Identification of pyrC gene as an immunosuppressive factor in Francisella novicida infection

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Francisella tularensis, a bacterial caustive agent of the zoonosis tularemia, is highly pathogenic to humans. The pathogenicity of this bacterium is characterized by intracellular growth in immune cells, like macrophages, and host immune suppression. However, the detailed mechanism of immune suppression by F. tularensis is still unclear. To identify the key factors causing Francisella-mediated immunosuppression, large-scale screening using a transposon random mutant library containing 3552 mutant strains of F. tularensis subsp. novicida (F. novicida) was performed. Thirteen mutants that caused stronger tumor necrosis factor (TNF-α) production in infected U937 human macrophage cells than the wild-type F. novicida strain were isolated. Sequencing analysis of transposon insertion sites revealed 10 genes, including six novel genes, as immunosuppressive factors of Francisella. Among these, the relationship of the pyrC gene, which encodes dihydroorotase in the pyrimidine biosynthesis pathway, with Francisella-mediated immunosuppression was investigated. The pyrC deletion mutant strain (ΔpyrC) induced higher TNF-α production in U937 human macrophage cells than the wild-type F. novicida strain. The ΔpyrC mutant strain was also found to enhance host interleukin-1β and interferon (IFN)-β production. The heat-inactivated ΔpyrC mutant strain could not induce host TNF-α production. Moreover, the production of IFN-β resulting from ΔpyrC infection in U937 cells was repressed upon treatment with the stimulator of interferon genes (STING)-specific inhibitor, H-151. These results suggest that pyrC is related to the immunosuppressive activity and pathogenicity of Francisella via the STING pathway.

KEYWORDS
Francisella, tularemia, pyrimidine, immune response, cytokine

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Introduction

Tularemia is a vector-borne zoonosis with severe symptoms, including fever, lymphadenitis, cutaneous lesions, and primary pulmonary involvement in humans (Petersen and Schriever, 2005). Francisella tularensis, the causative agent of tularemia, is a gram-negative, facultative, and intracellular bacterial pathogen. Tularemia is highly contagious, with subcutaneous infection with as few as 10 bacterial cells and aerosol-mediated inhalation of as few as 25 bacterial cells being able to cause human infection (Sjöstedt, 2007). Accordingly, highly virulent F. tularensis strains have been classified as a Category-A agents of potential bioterrorism by the Centers for Disease Control and Prevention and are required to be handled and contained in BSL-3 laboratories (Shapiro and Schwartz, 2002). F. tularensis is divided into four subspecies, subs. tularensis, holarctica, mediasiatica, and novicida. Among them, only Francisella tularensis subs. tularensis (F. tularensis) and Francisellatularensis subs. holarctica (F. holarctica) cause tularemia in human (Maurin, 2015). F. novicida infection in humans is considerably rare and can be handled in a BSL-2 laboratory. Most cases of human F. novicida infection have involved patients who were immunocompromised or had an underlying disease. However, F. novicida has over 98% identity to F. tularensis at the DNA level and shows many characteristics similar to F. tularensis with regards to its life cycle within macrophages and pathogenicity in mice (Rohmer et al., 2007; Kingry and Petersen, 2014). Therefore, F. novicida has been used as the model bacterium for research on Francisella pathogenicity.

While Francisella research has a long history, the detailed molecular mechanisms of infection by Francisella subspecies remain unknown. Francisella can replicate in immunocompetent cells, such as macrophages, neutrophils, and dendritic cells, which are essentially responsible for the elimination of pathogens from the body (Santic et al., 2006). To survive inside host cells, Francisella species employ various strategies. Francisella enters these cells via phagocytosis, escapes digestion by phagolysosomes and autophagosomes, and finally replicates in the cytoplasm (Cheng et al., 2008; Miller and Celli, 2016). Further, Francisella suppresses or evades host pattern recognition receptors (PRRs), which usually initiate the innate immune response to exclude pathogens (Jones et al., 2011; Gillette et al., 2014; Putzova et al., 2017). Although some of the factors responsible for Francisella’s immunosuppressive abilities were identified in previous research (Platz et al., 2010; Nallaparaju et al., 2011; Mahawar et al., 2012), the detailed mechanisms involved in Francisella-mediated immunosuppression remain to be elucidated.

