RESEARCH ARTICLE

N-Terminal Polypeptide of Annexin A2 Decreases Infection of Mycoplasma hyorhinis to Gastric Cancer Cells

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Abstract

Mycoplasma infection in human and its contamination in cell cultures are worldwide problems. The drugs currently available for preventing or treating mycoplasma infection suffer from low sensitivity, strong resistance and high toxicity. Our previous work showed that Mycoplasma hyorhinis (M. hyorhinis) infection was mediated by the interaction between p37 of M. hyorhinis and Annexin A2 (ANXA2) of host cells, however the translational value of this mechanism was unknown. Herein, we synthesized the N-terminal of ANXA2 polypeptide (A2PP) and found that A2PP could decrease the infection of M. hyorhinis to gastric cancer cells and block M. hyorhinis infection-induced cell migration. Furthermore, we found that A2PP could reduce M. hyorhinis contamination of passage cells. Moreover, compared with the commercial antibiotics commonly used in cell culture to prevent M. hyorhinis infection, A2PP demonstrated a more effectiveness but a low toxicity on cell growth. Thus, our study for the first time revealed A2PP’s potential for the treatment and prevention of M. hyorhinis infection.

Introduction

Pathogenic mycoplasmas, including Mycoplasma pneumoniae (M. pneumoniae), mycoplasma hyorhinis (M. hyorhinis), oral mycoplasma (M. orale) and Mycoplasma genitalium (M. genitalium), belong to class mollicutes, which is the smallest microorganism living in nature and can duplicate independently [1–2]. Many studies showed that infection with these mycoplasmas was associated with tumor development [3–8]. M. orale causes chromosome abnormality in human diploid cells and induces cell transformation [6]. Infection with M. hyorhinis or M. genitalium could increase the migration and invasiveness of prostate epithelial cells [3]. M. hyorhinis infection has been linked to arthritis, serositis, infertility and cancer of human [9–12].

Additionally, mycoplasma contamination in cell culture is a serious problem and the rate of passage cells infected by mycoplasma is high. Cell culture is widely used in life sciences, such as in the basic research, clinical trial research, development and production of biological products, as well as in the field of biopharmaceutical and vaccine production. Preventing the cells from
microbial contamination is critical to ensure the quality of research. The mycoplasma contamination is the most common problem in cell culture with an incidence of 30%-60% [1]. It was shown that four species of mycoplasmas account for more than 95% of infection in cell culture, including M. orale, arginine mycoplasma (M. arginini), M. hyorhinis, and Levin’s no Acholeplasma (A. laidlawii) [13,14,15,16]. Because of small size, mycoplasma is hardly to be found under the light microscope and is easily ignored by researchers. In recent years, numerous antibiotics used for mycoplasma treatment repeatedly resulted in the emergence of mycoplasma resistance. For these reasons, it is imperative to develop new agents for preventing or decreasing mycoplasma infection.

Our previous work showed that M. hyorhinis infection depends on the interaction of p37 (major membrane protein of M. hyorhinis) and host ANXA2 through their N-terminal domains. We also found polyclonal antibody to p37 could block the infection of M. hyorhinis and decrease M. hyorhinis-promoted migration of gastric cancer cell [17]. Based on these discoveries, we raised a hypothesis that the peptide, synthesized according to the amino acid sequence of ANXA2 N-terminal region (from 1st to 30th aa, here we named this peptide as A2PP), could block M. hyorhinis infection via its competition with ANXA2 and possibly be used as a drug for preventing M. hyorhinis infection in cell culture. In this study, we tested this hypothesis and also compared the effects of A2PP with other drugs in preventing M. hyorhinis infection.

Materials and Methods

Cell Culture

AGS gastric cancer cell line was from American Type Culture Collection (ATCC). BGC823 gastric cancer cell line was established by the Peking University People’s Hospital and was purchased from Cell Culture Center of Chinese Academy of Medical Sciences (Beijing, China). AGS and BGC823 cells were cultured in RPMI-1640 medium supplemented 10% fetal calf serum obtained from Invitrogen (Carlsbad, CA, USA). Mycoplasma test was implemented before each new experiment by PCR amplification of M. hyorhinis p37.

Mycoplasma Propagation and Co-Culture

Mycoplasma propagation and co-culture was carried out as previously reported [17,18,19].

Antibodies and Reagents

Anti-p37 monoclonal antibody PD4 was generated and characterized previously [5,20,21]. Polyclonal anti-p37 antibody was obtained from immunizing rabbit with GST-p37 fusion protein following the standard protocol. Anti-EGFR and Anti-phospho-EGFR was purchased from Cell Signaling Technology (CST, USA). Anti-ANXA2 was from Novus Biotechnology (Novus Biotechnology, USA). Anti-phospho-ANXA2 was from Santa Cruz (Santa Cruz, CA, USA). A2PP (1st to 30th aa) and various truncated A2PP peptides, including A2PP-Nd4 (5th to 30th aa), A2PP-Nd8 (9th to 30th aa), A2PP-Nd12 (13th to 30th aa), A2PP-Nd16 (17th to 30th aa), A2PP-Cd4 (1st to 26th aa), A2PP-Cd8 (1st to 22nd aa), A2PP-Cd12 (1st to 18th aa), A2PP-Cd16 (1st to 14th aa), together with random control peptides (ConP) were synthesized by Sbsbio (Beijing, China). Antimicrobials mycoplasma I (MYCO I) and mycoplasma II (MYCO II) were purchased from M&C GENE TECHNOLOGY (Beijing, China). Ciprofloxacin (CIP) was from Double-Crane Pharm (Beijing, China).
Detection of *M. hyorhinis* by Quantitative PCR (qPCR)

DNA was extracted from AGS or BGC823 cells after *M. hyorhinis* infection by DNA lysis buffer (50 mM Tris pH 8.5, 1 mM EDTA, 0.5% Tween-20, and 200 mg/L proteinase K) according to the standard protocol. qPCR was performed with 30 ng DNA and SYBR Green Real-time PCR 2×premix kit (Takara, Otsu, Japan) using Step One system from ABI (Foster City, CA, USA). The reaction programs and p37-specific primers (forward: 5′-TATCTCATTGACCTTGACTA-3′, reverse: 5′-ATTTCGCCCAATAGCATTTG-3′) were reported previously [22]. To compare *M. hyorhinis* DNA levels in cells, GAPDH (forward: 5′-TGAAGGTCGGAGTCAACGG-3′, reverse: 5′-CCTGGAAGATGGTGATGGG-3′) was amplified as control and the data were analyzed using the 2^ΔΔCt method. Primers were synthesized by Sangon (Shanghai, China).

Western Blotting

Cells were harvested from culture dish with 2×SDS loading buffer. Polyacrylamide gel electrophoresis (SDS-PAGE) and Western blotting were performed as previously described [23].

Cell ELISA

96-well were seeded with cells (1×10⁴/well), followed by treatment of indicated peptides and infection of *M. hyorhinis* for 24 hr. Cells were immobilized with 0.05% glutaraldehyde for 10 min, then cell ELISA was performed as described previously [24].

Solid-Phase Binding Assay and Pull-Down Assay

Recombinant GST-p37 and GST proteins were generated and purified as previously described [17]. GST-p37 and GST were diluted in buffer (0.1M Na₂CO₃, 0.1M NaHCO₃, pH 9.6) and coated in 96-wells plates at 4°C overnight. The plates were washed by PBS for three times and blocked by 5% skimmed milk/PBS at room temperature (RT) for 2 hr. After washing with PBS, indicated concentrations of biotin-conjugated A2PP (synthesized by Sbsbio) was added and incubated at RT for 2 hr. After washing with PBST for 3 times, streptavidin-conjugated HRP (Baltimore Pike, West Grove, PA, USA) was added and incubated at RT for 30 min. After color development with Ortho-Phenylenediamine (Sigma), optical density at 490 nm (OD490) was recorded with a Microplate reader (Bio-rad 550). For pull-down assay, 100 ng GST-p37 or GST protein was co-incubated with 20 μM biotin-A2PP and streptavidin beads (GE Healthcare, Pittsburgh, PA, USA) in binding buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 0.5% NP-40, 0.5 mM DTT, 1 mM PMSF, and 1× complete protease inhibitors) at 4°C overnight. The precipitates were washed with binding buffer for four times and analyzed by Western blotting.

Co-Immunoprecipitation

*M. hyorhinis* infected cells (BGC823 or AGS) were homogenized in lysis buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 1% Triton X-100, 0.5 mM DTT, 1 mM PMSF, and 1× complete protease inhibitors) at 4°C for 10 min. After 12,000 g centrifugation for 10 min at 4°C, supernatants were recovered. Protein lysates (500 μg) incubated with 20 μM A2PP or ConP, 1 μg anti-ANXA2 plus protein G sepharose beads (GE Healthcare) at 4°C overnight. Pre-immune IgG (1 μg) was used as control. Precipitated beads were washed with lysis buffer for four times, eluted in 2× loading buffer, boiled, and analyzed by Western blotting.
Immunofluorescence Assay
Cells were seeded on coverslips and cultured overnight. The next day, cells were treated with indicated peptides and infected with *M. hyorhinis* for 24 hr. Then cells were washed with ice cold PBS for three times, fixed in 4% paraformaldehyde for 15 min at RT. After blocking for 1 hr in 5% BSA/PBS, cells were incubated with indicated antibodies at RT for 1 hr. Next, cells were washed for 3 times again with PBST and incubated with FITC or TRITC-labeled secondary antibodies for 30 min at RT. Following washing with PBS and counterstaining with DAPI, cells were mounted on 50% glycerol/PBS. A ZEISS LSM780 confocal microscope (ZEISS Microsystems, Oberkochen, Germany) was used to observe the localization of indicated proteins.

Cell Migration Assay
Transwell chamber with 8.0 μm pore membranes (Corning, NY, USA) was used in the cell migration assay. The bottom chamber was filled with 800 μL medium containing 10% FBS as chemoattractant. Cells were resuspended in serum-free medium containing *M. hyorhinis* plus A2PP or ConP, then were carefully transferred onto the top chamber of each Transwell apparatus at a density of 2×10^5 cells/mL (120 μL/chamber). Cells were allowed to migrate for 24 hr at 37°C. The top surface of each membrane was cleared of cells with a cotton swab. Cells penetrated to the bottom side of the membrane were fixed in cold methanol, stained with 0.1% crystal violet, and counted in nine randomly selected microscopic fields per well. Each sample was prepared in triplicate chambers and each experiment was repeated for at least 3 times.

Cell Proliferation Assay
Gastric cancer cells were seeded in 96-well culture plates at density of 5×10^3/100 μL/well in triplicates, and were treated with indicated reagents. Proliferation of cells were quantified by the cell confluence with a CloneSelect Imager (Molecular Devices, Sunnyvale, CA, USA).

Microarray Analysis and Real-Time RT-PCR
AGS and BGC823 cells (1×10^6 per 10 cm^2 plate) were treated with A2PP, CIP, MYCO I and MYCO II for 24 hr, washed with PBS, and harvested in Trizol reagent. RNA samples were examined in OE Biotechnology (Shanghai, China) by using Affymetrix GeneChip™ Prime-View™ Human Gene Expression Array. Microarray data has been deposited in NCBI Gene Expression Omnibus (GEO) (accession no.GSE73777). Raw data was recorded by using Affymetrix GeneChip Command Console (version 4.0, Affymetrix). Next, Genespring software (version 12.5, Agilent Technologies) was employed to finish the basic analysis with the raw data. To begin with, the raw data was normalized with the RMA algorithm. Differentially expressed genes were then identified through fold change. The threshold set for up- and down-regulated genes was a fold change ≥2.0. Afterwards, GO and KEGG analysis were applied to select out genes that related to cell cycle and cell apoptosis. Real-time RT-PCR was used to validate the results of microarray and performed according to the manufacturers’ instruction (SYBR, TOYOBO). Expression levels of related genes were normalized through *GAPDH* and the 2^−ΔΔCT was used to calculate relative gene expression. The primers for real-time RT-PCR (ATF, forward, 5′-GAGTGGCGACAGGATAGAGC-3′, reverse, 5′-TTTAGCCTTCCTCCCTTTAGC-3′; DDIT3, forward, 5′-GGCGATGGAGAAAGAAC-3′, reverse, 5′-ACCATTCGGTCAAATCGAGGC-3′; CEBPB, forward, 5′-AGCGACGAGTACAGATTCCCG-3′, reverse, 5′-AGCTGCTCACCTTCTTTCTG-3′) were synthesized by Sangon.
Statistical Analysis
Data were presented as mean ± SD. The differences were analyzed by ANOVA using SPSS 11.0 software and P < 0.05 was considered statistically significant.

Microarray Accession Number
Microarray data has been deposited in NCBI Gene Expression Omnibus (GEO) (accession no. GSE73777).

