Effects of Calcium on the Interactions of Acinetobacter baumannii with Human Respiratory Epithelial Cells

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Abstract

Background Investigating the factors that influence Acinetobacter baumannii (Ab) adhesion/invasion into host cells is important to understand its pathogenicity. Metal cations have been shown to play an important role in regulating the biofilm formation and increasing the virulence of Ab; however, the effects of calcium on host-bacterial interactions have yet to be clarified. Here, the dynamic process of the interactions between Ab and human respiratory epithelial cells and the effects of calcium on host-bacterial interactions were explored using the technologies of microscopic imaging, quantitative PCR and real time cellular analysis (RTCA).

Results The concentration of calcium, multiplicity of infection and co-culture time were demonstrated to have effects on host-bacterial interactions. A unique "double peak" phenomenon changed to a sharp "single peak" phenomenon during the process of Ab infection under the effects of calcium were determined based on the time-dependent cell response profiles. Moreover, calcium can increase Ab adhesion/invasion of epithelial cells by regulating the expression of Ab-related genes (ompA, bfmRS, abal).

Conclusions Effective control of calcium concentrations can provide new ideas for the prevention and treatment of multi-drug resistant Ab.

Background

Over the last decades, Acinetobacter baumannii (Ab) has ranked among the most important Gram-negative nosocomial pathogens worldwide because of its increased rate of clinical isolation and the continuous emergence of multidrug-resistant strains[1, 2]. Ab is known for its prolonged survival throughout hospital settings, owing to its great capacity to adhere to abiotic or biotic surfaces (e.g., human respiratory epithelial cells) and its high tolerance of environmental conditions[3, 4]. Thus Ab is forecasted to become one of
the biggest challenges to medical care. Adhesion of Ab to epithelial cells is considered an essential first step in colonization and infection. Colonization or infection with Ab occurs mostly in critically ill patients and causes severe pneumonia or bloodstream infections that result in increased in morbidity and mortality in these patients, which is a troublesome problem for clinical diagnosis, treatment and prevention[1, 5, 6]. Moreover, it is important to emphasize that colonization with Ab is more common than infection, even in susceptible populations[5, 7-9]. Although the pathogenicity of Ab is generally low, bacterial colonization is the greatest risk factor for infection in hospitals. Once the balance between host and microbe tips towards the development of an infection, the result can be severe and possibly lead to an increased probability of Ab hospital outbreaks. Therefore, studies on controlling Ab infection and colonization are of great practical significance and value.

In view of the seriousness of the antibiotic resistance of and mass colonization by Ab on biotic surfaces, the pathogenicity of Ab has attracted wide attention, especially in regard to the host-bacteria interaction and the process and molecular mechanism of Ab adhesion/invasion into human respiratory epithelial cells. Host-bacterial interactions are influenced by many factors including biofilm formation and endotoxin production. In addition, biofilm formation represents an important factor associated with virulence and is affected by bacterial fimbriae, outer membrane proteins, adhesins, metal ions, quorum sensing, and complex regulatory networks (e.g., two-component regulatory systems), among others[10]. Investigating the factors that influence Ab adhesion/invasion into epithelial cells is important to understand its pathogenicity.

Metal cations have been shown to play an important role in regulating biofilm formation and the differential expression of Ab-related genes, as well as in increasing the virulence of Ab and its ability to adhere to epithelial cells; however, the effects of calcium on host-
bacterial interactions have yet to be clarified[11, 12]. Studies have shown that bacterial infection can lead to a destabilization of the cellular calcium homeostasis and the activation of the calpain system ultimately triggering cell death, which suggests that changing the concentration of calcium in the environment may have a significant impact on the pathogenicity of Ab[13]. Lee et al. showed that bacterial attachment and biofilm formation on human respiratory epithelial cells and plastic surfaces were markedly reduced in the presence of the chelating agent EDTA (low levels of metal cations such as Ca$^{2+}$ and Mg$^{2+}$) through the analysis of a group of multidrug-resistant Ab clinical isolates[14]. These data suggested that high concentrations of calcium may promote biofilm formation by Ab and enhance its ability to adhere to respiratory epithelial cells. However, this work did not specify the effects of calcium. It is important to explore the effects of calcium on host-bacterial interactions and to elucidate the functional mechanism, thus making it possible to take effective measures to control bacterial biofilm formation, adhesion and invasion and to ultimately provide new ideas for addressing the challenges of colonization and infection with multidrug-resistant Ab.

A label-free and noninvasive detection system (RTCA S16 system, ACEA Biosciences Inc.) based on dynamical and quantitative monitoring of cellular impedance in real time can produce specific time-dependent cell response profile (TCRPs) patterns. This approach can provide biological information related to cellular physiological function for studying host-bacterial interactions[15]. Thus, in this work, we developed microscopic imaging, quantitative PCR (qPCR) and TCRP methods for continuously monitoring the interactions between Ab and human respiratory epithelial cells and the effects of calcium on host-bacterial interactions. Our research can be used for the study of calcium-mediated signalling pathways in human respiratory epithelial cells infected with Ab, which provides a basis for studying the pathogenicity of Ab[16].
Results

Optimum multiplicity of infection (MOI) of Ab to human respiratory epithelial cells

The effects of Ab on the morphology and proliferation of human respiratory epithelial cells at different MOIs and co-culture time points were determined by inverted microscopy and are shown in Table 1 and Fig. 1A. After host-bacterial co-culture for 2 h, there were relatively few bacteria and almost none were adhered to epithelial cells. The differences among the groups were also relatively small (especially in the control group and the MOI 1 and MOI 10 experimental groups). After co-culture for approximately 4 h, the differences among the groups became more obvious. As the MOI increased, the effects of Ab on the epithelial cells were also increased (the differences between the MOI 50 and MOI 100 experimental groups were not obvious). Additionally, the changes in cell morphology and proliferation at 4 h of co-culture were more typical compared to other co-culture time points. Thereby, the co-culture time point of 4 h was selected as suitable for subsequent studies. In addition, at 6–8 h of co-culture, the quantity of bacteria was relatively large, and they were not suitable for further research and analysis.

Bacterial invasion (including strong adhesion) to epithelial cells at different MOIs was determined by qPCR. Host cells and bacteria were co-cultured for 4 h and $1 \times 10^5$ cells were isolated for qPCR. The qPCR results (threshold cycles, Ct values) are shown in Fig. 1B. With increased MOI, the invasion of Ab into human respiratory epithelial cells gradually increased (the smaller the Ct value was, the more bacterial invasion occurred). There was no invasion in the control group. The differences in the Ct values among all the experimental groups were statistically significant ($P < 0.05$). The MOI 100 group (Ct values: $23.56 \pm 0.04$) < the MOI 50 group ($24.31 \pm 0.05$) < the MOI 10 group ($27.36 \pm$
0.05) < the MOI 1 group (30.26±0.11). The differences in the Ct values between the MOI 10 and MOI 50 groups or between the MOI 1 and MOI 10 groups were approximately 3, while the differences between the MOI 50 and MOI 100 groups were relatively small (an approximately 0.8 difference in the Ct values).

The TCRPs were determined by real time cellular analysis. Ab infection induced the cell index (CI) to rise and then fall (Fig. 1C). As the MOI decreased, the time to reach the peak CI was gradually delayed and the peak CI was also higher. A unique “double peak” phenomenon emerged during the process of infection (the smaller the MOI was, the more obvious the phenomenon was). The bacterial concentration of the MOI 50 experimental group was most appropriate, and the phenomenon characteristics were more typical compared to the MOI 100 group. Therefore, the bacterial concentration (1 × 10^8 CFU/ml) corresponding to the MOI 50 was suitable for subsequent studies based on the above results.

The effects of Ab on different incubation states of human respiratory epithelial cells

Ab had an almost identical effect on the different incubation states of epithelial cells (Fig. 2A). In the early stages of host-bacterial co-culture, epithelial cells can adhere to the plastic surface, although bubble-like dead cells (nuclear pyknosis, cell swelling and dissolution, Additional file 1: Fig. S1) can be seen (after approximately 4 h). However, as the co-culture time is extended and the bacteria increase in quantity, the changes in cell morphology and proliferation, as well as the bacterial aggregation phenomenon became more typical (full field of bacteria after 8 h). It was observed that Ab had an effect on the adherent growth of epithelial cells. After 24 h, bacteria dominated the entire cell culture dish, while the host cells were all dead.
The dynamic processes (as determined based on the TCRPs) of the interactions between Ab and epithelial cells in different states of cell incubation were also similar (Fig. 2B). For instance, initial Ab infection had little effect on cell growth, and the “double peak” phenomenon emerged during Ab infection (this was not typical when the CI was about to enter the platform period). Almost identical cell growth curves were observed for the 0.45% sodium chloride solution (NaCl) and sterile distilled water groups. Compared to the untreated group, slight decreases or increases in the CI of the 0.45% NaCl and sterile distilled water groups may have been associated with the dilution of nutrients or metabolites in the culture medium. Therefore, the effect of 0.45% NaCl in the bacterial suspension on the adherent growth of epithelial cells could be considered negligible.

Effects of calcium on Ab proliferation and biofilm formation

As shown in Fig. 3 we found that high concentrations of calcium could contribute to the proliferation of Ab and that this effect was more pronounced with time. Additionally, the bacterial biofilm formation was more obvious as the concentration of calcium increased (Additional file 2: Table S1). Ab deficient in ompA may reduce biofilm formation.

Effects of calcium on the morphology and proliferation of human respiratory epithelial cells

It was difficult to distinguish differences among the groups using an inverted microscopy, so we next used TCRPs to evaluate the effects of calcium on the proliferation of human respiratory epithelial cells. With increasing calcium concentrations (≤4.4 mmol/L) and extended culture times (≤24 h), the CI of epithelial cells increased significantly (Additional file 3: Fig. S2). The CI values of each group under different calcium concentrations and culture times (0 h, 2 h, 4 h, 6 h, 8 h, 12 h and 24 h) were compared by multivariate ANOVA with repeated measures and the SNK test. The results showed that the
time as a factor was effective ($P = 0.00$), meaning that the CI changed over time. There was a positive interaction between the time and treatment (calcium concentrations) factors ($P = 0.00$). Exogenous calcium supplementation can promote the growth of human respiratory epithelial cells (Additional file 4: Table S2). The higher the calcium concentration was ($\leq 4.4$ mmol/L) and the longer the culture time was ($\leq 24$ h), the more significant the promoting effect on the CI values was.

**Effects of calcium on host-bacterial interactions**

The role of calcium in bacterial proliferation was the same as has been described above. With increasing co-culture time and increased calcium concentrations, the bacterial aggregation phenomenon (biofilm formation) became more obvious (Additional file 5: Fig. S3). After 8h, there was an obvious increase in the quantity of bacteria. After approximately 24h, the bacteria encompassed the full field, while massive numbers of host cells were dead. It is known that Ab can have effects on the adherent growth of epithelial cells; however, the effects of calcium on the interactions between Ab and epithelial cells could not be determined using an inverted microscopy.

Host and bacteria were co-cultured in calcium-supplemented medium for 2 h, 4 h and 6 h. The effects of calcium on bacterial invasion (including strong adhesion) of epithelial cells ($1 \times 10^5$ cells as the standard) were determined using qPCR and shown in Fig. 4. The results were negative for control group II. The Ct values were compared by univariate ANOVA with repeated measures and the SNK test. The time as a factor was significant with a $P = 0.00$, which meant that the amount of Ab epithelial cell invasion changed over time. Moreover, treatment (calcium concentrations) as a factor was effective ($P = 0.00$), meaning that the amount of Ab epithelial cell invasion differed depending on grouping, and this difference among the groups was statistically significant ($P < 0.05$). There was a
positive interaction between the time and treatment factors ($P = 0.00$). The role of time as a factor varied depending on the group. As a result, the higher the calcium concentration was and the longer the co-culture time was, the more frequently Ab epithelial cell invasion was observed.

The TCRPs showed that either initial Ab infection or the action of calcium can induce the CI to rise. With the increasing calcium concentration in host and bacteria culture medium ($\leq 4.4$ mmol/L), the faster the CI rose, the higher the peak CI was (6–8h), and the more significant cell growth stimulation was. By contrast, the CI declined rapidly with prolonged host-bacterial interactions (the higher the calcium concentration was, the faster the CI declined). Meanwhile, a sharp “single peak” phenomenon occurred during the infection (Fig. 5).

**Effects of calcium on the expression of Ab-related genes**

Calcium can affect the expression of Ab-related genes. The recA gene was used as an internal reference control. Both negative controls (I and II) had no amplification. Relative changes in the expression levels of target genes ($ompA$, $bfmRS$ and $abaI$) between experimental groups and control I group were calculated by the $2^{\Delta\Delta Ct}$ method. The trends were different between the abiotic and cellular environments that the bacteria were cultured in (Fig. 6).

There was no significant difference in the expression level of $ompA$ among the groups cultured in the abiotic environment ($P > 0.05$), whereas expression of this gene showed significant differences in the cellular environment ($P < 0.05$). In the cellular environment, relative changes in expression levels between group c and control I, as well as between group d and control I were small ($P > 0.05$); however, $ompA$ expression in group b was approximately 4-fold greater than that of the control I group.
In the abiotic environment, there were significant differences in the expression of *bfmRS* between the experimental groups and the control I group (*P* < 0.05). With the increase in calcium concentration in the culture medium, *bfmRS* expression in the experimental groups showed a decreasing trend; its expression in group d was approximately 0.31-fold lower than that of the control I group. In the cellular environment there was no significant difference in the expression levels between group a and the control I group (*P* > 0.05). Contrary to the expression trends in the abiotic environment, expression levels in groups b and c were approximately 2.3-fold higher than that of the control I group, while the level in group d was approximately 4-fold higher.

In the abiotic environment *abaI* displayed similar levels of expression among groups b, c and the control I group (*P* > 0.05). The expression of *abaI* in group a was approximately 0.5-fold lower than that of the control I group, while its expression in group d was approximately 2-fold higher. In the cellular environment, the *abaI* expression in groups a and b were approximately 0.4-fold lower than that of the control I group, while in group d its expression reached approximately 17-fold higher.

**Discussion**

Ab infection and colonization co-exist, mainly causing respiratory infections (such as ventilator-associated pneumonia)[17] that seriously endanger human life and quality of life and result in a major economic burden[18]. Elucidating the molecular mechanisms of the interactions between Ab and host cells is of great significance to further our understanding of the pathogenic mechanisms of this bacteria and to propose new prevention and treatment strategies.

Based on the normal blood calcium concentration of 2.25 - 2.75 mmol/L, the concentration of calcium in the media used in these experiments was controlled within 1.4 - 4.4 mmol/L to simulate the environment of the body. Our study found that exogenous calcium
supplementation can promote the proliferation of Ab and the adherent growth of human respiratory epithelial cells, as well as induce differential expression of Ab-related genes. In addition, calcium also played an important role in host-bacterial interactions, resulting in increased Ab adhesion/invasion of human respiratory epithelial cells and thereby increasing the degree of bacterial infection in the host cells. The higher the calcium concentration is (especially in the case of high calcium) and the longer the culture duration, the more severe the degree of host cells bacterial infection is. Calcium may affect the host-bacterial interaction by contributing to several factors.

RTCA detection is an important technique that can reflect changes in cell morphology (including size, shape, stretching, etc.), number and adhesion. Compared to traditional endpoint detection, RTCA has the advantages of being non-invasive and highly accurate, as well as providing real-time monitoring, complete TCRPs, and easy operation. It is widely used in cytology research, such as in cell migration and invasion assays, cytotoxicity tests, gene regulation and cell-microenvironment interactions[15, 19–21]. Therefore, the obtained TCRPs can provide better information on the effects of calcium on host-bacterial interactions, complementing the results of our microscopy observations.

Interestingly, RTCA detection in the present study found unique “double peak” (calcium-free, Fig. 1C and 2B) and sharp “single peak” (with calcium, Fig. 4) phenomena during bacterial infection of epithelial cells. The sharp “single peak” may suggest that during the initial stages of bacterial infection, calcium stimulation promoted the rapid growth of cells (the observed peak). When the amount of bacteria reached the critical point, calcium in turn enhanced bacterial adhesion/invasion of epithelial cells, thereby rapidly decreasing the cell index (CI). The mechanism of the “double peak” phenomenon needs to be further studied and analysed. Possible mechanisms include that cell death was caused by bacteria and cells stimulating, adapting and interacting with each other. Specifically, the bacteria
were added to the cell culture environment as exogenous foreign bodies, which stimulated the proliferation of the cells in the initial stage of infection (the appearance of the first peak), but then the bacteria inhibited host cell proliferation due to its own rapid proliferation (the decrease in the first peak). With prolonged host-bacterial interactions, the cells became tolerant to the bacterial inhibition (adaptation) and continued to proliferate (the appearance of the second peak), after which the two competed with one another in the nutrient-rich environment. Additionally, the increasing toxic effects of the bacteria on the epithelial cells resulted in a rapid reduction in the CI (the decrease in the second peak).

Ab-ompA, a highly conserved outer membrane protein, is also an important virulence factor that plays an important role in bacterial infection (which is also supported by our research) and causes an upregulation of epithelial cellular immune response signalling pathways[16, 17]. It has been shown that Ab-ompA secretes and transmits virulence factors through outer membrane vesicles. Translocation of ompA-containing vesicles into host cells can result in host cell apoptosis, whereas ompA mutants fail to induce cell death[22]. Thus, blocking or inhibiting the expression of Ab-ompA will greatly reduce Ab adhesion/invasion into host cells[23]. Our studies showed that compared with abiotic environments, Ab-ompA expression is obviously changed in cellular environments. Increased calcium content (especially at 2.4 mmol/L) may contribute to the expression of Ab-ompA, which is consistent with our finding that calcium can aggravate bacterial adhesion/invasion into host cells, while other factors may be involved under high calcium conditions. Our finding that Ab-ompA expression only changes significantly during host-bacterial interactions indicates that ompA is a virulence factor. Additionally, changing the calcium concentration of the environment may have a significant impact on Ab pathogenicity[13]. OmpA has a positive effect on biofilm formation was also proved in our
experiment. Based on the above findings, calcium may enhance bacterial virulence and promote a certain degree of ompA expression. Therefore, if calcium concentration is kept at the lower level of normal range (2.2 - 2.4 mmol/L), it may be possible to prevent Ab invasion caused by the upregulation of ompA expression.

Ab-bfmRS is a key factor in the survival of Ab in the environment and is comprised of a sensor kinase (bfmS) and a response regulator (bfmR). Inactivation of bfmRS not only reduces biofilm formation but also leads to the loss of bacterial adhesion to eukaryotic cells[24]. Interestingly, the composition of the culture medium and the interaction of Ab with abiotic surfaces play a significant role when the BfmRS system is not expressed[12]. In addition, environmental signals are important for the bfmRS regulatory system and biofilm formation, and are involved in inducing the bacterial morphology responsible for interacting with abiotic surfaces[12]. These observations may explain why exogenous calcium supplementation in cellular and abiotic environments has completely different effects on bfmRS expression in our research. In abiotic environments, calcium inhibits the expression of bfmRS, and the two are negatively correlated, while in the cellular environment, calcium promotes bfmRS expression. This is consistent with the results indicating that calcium may promote biofilm formation and increase Ab adhesion/invasion.

Calcium may be an environmental signalling molecule that affects the expression of Ab-bfmRS. As Ab-bfmRS expression is affected by many factors, how calcium causes a downregulation of bfmS expression in abiotic environments remains to be further studied. Thus, increased calcium concentrations may promote biofilm formation and enhance bacterial adhesion/invasion of epithelial cells.

The protein encoded by abal is a very important autoinducer synthase of Ab that produces distinct acyl-homoserine lactone (AHL) signals that can upregulate the expression of abal through a positive feedback loop to promote the secretion of AHLs and biofilm
maturation[25]. Our study found that calcium has a similar effect on Ab-abal expression in the cellular and abiotic environments. Low concentrations of calcium (<2.4 mmol/L) may inhibit abal expression, while high concentrations of calcium tend to promote its expression. The change in Ab-abal expression with high concentrations of calcium in the cellular environment was more significant than in the abiotic environment. Moreover, it is known that calcium may promote biofilm formation. These data suggest that bacterial biofilm formation may be reduced by controlling calcium concentrations (2.2 - 2.4 mmol/L), which may reduce Ab colonization or infection capacities in the environment.

There are many reports on the interaction between epithelial cells and Acinetobacter baumannii standard strains using either ATCC 19606 and/or ATCC 17978. However, in view of the low virulence and drug resistance of Acinetobacter baumannii standard strains, clinical multidrug-resistant strains were used to investigate the effects of calcium on the interaction between clinical strains and respiratory epithelial cells. The lack of experiments with standard strains is the shortcoming of this paper, so we have provided antimicrobial susceptibility testing results and genome sequencing results about the clinical multidrug-resistant isolate to make the experimental data comparable.

Conclusions

First, with increased MOIs, Ab adhesion/invasion of human respiratory epithelial cells was gradually increased. Second, in the initial stage of infection, low concentrations of Ab have no obvious effect on cell growth, and a unique “double peak” phenomenon emerged during the process of infection. Third, calcium may promote Ab biofilm formation, and ompA has a positive effect on biofilm formation; calcium may promote the proliferation of Ab and the adherent growth of human respiratory epithelial cells; furthermore, it can increase Ab adhesion/invasion of epithelial cells by regulating the expression of Ab-related genes (ompA, bfmRS, abal). Lastly, calcium also has an impact on the interactions
between Ab and human respiratory epithelial cells, and stimulates cell growth more significantly during early infection, resulting in a sharp “single peak” phenomenon. The higher the calcium concentration is and the longer the co-culture time is, the more severe the degree of host cells infection by the bacteria is. Based on the results of these experiments and the related literature[13, 14], it can be speculated that controlling calcium concentrations may play a role in the prevention and treatment of multi-drug resistant Ab colonization or infection.

Methods

Cells and bacteria

The human respiratory epithelial cells (HPAEpiC, Cat. No. 3200) were purchased from ScienCell Research Laboratories, Inc. (San Diego, California) and maintained in RPMI 1640 medium containing 10% fetal bovine serum and 100 U/ml penicillin/streptomycin (GIBCO) at 37 °C and 5% CO₂. The use of HPAEpiC cells can better reflect the colonization or infection of Acinetobacter baumannii in human respiratory tract and is close to the environment of the body. The clinical isolate of multidrug-resistant Ab (Additional file 6: Table S3) was originally collected from the sputum of patient hospitalized in Hangzhou First People’s Hospital, China. After identification of genome sequencing (Sangon Biotech Co., Ltd., Shanghai; the SRA accession: PRJNA523637), the bacteria were inoculated to blood agar plates for 18-20 h at 37 °C, and then single colonies were picked to suspend in 0.45% sodium chloride solution for preparing bacterial suspension. The bacteria per ml was calculated based on 1 McF = 2 × 10⁸ CFU/ml.

Host-bacterial co-culture

At different MOIs

The range of bacterial MOIs was selected according to previous studies[15]. A 1 ml aliquot
of HPAEpiC (2 × 10^5 cells/ml) was seeded into six-well plates containing RPMI 1640 medium. Immediately after, 100 μl of different concentrations of bacteria (2 × 10^6, 2 × 10^7, 1 × 10^8 and 2 × 10^8 CFU/ml, corresponding to the MOIs 1, 10, 50 and 100, respectively) were inoculated into each well, and 0.45% NaCl was used as the normal control. The effects of Ab on the morphology and proliferation of epithelial cells at different MOIs and co-culture time points (2 h, 4 h, 6 h and 8 h) were observed by an IX70 bright field inverted microscopy (Olympus Optical Co., Ltd., Japan) at a magnification of 30×. Meanwhile, we identified the live and dead epithelial cells according to the LIVE-DEAD viability/cytotoxicity assay kit instructions (Life Technologies, Grand Island, NY, USA) and detected using an IX71 inverted microscopy with Nomarski optics (Olympus Optical Co., Ltd., Japan). Based on these observations, a suitable time period for the host-bacterial co-culture was selected for subsequent analyses.

The liquid in each well of the six-well plates was carefully discarded at the end of the indicated host-bacterial co-culture time period. Before digestion with trypsin solution for 1–2 min at 37 °C, the cells were washed once with phosphate buffer saline (PBS). The cells were then harvested by centrifugation (100 × g, 5 min) and washed twice with PBS.

Finally, the bacterial DNA was extracted from 1 × 10^5 cells based on procedures described by Chen et al [26]. qPCR was performed with SYBR Premix Ex Taq II (TaKaRa Bio Inc., Shiga, Japan) on an ABI 7500 real-time PCR instrument (Applied Biosystems, United States) according to the manufacturer’s instructions. Each test (50-μl volume) was performed in triplicate. Sterile distilled water served as a negative control, and bacterial DNA served as a positive control. Primer sequences for the target gene (ompA, outer membrane protein A) were as follows: 5’-CACAGATAAACACTGGTCCACG–3’ and 5’-GAATACACGACGGTTCATAGC–3’. The changes in Ab invasion of epithelial cells (including
strong adhesion) at different MOIs were analyzed by comparing the resulting Ct values. Detailed RTCA (real time cellular analysis) experimental procedures have been previously described[15, 19, 20]. Briefly, 50 μl of medium was added to 16-well E-plates (specific cell culture plates for RTCA) to obtain background readings, which were followed by the addition of 100 μl of a cell suspension (2×10^5 cells/ml). After the E-Plate was incubated at room temperature for 30 min, and 10 μl of different concentrations of bacterial suspension or 0.45% NaCl was added to the wells containing cells. The E-Plates were placed onto the reader in the incubator for continuous recording of the CI. The cells were monitored every 5 min for 72 h to obtain TCRPs. The data were collected from three multiple-well duplicates and presented as the CI normalized (CI = 1.00) to the last time point before intervention. Additionally, a suitable bacterial concentration was selected for subsequent analyses based on the above results.

**In different states of cell incubation**

Cell suspension: to observe whether *Acinetobacter baumannii* has an effect on host cell adhesion; 30–40% cell confluence: to observe the effect of *Acinetobacter baumannii* on epithelial cells without growth space restriction; 80–90% cell confluence: close to the environment of the body. Based on the above considerations, the interaction between *A. baumannii* and epithelial cells was assessed at different degrees of epithelial cell confluency.

A total of 2 ml of cells (2 × 10^5 cells/ml) were seeded into six-well plates containing RPMI 1640 medium. The three experimental groups were as follows: cell suspension (0 h), 30–40% cell confluence (after 24 h) and 80–90% cell confluence (after 48 h). In other words, 200 μl of the indicated concentration of bacteria was added to wells containing cells in different incubation states and 0.45% NaCl was used as the normal control. An inverted
microscopy was used to observe the dynamic process of the interactions between Ab and epithelial cells at different states of cell incubation (2 h, 4 h, 6 h, 8 h and 24 h).

RTCA experimental procedures were as described above. A total of 10 μl of the abovementioned bacterial suspensions, 0.45% NaCl or sterile distilled water was added to wells containing cells at different incubation states. Wells without intervention were used as the blank control. The interactions between Ab and human respiratory epithelial cells were evaluated based on the TCRP results and microscopic imaging.

Effects of calcium on host-bacterial interactions

Ab growth assays

A total of 400 μl of each of the indicated concentrations of bacteria and different concentrations (12 mmol/L, 24 mmol/L, 36 mmol/L and 48 mmol/L) of a calcium chloride solution (CaCl₂) were added to wells containing 2 ml of RPMI 1640 medium. A 400 μl aliquot of each of the bacterial suspensions and EDTA solution (5 mmol/L) was used as the normal control I (considering RPMI contains calcium, the final calcium concentrations in experimental groups were 1.4 mmol/L, 2.4 mmol/L, 3.4 mmol/L and 4.4 mmol/L, respectively, and the normal control I was 0 mmol/L). The effects of different concentrations of calcium and culture times (2 h, 4 h, 6 h, 8 h, 16 h and 24 h) on the proliferation of Ab were measured at 600 nm using a microplate reader (Sunrise, Tecan, Switzerland). Each experiment was performed in triplicate.

Experiments on the differential expression of Ab-related genes

The sequences of Ab-related genes were selected according to previous studies[12, 16, 27, 28]. Specifically, the recA (internal reference), ompA (outer membrane protein A), bfmRS (two-component regulatory system) and abal (autoinducer synthase) sequences were downloaded from GenBank. The primers (Table 2) for reverse transcription
quantitative PCR (RT-qPCR) were designed using the Primer Premiers 5.0 software, selected based on a BLAST sequence comparison, and synthesized by Life Technology (Shanghai, China). After incubation for the indicated time period described above, all bacteria were collected and total RNA was isolated from each group using an AxyPrep Miniprep Kit (Corning Inc., NY, USA) according to the manufacturer’s instructions. The concentration and purity (OD_{260}/OD_{280} of approximately 2.0) of the RNA samples were measured using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, Wilmington, USA). RT-qPCR was performed with the One Step SYBR PrimeScript PLUS RT-PCR Kit (TaKaRa) on an ABI 7500 real-time PCR instrument according to the manufacturer’s instructions. Each test (50 μl volume, 20 ng of RNA template) was performed in triplicate. RNase free distilled water served as negative control I and templates treated with RNase served as negative control II. The differential expression of Ab-related genes (ompA, bfmRS and abal) under different calcium concentrations was analyzed by comparing the resulting Ct values.

**Ab biofilm assays**

Mutant strains of Ab deficient in *ompA* (Ab-ompA^-) were constructed and modified according to the conjugative transfer and parental conjugation methods[29, 30]. Briefly, a modified pMO130-TelR with the fragments corresponding to the regions up and downstream of the *ompA* gene generated from genome of Abwas constructed. The resultant plasmid was then transformed into the *E.coli* S17-1 λ pir competent cells. Then, trans-conjugation was performed between *E.coli* S17-1 λ pir donor strain and Ab recipient strain to transfer and integrate pMO130-TelR-ompA^[Up/Down] into the chromosome of Ab. Donor and recipient strains were plated onto LB agar containing tellurite (30 mg/L) and gentamicin (25 mg/L) and incubated at 37 °C overnight. 0.45 mol/L catechol solution was
sprayed on the surface of the plate, and yellow clones were picked for PCR identification. The second selection was then performed by incubating Ab on 10% sucrose with no salt LB agar plates, overnight at 37 °C to identify sucrose-resistant sensitive clones, then analyzed by PCR and sequencing to confirm that the target gene was excised, resulting in an unmarked in-frame deletion. A single clone with no ompA sequence was saved as Ab-ompA-. Ab and Ab-ompA- isolates were added separately to 96-well plates containing RPMI 1640 medium of different calcium concentrations (range from 0 to 4.4 mmol/L). The biofilm formation was measured based on procedures outlined by Selasi et al.\cite{31} with some modifications. After 24h, the liquid in each well was carefully discarded and washed three times with PBS. Then, the plates were air-dried and stained with crystal violet (0.1% v/v) for 20 min. Finally, the plates were washed three times with PBS, air-dried and decolorized with ethanol (95% v/v) for 20 min. The turbidity was measured at 600 nm using the Tecan Sunrise microplate reader. Each experiment was performed in triplicate.

**Experiments on the morphology and proliferation of epithelial cells**

A 2 ml cell suspension (2 × 10^5 cells/ml) was seeded into six-well plates and allowed to attach and grow for 48 h to reach the platform stage (80–90% cell confluence) before the addition of a 400 μl aliquot of each of sterile distilled water and different concentrations of CaCl_2. Additionally, a 400 μl aliquot of each of sterile distilled water and EDTA was used as the normal control. The effects of different concentrations of calcium and culture times (2 h, 4 h, 6 h, 8 h and 24 h) on the morphology and proliferation of epithelial cells were observed by inverted microscopy.

RTCA experimental procedures were as described above. After incubation for 48 h, cells were treated with a 20 μl aliquot of each of sterile distilled water and different
concentrations of CaCl$_2$ or a 20 µl aliquot of each of sterile distilled water and EDTA. A well without intervention was used as a blank control. The effects of calcium on the morphology and proliferation of epithelial cells were evaluated based on the TCRP results and microscopic imaging.

**Experiments on host-bacteria interactions**

A 2 ml cell suspension ($2 \times 10^5$ cells/ml) was seeded into wells and allowed to attach and grow for 48 h before the addition of a 400 µl aliquot of each of the indicated bacterial concentrations and different concentrations of CaCl$_2$. Additionally, a 400 µl aliquot of each of the bacterial suspensions and EDTA was used as the normal control I. The effects of Ab on the morphology and proliferation of epithelial cells with different calcium concentrations and co-culture times (2 h, 4 h, 6 h, 8 h and 24 h) were observed by inverted microscopy.

According to the procedures described above, total bacterial RNA was isolated from each group and RT-qPCR was performed after host cells and bacteria were co-cultured for the indicated time period. The expression of Ab-related genes in the biotic environment under different calcium concentrations was analyzed by comparing the resulting Ct values. In addition, a 400 µl aliquot of each of sterile distilled water and 0.45% NaCl was used as the normal control II. All the liquid in the wells was carefully discarded after a 2 h, 4 h or 6 h incubation, as described above. Before digestion, the cells were washed three times with PBS. Other procedures, including bacterial DNA extraction and qPCR were carried out as described above. The change in Ab invasion of epithelial cells (including strong adhesion) under different calcium concentrations and co-culture times was analyzed by comparing the resulting Ct values.

RTCA experimental procedures were as described above. After the cells (20,000 cells per
well) were incubated for 48 h, the two part experiment was carried out as follows:

1. A 20 μl aliquot of each of the abovementioned bacterial suspensions and different concentrations of CaCl2 was added to the wells; the bacterial suspensions and EDTA was used as the normal control I; sterile distilled water and 0.45% NaCl was used as the normal control II.

2. A 20 μl aliquot of each of different concentrations of CaCl2 and the bacterial suspension/0.45% NaCl/sterile distilled water was added to the wells; a total of 20 μl of sterile distilled water was added as the normal control. Wells without intervention were used as the blank control.

The effects of calcium on the host-bacterial interactions were evaluated based on the TCRP results, qPCR and microscopic imaging.

Statistical analysis

The data are presented as the means ± SD. One-way analysis of variance (ANOVA) and the Student-Newman-Keuls (SNK) test were applied to compare and analyse the changes in Ct values at different MOIs and the effects of calcium on the expression of Ab-related genes. The effects of different calcium concentrations and culture times on the CI of epithelial cells were analysed using multivariate ANOVA with repeated measures and the SNK test. Univariate ANOVA with repeated measures and the SNK test were used to compare and analyse the effects of different calcium concentrations and co-culture times on Ab invasion (including strong adhesion) of epithelial cells (Ct values). \( P < 0.05 \) was considered to be statistically significant.

Abbreviations

Ab, Acinetobacter baumanii; RTCA, real time cellular analysis; TCRP, time-dependent cell response profile; MOI, multiplicity of infection; CI, cell index; qPCR, quantitative PCR; RT-
qPCR, reverse transcription quantitative PCR; Ct, cycle threshold; ANOVA, analysis of variance; SNK, Student-Newman-Keuls; AHL, N-acyl homoserine lactone.

Declarations

Ethics approval and consent to participate: This study was approved by the Ethics Review Committee of Hangzhou First People’s Hospital, China. With permission from the patient, the clinical isolate of multidrug-resistant Ab was collected from his sputum.

Consent for publication: Not applicable.

Availability of data and materials: All data analysed during this study are included in this published article and its supplementary information files.

Competing interests: The authors declare that they have no competing interests.

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Authors’ contributions: Conceptualization, D. Y.; Methodology, D. Y. and Y. C.; Formal Analysis, Y. C., T. S., and S. F.; Investigation, Y. C., T. S., P. P., S. F., J. J., T. C. and D. Y.; Writing—Original Draft, Y. C. and D. Y.; Writing—Review & Editing, H. W. and S. F.; Project Administration, D. Y.; Funding Acquisition, D. Y. and S. F. All authors have read and approved the final manuscript.

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Tables

Table 1 Microscopy observations of the effects of Ab on epithelial cells at different MOIs
Co-culture time point | Control group | Experimental group |
|---------------------|---------------|---------------------|

| MOI 1 | MOI 10 | MOI 50 |
|-------|--------|--------|
| 2 h   | + / -  | + (-) / ±(-) | + (-) / ++ (-) | + (+) / & (+) |
| 4 h   | ++ / - | ++ (-) / + (-) | ++ (±) / +++ (-) | ++ (+++) / & & |
| 6 h   | +++ / -| +++ (-) / ++ (-) | +++ (+) / ++++ (-) | +++ (+++) / & & |
| 8 h   | ++++ / -| ++++ (-) / ++ (-) | ++++ (+) / ++++ (±) | +++ (+++) / & & & |

Control: 0.45% NaCl.
+: indicated the degree (the epithelial cells were expressed as the situation of adherent growth, bacteria were expressed as quantity, bubble-like dead cells and bacterial aggregation phenomenon were expressed as visibility); &: indicated the degree > ++++; - : indicated none.
For example: + / - indicated that a few of epithelial cells adhered to plates without bacteria; +++ (+++) / & & (++++) indicated that the bacterial amount increased and the bacterial aggregation phenomenon became more obvious compared to 4 h, a lot of epithelial cells adhered to plates and the bubble-like dead cells with bacterial adhesion visible.

Table 2 Primers for RT-qPCR
| Gene ID #   | Primers       | Sequences (5′–3′)                                           | Product length (bp) |
|------------|---------------|------------------------------------------------------------|--------------------|
| AF251469.1 | recA-F recA-R | ACGCCCTAGACCCTCAATAT AGAGTCACCCATCTCACCTTC                 | 197                |
| AY485227.1 | ompA-F ompA-R | CACAGATAACACTGGTCCACG GAATACACGACGGTTCATAGC                | 190                |
| AY838282.1 | bfmRS-F bfmRS-R | AACAAAGTTCCGGATTACGGG TCACTAAACGGGCAAGG                    | 128                |
| EU334497.1 | abal-F abal-R | CTATTCCCTGCTCACCAGA CCCGCAGCAGTAATAAAC                      | 208                |

#: GenBank.

Figures
Analysis of optimum MOI of Ab to human respiratory epithelial cells. Control: 0.45% NaCl. (A) Microscopy observations of the effects of Ab on epithelial cells (2 × 10^5 cells per well) at different MOIs (co-cultured for 4h). Scale bar = 30 μm.
Red arrows: bubble-like (denucleated) dead cells with bacterial adhesion visible around (especially in the MOI 50 and MOI 100 experimental groups); Yellow arrows: the phenomenon of bacterial aggregation. (B) The qPCR results indicating Ab invasion of epithelial cells (including strong adhesion) at different MOIs (host cells and bacteria were co-cultured for 4h and 1 × 105 cells were isolated). The differences in the Ct values (MOI 100 group < MOI 50 group < MOI 10 group < MOI 1 group) among all the groups were statistically significant, as determined by the SNK test (P < 0.05). The differences between the MOI 1 and MOI 10 groups and the MOI 10 and MOI 50 groups were greater than between the MOI 50 and MOI 100 groups. The higher the MOI was, the greater the Ab invasion of epithelial cells. (C) Ab infection TCRPs of epithelial cells (20,000 cells per well) at different MOIs. t: point at which bacteria or NaCl were added (after background readings for the E-Plate were obtained and the plate was incubated at room temperature for 30 min). The "double peak" phenomenon emerged during Ab infection. The smaller the MOI was, the later and higher the peak CI was, the more significant the phenomenon was. Representative curves are an average of three replicate wells.
The effects of Ab on human respiratory epithelial cells in different incubation states. (A) Microscopy observation of the effects of Ab on epithelial cells (4 × 105 cells per well) in different incubation states. Scale bar = 30 μm. Experimental group: addition of bacterial suspension; Control group: addition of 0.45% NaCl; Red arrows: bubble-like (denucleated) dead cells; Yellow arrows: the phenomenon of bacterial aggregation; Blue arrows: massive cell death. (B) Ab infection TCRPs of epithelial cells (20,000 cells per well) in different incubation states. I: addition of bacterial suspension (red arrow) at cell suspension (0 h); II: addition of bacterial suspension (green arrow) at 30-40% cell confluence (after 24 h); III: addition of bacterial suspension (blue arrow) at 80-90% cell confluence (after 48 h); IV: blank control (without intervention); t: treatment time point; t1: cell suspension (0 h); t2: 30-40% cell confluence (after 24 h); t3: 80-90% cell confluence (after 48 h). Ab infection caused the CI to rise and fall. A unique "double peak" phenomenon emerged during Ab infection. The addition of 0.45%
NaCl or sterile distilled water had no significant effect on the Cl.
Calcium supplementation final concentration:

- 1.4 mmol/L
- 3.4 mmol/L
- 2.4 mmol/L
- 4.4 mmol/L

Figure 3

The effects of calcium on Ab proliferation (growth curves). Control I: calcium final concentration was 0 mmol/L (with EDTA treatment). Calcium can promote the proliferation of Ab.
Figure 4

Calcium supplementation final concentration:
- 1.4 mmol/L
- 2.4 mmol/L
- 3.4 mmol/L
- 4.4 mmol/L
qPCR results of calcium effects on host-bacterial interactions. Ab and human respiratory epithelial cells were co-cultured in calcium-supplemented medium for 2 h, 4 h and 6 h, and $1 \times 10^5$ cells were isolated for qPCR detection. Control I: calcium final concentration was 0 mmol/L (with EDTA treatment). Calcium had positive effects on the interaction between Ab and epithelial cells. With increased calcium concentrations and prolonged co-culture times, the amount of Ab invasion into epithelial cells increased (the smaller the Ct value, the more bacterial invasion into cells).
Figure 5

The effects of calcium on Ab infection TCRPs of human respiratory epithelial cells. A total of 20,000 cells per well were seeded into E-plates. t: treatment time point (48 h, 80-90% cell confluence). With increased calcium concentrations (bacteria-infected cells), the CI increased or decreased faster and reached a higher peak value. A sharp "single peak" phenomenon occurred during the infection.
Effects of calcium on the expression of Ab-related genes. The calcium supplementation final concentrations were as follows: Group a, 1.4 mmol/L; Group b, 2.4 mmol/L; Group c, 3.4 mmol/L; Group d, 4.4 mmol/L; