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**Predominant mucosal *IL-8* mRNA expression in non-*cagA* Thais is riskfor gastric cancer**

Yamada S *et al*. *IL-8* mRNA expression and gastric cancer

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**Abstract**

**AIM:** To study mucosal *Interleukine-8* (*IL-8*) mRNA expression, the *cagA* mutation, and serum pepsinogen I/II ratio related risk in Thai gastric cancer.

**METHODS:** There were consent 134 Thai non-cancer volunteers who underwent endoscopic NBI examination, and 86 Thais advance gastric cancer patients who underwent endoscopic mucosal biopsies and gastric surgery. Tissue samples were taken by endoscopy with 3 points biopsies. The serum pepsinogen I, II, and *Helicobacter pylori* (*H. pylori*) IgG Antibody for *H. pylori* were tested by ELISA technique. The Histopathology description of gastric cancer and non-cancer with *H. pylori* detection was defined with modified Sydney Score System. Gastric Mucosal tissue *H. pylori* DNA was extracted and genotyped for *cagA* mutation. Tissue *IL-8* and COX-2 mRNA expression were conducted by Real Time relative quantitation polymerase chain reaction. From 17 Japanese advance gastric cancer and 12 benign gastric tissue samples, all were tested for genetic expression with same methods as well as Thai gastric mucosal tissue samples. The multivariate analysis was used for the risk study. Correlation and standardized T-test were done for quantitative data, *P* value <0.05 was considered as a statistically significant.

**RESULTS:** There is a high non *cagA* gene of 86.8 per cent in Thai gastric cancer although there are high yields of the East Asian type in the positive *cagA*. The H. pylori infection prevalence in this study is reported by combined histopathology and H. pylori IgG Ab test with 77.1% and 97.4% of sensitivity and specificity, respectively. The serum PGI/II ratio in gastric cancer is significantly lower than in the non-cancer group, *P* = 0.045. The serum PGI/II ratio of less than 3.0 and *IL-8* mRNA expression ≥ 100 or log10 ≥ 2 are significant cut off risk differences between Thai cancer and non-cancer, *P* = 0.03 and *P* < 0.001, respectively. There is a significantly lower PGI/II ratio in Japanese than that in Thai gastric cancer, *P* = 0.026. Serum PGI/II ratio at cut off less than 3.0 and *IL-8* mRNA expression Raw RQ > 100 or log10 > 2 are significantly difference between Thai cancer group when compared to non-cancer group, *P* = 0.013 and *P* < 0.001, respectively. In the correlation study, low PGI/II ratio does not associate with chronic atrophic gastritis severity score in Thais non-cancer cases. However, there is a trend, but not significant convert correlation between *IL-8* mRNA expression level and low pepsinogen I/II ratio in Thai positive *H. pylori* infection. The high expression of *IL-8* gene demonstrates a poorer prognosis by stage and histology.

**CONCLUSION:** Predominant gastric mucosal *IL-8* mRNA expression level, *H. pylori* infection, and low pepsinogen I/II ratio are relative risks for Thai gastric cancer without correlation with *cagA* mutation.

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**Key words:** Gastric cancer; *cagA* mutation; *Interleukine-8* mRNA expression; *Helicobacter pylori*; Pepsinogen I/II ratio

**Core tip:** A high level of *Interleukine-8* (*IL-8)* mRNA expression was detected in more than eighty percent of Thai gastric cancer patients and nearly two fold in the normal Thai population. The majority of Northern Thai gastric cancer patients who had negative *cagA* gene *Helicobacter pylori* infection even with or without its mutation, still have a high *IL-8* mRNA expression level. The pathogenesis of Thai gastric cancer may primarily involve another gate-way besides the bacterial factor. The results show that there is a predominantly cancer inflammation state regulated by *Il-8* mRNA expression level that can be detected in Thai gastric cancer patients.

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**INTRODUCTION**

Gastric cancer pathogenesis is a well-known worldwide multifactorial condition. The gastric cancer incidence rate in Thailand ranks ninth by 4.1:100 000 in males and 2.1:100 000 in females. Despite being a low incidence country, Northern Thailand has a higher gastric cancer incidence rate with 6.6:100 000 in males that ranks fifth of overall cancer in the Northern Thai region, and 4.5:100 000 in females [1]. The author was interested in the carcinogenesis of gastric cancer in Thais, and why the incidence in Thais is much different from other East Asian countries. The Interleukin-8 (*IL-8*) gene is one of the principal mediators for the inflammatory response gate way that was first reported in 1970s, and it is one of factors that are possible to affect gastric cancer carcinogenesis[2]. A recent case-controlled surveillance study in Northern Thailand on cytokine gene *Il-1b-511* mutations in three East Asian populations showed no predominantly correlated specific causative factor responsible for differences among ethnics and histologic types[3, 4]. Therefore, the author proposed the study on other gate-ways of cytokine expression in the human gastric mucosal cell.

Recently,an *in vitro* study showed the association of the mucosal tissue *IL-8* mRNA expression related to the *Helicobacter pylori (H. pylori),* positivity *cagA* gene. The East Asian genotype was reported in Japanese gastric cancer in about 85 percent of the cases. Many *in vitro* studies showed this toxicity gene related to gastric mucosal cell injury, inflammation, and oncogenic potential[5-7]. The *cagA*, East Asian genotype is commonly detected in chronic gastritis and gastric cancer of the Japanese [8,9].There is reported data that a low serum pepsinogen I/II ratio of less than 3.0 with a pepsinogen I level of less than 70ng/dl was considered as a high risk factor for Japanese gastric cancer[10, 11]. There is no recent *in vivo* study reporting a correlation among these above factors, especially *IL-8* and COX-2 mRNA expression level in Thais.

The author hypothesized that gastric mucosal tissue *IL-8* mRNA expression may be different among ethnicities, and it may correlate to other reference pathogenesis factors. This study aimed to look for the risk and correlation of these factors in Thai gastric cancer. The level of *IL-8*, COX-2 mRNA expression, and *cagA* gene mutation distribution were also to be the first report in Thai gastric cancer.

