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Review Article

HELICOBACTER PYLORI IS A MAJOR HUMAN PATHOGEN AND IS ASSOCIATED WITH CHRONIC GASTRIC INFLAMMATION OF THE GENE HP0102 ENCODING A CONSERVED LPS GLYCOSYLTRANSFERASE IN THE PATHOGENESIS OF *HELICOBACTER PYLORI*

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Abstract:

Peptic ulcer disease and gastric cancer. Contact with host cells is recognized as a signal capable of triggering expression of bacterial genes important for host pathogen interaction. Adherence of H. pylori to the gastric epithelial cell lines AGS and MKN45 strongly upregulated expression of a gene HP0102 in the adhered bacteria as determined by qRT-PCR. In silico analysis suggested that HP0102 shows tremendous sequence conservation among different strains of H. pylori including several Indian clinical isolates and was predicted to encode for a glycosyltransferase enzyme. To elucidate the role of HP0102, a HP0102 knockout strain was constructed (Δ HP0102) and analyzed. The gene was found to be associated with two distinct phenotypes related to pathogenicity. In AGS cell-adhered H. pylori, it has a role in upregulation of cagA, a major virulence factor and consequent induction of the hummingbird phenotype in the infected AGS cells. HP0102 was also found to be involved in the glycosylation of bacterial lipopolysaccharides (LPS) by glycostaining analysis. Bacterial LPS is a major virulence factor triggering the expression of cytokines via TLR 2 and TLR4 dependent signaling cascades.

Results of a cytokine array using the cell culture supernatants of MKN45 cells (expressing both TLR2 and TLR4) infected with either H. pylori wild type or the Δ HP0102 strain suggested that the HP0102 mutant was impaired in inducing the expression of several cytokines including the proinflammatory cytokine IL-8. Further work is under progress to identify more specific functions of HP0102 and also identify other signaling networks that may be affected by LPS and its glycosylation state during the pathogenesis of H. pylori.

Keywords: *Helicobacter pylori, inflammation, peptic ulcer disease and gastric cancer, AGS and MKN45, HP0102 and MKN45 cells.*

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INTRODUCTION:

Helicobacter pylori is a Gram-negative, microaerophilic bacterium which is specialized in colonization of the gastric mucosa.(1) Between 1979-82, Australian pathologist, Robin Warren and Australian gastroenterologist, Barry Marshall, identified *H. pylori* and suggested a link to the development of stomach ulcers. *H. pylori* are slow growing microorganisms but act as a dominant pathogen that can cause peptic ulcers and gastritis that can lead to gastric cancer.(2) Human stomach acts as a reservoir providing most suitable environment for the growth of the *H. pylori*. It's name refers to both its spiral shape and the area of the lower stomach which it habitually colonizes between the stomach and small intestine.(3) The adaptation to such a harsh and acidic environment in human stomach reduces its competition with other bacterium and enhances its ability to cause chronic infection.(4) This makes *H. pylori* one of the most successful human bacterial parasites, which colonizes more than 50% of the human population.

The link between *Helicobacter pylori* and peptic ulcer, first recognized by Barry Marshall and Robin Warren in 1982, provided major insight into human gastric pathology. Colonization ranges from 50-100%, making *Helicobacter pylori* the most common infectious agent of humans in the world today.(4) Because of the prevalence and importance of *Helicobacter pylori* infection, understanding the mechanisms by which it colonizes the gastric mucosa and causes disease has received intense interest. *Helicobacter pylori* possess several putative colonization factors, including urease, various adhesins and flagellar motility, some of which have been shown to be necessary for gastric colonization. The most severe *Helicobacter pylori* mediated disease states are attributed to strains harbouring the cag pathogenicity island (cag PAI), a 40 kb DNA element. Analysis of the cag PAI sequence suggested that it encodes a putative secretion apparatus with homology to type-IV secretion system, which are involved in the transfer of effector macromolecules into host cells.(5,6) In 1994, the National Institute of Health (USA) published an opinion stating that most recurrent duodenal and gastric ulcers were caused by *H. pylori* and recommended that antibiotics be included in the treatment regimen.(7,8) Warren and Marshall were awarded the Nobel Prize in Medicine in 2005 for their work on *H. pylori*. In silico analysis of HP0102 gene product and prediction of its function by checking for conserved domains. Importance of host cell contact in determining the expression levels of HP0102. Construction of ΔHP0102 mutant of *H. pylori* by non vectorial allelic replacement method for

deciphering their role in *H. pylori* pathogenesis. Characterization of the mutant with respect to their adherence (to AGS cell line) and virulence as compared to the parent (wild type) strain. Gene expression and phenotype analysis of HP0102, major virulence genes (wild type and HP0102 mutant) by qRT-PCR following adherence to AGS cell line.

MATERIALS AND METHODS:**Media preparation (9)****Brain Heart Infusion (BHI) Agar medium (300ml)****Components**

Quantity-BHI powder 16.8g, Bacto agar 4.5g
Distilled water Upto 285 ml

The above all components were weighed by using a weighing balance and dissolved in 285ml of double distilled water in a 500ml conical flask and mixed gently.

Then the flask was autoclaved at 121°C, 15lb/sq inch for 15 minutes.

After autoclaving the flask was allowed to cool at room temperature.

Then 5% (15ml) horse serum (GIBCO) was added to the flask.

15ml of media was poured on each sterile petri plates and gently swirled in a laminar air flow chamber.