Francisella grows in nutrient-limited host cells, and this nutrient limitation is closely related to its pathogenicity (Best and Abu Kwaik, 2019). To date, a lot of genes responsible for the uptake or biosynthesis of nutrients, such as amino acids, carbon, vitamins, and bases, were reported as crucial factors for the intracellular replication of Francisella (Meibom and Charbit, 2010; Santic and Abu Kwaik, 2013; Feng et al., 2014). Especially, nucleotide biosynthesis is essential for the survival and virulence of bacterial pathogens, including intracellular bacteria, such as Salmonella, Listeria, Brucella, and Francisella (Goncheva et al., 2022). Since pyrimidine nucleotides are essential for all organisms, almost all bacterial species have a de novo pyrimidine biosynthesis pathway, which has highly conserved enzymatic steps. The de novo pyrimidine biosynthesis pathway for synthesizing uridine 5’-monophosphate consists of six steps and employs enzymes encoded by carA/B and pyrB-F (Turnbough Charles and Switzer Robert, 2008). The pyrC gene encodes a putative dihydroorotase that converts carbamoyl N-Carbamoyl-L-aspartate into 4,5-dihydroorotate in the Francisella pyrimidine biosynthesis pathway. Although some reports suggest pyrimidine biosynthesis is important for the intracellular growth of Francisella, it is not well understood how pyrimidine biosynthesis is involved in immunosuppression by Francisella (Qin and Mann, 2006; Horzempa et al., 2010).

In this study, we performed large-scale screening of a F. novicida transposon mutant library to search for the key factors involved in the immunosuppression mechanisms of Francisella. We identified pyrC as a novel F. novicida factor suppressing host innate immune responses and evaluated the immunological characteristics of host cells infected with F. novicida.

Materials and methods

Bacterial strains and culture conditions

F. novicida U112 was obtained from the Pathogenic Microorganism Genetic Resource Stock Center (Gifu University, Gifu, Japan). F. novicida was cultured aerobically at 37°C in a chemically defined medium (CDM) (Nagle et al., 1960) or brain heart infusion broth (Becton, Dickinson and Company, NJ, USA) supplemented with 0.1% cysteine (BHIc) (Mc Gann et al., 2010) or BHIc containing 1.5% agar (Wako Laboratory Chemicals, Osaka, Japan). All experiments were conducted in compliance with the institutional biosecurity guidelines and were approved by Yamaguchi University.

Cell culture

Human monocytic U937 cells were grown in Roswell Park Memorial Institute (RPMI) 1640 medium (Sigma-Aldrich, St Louis, MO, USA) supplemented with 10% heat-inactivated fetal bovine serum at 37°C in an atmosphere containing 5% CO₂.
Plasmid construction, transformation, and transfection

Supplementary Table 1 lists the primer sets and templates used to construct the plasmids used in this study. Polymerase chain reaction (PCR) was performed using KOD-Plus-Neo polymerase (Toyobo, Osaka, Japan), and ligation was performed using an In-Fusion HD Cloning Kit (Takara Bio, Shiga, Japan). Plasmids were used to transform *F. novicida* via electroporation. Specifically, bacterial cells were suspended in 0.5 M sucrose with 2 μg of plasmid DNA and were electroporated using a Bio-Rad micropulser (Bio-Rad, Hercules, CA, USA) at 3.0 kV, 10 µF, and 600 Ω with a 0.2 cm cuvette. The transformants were pre-incubated in BHlc medium overnight. To select transformed bacteria, the pre-incubated bacteria were cultured on BHlc agar plates containing 30 μg/ml kanamycin or 2.5 μg/ml chloramphenicol.

Construction of a transposon mutant library

The transposon mutant library was constructed using the Ez-Tn5 transposon system (Epicentre Lucigen, Madison, WI, USA), as previously reported (Nakamura et al., 2019). Briefly, the multiple cloning site of pMOD3 was linearized by digestion with Hind III and EcoRI, and the kanamycin resistance cassette of pKEK1140 (Rodriguez et al., 2008) was ligated into the Hind III and EcoRI sites to generate pMOD3-FtKm. The transposon moiety of pMOD3-FtKm was amplified by PCR, purified, mixed with transposase according to the manufacturer’s instructions, and then used to transform *F. novicida* via electroporation. Transformed bacteria were cultured on BHlc plates containing 30 μg/ml kanamycin.