**MATERIALS AND METHODS**

Research methodology was considered and permitted by Thai and Japanese local ethical committees, the NRCT and Japan Society for the Promotion of Science (JSPS) code ID-NRCT 10726.

***Patient characteristics and volunteer selection***

An experimental based cross-sectional study was conducted in the Gastrointestinal Surgery and Endoscopy Unit, Chiang Mai University Hospital from 2007 -through 2010. Informed consents were obtained from 86 Thai gastric cancer patients who underwent NBI endoscopy and gastric surgery during year 2007-through 2010, and 134 Thai non-cancer volunteers who underwent NBI endoscopic examination from 2006-through 2008. All gastric cancer patients in this study had locally advanced gastric cancer, and underwent examinations by endoscopy before curative gastric resection. Seventeen advanced stage Japanese gastric cancer and 12 non-cancer surveillance patients were recruited. Peptic ulcer disease was excluded in this study. Gastric mucosal tissue samples were taken by endoscopy with three biopsy sites for pathology and bimolecular genetic tests before surgical treatment. In cancer cases, biopsy points were specified from non-necrotic areas of the tumor. The histopathology description of the tumor and histologic type were defined. For pathological examination in both groups, chronic gastritis and metaplasia with *H. pylori* detection were classified with a modified Sydney Score System.

***Serum pepsinogen I and II level, and H. pylori IgG antibody******test***

A five cc sample of venous blood was collected from each study participant. The red blood cell and serum separation was done, and preserved at -20°C. The serum pepsinogen I, II, and *IgG* Antibody for *H. pylori* were tested by the standard ELISA technique. The standard cut off value used was a PGI level of more than 70 ng/mL or PGI/II ratio more than 3.0 for no atrophy or positive Grade 1, PGI <70 ng/mL and PGI/II ratio < 3.0 excluding severe atrophy for moderate atrophy or positive Grade 2, and PGI < 30 ng /mL and PGI/II ratio < 2.0 for severe atrophy or positive Grade 3, respectively[**10,11]**.All samples were tested twice for reliability confirmation (Toyobo, co, Ltd.)

***Tissue H. pylori DNA extraction and cagA genotypting method***

The tissue *H. pylori* DNA extracted from the lower antral position in the stomach was examined by the polymerase chain reaction method, and genotyped for *cagA* mutation in all samples by the author (Samples were also examined by double blinded test by Toyobo, co, Ltd). The *H. pylori*positive control of *cagA* positive strain number 11638 (Western), 26695 (Western), and F57 (East Asian) were provided by the collaborative institute. The bacterial tissue DNA and genotyping method with primers used in this study were conductedas recently described. The specific oligonucleotide primers Forward (5’- AAAAGCGACCTTGAAAATTCC-3’; nucleotides 2299-2319), Reverse-1 (5’-CTTCATTTTTTTGAGCTTGTTGAGC-3’; nucleotides 2488-2463) and Reverse-2 (5’-ATTAATGCGTATGTGGCTGTTAGTAGC-3’; nucleotides 3222-3195, were originally described by Azuma *et al*[12].

***Cell line culture and gastric mucosal total mRNA extraction with reverse transcriptase reaction for cDNA synthesis***

The AGS cell line was grown before cell collection for mRNA extraction at a cell count of 2-4 × 106. They underwent a total mRNA extraction protocol. The technique followed was a reverse transcriptase reaction using a commercial high capacity RNA-to-cDNA kit (Applied Biosystems) [13].

***Gastric mucosal IL-8 and COX-2 mRNA expression by Real time RT-PCR (Relative quantification real time RT-PCR)***

We conducted the experiment from three positions of gastric mucosal biopsies in all Thais and Japanese study participants. All of gastric mucosal tissue samples were transformed to cDNA after total mRNA extraction. The analysis was substantially correctable by analysis both in raw relative quantitation (raw RQ) and log10 value for adjusted normal distribution curve. All Human TaqMan probe primer express that was used in this study had 81-base pairs (bp) *IL-8* specific human primer Assay ID number Hs99999034\_m1, 111- base pairs (bp) COX- 2 Assay ID number Hs01573471\_m1, and 121- base pairs (bp) specific human Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) Hs99999905\_m1 those designed and supplied by Applied Biosystems, United States. The internal control was performed by GAPDH of a matched number template. The Real-Time relative quantitation value was measured by comparing to the base line value of AGS cell line subject control before making the analysis.

***Statistical analysis***

A student t-test was used for quantitative data, *IL-8* and COX-2 mRNA expression level, and pepsinogen level. The 2 test was used for qualitative data. The correlation study for pair factors was done in subgroup analysis for defined groups of ethnic, cancer and non-cancer populations. The multivariate analysis was used for risk study for both non-normal distribution and normal distribution curve data bases. STATA 11.0, United States and SPSS 16, United States were used for statistical analysis,and the *P-* value of less than 0.05 was considered statistically significant.

**RESULTS**

There were 86 cases of advanced gastric cancer and 45 (33.8%) normal control cases, 46 (34.6%) non-peptic disease benign lesions without recent history of any treatment, and 42 (31.6%) chronic gastritis cases among 134 non-cancer control cases who were included in the genetic expression experiment. Thai male and female cancer incidences are 60.5% (52/86) and 34% (39/86), respectively. Males are also the predominant gender in Japanese. Both nations have significantly high incidence of gastric cancer at age 40 years old or above.

The *H. pylori* infection prevalence is reported by combined histopathology, *H. pylori* *IgG* Ab level, and 23SrDNA results that have 77.1% and 97.4% of sensitivity and specificity, respectively. Among Thai cancer patients and non-cancer volunteers, *H. pylori* prevalence was 72.1% and 71.6%, respectively. Meanwhile, Thai gastric cancer cases had a *cagA* genotype demonstrated in only 7/62 (12.3%) in positive *H. pylori* infection cases by 23SrDNA that yields six cases of East Asian type and one case of Western type. In non-cancer volunteers, there were 62/98 (63.9%) of positive *cagA* and 34/98 (36.1%) of negative *cagA* genotyping in positive *H. pylori* infection cases that yielded 47.7% of East Asian, 27.4% of Western, and 24.9% of Mixed genotype. For the six year follow up of 18 cases of high grade chronic atrophic gastritis (CAG group) in non-cancer Thais who had long term *H. pylori cagA* East Asian type infection, no one has developed gastric cancer.