The plates were left aside over night for solidification and then stored at 4°C.

GC Blood Agar Medium(200ml) (10)**Components**

Quantity- GC agar base 4g, Bacto agar 480mg, Cysteine HCl 80mg, Distilled water Upto 170ml.

The above all components were weighed by using a weighing balance and dissolved in 170 ml of distilled water taken in a 250ml conical flask and mixed gently.

Then the flask was autoclaved at 121°C, 15lb/sq inch for 15 minutes.¹⁷

After autoclaving 13.6ml defibrinated horse blood (Remel) was added to the flask and

incubated at 80°C for 20 minutes and then allowed to cool at 50°C.

After cooling 16ml horse serum (GIBCO) and 560µl isovitalax (BD BBL) was added to the flask and mixed properly. [20µg/ml Chloramphenicol was added to the flask when required.] Then 15ml of media was poured on each sterile petri plates and gently swirled. The plates were left aside over night for solidification and then stored at 4°C.

Brucella Agar(300ml) (11)**Components**

Quantity-Brucella Broth mixture 8.4g, Agar 4.5g(1.5%), Cysteine HCl 120mg, Distilled water Upto 285ml.

Above all the components were weighed by using a weighing machine and dissolved in 170ml of distilled water taken in a 250ml conical flask and mixed gently.

Then flask was autoclaved at 121°C, 15lb/sq inch for 15 minutes.

After autoclaving the flask was allowed to cool at 50°C.

Then 5% (15ml) horse serum (GIBCO) was added to the flask.

15ml of media was poured on each sterile petri plates and gently swirled.

The plates were left aside over night for solidification and then stored at 4°C.

Brucella soft agar plates(12)

Brucella agar also prepared by following the above mentioned protocol and here only the amount of agar added was restricted to 0.3%.

RPMI-1640 Complete Medium (500ml)(13)

Components

Quantity-RPMI-1640 5.194g, **NaHCO₃** 1g, **Fetal Bovine Serum(FBS)** 50ml(10%), **Penstrep** 5ml, **Distilled water** Upto 500ml. 5.194g of RPMI-1640(GIBCO) and 1gm **NaHCO₃**(2%) were weighed and dissolved in 50ml (10%) heat inactivated **Fetal Bovine Serum (FBS)** [Pan Biotech] in a clean 500ml conical flask. 5ml of **Penstrep**(5,000units/ml **penicillin G sodium salt** and 5000µg/ml **streptomycin (monosulfate)**(GIBCO) was added to the flask.

Then the final volume was made upto 500ml by adding required amount of distilled water. The components were mixed properly and filtered with 0.22µm filter units, collected in A50ml sterile falcon tubes and stored at 4°C.

Urea Agar Medium (20ml)

Components

Quantity-Urea agar base 0.42g, **Agar** 0.2g(1%), **Urea**(40% stock) 0.8ml, **Distilled water** Upto 20ml.

Maintenance of *Helicobacter pylori* culture

A heavy loopful of *H. pylori* culture was scrapped out from 48 hour old GC blood agar plate and streaked onto a fresh GC agar plate with a sterile inoculating loop for mass culturing of the bacteria. □ The plate was then incubated at 37°C under microaerophilic conditions(10% CO₂) for 48 hours and subcultured every 2 days.

Maintenance of AGS cell line

A confluent T-25 tissue culture flask was taken and the media was discarded completely.

The flask was washed twice with 3ml 1X PBS.

1ml Trypsin-EDTA(0.5%) was added and incubated at 37°C.

3ml of complete RPMI-1640 medium was added and the bottom of the flask was washed gently.

The media was collected in a 15ml tube and centrifuged at 1200rpm for 5 minutes.

The supernatant was discarded and the pellet was resuspended in 1ml of RPMI-1640

complete media (including 10% FBS and Pencillin+streptomycin).

Adherence of *H. pylori* with AGS cell line

A heavy loopful of bacterial culture was scraped out from a 48 hour cultured GC plate

and suspended in 1ml filtered PBS in a 1.5ml microcentrifuge tube.

The tube was centrifuged at 4000 rpm for 5mins at room temperature.

The pellet was washed with 1X PBS and the pellet was re-suspended in RPMI-1640

incomplete medium(without serum and antibiotics).

The Optical Density of the cells were measured at 600nm by UV spectrophotometer and the volume for 1 O.D cells were calculated (1 O.D=5×10⁸ cells).

The required amount of cells (5×10⁸) was added to the AGS monolayer at multiplicity of infection(MOI) of 50 in 0.5ml of RPMI media.

The T-25 flask was incubated at 37°C under microaerophilic condition as described above.

The cultured AGS cell line in T-25 flask was observed under phase contrast microscope (Zeiss) for confluency.

The experimental flask was washed with 3ml of filtered 1X PBS and then PBS was pipette out completely.

4ml of incomplete media was added in the experimental and control flask(without the cell line).

5×10⁸ cells(*H. pylori*) added in both experimental and control flask.

1920 500µl of FBS(Pan Biotech) was added to both the experimental and control flask.

The flasks were incubated under microaerophilic condition for 2 hours.

After incubation 10µl of supernatant was taken from both the flasks for CFU(Colony

Forming Unit) assay. This was considered as CFU taken before wash.

The control flask was kept for RNA isolation.

Isolation of adhered bacteria flask (14)

The media was fully decanted from the flask and washed thrice with 3ml 1X PBS.

