., 1993). suggested that certain regions may harbor genetic elements lethal to E. coli or may be regions where the effects of E. coli promoter-like sequences were of sufficient strength to interfere with clone stability.

E. Conclusion

Using the lambda vector λ -DASH II, genomic libraries of H. pylori UA802 DNA were constructed. These libraries will serve as a source of cloned H. pylori for the isolation and characterization of genes from this organism. The construction of genomic libraries of H. pylori DNA was also attempted using other vectors. The results of these attempts indicated that restriction modification was a factor that needs to be addressed for the production of genomic libraries of H. pylori DNA.

Chapter 4

Isolation and characterization of the P-type ATPase and rRNA genes from the genomic library of H. pylori strain UA802

A. Introduction

A.1. P-type ATPase of H. pylori

mentioned in Chapter 1, several substituted benzimidazole compounds have been used in the treatment of Tn se compounds block the peptic and duodenal ulcers. sulfhydral-groups of gastric P-type ATPases and act as proton pump inhibitors to reduce the severity of ulceration (Fellinius et al., 1981; Lindberg et al., 1986; Iwahi et al., The bacteriostatic, and in some instances 1991). bactericidal, activity of these compounds on H. pylori suggested the presence of a P-type ATPase. Described in this chapter are the results of research showing the existence of a P-type ATPase in H. pylori and the cloning of the genes encoding this ATPase from genomic libraries of H. pylori UA802 DNA.

A.2. The 23S and 5S genes of H. pylori UA802

Results for the isolation of rRNA genes and characterization of the 23S and 5S genes of *H. pylori* UA802 are also presented in this chapter. Sequence data for the complete 16S rRNA genes of several *H. pylori* strains have already been published (Eckloff et al., 1994). The 23S and 5S genes of *H. pylori* have not been extensively

characterized. The phylogenetic relationships of a variety of Campylobacter spp. and related organisms, including H. pylori, were investigated by Vandamme et al. (1991). In that study, DNA-rRNA hybridizations using radioactively-labelled 23S rRNA molecules were carried out. Vandamme et al. gave indirect evidence that H. pylori was included in the epsilon subdivision of the class Proteobacteria, along with Campylobacter spp., Arcobacter spp., Wolinella spp, Flexispira spp. and other Helicobacter spp. The phylogenetic position of H. pylori was further investigated in this research through the characterization of the 23S and 5S genes.

B. Materials and Methods

Please refer to Chapter 2 for detailed methodology.

C. Results

C.1. Cloning of a P-type ATPase of H. pylori UA802 Evidence for the existence of a P-type ATPase in H. pylori

Before cloning of genes encoding the P-type ATPase of H. pylori was begun, preliminary hybridizations were carried out to show that such genes existed in the genome of H. pylori. DNA from H. pylori strains UA802 and UA763, digested with BglII, was fractionated on agarose gels and transferred to nitrocellulose as described (Chapter 2, B.2). The membranes were probed with cloned radioactively-labelled

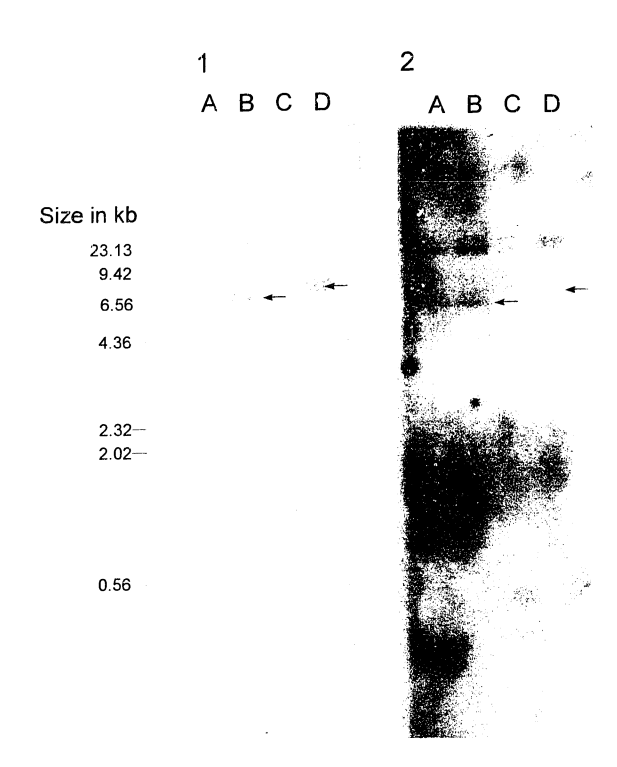
P-type ATPases from three heterologous sources (Figure 4-1). The cloned genes used as probes were the E. coli kdpABC genes (Hesse et al., 1984), the Enterococcus hirae (formerly Streptococcus faecalis) K+-ATPase gene (Solioz et al., 1987), and a rat stomach (H+-K+)-ATPase gene (Shull and Lingrel, 1986). Figure 4-1. shows Southern analysis of two strains of H. pylori, UA763 and UA802, probed with the cloned E. coli kdpB gene and E. hirae K+-ATPase gene. Hybridization was very weak and X-ray film had to be exposed to the probed blots for 2 weeks to obtain the results shown in Figure 4-1. Similar results were obtained with the rat (H++K+)-ATPase probe (data not shown). These results indicated that both H. pylori UA802 and UA763 encoded P-type ATPAses in their genomes.

Screening of the $\lambda\text{-DASH}$ genomic libraries of H. pylori UA802 with P-type ATPase probes

Based on the hybridization data, the presence of genes encoding a P-type ATPase in *H. pylori* was established. In order to isolate these genes, PCR probes were produced for screening of the λ-DASH genomic libraries of *H. pylori* UA802 DNA. From amino acid sequence alignments of several P-type ATPases, some regions were observed to be highly conserved (Figure 4-2). Primers for PCR were designed from two of these regions. The primers are shown in Figure 4-2. These primers were used to amplify a PCR probe from *H. pylori* UA802 genomic DNA. A PCR product of approximately 600 bp was amplified from genomic DNA of *H. pylori* UA802

Figure 4-1. Hybridization of restriction digestions of *H. pylori* strains UA763 and UA802 with cloned P-type ATPases

Panel 1 shows hybridization using the cloned *E. hirae* K⁺-ATPase probe. Panel 2 shows hybridization using the cloned *E. coli kdp*B probe. Lanes A and B in both panels were 1 μ g and 5 μ g respectively of *H. pylori* UA802 digested with *BglII*. Lanes C and D in both panels were 1 μ g and 5 μ g respectively of *H. pylori* UA763 digested with *BglII*. Arrows indicate hybridizing fragments. Larger hybridizing bands were undigested chromosomal DNA. Both probes hybridized to a fragment of approximately 7 kb in *H. pylori* UA802 and a fragment of approximately 7.5 kb in *H. pylori* UA763. The probes were labelled with [α^{32} P]-dATP. Hybridizations were carried out at 37°C in the presence of 50% formamide.



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amino acid sequence

E.coli VLLLDKTGTI TLG 315

E. hirae VIMLDKTGTL TQG 286

Rat VICSDKTGTL TQN 393

Consensus VI-LDKTGTL TQG

DNA sequence

E.coli 5'-GTTCTGCTAC TEGATAAAAC CGGCACCATC ACACTCGGT-3'

E. hirae 5'-GTGATCATGT TAGATAAAAC AGGCACCTTG ACCCAAGGA-3'

Rat 5'-GTCATCTGCT CAGACAAGAC AGGAACTCTT ACTCAGAAC-3'

Consensus 5'-GT-ATC-T-T TAGATAAAAC AGGCACC-T- AC-CA-GG-3'

5'-primer 5'-(C/A) AGT TAGATAAAAC AGGCACG-3'
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amino acid sequence

E.coli

VAMTGDGTND APALAQ 528

E. hirae

VIMVGDGIND APSLAR 486

Rat

VAVTGDGVND SPALKK 736

Consensus

VAMTGDG-ND APALA-

DNA sequence

E.coli

5'-GTAGCGATGA CCGGCGACGG CACCAACGAT GCTCCGGCGC TGGCGCAG-3'

E. hirae

5'-GTAGCGATGA CAGGGGATGA CACCAATGAT GCCCAAGCT TAGCACGG-3'
Rat

5'-GTGGCTGTAA CAGGGGATGG TGTGAATGAT GCCCAAGCC TGAACGAG-3'

Consensus

5'-GTGGC-ATGA C-GG-GACGG CATCAATGAT GC-CCAGCC TGAACGAG-3'

3'-CACCG-TACT G-CC-CTGCC GTAGTTACTA CG-GGTCGGG TGGCGCAG-5'

3'-Primer

3'-ACCACTGCC GTAGTTACTA CGC-5'
```

Figure 4-2. Regions of conserved amino acid sequence in P-type ATPases used for designing of PCR primers

The top panel shows the region of conserved amino acid sequence used to design the 5' primer. The bottom panel shows the conserved region used to design the 3' primer. a- from the Escherichia coli kdpB sequence, Hesse et al., 1984
b- from the Enterococcus hirae K+-ATPase sequence, Solioz et

al., 1987

c- from the rat stomach (H++K+)-ATPase sequence, Shull and Lingrel, 1986

(Figure 4-3). Approximately 2000 plaques from each of the λ -DASH H. pylori UA802 genomic DNA BglII and Sau3AI libraries were screened using both the cloned E. hirae K⁺-ATPase gene and the 600 bp PCR product amplified from H. pylori UA802. No positive plaques were observed using the E. hirae probe. Twenty-three positive plaques, 10 from the BglII library and 13 from the Sau3AI library, were observed using the PCR probe. DNA isolation was carried out on two of these positive plaques, designated $\lambda P9$ (Sau3AI library) and $\lambda P14$ (BglII library). Restriction endonuclease digestion analyses were carried out on these clones and the regions containing the P-type ATPase sequences were localized to a 1.3 kb EcoRI-HindIII fragment of clone $\lambda P9$ and a 2.7 kb EcoRI-HindIII fragment of clone $\lambda P14$ (Figure 4-4).

C.2. PCR amplification of rRNA probes and cloning of the rRNA genes of H. pylori UA802

The rRNA PCR probes were amplified from *H. pylori* UA802 using the primers listed in Table 4-1. For the 16S primers, primer 16.5 in combination with primer 16.3a gave a product of approximately 700 bp (Figure 4-5). The combination of primer 16.5 and primer 16.3b gave an almost full-length 16S product of approximately 1500 kb (Figure 4-5).

Nearly full-length 23S PCR products of approximately 2700 bp were amplified using primer 23.5a and primer 23.3a (Figure 4-5), while a product of approximately 650 bp was amplified using the 23.5b primer and 23.3b primer (Figure 4-5). Other combinations of primers (23.5a-23.3b, 23.5b-23.3a) gave

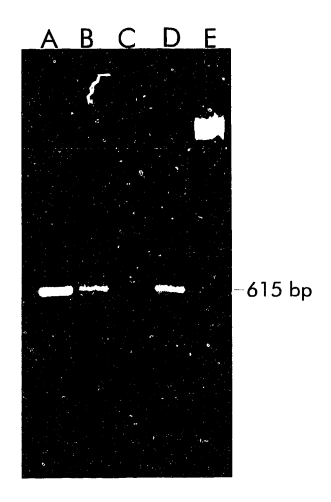


Figure 4-3. Agarose gel analysis of P-type ATPase PCR probe amplified from H. pylori UA802

Lanes A-D were 5 μ l samples of four separate PCR amplifications of H. pylori UA802 using the P-type ATPase primers. Lane E shows the 123 bp ladder size marker. The products were amplified from H. pylori UA802 chromosomal DNA using the primers shown on Figure 4-2 in standard PCR reaction conditions (Chapter 2) for 40 cycles of 94°C for 40s, 45°C for 40s and 72°C for 90s. The PCR product in lane C appears to be degraded or did not amplify. The variations in intensities of the other bands reflect the minor variations in template, primer and buffer concentrations which normally occur in the preparation of PCR reactions. The gel used was a 1% agarose gel run in 0.5 % TBE-buffer, set at a constant voltage of 100v.

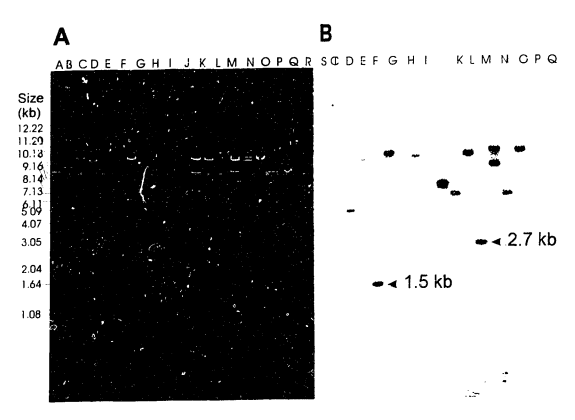


Figure 4-4. Restriction endonuclease digestion analysis and hybridization of $\lambda P9$ and $\lambda P14$

Panel A shows the restriction digestions of $\lambda P9$ and $\lambda P14$ fractionated on a 0.7% agarose gel; Panel B shows the Southern blot of the gel pictured in panel A, probed with the P-type ATPase PCR probe labelled with [$\alpha^{32}P$]-dATP. Hybridization was carried out at 37°C in the presence of 50% formamide.

rormanicue.		
$A-\lambda$ AccI size std.	H- λP9 <i>Xba</i> I	O- λP14 HindIII
$B-\lambda$ Aval size std.	I- λP9 SalI	P- λP14 XbaI
C- λP9 BglII	J- 1 kb ladder	Q- λP14 SalI
	K- λP14 BglII	R- 1 kb ladder
		s- λ NcoI size std.
E- λP9 EcoRI	n. Will Darre moore	
F- λP9 EcoRI-HindIII	M- λP14 ECORI	$T-\lambda$ XhoI size std.
G- λP9 HindIII	N- λP14 EcoRI-HindIII	

Arrows show the 1.5 kb λ P9 EcoRI-HindIII fragment and the 2.7 kb λ P14 EcoRI-HindIII fragment. The 2.7 kb λ P14 EcoRI-HindIII fragment was subsequently subcloned and characterized by Ge et al., 1995.

Table 4-1. Primers used for PCR amplification of 16S and 23S rRNA probes

Gene	Primer sequence	Reference
16S	5'-primer	
	16.5 5'-GTCGGGTAAGTTCCGACCTG-3'	
1	3'-primers	Paster <i>et al.</i> ,
	16.3a 5'-CCGCTTGTCGGTATGGGAAC-3'	
	16.3b 5'-CCAGTCGCTGTGTG	
	TGCCGTGGGCAGTAGC-3'	
235	5'-primers	
	23.5a 5'-GGTGGATGCCTTGGCA(A/C)T-3'	
<u> </u> 	23.5b 5'-GTCGGGTAAGTTCCGACCTG-3'	Egebjerg et
<u> </u> 	3'-primers	
	23.3a 5'-GCTTAGATGCTTTCAGC-3'	
	23.3b 5'-GGCGAACAGCCATACCCTTG-3'	

3.1 kb 2.0 kb 1.0 kb

Figure 4-5. Agarose gel analyses of rRNA probes amplified from H. pylori UA802 using PCR. Five μ l samples of each amplification reaction were run on a 1% agarose gel in 0.5 X TBE-buffer.

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Lanes A-C: product amplified with primers 23.5b-23.3b
Lanes D-F: product amplified with primers 23.5b-23.3a
Lanes H-I: product amplified with primers 23.5a-23.3b
Lanes J-L: product amplified with primers 23.5a-23.3a
Lanes M,R,U: 1 kb ladder
Lanes N,O: product amplified with primers 16.5-16.3a
Lanes P,Q,S,T: product amplified with primers 16.5-16.3b.
```

The products shown in lane J-L represent almost the entire 23S gene (full-length 23S genes are approximately 2900 bp), while the products in lanes P, Q, S, and T represent almost the entire 16S gene (full length 16S genes are approximately 1600 bp).

products of sizes intermediate to the 650 bp and 2700 bp products (Figure 4-5).

Screening of the λ -DASH genomic libraries of H. pylori UA802 with rRNA probes

 λ -DASH genomic libraries (*BylII* and *Sau3AI*) of The H. pylori UA802 DNA were screened with the 700 bp 16S-PCR probe for the 16S rRNA genes and the 650 bp 23S-PCR probe for the 23S rRNA genes. Approximately 2000 plaques were screened from each library. With the 16S probe approximately 18% of the clones were positive, while with the 23S approximately 7.5% of the clones were positive. Phage stocks were made from 16S and 23S clones randomly chosen from the BglII and Sau3AI libraries. DNA was isolated from ten of the 16S stocks and ten of the 23S stocks. Figure 4-6 shows the restriction analyses of one of the 16S clones (λ 16.4) and one of the 23S clones (λ 23.54). Low resolution restriction maps of $\lambda 16.4$ and $\lambda 23.54$ were made by partial digestion mapping The insert in $\lambda 16.4$ was found to be (Figure 4-7). appproximately 16.4 kb while the insert in $\lambda 23.54$ was found to be approximately 20 kb. The 600 bp 16S probe was shown to hybridize to the 1 kb EcoRI fragment of λ 16.4. 23S probe hybridized to the 10 kb Xba \bar{i} fragment of λ 23.54.

Isolation of each individual copy of the 23S genes was carried out using $\lambda 23.54$ as a starting point. The 23S gene was localized to a 10 kb XbaI fragment of this clone (Figures 4-6 and 4-7). The 3 kb EcoRI-XbaI fragment did not hybridize with the 23S probe and was localized to the opposite end to

Figure 4-6. Agarose gel electrophoresis of EcoRI and XbaI restriction digestions of clones $\lambda 16.4$ and $\lambda 23.54$, and hybridization of Southern transfers of the gels with either the 16S or 23S PCR probes.

Lane A- $\lambda 16.4$ EcoRI

Lane B- $\lambda 16.4 XbaI$

Lane C- lane A hybridized with the 700 bp 16S PCR probe

Lane D- lane B hybridized with the 700 bp 16S PCR probe

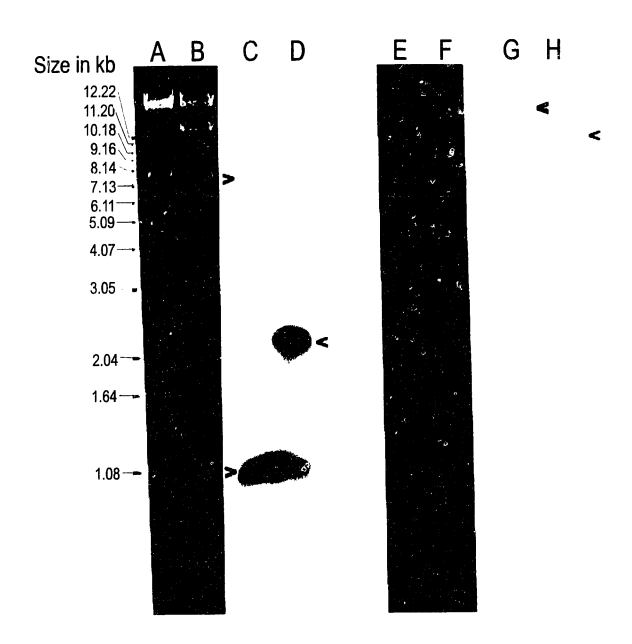
Lane E- $\lambda 23.54$ EcoRI

Lane F- $\lambda 23.54$ XbaI

Lane G- Lane E hybridized with the 650 bp 23S PCR probe

Lane H- Lane F hybridized with the 650 bp 23S PCR probe

The probes were labelled with $[\alpha^{32}P]$ -dATP. Hybridizations were carried out at 37°C in the presence of 50% formamide.