Sequence analysis of transposon mutants

pMOD3 harbors the *E. coli* R6Kγ origin of replication. The genomes of *F. novicida* transposon mutants were purified using a PureLink Genomic DNA Mini Kit (Thermo Fisher Scientific, MA, USA) and digested with a combination of restriction enzymes, including XhoI, BglII, EcoRI, Sall, NotI, and BamHI. The ends of the digested DNAs were then blunted using a DNA Blunting Kit (Takara Bio) and ligated using Ligation High Ver. 2 (Toyobo). The ligated DNA was used to transform λpir Chemically Competent *E. coli* (Thermo Fisher Scientific). The transformed *E. coli* were selected for kanamycin resistance, and plasmid DNAs were purified. Sequence analysis was performed using the primer described in the manufacturer’s instructions for the Ez-Tn5 transposon system.

Construction of *F. novicida* mutants

The dotU homolog (*FTN_1316*) deletion mutant (ΔdotU) was previously constructed (Shimizu et al., 2019) through group II intron insertion using a TargeTron Gene Knockout System (Sigma-Aldrich), which was modified for Francisella species (Rodriguez et al., 2008). The pyrC gene (*FTN_0024*) deletion mutant (ΔpyrC) was generated via homologous recombination using the Francisella suicide vector pFRSU (Shimizu et al., 2019). The upstream and downstream regions of pyrC (1.5 kb each) were cloned into the BamHI site of pFRSU to generate pFRSU-pyrC. The pFRSU-pyrC vector (2 μg) was used to transform *F. novicida*; transformants were selected on BHlc plates containing 30 μg/ml kanamycin. Isolated bacteria were cultured in BHlc without antibiotics overnight and then plated on BHlc plates containing 5% sucrose. The deletion of the pyrC gene was confirmed via PCR.

Green fluorescent protein- and PyrC-expressing *F. novicida* strains

A GFP-expressing plasmid, pOM5-GFP, was constructed according to published procedures (Shimizu et al., 2019). The *F. novicida* chromosomal pyrC gene was cloned into pOM5 to generate pOM5-pyrC. To construct GFP-expressing strains and pyrC complemented strains, pOM5-pyrC and pOM5-GFP were used to transform the wild-type strain or the ΔpyrC mutant strain of *F. novicida* via electroporation.

Intracellular growth assay

U937 cells (1 × 10^5 cells/well) were incubated in a 48-well tissue culture plate with 100 nM phorbol myristate acetate (PMA) for 48 h. Then, *F. novicida* strains were added at a multiplicity of infection (MOI) of 1. Next, the plates were centrifuged for 10 min at 300 × g and incubated for 1 h at 37°C. The cells were then washed three times with RPMI 1640 medium, and extracellular bacteria were killed with gentamicin at 50 μg/ml for 1 h. The cells were then incubated in fresh medium at 37°C for the indicated time durations in figure legends. To measure intracellular growth, the cells were washed with PBS and then lysed with 0.1% Triton X-100 in CDM. The CFUs were determined on BHlc agar plates by plating serial dilutions of cultures.

Fluorescence microscopy

U937 cells (1 × 10^5 cells/well) were incubated with 100 nM PMA for 48 h on 12 mm glass coverslips in 24-well tissue culture
plates. GFP-expressing *F. novicida* strains were infected at an MOI of 1. Plates were then centrifuged for 10 min at 300 × g and incubated for 1 h at 37°C. The cells were washed three times with RPMI 1640 medium, and extracellular bacteria were eliminated using gentamicin at 50 μg/ml for 1 h. The cells were then incubated in fresh medium at 37°C for the indicated time durations in figure legends. Cells were fixed with 4% paraformaldehyde at room temperature for 30 min. A FluoView FV1000 confocal laser scanning microscope (Olympus, Tokyo, Japan) was used to obtain images of the cells.