Results
A2PP Has No Effect on the Viability or Migration of the Gastric Cancer Cells
In order to evaluate the role of ANXA2’s N-terminal domain in M. hyorhinis infection, we firstly synthesized A2PP peptide corresponding to N-terminal of ANXA2 (Fig 1A). Biotin labeled A2PP was shown to specifically interact with GST-p37 in the solid phase binding assay (Fig 1B) and pull-down assay (Fig 1C). We noticed that A2PP didn’t affect the morphology of AGS and BGC823 cells (Fig 1D). In addition, in concentrations less than 20 μM, A2PP didn’t inhibit cell proliferation (Fig 1E), suggesting that A2PP has minimal toxicity to cells. EGFR has been implicated in regulating ANXA2 phosphorylation [17,25,26], while A2PP had no effects on the phosphorylations or protein levels of EGFR and ANXA2 [Fig 2A]. Moreover, A2PP had no effects on the migration of AGS and BGC823 cells (Fig 2B and 2C). Cellular localization of ANXA2 is regulated by its phosphorylation, which is critical for its function [17,27,28]. We found A2PP didn’t change the subcellular localization of endogenous ANXA2 (Fig 2D). These results indicate that A2PP alone has no effects on malignant phenotypes or EGFR-ANXA2 signaling of gastric cancer cells.

A2PP Decreases M. hyorhinis Infection
Our previous work has shown that N-terminal of ANXA2 mediates M. hyorhinis infection [17]. As shown by the result of cell ELISA assay, we found that M. hyorhinis infection was blocked by A2PP in a dose-dependent manner in BGC823 and AGS cells, but ConP-treated cells were still highly infected by M. hyorhinis (Fig 3A). These results were supported by qPCR assays (Fig 3B). Consistently, A2PP also decreased protein levels of p37 protein in the Western blotting analysis (Fig 3C and 3D). Meanwhile, levels of phophos-AXNA2 and phophos-EGFR in the infected cells were down-regulated by treatment with A2PP (Fig 3C and 3E), while total levels of AXNA2 and EGFR were relatively stable (Fig 3C). In the immunofluorescence analysis, A2PP was found to inhibit M. hyorhinis infection-induced p37 accumulation and co-localization between p37 and ANXA2 (Fig 4A). In the co-immunoprecipitation assay with protein lysates from M. Hyorhinis infected cells, A2PP decreased the interaction between p37 and ANXA2 (Fig 4B). Together, these evidences supported the idea that A2PP could decrease M. hyorhinis infection and confirmed our previous discovery that the N-terminal of ANXA2 is required for mediating M. hyorhinis infection [17].

A2PP Suppresses M. hyorhinis-Induced Migration
Our previous studies suggested that M. hyorhinis infection could promote migration of gastric cancer cells [17, 23]. In this study, we noticed that M. hyorhinis-promoted migration of gastric cancer cells was not affected by ConP, but was dose-dependently inhibited by A2PP (Fig 5A and 5B). These results suggest that A2PP inhibited M. hyorhinis-induced migration, which was
Fig 1. A2PP binds to p37 of M. hyorhinis and has minimal effect on the proliferation of gastric cancer cell. (A) Schematic diagram of amino acid sequence of N-terminal of ANXA2 polypeptide (A2PP). (B) Solid-phase binding assay. OD490 of 15 nM biotin-A2PP to GST was set as 1 and relative bindings were calculated. Mean ± SD of three independent assays with triplicate samples. *** P < 0.001. (C) Streptavidin pull-down assays identified Biotin-A2PP as a GST-p37 binding polypeptide. (D) Cell morphology of AGS and BGC823 following indicated concentrations of A2PP treatment for 24 hr and 48 hr. (E) Proliferation of gastric cancer cell lines (AGS and BGC823) treated with increasing concentration of A2PP for 72 hr. Mean ± SD from three experiments with triplicate samples.

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associated with its ability to decrease infection-promoted phosphorylations of EGFR and ANXA2 (Fig 3C and 3E).

**A2PP Has Essential Motif to Decrease *M. hyorhinis* Infection**

To characterize the critical motif of A2PP, we sought to determine whether diverse truncated peptides of A2PP (Fig 6A) could reduce *M. hyorhinis* infection. In the qPCR analysis, A2PP...
and truncated peptides A2PP-Nd4, Nd8, Nd12, Cd4, Cd8, and Cd12 reduced *M. hyorhinis* infection in gastric cancer cells, but A2PP-Nd16 and A2PP-Cd16 failed to decrease *M. hyorhinis* infection (Fig 6B). Similar pattern of inhibition was obtained from Western blotting assay (Fig 6C). These results suggest the existence of a specific motif of A2PP is essential for its ability to reduce *M. hyorhinis* infection. To further map the core sequences of this potential motif, we synthesized the two peptides corresponding to 11th-20th aa (termed as A2PP-10aa) and 13th-18th aa (termed as A2PP-6aa) (Fig 7A). In qPCR assay, both peptides could decrease *M. hyorhinis* infection (Fig 7B), which was supported by Western blotting assay (Fig 7C). Notably, the inhibitory effects of A2PP-10aa were better than those of A2PP-6aa, but the effects of both

**Fig 3.** A2PP reduces *M. hyorhinis* infection. (A) Cell ELISA analysis of p37 (OD 490 nm) of p37 protein in AGS and BGC823 cells infected with 10^5 CCU (color changing units)/ml of *M. hyorhinis* and treated with A2PP or ConP for 24 hr. *M. hy., M. hyorhinis*. Mean ± SD from 3 experiments with triplicate for each sample. (B) Quantitative PCR (qPCR) analysis of p37 in AGS and BGC823 cells infected and treated as in (A). Mean ± SD from 3 experiments with triplicate for each sample. (C) Western blotting of p37, p-EGFR, EGFR, p-ANXA2 and ANXA2 from AGS and BGC823 cells treated as in (A). (D) Quantification of p37 protein levels of in (C). Levels of p37 were normalized to those of GAPDH. Mean ± SD from 3 independent experiments. (E) Quantification of p-EGFR and p-ANXA2 levels in (C). Levels of p-EGFR or p-ANXA2 were normalized to those of EGFR or ANXA2. Mean ± SD from 3 independent experiments. **P < 0.01; *** P < 0.001; n.s, no significance.

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peptides were less robust as those of A2PP (Fig 7B and 7C). Therefore, the central sequences (11th - 20th) of A2PP could be the essential motif to inhibit M. hyorhinis infection.

Comparison of A2PP with Other Drugs in Preventing M. hyorhinis Infection

M. hyorhinis was one of the main pollution sources of cell culture [13,14,15]. The best way to eliminate M. hyorhinis infection is to discard cells and quickly sterilize all the contacted vessels, but some infected cells couldn’t be replaced in several cases. Consequently, it is imperative to use specific antimicrobial approaches to prevent or eliminate M. hyorhinis infection. There have been different antimicrobial agents for preventing M. hyorhinis from infecting. For example, Ciprofloxacin (CIP) could eradicate M. hyorhinis in a suitable concentration [29]. Two antimicrobial components named as mycoplasma I (MYCO I) and mycoplasma II (MYCO II) could reduce M. hyorhinis infection by blocking its protein and nucleic acid synthesis. However, some concerns such as low sensitivity, strong resistance and high toxicity may limit their applications in cell culture. Based on these considerations, we believe it is necessary to compare the efficiencies of A2PP, CIP, MYCO I and MYCO II in preventing M. hyorhinis infection. By Western blotting and qPCR assays, we found that A2PP’s inhibitory effect on M. hyorhinis infection was similar to that of MYCO I, but was better than those of CIP and MYCO II (Fig 8A and 8B). Interestingly, we found that A2PP could also efficiently eliminate the M. hyorhinis...
infection in heavily infected cells in the process of passages from P1 to P3 (Fig 8C and 8D). Additionally, compared with MYCO I and MYCO II, A2PP had less inhibitory effect on the cells proliferation (Fig 9A). To explore the mechanisms of A2PP, CIP, MYCO I and MYCO II’s effects on cells proliferation, we performed microarray analysis and screened a subset of differentially expressed genes in A2PP, CIP, MYCO I and MYCO II treated cells. Quantitative RT-PCR analysis showed increased expression of apoptosis-related genes (ATF5, DDIT3, CEBPB) by MYCO I and MYCO II treatment, but little changes were observed in A2PP and CIP groups (Fig 9B). These results indicate that A2PP has less toxicity and better preventative potential in blocking *M. hyorhinis* infection in cultured cells.