The enzyme pepsinogen results, showed a significantly lower PGI/II ratio with a mean of 3.3 ± 1.7 in gastric cancer patients than one in non-cancer volunteers*, P* = 0.045, and of other chronic active gastritis (CAG), *P* = 0.002. There is a significantly lower PG I/II ratio in Japanese gastric cancer than in Thai gastric cancer, *P* = 0.026.

For *IL-8* and COX-2 mRNA expression results, 86 Thai gastric cancers were tested successfully in comparison with 134 Thai non-cancer volunteers. The detection rates of *IL-8* mRNA expression were 77/86 (89.5%) in Thai gastric cancer and 102/134 (74.6%) in Thai non-cancer volunteers. Thai population characteristic data that was examined for *IL-8* mRNA expressionisdemonstrated inTable 1. Serum enzyme pepsinogen I, II level, and *H. pylori* infection status are demonstrated in the cancer population and non-cancer volunteers in Thais is demonstrated in Table 2. We found a remarkable number of Thai gastric cancers with a negative *cagA*; therefore, *IL-8*mRNA expression was examined and the cut-off point of expression value difference is demonstrated in Table 3. Serum PGI/II ratio at cut-off point of less than 3.0 and Raw RQ ≥ 100 or log10 ≥ 2 of *IL-8* mRNA expression level showed the significantly different between the Thai gastric cancer group and the non-cancer group, *P* = 0.045 and *P <* 0.001, respectively. In the multivariate analysis application, the four co-factors related to gastric cancer risk including *IL-8* mRNA expression in Thais are shown in Table 4.

At the same stage of advanced gastric cancer, the mean levels of *IL-8* mRNA expression in Thai cancer and Japanese cancer were 9,615.65 (log 10 = 2.62) and 1509.11 (log 10 = 2.17), respectively, *P* = 0.014. For gastric cancer risk at cut-off *IL-8* expression level by log10 greater than two 2 in Thais and Japanese, Odds ratio = 7.97 (95%CI: 3.75-16.97, *P* < 0.001) and Odd ratios = 4 (95%CI: 1.29-12.40), respectively**.** In the non-cancer group, we found that the *IL-8* mRNA expression level was lower than cancer population with a significant difference, *P* < 0.001. The total mean *IL-8* mRNA expression in non-cancer Thais was 2,262 (log 10= 1.49) while that in Japanese non-cancer was 10.79 (log10 = 0.69), *P* < 0.001. In comparison within the same ethnic group, the mean levels of *IL-8* mRNA expression in Thai and Japanese cancer were higher than those in non-cancer, *P* = 0.05 as showed in Figure 1.

The COX-2 mRNA expression did not indicate significant rising level with detection rate of 65 percent in Thai and Japanese gastric cancer. In comparison with *IL-8* mRNA expression, although the level of COX-2 mRNA expression was slightly higher in gastric cancer than normal gastric mucosal tissue, there were much lower levels than those of *IL-8* mRNA expression.

In the correlation study, low PGI/II ratio was not associated with the chronic atrophic gastritis (CAG) severity score in Thai non-cancer cases because of a few number of CAG in both Thai gastric cancer and non-cancer populations in this study. There was no significant difference for the *IL-8* mRNA expression level in cancer between positive and negative *H. pylori* infection. There was no direct correlation of *IL-8* mRNA expression level and serum *IgG* levels. In subgroup analysis, there was a significant difference of higher levels in groups of poorly differentiated histopathology in comparing both nations. For the diffuse histologic type, the *IL-8* mRNA expression level is about 1.5 times higher than that of intestinal histologic type with a statistically significant difference in Japanese.

High *IL-8* mRNA expression was primarily found in the non-*cagA* Thai gastric cancer population. There was a significantly different mean *IL-8 mRNA* expression level between groups of negative *cagA* by log10 = 2.46 (±1.04) and positive *cagA* by log 10 = 3.29 (±1.68) in the Thai gastric cancer group. However, there were few numbers of Thai gastric cancers with positive *cagA*. In other subgroup analysis of 18 Thais who had high grade chronic atrophic gastritis, some level of *IL-8* mRNA expression in the 12 Japanese non-cancer patients appeared which an equivalently lower level.

Gastric cancer mucosal tissue *IL-8* mRNA expression in the cancer position had a significantly higher mean level than its level at the non-cancer background position in both Thai and Japanese shown in Figure 2. There was significantly different *IL-8* mRNA expression level between intestinal (favorable) and diffuse (unfavorable) histologic cell types. In Thai gastric cancer, the poorly differentiated gastric adenocarcinoma and signet ring cell were predominantly found in this study. The log10 *IL-8* mRNA mean expressions in unfavorable cell type were 2.55 and 2.85 in Japanese and Thais, respectively, as shown in Table 5. There is a significantly higher level of *IL-8* mRNA expression in diffuse cell type than that in a differentiated histologic cell type, *P* = 0.04. The differentiated histologic cell type in Thais has higher expression level than that in Japanese with a statistically significant difference, *P* = 0.013 The RT-PCR results of gastric mucosal tissue and AGS cell line *IL-8* mRNA expression were demonstrated in Figure 3. and Figure 4**.** However, there was no significant difference of *IL-8* mRNA expression level between positive and negative *H. pylori* infection in subgroup analysis of non-cancer background positions**.**

In summary, *IL-8* mRNA expression level is predominantly found, and trend toward an inverted correlation to PGI/II low ratio in Thai gastric cancer patients. There is no direct correlation of *IL-8* mRNA expression level with the *cagA* gene mutation in Thai gastric cancer.