**RNA isolation and qPCR analysis**

U937 cells (4 × 10⁵ cells/well) were incubated in a 12-well tissue culture plate with 100 nM PMA for 48 h. The medium was exchanged with fresh pre-incubated RPMI 1640 medium one-hour prior to infection. Cells were infected with *F. novicida* strains at MOI = 1 or stimulated with 100 ng/ml of lipopolysaccharide (LPS) derived from *E. coli* (O127:B8) (Sigma-Aldrich), or 10 ng/ml 2′-3′-cGAMP (InvivoGen, CA, USA). The plates were centrifuged for 10 min at 300 × g and incubated for indicated time the indicated time durations in figure legends. Cells were carefully washed twice with PBS, and total RNA was collected using NucleoSpin RNA kit (Takara Bio). RNA was quantified by determining absorption at 260 nm using NanoDrop 2000 (Thermo Fisher Scientific). Next, qPCR was performed using the RNA-direct Realtime PCR Master Mix (Toyobo) with an RNA concentration of 50 ng per 20 μl reaction. The HPR71 ampiclon was used as an endogenous control to normalize all mRNA expression data. The relative expression levels of genes in various conditions compared with those in the BHIC medium-treated control were calculated using the relative quantification method (ΔΔCt method) (Pfaffl, 2001). Used primer sets are shown in Table S1.

**ELISA**

U937 cells (1 × 10⁵ cells/well) were pre-incubated in a 48-well tissue culture plate with 100 nM PMA for 48 h. After exchanging the medium with pre-incubated fresh RPMI medium 1 h prior to infection, cells were infected with bacterial strains (*F. novicida* transposon mutant strains or deletion mutant strains) at an MOI of 1 or stimulated with 100 ng/ml of LPS derived from *E. coli* (O127:B8) or 10 ng/ml of 2′-3′-cGAMP (InvivoGen). Heat-inactivation of each strain was performed by incubating the bacterial suspension in a heat block at 90°C for 5 min. In the case of STING inhibition, a STING-specific inhibitor, H-151 was used (Haag et al., 2018). U937 cells were treated with H-151 (final concentration 0.5 μM) or the same volume of DMSO 2 h prior to infection. After incubation for the indicated time durations in figure legends, concentrations of tumor necrosis factor (TNF)-α and interleukin (IL)-1β in the supernatants were measured using ELISA MAX Standard Kit (Biolegend, CA, USA) according to the manufacturer’s instructions. Interferon (IFN)-β in the supernatants were measured using VeriKine Human Interferon Beta ELISA Kit (PBL Assay Science, Piscataway, NJ, USA).

**Statistical analysis**

Student’s *t* test or multiple comparisons using the Tukey-Kramer test and Dunnett’s test were used to evaluate the significance of differences compared with the wild-type strain; *P* < 0.05 indicates a significant difference.

**Results**

**Ten genes were identified as immunosuppressive factors of *F. novicida***

To identify novel immunosuppressive factors of *F. novicida* in human macrophages, we expanded a previously constructed *F. novicida* transposon mutant library consist of 750 strains (Nakamura et al., 2019) up to 3552 strains. TNF-α is a cytokine produced through a broad range of innate immune signaling pathways, including Toll-like Receptor (TLR) 4- and TLR2-mediated pathways, and is reported to be suppressed by Francisella infection (Telepnev et al., 2003; Butchar et al., 2008). To identify genes responsible for immunosuppression by *F. novicida*, U937 cells were infected with a mutant library, and the transposon mutants inducing excessive TNF-α production compared with the wild-type strain were selected through ELISA using the culture supernatant. In the 1st screening, the cut-off was set to a 1.5-fold increase in TNF-α production. In the 2nd screening, U937 cells were infected with mutant strains selected via the first screening, and their TNF-α production was measured three times. Finally, 13 mutants that increased TNF-α production in U937 cells were identified (Figure 1). The Δft mutant strain, lacking the gene encoding soluble lytic transglycosylase, was used as a positive control (Nakamura et al., 2019). To determine the genes responsible for Francisella-mediated immunosuppression, the transposon insertion sites of the selected mutant strains were evaluated by sequence analysis; 10 unique genes were identified (Table 1). In this study, we focused on *pyrC* (*FTN_0024*), the gene putatively encoding dihydroorotase, and the effect of this gene on immunosuppression by Francisella.

**The *pyrC* deletion mutant evokes the innate immune responses of host U937 macrophage cells**

To estimate the effect of *pyrC* in immunosuppression by *F. novicida*, we constructed a Δ*pyrC* mutant strain of *F. novicida*. 