Again 1ml of 1X PBS was added and 10 µl was taken from it and mixed with 990 µl

1XPBS for CFU assay. This was considered as CFU taken after last wash.

The remaining PBS was decanted and 1ml of 0.25% Triton X 100 was added and incubated for 2 minutes at room temperature.

The lysed mixture was taken in an eppendorf tube and vortexed vigorously for 1-2

minutes and then centrifuged at 8000rpm for 2 minutes.

Triton X 100 (which had no effect on the viability of *H. pylori*) was decanted and 1ml of PBS was added and vortexed vigorously.

10 µl of the above mixture was taken for CFU assay (this was considered as CFU taken for adhered bacterial cells) and the remaining solution was kept for RNA isolation.

0.5µl seeded in one T25 flask so that the cells were expected to get fully confluent after 48 hours of incubation at 37°C in a 10% CO₂ incubator.

RNA isolation from *H. pylori* cells by Trizol method(15)

A loopful of *H. pylori* cells were scraped out from a 48 hour old cultured GC blood

agar plate and suspended in 1ml of filtered 1X PBS in a 1.5ml eppendorf tube.

The tube was centrifuged at 8000rpm for 2 minutes at room temperature.

Supernatant was discarded and pellet was resuspended in 1ml Trizol.

The samples were incubated and vortexed vigorously for 15 minutes in ice to permit the complete lysis of bacterial cells. 200µl of Chloroform per ml of Trizol was added, shaken vigorously by hand and vortexed for 15 minutes and intermittently kept in ice. Then the samples were incubated in ice for 15 minutes, then at room temperature for 2-3 minutes.

Then the samples were centrifuged at 12000g for 15 minutes at 4°C.

400µl of the aqueous phase was transferred into a fresh tube.

400µl of Isopropanol was added and mixed by inversion for precipitating the RNA from the aqueous phase.

The sample was vortexed and incubated at room temperature for 10 minutes.

The samples were centrifuged at 12000g for 10 minutes at 4°C.

The supernatant was discarded and the pellet was washed in 1ml of 75% ethanol and stored at -70°C overnight. RNA pellet in 1ml of 75% ethanol was mixed by vortexing and centrifuged by 7500g for 10 minutes at 4°C.

The RNA pellet was dried briefly for 5-10 minutes and dissolved in 10µl DEPC (Diethyl pyrocarbonate) water then vortexed for 10 minutes.

Then heated at 65-70°C for 5-10 minutes.

Vortexed for 2 minutes and kept in ice.

From the total content 2µl of the sample was taken and mixed with 998µl of DEPC water. The absorbance (O.D) of the solution was taken at 260nm.

Isolation of chromosomal DNA by Sodium Perchlorate (NaClO₄) Method(16)

A loopful of *H. pylori* culture was scrapped out from the 48 hour old GC blood agar plate and suspended in 1ml of 1X filtered PBS.

The sample was then centrifuged at 6000rpm for 3mins.

The pellet was dissolved in 100µl of Sodium chloride Tris EDTA (STE) (0.1M)

0.5% SDS from 10% SDS stock and 5µl of Proteinase K (50µg/ml) was added to the sample.

The sample was incubated at 37°C for overnight.

Next day the samples were brought at room temperature and again 5µl 10% SDS was added.

Equal volume of Tris saturated phenol (120µl) was added and kept on ice for 10 minutes. The sample was centrifuged at 6000rpm for 5-10 minutes and the aqueous supernatant was taken in a fresh eppendorf tube.

Then equal volume of Phenol chloroform (1:1) was added and kept on ice for 10 minutes.

Again it was centrifuged at 6000rpm for 10 minutes and the aqueous supernatant was taken in a fresh eppendorf tube.

1M sodium perchlorate (NaClO₄) was added and then equal volume of chloroform was added to the sample and then kept on ice for 10 minutes.

The sample was centrifuged at 6000rpm for 10 minutes and the supernatant was taken in a fresh eppendorf tube.

4µl RNase (composition: 10mg RNase, 15mM NaCl, 5mM Tris pH8, volume adjusted to 1ml) was added to the sample and incubated at 37°C for 30 minutes.

Then equal volume of chloroform was added to the sample and kept on ice for 10 minutes.

The sample was centrifuged at 6000rpm for 10 minutes. This step was again repeated.

The supernatant was taken in a fresh eppendorf tube and 1ml of cold ethanol was added to the sample.

The sample was then centrifuged at 13000rpm for 15 minutes.

The pellet was dissolved in 200µl distilled water and stored at 4°C.

Isolation of plasmid DNA (*pRY109*) from *Escherichia coli*(17)

1.5ml overnight culture was taken in a eppendorf tube centrifuged at 8000rpm for 2 minutes.

The supernatant was discarded and another 1.5ml overnight culture was added to the pellet.

Centrifuged at 8000 rpm for 2 minutes and supernatant was discarded.

Pellet was resuspended in 100 µl solution-I

Composition (18)

50mM glucose, 25mM Tris chloride (pH-8), 10mM EDTA] and kept in ice for 5 minutes.

200 µl freshly prepared solution-II [composition: 4M NaOH and 10% SDS] was then added and kept in ice for 5 minutes.

150 µl solution-III [composition: 5M CH₃COOK (pH4.8) and glacial acetic acid] was then added and kept on ice for 20 minutes. The mixture was then centrifuged at 10,000 rpm for 10 minutes.