**DISCUSSION**

In the present *in vivo* study, there is a significant risk of *IL-8* mRNA expression level predominantly found rising up more than 80% of Northern-Thai gastric cancer. There is significantly higher level of *IL-8* mRNA expression in poorly differentiated than in well differentiated carcinoma in both Thailand and Japan. There is a trend of converted independent correlation with the very low PGI/II ratio in gastric cancer as well as a few numbers of Thai severe CAG, but no direct correlation with positive *H. pylori* infection or *cagA* genotypes.

Higher level *IL-8* mRNA expression in non-cancer Thais comparing with non-cancer Japanese demonstrated the difference of gastric mucosal defense and genetic expression between the two nations. Thai gastric cancer has less background of chronic atrophic gastritis. In this study, there is no evidence that showed a direct correlation of *IL-8* mRNA expression with *cagA* mutation genotype in *H. pylori* positive cases. Predominant *IL-8* mRNA expression level resulted in non-atrophic mucosa of both gastric cancer and non-cancer Thais. The result is different from previous *in vitro* or some *in vivo* studies in high incidence gastric cancer countries, such as Japan, China, and Korea.

In Thai gastric cancer patients, *IL-*8 mRNA expression level at the lessor curvature is the most represented location. Its predominantly high level may represent relatively vascular invasion as well as major gastric mucosal inflammation. However, the area of gastric antrum in Thai gastric cancer patients has less activity than the lessor curvature because atrophy occurs more frequently.

Also, a high level of *IL-8* mRNA expression is matched with the poor prognosis by histopathology cell type and tumor stage. Long-term bacterial infection has less effect to change the gastric mucosa into chronic atrophic gastritis. In our recent six year followed up study in non-cancer cases controlled with positive *H. pylori* infection and without eradication of 500 new non-cancer cases, a few people developed severe gastritis and still have a high PGI/II ratio. In this study, the number of high pepsinogen I/II ratio in Thai cancer is still about 45% which is nearly the same percentage in our recently published study [4].

A few gastric cancer preventive models on natural Thai products were reported[14]. There is one study on diet consumption in Thais showing a linkage of gastric cancer risk. The factors which were found to be a higher risk but not statistically significant were low intake of vegetables and fruits (OR=1.2, 95%CI: 0.74-1.96) and Jeaw prik (mainly chilly with Plara broth or pickled fish), a kind of preserved food in North and North-eastern regions of Thailand (OR = 1.2, 95%CI: 0.76-2.01)[15]. The consensus of the Asian Pacific guideline on gastric cancer prevention is still debated in some experts’ opinions[16]. *H. pylori* infection screening in a low risk gastric cancer population is not recommended, but serum pepsinogen may be helpful to screen the high risk population in Northern Thailand. We reported its different characteristics that rely on *H. pylori* *IgG* antibody and pepsinogen I/II ratio in our cancer populationscomparing to the data in a Japanese report[17].

The Japanese study primarily reported the relation of *cag*A genotype and a low pepsinogen I/II ratio. Nevertheless, approximately half of the Thai cancer population demonstrates a low ratio of pepsinogen I/II similar to the Japanese. There are still a small number of Thais who had a severe atrophy score related *to H. pylori* infection though in the positive *cagA* Thai population*.*

In this study, the negative *cagA* gene is found in the majority of Thai gastric cancer unlike the Japanese. In Thais, the poorly differentiated cell type gastric adenocarcinoma occurred mainly in the negative *cagA* gene *H. pylori* infection, not in Western type *cagA*.

There is a small number of the Thai population who had a severe atrophy score of gastric mucosa found in our recent and present collaborative study [18]. Reduction of the fundic gland in chronic gastritis was also related to the low level of PG I/II ratio in Japanese[19, 20]. In our recent study of subgroup analysis on the PGI/II ratios, there was no significant difference between chronic atrophic gastritis and gastric cancer group, *P =* 0.12. However, the low PGI/II ratio was significantly related to gastric cancer when compared to normal population in a recent match-case control study by Odds ratio of 2.3 (95%CI: 1.10 - 4.80), *P* = 0.025[4]. The tumor location demonstrated locations mainly at the upper portion and corpus in both of our studies. In the present study, the author found risk for cancer by Odds ratio 2.06 (95%CI: 0.94-4.47), *P* = 0.059 that seems to be close to the result of our recent study. The low PGI/II ratio was not found to be a high percentage in Thais unlike in Japanese gastric cancer[21].

For other genetic host factors, in one interesting study, two of the four gastric carcinoma cell lines expressed Vascular Endothelial growth factor (VEGFR-3) mRNA. In 17 of 36 gastric carcinoma specimens, VEGFR-3-specific immune activity was detected in tumor cells. These angiogenesis and lymphangiogenesis were also detected in VEGF-C-transfected tumors than in control tumors[22**]**.*IL-8* mRNA expression is found to be the gate way mechanism of the vascular epithelial growth factor. For *IL-1* gene, it is a pro-inflammatory cytokine, and the T/T genotype of IL-1β-511 is suspected as the risk factor of both hypochorhydria related *H. pylori* infection and gastric cancer in a case-control population in the United States[23].The author reported that c/c genotype was a risk in Japanese, and a lower number of c/c genotype wasfound as a minor risk related to Thai gastric cancer[4].

Recent *in vivo* animal models studies showed the expressions of *IL*-8 and COX-2 had linkage to the epithelial cell which was co-infected with *H. pylori*. However, our preliminary study reported that there was no difference of *IL-8* mRNA expression level between a cell line which was co-infected with *H. pylori* and the Thai gastric mucosa tissues which had positive or negative *H. pylori*[24,25]. The toleration, remarkable host response to cancer inflammatory process, healing of stomach mucosal turnover rate, and re-healing process of ulcer in Thais may be different and caused by host susceptible differences to the virulence bacteria.

The theory regarding inflammatory cytokine’s influence on cancer development was first contributed by Rudolph Virchow around 150 years ago[26]. Many studies regarding *IL-8* gene expression remarkably found significant relation with *H. pylori* infection and many *cag* PAIboth *in vitro*[27,28] and in some number in *in vivo* study of Japanese cancer[29,30].However, no study has demonstrated differences of expression level in the individualized host [31].

