Helicobacter pylori Infection Increased Anti-dsDNA and Enhanced Lupus Severity in Symptomatic FcγRIIb-Deficient Lupus Mice

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The defect on Fc gamma receptor IIb (FcγRIIb), the only inhibitory FcγR, has been identified as one of the genetic factors increasing susceptibility to lupus. The prevalence of Helicobacter pylori (HP) and FcγRIIb dysfunction-polymorphisms are high among Asians, and their co-existence is possible. Unfortunately, the influence of HP against lupus progression in patients with lupus is still controversial. In this study, the interactions between these conditions were tested with HP infection in 24-week-old FcγRIIb−/− mice (symptomatic lupus). HP induced failure to thrive, increased stomach bacterial burdens and stomach injury (histology and cytokines) in both wild-type and FcγRIIb−/− mice. While the severity of HP infection, as determined by these parameters, was not different between both strains, antibodies production (anti-HP, anti-dsDNA and serum gammaglobulin) were higher in FcγRIIb−/− mice compared to wild-type. Accordingly, HP infection also accelerated the severity of lupus as determined by proteinuria, serum creatinine, serum cytokines, renal histology, and renal immune complex deposition. Although HP increased serum cytokines in both wild-type and FcγRIIb−/− mice, the levels were higher in FcγRIIb−/− mice. As such, HP also increased spleen weight and induced several splenic immune cells responsible for antibody productions (activated B cell, plasma cell and follicular helper T cell) in FcγRIIb−/− mice, but not in wild-type. These data describe the different systemic responses against localized HP infection from diverse host genetic background. In conclusion, the mutual interactions between HP and lupus manifestations of FcγRIIb−/− mice were demonstrated in this study. With the prominent immune responses from the loss of inhibitory signaling in FcγRIIb−/− mice, HP infection in these mice induced intense chronic inflammation, increased antibody production, and enhanced lupus severity. Thus, the increased systemic inflammatory responses due to localized HP inducing gastritis in some patients with lupus may enhance lupus progression. More studies are needed.

Keywords: Helicobacter pylori, FcγRIIb-deficient mice, lupus, susceptibility, murine model
INTRODUCTION

*Helicobacter pylori* (HP), microaerophilic, spiral-shaped gram-negative bacteria, are organisms that can survive in the highly acidic stomach environment, and are known to cause chronic gastric inflammation and cancer (Mahachai et al., 2016). The infection is very common among Asians, with a prevalence rate of up to 50–80% in some countries (Thirumurthi and Graham, 2012; Xie and Lu, 2015). Interestingly, eradication of HP in some patients with associated autoimmune diseases leads to long-term remission of the autoimmune disease (Fujimura et al., 2005; Kuwana, 2014). Moreover, HP infection down-regulates the expression of Fc gamma receptor IIb (FcγRIIb), the only inhibitory FcγR (Bolland and Ravetch, 2000) on circulating monocyte of patients with autoimmune diseases (Asahi et al., 2008; Wu et al., 2012). As Fcγ receptors (FcγR) is the immunoglobulin superfamily that contributes to the protective functions, in part, by inducing phagocytosis of opsonized microbes, loss of the inhibitory FcγR results in effective organism control but enhances the risk of autoimmune diseases (Ravetch and Bolland, 2001). Although HP infection has shown a protective effect on the development of lupus in a case control study, especially among African-American patients, the relationship of lupus-HP is still intriguing (Sawalha et al., 2004; Hasni et al., 2011).

Inadvertently, FcγRIIb dysfunction polymorphisms are common in Asia (Chu et al., 2004), partially due to the genetic pressure from malarial infection (Clatworthy et al., 2007). Although FcγRIIb dysfunction protects against malaria, the insufficient inhibitory signaling increases the risk of autoimmune activation. Indeed, the association between FcγRIIb polymorphisms and systemic lupus erythematosus (lupus) in patients has been reported (Tsuchiya and Kyogoku, 2005). Both FcγRIIb dysfunction polymorphisms and HP infection are common in the Asian population (Smith and Clatworthy, 2010; Hooi et al., 2017). While FcγRIIb loss-of-function is associated with lupus (Siriboonrit et al., 2003; Tsuchiya and Kyogoku, 2005; Jakes et al., 2012), HP infection has been associated with other autoimmune diseases such as immune thrombocytopenic purpura and membranous nephropathy (Hasni et al., 2011). As chronic inflammation accelerates lupus (Hasni et al., 2011) and the co-existence of FcγRIIb dysfunction polymorphisms with HP infection are possible, information on the responses of FcγRIIb−/− mice to HP infection in patients with lupus. Thus this study tested HP infection in FcγRIIb−/− condition, *in vivo* and *in vitro*.