The supernatant was transferred to a fresh eppendorf tube and equal volume of phenol-chloroform was added to the tube mixed well and kept in ice for 3 minutes.

The mixture was centrifuged at 10,000 rpm for 10 minutes.

Aqueous layer was transferred in a fresh eppendorf tube and equal volume of chloroform was added to the tube mixed well and kept on ice for 3 minutes.

Then the mixture was centrifuged at 10,000 rpm for 10 minutes twice.

Aqueous layer was transferred in a fresh eppendorf tube. 0.3M NaCl was added to each eppendorf tube. Equal volume of isopropanol was added mixed well and kept at room temperature for 1 hour. Centrifuged at 13,000 rpm for 20 minutes.

Supernatant was discarded and the pellet was resuspended in 1 ml of 80% ethanol and kept in ice for 20 minutes. Centrifuged at 13,000 rpm for 20 minutes.

Supernatant was discarded and the pellet was allowed to dry at room temperature until it became transparent. 4 µl RNase (50-100 µg/ml) was added to the pellet and incubated at 37°C for 30 minutes. The pellet was then dissolved in 10-20 µl of millipore water or in TE buffer.

PCR-Polymerase chain reaction(19)

PCR (Polymerase Chain Reaction) is a revolutionary method developed by Kary Mullis in the 1980s. PCR is based on using the ability of DNA polymerase to synthesize new strand of DNA complementary to the offered template strand. Because DNA polymerase can add a nucleotide only onto a pre-existing 3'-OH group, it needs a primer to which it can add the first nucleotide. This requirement makes it possible to delineate a specific region of template sequence that the researcher wants to amplify. At the end of the PCR

reaction, the specific sequence will be accumulated in billions of copies. 2324

Quantitative Real-Time PCR(qRT-PCR) (20)

The purity of the RNA sample was checked by spectrophotometer.

2 µg of RNA isolated from *H. pylori* cells incubated under different conditions using Trizol reagent (Invitrogen) was taken for the quantitative real time analysis.

Samples were treated with RNase free DNase I (1U/ µg, Roche) in the presence of an RNase inhibitor (RNasin, Invitrogen).

DNase Step(21)

Sample

Quantity/Volume-DNase (1U/ µl) 2 µl 10X DNase Buffer 2 µl RNA 2 µg DEPC treated water Up to 20 µl Samples were then incubated for 40 minutes at room temperature.

2 µl (25mM) EDTA was added to the samples.

Samples were incubated in a water bath at 75°C for 10 minutes.

After 10 minutes samples were taken out and immediately kept in ice for „heat chill“ shock.

Quantitative real time PCR (qRT-PCR) was performed using SYBR green one step RT-PCR kit (Takara) in an iCycler IQ5, Real-time PCR detection system (Roche).

The wells of the plate were loaded with the above reagents and with RNA and primers of choice, covered with a piece of transparent sealing tape and the plate was then rotated for 5 minutes. The plate was then placed on the Roche Real-time PCR detection system after setting the following thermal cycling parameter. A melting curve was also generated to verify that a single product was amplified. From threshold cycle (CT) values, the relative levels of expression of the genes under different conditions were calculated using the 2-ΔΔCT method using 16S rRNA as control. 26

Construction of Chloramphenicol acetyl transferase(*cat*) (22)

Cassette for vector free allelic replacement mutagenesis

The vector-free allelic-replacement mutagenesis strategy used to generate HP0102 deficient mutant. Chloramphenicol resistance (Cmr) cassette *cat* originating from was amplified with the primers *cat* F and *cat* R (listed in Table-2) and the plasmid *pRY109* was used as the template. *H. pylori* SS1 genomic DNA was isolated as previously described. Four pairs of gene specific primers designated HP0102 P1/HP0102 P2 (for HP0102) and HP0102 P3/HP0102 P4 (for HP0102) were used to PCR amplify the up- and downstream regions, respectively, of each target gene in order to produce fragments of approximately 500 base

pairs (bp) flanking the region to be deleted. The P2 and P3 primers contained 5' overhangs complementary to the cat gene sequence. Amplicons generated with each primer pair (Left flank, Right flank, and cat) were amplified in a second PCR with P1 and P4 primers to generate the long PCR product (mutation construct or allelic replacement cassette). This product was used for natural transformation of *H. pylori*, followed by selection on the appropriate antibiotic chloramphenicol in this case. 2728

Purification of the PCR products using kit(QIAGEN)(23)

The products obtained from the above PCR reaction were subjected to purification using QIAGEN PCR purification kit. 5 volumes Buffer PB was added to 1 volume of the PCR product mixture.

A QIAquick column was placed in a provided 2 ml collection tube

To bind DNA, the sample was applied on to the QIAquick column, allowed to stand for 5 minutes and centrifuged at 13,000rpm for 30–60 seconds.

The flow-through was discarded and the QIAquick column was placed back in the same tube.

To wash, 0.75 ml Buffer PE was added to the QIAquick column and centrifuged at 13,000 rpm for 60 seconds, The flow-through was discarded and the QIAquick column was placed back in the same tube.

The QIAquick column was centrifuged once more at 13,000 rpm for 60 seconds in the provided 2 ml collection tube to remove residual wash buffer.

Then each QIAquick column was placed in a clean 1.5 ml microcentrifuge tube.

To elute DNA, 50 µl milliQ water (pH 7.0–8.5) was added to the center of the QIAquick membrane, was allowed to stand for 5 minutes and the column was centrifuged at 13,000rpm for 60 seconds.