In this study, Japanese gastric cancer has a lower *IL-8* mRNA expression on average than that in Thai gastric cancer patients at the same stage of disease and *H. pylori* infection status. However, Japanese gastric adenocarcinoma cases are mostly an intestinal type and infected by positive *cagA* strain *H. pylori*. In contrast with Northern Thai gastric adenocarcinoma cases, they are mostly diffuse histologic cell type, and infected by negative *cagA* strain *H. pylori*. Therefore, the authors speculate that the predominant level of *IL-8* mRNA expression found in non-*cagA* gene *H. pylori* infection is not directly related to atrophic gastritis mucosa in Thai gastric cancer.

Some results in this study are unexpected outcomes and different from our recent knowledge of *in vitro* and *in vivo* study in Japanese atrophic gastric mucosa which almost easily infected by positive *cagA* *H. pylori* infection. The *cagA* was also the suspected cause of the *IL-8* gene expression rising*.*

This result in Thais showed that gastric mucosal tissue *IL-8* mRNA expression has a higher level in the advanced stage and poorer differentiated cell type than in favorable histology or differentiated cell type. The author was suspicious that the less atrophic background of Thai stomach cancer and non-cancer gastric mucosa may be caused by non-*cagA* *H. pylori* infection. However, the high level of mRNA *IL-8* gene expression in Thai gastric cancer cases may be explained by the cancer inflammation carcinogenesis that may not be directly related to only *H. pylori* infection in Thais.

The level of *IL-8* mRNA expression in Thai gastric cancer or poorly differentiated gastric carcinoma may be regulated by other factors besides of *H. pylori* infection. Also, unknowns remain regarding how long *IL-8* mRNA expression has been high before the occurrence of gastric cancer or after becoming a more advanced stage. Although the author analyzed the level of *IL-8* and *COX-2* mRNA expression level in normal mucosa and of advanced gastric cancer, this occurrence could not be shown in early gastric cancer. The environmental factors and bacterial virulence effect cannot be excluded.

In this study, the signet ring cell type is predominantly found in Thai gastric cancer population. The poor prognostic histological cell type may have different disease carcinogenesis related to gastric mucosal tissue *IL-8* mRNA high expression level and severity of the disease. The COX-2 mRNA expression level is directly correlated with only the *H. pylori* infection, and tended to be suppressed unlike *IL-8* mRNA expression. The author supposed that extremely high gastric mucosal *IL-8* expression level may relate to other factors, such as VEGF that could not be demonstrated in this study and should be explored further.

In conclusion, this present *in vivo* study shows results of new factual data on predominant gastric mucosal *Il-8* mRNA expression level in Thai gastric mucosal biopsy tissues in both non-cancer and gastric cancer volunteers. In the present study, the Northern Thai gastric cancer population has a high incidence of signet ring cell by the nature of histologic type. The positive *H. pylori* may be one of the co-factors, though the host is infected with non-*cagA* gene and still has an extremely high *IL-8* mRNA expression level. This cytokine expression may represent the individual host defense in both high and low incidence gastric cancer ethnics. The factual results in an experimental based study demonstrated how prevalence of the Northern Thai gastric cancer host had active co-infection or had been recently infected by non-*cagA* gene *H. pylori* infection. The *IL-8* mRNA expression level does not directly correlate to non-*cagA* *H. pylori* infection in Thai gastric cancer*.* However, there is a trend of converted correlation between *IL-8* mRNA expression and low ratio pepsinogen I/II without statistical significance, and it seems to be an independent correlation. By these preliminary results, the author expected to do further study on *IL-8* mRNA expression that may act as one of prognostic genetic biomarkers in clinical practice and for chemotherapy application in the nearby future. A study on the current cancer chemotherapy with Northern Thai gastric cancer population is on-going.

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**COMMENTS**

***Background***

The incidence of gastric carcinoma is very low although the incidence of *Helicobacter pylori (H. pylori)* seemed not to be low in Thais. However, the Northern Thai population still has the highest gastric cancer incidence in Thailand. Currently, the disease incidence is rising faster than in the past and still the second cause of cancer death worldwide. The cause and risk factors for gastric cancer carcinogenesis in Thais is unclear especially the risk related with *H. pylori* infection or other cofactors.

***Research frontiers***

*Interleukine-8* (*IL-8*) mRNAexpression is a common event found in some epithelial malignancies and in gastric adenocarcinoma either due to *H. pylori* caused chronic inflammation or other causes by unrelated carcinogens. It is not clear how the level of this expression related to gastric adenocarcinoma. The authors demonstrated that the predominant overexpression of *IL-8* mRNAcould be a potential relative risk for gastric adenocarcinoma in Thais and demonstrated the difference of its level in relationship with the histologic type of gastric cancer.

***Innovations and breakthroughs***

Recent reports have highlighted the importance of *cagA* gene *H. pylori* infection and its mutation type that is shown predominantly in Japanese gastric cancer carcinogenesis. Particularly in the well differentiated histologic type, the *IL-8* mRNA expression level in the Japanese seems to be much lower than in Thais. However, its level in poorly differentiated cell type of Thai gastric adenocarcinoma has less atrophic background and a higher level of expression than in well differentiated gastric adenocarcinoma. This is the first study to report how measurement of *IL-8* mRNA expression level demonstrates risk in *non-cagA* *H. pylori* infection of Thai gastric cancer and the trend of differences in carcinogenesis related to *H. pylori* infection between high and low incidence ethnics. Furthermore, our *in vivo* studies would suggest that the *IL-8 mRNA* expression level yields high prevalence detection in gastric adenocarcinoma and may be a useful tool for gastric cancer prognostic or therapeutic study.

***Applications***

By understanding different *IL-8* mRNA expression levels, this study may represent a future study with a tissue molecular biomarker for gastric cancer.

***Terminology***

*IL-8 mRNA* expression is pro-inflammatory cytokines that is detected in gastric epithelial mucosa and gastric cancer cell lines, such as Kato III and AGS cells.