**Discussion**

Mycoplasma infection and contamination are still prevalent today and bring considerable risks to patients and research quality. In recent years, several studies suggested that mycoplasma infection was also associated with tumorigenesis [3–8]. Mycoplasma infection could cause the DNA damage, affect gene expression, and disrupt the cell cycle checkpoint and apoptotic response [30]. *M. hyorhinis* was found in 56% of gastric cancers, 55.1% of colon cancers and 39.7% breast cancers tissues [20]. Besides, serological testing showed that 36% benign prostatic hyperplasia (BPH) and 52% prostatic cancer tissues were *M. hyorhinis* positive [4]. These studies suggest a possible association between *M. hyorhinis* infection and the occurrence of tumors.
What’s more, in the process of cell culture, the most common contamination sources are mycoplasma, bacteria and mold, but the contamination rate of mycoplasma is relatively higher [13]. Results from various groups have shown that the rate of mycoplasma contamination varied from 15 to 80%, some even reached 100% [15,31]. An analysis of DNA sequences

Fig 6. Effects of truncated A2PP peptides on *M. hyorhinis* infection. (A) Schematic diagram of truncated peptides of A2PP. (B) qPCR analysis of p37 in AGS and BGC823 cells infected with 10^5 CCU/ml of *M. hyorhinis* and treated with 20 μM indicated peptides for 24 hr. Mean ± SD from 3 experiments with triplicate for each sample. (C) Western blotting of p37 from AGS and BGC823 cells infected and treated as in (B). Mean ± SD from 3 independent experiments. doi:10.1371/journal.pone.0147776.g006
from 1000 genomes project implied that 7% of the samples were contaminated by mycoplasma [32]. However, preventing mycoplasma infection and contamination is difficult. Firstly, because of pleomorphic, plasticity, filterability and easy solubility, mycoplasmas could pass through microfiltration membrane easily, rendering them exist on the host cell surface or to be endocytosed by cells [1,13,33]. Secondly, mycoplasma lacks cell walls, rendering them insensitive to antibiotics which inhibit cell wall synthesis, such as penicillin. Effective antimicrobials do exist, but their continuous applications in cell culture is not recommended because of the toxicity to cells. Finally, mycoplasma is a atypical bacteria, causing difficulties in correct diagnosis [34].

Numerous anti-mycoplasma drugs have been developed, such as Erythromycin, Leucomycin, Roxithromycin, Ofloxacinc, and Ciprofloxacin. However the adverse effects, toxicity, and resistance still bring trouble for pollutio abatement [10,11,35–38]. Besides, continuous administration of these antibiotics couldn’t completely eliminate mycoplasma for one or even more weeks [39,40]. Therefore, it is critical to find new ways that can eliminate the mycoplasma infection or contamination effectively and timely. The solution to this aim can probably be found through the characterization of molecular mechanisms underlying mycoplasma infection.

Our previous study found that the interaction of P37 protein with AXNA2 mediated M. hyorhinis infection [17]. In this study, we firstly synthesized the N-terminal polypeptide of ANXA2 (A2PP) and validated its specific interaction with GST-p37 protein in the solid-phase binding and streptavidin pull-down assays. Inspired by such interaction, we wondered whether A2PP could antagonize infection of M. hyorhinis. We found that A2PP, in an appropriate concentration (20 μM), decreased M. hyorhinis infection, inhibited infection-promoted migration, and reduced infection-provoked the phosphorylations of EGFR and ANXA2 in gastric cancer cells. These effects were associated with decreased interaction between p37 and ANXA2. It...
should be noted that A2PP alone exhibited minimal effects on proliferation of cells and phosphorylations of EGFR and ANXA2, suggesting that A2PP is likely to be a useful reagent to reduce M. hyorhinis infection. This notion was supported by comparisons of A2PP’s and commercial antibiotics’ effects on cell proliferation and M. hyorhinis infection, which showed that A2PP indeed had less toxicity to cells but strong ability to counteract infection.

Despite that A2PP, a 30 aa polypeptide, could decrease M. hyorhinis infection effectively, we still need to understand whether truncated forms of A2PP could achieve similar anti-M. hyorhinis ability. By using several truncated peptides of A2PP, we found that the central sequences...
(11<sup>th</sup> -20<sup>th</sup>) of A2PP could be the essential motif to inhibit <i>M. hyorhinis</i> infection. However, the inhibitory effects of truncated peptides were still not as strong as that of full length A2PP, suggesting that flanking sequences are essential to maintain proper structure to achieving efficient binding with p37. Further studies are required to find the effects of modification or/and mutation of this motif on inhibiting <i>M. hyorhinis</i> infection.

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**Author Contributions**

Conceived and designed the experiments: SY CS. Performed the experiments: SY. Analyzed the data: SY LQ CS. Wrote the paper: SY LQ.

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