MATERIALS AND METHODS

Animal

FcγRIIb−/− mice on C57BL/6 background were kindly provided by Dr. Silvia Bolland (NIAID, NIH, Maryland, United States). Wild-type female C57BL/6 mice were purchased from the National Laboratory Animal Center in Nakhon Pathom Province, Thailand. The animal protocols, as per NIH criteria, were approved by the Faculty of Medicine, Chulalongkorn University. Due to lupus manifestations’ age-dependency, female symptomatic lupus (24-week-old with kidney injury) mice or age-matched wild-type control groups were used. Serum samples were collected through tail-vein nicking and through cardiac puncture at sacrifice for time-course analysis. Mice were sacrificed with cardiac puncture under isoflurane anesthesia and internal organs were collected, fixed in 10% formalin and embedded in paraffin. Staining in 4-µm sections with haematoxylin and eosin color (H&E) were used for further evaluation.

*Helicobacter pylori* Administration Model

HP ATCC 43504 (ATCC, Manassas, VA, United States) was cultured on supplemented Columbia agar (Oxoid, Hampshire, United Kingdom) under microaerophilic conditions (6–12% O2, 5–8% CO2) at 37°C for 48 h before use. The mouse model for HP infection was modified from a previous study (Konturek et al., 1999). Briefly, HP at 2 × 109 CFU/ml in 0.5 ml or phosphate buffer solution (PBS) control were orally administered twice daily for 2 weeks and once daily 3 weeks after. Mice were sacrificed at 1 week after the last administration of HP. Mouse blood was centrifuged and serum was kept at −80°C until analysis. Stomach was divided longitudinally through the greater and lesser curvature into several parts, washed with PBS, weighed and used to test HP burdens (i) by urease test, polymerase chain reaction (PCR) and direct culture, (ii), histopathology (fixed in 10% formaldehyde) and (iii) cytokine analysis from gastric tissue.

Urease Test, a Semi-Quantitative Analysis of Gastric *Helicobacter pylori* Burden

The principle of urease test is based on HP’s urease enzyme production (Midolo and Marshall, 2000). Urease enzyme splits urea metabolites into ammonia and carbon dioxide, and ammonia alkalinizes the culture media (the media color turns from yellow to pink). The stomach specimens (one-fourth of the total stomach tissue; 50 mg) were minced and directly put onto urea agar slant, and incubated at 37°C. The media color was observed at 24 h after incubation. As the color media alteration starts from the top to the bottom (*Figure 3*), the ratio of the pink color to the total depth of the media was used as a semi-quantitative measurement of HP burden.

Polymerase Chain Reaction (PCR), a Quantitative Analysis of Gastric *Helicobacter pylori* Burden

Quantitative real-time PCR of HP from gastric tissue were performed as previously described (He et al., 2002). In short, genomic DNA was extracted by High Pure PCR Template Preparation Kit (Roche, United States), and quantified by spectrophotometry (NanoDropTM, Thermo Fisher Scientific, United States). The primers of *UPEC* gene-fragment were HP-FOR (5′- TTATCGGTTAAAGACACCGAAGA-3′) and HP-REV (5′- ATCACAGCGCATGTCTTC -3′). The amplification product was 132 bp and the genome size of HP ATCC 43504 (also designated HP CCUG 17874) was 1,615,763 bp (Clancy et al.,...
2012). Bacterial genome is approximately 1.06 × 10^9 g/mol and contains 6.02 × 10^{23} molecules/mol. One bacterium corresponds to 1.8 fg of DNA. The constructive of standard curve was created by the LightCycler software using 10-fold serial dilution (3.6 fg-360 pg) per 5 µl of HP DNA, with bacterial concentrations ranging from of 2 to 2 × 10^5 bacteria. The profiling standard curve was indicated as a graph of crossing point (Cp) vs. bacterial number (CFUs). HP quantification was calculated by the standard curve and shown in bacterial number (CFUs).