***Peer review***

The authors examined the expression of *IL-8* *and Cox-2* in AGS cell line, normal gastric mucosa, gastritis, and gastric adenocarcinoma tissues. It revealed that *IL-8* mRNA expression predominantly increased in poorly differentiated or signet ring cell gastric adenocarcinoma that showed the trend of a poorer prognosis. The expression was not directly correlated to *CagA H. pylori* infection and its mutation type. The results are interesting and may represent a different carcinogenesis of Thai gastric cancer in comparison to recent Japanese studies.

**REFERENCES**

1 **Khuhaprema T**, Srivatanakul P, Stomach. In: Khuhaprema T, Srivatanakul P, Sriplung H, Wiangnon S, Sumitsawan Y, Attasara P, editors. Gastric Cancer. In: Cancer in Thailand Vol. IV, 1998–2000. Bangkok Medical Publisher; Bangkok: 2007; 32-33

2 **Kozlov SV**. Experimental models and practical approaches vol.1 In: Inflammation and cancer SAIC- Frederick, Inc. and National Cancer Institute at Federick, MD 2009; V-VIII

3 **Matsukura N**, Yamada S, Kato S, Tomtitchong P, Tajiri T, Miki M, Matsuhisa T, Yamada N. Genetic differences in interleukin-1 betapolymorphisms among four Asian populations: an analysis of the Asian paradox between H. pylori infection and gastric cancer incidence. *J Exp Clin Cancer Res* 2003; **22**: 47-55 [PMID: 12725322]

4 **Yamada S**, Matsuhisa T, Makonkawkeyoon L, Chaidatch S, Kato S, Matsukura N. Helicobacter pylori infection in combination with the serum pepsinogen I/II ratio and interleukin-1beta-511 polymorphisms are independent risk factors for gastric cancer in Thais. *J Gastroenterol* 2006; **41**: 1169-1177 [PMID: 17287896 DOI: 10.1007/s00535-006-1951-6]

5 **Crabtree JE**, Wyatt JI, Trejdosiewicz LK, Peichl P, Nichols PH, Ramsay N, Primrose JN, Lindley IJ. Interleukin-8 expression in Helicobacter pylori infected, normal, and neoplastic gastroduodenal mucosa. *J Clin Pathol* 1994; **47**: 61-66 [PMID: 8132812]

6 **Wu K**, Crusius JB, Fan D, Peña AS. The immunogenetics and pathogenesis of gastric cancer. Highlights of the First Sino-European Workshop on the Immunogenetics and Pathogenesis of Gastric Cancer. *Drugs Today (Barc)* 2002; **38**: 391-417 [PMID: 12532177]

7 **Aihara M**, Tsuchimoto D, Takizawa H, Azuma A, Wakebe H, Ohmoto Y, Imagawa K, Kikuchi M, Mukaida N, Matsushima K. Mechanisms involved in Helicobacter pylori-induced interleukin-8 production by a gastric cancer cell line, MKN45. *Infect Immun* 1997; **65**: 3218-3224 [PMID: 9234778]

8 **Sasazuki S**, Inoue M, Iwasaki M, Otani T, Yamamoto S, Ikeda S, Hanaoka T, Tsugane S. Effect of Helicobacter pylori infection combined with CagA and pepsinogen status on gastric cancer development among Japanese men and women: a nested case-control study. *Cancer Epidemiol Biomarkers Prev* 2006; **15**: 1341-1347 [PMID: 16835334 DOI: 10.1158/1055-]

9 **Kikuchi S**, Wada O, Miki K, Nakajima T, Nishi T, Kobayashi O, Inaba Y. Serum pepsinogen as a new marker for gastric carcinoma among young adults. Research Group on Prevention of Gastric Carcinoma among Young Adults. *Cancer* 1994; **73**: 2695-2702 [PMID: 8194008 DOI: 10.1002/1097-0142(19940601)]

10 **Azuma T**, Yamazaki S, Yamakawa A, Ohtani M, Muramatsu A, Suto H, Ito Y, Dojo M, Yamazaki Y, Kuriyama M, Keida Y, Higashi H, Hatakeyama M. Association between diversity in the Src homology 2 domain--containing tyrosine phosphatase binding site of Helicobacter pylori CagA protein and gastric atrophy and cancer. *J Infect Dis* 2004; **189**: 820-827 [PMID: 14976598 DOI: 10.1086/381782]

11 **Jones KR**, Joo YM, Jang S, Yoo YJ, Lee HS, Chung IS, Olsen CH, Whitmire JM, Merrell DS, Cha JH. Polymorphism in the CagA EPIYA motif impacts development of gastric cancer. *J Clin Microbiol* 2009; **47**: 959-968 [PMID: 19158258 DOI: 10.1128/JCM.02330-08]

12 **Azuma T**. Helicobacter pylori CagA protein variation associated with gastric cancer in Asia. *J Gastroenterol* 2004; **39**: 97-103 [PMID: 15069615 DOI: 10.1007/s00535-003-1279-4]

13 **Keith WN**, Hoare SF. Detection of telomerase hTERT gene expression and its splice variants by RT-PCR. *Methods Mol Med* 2004; **97**: 297-309 [PMID: 15064501 DOI: doi: 10.1385/1-59259-760-2: 297]

14 **Jagetia GC**, Aggarwal BB. "Spicing up" of the immune system by curcumin. *J Clin Immunol* 2007; **27**: 19-35 [PMID: 17211725 DOI: 10.1007/s10875-006-90667]

15 **Suwanrungruang K**, Sriamporn S, Wiangnon S, Rangsrikajee D, Sookprasert A, Thipsuntornsak N, Satitvipawee P, Poomphakwaen K, Tokudome S. Lifestyle-related risk factors for stomach cancer in northeast Thailand. *Asian Pac J Cancer Prev* 2007; **9**: 71-75 [PMID: 18439078]

16 **Fock KM**, Talley N, Moayyedi P, Hunt R, Azuma T, Sugano K, Xiao SD, Lam SK, Goh KL, Chiba T, Uemura N, Kim JG, Kim N, Ang TL, Mahachai V, Mitchell H, Rani AA, Liou JM, Vilaichone RK, Sollano J. Asia-Pacific consensus guidelines on gastric cancer prevention. *J Gastroenterol Hepatol* 2008; **23**: 351-365 [PMID: 18318820 DOI: 10.1111/j.1440-1746.2008.05314.x.]