Visualization of the PCR product by Agarose gel electrophoresis(24)

10 µl of the purified PCR product was mixed with 5 µl 6X gel loading dye and was run on a 2% agarose gel along with a 100bp ladder and then the bands were visualized under gel doc (Biorad).30

Splicing by Overlap Extension(SOE) PCR(25)

The corresponding flanking sequences of HP0102 was taken and mixed with the amplified cat gene in a 1:1:1 proportion.

All the reagents except the primers P1 and P4 (corresponding to HP0102) were added and the reaction mixtures were placed in a thermal cycler after setting the following

thermal cycling parameter.

After 6 cycles the thermal cycling was paused and primers P1 and P4 were added.

Then again thermal cycling was continued abiding the following protocol.

Components

Volume-Template 2 µl(left flank)+0.5 µl(cat)+1 µl(right flank) 10X Taq buffer(Thermo scientific) 5 µl 10mM dNTPS 2 µl Taq polymerase(5U/µl)(Thermo scientific) 0.5 µl Forward primer (10pm)2 µl Reverse primer (10pm) 2 µl MilliQ water

Upto 50 µl Visualization of the PCR product by Agarose gel electrophoresis(26)

10 µl of the purified PCR product was mixed with 5 µl 5X gel loading dye and was run on a 1.5% agarose gel along with a 1kb ladder and then the bands were visualized under gel doc (Biorad).

Transformation of *Helicobacter pylori* cells with the mutant Constructs(27)

DAY-1

A heavy loopful of *H. pylori* wild type cells were scraped out from a GC blood agar plate and washed with 1X PBS.

Then it was divided equally in two 1.5ml microcentrifuge tubes and centrifuged at 8000rpm for 2 minutes.

The supernatant was discarded and the pellet in each tube was resuspended in 100 µl of BHI broth.

100 µl of the each of the mutant constructs(for ΔHP0102) prepared by SOE PCR

and was added in one microcentrifuge tube containing the resuspended culture.

The entire mixture from microcentrifuge tube was dropped on the centre of GC blood agar plate and was then incubated for 24 hrs under microaerophilic conditions for uptake of DNA by bacterial cells.

DAY-2

After 24hrs of incubation the, bacterial cultures were streaked onto chloramphenicol(cam) GC blood agar plate to obtain isolated colonies.

Then the plates were incubated for 48hours at 37°C under microaerophilic condition.

DAY-4

After 48 hours of incubation 10 single colonies from mutant was picked with sterile toothpicks and then streaked onto chloramphenicol (20 µg/ml) GC blood agar plate.

Then the plate was incubated for 48hrs at 37°C under microaerophilic condition.

Microscopic analysis of AGS cell lines following its adherence with *H. pylori* wild type and *H. pylori* ΔHP0102 mutant cells.

A heavy loopful of bacterial cultures(wild type, ΔHP0102)were scraped out from 48 hour cultured GC plates and suspended in 1ml filtered PBS in a 1.5ml microfuge tubes.

The tubes were centrifuged at 4000 rpm for 5mins at room temperature.

The pellets were washed with 1X PBS and the pellet were resuspended in RPMI- 1640 incomplete medium.(without serum and antibiotics).

The O.D of the cells were measured at 600nm by UV spectrophotometer and the volume for 1 O.D cells were calculated (1 O.D=5×10⁸ cells).

The required amount of cells (5×10⁸) was added to the AGS cell line monolayer of 50% confluency grown in T-25 flasks at multiplicity of infection(MOI) of 50 in 0.5ml of RPMI media. One of the flasks was incubated with 5x10⁸ wild type cells,

one with 5x10⁸ ΔHP0102 mutants and the third one with 5x10⁸ mutants. Another T- 25 flask was kept as control in which no bacterial cells were added.

The flasks were then incubated under microaerophilic conditions (10% CO₂).

After every one hour following infection the flasks were checked under the microscope for visible morphological changes in the AGS cell line.

Images of the infected AGS cell line were taken after 5 hours following infection.

AGS cell line fixation for microscopy

Materials- 1X PBS 25% Glutaraldehyde 40% Formalin

Method

5ml of cell fixation solution (PBS-4.25ml, 40% Formalin-250μl, 25% Glutaraldehyde-500μl) was prepared for cell fixation.

The tissue culture flask was taken, the media fully decanted from it and then washed with 1X PBS twice.

1ml of fixative solution was then added to the flask and allowed to stand for atleast 10 minutes.

3233 Cell fixative solution was fully decanted from the flask and washed with 1X PBS twice.

2ml of 1X PBS was added to the flask to prevent the cells from drying.

Finally images were taken under phase contrast microscope as necessary.

Isolation of *H. pylori* Lipopolysaccharides (LPS)(28)

H. pylori strains grown in GC agar plates for 48 hours were harvested (1 O.D),resuspended in 200 μl of 1X SDS-buffer (2% β-mercaptoethanol, 2% SDS and 10% glycerol in 0.1M Tris-HCl, pH 6.8, pinch of bromophenol blue), boiled for 15 minutes, followed by incubation at room temperature for 15 minutes.

Next, 5 μl of both DNaseI (10mg/ml) and RNaseA (10mg/ml) were added and incubated at 37° C for 30 minutes, followed by addition of 10 μl of

Proteinase K (10mg/ml) and incubation at 60°C for 3 hours.