**Direct Culture, a Quantitative Analysis of Gastric Helicobacter pylori Burden**

As an additional method, burdens of HP from gastric tissue were performed in a selective media (Skene et al., 2007). In brief, gastric samples were weighed, homogenized in 1 ml PBS and serially plated onto the supplemented Columbia agar (Oxoid, Hampshire, United Kingdom) with 10 mg/l vancomycin, 5 mg/l amphotericin B under microaerophilic conditions at 37°C for 48 h before bacterial enumeration.

**Stomach Histology and Anti-Helicobacter pylori Antibody**

Inflammatory response in stomach was scored by 2 blinded observers according to the Sydney Scoring System at 200 × magnification in 10 randomly selected fields of each sample (Dixon et al., 1996). The scoring system was determined by leukocyte mucosal infiltration with the following criteria: Score 0, normal; Score 1, mild; Score 2, moderate; Score 3 marked histopathological changes. In addition, serum anti- H. pylori IgG antibody (anti-HP) was measured by ELISA (My BioSource Inc, SanDiego, CA, United States).

**Cytokines Measurements in Gastric Tissues and Serum**

Cytokine in gastric tissue was measured using the same method as in a previous publication (Ferrero et al., 2008). Briefly, gastric tissues were homogenized (Ultra-Turrax homogenizer, IKA, Staufen, Germany) in 500 µl of PBS containing protease inhibitor, centrifuged at 12,000 × g for 10 min at 4°C. The supernatant was collected and stored at −80°C until analyzed. Quantikine ELISA (ReproTech, NJ, United States) was used to measure CXC chemokines (MIP-2 and KC) and pro-inflammatory cytokines (IL-1β and TNF-α) in the supernatant and the serum.

**Lupus Characteristics (Anti-dsDNA, Proteinuria, Serum Creatinine, and Renal Histology)**

Lupus characteristics were analyzed as per the protocol in previous publication (Sarawut et al., 2017). In brief, serum anti-dsDNA measurement was based on the ELISA assay of calf DNA (Invitrogen, Carlsbad, CA, United States) coated on 96-well plates (Surawut et al., 2017). Urine protein creatinine index (UPCI), a representative of 24 h proteinuria, was determined by the equation; spot urine protein/spot urine creatinine

An Analysis of Serum Gamma Globulin

For the evaluation of antibody responses, mouse serum was analyzed for total immunoglobulin by capillary protein electrophoresis (MINICAP-2 Sebia, Evry Cedex, France). The percentage of protein in the gamma zone of protein electrophoresis was converted into total immunoglobulin level by multiplying the ratio of protein at the gamma zone by serum total protein. Serum total protein and urine protein were measured by Bradford protein assay.

**Flow Cytometry Analysis of Spleen**

Flow cytometric analysis was performed following a standard protocol. In brief, spleens were minced in supplemented RPMI-1640 (Roswell Park Memorial Institute media), and the cells were centrifuged at 300 g for 5 min at 4°C. Red blood cells were removed using lysis buffer (ACK buffer: NH4Cl, KHCO3 and EDTA) and the splenocytes were washed twice in supplemented RPMI-1640. Subsequently, the splenocytes were resuspended in staining buffer (0.5% BSA and 10% FBS in PBS), and then were stained with fluorochrome-conjugated antibodies against different mouse immune cells including; CD19, CD80, CD138, CD3, CD4, CXCR5, B220, CD19, GL-7 and F4/80 (BioLegend). All stained cells were analyzed by flow cytometry using BD LSR-II (BD Biosciences) and data analysis by FlowJo software (version 10).