17 **Ohata H**, Kitauchi S, Yoshimura N, Mugitani K, Iwane M, Nakamura H, Yoshikawa A, Yanaoka K, Arii K, Tamai H, Shimizu Y, Takeshita T, Mohara O, Ichinose M. Progression of chronic atrophic gastritis associated with Helicobacter pylori infection increases risk of gastric cancer. *Int J Cancer* 2004; **109**: 138-143 [PMID: 14735480 DOI: 10.1002/ijc.11680]

18 **Matsuhisa TM**, Yamada NY, Kato SK, Matsukura NM. Helicobacter pylori infection, mucosal atrophy and intestinal metaplasia in Asian populations: a comparative study in age-, gender- and endoscopic diagnosis-matched subjects. *Helicobacter* 2003; **8**: 29-35 [PMID: 12603614 DOI: 10.1046/j.1523-5378.2003.00121.x]

19 **Miki K**, Ichinose M, Shimizu A, Huang SC, Oka H, Furihata C, Matsushima T, Takahashi K. Serum pepsinogens as a screening test of extensive chronic gastritis. *Gastroenterol Jpn* 1987; **22**: 133-141 [PMID: 3596151 DOI: 0.1007/BF02774209]

20 **Matsuhisa T**, Yamada S. In: the Helicobacter Pylori Infection in Asia( Japanese- English), Nishinura Shoten, Chiyoda-ku, Tokyo 2009; 58-59 , 167-176

21 **Kikuchi S**, Wada O, Miki K, Nakajima T, Nishi T, Kobayashi O, Inaba Y. Serum pepsinogen as a new marker for gastric carcinoma among young adults. Research Group on Prevention of Gastric Carcinoma among Young Adults. *Cancer* 1994; **73**: 2695-2702 [PMID: 8194008]

22 **Kodama M**, Kitadai Y, Tanaka M, Kuwai T, Tanaka S, Oue N, Yasui W, Chayama K. Vascular endothelial growth factor C stimulates progression of human gastric cancer via both autocrine and paracrine mechanisms. *Clin Cancer Res* 2008; **14**: 7205-7214 [PMID: 19010837 DOI: 10.1158/1078-]

23 **El-Omar EM**, Carrington M, Chow WH, McColl KE, Bream JH, Young HA, Herrera J, Lissowska J, Yuan CC, Rothman N, Lanyon G, Martin M, Fraumeni JF, Rabkin CS. Interleukin-1 polymorphisms associated with increased risk of gastric cancer. *Nature* 2000; **404**: 398-402 [PMID: 10746728]

24 **Yamada S,** Makonkawkeyoon L, Jukrabandhu T, Lertprasertsuk N, Matsuhisa T, Azuma T. Correlation of cytokine gene IL-8 expression, cag A mutation of *H. pylori* infection and pepsinogen I/II ratio result: Reflection of host response in proximal gastric cancer. Abstract in Ann Oncol 2008; **19** (Suppl 6): vi29-vi105 [DOI: 10.1093/annonc/mdn361]

25 **Sutharat P**, Kato S, Yamada S, Matsuda N, Matsukura N, Sandhu T, Tajiri T. Racial variations for the risk of stomach carcinogenesis depend on Helicobacter pylori infection and mucosal conditions of stomach (oral presentation abstract symposium) abstract in AACR 100 meeting proceeding 2009; 09-AB

26 **Coussens LM**, Werb Z. Inflammation and cancer. *Nature* 2002; **420**: 860-867 [PMID: 12490959 DOI: 10.1038/nature01322]

27 **Chiou CC**, Chan CC, Sheu DL, Chen KT, Li YS, Chan EC. Helicobacter pylori infection induced alteration of gene expression in human gastric cells. *Gut* 2001; **48**: 598-604 [PMID: 11302954 DOI: 10.1136/gut.48.5.598]

28 **Boonjakuakul JK**, Canfield DR, Solnick JV. Comparison of Helicobacter pylori virulence gene expression in vitro and in the Rhesus macaque. *Infect Immun* 2005; **73**: 4895-4904 [PMID: 16041003 DOI: 10.1128/IAI.73.8.4895-4904.2005]

29 **Lee KH**, Bae SH, Lee JL, Hyun MS, Kim SH, Song SK, Kim HS. Relationship between urokinase-type plasminogen receptor, interleukin-8 gene expression and clinicopathological features in gastric cancer. *Oncology* 2004; **66**: 210-217 [PMID: 15218312]

30 **Park MJ**, Kim KH, Kim HY, Kim K, Cheong J. Bile acid induces expression of COX-2 through the homeodomain transcription factor CDX1 and orphan nuclear receptor SHP in human gastric cancer cells. *Carcinogenesis* 2008; **29**: 2385-2393 [PMID: 18775915 DOI: 10.1093/carcin/bgn207]

31 **Subramaniam D**, Ramalingam S, May R, Dieckgraefe BK, Berg DE, Pothoulakis C, Houchen CW, Wang TC, Anant S. Gastrin-mediated interleukin-8 and cyclooxygenase-2 gene expression: differential transcriptional and posttranscriptional mechanisms. *Gastroenterology* 2008; **134**: 1070-1082 [PMID: 18395088 DOI: 10.1053/j.gastro.2008.01.040.]

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**Figure 1 Mean *interleukine-8* mRNA expressio*n* level measurement of Relative Quantitation by real time real-time reverse transcription-polymerase chain reaction study in gastric cancer comparing with non-cancer population both Japanese and Thais.** In Japanese and Thais, gastric mucosal *Interleukine-8* (*IL-8*) mRNA expression in cancer is higher than in non-cancer with *P* = 0.05. The mean level of *IL-8* mRNA expressions in Thai cancer and Japanese cancer were 9615.65 (log 10 = 2.62) and 1509.11 (log10 = 2.17), respectively, *P* = 0.014. The mean level of *IL-8* mRNA expression in non-cancer Thais is 2,262 (log10 =1.49) while that in non-cancer Japanese is 10.79(log10 = 0.69), *P* < 0.001.