 $\lambda 23.54$



λ16.4

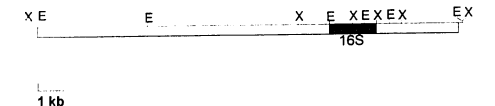


Figure 4-7. Low resolution restriction maps of clones $\lambda 16.4$ and $\lambda 23.54$ constructed using partial digestion mapping, showing the location of the 16S and 23S genes

E- ECORI

X- XbaI

The rRNA genes were localized using data obtained in the restriction endonuclease digestion analyses of these clones (Figure 4-6) and are indicated by shading.

the 23S gene on the inserted H. pylori UA802 DNA fragment of λ23.54. The 3 kb EcoRI-XbaI fragment was isolated from low melting point agarose and hybridized to restriction digestions of nine 23S DNA λ -clones (Figure 4-8). isolated from the same region of the genome of H. pylori UA802 might also carry this fragment. Thus all clones hybridizing to this fragment could be eliminated and clones which did not hybridize could have originated from the other 23S region on the H. pylori UA802 genome. By using this hybridization data in combination with comparisons of restriction endonuclease digestion patterns, clones carrying the other copy of the 23S were selected. The lambda vector was expected to give fragments of 9 and 22 kb in the XbaI The 3 kb EcoRI-XbaI fragment appeared to have digests. homology with lambda, as the 9 and 22 kb lambda bands hybridized to the probe (Figure 4-8). However, Figure 4-8 also shows that the inserted H. pylori UA802 DNA of clones $\lambda 23.B1$, $\lambda 23.B2$, $\lambda 23.S1$ and $\lambda 23.S4$ had no homology with the probe, indicating that the clones may have originated from a different region of the genome than $\lambda 23.54$. Comparisons of restriction endonuclease patterns also suggested that these clones had different chromosomal origins than $\lambda 23.54$. scale DNA isolation was carried out on $\lambda 23.B1$ for further characterization. No further characterization was done with clones $\lambda 23.B2$, $\lambda 23.S1$ and $\lambda 23.S4$. With clones $\lambda 23.54$ and λ23.B1, two copies of the 23S genes of H. pylor. UA802 were

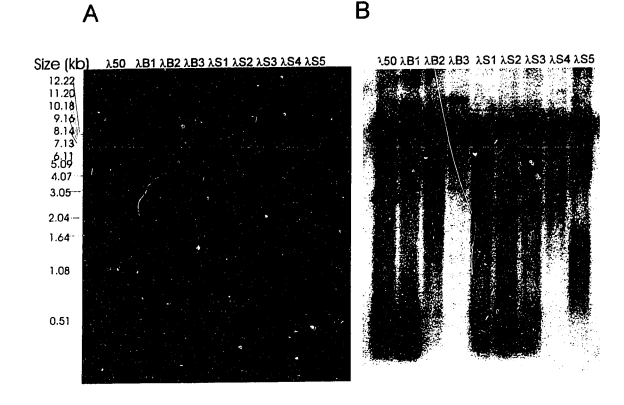


Figure 4-8. Agarose gel electrophoresis of XbaI digestions of nine 23S λ -clones and hybridization of these clones with DNA from the region flanking the 23S gene of $\lambda 23.54$

Panel A shows XbaI digestions of 23S lambda clones fractionated on 0.6% agarose. With XbaI, the lambda vector gives fragments of 9 kb and 22 kb.

Panel B shows a Southern blot of the gel hybridized with the 2 kb <code>EcoRI-XbaI</code> fragment of $\lambda 23.54$. The probe hybridized with the lambda bands and did not hybridize with the insert DNA of $\lambda 23.B1$, $\lambda 23.B2$, $\lambda 23.S1$, and $\lambda 23.S4$. The probes were labelled with $[\alpha^{32}P]$ -dATP. Hybridizations were carried out at 42°C in the presence of 50% formamide.

represented and the 23S sequences from these clones were subcloned and characterized.

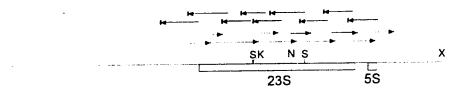
Several attempts to amplify probes for the 5S gene of H. pylori UA802 were carried out in our laboratory but were not successful (E. Newnham, personal communication). However, the 5S genes are normally found immediately 3' to the 23S genes in procaryotes (Nomura et al., 1984) and it was likely that the 5S gene would be present, along with the 23S genes, in clones $\lambda 23.54$ and $\lambda 23.B1$.

Subcloning of the 23S and 5S rRNA genes

For further characterization of the rRNA genes the DNA fragments harboring the 23S genes were further subcloned into the plasmid pBluescript SK- (Stratagene). The 23S genes of both $\lambda 23.54$ and $\lambda 23.B1$ were localized to XbaI fragments of approximately 10 kb in size (data not shown). fragments were cloned into the pBluescript XbaI site to give the plasmids p54X1.1 and pB1X1.1. Restriction maps of these clones were made with the enzymes KpnI, SstII and NruI (Figure 4-9). Restriction digestions of these clones were blotted onto nitrocellulose and hybridized with the 700 bp A 1 kb SstII fragment was and 2800 bp 23S PCR probes. located within the 23S genes of both p54X1.1 and pB1X1.1. These 1 kb SstII fragments were further subcloned into pBluescript as were the SstII-XbaI fragments flanking the 1 kb fragments. Subcloning of the NruI-XbaI fragments was also done.

p54X1.1

×



pB1X1.1

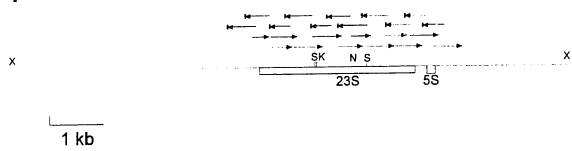


Figure 4-9. Restriction maps of p54X1.1 and pB1X1.1 subcloned from $\lambda 23.54$ and $\lambda 23.B1$ respectively, showing the location of the 23S and 5S genes and sequencing strategy used to determine rRNA gene sequences

S- SstII

K- KpnI

N- NruI

X- XbaI

p54x1.1 was the cloned 10 kb <code>XbaI</code> fragment of $\lambda 23.54$ and pBlX1.1 was the cloned 10.2 kb <code>XbaI</code> fragment of $\lambda B1$. The figure also shows the sequencing strategy used. Arrows above the map show the length and direction of segments sequenced.

Sequencing of the 23S and 5S genes of H. pylori UA802

DNA sequence was obtained from all of the plasmids described above using cycle sequencing and either T7 or T3 Sequential deletions of some of these sequencing primer. Custom sequencing primers were subclones were made. constructed as sequence became available. These primers are listed on Table 4-2 and are shown on Figure 4-10. The sequencing strategy used is shown in Figure 4-9. The complete sequences of the 23S and 5S genes as well as sequence from both the 5'- and 3'-flanking regions and the intervening sequence between the 23S and 5S genes of both Both sequences were identical clones were determined. starting from 373 bp 5' to the 23S gene, through the 228 bp intervening sequence and up to 61 bp beyond the 3' end of the 5S gene, at which point the sequences diverged (Figure 4-10).

Primer extension of 23S and 5S rRNA

Total RNA was extracted from *H. pylori* UA802, and primer extensions were carried out using the primers listed in Table 4-2. From this analysis the starting points of the 23S and 5S rRNA molecules could be determined. The start point of the 23S rRNA molecule was found to be an A nucleotide at position 373 of the sequence (Figures 4-10 and 4-11). Five different start sites were found for the 5S rRNA molecule, four T nucleotides at positions 3568,3572, 3573, and 3574, and one C nucleotide at position 3571 (Figures 4-10 and 4-11).

Table 4-2. Primers used for sequencing and primer extension of the 23S and 5S genes of *H. pylori* UA802

Primer	Application	Sequence (5'-3')	Position of 5'-end*
5b	sequencing	CGGGGAAAAGGGCAAACC	base 699 (s)
5c	sequencing	GGTTTGCCCTTTTCCCCG	base 716 (as)
5d	sequencing/ primer extension	CTTATCGCAGTCTAGTAC	base 533 (as)
5e	sequencing	GTACTAGACTGCGATAAG	base 516 (s)
5 f	sequencing	GAGTGATTATAGCAAG	base 264 (s)
5g	sequencing	GTCAGGGTGATGGACTG	base 1026 (s)
5h	sequencing	GACTATGTGCTTGATAG	base 3257 (s)
5i	sequencing	CCATTCCGAACCTG	base 3611 (s)
5 j	sequencing	GGACTGAACTCCTACCC	base 1207 (s)
3a	sequencing/	GGCGAACAGCCATACCCTTG	base 3014 (as)
3b	sequencing/ primer extension	GAGCAGTATTATCAG	base 3651 (as)
3c	sequencing	CGACATCGAGGTGCC	base 2963 (as)
3d	sequencing	CCAGCACTCCTTACC	base 1711 (as)
3e	sequencing	CTAGCCCAATCAGCGC	base 1358 (as)

^{*-} position of the first nucleotide of the primer on the sequence shown in Figure 4-10. Whether the primer is sense (s) or anti-sense (as) is indicated in brackets.

Figure 4-10. Complete nucleotide sequences of the 23S and 5S genes as well as part of the flanking region from clones p54X1.1 and pB1X1.1

-Translation signals (Pribnow boxes) are marked by underlining and are highlighted.

-Start positions of the 23S and 5S rRNA molecules as determined by primer extension are in bold and are highlighted.

-Putative end positions are in bold and are highlighted.

-The 3' sequences flanking the rRNA genes are shown up to the point where the sequences of the two clones diverge.

-Primers used for sequencing and primer extension, listed in Table 4-2 are underlined and labelled. The direction of the primers are indicated by arrows.

-The DNA sequence shown in this figure has been submitted to Genbank: accession number U27270.

Figure 4-1	2.0	50
10 AAAGCTTCAT		CCAATCACTTTTATCCATTTCTTTCAA
72410011011		
70		P3 110 TATAAT TACATTTCGTTTTAAAGACAA
ACCCAAAAAC	TITAAGCAAACTTTAAGCATGT	ALBORNA
130		170
GCTTTAAAAG	TCTTTAATTGAACCACTCAAAC	AGTTCTACAAGCTAAAGCTTTAAATAA
190		
AACCCACCAG	CTGGTAAAACTTGAGTGT TATA	AAGATTAGGGATCAAGCATTTTTAGT
250	270	290
CTTCTTTAAG		'AGCAAGTTTTTAAAGAAAAACGAAGTT
310	5f → 330	350
ATTTGATTTA	ACATTGTTAATAGCCTATGTAA/	AGTAAAGTAAAACTACAATAACTCTGT
P1 370	23S start 390	410
CT TATATT CA		AATGTTCGTGCAATTGTCGTTATTCAT
		470
430		ATTTAAAACAAGCTTTTAAGAGCAGAT
490		530 AC <u>GTACTAGACTGCGATAAG</u> CTATGCG
GGCGGATGCC	TIGE CAAAGAGAGGCGAIGAAGC	ACOINCIACIO CONTINUE
	1100011	5e → ← 5d
550	570	5e → ← 5d 590
	570	5e → ← 5d
GAGCTGTCAA	570 GGAGCTTTGATGCGTAGATGTCC	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650
GAGCTGTCAA	570 GGAGCTTTGATGCGTAGATGTCC	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT
GAGCTGTCAA 610 ATTAGTTACT	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710
GAGCTGTCAA 610 ATTAGTTACT	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA	5e → 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 GCGAACGGGGAAAGGGCAAACCGAGT
GAGCTGTCAA 610 ATTAGTTACT	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710
GAGCTGTCAA 610 ATTAGTTACT 670 AAGAAATCAA 730	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 GCGAA <u>CGGGGAAAGGGCAAACC</u> GAGT 5b → ← 5c'
GAGCTGTCAA 610 ATTAGTTACT 670 AAGAAATCAA 730 GCTTGCATTC	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACCTACCA	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 AGCGAACGGGGAAAGGGCAAACCGAGT 5b → ← 5c' 770
GAGCTGTCAAGATTACTGAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGATTACCGCTTGCATTCGATTCGATTCGATTCGATTCAAGAAAAGAAATCAAGAAAATCAAGAAAATCAAGAAAATCAAGAAAATCAAGAAATCAAGAAAATCAAGAAAATCAAGAAAATCAAGAAAATCAAGAAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAATCAAGAAAATCAAGAAAATCAAGAAAATCAAGAAATCAAGAAATCAAGAAAATCAAGAAAATCAAGAAAAAAAA	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACCTACCA	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 GCGAACGGGGAAAGGGCAAACCGAGT 5b → ← 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG
GAGCTGTCAAGATTACTGAAGAAATCAAGAAATCAAGGTTACTGCATTCGAAGGTAAGCGAAGGTAAGCGAAAGGTAAGCGAAGAAAGGTAAGCGAAGGTAAGCGAAAGGTAAAGCGAAAGGTAAAGCGAAAGGTAAAGCGAAAAGGTAAAGCGAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGTAAAGCGAAAAGGAAAAGGAAAAGGAAAAGCGAAAAGGAAAAGCAAAAGGAAAAGGAAAAGGAAAAGGAAAAGGAAAAGGAAAA	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACTGCAACATCCA 810 CATAGAAAGTGATAGCCTTGTAT	5e → 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 GCGAACGGGGAAAGGGCAAACCGAGT 5b → ← 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG 830 GCGACAAGGCGTTTTAGGTAGCAGTA
GAGCTGTCAA 610 ATTAGTTACT 670 AAGAAATCAA 730 GCTTGCATTC 790 AAAGGTAAGC	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACTGCAACATCCA 810 CATAGAAAGTGATAGCCTTGTAT	590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 AGCGAACGGGGAAAGGGCAAACCGAGT 5b → ← 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG 830 AGCGAAAGGCGTTTTTAGGTAGCAGTA 650 830 AGCGAAAGGCGTTTTTAGGTAGCAGTA
GAGCTGTCAA 610 ATTAGTTACT 670 AAGAAATCAA 730 GCTTGCATTC 790 AAAGGTAAGC	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACTGCAACATCCA 810 CATAGAAAGTGATAGCCTTGTAT	5e → 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 GCGAACGGGGAAAGGGCAAACCGAGT 5b → 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG 830 GCGACAAGGCGTTTTTAGGTAGCAGTA 690 TGAAGCCGGGGAGACCACTCTCCAACT
GAGCTGTCAAG 610 ATTAGTTACTG 670 AAGAAATCAAG 730 GCTTGCATTCG 790 AAAGGTAAGCG 850 TCCAGAGTAGC	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACTGCAACATCCA 810 CATAGAAAGTGATAGCCTTGTAT	5e → 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 GCGAACGGGGAAAGGGCAAACCGAGT 5b → 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG 830 GCGACAAGGCGTTTTTAGGTAGCAGTA 1890 TGAAGCCGGGGAGACCACTCTCCAACT 950
GAGCTGTCAAG 610 ATTAGTTACTG 670 AAGAAATCAAG 730 GCTTGCATTCG 790 AAAGGTAAGCG 850 TCCAGAGTAGC	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACTGCAACATCCA 810 CATAGAAAGTGATAGCCTTGTAT	5e → 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 GCGAACGGGGAAAGGGCAAACCGAGT 5b → 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG 830 GCGACAAGGCGTTTTTAGGTAGCAGTA 690 TGAAGCCGGGGAGACCACTCTCCAACT
GAGCTGTCAAG 610 ATTAGTTACT 670 AAGAAATCAAG 730 GCTTGCATTCG 790 AAAGGTAAGCG 850 TCCAGAGTAGCG 910 CTAAATACTAG	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACTGCAACATCCA 810 CATAGAAAGTGATAGCCTTGTAT 870 GCCAGGACACGAGGAATCCAGGT	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 AGCGAACGGGGAAAGGGCAAACCGAGT 5b → ← 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG 830 AGCGACAAGGCGTTTTAGGTAGCAGTA AGAGACCGGGGAGACCACTCTCCAACT 950 ATACCGTGAGGGAAAGGAAACCC 1010
GAGCTGTCAAG 610 ATTAGTTACT 670 AAGAAATCAAG 730 GCTTGCATTCG 790 AAAGGTAAGCG 850 TCCAGAGTAGCG 910 CTAAATACTAG	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACTGCAACATCCA 810 CATAGAAAGTGATAGCCTTGTAT 870 GCCAGGACACGAGGAATCCAGGT	590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 AGCGAACGGGGAAAGGGCAAACCGAGT 5b → ← 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG 830 AGCGACAAGGCGTTTTAGGTAGCAGTA CGCGACAAGGCGTTTTAGGTAGCAGTA 890 ATGAAGCCGGGGAGACCACTCTCCAACT 950 ATACCGTGAGGGAAAGGAAACCC
GAGCTGTCAAG 610 ATTAGTTACTG 670 AAGAAATCAAG 730 GCTTGCATTCG 790 AAAGGTAAGCG 850 TCCAGAGTAGCG 910 CTAAATACTAG 970 GCAGTGAGCGG	570 GGAGCTTTGATGCGTAGATGTCC 630 CTAACAGAGAGCGAACCTAGTGA 690 CGAGATTCCCTAAGTAGTGGCGA 750 GGGGTTGAGGACTGCAACATCCA 810 CATAGAAAGTGATAGCCTTGTAT 870 GCCAGGACACGAGGAATCCAGGT 930 CTCTTTGAGCGATAGCGAACAAC	5e → ← 5d 590 GAATGGGGCAACCCAACTAATAGAGAT 650 AGTGAAACATCTCAGTAACTAGAGGAA 710 AGCGAACGGGGAAAGGGCAAACCGAGT 5b → ← 5c' 770 AGAGAACGCTTTAGCAGAGTTACCTGG 830 AGCGACAAGGCGTTTTAGGTAGCAGTA AGAGACCGGGGAGACCACTCTCCAACT 950 ATACCGTGAGGGAAAGGAAACCC 1010

->		
5g→ 1090	1110	1130
		TCTTAATAGGGCGAACAAGTCAGA
1150	1170	1190
		CCAAGTTGAAACGCGTGTAATAGCG
1210	1230	1250
CGTGGAGGACTGAACTCCTA	<u>\CCC</u> ATTGAAACGGGTT	GGGATGAGCTGTGGATAGGGGTGA
1270	1290	1310
AAGGCCAAACAAACTTAGTG	;ATAGCTGGTTCTCTTC	GAAATATATTTAGGTATAGCCTCA
1330	1350	1370
		AGGGCTGCTCGCCGCGGTACCAAA ←3e
1390	1.410	1430
CCCTATCAAACTTCGAATAC	CTTTTATCGTATCTTG	GGAGTCAGGCGGTGGGTGATAAAA
1450	1470	1490
	CAACCCAGACTACCAAA	TAAGGTCCCTAAGTTCTATTCTGA
1510	1530	1550
GTGGAAAAAGATGTGTGGCT	'ACTAAAACAACCAGGA	AGGTTGGCTTAGAAGCAGCCATCCT
1570	1590	1610
TTAAAGAAAGCGTAACAGCT	CACTGGTCTAGTGGTC	CATGCGCTGAAAATATAACGGGGCT
1630	1650	1670
AAGATAGACACCGAATTTGI	AGATTGTGTTAAACAC:	AGTGGTAGAAGAGCGTTCATACCA
1690	1710	1730
	← 3d	STATGAAGTGAGCATGCAGGAATGA
1750	1770	1790
GTAACGATAAGATATATGAG	GAATTGTATCCGCCGTA	AATCTAAGGTTTCCTACGCGATGG
1810	1830	1850
TCGTCATCGTAGGGTTAGTC	GGGTCCTAAGCCGAGT	CCGAAAGGGGTAGGTGATGGCAAA
1870	1890	1910
TTGGTTAATATTCCAATACC	GACTGTGGAGCGTGAT	rGGGCGGACGCATAGGGTTAAGCGA
1930 GCTAGCTGATGGAAGCGCTA	1950 AGTCTAAGGGCGTAGAT	1970 TTGGAGGGAAGGCAAATCCACCTCT
1990	2010	2030
GTATTTGAAACCCAAACAGG	CTCTTTGAGTCCTTTT	TAGGACAAAGGGAGAATCGCTGATA
2050	2070	2090
CCGTCGTGCCAAGAAAAGCC	CTCTAAGCATATCCATA	AGTCGTCCGTACC(;CAAACCGACAC
2110	2130	2150
AGGTAGATGAGATGAGTAT7	CTAAGGCGCGTGAAA	BAACTCTGGTTAAGGAACTCTGCAA