**Macrophages Phagocytosis and Macrophage Killing Activity**

Macrophages were derived from the bone marrow (BM) (Ondee et al., 2017) and phagocytosis assay was performed following an established protocol (Chmiela et al., 1997; Miliukene et al.,
In short, $1 \times 10^9$ cell/ml heat-killed HP (60°C for 30 min) was incubated with 100 µg/ml fluorescein isothiocyanate (FITC) (Sigma, United States) in PBS at 35°C for 30 min for FITC-labeling. This was then mixed with activated macrophages at multiplicity of infection (MOI) 500:1 (HP-FITC: macrophage) in DMEM complete media with 5% mouse normal serum (as opsonin) in 96-well polystyrene tissue culture plates. Of note, the high MOI was necessary for the adequate fluorescent intensity for the detection of phagocytosis activity. The incubation with 10 ng/ml IFN-γ (Biolegend, United States) for 17 h followed by 100 ng/ml of lipopolysaccharide (LPS; Sigma, United States) for 24 h was performed for macrophage activation. Supernatant IL-12p70 was measured by ELISA (eBioscience, United States) to support the activated-state of macrophage.

Macrophages and HP were allowed for phagocytosis for 0.5, 1, and 2 h. At each time point, all media was removed and 100 µl/well of 0.2% trypan blue in PBS was added to quench extracellular FITC labeled-bacteria. Phagocytosis activity was determined by the detection of intracellular bacteria with fluorescent intensity read at 492 nm excitation and 518 nm emission wavelengths. On the other hand, the killing assay was performed as previously published (Keep et al., 2010). In brief, live HP to macrophage at MOI 500:1, as described above, was allowed for phagocytosis for 0.5 h. Then, the media and extracellular bacteria were removed, 100 µg/ml gentamycin was added for 1 h at 37°C, 5% CO₂ to eliminate the remaining extracellular bacteria. Cells were washed with PBS 5 times, the final wash was plated in Columbia agar (Oxoid, United Kingdom) to ensure that no extracellular bacteria remained after washing. Then the remaining HP-phagocytized cells were used to determine the intracellular bacteria at 0.5 h phagocytosis, while some of the wells with HP-phagocytized cells were further incubated for 2 and 6 h after extracellular bacteria removal process before bacterial determination. For intracellular bacterial determination, 200 µl/well of 0.1% saponin was added for 15 min at 37°C, 5% CO₂ to release intracellular bacterial burden.

![FIGURE 1](image-url)
bacteria, serially diluted and plated on Columbia agar (Oxoid, United Kingdom) incubated under micro-aerophilic conditions at 37°C up to 5 days for colony enumeration. The macrophage killing activity was determined by intracellular proliferation rate at each time-point by the ratio of colony forming unit (CFU) of bacteria at the specific time-point divided by CFU at 0.5 h phagocytosis.

Since immune complex (IC) can affect phagocytosis and killing activity of macrophage, the functions with and without IC treated conditions were tested. The ICs were generated as described previously (Ding et al., 2015). Briefly, mouse IgG1 anti-ovalbumin (anti-OVA; Sigma-Aldrich) was mixed with ovalbumin (OVA; Sigma-Aldrich) followed by incubation at room temperature for 30 min to allow IC formation. Then the ICs were challenged with macrophages for 24 h before performing phagocytosis and killing assay. Macrophages without ICs activation was used as a control condition. In addition, to test if serum of symptomatic FcγRIIb−/− mice influenced macrophage functions, 10% mouse serum (in DMEM) from FcγRIIb−/− (symptomatic lupus) or wild-type (control) was incubated with macrophage for 24 h before performing phagocytosis and killing activity assay.

**Statistical Analysis**

Mean ± standard error (SE) was used for data presentation. Unpaired Student’s t-test or one-way analysis of variance (ANOVA) with Tukey’s comparison test was used for the analysis of experiments with groups 2 and 3, respectively. The repeated measures ANOVA with Bonferroni post hoc analysis was used for the analysis of data with several time-points. P-values < 0.05 were considered statistically significant. SPSS 11.5 software (SPSS Inc., Chicago, IL, United States) was used for all statistical analysis.

**RESULTS**

The Prominent Anti-*Helicobacter pylori*, With Similar Disease Severity to Wild Type, in *Helicobacter pylori* Infection of Symptomatic Lupus FcγRIIb Deficient-Mice

Lupus characteristics as determined by serum creatinine, proteinuria, and anti-dsDNA in FcγRIIb−/− mice after 24-week-old is previously described (Ondee et al., 2017; Surawut et al., 2017). Hence, HP administration in 24-week-old mice represents HP infection in patients with symptomatic lupus. Indeed, HP infection caused significant weight loss in both wild-type and FcγRIIb−/− mice (Figures 1A,B) and the wild-type demonstrated a little bit more prominent weight loss. In addition, similar disease severity between both strains were demonstrated with (i) bacterial burdens in stomach with semi-quantitative assay (urease test) (Figures 1C, 2A) and quantitative methods (real-time PCR and direct culture from stomach tissue) (Figures 1D, E), (ii) histological inflammation in stomach (Figures 1F, 2B–D) and (iii) macrophage inflammatory protein-2 (MIP-2) and keratinocyte-derived cytokine (KC), the important chemokines of HP pathogenesis (Sgouras et al., 2005; De Filippo et al., 2008), in gastric-tissue (Figures 1G, H). Despite the similar severity of HP infection between both strains, FcγRIIb−/− mice had the higher serum anti-HP IgG than wild-type (Figure 1I).

*Helicobacter pylori* Infection Enhanced Anti-dsDNA Levels and Clinical Characteristics of Lupus in 24-Week-Old FcγRIIb Deficient-mice

HP infection has been shown to enhance auto-antibodies (Yamanishi et al., 2006; Hasni et al., 2011), thus this study specifically looked at anti-dsDNA levels, a specific auto-antibody of lupus, which also helps determine disease severity. Indeed, HP induced anti-dsDNA in wild-types and accelerated anti-dsDNA level in FcγRIIb−/− (Figure 3A). Although HP induced anti-dsDNA in both mouse strains, anti-dsDNA levels were more prominent in FcγRIIb−/− group (Figure 3B), implying the autoimmune inducibility of FcγRIIb−/− mice. As most of the serum protein in gamma zone of serum protein electrophoresis analysis is immunoglobulin, serum gamma globulin is measured as a representative of total immunoglobulin. Interestingly, HP
infection enhanced immunoglobulin production in both strains but more predominantly in FcγRIIb−/− mice (Figure 3B and Supplementary Figure S1). This is in concordance with the higher level of serum anti-HP and anti-dsDNA in FcγRIIb−/− mice (Figures 1I, 3A). In addition, there was prominent immunoglobulin deposition in the glomeruli of FcγRIIb−/− mice (Figure 4) with the enhanced severity of lupus nephritis as demonstrated with serum creatinine, urine protein (urine protein creatinine index; UPCI), renal histology and serum cytokines (IL-6, MIP-2 and KC; the representatives of systemic inflammatory responses) (Figures 3C–H, 5). HP induced chemokine responses both locally (gastric tissue) (Figures 1H,I) and systemically (Figures 3H,I). With HP infection, gastric cytokines were not different between wild-type and FcγRIIb−/− mice but serum cytokines were more prominent in the FcγRIIb−/− group.

**Enhanced Immune Cells in the Spleen of FcyRIIb Deficient-Mice With Helicobacter pylori Infection**

Due to the prominent immunoglobulin production in FcγRIIb−/− mice with HP infection, lymphoid organs in these mice were examined. It is surprising that there was no difference in gastric weight and mucosal associated lymphoid tissue morphology between wild-type and FcγRIIb−/− mice with or without HP infection (data not shown). Spleen weight among these mice was different. Spleen weight without HP infection of FcγRIIb−/− mice was higher than wild-type and the infection enhanced FcγRIIb−/− spleen weight but not wild-type spleen (Figure 6A). Thus, we examined splenocyte by flow cytometry analysis (Figures 6B–G, 7). Without HP, plasma cell (CD138 +) was the only
cell population that was different between wild-type and FcyRIIB−/− mice (Figure 6B). In FcyRIIB−/− mice with HP infection, there was an increase in plasma cells, activated B cells (CD19 + and CD80 +) and follicular helper T cells (CD3 +, CD4 + and CXCR5 +) but not follicular B cells (CXCR5 + and B220 +), germinal center B cells (CD19 + and GC +) and macrophages (F4/80 +) in comparison with FcyRIIB+/− without HP or wild-type with HP (Figures 6B–G). Only follicular helper T cells were increased in wild-type with HP compared to wild-type without HP infection (Figure 6F). This data supports the hyper-immune response of FcyRIIB−/− against HP compared with wild-type.