**Figure 2 The mean *interleukine-8* mRNA expression in Thais divided by histology and cancer position Middle is the lessor curvature and non- cancer position, Lower is the antrum, and Upper is the greater curvature.**

**Figures 3 Basic experiment result on real-time reverse transcription-polymerase chain reaction of *interleukine-8* mRNA expression with AGS, macrophage cell line, normal gastric mucosal cell, AGS cancer cell line co-culture with two strains of cagA *Helicobacter pylori*, and positive *cagA*, Thai non-cancer samples sequences showed on 12 lanes.**

**Figure 4 Real-time reverse transcription-polymerase chain reaction result of *interleukine-8* mRNA expression from AGS, Negative control, and Thai gastric cancer mucosal tissues.** The positive results of *IL-8* mRNA expression appeared at 320bp band comparing with 300bp band of marker in lane 1.

**Table 1 Characteristics of 220 Thais examined for *interleukine-8* mRNA expression *n* (%)**

|  |  |  |
| --- | --- | --- |
| **Variable** | **Cancer**  ***n* =86** | **Benign**  ***n* =134** |
| Sex  Male  Female | 52(60.5)  34(39.5) | 41(30.6)  93(69.4) |
| Age (yr)  < 40  > 40  mean ± SD | 5(5.8)  81(94.2)  56(11.3) | 28(20.9)  106(79.1)  48.5(11.2) |
| Alcohol drinking  no  yes | 50(58.1)  36(41.9) | 80(59.7)  54(40.3) |
| Smoking  no  yes | 62(72.1)  24(27.9) | 122(91)  12(9) |
| Diseases  Normal  Benign lesion(polyps, erosion, mild superficial gastritis)  Chronic active gastritis  Cancer | -  -  -  86(100) | 45(33.8)  46(34.6)  42(31.6)  - |

**Table 2 Serum enzyme pepsinogen I, II level and *Helicobacter pylori* infection detection results in Thais.**

|  |  |  |  |
| --- | --- | --- | --- |
| **Variable** | **Cancer**  ***n* = 86** | **Benign**  ***n* = 134** | ***P*-value** |
| PG I (ng/microliter)  mean ±SD | 57.39± 46 | 54.86±68.5 | 0.78 |
| PG II (ng/microliter)  mean ± SD | 19.42± 21 | 15.42±11.7 | 0.09 |
| PG I/II ratio  ≤ 3  >3 | 28(39.4)  43(60.6) | 34(27.9)  88(70.1) | 0.0451 |
| *H. pylori* pathology  Negative *n* (%)  Positive *n* (%) | 31(36.8)  55(63.2) | 60(44.8)  74(55.2) | 0.001 |
| Serum*IgG*  Negative  Positive | 31(46.3)  36(53.7) | 57(44.2)  72(55.8) | 0.821 |
| *Cag A* genotyping in  positive 23SrDNA  Negative  Positive | 55(88.7)  7(12.3) | 62(64.6)  34(35.4) | <0.0011 |
| *H. pylori* Infection status  True Negative  True Positive | 20(24.4)  62(75.6) | 37(36.0)  96(64.0) | 0.121 |

1 Some numbers are not included in the analysis due to missing laboratories entries.

The statistical analysis was performed by 2 of each factor. *H. pylori*: ***Helicobacter pylori*.**

**Table 3 Molecular genetic results of COX-2 and *interleukine-8* mRNA expression in Thais *n* (%)**

|  |  |  |  |
| --- | --- | --- | --- |
| **Variable** | **Cancer**  ***n* = 86** | **Benign**  ***n* = 134** | ***P*-value** |
| *COX2* Raw RQ  No expression detection  expression detection | 30(34.9)  56(65.1) | 43(53.1)  38(46.9) | <0.0011 |
| *COX2* Raw RQ  Mean ±SD | 41.69(4.9) | 5.37(4.2) | <0.0011 |
| COX2 Log 10 (N, %)  Mean ±SD | 1.62(0.96) | 0.73(0.62) | <0.0011 |
| *IL-8* Raw RQ  No expression detection  expression detection | 9(10.47)  77(89.53) | 32(25.37)  102(74.63) | <0.001 |
| *IL-8* Raw RQ  <100 or undetected  >100  Mean± SD | 33(38.37)  53(61.63)  9615.64± 49715.0 | 105(78.36)  29(21.64)  2262.29± 10454.6 | <0.001  <0.01 |
| IL-8 Log 10  <2 or undetected  >2  Mean± SD | 32(37.21)  54(62.79)  2.62±1.1 | 105(78.36)  29(21.64)  1.49±1.2 | <0.001  <0.01 |

1Some numbers are not included in the analysis due to missing laboratory data.

**Table 4 Multivariate risk analysis for Thai gastric cancer**

|  |  |  |  |
| --- | --- | --- | --- |
| **Variable** | **OR** | **95%CI** | ***P*-value** |
| Male | 4.32 | 2.06-9.04 | <0.001 |
| *H. pylori* infection status | 0.98 | 0.96-0.99 | 0.02 |
| PG II/I ratio ≤ 3 | 2.06 | 0.94-4.47 | 0.06 |
| *IL-8* mRNA expression | 7.97 | 3.75-16.97 | <0.001 |

OR: Odds ratio

**Table 5 Comparative means *interleukine-8* mRNA expression level detection between Thai and Japanese Cancer populations divided by histopathology *n* (%)**

|  |  |  |  |
| --- | --- | --- | --- |
| **Histopathology** | **Thai**  **(*n* = 77)** | **Japanese**  **(*n* = 17)** | ***P*-value** |
| Diffuse type  Intestinal type  Mean *IL-8* Log 10 mRNA expression ± SD  Diffuse type  Intestinal type | 55(71.4)  22(28.6)  2.85 ± 1.10  2.52 ± 1.11 | 4(23.5)  13(76.5)  2.55 ± 0.52  1.56 ± 1.06 | 0.011  0.95  0.01  0.041 |

1Statistical difference between groups.