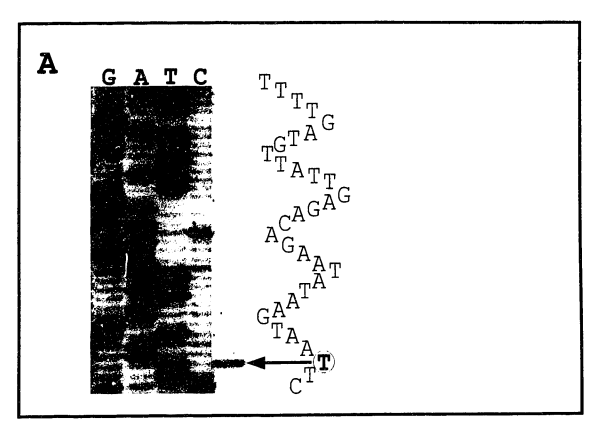
2170	2190	2210
	SATA AGGTGTGCCACAGC	GATGTGGTCTCAGCAAAGAGTCC
ACTAGCACCGTAAGTTCGC	·	
		0.050
2230	2250	2270
CTCCCGACTGTTTACCAAA	AACACAGCACTTTGCCAA	CTCGTAAGAGGAAGTATAAGGTG
C1CCC0		
	2210	2330
2290	2310	
TGACGCCTGCCCGGTGCTCC	JAAGGTTAAGAGGATGCG	TCAGTCGCAAGATGAAGCGTTGA
2350	2370	2390
2330		GTCCTAAGGTAGCGAAATTCCTT
	3CGGCCGTAAC.(ATAACG	GICCIAAGGIAGCOARIIICGII
Figure 4-10 cont'd		
2410	2430	2450
	CATGAATGGCGTAACGA	GATGGGAGCTGTCTCAACCAGAG
GICGGITAAP.INCCOACCIN	307110121100001111001	
		0.51.0
2470	2490	2510
ATTCAGTGAAATTGTAGTG	GAGGTGAAAATTCCTCCT	ACCCGCGGCAAGACGGAAAGACC
	2550	2570
2530	2550	
CCGTGGACCTTTACTACAA	CTTAGCACTGCTAATGGG	AATATCATGCGCAGGATAGGTGG
2500	2610	2630
2590		— -
GAGGCTTTGAAGTAAGGGC'	PTTGGCTCTTATGGAGTC	ATCCTTGAGATACCACCCTTGAT
2650	2670	2690
	ついかのかんかでんなかんなん	AGGACAATGCTTGGTGGCTAGTT
GTTTCTGTTAGCTAACTGG	_CIGIGITATCCACAGGC	AGGACATIGGITGGITGGI
2710	2730	2750
TCACTCCCCCCCCTCCCTCC	raaaagtaacggaggcT	TGCAAAGGTTGGCTCATTGCGGT
104610000001666166		
		2010
2770	2790	2810
TGGAAATCGCAAGTTGAGT	STAATGGCACAAGCCAGC	CTGACTGTAAGACATACAAGTCA
0030	2850	2870
2830		
AGCAGAGACGAAAGTCGGT(CATAGTGATCCGGTGGTT	CTGTGTGGAAGGGCCATCGCTCA
2890	2910	2930
		COCCON NONCOTON CATOCA CCC
AAGGATAAAAGGTACCCCG	GGATAACAGGCTGATCT	CCCCCAAGAGCTCACATCGACGG
2950	2970	2990
	こ こ こ こ こ こ こ こ こ こ こ こ こ こ こ こ こ こ こ	GGGGCTGGAGCAGGTCC <u>CAAGGG</u>
GGAGGTTT <u>GGCACCTC</u> GAT		GGGGCIGGAGCAGGICG
	←3c	3.850
3010	3030	3050
TATGGCTGTTCGCCATTTAL	AAGCGGTACGCGAGCTGG	GTTCAGAACGTCGTGAGACAGTT
4 -3a		
3070	3090	3110
CGGTCCCTATCTGCCGTGG	200m2 002 2 2 0mm02 002	CACCTCTCCCTACTACCACACCA
	GCGTAGGAAAGTTGAGGA	GAGCTGTCCCTAGTACGAGAGGA
	GCGTAGGAAAGTTGAGGA	GAGCTGTCCCTAGTACGAGAGGA
3130	GCGTAGGAAAGTTGAGGA 3150	GAGCTGTCCCTAGTACGAGAGGA 3170
	3150	3170
	3150	
	3150 GGTGCACCAGTTGTCTGC	3170 CAAGAGCATCGCTGGGTAGCACA
CCGGGATGGACGTGTCACTC	3150 GGTGCACCAGTTGTCTGC 3210	3170 CAAGAGCATCGCTGGGTAGCACA 3230
CCGGGATGGACGTGTCACTC	3150 GGTGCACCAGTTGTCTGC 3210	3170 CAAGAGCATCGCTGGGTAGCACA

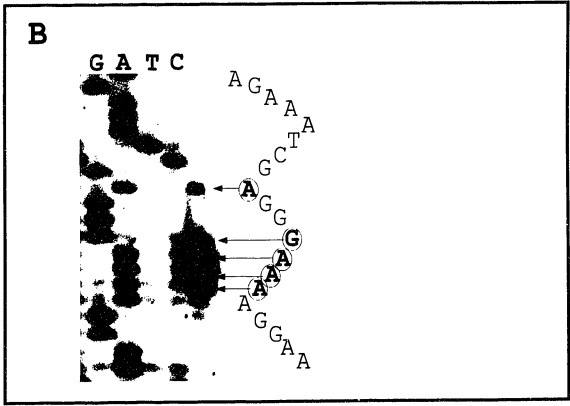
2250	3270	3290
3250		"
CTGAAGCTCGCACAAA <u>GACTAT</u>	GIGCIIGAIAGGGIAGAIG	IGIGAGCGCAGIAAIGCGI
3310	3330 235 ends	3350
-		
TTAGCTGACTACTACTAATAGA	GCG111GGC11G1111E1G	CITITIGATAAGATAACGG
	2200	3410
3370	3390	
CAATAAGCGCGAATGGGTTACC	ACTGCCTTACTGAGTGTAA	GAGAGTTGGAGTTTTATGA
3430	3450	3470
AGACTTTTATAAGATTAAACTT	TAATGAGGAATGAGATACC	ATCTCAATGGTTTAAAGTT
3490	3510	3530
AAAGGCTATTAACGATCTTCTT	TGTTAAAAACAGCTCCCCT	'ATAAAGAGAAAGGGGAGT'T
3550	3570 55 begins	3590
AAGGGTAAATGCGTTTTTATCT	TTAGOTICOCTTTTTCCTTGT	GCCTTTAGAGAAGAGGAAC
Figure 4-10 cont'd		
119010 1 10 00110 0		
3610	3630	3650
TACCCAGTTAACCATTCCGAAC		TCCCTGATAATACTCCTCT
5i →	CIGGRAGICARGCICIICA	4- 3b
3670	3690 5S ends	3710
TTTCAAGAGTGGGAATGTAGGT	· · · · · · · · · · · · · · · · ·	THETTTTTTAGTCTTGCTT
ITICAAGAGIGGGAAIGIAGGI	CGG1GCAGGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	
2522	3750	
3730		an CC
TTTTATTTGATTTCATTATTGA	CTCATTGTTTTGTTTGTTT	AGG

Figure 4-11. Primer extension of total RNA of *H. pylori* UA802 to determine the start points of the 23S and 5S rRNA molecules

Panel A shows primer extension using primer 5d to determine the start position of the 23S rRNA molecule.

Panel B shows primer extension using primer 3b to determine the start position of the 5S rRNA molecule.





PCR from divergent sequence

Because the sequences of the two copies of the 23S/5S genes showed so much identity, it was possible that only one copy of the genes was represented by these two clones. At some point during the cloning of the genes some rearrangement may have occurred to give the appearance that two copies diverging 61 bp from the end of the 5S gene that had been In order to show that two copies had in fact been cloned, primers were designed from the sequence of the clones within the regions where they were different (Figure 4-12). These primers, 54.ext and Bl.ext, were then used to amplify products from the plasmid clones (p54x1.1 and pB1x1.1), the lambda clones ($\lambda 23.54$ and $\lambda B1$) and H. pylori UA802 chromosomal DNA in combination with the primers 5h and 5i The primer 5h starts 82 bp from the end of the (Table 4-2). 23S gene and primer 5i starts 90 bp from the end of the 5S In combination with primer 5h, a product of 655 bp is expected with primer 54.ext and a product of 569 bp is expected with primer Bl.ext. In combination with primer 5i, a product of 300 bp is expected with primer 54.ext and a product of 214 bp is expected with primer Bl.ext. clones do represent both copies of the 23S/5S genes, then products amplified using the 54.ext primer should only be amplified from p54x1.1, λ 23.54 and the *H. pylori* UA802 chromosomal DNA. Likewise, using the primer Bl.ext, products should only be amplified from pBlX1.1, λ Bl and the *H. pylori* UA802 chromosomal DNA. Rearranged clones would either have a



Figure 4-12. Sequence of DNA from the points where p54X1.1 and pB1X1.1 diverge.

Shaded boxes show end of sequence up to where the sequences of the two clones were identical. The last nucleotide in the boxes corresponds to the last nucleotide of the sequence shown in Figure 4-10. Underlined sequences indicate sequence used as PCR primers and the direction of these primers is indicated with an arrow.

different-sized product or no product amplified from the *H*. pylori UA802 chromosomal DNA. Figure 4-13 shows that for all primer combinations, the expected products were amplified from the specific clones and from the *H*. pylori UA802 chromosomal DNA.

Phylogenetic comparisons using the 23s and 5s genes of H. pylori

Sequence analyses and phylogenetic comparisons were done using the Wisconsin Sequence Analysis Package (GCG) as outlined in Chapter 2. The complete sequence of the 23S gene of H. pylori UA802 was compared to the complete sequences of the 23S genes of C. jejuni, C. coli, Pseudomonas cepacia (accession #X16368, Hopfl et al., 1989), E. coli (accession #V00331, Brosius et al., 1980), Rhodobacter capsulatus (accession #X06485, Hopfl et al., 1988), Anacysistis nidulans (accession #X00512, Douglas and Doolittle, 1984), Bacillus subtilis (accession #D11460, Wawrousek and Hansen, 1983) and Micrococcus luteus (accession #X06484, Regensburger et al., 1988) using the pileup program of the Wisconsin Sequence Figure 4-14 shows the dendogram Analysis Package. constructed from this comparison and represents the graphical output of the pileup program.

Phylogenetic comparisons were also carried out with the partial 23S sequences of several Campylobacter species (Figure 4-15)(van Camp et al., 1993), again using the pileup program of GCG. For these comparisons, only the regions of the H. pylori UA802, C. jejuni (Kim et al., 1994) and C. coli

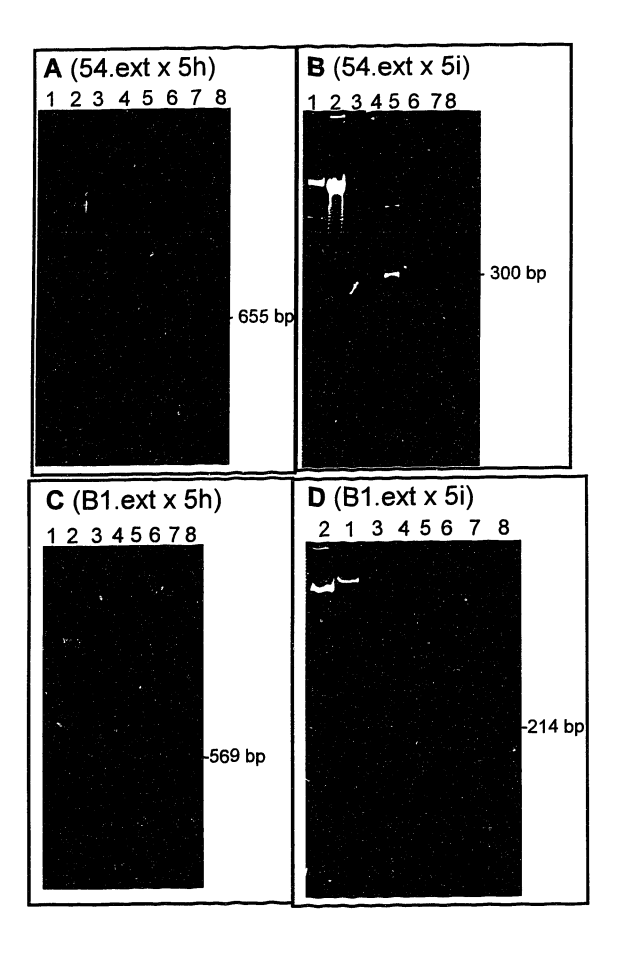
Figure 4-13. PCR amplification from p54X1.1, pB1X1.1, λ 23.54, λ B1 and *H pylori* UA802 chromosomal DNA using primers designed from sequences 3' to the 5S gene where the p54X1.1 and pB1X1.1 sequences differ.

```
For all panels:
1- 1 kb ladder (BRL)
2- 123 bp ladder (BRL)
3- product amplified from p54X1.1
4- product amplified from \(\lambda 23.54\)
5- product amplified from \(H \) pylori UA802 chromosomal DNA
6- product amplified from pB1X1.1
7- product amplified from \(\lambda B1\)
8- no DNA; negative control

Panel A- primers 54.ext and 5h
Panel B- primers 54.ext and 5i
Panel C- primers B1.ext and 5h
Panel D- primers B1.ext and 5i
```

Panels A and C show $5\mu l$ aliquots of the PCR products electrophoresed through 1.5% agarose gels which were run with 100v constant voltage in 0.5 X and then stained with ethidium bromide.

Panels B and D show $5\mu l$ aliquots of the PCR products electrophoresed through 5% acrylamide gels which were run with 300v constant voltage in 0.5 X and then stained with ethidium bromide.



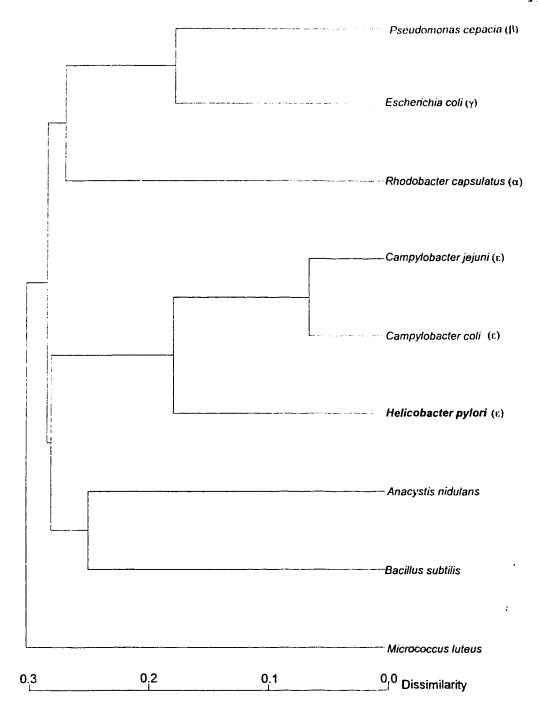


Figure 4-14. Phylogenetic tree based on comparisons of complete 23S rRNA gene sequences, including representative organisms from four of the five Proteobacteria subdivisions and representative organisms from the Gram-positive (B. subtilis and M. luteus) and cyanobacteria (A. nidulans) phyla

Proteobacteria subdivisions are shown in brackets.

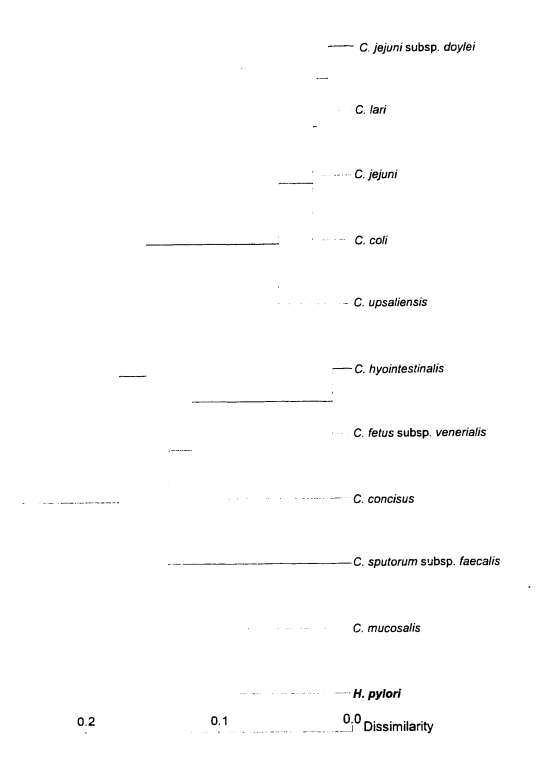


Figure 4-15. Phylogenetic tree based on comparisons of available partial 23S rRNA gene sequences of Campylobacter species and H. pylori

(Trust, et al., 1994) 23S genes homologous to the available Campylobacter partial sequences were used.

The pileup program of GCG was also used to construct the dendogram comparison of the complete 5S gene sequence of *H. pylori* UA802 with the complete 5S gene sequences from representative organisms of all 5 subdivisions of the class *Proteobacteria*, as well as eight out of ten of the other Eubacterial classes described by Woese (1987). Figure 4-16 shows the dendogram constructed for this comparison.

D. Discussion

D.1. Evidence for a P-type ATPase in H. pylori UA802 and UA763

The existence of a P-type ATPase in H. pylori was shown by the hybridization of chromosomal digestions of two strains of H. pylori, UA802 an UA763, with cloned P-type ATPases from the E. hirae (K⁺-ATPase), E. coli (kdpb) and rat ((H⁺-K⁺)-ATPase gene)(Figure 4-3). This suggested that H. pylori P-type ATPases. element homologous to an Hybridization was very weak, which was not surprising as the homology at the DNA level between these ATPases was low, even in regions which showed extensive amino acid conservation (Figure 4-1). The sizes of the hybridizing bands in the two Other genes in H. pylori have shown strains differed. differences in restiriction patterns. Both urease genes (ureABCD) and flagellin genes (flaA) have shown restriction fragment length polymorphism (RFLP) by PCR (Foxall et al.,

Figure 4-16. Phylogenetic tree based on comparisons of complete 5S rRNA gene sequences from representative organisms of eight of the ten Eubacterial phyla

Proteobacteria subdivisions are indicated in brackets.

Sequences used in comparisons:

Flexibacter flexis 55- Van den Eynde et al., 1990.

Accession #M33889

Leptospira interrogans 5S- Fukunaga et al., 1990.

Accession #D90074

Thermus thermophilus 5S- Komiya et al., 1983.

Accession #X01554

Bacillus subtilis 5S- Marotta et al., 1976.

Accession #M10815

Escherichia coli 5S- Brownlee et al., 1967.

Accession #K00609

Desulfovibrio vulgaris 5S- Miura et al., 1986.

Accession #M32519

Borrelia burgdor eri 5S- Gazumyan et al., 1994.

Accession #U03396

Pseudomonas cepacia 5S- Vandenberghe et al., 1.35.

Accession #X02629

Rhodobacter capsulatus 5S- Kato and Komagata, 1986.

Accession #X03904

Micrococcus luteus 5S- Hori et al., 1980.

Accession #J10868

Chlorobium limicola 5S- Van den Eynde et al., 1990.

Accession #MJ3893

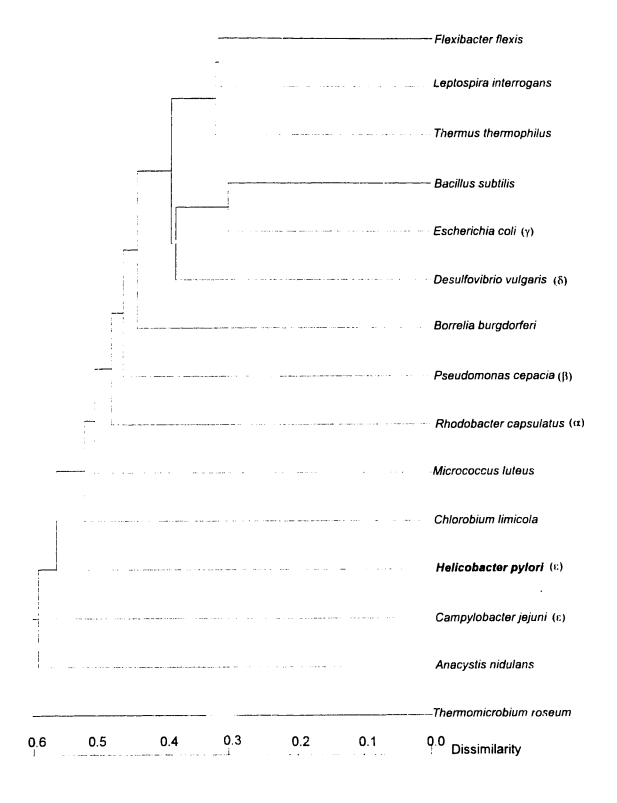
Campylobacter jejuni 5S- Kim et al., 1994.

Accession #229326

Anacystis nidulans 5S- Corry et al., 1974.

Accession #X00757

Thermomicrobium roseum 5S- Van den Eynde et al., 1990.
Accession #M33891



1992; Akopyants et al., 1995). In addition, Cover et al (1994) showed that there could be variation in the vacA sequences from stains which produced active forms of the VacA protein and those which produced inactive forms. The hybridization data presented here using the P-type ATPase probes suggest that similar variations may occur in these genes between strains of H. pylori.

Cloning of a P-type ATPase from H. pylori UA802

Amino acid comparisons have shown that certain regions of P-type ATPases are highly conserved (Hesse et al., 1984; Solioz et al., 1987; Shull and Lingrel, 1986). PCR primers were designed from two of these conserved regions (Figure The primers were designed from the nucleotide 4-2).sequences shown. At positions where the nuclotide sequences sequence was used, as this were variable, the E. hirae organism has similar a G +C ratio as H. pylori. A 600 bp P-type ATPase PCR probe was amplified from H. pylori UA802 using these primers and was used to screen the BglII and Sau3AI $\lambda-DASH$ II genomic libraries of H. pylori UA802. Twenty-three clones were identified with this probe and two of these clones, $\lambda P9$ and $\lambda P14$, were further characterized by restriction endonuclease digestion analysis and hybridization. The PCR probe hybridized to a 1.5 kb EcoRI-HindIII fragment from λP9 and a 2.7 kb EcoRI-HindIII fragment The 2.7 kb EcoRI-HindIII fragment from from clone λ P14. clone $\lambda P14$ was subcloned and used in the characterization of the putative P-type ATPase by Ge et al., 1995. Ge et al.

found that the cloned H. pylori UA802 DNA encoded 3 open reading frames (ORF), designated ORF1, ORF2, and ORF3. polypeptide encoded by ORF1 was found to have extensive homology with several bacterial and eucaryotic P-type ATPases, most strikingly with ATPases involved in copper transport (Vulpe et al., 1993; Odermatt et al., 1993), and was designated hpCopA. The polypeptide encoded by ORF2 was found to have homology with the mercury binding protein merP (Barrineau et al., 1984) and was designated hpCopP. homology could be determined for the ORF3 polypeptide. knock-out mutant of the hpCopA gene was constructed by insertion of a chloramphenicol-resistance cassette, disrupting the gene. This knock-out mutant was susceptible to omeprazole and a number of metal ions at levels similar to those of the wild-type, but was shown to have increased sensitivity to Cu^{2+} ions. Pased on the experimental results and the homology of hpCopA to a copper transport gene, it was concluded by Ge et al. (1995) that this P-type ATPase is probably involved in copper transport in H. pylori UA802.

D.2. Cloning and characterization of the rRNA genes of H. pylori UA802

When the genomic libraries of *H. pylori* UA802 were screened for the 16S and 23S genes with PCR probes, 18% and 7.5% of the plaques hybridized with the 16S and 23S probes respectively. This was significantly higher than the expected frequency of approximately 2% calculated based on an average insert fragment size of 17 kb, a genome size of 1700

kb and the existence of 2 copies of the rRNA genes. suggested that clones carrying the rRNA genes or regions flanking the rRNA genes had some selective advantage over Similar results were obtained in the other clones. construction of genomic libraries of the Caenorhabditis elegans, where ribosomal DNA (rDNA) sequences were cloned at a frequency higher than expected using both lambda vectors (Ellis et al., 1986) and cosmid vectors (Gibson et al., 1987). Gibson et al. suggested that the rDNA had greater stability when cloned in E. coli, as it does not encode a protein and thus is less likely to be toxic. addition, Gibson et al. found that the G + C content of the rDNA (49.3 mol%) was much higher than the rest of the genome mol%), which may have reduced sequence-related instability of the cloned DNA. The cloned 23S-5S regions of H. pylori UA802 also showed a higher G + C ratio (45.3 mol%) than the ratio calculated for the genome (35.2 mol%; Béji et al., 1988). As a result, clones harboring the rRNA genes may have been more stable than other clones. Since the plates that were screened were not primary plates, but were plates made using pooled overlays from primary plates, this phenomenon may have been amplified. Although it was not tested, the frequency of rRNA clones on the primary plates may have been closer to the expected number.

Characterization of the rRNA genes of H. pylori UA802

The 16S rRNA sequences of many diverse organisms have been extensively studied and the phylogenetic relationships

between many organisms have been elucidated by 16S rRNA sequence comparisons (Woese, 1987). In contrast, there have been relatively few studies comparing 23S. As of 1992, only twenty-one complete eubacterial 23S sequences had been reported (Gutell et al., 1992) as opposed to 625 complete eubacterial 16S sequences (de Rijk et al., 1992). Although quite a few 5S sequences have been reported (Erdmann et al., 1985), the small size of the 5S molecules makes them less suitable for phylogenetic comparisons (Woese, 1987). Since the partial and complete sequences of the 16S rRNA genes of a number of strains of H. pylori have already been determined (Romaine et al., 1987; Eckloff et al., 1994) my work was focussed on the characterization of the 23S and 5S rRNA genes of H. pylori UA802.

The complete nucleotide sequences of both copies of the 23S and 5S genes of *H. pylori* UA802 were determined and are shown in Figure 4-10. The fact that the sequences of both copies of the 23S and 5S genes, as well as the flanking and intervening regions, from *H. pylori* UA802 were identical is not unusual. Comparisons of the sequences of 23S and 5S genes from six different *rrn* operons from *Bacillus subtilis* showed very little variation in the 23S and 5S sequences and the intervening sequences (Appendix C). All six copies of the 23S and 5S genes were nearly identical with minor variations in 2 copies (*rrnJ* and *rrnW*). The 5'-flanking regions to the 23S genes for the *rrnO* and *rrnB* were identical at for at least 327 bp, as were the 5' flanking regions of

rrnA and rrnJ (Appendix C). The regions 3' to the 5S gene showed more variation, but the rrnI and rrnO sequences were identical up to 39 bp past the end of the 5S gene (Appendix Similarly, the regions 5' to the 23S genes of H. pylori UA802 were identical for at least 372 bases before the 23S gene and, by comparison of patterns on the sequencing gels (not shown), were similar several hundred bases beyond the point to which sequences were determined. The regions 3' to the 5S genes were identical up to 61 bp from the end of the 5S genes, at which point the sequences diverged. Although the sequences of individual copies of the 16S rRNA genes of H. pylori have not been compared, sequencing data obtained by Eckloff et al. (1994) would suggest that very little variation occurs between individual copies of the 16S genes in H. pylori. In this study the DNA sequence of the 16S genes of five different strains of H. pylori was determined. The 16S genes were first amplified from the chromosomes by PCR and the sequence was determined from these PCR products. If any variation was present in the individual copies of the 16S gene, this would become apparent, as ambiguous sequence would be obtained. No such ambiguities were observed for any of the 16S sequences obtained. In addition, although some variations were observed between the 16S genes from different strains of H. pylori, these variations were small (0.2-0.5%). As Eckloff et al. state, this would indicate that the genomic variability observed between strains of H. pylori does not extend to the 16S genes. The data presented here for the 23S and 5S genes from *H. pylori* UA802 also suggest that variability is not likely to occur in these regions.

Generally, the rRNA genes of procaryotes are found clustered together in operons. In some organisms, as mentioned in Chapter 1, the rRNA genes are not linked. Some strains of Borrelia spp.(Schwartz et al., 1992), C. jejuni and C. coli (Taylor et al., 1992b), Mycoplasma hyopneumoniae (Taschke et al., 1986), Mycoplasma gallisepticum (Chen and Finch, 1989), Leptospira interrogans (Fukunaga and Mifuchi, 1989), Pirellula marina (Liesack and Stackebrandt, 1989), Thermoplasma acidophilum (Ree and Zimmermann, 1990), Thermus thermophilus (Hartman and Erdmann, 1989) and H. pylori (Taylor et al., 1992a; Bukanov and Berg, 1993) are examples of procaryotes in which the rRNA genes are not linked.

Sequence of the 5' region of the 23S genes of *H. pylori* UA802 did not reveal the presence of a 16S gene (Figure 4-10). Thus the rRNA genes of *H. pylori* UA802 do not appear to be linked. This was consistent with results obtained for the construction of the genomic restriction map of *H. pylori* UA802 (Taylor et al., 1992a; Chapter 5, this thesis) where the 16S rRNA genes were not linked with the 23S genes. Coordination of the amounts of rRNA species in organisms where the rRNA genes are linked appears to be governed by the co-transcription of the genes in operons, followed by post-transcriptional processing to give the individual species of rRNA (Nomura et al., 1984). In *E. coli*, transcription of the rRNA operons is regulated by two tandem promoters (Nomura et

al., 1984). These promoters are found approximately 200-300 bases upstream from the 16S rRNA gene and are normally found approximately 120 bp apart (Gilbert et al., 1979). Other regions surrounding the promoters, such as the upstream activation region (UAR) and the factor-dependent and factor-independent regions, are also important in regulating the transcription of rRNA in *E. coli* (Condon et al., 1992).

Three Pribnow boxes (consensus sequence is TATAAT) could be found 5' to the start of the H. pylori UA802 23S gene (Figure 4-10). No -35 region (consensus sequence is GTTGAC) could be clearly defined 5' to any of these Pribnow boxes. sequences homologous to the UAR, factor-dependent or factor-Thus, the control independent regions could be found. mechanisms for H. pylori UA802 rRNA genes must differ from in E. coli and would require those found investigation. It is also possible that sequences similar to the UAR, factor-dependent or factor-independent regions existed beyond the regions which were sequenced for this research. Cloning and sequencing the 5' regions of the 23S genes and in vitro transcription would provide more information about the transcriptional control of this operon. It is interesting to note that the Pribnow box designated Pl was exactly ten bases from the start of the mature 23S sequence, which was determined by primer extension (Figure 4-11). In E. coli, transcription begins almost 300 bp upstream from the 16S gene (de Boer et al., 1979). This suggests that, unlike E. coli, the 5'-end of the 23S-5S transcript is

not processed and transcription, at least from the Pl promoter, begins at position 373 (Figure 4-10), which correlates with the primer extension data (Figure 4-11).

The end of the 23S gene and the start of the 5S gene are separated by 229-235 bp (Figure 4-10) and no promoter sequences could be identified in this region. The 23S and 5S genes must therefore be co-transcribed and post-transcriptionally modified to give the mature 23S and 5S rRNA molecules. It appears that the processing mechanism is not highly specific, as a total of 5 primer extension products were observed for the 5'-end of the 5S rRNA molecule (Figure 4-11). Putative end sites were determined by sequence alignments of the 23S and 5S sequences of a number of other procaryotes with the sequences obtained here (data not shown).

Phylogenetic comparisons of 23S and 5S sequences

As mentioned in Chapter 1, *H. pylori* has been placed in the epsilon subdivision of the class *Proteobacteria* along with *Campylobacter* spp., *Arcobacter* spp., *Wolinella* spp., and *Flexispira* spp. (Vandamme et al., 1991). Phylogenetic comparisons based on 23S sequence are limited by the fact that, compared to 16S rRNA sequences, relatively few complete 23S sequences have been determined. Recently the complete sequences of the 23S genes of *C. jejuni* (Kim et al., 1993, GenBank accession number Z29326) and *C. coli* (Trust et al.,

1994; GenBank accession number U09611) have been determined. Partial sequences of the 23S genes of a number of species of Campylobacter are also available (Van Camp et al., 1993).

The results of the comparisons of the H. pylori UA802 23S gene with the complete sequences of the 23S genes of C. jejuni, C. coli and representative organisms from the other subdivisions of the phylum Proteobacteria (Figure 4-12) were consistent with earlier phylogenetic classifications of The comparisons showed that H. pylori, along with H. pylori. and C. coli, belongs in a separate subdivision C. jejuni Gram-positive bacteria al., 1994). еt cyanobacteria classes are most closely related to the Proteobacteria (Woese, 1987). Therefore, Bacillus subtilis (Gram-positive, low G + C), Micrococcus luteus (Grampositive, high G + C) and Anacystis nidulans (cyanobacteria) were included in the phylogenetic tree (Figure 4-12).

The results of the comparison of the partial 23S sequence of *H. pylori* UA802 with the partial 23S sequences of several Campylobacter species determined by van Campe et al. (1993) were consistent with a previous study which showed that *H. pylori* was distinct from the Campylobacter spp. (Vandamme et al., 1991). In this study, phylogenetic relationships were based on melting temperatures and percent rRNA binding in DNA-rRNA hybrids. The phylogenetic positioning of *H. pylori* was based on DNA-rRNA hybrids using labelled-rRNA from *C. fetus* subsp. fetus, *C. concisus*, *C. coli*, and Wolinella succinogenes. Reciprocal hybrids using rRNA from

H. pylori were not performed. The phylogenetic comparisons done for this research provide direct evidence of the relationships H. pylori has with several members of the genus Campylobacter and supports the findings of Vandamme et al. (1991) that H. pylori belongs in a separate phylogenetic group from the Campylobacter spp..

Comparisons of 5S gene sequences showed that H. pylcri was phylogenetically distinct from the other members of the class Proteobacteria (Figure 4-14). Although a close association between H. pylori and C. jejuni could not be established, H. pylori and C. jejuni were shown to be equally dissimilar to other organisms based on the 5S comparisons. Complete 5S gene sequences from organisms of eight of the ten phyla, described by Woese (1987) are also represented on this phylogenetic tree, including representative organisms from all 5 of the subdivisions of the phylum Proteobacteria. The relationships between the Proteobacteria, Gram-positives (B. subtilis, M. luteus), and cyanobacteria (A. nidulans) observed based on the 23S rRNA sequences could not be demonstrated with the phylogenetic comparisons based on the 5S rRNA sequences. This reflects the lower resolving power of comparisons using the 5S gene sequences as compared to the use of 16S or 23S gene sequences. Because of their relatively small size compared to the 16S and 23S rRNA, differences in 5S rRNA sequences would be more useful for resolution of more recent evolutionary change (Woese, 1987). Similarly, since 23S rRNA molecules are much larger than the 16S rRNA and have approximately twice the number of domains (stem-loop structures), they provide more accurate phylogenetic comparisons than the 16S rRNA (Woese, 1987).

Chapter 5

Pulsed field gel electrophoresis analysis and revision of the genomic map of H. pylori strain UA802

A. Introduction

A genomic restriction map of *H. pylori* UA802 (Figure 5-1) was constructed in 1992 using pulsed-field gel electrophoresis with the enzymes *NotI* and *NruI* (PFGE, Taylor et al., 1992a). PFGE analyses done for this research led to the revision of this map and the addition of *SfiI* sites.

Genomic restriction maps, in general, are useful for studies of genomic variability between strains of bacteria, as well as for the localization of genes and mutant phenotypes (Weinstock, 1994). The genomes of different H. pylori strains have been shown to be highly variable (Majewski et al., 1988; Clayton et al., 1989; Akopyantz et al., 1992; Taylor et al., 1992a). Therefore, genomic restriction maps of H. pylori, such as the map constructed for this work, can be used for comparisons of genetic organization of the chromosomes of different strains. Genomic restriction maps of a number of strains of H. pylori have been constructed (Bukanov and Berg, 1994; Jiang, 1994). Alignment of these maps with the map of H. pylori UA802 constructed for this research may give insights about the genomic variability observed between strains of H. pylori.

H. pylori UA 802

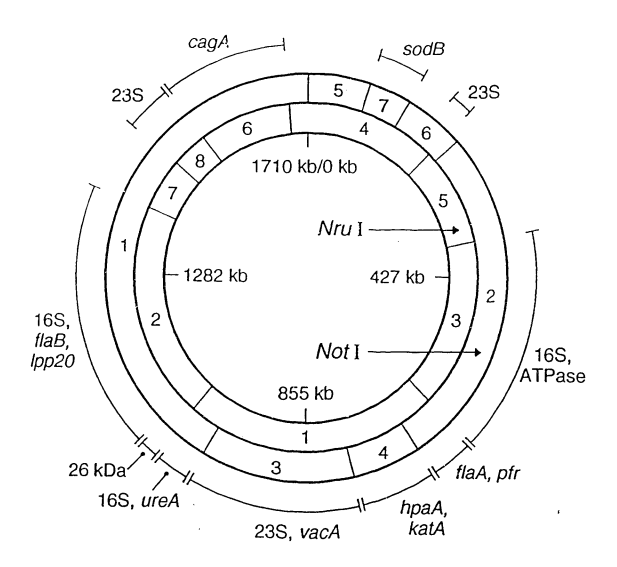


Figure 5-1. Restriction map of H. pylori UA802 constructed by Taylor et al., 1992a

B. Materials and Methods

Please refer to Chapter 2 for detailed methodology used for this chapter.

C. Results

C.1. Construction of the *H. pylori* UA802 restriction map

PFGE analyses

PFGE analysis of H. pylori UA802 chromosomal DNA was the initial step in the construction of the genomic restriction map. Embedded H. pylori UA802 DNA was restricted with NotI, NruI, and SfiI as described in Chapter 2 (Section D). of the digested DNA was carried out with both regular agarose and LMP agarose. Complete digestion of the H. pylori UA802 DNA with these enzymes gave nine NruI fragments, eight NotI fragments and three SfiI fragments. The sizes of the various restriction fragments were estimated from the PFGE analyses of these complete digestions (Table 5-1). The size of the H. pylori UA802 genome was estimated by totaling the sizes of the fragments for each enzyme. The average size estimated from these totals was 1695 ± 17 kb and was in agreement with the size estimated previously for H. pylori UA802 (Taylor et al., 1990) and with estimated sizes of other strains of H. pylori (Bukanov and Berg, 1994; Jiang, 1994), which ranged from 1674 kb to 1730 kb.

Genomic restriction map construction

To determine the order of the fragments in the genomic

Table 5-1. Restriction fragment sizes of H. pylori UA802 DNA

Frag. #	NotI fragments	NruI fragments	Sfil fragments
1	710.0 kb ± 15.0	420.0 kb ± 6.0	870.0 kb ± 15.0
2	458.0 kb ± 8.0	351.0 kb ± 5.0	458.0 kb ± 7.0
3	200.0 kb ± 4.0	253.0 kb ± 8.0	340.0 kb ± 9.0
4	89.0 kb ± 4.0	222.0 kb \pm 3.0	
5	75.0 kb ± 2.0a	152.0 kb ± 6.0	••
6	75.0 kb ± 2.0a	123.0 kb ± 8.0	
7	$54.0 \text{ kb} \pm 1.0$	87.0 kb ± 2.0	-
8	50.0 kb ± 3.0	55.0 kb ± 3.0	
9		38.0 kb ± 2.0	
Total	1711.0 kp	1704.0 kb	1668.0

Average size 1695.0 kb ± 17

a- fragments 5 and 6 formed a doublet band

restriction map, Southern blots of PFGE gels of complete digestions of H. pylori DNA were hybridized with restriction digestion fragments isolated from LMP agarose. The results of the fragment hybridizations are shown in Table 5-2. hybridization patterns of some of the fragments differed from the data obtained in earlier map construction of H. pylori These differences are as UA802 (Taylor et al., 1992). follows and are also indicated on Table 5-2. NruI fragment 1 hybridized with NotI fragments 1, 2, 4, and 6, while on the earlier map NruI fragment 1 hybridized with NotI fragments 1, 2, 3, and 4. NruI fragment 3 hybridized with Not I fragments 2 and 5, while on the earlier map NruI fragment 3 hybridized only with NotI fragment 2. NruI fragment 4 hybridized with NotI fragments 1, 3, 5, and 8 (formerly 7), while on the earlier map NruI fragment 4 hybridized with NotI fragments 1, 5, 6 (7 on revised map), and 7 (8 on the revised map). NruI fragment 5 hybridized with NotI fragment 3, while on the earlier map NruI fragment 5 hybridized with NotI fragments 2 and 6 (7 on the revised map). NruI fragment 6 hybridized with NotI fragment 2, while on the earlier map NruI fragment NruI fragment 8 6 hybridized with NotI fragment 1. hybridized with NotI fragments 3 and 6, while on the earlier map NruI fragment 1 hybridized with NotI fragment 1. fragment lhybridized with Nrul fragments 1, 2, 4, and 7, while on the earlier map NotI fragment 1 hybridized with NotI fragments 1, 2, 4, 6, 7, and 8. NotI fragment 2 hybridized

Table 5-2. H. pylori UA802 fragment localization

			
Digest	NruI*	NotI*	SfiI**
Fragment	NL U I		
NruI 1	1	1,2,4,7	1
		(1,2,3,4)	
NruI 2	2	1	1,2
NruI 3	3	2,5 (2)	1
NruI 4	4	1,3,5,8 (1,5,6,7)	2,3
NruI 5	5	3 (2,6)	1,3
NruI 6	6	2 (1)	1
NruI 7	7	1	2
NruI 8	8	3,5 (1)	1
NruI 9**	9	3	3
Notl 1	1,2,4,7 (1,2,4,6,7,8)	1	1,2,3
NotI 2	1,3,6 (1,3,5)	2	1
NotI 3	4,5,8,9 (1)	3	1,3
NotI 4	1	4	1
NotI 5	4	5	3
NotI 6	3,8	5	1
NotI 7	1 (4,5)	6	1
NotI 8	4	7	3
Sfil 1**	1,2,3,5,6,8	1,2,3,4,6,7	1
SfiI 2**	2,4,7	1	2
Sfil 3**	4,5,9	1,3,5,8	3

^{*} Numbers in brackets indicate fragments which hybridized in the original map (Taylor et al., 1992)

^{**} Did not appear on original map

with NruI fragments 1, 3, and 6, while on the earlier map NotI fragment 2 hybridized with NotI fragments 1, 3, and 5.

NotI fragment 3 hybridized with NruI fragments 4, 5, 8, and 9, while on the earlier map NotI fragment 3 hybridized with NotI fragment 1. NotI fragment 7 (6 on the earlier map) hybridized with NruI fragment 1, while on the earlier map NotI fragment 7 hybridized with NotI fragments 4 and 5. In addition, NruI fragment 9 was not included in the original map and NotI fragment 5 from the original map was shown to be a doublet. Figure 5-2 shows hybridization data confirming the results presented in Table 5-2. The data from the fragment hybridizations were compiled to construct the revised restriction map of H. pylori UA802 (Figure 5-3).

On the revised map, the relative positions of NotI fragments 3 and 7 (previously 6) are different from the original map. As well, the relative positions of NruI fragments 6 and 8 are different from the original map.

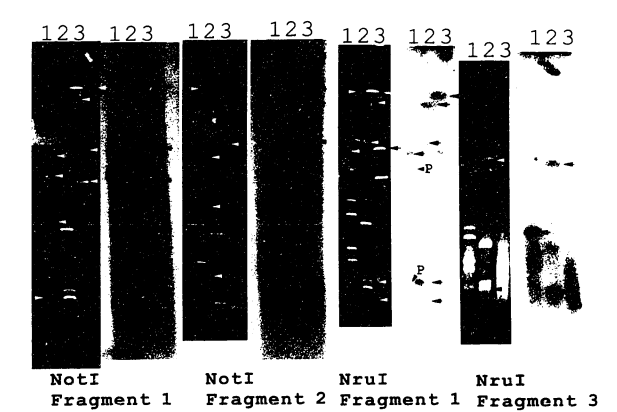
Localization of genes on the H. pylori UA802 map

The Southern blots were also probed with a variety of genetic elements to localize them on the map. These probes and their sources are listed in Table 5-3. Hybridization data confirmed that *H. pylori* UA802 has two copies of the 16s and 23s rRNA genes. This is in agreement with the findings of Bukanov and Berg (1994) and Jiang (1994), who found that other strains of *H. pylori* had two copies of each rRNA gene. Hybridization data combined with sequence data (Chapter 4) showed that the two 23s genes straddled the *NruI* sites

Figure 5-2. PFGE of *H. pylori* UA802 DNA digested with *Not* I, *NruI*, and *SfiI*, and Southern blots of the gels probed with α -[32 P]-dATP-labelled PFGE restriction digestion fragments isolated using LMP-agarose

- 1- NruI digest of H. pylori UA802
- 2- NotI digest of H. pylori UA802
- 3- SfiI digest of H. pylori UA802

Each panel represents an agarose gel on the left and the Southern transfer of that gel, probed with the PFGE fragment indicated, on the right. Arrows on the agarose gel show fragments which hybridized Fragments on the autoradiograms which are not indicated with an arrow were presumed to be For the Nrul fragment partial digest fragments. hybridization, the probe itself contained a partial digest fragment, accounting for the hybridization of Nrul fragments 2 and 7 (indicated with a P on the Southern blot). PFGE was carried out in 0.75% agarose at 8°C with a constant voltage of 190v, with pulse times of 8h at 10s, 8h at 30s and 8h at 60s for a total running time of 24h. PFGE fragments used as probes were run under the same conditions through 1% LMPagarose, visualized by ethidium bromide staining, excised from the gels and labelled with $\alpha \text{-}[^{32}P]\text{-}dATP$ by nick translation. Hybridizations were carried out at 37°C in the presence of 50% formamide.



123
123
123
123
123
NruI
Fragment 4
Fragment 5

123

NruI
Fragment 6

H. pylori UA 802

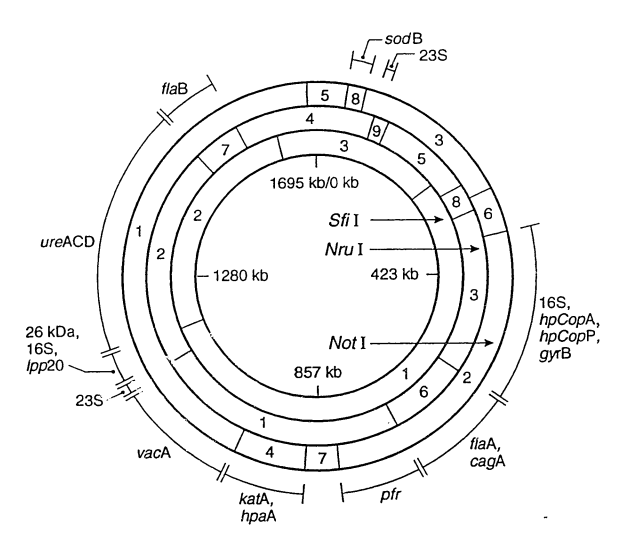


Figure 5-3. Revised genomic restriction map of H. pylori UA802

Table 5-3. Probes used for localization of genes in the $\emph{H.}$ \emph{pylori} UA802 genome

Gene	Plasmid or PCR probe	Origin	Source
23S rRNA	PCR	H. pylori UA802	this work
16S rRNA	PCR	H. pylori UA802	this work
ureACD	pILL594	H. pylori 85P	Labigne et al., 1991
26 kDa protein	₽26K	H. pylori 915	O'Toole et al., 1991
vacA	pCTB1	H. pylori 60190	Cover et al., 1994
- Con	PCR	H. pylori UA802	this work
prf	PCR	H. pylori UA802	Jiang, 1994
flaA	PCR	H. pylori UA802	Jiang, 1994
flaB	PCR	H. pylori UA802	Jiang, 1994
hрСорА	PCR	H. pylori UA802	Jiang, 1994; this work
sodB	pCS188	H. pylori	Spiegelhalder et al., 1993
cagA	PCR	H. pylori UA802	Jiang, 1994
1pp20	PCR	H. pylori UA802	Jiang, 1994
gyrA	plasmid	H. pylori	Moore et al., 1995
gyrB	PCR	H. pylori UA802	Jiang, 1994
katA	PCR	H. pylori UA802	Jiang, 1994
ћраА	PCR	H. pylori UA802	Jiang, 1994

For probes which have the source indicated as "this work" and "Jiang, 1994", the PCR probes were designed from published sequences of the genes. See Appendix B for the primers used.

between NruI fragments 1 and 2 and NruI fragments 4 and 9 (Figure 5-3). During the construction of the original map, where other genes (ureACD, 16S and 26 kDa-protein genes) all appeared to straddle the site between NruI fragments 1 and 2. It was found that NruI fragments 2 and 7 often form a partially digested fragment, particularly resistant to complete digestion, which is the same size as NruI fragment 1. Hybridization data later showed that the ureACD genes were localized to fragment NruI 2 and that the 16S and 26 kDa-protein genes were localized to NruI fragment 1 (Figure 5-4, 26 kDa-protein data not shown). The gyrA gene could not be localized for undetermined reasons.

Assignments of map positions of the various genetic elements, shown in Figure 5-3, were based on hybridization data presented in Table 5-4. The positions of some of the genetic elements on the revised map differed from their positions on the previous map. As mentioned earlier, the positions of the 23S, 16S, ureACD and the 26 kDa protein genes were revised. The positions of the vacA, flaA, and flaB genes were also different. The vacA gene was localized to NruI fragment 1 and NotI fragment 3 on the original map, but localized to NruI fragment 1 and NotI fragment 1 on the revised map. The flaA gene originally was localized to NruI fragment 1 and NotI fragment 2, but on the revised map was found to hybridize to NruI fragment 6 and NotI fragment 2. The flaB gene was mapped to NruI fragment 2 and NotI fragment 1 on the original map, but was later mapped to NruI

Figure 5-4. PFGE of *H. pylori* UA802 DNA digested with *Not* I, *Nru*I, and *Sfi*I, and Southern blots of the gels hybridized with various α -[32 P]-dATP-labelled gene probes

- 1- NruI digest of H. pylori UA802
- 2- NotI digest of H. pylori UA802
- 3- Sfil digest of H. pylori UA802

Each panel represents an agarose gel on the left and a Southern transfer of that gel, screened with the probe indicated below the panel, on the right. Arrows on the agarose gel show fragments which hybridized. Partial digest fragments are indicated with a "P". PFGE was carried out in 0.75% agarose at 8°C with a constant voltage of 190v, with pulse times of 8h at 10s, 8h at 30s and 8h at 60s for a total running time of 24h. Probes were labelled with α -[32 P]-dATP by nick translation. Hybridizations were carried out at 37°C in the presence of 50% formamide.

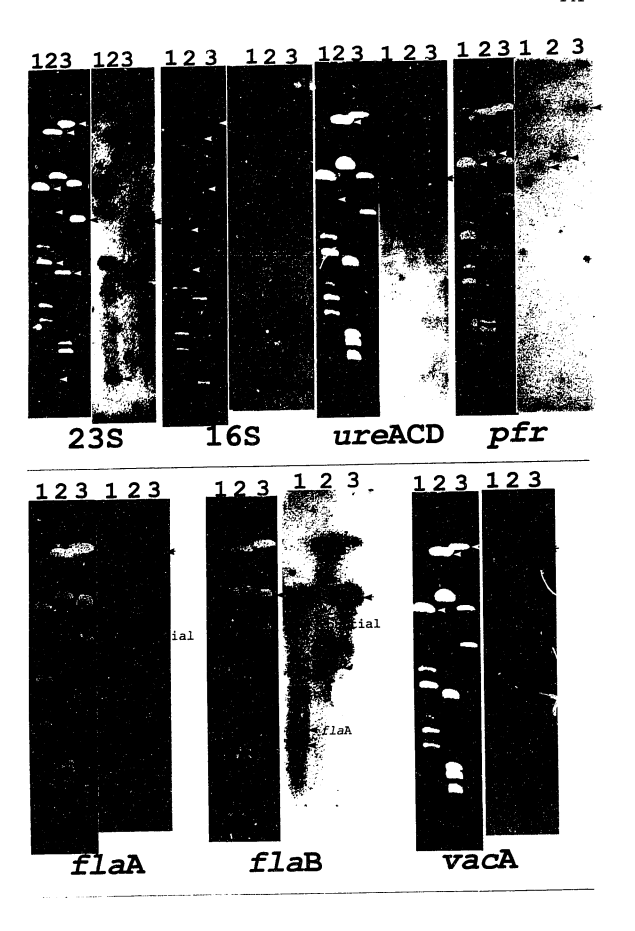


Table 5-4. Localization of genes to the $\emph{H. pylori}$ UA802 genomic map

Gene	NruI	NotI	SfiI			
23S rRNA	1,2,4,9	1,3	1,3			
16S rRNA	2,3	1,2	1			
UreACD	2	1	2			
26 kDa-protein	2	1	1,2			
vacA	1 (4)	1 (3)	1			
pfr	1	2	1			
flaA	6	2	1			
flaB	7	1	2			
hpCopA ^a	3	2	ND			
sodB	4	7	3			
caqA	6	2	1			
1pp20	2	1	ND			
gyrB	3	2	1			
katA ^a	1	6	ND			
hpaA ^a	1	4	ND			

ND- Not done a- assigned to these fragments by Jiang, 1994

fragment 7 and NotI fragment 1. The gyrB gene, which did not appear on the original map, was added to the revised map. The data used for the revisions of the H. pylori UA802 genomic map were presented to and accepted by the authors of the original map.

Alignment of H. pylori genomic maps

In order to compare the arrangements of the genes in different strains of *H. pylori*, the genomic map of *H. pylori* UA802 was aligned with genomic restriction maps constructed by Jiang (1994) of *H. pylori* strains UA861 and NCTC11639 and with the physical map of strain NCTC11638 produced by Bukanov and Berg (1994)(Figure 5-5). The maps were aligned with the vacA gene as the starting point.

D. Discussion

D.1. PFGE analysis and construction of a genomic restriction map of *H. pylori* UA802

The genomic restriction map of *H. pylori* UA802 constructed for this research differed from the previously reported map (Taylor et al., 1992a), although the PFGE patterns obtained were identical. Both the arrangement and number of the restriction fragments and the arrangements of genes have been changed. Partial digestions, difficulties in isolating restriction fragments for probes, and inconsistencies in DNA isolation for PFGE have hampered studies of *H. pylori* UA802 and are likely causes for the differences observed. Improvements in preparation of embedded DNA for PFGE, with

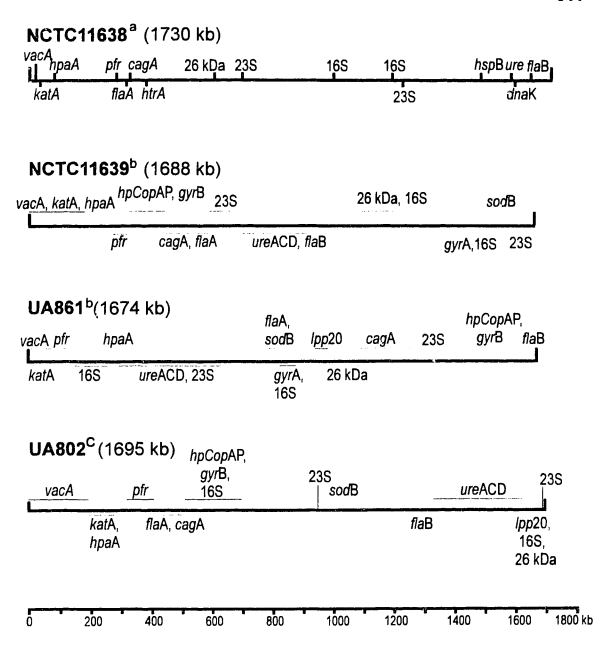


Figure 5-5. Alignment of genomic maps of H. pylori strains NCTC11638, NCTC11639, UA861 and UA802

The vacA gene was used as a starting point for the alignment of the maps.

- a- map constructed by Bukanov and Berg, 1994
- b- map constructed Jiang, 1994
- c- map constructed for this work

the addition of higher EDTA concentrations and the addition of a lysozyme step, as well as refinements in conditions for PFGE (adjustments of pulse times, etc.), have improved the resolution and detection of restriction fragments. This has made the construction of the revised map possible. The additional data obtained with the Sfil restriction digestions have also helped to confirm some of the revisions.

D.2. Comparisons of genome arrangements

Several genes were localized on the genomic restriction map These genes, listed on Table 5-4, of H. pylori UA802. represent the majority of genes which have been cloned and characterized from H. pylori. Not surprisingly, most of these genes are thought to play a role in the pathogenesis of H. pylori. Urease (ureACD) and flagella (flaA and flaB) have been shown to be important colonization factors (Eaton et Superoxide dismutase 1989 and 1991). Spiegolaa der et al., 1993) and catala (katA; Lior & Johnson, 1985) could be involved in resistance to phagocytic killing (Cover and Blaser, 1995). The cytotoxin encoded by the vacA gene causes damage to epithelial cells and, along with the cytotoxin-associated gene cagA, has been associated with the development of ulcers. The rRNA genes (16S and 23S) and gyrase genes (gyrA and gyrB), while not directly involved in pathogenesis, may be important in specifying antibiotic sensitivities or resistances.

On aligning the genomic map of *H. pylori* UA802 with maps of other *H. pylori* strains, considerable variability in gene

arrangement was observed (Figure 5-5). However, some genes were observed to be loosely associated in all the strains. The gyrase gene, gyrB, and the P-type ATPase genes, hpCopA and hpCopP, localized to the same regions on all of the strains for which they were mapped. The vacA, katA, hpaA, and pfr also appeared to be clustered together in all of the strains. With the exception of strain UA861, cagA and flaA genes also appeared to have similar arrangements in the It would appear from these clustered genes that certain regions may be conserv to at least are less susceptible to rearrangement. Krawec and Riley (1990) suggest that nonrandom arrangements, or clustering, of genes suggests that the function or regulation of the genes may be affected by their organization. Since many of the virulence factors of H. pylori, such as vacA and cagA, are variably expressed from strain to strain, it is possible that their arrangement in the chromosome may be a determing factor as to whether they are expressed or not.

Variability has also been observed in restriction digestion patterns of genomic DNA of various strains of *H. pylori* using conventional agarose gels (Majewski et al., 1988; Clayton et al., 1989) and PFGE analyses (Taylor et al., 1992), as well as by arbitrary PCR fingerprinting (Akopyantz et al., 1992). A number of hypotheses for this variability were mentioned in Chapter 1, including differences in methylation patterns, a high frequency of silent mutations, genetic uptake or exchange of DNA and recombination with other strains of *H.*

rearrangements of the chromosome, as well as environmental pressures (host factors). The wide variation of gene arrangements would support the hypothesis that the genomic diversity was the result of chromosomal rearrangements. The mechanisms by which such rearrangements occur are unclear. Through the course of this study, the *H. pylori* strains used were passaged several times without any observed alteration in PFGE patterns, indicating that such rearrangements were not triggered simply by passaging the organism. Of course, the *in vitro* growth environment differs considerably from *in vivo* conditions where host factors and other factors could trigger rearrangements.

It is unlikely that genomic rearrangement is the sole reason for genomic variability. Although variation in restriction endonuclease profiles would be expected with genomic rearrangement, the extent of variability observed also supports the hypothesis that part of the genomic variability is due to spontaneous mutation and/or genetic exchange (Majewski and Goodwin, 1988). Considerable variations in PCR-based fingerprint analyses further support this hypothesis (Foxall et al., 1992; Fujimoto et al., 1994; Because H. pylori Akopyanz et al., 1992a and 1992b). inhabits an environment which is relatively free of other organisms, the possibility of genetic exchange with other organisms is low. However, Akopyants et al. (1995) ha e demonstrated the ability of two strains of H. pylori to cocolonize the stomachs of gnotobiotic piglets, thus creating the possibility for intra-species exchange of DNA. Most likely, a combination of genomic rearrangements, spontaneous mutations and genetic exchange contribute to the genomic variability observed among *H. pylori* strains. Further research is needed to elucidate the exact mechanisms by which genomic variability occurs in *H. pylori*.

Chapter 6

Campylobacter jejuni outer membrane proteins

A. Introduction

Campylobacter species-specific DNA probes were developed in 1990 in this lab (Taylor and Hiratsuka, 1990). One probe, pDT1719, specifically hybridized to DNA from C. jejuni and C. coli. When a lower hybridization temperature was used to reduce stringency (37°C instead of 42°C), this probe was also able to detect Campylobacter lari DNA. The second probe, pDT1720, could only detect C. jejuni DNA. Both probes were initially isolated from a λ-gtll expression library of C. jejuni UA580. This library was screened with antiserum directed against the C. jejuni UA580 major outer membrane protein (MOMP). Two positive lambda clones were isolated and the C. jejuni UA580 DNA fragments from these two clones were further subcloned into pUC13. These plasmid sub-clones, pDT1719, and pDT1720, were used as probes.

To date no genetic characterization of the MOMP of C. jejuni has been published although some characterization of the MOMP protein of C. jejuni has been reported (Huyer et al., 1986; Page et al., 1989). The porin protein has a molecular weight of approximately 45 kDa and was shown to form cation-selective pores in liposome vesic as that allow passage of molecules smaller than ~360 Da.

Since the two cloned DNA probes were isolated from a λ -gtll expression library using anti-sera directed against

the MOMP, it was possible that all or part of the MOMP gene of this organism had been cloned in these two DNA probes. However, results of characterization of the DNA fragments cloned in these probes, described in this chapter, revealed that the MOMP gene had not been cloned. In this chapter the analysis of the nucleotide sequence of the cloned DNA is described. The N-terminal amino acid sequence for purified MOMP protein was also obtained.

B. Materials and Methods

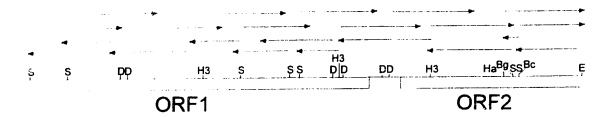
Please refer to Chapter 2 for detailed methodology used for this chapter.

C. Results

C.1. DNA sequence analysis of pDT1719 and pDT1720

Nucleotide sequencing was carried out to determine if the DNA cloned in the two Campylobacter DNA probes encoded all or part of the MOMP of C. jejuni UA580. Restriction maps produced during the initial characterization of the two Campylobacter probes, pDT1719 and pDT1720, are shown in Figure 6-1. Utilizing the restriction sites available in these plasmids, restriction fragments were sub-cloned into pUC13 and sequenced directly by double stranded DNA sequencing (Wang, 1988) using universal forward and reverse primers. Figure 6-1 shows the direction and length of sequence obtained using these sub-clones. The complete sequence of the C. jejuni UA580 DNA fragments cloned in

pDT1719



pDT1720

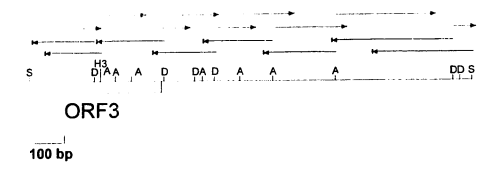


Figure 6-1. Restriction maps of pDT1719 and pDT1720, and sequencing strategy used to determine the sequence of the cloned DNA

A- AluI.

Bc- BclI

Bg- BglII

D- DraI

H3- HindIII

Ha- HaeII

S- Sau3AI

Arrows represent size and direction of segments sequenced. Open boxes indicate position of the open reading frames (ORF).

pDT1719 and pDT1720 were obtained. Figure 6-2 shows the DNA sequence for the pDT1719-cloned DNA and Figure 6-3 shows the sequence for pDT1720-cloned DNA. Neither plasmid was found to encode an entire open reading frame (ORF). Instead, pDT1719 was found to encode two partial ORFs, while pDT1720 encoded one partial ORF. The two ORFs found in pDT1719, designated ORF1 and ORF2, were separated by 78 bp. A ribosome binding site was found in this intervening sequence immediately 5' to ORF2 (Figure 6-2). The ORF found in pDT1720, ORF3, encoded only the C-terminal portion of a polypeptide (Figure 6-3).

The DNA and predicted amino acid sequences of the three ORFs were compared to sequences found in the computer database GenBank to find homologous structures in other organisms. No sequences were found which had homology with ORF3 for either the DNA or predicted amino acid sequence. A partial DNA sequence (150 bp) of a putative *C. jejuni* serine protease appeared in GenBank as a direct submission (accession number X82628) which has 97% homology with ORF1. The deduced amino acid sequence of ORF1 also has homology to the HtrA serine protease/heat shock proteins of Salmonella typhimurium and *E. coli* (Figure 5-4). ORF1 has 63% and 64% homology respectively with the HtrA proteins of *S. typhimurium* and *E. coli*. Because of this homology, ORF1 was designated *C. jejuni htr*A.

ORF2 shows homology with regulatory-signalling elements of a number of procaryotic two-component regulatory systems

Figure 6-2. The complete nucleotide sequence of the *C.jejuni* DNA cloned in the *Campylobacter* DNA probe pDT1719 and the predicted amino acid sequences encoded by the DNA. The DNA sequence has been submitted to GenBank: accession number U27271

Potential ribosome binding site at position 1176 is underlined and highlighted. Asterisk indicates termination codon.

```
Figure 6-2
                    30
GATCAGGGGTGATTATTTCAAAAGATGGTTATATAGTAACAAATAATCACGTTGTAGATG
 S G V I I S K D G Y I V T N N H V V D D
                    90
ATGCTGATACGATTACAGTGAATTTACCAGGAAGCGACATAGAATATAAAGCAAAACTTA
 A D T I T V N L P G S D I E Y K A K L I
                   150
TAGGTAAAGATCCAAAAACAGATTTGGCTGTTATAAAAATAGAGGCTAATAATCTTTCAG
 G K D P K T D L A V I K I E A N N L S A
                   210
CTATTACTTTTACAAATTCTGATGATTTAATGGAAGGAGATGTTGTTTTTTGCACTTGGAA
 I T F T N S D D L M E G D V V F A L G N
                   270
P F G V G F S V T S G I I S A L N K D N
                   330
     310
ATATAGGTTTAAATCAATATGAAAATTTTATACAAACAGATGCTTCTATCAATCCAGGAA
 I G L N Q Y E N F I Q T D A S I N P G N
                   390
ATTCAGGTGGAGCTTTGGTGGATAGTCGCGGATATT'1AGTAGGTATTAATTCAGCTATTC
 S G G A L V D S R G Y L V G I N S A I L
                   450
TTTCTCGTGGTGGTAAATAACGGCATAGGTTTTGCCATACCTTCAAATATGGTTAAAG
 S R G G G N N G I G F A I P S N M V K D
                   510
     490
ATATAGCTAAAAAACTTATTGAAAAAGGCAAGATTGATAGAGGATTTTTAGGTGTGACTA
 I A K K L I E K G K I D R G F L G V T I
     550
                   570
TTTTAGCTTTGCAAGGTAATACTAAAAAAGCTTACAAAAATCAAGAAGGAGCTTTAATCA
 LALQGNTKKAYKNQEGALIT
                                  650
                   630
CTGATGTTCAAAAAGGTTCAAGTGCTGATGAAGCAGGGCTTAAGCGTGGAGATTTAGTTA
 D V Q K G S S A D E A G L K R G D L V T
     670
                   690
CTAAAGTTAATGATAAGGTTATAAAAAGTCCTATTGATCTTAAAAATTATATAGGAACTT
 K V N D K V I K S P I D L K N Y I G T L
                    750
     730
EIGQKISLSYERDGENKQAS
                    810
F . K G E K E N P K G V Q S D L I D G
                    870
     850
GTTTGAGTTTGAGAAATTTAGATCCAAGACTTAAGGATCGTTTGCAAATTCCAAAAGATG
 LSLRNLDPRLKDRLQIPKDV
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Figure 6-2 continued
                    930
N G V L V H S V K E K S K G K N S G F Q
                     990
                                    1010
AAGAGGGTGATATTATCATAGGTGTTGGACAAAGTGAAATTAAAAATTTAAAAGATTTAG
 E G D I I G V G Q S E I K N L K D L E
                    1050
A ACAAGCTTTAAAACAAGTTAATAAAAAAGAATTTACCAAAGTTTGTGTATATCGCAATG
 Q A L K Q V N K K E F T K V C V Y R N G
                    1110
                                    1130
    1090
GTTTTGCGACTTTGCTTGTGCTTAAATAAAATAACGGAAGGATTTTCCTTTCGTTATAA
 F A T L L V L *
                    1170
                          RBS
                                    1190
    1150
ATTTTAGTTTAAAATTAAAATTAAGTTTAAAATAA<u>AGGA</u>AAACTATGACAAATATTCTTA
                                  MTNILM
                                    1250
                    1230
    1210
TGATAGAAGATGATTTAGAATTAGCAGAAATTACAGCTGAATATTTGGAAAAATTTGATA
 I E D D L E L A E I T A E Y L E K F D M
                                    1310
                    1290
    1270
TGAAAGTTGATATAGCTCATGAACCTTATATAGGTCTTTCTAAGCTTGCATTAAAAGAAT
 K V D I A H E P Y I G L S K L A L K E Y
                    1350
    1330
ATCAGCTTATCATTTTAGACCTTTCTTTGCCAGGGCTTGATGGGCTTGAAGTGTGTGAAG
 Q L I I L D L S L P G L D G L E V C E E
                    1410
                                    1430
    1390
AGATTCGTAAAAAGTATGATACGCCTATTATTGTTTCAAGCGCAAGACATGATATTACAG
 IRKKYDTPIIVSSARHDITD
                    1470
                                    1490
    1450
K V N A L E L G A D D Y L P K P Y N P K
                                    1550
                    1530
    1510
AAGAATTACAAGCACGTATTAAAAGTCATTTAAGACGTATTTCAAATACAAAAAGCGCCA
 ELQARIKSHLRRISNTKSAI
                    1590
                                    1610
    1570
TAGCAAAAAGTGTAAAAGATCTTGTTTATGATCAATATAAGCATATTATTACCATGAAAG
 A K S V K D L V Y D Q Y K H I I T M K G
                                    1670
    1630
                    1650
GACAAGAGCTTACTTTAACTAATGCTGAATTTGATATTTTAAGCTATTTGATTAAAAAAG
 Q E L T L T N A E F D I L S Y L I K K E
                                    1730
                    1710
    1690
AGGGTGGAGTGGTAAGTCGTGAAGAGCTTGTTTATAATTGTTCTTCTATTAGTGAGGATT
 G G V V S R E E L V Y N C S S I S E D S
                    1770
    1750
CCAGTAATAAAAGTATAGATGTTATTATCAGAGAATTC
```

SNKSIDVIIREF

156

Figure 6-3. The complete nucleotide sequence of the *C.jejuni* DNA cloned in the *Campylobacter* probe pDT1720 and the predicted amino acid sequences encoded by the DNA. The DNA sequence has been submitted to GenBank: accession number U27272. Asterisks indicate termination codons.

Figure 6-3

			10							0						50			
					_														CGCT
D	P	S	N	K	N	L	L	E	Q	L	K	N	K	N	Т	N	L	Y	A
			70							0						110			
																			ACTA
Ι	F	L	L	K	E	N	I.	N	D	F	N	N	T	Т	L	Q	N	E	L
			30						15							170			
													-						ATCT
K	Q	Ι	Y	N	N	A	Q	T	N	Т	L	L	K	N	I	I	Α	L	S
		1	90						21	0						230			
-																			ACTT
L	G	D	K	S	I	F	L	K	N	Y	D	K	L	L	E	A	Y	K	L
			50						27	_						290			
																			TTCA
L	E	Q	N	K	I	E	E	Α	Ñ	V	L	L	S	Q	Ι	K	E	N	S
		3	10						33	0						350			
AG'	_		TCA																ATTG
S	L	N	Q	I	A	K	N	L	N	F	Q	A	S	М	K	I	7	Т,	L
		3	70						39	0						410			
TA!	rtc'	rtc	ترG	TGG	CCC	AAA	GCA		TTT										GTCA
Y	S	W	R	G	P	K	Q	Y	L	Т	K	T	N	R	W	R	F	K	S
		4	30						45	0						470			
TAI	ATA	GCA.	TTA	TAA	AAA	GCA	'AA	TCG'	TTG	TTA	GGA	A'I'C	TAG'	TTT(CTG	CAA	AAC'	TTA	GCAA
*	*																		
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IAA	41M	-84	LIA	101	IMM	WWW	-MAT	-74-11	MIC.	100	41W	JAC.	AAG.	1111	-1-1-1-1	CII	1	~~~	WII
		5 !	50						570)						590			
ATA	ATGO	TTC	CTA	GCT'	TAT	CAA	SAT	GGG	GA:	rat:	rtt <i>i</i>	ACT	GCA	GAT	TAP	TAA	GA.	AA'T'	AATT
			10						630							650			
AAA	ATTI	TTC	3AC	AAT'	TCT"	TAT	AATO	JAAA	ATT	ГАТ	AAT'	l'T'T	CAA'	rtt(3AC	GCTA	AGT(TΤυ	GTAG
		67	7.0						690	1						710			
GAG	TAG			TAP	GGA(GATO	JAT'	TAC			TAC	CTT	GCA/	\AC!			ATA	GTT(CTTG
		73	30						750)						770			
CAA	ATC	GT7	CT.	rtg	GGT	ATTA	'AA	rtc <i>i</i>	AGTO	CAA	ACCO	CTA	ACC:	CAC	GCT(CCAC	3CC	CAAC	GA CA
		-																	
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		91	.0						930)					:	950			

 ${\tt AAAATTTTTCAACAATGTAATCCATCTAAGTGTAATCGGAGATAAACTTATAGCAGCAA}$

Figure 6-3 continued

rigule 6-3 concil	iueu							
970	990	1010						
CAGCTAAAAAATCATTGTTGTAAGCCCTGCTAGAAACTCTTTATCT'IGACGC'IGACATAA								
1030	1050	1 170						
AAGATGTTGCACTTAGCGAT	GATGGTATTTTTATACTA	AGAAAAAGA. GCCACTATTATTA						
1090	1110	1130						
<i>L</i> AACTGATTATAACTTAAG <i>A</i>	\AAAATTGCGGAAAAAAA	ATTTGAATTTGCTATTTTTGTAA						
1150	1170	1190						
	TATTTATATATTTTTGA	AAAAACAGGCTATCTTATCAAAA						
1210	1230	1250						
TGAATCTGAATTTAGATAATGCACAAG1ATTTAAACTTTCAGAAGCTGTTGATAAAATTT								
1270	1290	1310						
CTTTCATGGGAAATGGTAAA	ATTTTACTACGGAGATAAA	ATTTTAGATTTGCTATGATAAA						
1330	1350	1370						
AATTTTAGAACAAACAATAAAGGCTTTAAAATTTAAATTTAAAGCCCTATGACTTAAGTAT								
1390	1410							
GTTGACTAGAAAAAAAAGTT		2						

Figure 6-4. Amino acid sequence comparisons of the putative C. jejuni HtrA protein with sequences found in GenBank

- a- predicted amino acid sequence of the putative htrA gene of C. jejuni UA580 determined in this research. Submitted to GenBank: Accession # U27271.
- b- translation of DNA sequence determined by Lipinska et al., 1988. Accession #X12457
- c- translation of DNA sequence from GenPank direct submission 1994. Accession #A19802
- d- translation of DNA sequence from GenBank direct submission 1994. Accession #X82628

Figure 6-4

	1				50
C. jejuni HtrA ^a F.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d	MKKTTLAMSA	LALSLGLALS LALSLGLALS	PLSATAAE: S	SSATTAQQMP SSATTAÇQMP	STT EMLEKVM SLAPMLEKVM
Consensus		LALSLGLALS			
a deduced transa	51				100
C. jejuni HtrA ^a E.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d	PSVVSINVEG	STTVNTPRMP ST /NTPRMP	RNFCQFFCDD	_	QNSPFCQGGG
Consensus		STTVNTPRMP			
	101				150
C. jejuni HtrA ^a E.coli HtrA ^b		SGVI FMALGSGVII			
S. typhimurium HtrAc		FMALGSGVII			
C. jejuni HtrA ^d					
Consensus		FMALGSGVII	DA-KGYV /TN	NHVVDNADVI	
C. jejuni HtrAª	151 YKAKLIGKDP	KTDLAVIKÏE	.ANNLSAITF	TNSDDLMEGD	VVFALGNPFG
E.coli HtrAb	FDAKMVGKDP	RSDIALIQIQ	NPKNLTAIKM	ADSDALRVGD	YTVGIGNPFG
S. typhimurium HtrA ^c		RSDIALIQIQ			
C. jejuni HtrA ^d Consensus		KTDLAVIKIE			
Consensus	201	DAII	n ne ne .	00 2 00	250
C. jejuni HtrAª		SALNKDNIGL	NOYENFIOTD	ASINPGNSGG	
E.coli HtrAb		SALGRSGLNA			
S. typhimurium HtrA ^c C. jejuni HtrA ^d		SALGRSGLNV SALNKDNIGL			
Consensus		SALL L			
			-		
	251				300
C. jejuni HtrA ^a	251 GINSAILSRG	GGNNGIGFAI	PSNMVKDIAK	KLIEKGKIDR	
E.coli HtrA ^b	GINSAILSRG GINTAILAPD	GGNIGIGFAI	PSNMVKNLTS	QMVEYGQVKR	GFLGVTILAL GELGIMGTEL
E.coli HtrA ^b S. typhimurium HtrA ^c	GINSAILSRG GINTAILAPD GINTAILAPD	GGNIGIGFAI GGNIGIGFAI	PSNMVKNLTS PSNMVKNLTS	QMVEYGQVKR QMVEYGQVKR	GFLGVTILAL GELGIMGTEL GELGIMGTEL
E.coli HtrA ^b	GINSAILSRG GINTAILAPD GINTAILAPD GINSAIFLVV	GGNIGIGFAI	PSNMVKNLTS PSNMVKNLTS PSNMVKDLQA	QMVEYGQVKR QMVEYGQVKR SP	GFLGVTILAL GELGIMGTEL GELGIMGTEL
E.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d	GINSAILSRG GINTAILAPD GINTAILAPD GINSAIFLVV GIN-AILAPD 301	GGNIGIGFAI GGNIGIGFAI GGNNGIGFAI GGN-GIGFAI	PSNMVKNLTS PSNMVKDLQA PSNMVK-LTS	QMVEYGQVKR QMVEYGQVKR SP QMVEYGQVKR	GFLGVTILAL GELGIMGTEL GELGIMGTEL GELGIMGTEL 350
E.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d Consensus C. jejuni HtlA ^a	GINSAILSRG GINTAILAPD GINTAILAPD GINSAIFLVV GIN-AILAPD 301 QGNTKKAYK.	GGNIGIGFAI GGNIGIGFAI GGN-GIGFAI . NQEGALITD	PSNMVKNLTS PSNMVKNLTS PSNMVKDLQA PSNMVK-LTS VQKGSSADEA	QMVEYGQVKR QMVEYGQVKR SP QMVEYGQVKR GLKRGDLVTK	GFLGVTILAL GELGIMGTEL GELGIMGTEL GELGIMGTEL 350 VNDKVIKSPI
E.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d Consensus C. jejuni HtlA ^a E.coli HtrA ^b	GINSAILSRG GINTAILAPD GINTAILAPD GINSAIFLVV GIN-AILAPD 301 QGNTKKAYK. NSELAKAMKV	GGNIGIGFAI GGNIGIGFAI GGN-GIGFAI . NQEGALITD DAQRGAFVSQ	PSNMVKNLTS PSNMVKNLTS PSNMVK-LTS PSNMVK-LTS VQKGSSADEA VLPNSSAAKA	QMVEYGQVKR QMVEYGQVKR SP QMVEYGQVKR GLKRGDLVTK GIKAGDVITS	GFLGVTILAL GELGIMGTEL GELGIMGTEL GELGIMGTEL 350 VNDKVIKSPI LNGKPISSFA
E.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d Consensus C. jejuni HttA ^a E.coli HtrA ^b S. typhimurium HtrA ^c	GINSAILSRG GINTAILAPD GINTAILAPD GINSAIFLVV GIN-AILAPD 301 QGNTKKAYK. NSELAKAMKV	GGNIGIGFAI GGNIGIGFAI GGN-GIGFAI . NQEGALITD	PSNMVKNLTS PSNMVKNLTS PSNMVK-LTS PSNMVK-LTS VQKGSSADEA VLPNSSAAKA	QMVEYGQVKR QMVEYGQVKR SP QMVEYGQVKR GLKRGDLVTK GIKAGDVITS GIKAGDVITS	GFLGVTILAL GELGIMGTEL GELGIMGTEL GELGIMGTEL 350 VNDKVIKSPI LNGKPISSFA
E.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d Consensus C. jejuni HtlA ^a E.coli HtrA ^b	GINSAILSRG GINTAILAPD GINTAILAPD GINSAIFLVV GIN-AILAPD 301 QGNTKKAYK. NSELAKAMKV NSELAKAMKV	GGNIGIGFAI GGNIGIGFAI GGN-GIGFAI . NQEGALITD DAQRGAFVSQ	PSNMVKNLTS PSNMVKNLTS PSNMVKDLQA PSNMVK-LTS VQKGSSADEA VLPNSSAAKA VMPNSSAAKA	QMVEYGQVKR QMVEYGQVKR SP QMVEYGQVKR GLKRGDLVTK GIKAGDVITS GIKAGDVITS	GFLGVTILAL GELGIMGTEL GELGIMGTEL GELGIMGTEL 350 VNDKVIKSPI LNGKPISSFA LNGKPISSFA
E.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d Consensus C. jejuni HtrA ^a E.coli HtrA ^b S. typhimurium HtrA ^c C. jejuni HtrA ^d Consensus	GINSAILSRG GINTAILAPD GINTAILAPD GINSAIFLVV GIN-AILAPD 301 QGNTKKAYK. NSELAKAMKV NSELAKAMKV NSELAKAMKV	GGNIGIGFAI GGNIGIGFAI GGNNGIGFAI GGN-GIGFAI .NQEGALITD DAQRGAFVSQ DAQRGAFVSQ DAQRGAFVSQ	PSNMVKNLTS PSNMVKNLTS PSNMVKDLQA PSNMVK-LTS VQKGSSADEA VLPNSSAAKA VMPNSSAAKA V-PNSSAAKA	QMVEYGQVKR QMVEYGQVKR SP QMVEYGQVKR GLKRGDLVTK GIKAGDVITS GIKAGDVITS GIKAGDVITS	GFLGVTILAL GELGIMGTEL GELGIMGTEL 350 VNDKVIKSPI LNGKPISSFA LNGKPISSFA LNGKPISSFA
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E.coli Htrab S. typhimurium Htrac C. jejuni Htrad Consensus C. jejuni Htrad E.coli Htrad S. typhimurium Htrac C. jejuni Htrad Consensus C. jejuni Htrad E.coli Htrad E.coli Htrad S. typhimurium Htrac C. jejuni Htrad	GINSAILSRG GINTAILAPD GINTAILAPD GINSAIFLVV GIN-AILAPD 301 QGNTKKAYK. NSELAKAMKV NSELAKAMKV 351 QGNTKKAYK. ALRAQVGTMP ALRAQVGTMP ALRAQVGTMP SLRNLDPRLK EGAEMSNKGK EGAEMSNKGK EGAEMSNKGK 451 EIKNLKDLEQ AVKNIAELRK PVKNIAELRK	GGNIGIGFAI GGNIGIGFAI GGNIGIGFAI GGNNGIGFAI .NQEGALITD DAQRGAFVSQ DAQRGAFVSQ DAA: GAFVSQ **YOEGALITD VGSKLTLGLL VGS; SLGLL **JQ DK DK DK DK **DVGSKISIGLL **D	PSNMVKNLTS PSNMVKNLTS PSNMVKDLQA PSNMVK-LTS VQKGSSADEA VLPNSSAAKA V-PNSSAAKA V-PNSSAAKA V-PNSSAAKA VCKGSSADEA RDGKQVNVNL REGKAITVNL GVLVHSVKEK GVVVNNVKTG GVVVSSVKAN GVVV-SVK TKVC/TkNGF LNICRODRHL	QMVEYGQVKR QMVEYGQVKR SP QMVEYGQVKR GLKRGDLVTK GIKAGDVITS GIKAGDVITS GLKRGDLVTK ELQQSSQNQV ELQQSSQNQV ELQQSSQNQV SKGKNSGFQE TPAAQIGLKK SPAAQIGLKK SPAAQIGLKK ATLLVLK LVNAVISLNPSI.	GFLGVTILAL GELGIMGTEL GELGIMGTEL 350 VNDKVIKSPI LNGKPISSFA LNGKPISSFA LNGKPISSFA LNGKPISSFA LNGKPISSFA 400 VNDKVIKSPI D.SSSIFNGI D.SSSIFNGI D.SSTIFSGI 450 GDIIIGVGQS GDVIIGANQQ GDVIIGANQQ GDVIIGANQQ GDVIIGANQQ FLICKERGSPY

(Figure 6-5). Because ORF2 was found adjacent to the C. jejuni htrA, and appears to be expressed with htrA in an operon, it was designated ntrB.

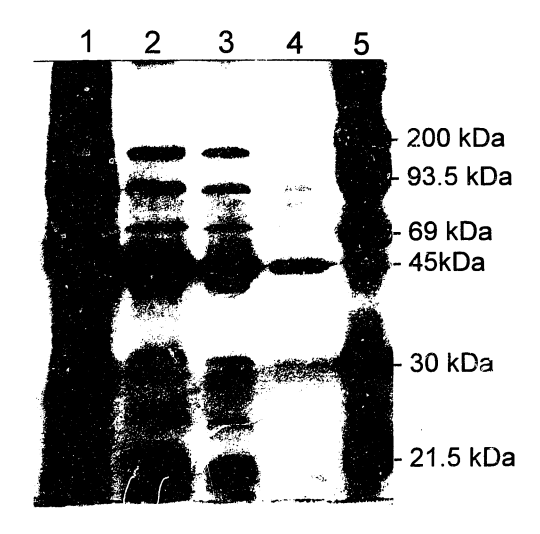
C.2. N-terminal amino acid sequencing of the MOMP of C. jejuni UA580

By obtaining the N-terminal sequence of the purified MOMP protein, it would be possible to design oligonucleotide probes to identify clones which carry the MOMP gene. SDS-PAGE of MOMP preparations and Western blots of these gels to PVDF membranes were carried out for N-terminal sequence analysis. SDS-PAGE revealed that the MOMP preparations were not pure (Figure 6-6). These were the same MOMP preparations used to inoculate rabbits in the preparation of antiserum against the MOMP protein (Taylor and Hiratsuka, 1990) which was subsequently used for the screening of a λ -gtll library The MOMP band at approximately 45 of C. jejuni UA580 DNA. kDa was excised from Western blots of the protein on PVDF membranes for N-terminal sequencing analysis. The first 30 amino acids of the MOMP protein were determined (Figure 6-7). This amino acid sequence was capared with those found in the SwissPro protein database and was found to have 59% identity with the N-verminal sequence of two porin proteins from the organism Campylobacter rectus, formerly Wolinella recta (Figure 6-7).

The amino-acid sequence was intended to be used for designing oligonucleotide probes for detection of MOMP clones of *C. jejuni*. However, the focus of my research was shifted

Figure 6-5. Amino acid sequence comparisons of the putative C. jejuni HtrB protein with sequences found in GenBank

- a- Campylobacter jejuni "HtrB". Translation of the DNA sequence of ORF2 of plasmid pDT1719. This thesis. Accession #U27271
- b- Bacillus subtilis Phop. Translation of DNA sequence determined by Lee and Hullet, 1992. Accession #X67676
- c- Porphyridium aerugineum OmpR. Translation of DNA sequence determined by Kessler et al., 1992. Accession #X62579
- d- Escherichia coli OmpR. Translation of DNA sequence determined by Wurtzel et al., 1982. Accession #J01656
- e- E. coli CpxR. Translation of DNA sequence determined by Dong et al., 1993. Accession #L14579
- f- B. subtilis "OmpR". Translation of DNA sequence
 determined by Yamamoto et a! , 1991. Accession #L09228
- g- E. coli BaeR. Translation of DNA sequence determined by Nagasawa et al., 1993. Accession #D14054
- h- Streptomyces lividans CutR. Translation of DNA sequence determined by Tseng and Chen, 1991. Accession #S67525
- i- Salmonella typhimurium TctD. Translation of DNA sequence determined by Widenhorn, et al., 1989. Accession #28368
- j- Streptococcus pneumoniae CiaR. Translation of DNA sequence determined by Guenzi, et al., 1994. Accession #X77249



SDS-PAGE of purified MOMP samples from C. Figure 6-6. jejuni UA580

Lane 1- C. jejuni UA580 sonicate Lane 2- 10 μ l of MOMP preparation Lane 3- 5 μ l of MOMP preparation Lane 4- 1 μ l of MOMP preparation Lane 5- molecular weight marker

C. rectus 43 kDa OMP TPLEEAIKDV DFSGFAAYXY TGNKLXVNNK
C. rectus 51 kDa OMP TPLEEAIKDV DLSGYAXYXY NNITVKKNPG
C. jejuni 45 kDa MOMP TPLEEAIKDV DVSGL.RYRY DTGNFDKNFV
Cc. sensus TPLEEAIK-V D-SG-ARY-Y --GT-DKN--

Figure 6-7. N-terminal amino acid sequence of the *C. jejuni* UA580 45 kDa MOMP and comparison with the N-terminal sequence of the *C. rectus* 43 kDa and 51 kDa membrane proteins

The $\mathcal{C}.$ rectus sequences were determined by Kennell and Holt, 1991.

to work on *H. pylori* and no further research was conducted on the *C. jejuni* MCMP.

D. Discussion

D.1. Characterization of pDT1719 and pDT1720

DNA sequencing of the C. jejuni UA580 DNA fragments cloned in pDT1719 and pDT1720 was carried out for this research. Amino acid and DNA homology analyses were carried out on the DNA sequences and deduced amino acid sequences of the three ORFs encoded by pDT1719 and pDT1720 against the GenBank database. Because of strong homology observed with the htrA genes of E. coli and S. typhimurium, ORF1 was designated In E. coli this protein is expressed in response to environmental stress and is necessary for survival at elevated temperatures (Lipinska et al., 1989). The exact role of these proteins is not clear but they are believed to function as proteases which specifically degrade protein byproducts and abnormal proteins which accumulate in the periplasm or inner membrane during heat shock (Lipinska et al., 1990). In A. coli, expression of the htrA gene is controlled the protein (Erickson and Gross, 1989). Because the 5% end of the OPF1 sequence was not cloned, the control of expression of the putative HtrA protein in C. jejuni remairs clear.

ORF2, which was designated htrB since it appeared to be linked to htrA in an operon, showed homology with several response-regulatory elements involved in two-component

In these two-component regulatory regulatory systems. systems, one component, the s asory receptor, responds to environmental factors, activating the second component, the response-regulatory element (Stock et al., 1990). In most instances, the sensory receptor is a protein kinase which first activates itself by auto-phosphorylation, then transfers the phosphoryl group to the response-regulatory element thereby activating it. This response-regulatory element then goes on to specifically activate or repress transcription of a gene or set of genes to respond to the environmental conditions (Stock et al., 1990). The environmental factors which lead to the activation of the putative regulatory element I have cloned, htrB, could not be Since htrB was linked to htrA, a serine determined. protease/Lest shock protein, it was possible that htrB was involved in the heat shock response of C. jejuni. Ιn addition, since these two genes appear to be linked together, it would be interesting to determine if and how their products interact. The HtrA protein may have kinase activity and may activate the HtrB protein in a two-component regulatory system responding to heat shock. It is also possible that the HtrA protein estivates the HtrB protein by specific protease activity. A third possibility is that the HtrB protein is activated by a protein other than HtrA. Further research is needed to answer these questions.

Since the preparation of MOMP used to raise antibodies was shown to contain other proteins (Figure 6-6), the HtrA or B

proteins, as well as the protein encoded by ORF3, were likely present in the "MOMP" preparation, and antibodies against these proteins were raised. This would account for their detection and isolation from the λ -gtl1 expression library. Since porins are identified by their secondary and tertiary amino acid structure, even though ORF3 did not show homology to any DNA or protein sequence in the GenBank database, it was still possible that it represented the C-terminal portion of the MOMP protein. If this was the case, then the complete sequence of this gene would have to be determined and the three-dimensional structure of the protein it encodes would identify it as a porin.

D.2. Comparison of the N-terminal sequence of the c. jejuni MOMP to the N-terminal sequence of the porin protein of Campylobacter rectus

The results of the DNA sequencing of the two Campylobacter probes revealed that no complete open reading frames had been cloned. The sequence also showed that none of ORFs had homology with any know porin proteins. It was possible that the MOMP protein could not be expressed in the expression vector system and perhaps a different strategy was needed to identify MOMP clones. By determining the N-terminal amino acid sequence of the MOMP protein of C. jejuni UAD80, oligonucleotide probes could be designed based on this sequence and and for identification of MOMP clones.

The first thirty amino acids of the N-terminal amino acid sequence of the 45 kDa MOMP of C. jejuni UA580 were determined and represented the first reported sequence for This amino acid sequence was found to have this protein. extensive homology with two membrane proteins (45 and 51 kDa) of C. rectus (Kennell and Holt, 1991). The 51 kDa protein was shown to be a porin, while the function of the 45 kDa protein could not be determined. The MOMP of C. jejuni has also been demonstrated to be a porin (Huyer et al., 1986). The extensive homology observed between the N-terminal amino acid sequences of the outer membrane proteins of C. rectus and C. jejuni was consistent with the observed similarities in rRNA sequences between C. rectus and other Campylobacter species and supports the reclassification of C. rectus from its former genus, Wolinella, to the genus Campylobacter. This comparison demonstrates the possible use of the MOMP Nterminal amino acid sequence as a taxonomic marker for the If the gene for this MOMP could be genus Campylobac'-cloned in the files: all or part of the gene may be useful as a genus-specific probe for Campylobacter species.

Chapter 7

General Discussion

A. Helicobacter pylori genetics

A.1. Genomic libraries of H. pylori UA802

An objective of this research was to construct genomic DNA libraries of H. pylori UA802 DNA. This was accomplished using the lambda cloning vector λ -DASH. Less efficient libraries were also constructed with other vectors (pUC20, pHC79, pSa747, and λ -GEM11). With all of these different vectors, restriction modification was one of the main barriers which had to be overcome to enable efficient library The λ -DASH libraries were constructed using construction. the host strain SRB which, although highly mutated for restriction modification functions, had been found to be less efficient than other strains, such as E. coli ER1792 (Phadnis et al., 1993). E. coli ER1792 and E. coli SRB share the same restriction modification phenotype, but Phadnis et al. showed E. coli ER1792 to be almost one hundred times more efficient than SRB as an H. pylori lambda host. Both strains carry deletions of the mcrc-hsdRMS-mmr region, but at different points. Phadnis et al. speculated that another restriction modification locus specific for H. pylori modified DNA may have been affected by the deletion in E. coli ER1792 but not in E. coli SRB, or that E. coli ER1792 had an H. pylorispecific modification function not found in E. coli SRB. combined effects of a number of factors, such as sequencerelated factors (*E. coli* promoter-like sequences), and lethal genetic elements, in addition to restriction modification, may also have contributed to the low efficiencies observed in library construction (Clayton *et al.*, 1989a; Morrison and Jaurin, 1990; Covacci *et al.*, 1993).

My results show that the libraries were not representative of the entire genome. When screened with either the 16S or 23S probes, 7.5-18% of the plaques were positive. d the expected 2% value estimated for a greatly ibrary. This suggests that these clones must represe have had some selective advantage over the other clones or that conditions selected for these clones or selected against other clones. This phenomenon was also observed during the construction of genomic libraries of the nematode Caenorhabditis elegans (Ellis et al., 1986; Gibson et al., 1987), where rDNA sequences were cloned at a frequency much higher than expected using both lambda vectors and cosmid vectors. Two possible reasons were advanced for this phenomenon (Gibson et al., 1987). First, since rDNA did not encode a protein, it was less likely to produce toxic products and may have had greater stability. Second, the G + C content of the rDNA (49.3 mol%) was much higher than the rest of the genome (36 mol%), reducing sequence-related instability. Similarly, the DNA sequence of the cloned 23S-5S genes of H. pylori UA802 also showed a higher G + C ratio (45.3 mol%) than the ratio calculated for the H. pylori genome (35.2 mol%; Béji et al., 1988). Other genes or

regions of the chromosome with higher localized G + C ratios may have a similar advantage over regions with lower G + C be represented at a higher than normal and thus may a frequency. Co. sely, regions which have a lower G + C content may be poorly represented in such libraries. hypothesis that localized G + C ratios can affect the stability of cloned DNA is corroborated by the results of genomic library construction by Bukanov and Berg (1994), who produced a genomic library of H. pylori NCTC11368 using the This vector was designed to cosmid vector Lorist6. counteract the effects of sequence-related instabilities associated with AT-rich DNA. With this vector, 16S and 23S genes were cloned at the expected frequencies. For this research screening of the libraries was not carried out on primary plates, but remer on plates made from overlays of the primary plates. It was possible that the frequency of 23S and 16S clones on the palmary plates may have been closer to the expected frequency and any advantage that these clones may have had were amplified in the overlay plates.

Other researchers have shown that genomic libraries are not representative of entire genomes. In particular, certain regions of the *H. pylori* genome appear to be toxic to *E. coli* host strains and appear to be poorly represented or unclonable. The cagA (Covacci et al., 1993) and vacA (Telford et al., 1994) genes were examples of this phenomenon. In addition, three regions of the *H. pylori* genome appeared as gaps in the ordered cosmid map constructed

by Bukanov and Berg (1994). Similar gaps have also been observed during the construction of ordered genomic libraries of *E. coli* (Kohara *et al.*, 1987) and *Mycobacterium leprae* (Eiglmeier *et al.*, 1993).

I have successfully constructed genomic libraries of H. pylori UA802 DNA using the lambda vector λ -DASH II in the host strain E. coli SRB. Although my results indicate that these libraries are not representative of the entire genome, they were used in the isolation of genes encoding a P-type ATPase and the rRNA of H. pylori UA802. To circumvent problems associated with variable representation of certain regions of the genome, the concurrent construction of libraries with other restriction enzymes or other vector systems such as Lorist6 (Bukanov and Berg, 1994) could be Amplification of these libraries through an done. alternative host strain, such as E. coli ER1792, may improve efficiency by reducing the effects of the restriction modification barrier.

A.2. Cloning and characterization of the H. pylori UA802 hpCopA, hpCopP, and rRNA genes

The P-type ATPase genes

Another of the objectives of my research was to clone the putative P-type ATPase of H. pylori UA802. Twenty-three clones were isolated which hybridized to a probe specific for the H. pylori P-type ATPase, from the λ -DASH II genomic libraries, as described in Chapter 4. A 2.7 kb fragment from

one of these clones, $\lambda P14$, was subcloned in pBluescript SK+ and characterized by Ge et al., 1995. Three open reading frames were found by Ge et al.. The first ORF encoded a protein, designated HpCopA, which showed extensive homology with a copper transport protein, CopA, from Enterococcus hirae (Odermatt et al., 1993). The second ORF encoded a protein, designated HpCopP, which had homology with the heavy-metal-binding protein MerP (Barrineau et al., 1984). No homology could be determined for the protein encoded by the third ORF. Knock-out mutants of hpCopA and hpCopP were constructed by Ge et al. and showed increase nsitivity to copper ions, while having similar susception that to other The knock-out mutants were heavy metals and omeprazole. susceptible to omeprazole at levels similar to that of the wild-type, which was not expected. Since omeprazole targets the gastric human P-type ATPases, it was thought that omeprazole would also target the P-type ATPase of H. pylori. Data obtained by Ge et al. showed that the HpCopA/P proteins were not affected by omeprazole in vitro and that these proteins were involved in copper transport.

Ribosomal RNA genes

The cloning and characterization of the rRNA genes of $H.\ pylori$ UA802 were also objectives of this research. A 16S gene and both copies of the 23S and 5S genes were cloned from the λ -DASH II libraries of $H.\ pylori$ UA802 DNA. Extensive characterization of the 16S gene of $H.\ pylori$ has already been published (Romaniuk et al., 1987; Eckloff et al., 1994),

so the focus of my work was the characterization of the 23S and 5S genes. Sequence analysis showed that the 23S and 5S genes were closely linked and appeared to be part of an operon, separate from the 16S gene. Three Pribnow box sequences were located in the region 5' to the 23S gene (Figure 4-10). In E. coli, two tandem promoters are present for each rRNA operon (Young and Steitz, 1979; de Boer et al., 1979; Jinks-Robertson and Nomura, 1992). Experimental data with promoter-terminater fusion plasmids indicated that, of the two promoters, the Pl promoter was responsive to growth rate control, while transcription from the P2 promoter was constitutive. In vivo studies have indicated that initiation of transcription originates mostly from the Pl promoter (Sarmientos and Cashel, 1983), although in vitro the promoters have been observed to have relatively equal efficiency (Glaser and Cashel, 1979). This has led to speculation that the two promoters are differentially regulated and respond to different growth conditions (Jinks-Robertson and Nomura, 1992). The fact that three Pribnow boxes could be identified in the 5'-region of the 23S/5S pylori UA802 suggests that H . genes transcriptional control may exist in H. pylori. Further study of the 5'-region of the H. pylori UA802 23S-5S operon, perhaps using promoter fusion plasmids as were used by Condon et al. (1992) with E. coli, are needed to determine how transcription is regulated, where it is initiated and what other regions in the 5'-region are important for transcription.

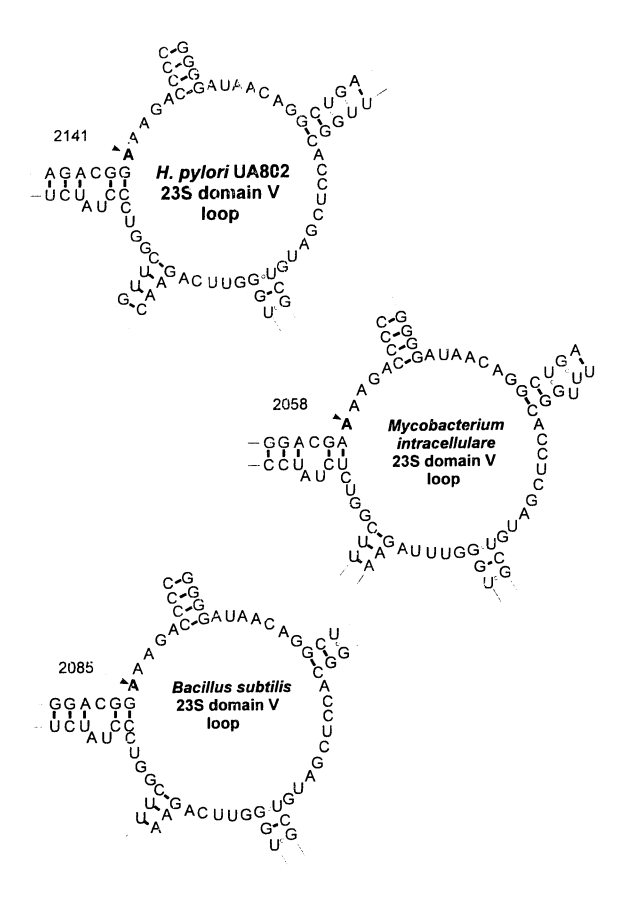
Several antibiotic resistances are specified in rRNA An example of this is the resistance to the macrolide-lincosamide-streptogramin B (MLS)-group antibiotics such as clarithromycin and erythromycin (Skinner et al., 1983; Meier et al., 1994; Zhong et al., 1995). E. coli (Skinner et al., 1983) and B. subtilis (Zhong et al., 1095), the methylation of a specific A-residue in a conserved loop of the 23S rRNA molecule, designated domain V, has been shown to confer resistance to the MLS-group of antibiotics. The methylation is carried out by the methyltransferase ErmC', which can be acquired on a plasmid. The A-2058 residue of E. coli (Skinner et al., 1983) and the A-2085 residue of B. subtilis (Zhong et al., 1995) have been shown to be the methylation site of this enzyme. Recently, resistance to the MLS-group of antibiotics has been shown to arise from the mutation of the A-residue of M. intracellulare (equivalent to the A-2058 in E. coli) to G,C, or T (Meier et The domain V loop in the H. pylori 23S gene al., 1994). sequence is almost identical to those found in B. subtilis and

M. intracellulare (Figure 7-1). The A-2141 residue (2141 bp from the start of the 23S gene or position 2514 on Figure 4-10) is equivalent to the A residues found in the domain V loops of the other organisms. Thus, it is possible that H. pylori could acquire resistance to the MLS-group of

Figure 7-1. Comparison of the domain V loops of the 23S IRNA molecules from H. pylori UA802, Bacillus subtilis and Mycobacterium intracellulare

Arrow indicate the A residues which can be methylated by ErmC'-methylases or mutated to confer resistance to the MLS-group of antibiotics.

The B. subtilis loop was from Zhong et al., 1995. The M. intracellulare loop was from Meier et al., 1994.



antibiotics by the acquisition of the specific methylase (ErmC') or by mutation of residue A-2141.