The Hyper-Responsiveness of FcyRIIB−/− Macrophages to Helicobacter pylori Activation and the Loss of Response With Immune Complex Treatment

We examined macrophage functions, in vitro, as a pilot experiment because (i) the influence of macrophage against HP in lupus and wild-type might be different, (ii) the possible effect of high circulating immune complex against macrophage functions (Bolland and Ravetch, 2000; Clatworthy et al., 2007; Surawut et al., 2017) and (iii) the lack of data on FcyRIIB−/− macrophage against HP. Interestingly, FcyRIIB−/− macrophage showed more prominent phagocytosis than wild-type after 0.5 h of incubation, however, this was not the case after 1 or 2 h of incubation (Figures 8A,C). In addition, the IC incubation reduced phagocytosis at 0.5 h in both groups of macrophage and at 1 h in FcyRIIB−/− group (Figure 8A). Further, FcyRIIB−/− macrophage showed prominent killing activity over wild-type at both time-points (lower intracellular bacteria) (Figures 8B,D). After IC-incubation, the killing activity worsened in both strains of macrophage, but predominantly in the wild-type cells (Figure 8B). Interestingly, pre-treatment macrophage with serum from FcyRIIB+/− mice, but not wild-type serum, reduced macrophage killing activity without the influence on phagocytosis activity (Figures 8C,D). This implies that there are some macrophage-neutralizing factors in serum of symptomatic lupus mice.

DISCUSSION

Although a previous case-control study demonstrates low prevalence of HP infection in patients with lupus (Sawalha et al., 2004), there has been no other studies exploring the lupus-HP relationship. Despite the case-control study in that report, it is still unclear if the immune response of patients with lupus prone genes is effective against HP or HP infection is protective for lupus. Hence, we explored lupus-HP relationship in FcyRIIB−/− lupus mice in this study.

There was prominent antibody production but similar severity of HP gastritis in FcyRIIB−/− mice in comparison with wild-type. Enhanced immune responses and effective organism control due to the defect of FcyRIIB, the only inhibitory receptor of the FcyR family, is well-known (Bolland and Ravetch, 2000). Despite these benefits, the severity of HP infection, demonstrated by survival, bacterial burdens and gastric-cytokines, were similar between FcyRIIB−/− and wild-type. Surprisingly, anti-HP antibody was more prominent in FcyRIIB−/− mice. Despite the beneficial effects of anti-HP antibody against HP in wild-type mice (Kaparakis et al., 2008), the antibody in FcyRIIB−/− mice seemed to be less effective.
There was enhanced antibody production and lupus acceleration in FcγRIIb−/− mice with HP gastritis, a localized infection. There was not only the increase in anti-HP, a specific antibody against HP, but also anti-dsDNA and serum gamma immunoglobulin in FcγRIIb−/− mice with HP. Interestingly, it seems that the spleen, but not the gastric lymphoid tissue, was responsible for the prominent antibody production in FcγRIIb−/− with HP because of the increased weight of the spleen, not the stomach. HP infection induced several immune cells that associated with the antibody production process in the spleen, including activated B cell, plasma cell and follicular helper T-cell. In the wild-type with HP, anti-HP increased without spleen activation and serum immunoglobulin. Indeed, HP infection activated only gastric lymphoid organ and lymphocyte but not the spleen in the wild-type mice (Bussiere et al., 2006; Floch et al., 2015). The systemic response against HP (serum cytokines) was found only in FcγRIIb−/− mice but not wild-type supporting the hyper-inflammatory responses of FcγRIIb−/− mice. It is interesting to note that increased serum cytokines after HP infection is demonstrated in most of the studies on mouse model (Kodama et al., 2005) and juvenile patients (Khaiboullina et al., 2016) but not in other group of patients (Bayraktaroglu et al., 2004). Although there is no data on the influence of increased systemic cytokines in patients with HP, our study implicated the importance of increased systemic cytokines in lupus with HP infection.