To each sample 200 μl of ice cold Tris saturated phenol was next added and incubated

at 65°C for 15 minutes with occasional vortexing.

The samples were next mixed with 1 ml of diethyl ether at room temperature and centrifuged at 20,600 ×g for 10 minutes.

The bottom blue layer was carefully extracted and 15 μl of the sample was separated on a 12% SDS-polyacrylamide gel and visualized either by silver staining (Pierce) or by using the Pro-Q Emerald 300 Lipopolysaccharide gel staining Kit (Molecular Probes) as per manufacturer's instructions.

RESULTS AND DISCUSSION:

Identification of genes of *H. pylori* which are regulated by attachment to gastric epithelial cells and understanding their role in virulence

Contact with host cells is recognized as a signal capable of triggering expression of bacterial genes important for host pathogen interaction. Previously, it has been demonstrated that the expression of *H. pylori* virulence genes, *cagA* is strongly induced in bacteria following adherence to epithelial cell lines (34). In an endeavor to identify factors necessary for host cell contact dependent virulence gene expression, an adherence assay was utilized where *H.pylori* can adhere to gastric epithelial cells and express its virulence properties under laboratory conditions.

Adherence of *H. pylori* to the gastric epithelial cell line AGS

Different strains of *H. pylori* have been reported to efficiently adhere to AGS cells in-vitro.

To test whether, *H. pylori* strain SS1 can also efficiently adhere to AGS cells an adherence assay was performed. *H. pylori* grown to the mid logarithmic phase of growth were incubated with AGS cell line for different time intervals and the number of AGS-adhered bacteria was determined by CFU assay on BHI agar plates. Results of the CFU analysis suggested that the number of adhered bacteria increased gradually up to 12 hours but did not increase any further when examined up to 24 hours. Adherence of *H. pylori* strain SS1 to the gastric epithelial cell line AGS. (A) Schematic description of adherence assay. (B) *H. pylori* SS1 WT and ΔHP0102 strains were added to semi confluent AGS monolayers at MOI 50 and CFU of the adhered bacteria was estimated at different time intervals.

Previous studies have identified genes that are essential for colonization of *H. pylori* In a mice model (29). The functional roles of only a sub-set of genes identified in these studies are known, while most of the genes lack functional annotation. Moreover, a definite correlation could not be

established between any of these genes and regulation of virulence gene expression.

HP0102 has been identified as factor essential for colonization of *H. pylori* (68). Expression of HP0102 following host cell contact and the role of this gene in *H. pylori* virulence has been investigated in this study and the results are described in the following sections.

Expression of the HP0102 gene is upregulated in *H. pylori* following adherence to AGS cells

Semi-confluent monolayers of AGS cells were infected with *H. pylori*, and expression of the HP0102 gene was examined in the adhered bacteria by qRT-PCR at different time points 35 after adherence. HP0102 expression was also examined in unadhered bacteria grown under identical conditions without cell line or isolated from the supernatant of *H. pylori* infected AGS monolayer. Significant increase in HP0102 expression was consistently observed in AGS adhered *H. pylori* as compared to the unadhered controls. No difference in HP0102 expression was observed between unadhered bacteria grown without cell line and those isolated from the supernatant of infected AGS monolayers, suggesting that HP0102 induction required direct contact of the bacteria with the AGS cells and was not due to a component secreted by the cell line.

Expression of HP0102 in AGS-adhered and unadhered *H. pylori* strains.

Expression of HP0102 at different time points was estimated in unadhered and AGS-adhered *H. pylori* by qRT-PCR. Results represent the average of at least three independent experiments. Transcript levels are expressed relative to that in unadhered WT strain at 2 hours arbitrarily taken as 1. Error bars represent SD. The statistical significance (P values) of the differences in gene expression between AGS adhered and unadhered *H. pylori* strains are indicated. * $P \leq 0.05$; ** $P \leq 0.01$; NS, Not significant.

36 Prevalence and conservation of HP0102 gene in *H. pylori* strains

To detect the prevalence of the HP0102 gene in *H. pylori* strains, the gene sequence of strain 26695 was used as a query to search for homologous sequences in *H. pylori* genomes. The gene was present in all the 69 *H. pylori* strains whose genome sequences are available and a high degree of sequence conservation of the gene (more than 93%) was observed among the different strains. Since it is not known if any of the 69 strains were Indian isolates, 15 *H. pylori* clinical strains isolated from India were examined for the presence of the HP0102 gene. PCR using primers flanking the HP0102 ORF as well as gene internal primers

clearly indicated the presence of HP0102 in all isolates (data not shown). Furthermore, sequencing of the PCR products revealed a high degree of conservation of the gene sequence (Table-4). These results indicated that the HP0102 gene is almost ubiquitously present and highly conserved in *H. pylori* strains.

pylori strains. Sequence conservation was analyzed considering the HP0102 gene sequence of *H. pylori* strain 26695 as reference.

Interestingly, NCBI BLAST (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>) analysis also suggested that HP0102 protein shares homology with glycosyl transferase enzymes that are found in various other bacterial species and contains a conserved glycosyl transferase (GT-2) domain. Further analysis suggested that HP0102 shares similarity with members of protein super-families which are involved in the glycosylation of bacterial Lipopolysaccharide (LPS).

Next, to investigate the role of HP0102, a Δ HP0102 deletion mutant was constructed and analyzed.

Prediction of a putative glycosyl transferase domain in HP0102 protein by NCBI BLAST.