The complete and partial sequences of the 23S gene and the complete sequences of the 5S gene, determined for this research, were compared with 23S and 5S sequences of several other procaryotes and firmly established the phylogenetic position of H. pylori relative to other Eubacteria. The comparisons of the complete 23S and 5S genes confirmed that, phylogenetically, H. pylori and Campylobacter spp. were similarly distant from other Eubacteria. In addition, the comparisons of the partial 23S sequences confirmed that H. pylori belonged in a sub-class distinct from Campylobacter spp.. These findings correlated with earlier studies of the phylogeny of Campylobacter spp. and related organisms, including H. pylori (Lau et al., 1987; Vandamme et al., 1991; Vandamme and de Ley, 1991, Trust et al., 1994).

A.3. Genomic restriction map and PFGE analyses of H. pylori UA802

Genomic restriction map of H. pylori UA802

A genomic restriction map of *H. pylori* UA802 was constructed which differed from a previously published map of Taylor et al. (1992a). The revised map included *SfiI* restriction digestion data and had an additional *NruI* fragment (*NxuI* fragment 9, Figure 5-3) and an additional *NotI* fragment (*NotI* fragment 5b, Figure 5-3). In addition, the relative positions of some of the fragments were different. The size of the genome of *H. pylori* UA802, estimated from the

previously estimated size of 1710 kb ± 12 for *H. pylori* UA802 determined by Taylor et al. (1992a), as well as with the sizes of the genomes of other *H. pylori* strains which ranged in size from 1674-1730 kb (Bukanov and Berg, 1994; Jiang, 1994).

When the H. pylori UA802 map was compared with genomic maps of three other H. pylori strains (Figure 5-6), significant variability in the arrangement of genes was observed. with this variability, some genes appeared to be clustered together. The hpCopAP genes and the gyrB genes appeared to cluster together in all the strains for which they were mapped, and the vacA, katA, hpaA, pfr, and, with the exception of strain UA861, cagA and flaA genes had similar arrangements in all strains. The vacA, katA, hpaA, cagA and and flaA have been implicated in the pathogenesis of H. pylori (Telford et al., 1994; Tummuru, et al., 1993; Evans et al., 1993; Cover and Blaser, 1995). In a review of chromosome organization, Krawec and Riley (1990) suggest that clustering of genes may indicate that their arrangement may affect the function or regulation of the genes. This may be of particular importance for genes such as vacA and cagA in H. pylori, which are both variably expressed from strain to strain. Through the use of genomic restriction maps, regions which may be important for rearrangement could be localized and these regions could be isolated from genomic libraries to This may lead to an further dissect these regions.

understanding of some of the mechanisms of genomic variability, as well as variability in the mechanisms of pathogenesis of different strains of *H. pylori*.

B. Analysis of the *C. jejuni* DNA probes and MOMP B.1. Characterization of the *C. jejuni* DNA probes

Characterization of the two Campylobacter diagnostic probes (Taylor and Hiratsuka, 1990) was another objective of this research. Sequencing analyses revealed that pDT1720 encoded a partial open reading frame, ORF3. No function could be determined for this partial protein sequence. Sequencing of the cloned DNA of pDT1719 showed that two partial ORFs, ORF1 and ORF2, were encoded by this plasmid. ORF1 had homology with the heat shock/serine protease genes of a number of organisms. ORF 2 exhibited homology with the ompR and ompRlike genes of the response regulators of two-component signalling systems. To my knowledge, no htr/serine protease activity has been associated with a response regulatory element as part of a two-component regulatory system. most instances the sensory element is a protein kinase (Stock et al., 1990).

The gene sequence of the *C. jejuni* MOMP has yet to be determined. The MOMP gene has not been cloned and may be an example of a toxic element (Taylor and Hiratsuka, 1990). As mentioned earlier, the involvement of the MOMF porin in the virulence of *C. jejuni* is not clear, although it has been

implicated in adhesion to eucaryotic cells (de Melo and Pechere, 1991).

The N-terminal sequence of the first 30 amino acids of the purified MOMP was determined. This sequence had significant homology with the *C. rectus* 43 kb and 51 kb major outer membrane proteins. *C. rectus* was formerly called *Wolinella recta*. This organism was recently reclassified and found to belong to *Campylobacter* spp. (Vandamme et al., 1991). The comparisons of the N-terminal amino acid sequence of the *C. jejuni* MOMP to the *C. rectus* MOMPs (Figure 6-5), supported the inclusion of *C. rectus* in the family *Campylobacteriaceae* (Vandamme et al., 1991). In addition, because the *C. rectus* was the only sequence in the database which showed such a degree of homology with the *C. jejuni* MOMP, the N-terminal amino acid sequence of the MOMP may be a possible candidate for a species-specific sequence which can be used for classification of *Campylobacter* species.

C. Future study

There are several possibilities for further research stemming from the results presented in this thesis. The genomic libraries of H. pylori UA802 DNA that I constructed using the vector λ -DASH will be useful as a source of cloned DNA for future work.

The study of the transcriptional control mechanisms of the rRNA genes of *H. pylori* would be interesting. As mentioned earlier, the rRNA genes of most procaryotes are co-

transcribed from a single operon and this large transcript is then processed to give the individual species of rRNA. In this way, the relative amount of each rRNA species is kept equal. In *H. pylori*, the 23/5S genes and the 16S genes are transcribed separately. Thus, the amount of each rRNA species must be regulated by the transcriptional control mechanism. Cloning and characterization of the 5'-flanking region of the 23S/5S operons and the 16S genes would help in determining how transcription of these rRNA genes is regulated in *H. pylori*.

Finally, sequencing of the htrAB genes of C. jejuni UA580 could be completed. The clone pDT1719 could be used to screen genomic libraries of C. jejuni UA580 to obtain the Since the HtrA protein showed significant complete genes. homology to the HtrA heat shock proteins from E. coli and S. typhimurium, it is possible that this protein responds to heat shock. Does this element respond directly to heat shock or is it part of a cascade of reactions responding to heat Transcription of the htrA gene of E. coli is shock? controlled by a novel sigma factor, σ^{E} (Erickson and Gross, 1989), which is expressed in response to heat shock. would be interesting to determine if the C. jojuni htrA gene is under similar control or if it responds directly to some environmental stimulus.

If these two proteins are involved in a two-component regulatory system, it would also be interesting to determine how the signal is transduced from the sensory element (htrA)

to the response element (htrB). The C. jejuni HtrA protein exhibits homology with serine proteases, so it may activate the response element by proteolytic cleavage. It would be interesting to determine what functions are controlled by the response element once it is activated. The complete sequences of these genes may provide more information about the function of these proteins.

The N-terminal sequence of the C. jejuni UA580 MOMP can be used to design oligonucleotide probes to screen DNA libraries of C. jejuni for the isolation of a gene encoding this An alternative to this would be to use the protein. oligonucleotide as a PCR primer, using random primers in the opposite direction. Cloning and sequencing of any products of this PCR would reveal if sections of the MOMP gene were cloned, and these cloned PCR products could then be used to probe C. jejuni DNA libraries to isolate the remainder of the Part or all of the cloned gene of the MOMP may be useful as a genus-specific probe for Campylobacter species, similar to the C. jejuni DNA probe described by Strucki et Strucki et al. described the cloning of a al. (1995). C. jejuni membrane protein gene, which encoded a 24 kDa protein. This gene, when used as a DNA probe, could be used to distinguish C. jejuni from other Campylobacter species.

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Appendix A

Composition of media and solutions

1.2% denaturing agarose gels (Section F.2)	<pre>1.2% (w/v) agarose, 1 X MOPS buffer, 2.2 M formaldehyde</pre>
5% stacking gel (Section G)	5% acrylamide (30:0.8(acrylamide:bis-acrylamide)), 50 mM Tris·HCl, pH 6.8
10% separating gel (Section G)	10% acrylamide (30:0.8(acrylamide:bis-acrylamide)), 180 mM Tris·HCl, pH 8.8
1 X MOPS buffer (Section F.2)	40 mM morpholinopropanesulfonic acid, 10 mM NaOAc, 1 mM EDTA, pH 7.0
10 X PM (Section C.4)	0.2% ficoll(Sigma), 0.2% BSA fraction V (Sigma), 0.2% polyvinyl-pyrrolidone (Sigma) in 10 X SSC
20 X SSC	3.0 M NaCl, 3.0 M Na-citrate
Birnboim and Doly solution 1 (Section B.9)	25 mM Tris-HCl pH 7.5, 2% (w/y) glucose, 10 mM EDTA , 50 μg/ml RNase, 10 mg/ml lysozyme
Birnboim and Doly solution 2 (Section B.9)	0.2N NaOH, 1% (w/v) SDS
Birnboim and Doly solution 3 (Section B.9)	3.0 M NaOAc, pH 4.8
blocking buffer (Section C.4)	maleic acid buffer + 1% blocking reagent (BMC)1%
Ca-buffer (Section B.6)	50 mM CaCl ₂ , 10 mM Tris·HCl pH 7.5
CAPS buffer (Section G)	10 mM 3-[cyclohexylamino]-1- propanesulfonic acid (CAPS), pH 11
DEPC-water (Section F)	200 μ l of diethyl pyrocarbonate (DEPC; Aldrich) were added to 100 ml of dH ₂ O which was incubated at 65°C overnight
denaturing mix (Section F.2)	10% formamide, 2.2 M formaldehyde, 1 X MOPS buffer
denaturing solution (Section B.8)	0.5 M NaOH, 1.5 M NaCl

deproteination solution (Section D.1)	1 mg/ml proteinase K in a solution of 100 mM EDTA, 0.2% (w/v) sodium deoxycholate, 1% (w/v) sarkosyl, 1%(w/v)SDS
destaining solution (Section G)	50% methanol, 10% glacial acetic acid
dNTP mix (Section C.3)	1 mM each of dATP, dCTP, dGTP; 0.65 mM dTTP: 0.35 mM DIG-dUTP; pH 6.5
hybridization solution (Section C.4)	50% formamide, 5 X SSC, 1.0 mM EDTA, 1 x PM, 0.1% SDS, 1% blocking reagent (BMC), 250 µg/ml denatured herring sperm DNA (BMC)
isolation buffer (Section B.1)	0.15 M NaCl, 0.1 M EDTA pH 8.0
LBM-broth (Section B.10)	Luria broth, 10 mM MgSO4, 0.5% (w/v) maltose
loading dye	48% (w/v) sucrose, 0.25% bromophenol blue, 12 mM EDTA, pH 8.0
lysozyme buffer (Section D.1)	10 mg/ml lysozyme in a solution of 100 mM EDTA, 50 mM NaCl, 10 mM Tris·HCl, pH 7.5, 0.2% (w/v) sodium deoxycholate (BDH), 1% (w/v) sarkosyl (BDH)
maleic acid buffer (Section C.4)	100 mM maleic acid, 150 mM NaCl, pH 7.5 (pH adjusted with solid NaOH)
neutralizing solution (Section B.8)	0.5 M Tris·HCl, pH 7.5, 1.5 M NaCl
NT buffer (Section C.3)	25 mM Tris·HCl pH 7.0.5, 5 μM β-mercaptoethanol,20μM dCTP, 20μM dGTP, 20μM dTTP
NT buffer (10 x) (Section C.3)	250 mM Tris·HCl pH 7.0.5, 50 μ M ß-mercaptoethanol, 0.2 mM dCTP, 0.2 mM dGTP, 0.2 mM dTTP
PFGE block washing buffer (Section D.1)	10 mM Tris·HCl, pH 7.0.5, 50 mM EDTA
PFGE running buffer (Section D.3)	0.05 M Tris-base, 0.05 M boric acid, 1 mM EDTA, pH 8.3
PMSF-buffer (Section D.1)	1 mM phenylmethylsulfonyl fluoride (PMSF, Sigma) in PFGE block washing buffer
PPD-buffer (Section C.4)	100 mM Tris·HCl, pH 9.5, 100 mM NaCl, 50 mM MgCl ₂

protoplast buffer (Section F.1)	15 mM Tris·HCl, pH 8.0, 3.6% sucrose, 8 mM EDTA
SDS-PAGE running buffer (Section G)	25 mM Tris-base, 0.192 M glycine, 0.1% (w/v) SDS
SDS-PAGE running dye (Section G)	2%(w/v) SDS, 5%(w/v) B- mercaptoethanol, 10%(w/v) glycerol, 62.5 mM Tris·HCl pH 8.3, 0.1%(w/v) bromophenol blue
staining solution (Section G)	0.1% Coomassie blue, 50% methanol
suspension buffer (Section D.1)	0.25 M EDTA, 10 mM Tris·HCl pH 7.5
TAFE running buffer (Section D.3)	5 mM Tris-base, 5 mM boric acid, 0.1 M EDTA, pH 8.3
TBE-buffer (Section E.3)	0.18M Tris-base, 0.18 M boric acid, 0.2 mM EDTA, pH 8.3
TE	10 mM Tris·HCl, pH 7.5, 1 mM EDTA
TM (Section 2.10)	10 mM Tris·HCL pH 7.0.5,10 mM MgSO4
Tris-borate (TB) buffer (0.5 x) (Section C.1)	45 mM Tris base, 45 mM boric acid, 1 mM EDTA, pH 8.3
wash buffer (Section C.4)	0.3% (v/v) Tween 20 in maleic acid buffer

Appendix B

PCR primers used to amplify gene probes from H. pylori UA802

The primer sequences are shown from 5'-3'. For each gene probe, the top primer represents the forward (sense) primer and the bottom primer represents the reverse (anti-sense) primer.

Gene	Primer sequence	Reference		
vacA	-ATGGAAATACAAACACAC-	Cover et al.,		
	-CGCGCTTTGCACAAATCCAG-	1994		
pfr	-CAACACAAAGGAGATACTATG-	Pfeifer et al.,		
·	TGCACTCAATGACCCTAAAAT-	1993		
flaA	-CGGCCCTGAAGCTGTGAGTGAG-	Leying et al.,		
	-TAACACTTGCAGGATCTATTAC-	1992		
flaB	-CAGCGCTCGTAGAAATGCGCACGG-	Suerbaum et al.,		
	-GTTTATACCTATTAATGAATG-	1993		
cagA	-GATAACGCTGTCGCTTCATACG-	Tummuru et al.,		
	-ACCTGCTATGACTAAGCCACTG-	1993		
1 <i>pp</i> 20	-TTAGGGTAGAGTGTGGTAGCT-	Kostrzynska et		
	-ACCTGCTATGACTAAGCCACTG-	al., 1994		
gyrB	-ARMGNCCNGGNATGTAYATHGGN-	Huang, 1992		
	-NGGDATNCCRCKNCCRTCRTC-			
katA	-GTAGTGCATGCTAAAGGAAGC-	Newell et al.,		
	-GGTAACATGAGCTAAAGACTC-	1991		
hpaA	-GATAGAACGATGAAACAAATG-	Evans et al.,		
	-GTCTAACTCACTCAACTCCAT-	1993		

R-A/G Y-T/C M-A/C K-G/T S-G/C W-A/T H-A/C/T B-C/G/T V-A/C/G D-A/G/T N-A/C/G/T

70

Ammendix C

Comparisons of rRNA gene sequences from Bacillus subtilis

Sequences shown below, show comparisons of the 23S and 5S genes from six different rRNA operons from B. subtilis, as well as the flanking regions and intergenic spacer. The 5'-flanking region and first 107 bp of the 23S gene from the rrnI operon were not available.

References: rrnI - Warousek and Hansen, 1983

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rrnO, rrnA, rrnJ and rrn W - Ogasawara et al., 1994 rrnB - Green et al., 1985

The start and end sites of the 23S and 5S genes are indicated by shaded boxes and are labelled.

```
rrnl
    rrno AGACCTTGGG TCTTATAAAC AGAACGTTCC CTGTCTTGTT TAGTTTTGAA GGAACTTTGT TCCTTGAATA
rrna GCCTTGTACA CACCGCCCGT CACACCACGA GAGTTTGTAA CACCCGAAGT CGGTGAGGTA ACCTTTTAGG
rrnj GCCTTGTACA CACCGCCGT CACACCACGA GAGTTTGTAA CACCCGAAGT CGGTGAGGTA ACCTTTTAGG
rrnW GCCTTGTACA CACCGCCCGT CACACCACGA GAGTTTGTAA CACCCGAAGT CGGTGAGGTA ACCTTTTAGG
rynb AGACCTTGGG TCTTATAAAC AGAACGTTCC CTGTCTTGTT TAGTTTTGAA GGAACTTTGT TCCTTGAATA
rrnI ......
rrno AGTTAAGATG GGCCTGTAGC TCAGCTGGTT AGAGCGCACG CCTGATAAGC GTGAGGTCGG TGGTTCGAGT
rrna AGCCAGCCGC CGAAGGTGGG ACAGATGATT GGGGTGAAGT CGTAACAAGG TAGCCGTATC GGAAGGTGCG
rrnJ AGCCAGCCGC CGAAGGTGGG ACAGATGATT GGGGTGAAGT CGTAACAAGG TAGCCGTATC GGAAGGTGCG
rrnW AGCCAGCCGC CGAAGGTGGG ACAGATGATT GGGGTGAAGT CGTAACAAGG TAGCCGTATC GGAAGGTGCG
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rrna GTTGGAAATC ATTCGCAGAG TGTAAAGGCA CAAGGGAGCT TGACTGCGAG ACCTACAAGT CGAGCAGGGA
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rinj GTTGGAAATC ATTCGCAGAG TGTAAAGGCA CAAGGGAGCT TGACTGCGAG ACCTACAAGT CGAGCAGGGA
rinw GTTGGAAATC ATTCGCAGAG TGTAAAGGCA CAAGGGAGCT TGACTGCGAG ACCTACAAGT CGAGCAGGGA
      GTTGGAAATC ATTCGCAGAG TGTAAAGGCA CAAGGGAGCT TGACTGCGAG ACCTACAAGT CGAGCAGGGA
rrni CGAAAGTCGG GCTTAGTGAT CCGGTGGTTC CGCATGGAAG GGCCATCGCT CAACGGATAA AAGCTACCCC
      CGAAAGTCGG GCTTAGTGAT CCGGTGGTTC CGCATGGAAG GGCCATCGCT CAACGGATAA AAGCTACCCC
rrnO
      CGAAAGTCGG GCTTAGTGAT CCGGTGGTTC CGCATGGAAG GGCCATCGCT CAACGGATAA AAGCTACCCC
rrnA
      CGAAAGTCGG GCTTAGTGAT CCGGTGGTTC CGCATGGAAG GGCCATCGCT CAACGGATAA AAGCTACCCC
rrnJ
rrnw CGAAAGTCGG GCTTAGTGAT CCGGTGGTTC CGCATGGAAG GGCCATCGCT CAACGGATAA AAGCTACCCC
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rrnB
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rrnI
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rrnO
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rrnJ
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rrnW
rinb GGGGATAACA GGCTTATCTC CCCCAAGAG. TCCACATCGA CGGGGAGGTT TGGCACCTCG ATGTCGGCTC
                                                                              2940
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rrnW
      TAGTACGAGA GGACCGGGAT GGACGCACCG CTGGTGTACC AGTTGTTCTG CCAAGGGCAT CGCTGGGTAG
rrnB
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rrnJ
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                                         23S gene ends
rrni GAGCTGACAG ATACTAATCG ATCGAGGACT TAACCATATT TTTGAATGAT GTCACACCTG TTATCTAGTT
rrno GAGCTGACAG ATACTAATCG ATCGAGGACT TAACCATATT TTTGAATGAT GTCACATCTG TTATCTAGTT
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rrnJ rrnW rrnB	GAGCTGACAG	ATACTAATCG ATACTAATCG ATACTAATCG	ATCGAGGACT	TAACCATATT	TTTGAATGAT	GTCACACCTG	TTATCTAGTT	
	3291						3360	
rrnI	TTGAGAGAAC	AC						
rrnO	TTGAGAGAAC	AC						
rrnA	TTGAGAGAAC	AC						
rrnJ	TTGAGAGAAC	AC						
rrnW	TTGAGAGAAC	AC						
rrnB	TTGAGAGAAC	ACCTCTCTAA	AGGCGGAAGG	TAAGGAAACT	CCGCTAAGGG	CTCTCACATC	CTGTGAGAAA	
			5S gene 1	oegins				
	3361		*				3430	
rrnI			- F-1					
rrnO		,						
rrnA			F-1					
rrnJ			r i					
rrnW			· F3		•			
rrnB	CGCCCAGTAC	CTTCATCCTG	AAGGCATTTG	TTTGGTGGCG	ATAGCGAAGA	GGTCACACCC	GTTCCCATAC	
	3431						3500	
rrnI		AGTTAAGCTC						
rrnO		AGTTAAGCTC						
rrnA		AGTTAAGCTC						
rrnJ		AGTTAAGCTC						
rrnW		AGTTAAGCTC						
rrnB		AGTTAAGCTC	TTCAGCGCCG	ATGGTA	• • • • • • • • • •	• • • • • • • • • •		
	5S ends							
	35 <u>0</u> 1	3544						
rrnI	- T1	TTAAACCCAG						
rrnO	- 11	TTAAACCCAG						
rrnA		GCACGTTAGT						
rrnJ		TTATTAATTC						
rrnW	CCMAGCAATT	GCACG.TTAG	TGCAATATGG	AGGATTAGCT	CAGC			
rrnB								