It is well known that chronic infection and inflammation initiates and accelerates lupus (Munoz et al., 2010; Esposito et al., 2014; Podolska et al., 2015; Rigante and Esposito, 2015) and the hyper-inflammatory responses in FcγRIIb−/− mice due to the inhibitory signaling defect (Clatworthy et al., 2007) might enhance this effect. Thus, increased antibody production in FcγRIIb−/− mice with HP might be due to the prominent systemic responses against HP infection. Perhaps, the persistent gastric inflammation from HP infection induces epitope spreading and/or bystander activation in...
FIGURE 7 | The representatives of flow-cytometric analysis were demonstrated; spleen plasma cell (A) and spleen follicular helper T cell (B).
FcyRIIb−/− mice resulting in the increased auto-antibody. As such, HP enhanced anti-dsDNA levels in both strains, but more predominantly in FcyRIIb−/−, supports the well-known HP-induced autoimmunity hypothesis (Hasni et al., 2011). In addition, the increased anti-dsDNA in FcyRIIb−/− mice also enhanced lupus severity, supporting its role in lupus pathogenesis (Giles and Boackle, 2013).

Regarding the increased anti-HP antibody (Figure 11) and the prominent macrophage functions (Bolland and Ravetch, 2000; Clatworthy et al., 2007; Surawut et al., 2017) in FcyRIIb−/− mice, the lower bacterial burdens and the more severe gastritis were expected in these mice compared with wild-type. This is due to the attenuation property of anti-HP and the macrophage enhanced gastritis in macrophage depleted wild-type mice (Kaparakis et al., 2008). Surprisingly, bacterial burdens and gastritis severity were not different between the 2 groups. This suggests different influence of immune responses or different neutralizing factor of antibody between wild-type and lupus mice. Moreover, infection susceptibility and high circulating immune complex (CIC) in symptomatic lupus is well-known, thus the negative influence of CIC in lupus against infection is possible. Indeed, we demonstrated the inhibitory effect of IC and lupus mouse serum against macrophage phagocytosis and killing activity, in vitro, in both wild-type and FcyRIIb−/− macrophage which, at least in part, explained the infection susceptibility of lupus. Despite several macrophage-neutralizing factors (e.g., uremic toxins) in FcyRIIb−/− mouse serum, CIC might contribute some influences. More studies on this subject are necessary.

Several limitations of translation research should be considered before applying any clinical translation such as; (i) mouse is not the natural host of HP and repeated gavage of HP is different from the disease’s natural course, (ii) there are some different properties and expressions of FcyRIIb receptor between human and mouse (Bruhns, 2012; Hussain et al., 2015), (iii) unmeasured factors might also affect the model due to other FcyRIIb−/− characteristics, and (iv) the opsonin used for macrophage activity assays, in vitro, might not resemble the in vivo physiology.

CONCLUSION

Prominence of anti-HP, anti-dsDNA and increased serum immunoglobulin despite the similar disease severity of HP
infection in FcγRIIb−/− mice compared with wild-type were demonstrated. HP infection in FcγRIIb−/− mice enhanced systemic inflammation, induced antibody-producing immune cells in the spleen and enhanced lupus disease severity. Thus, the localized HP gastritis may induce the systemic inflammatory responses and enhance lupus progression in some patients with lupus. Thus patients with dyspepsia or increased systemic cytokine of unknown causes should be further investigated for HP-induced chronic gastritis. Clinical studies to confirm these findings in humans are necessary, which may change our current approach to clinical management of lupus.

**AUTHOR CONTRIBUTIONS**

SS and WP designed and coordinated all the experiments, performed in vitro and in vivo experiments, and wrote the manuscript and approved. JM, PT, and AT-U performed in vitro experiments, and approved the manuscript. JW performed histopathology and approved the manuscript. ST designed experiment and approved the manuscript. AL designed and coordinated all the experiments, analyzed all of these experiment and wrote the manuscript and approved.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2018.01488/full#supplementary-material

**FIGURE S1** The representative patterns of capillary protein electrophoresis from wild-type (FcγRIIb +/+ ) and FcγRIIb−/− mice with phosphate buffer solution (PBS) gavage or H. pylori administration were demonstrated (A–D).

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