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via homologous recombination. ELISA showed that the ΔpyrC mutant strain induced significantly higher levels of TNF-α production in the cell culture supernatant of U937 cells than the wild-type strain (Figure 2A). Although not significantly different, the pyrC complemented strain tended to show decreased levels of TNF-α induction compared with the ΔpyrC mutant strain. The pyrC complemented strain showed equivalent levels of TNF-α induction to the wild-type strain (Figure 2A). Next, we measured the induction of IL-1β and IFN-β, which are important cytokines for Francisella infection, in the ΔpyrC mutant strain (Gavrilin and Wewers, 2011). As in the case of TNF-α, the ΔpyrC mutant strain-infected U937 cells showed significantly higher levels of IL-1β and IFN-β production than the wild-type strain, and pyrC complementation decreased these levels to that observed upon infection with the wild-type strain (Figures 2B, C). The mRNA expression levels of TNF, IL1B, and IFNB1 in Francisella-infected U937 cells were also examined using real-time PCR. Similar results to those obtained via ELISA were observed for the mRNAs of all these genes (Figures 2D–F). These results indicate that the ΔpyrC mutant strain strongly evokes host innate immune responses compared to the wild-

### TABLE 1 The results of screening and sequence analysis of transposon mutants.

| Strain | TNF-α (pg/ml) ± S.D. | Locus | Name | Putative protein |
|--------|----------------------|-------|------|------------------|
| Wild-type (WT) | 432.39 ± 131.00 | – | – | – |
| A11-6 | 2912.36 ± 1332.87 | FTN_0756 | fopA | OmpA family protein |
| B22-5 | 2489.22 ± 1021.12 | FTN_1641 | ampG | Major Facilitator Superfamily protein |
| C15-5 | 2635.81 ± 1576.51 | FTN_1286 | mltA | MltA specific insert domain protein |
| E12-3 | 1153.80 ± 797.89 | FTN_1199 | capA/B | Poly-gamma-glutamate system family protein |
| H31-3 | 946.36 ± 303.20 | FTN_057 | cas9 | CRISPR-associated endonuclease Cas9 |
| J11-2 | 1138.33 ± 445.28 | FTN_1548 | yfgL | Outer membrane assembly lipoprotein YfgL |
| J7-1 | 2068.55 ± 942.58 | FTN_0917 | dacB | D-alanyl-D-alanine carboxypeptidase/D-alanyl-D-alanine-endopeptidase |
| J21-3 | 2925.88 ± 1726.95 | FTN_0496 | slt | Transglycosylase SLT domain protein |
| L8-8 | 3024.12 ± 1718.24 | FTN_0496 | slt | Transglycosylase SLT domain protein |
| L32-4 | 2094.21 ± 1281.19 | FTN_0611 | kdhA | 3-deoxy-D-manno-octulosonic acid 8-phosphate synthase |
| M14-8 | 970.04 ± 320.70 | FTN_0757 | cas9 | CRISPR-associated endonuclease Cas9 |
| M20-7 | 1290.48 ± 310.36 | FTN_0024 | pyrC | Dihydroorotase, multifunctional complex type domain protein |
| O7-8 | 2285.06 ± 352.58 | FTN_0496 | slt | Transglycosylase SLT domain protein |
| Δslt | 1767.43 ± 351.55 | FTN_0496 | slt | Transglycosylase SLT domain protein |
| LPS (O127:B8) | 1491.96 ± 455.53 | – | – | – |
type strain, suggesting that pyrC is a critical factor for the immunosuppression of F. novicida.

**The pyrC gene is important for the intracellular growth of F. novicida in U937 cells**

Next, we examined whether ΔpyrC is involved in the intracellular growth of F. novicida. The ΔpyrC mutant strain entered the stationary phase slightly earlier than wild-type strain in BHIc medium and failed to grow in CDM (Supplementary Figure 1). The GFP-expressing wild-type strain of F. novicida grew intracellularly in U937 cells from 2 to 48 h post infection, while the GFP-expressing ΔpyrC mutant strain did not show remarkable intracellular growth during the same period through fluorescence microscopy (Figure 3A). To support this finding, the number of ΔpyrC and transposon mutant strain cells were significantly decreased to approximately 1/10th that of wild-type strain cells within U937 cells. On the contrary, the ΔpyrC mutant...
strain showed higher intracellular growth compared to the strain deficient in the type VI secretion system (ΔdotU), which was used as a negative control. The complemented strain restored the ability of the ΔpyrC mutant strain to grow intracellularly (Figure 3B). These results suggest that pyrC is not essential but is important for the intracellular growth of F. novicida in host cells.

**TNF-α induction by the ΔpyrC mutant strain is abolished by heat treatment**

Host cells infected with bacteria produce TNF-α due to the recognition of various bacterial ligands, such as LPS, peptidoglycan, and nucleotides, by PRRs (Kawai and Akira, 2006; Li and Wu, 2021). To identify ΔpyrC mutant strain
ligands responsible for the induction of TNF-α in the ΔpyrC mutant strain infected U937 cells. U937 cells were treated with heat-inactivated F. novicida mutant strains. TNF-α induction by all heat-inactivated F. novicida strains, including the ΔpyrC mutant strain, was decreased to the same level as that by the negative control; however, LPS retained the ability to induce TNF-α with or without heat inactivation (Figure 4). This result suggests that TNF-α production by F. novicida strains, including the ΔpyrC mutant strain, is induced through biological activities of F. novicida, such as internalization by or proliferation in host cells, and not through heat-stable ligands, such as LPS, peptidoglycan, and nucleotides.

**IFN-β induction in U937 cells infected with the ΔpyrC mutant strain is mediated by the STING pathway**

Because IFN-β production is induced by recognition of cytosolic DNA by the cyclic-di-nucleotide sensor STING (Jones et al., 2010; Storek et al., 2015) pathway, we next examined whether F. novicida pyrC affects the STING pathway. U937 cells treated with a STING inhibitor H-151 were infected with the ΔpyrC mutant strain, and the IFN-β levels induced in them were measured using ELISA. H-151 showed no significant effect on the growth of both the wild-type and the ΔpyrC mutant strain in BHIc medium (Supplementary Figure 2). IFN-β levels in the supernatant of H-151-treated U937 cells infected with the ΔpyrC mutant or transposon mutants and stimulated with the STING agonist 2’3’-cGAMP were significantly decreased compared to those in the supernatant of the DMSO control cells (Figure 5). Contrarily, H-151-treated U937 cells showed no significant difference in IFN-β induction upon infection with ΔpyrC mutant or transposon mutants compared to the wild-type and pyrC complemented strains. These results indicate that pyrC is involved in the suppression of IFN-β through the STING pathway.

**Discussion**

Intracellular bacteria, including Francisella, have refined their strategy to escape the host immune system and survive in host cells. To date, the importance of immunosuppression and immune evasion in Francisella infection has been well recognized, but their detailed mechanisms are poorly understood. To our knowledge, a large-scale gene screening of Francisella mutants focusing on Francisella immunosuppressive properties has not been performed yet. Therefore, here, we developed a transposon mutant library of F. novicida consisting of 3552 mutants. This library seems to cover the 1731 protein coding genes of the F. novicida U112 strain (Rohmer et al., 2007). Among the 3552 mutant strains in our library, strains that induced higher levels of host immune response than the wild-type F. novicida strain were isolated. Ten genes were determined to be immunosuppression-related. Among these 10 genes, four genes, FTN_0756 (fopA), FTN_1286 (mltA), FTN_0757 (cas9), and FTN_0496 (slt), were previously reported as immunosuppressive factors of Francisella (Peng et al., 2011; Sampson et al., 2013; Nakamura et al., 2019; Nakamura et al., 2021), ensuring the reliability of the screening method employed in this study. The remaining 6 genes, FTN_1641 (ampG),
F. novicida. TNF-α is produced through a broad range of innate immune signaling pathways, including TLR signaling pathways activated by various bacterial ligands, such as LPS, peptidoglycan, and nucleotides (Kawai and Akira, 2006; Li and Wu, 2021). In host cells infected by Francisella, the production of inflammatory cytokines, such as TNF-α and IL-6, is induced by the recognition of Francisella DNA by TLR9 (Jones et al., 2012). Because pyrC is related to pyrimidine biosynthesis, it may suppress immune responses by modifying nucleotides recognized by TLRs, such as TLR9. However, as discussed below, heat-inactivated F. novicida strains, including the ΔpyrC mutant strain, failed to induce TNF-α production, indicating that biological activities of F. novicida, such as its phagosomal escape, are necessary for it to induce TNF-α and other cytokines.

Our data also revealed that the disruption of the pyrimidine biosynthesis pathway by F. novicida induces higher levels of IL-1β and IFN-β production in U937 macrophage cells. IL-1β is secreted when its precursor is expressed through the activation of TLR or type I IFN signaling followed by cleavage with caspase-1 activated through recognition by inflammasomes (Henry et al., 2007; Man and Kanneganti, 2015). In F. novicida infections, infected host cells exhibit robust immune responses while infected host cells infected with wild-type F. novicida and increased in host cells infected with the ΔpyrC mutant strain. This result is inconsistent with those of previous reports by Horzempa’s and Schulte, which indicate that several genes of F. tularensis Schu S4 and F. holarctica LVS strains are related to the suppression of IL-1β production (Schulte et al., 2015).
Previous report on *Pseudomonas aeruginosa* indicated that uracil controls biofilm formation via quorum-sensing, and *P. aeruginosa* mutants lacking genes involved in uracil biosynthesis could not form biofilms (Ueda et al., 2009). A biofilm is a structured community of microbial cells in a matrix formed by extracellular polymeric substances (EPS). The EPS consist of polysaccharides, nucleic acids (extracellular DNA and RNA), proteins, lipids, and other biomolecules (Karygianni et al., 2020). It has also been shown that biofilms formed by *Mycobacterium avium* and *P. aeruginosa* and the extracellular DNA of these bacteria have the potential to induce TNF-α production in host cells infected by them (Rose and Bermudez, 2014; Ramirez et al., 2019). In the ΔpyrC mutant strain-infected U937 cells, the production of the TNF-α, IL-1β, and IFN-β cytokines was increased in response to the recognition of bacterial nucleic acids compared with that in wild-type strain-infected U937 cells. These results suggest that pyrC may be involved in the coordination or modification of ligands, such as extracellular DNA in host cytosol, protecting *Francisella* from TLR or STING recognition and allowing it to grow intracellularly by suppressing immune responses.

Intracellular growth is one of the most important abilities determining *Francisella*’s pathogenicity, a lot of genes involved in the intracellular growth of *Francisella* have been identified in previous studies (Qin and Mann, 2006; Su et al., 2007). PyrC is required for *de novo* pyrimidine biosynthesis (Choi and Zalkin, 1990). Mutants of *Francisella* genes involved in the pyrimidine pathway (e.g., *carA*, *pyrB*, *pyrD*, and *pyrF*) become uracil auxotrophs and show deficient growth on complete medium (Maier et al., 2006; Qin and Mann, 2006; Horzempa et al., 2010). Several reports indicate that these pyrimidine pathway-related mutants can grow within epithelial cells but not macrophage cells (Qin and Mann, 2006; Horzempa et al., 2010). In addition, Schurert et al. showed that a pyrimidine biosynthesis pathway transposon mutant of *F. novicida* was eliminated by monocyt-derived macrophages, in part via phagosomes (Schurert et al., 2009). In our study, the ΔpyrC mutant showed decreased growth in culture medium, and decreased but constant intracellular growth in U937 cells compared with that of the wild-type strain. *Francisella* mutants that is deficient in the phagosomal escape-related factor such as mglA or type VI secretion system, have no ability to induce IL-1β secretion (Mariathasan et al., 2005; Gavrili et al., 2006; Jones et al., 2010). These results indicate that the ΔpyrC mutant strain can enter the host cytoplasm and grow intracellularly. Although the intracellular bacterial number of the ΔpyrC mutant strain was relatively low, it increased the production of cytokines compared with that in cells infected with the wild-type strain, suggesting strongly that pyrC contributes to the suppression of host immune responses.

In summary, we performed here, a large-scale screening to search for factors responsible for immunosuppression by *F. novicida* in human macrophage cells. Ten genes were determined...
to be responsible for the immunosuppression. Among them, prpC was identified as a novel F. novicida immunosuppressive factor and was immunologically characterized. Although further studies are needed to elucidate the detailed mechanisms by which prpC is involved in host immunosuppression by Francisella, research on pyrimidine metabolic pathways involving prpC may provide new insight into Francisella immunosuppression and pathogenicity and into the mechanisms by which host cells recognize intracellular bacteria.

Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

Author contributions

TN contributed to laboratory analysis, investigation, and writing of the original draft. TS contributed to study conceptualization, laboratory analysis, investigation, and writing of the original draft. RI contributed to laboratory analysis. AU contributed to methodology and resources. KW contributed to laboratory analysis, investigation, and validation. MW was involved in study conceptualization, laboratory analysis, supervision, manuscript review, and manuscript editing. All authors contributed to the article and approved the submitted version.

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Supplementary material

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