Construction of Δ HP0102 mutant by non vectorial allelic replacement method

The left flanks and right flanks of the genes to be mutated i.e. HP0102 was amplified and the products were observed in an agarose gel. The flanks had sizes around 500bps and distinct band of the expected size was observed in respective lanes denoting that the flanks were properly amplified. The *cat* gene was also amplified from the plasmid *pRY109*. A product of 738bp (between the 700 and 800 bp of the marker ladder) suggested that the *cat* gene was properly amplified. The left and right flank regions of the gene *repG* was amplified as positive controls. Showing the amplified left and right flanks of *repG* (positive control) HP0102 and also the amplified *cat* gene.

Result of mutation cassette construction by SOE PCR

The chloramphenicol resistant cassette (*cat* gene) fused to the left and right flanks of the gene HP0102 was constructed by SOE PCR and was run on a 1.5% Agarose gel with a 1kb ladder as the marker. Presence of a prominent band between 1.5kb and 2kb in respective lanes signified that the SOE PCR was successful. Thus the obtained product was used for transformation. The construct of *cat* flanked by the flanking sequence of *cat* was denoted as *cat* mutation construct and the construct

of cat flanked by the flanking sequences of HP0102 was denoted as Δ HP0102 mutant construct. The exact size of Δ HP0102 mutation construct was 1.76kb. Gel picture showing the mutation constructs, the third lane from the left contains the constructs containing the chloramphenicol resistant cassette with flanks of HP0102 gene.

Mutation constructs for the gene *repG* was generated as a positive control of SOE PCR.

Confirmation of mutant by PCR

To confirm the successful construction of mutant in HP0102 the DNA isolated from the wild type and mutant (Δ HP0102) were subjected to PCR and analyzed by agarose gel electrophoresis. PCR products with primers P1-P4 shows a shift in the HP0102 mutant (Δ HP0102) with respect to the wild type *H. pylori*. (Lane 2-3). Presence of cat gene in the mutant and its absence in the wild type strain (lane 4-5) and presence of HP0102 in the wild type and its absence in the mutant (lane 6-7) proved that the cat cassette successfully replaced the HP0102 gene. Successful amplification of *cagA* for both the wild type and mutant (Δ HP0102) acted as the positive control lane (lane 8-9).

Agarose gel electrophoresis of the PCR samples for mutant confirmation. Lane2-Wildtype genome amplified with primers HP0102 P1 and HP0102 P4. Lane3-genome of Δ HP0102 mutant amplified with primers HP0102 P1 and HP0102 P4. Lane 4-5 wild type and Δ HP0102 mutant amplified with primers *cat* internal F and HP0102 P1. Lane 6-7 wild type and Δ HP0102 mutant amplified with the primers of HP0102 F and HP0102 R. Lane 8-9 wild type and Δ HP0102 mutant amplified with the primers of *cagA* (*cagA* F and *cagA* R).

Following mutant construction the relative growth rates of *H. pylori* WT and Δ HP0102 mutant was estimated by CFU analysis. Results of the CFU analysis suggested that both the WT and Δ HP0102 strains demonstrated similar growth kinetics and further suggested that HP0102 was not essential for growth of *H. pylori* under the growth conditions used in our laboratory.

Growth kinetics of HP0102 wild type and mutant. **HP0102 is required for the glycosylation of *H. pylori* LPS** LPS isolated from both the *H. pylori* SS1 WT and Δ HP0102 strains were separated by SDS PAGE and visualized either by silver staining or by using a glycolipid staining kit. A similar pattern of bands corresponding to LPS could be observed by silver staining for both the strains (Fig.12A). Upon staining with the glycolipid staining kit, bands corresponding to the glycosylated LPS could be observed in the WT sample. However, no visible bands could be

observed for LPS isolated from Δ HP0102 strain when stained using the glycolipid staining kit, suggesting a role of HP0102 in LPS glycosylation. Bacterial LPS is known to be involved in the process of adhesion to host cells in the early stages of infection. Taken together with the fact that the gene HP0102 is also strongly upregulated in AGS-adhered *H. pylori*, the possibility that the gene might have a role in pathogenesis was next investigated.

Marker Δ HP0102 WT Δ HP0102 WT HP0102 is required for the glycosylation of *H. pylori* LPS. Analysis of *H. pylori* LPS isolated from SS1 WT and Δ HP0102 by (A) Silver staining and (B) Glycolipid staining.

Role of HP0102 in induction of the hummingbird phenotype in AGS cells

H. pylori SS1 WT and SS1 Δ HP0102 strains were added to AGS monolayers at MOI 50 and the number of adhered bacteria was enumerated at different times after adherence. No difference in the number of AGS-adhered bacteria was observed between the WT and Δ HP0102 strains upto 24 hours examined, indicating that HP0102 might have no role in the adherence of *H. pylori* to gastric cells. However, a remarkable difference in morphology was observed between AGS cells infected with WT or Δ HP0102 strain.

When infected with the *H. pylori* SS1 WT strain, AGS cells exhibited the characteristic elongation and scattering, known as the hummingbird phenotype. However, much lower scattering and elongation was observed in AGS cells after adherence of the Δ HP0102 strain.

Morphology of AGS cells following adherence of *H. pylori* strains. (A) AGS monolayers were incubated with *H. pylori* SS1 (WT) and Δ HP0102 for 5 hours, washed to remove unadhered bacteria and observed on an inverted microscope in phase contrast at 20 45magnification. The number of elongated cells were counted and expressed as percent of total cells in each sample. Bar diagram represents the means \pm SD of 3 experiments. The statistical significance of the differences (P values) in number of elongated cells between WT and the mutant strain is indicated. * $P \leq 0.05$; *** $P \leq 0.001$.

Since the hummingbird phenotype is associated with production and subsequent translocation of CagA from adhered *H. pylori* to the gastric cells, *cagA* expression was next examined in AGS-adhered SS1 WT and Δ HP0102 cells.

Role of HP0102 in *cagA* induction in AGS-adhered *H. pylori*

Expression of the *cagA* gene was observed in WT and Δ HP0102 strains following adherence to AGS cells and compared to that in unadhered bacteria. As reported earlier (Raghwan et al., 2013), about 4 fold upregulation of *cagA* expression was consistently observed in the WT strain within 2 hours of adherence. However, consistent with the lack of hummingbird formation practically *cagA* expression was not observed in AGS-adhered Δ HP0102 strain upto 8 hours examination period (In the complement Δ HP0102/HP0102 strain, *cagA* expression was similar to the WT strain. The difference was not observed in *cagA* expression of unadhered bacteria between WT and Δ HP0102 strains.

Expression of *cagA* in AGS-adhered and unadhered *H. pylori* strains. Expression of *cagA* at different time points was estimated in unadhered and AGS-adhered WT and Δ HP0102 mutant by qRT-PCR. Results represent the average of at least three independent experiments. Transcript levels are expressed relative to that in unadhered WT strain arbitrarily taken as 1. Error bars represent SD. The statistical significance (P values) of the differences in gene expression between AGS adhered and unadhered *H. pylori* strains are indicated. * $P \leq 0.05$; ** $P \leq 0.01$; NS, Not significant.

Next, Western blot analysis determined that the levels of *CagA* protein also increased in AGS adhered WT bacteria but not in the mutants of HP0102 (Fig.15). Hence, significantly *CagA* phosphorylation was observed in WT infected AGS cells. While, only minor amounts were detected in AGS cells infected with Δ HP0102. These results suggested that the Δ HP0102 mutant was specifically defective in induction of *cagA* following adherence to AGS cells, indicating that HP0102 might have a role in host cell contact dependent upregulation of *cagA* expression in *H. pylori*.

Expression of HP0102 and *cagA* Is Upregulated in *H. pylori* Following Adherence to MKN-45 Cells

H. pylori LPS has been reported to induce a cytokine response in host cell upon infection via the receptors TLR2 and TLR4. Since AGS do not express these receptors, we next examined whether HP0102 has a role in *cagA* upregulation during *H. pylori* infection of MKN-45 cells which is a gastric epithelial cell line expressing TLR2 and TLR4. Upon adherence of *H. pylori* WT to MKN-45 cells, HP0102 and *cagA* were both upregulated, but the upregulation was not observed upon infection with the Δ HP0102 strain (Figure 16), similar to the results obtained with the AGS cell line.

CONCLUSION:

The ubiquitous presence and high degree of conservation of the HP0102 gene in all the 84 *H. pylori* strains including 15 Indian isolates examined, as well as upregulation of expression of the gene following adherence to gastric cells suggested that the gene might have an important function in pathogenicity. Indeed, it has been demonstrated in this study that HP0102 has a role in two different processes related to pathogenicity, a) induction of the hummingbird phenotype in AGS cells following *H. pylori* adherence and b) LPS glycosylation.

Initial analysis by qRT-PCR revealed that the mRNA levels of HP0102 increases after adherence to AGS and MKN-45 cells, further suggesting that HP0102 gene product must have some role after it infects AGS cells. Next, for understanding the role of HP0102, a deletion mutant was constructed by non vectorial allelic replacement method in *H. pylori* strain G27. The growth kinetics and the rate of adherence to AGS cells of the HP0102 mutants was similar to that of the wild type. However, it was also seen that the severity of infection in AGS cells (demonstrated by the typical humming bird phenotype) was much less when AGS cells were infected with Δ HP0102 mutants as compared to the wild type strain. Unlike wild type *H. pylori*, expression of the gene *cagA*, responsible for this phenotype was not found to increase in HP0102 mutants following host cell contact. Further investigation is required to understand how HP0102 influences *cagA* expression upon host cell contact and whether the results obtained in a cell culture model also hold true in vivo.

HP0102 was also found to be involved in the glycosylation of bacterial lipopolysaccharides (LPS) by glycostaining analysis. Bacterial LPS is a major virulence factor triggering the expression of cytokines via TLR 2 and TLR4 dependent signaling cascades. Absence of the gene HP0102 led to impaired production of several cytokines including CCL20, CXCL1, IL8, CD147, CCL3/4, CXCL10, etc in MKN-45 cells. The results indicate that HP0102 seems to have a significant role in inducing the inflammatory response in host cells upon infection with *H. pylori*. Further work is under progress to identify more specific functions of HP0102 and also identify other signaling networks that may be affected by LPS and its glycosylation state during the pathogenesis of *H. pylori*. Overall, HP0102 appears to be an important factor involved in the regulation of LPS structure and the induction of *cagA* expression (a major determinant of gastric cancer) upon host cell contact. Since, *cagA* is the major toxin responsible for inducing gastric cancer, decrease in levels of

cagA in the absence of HP0102 gene product might be of importance in developing therapeutic approaches to cure infection or cancer caused by *H. pylori*.

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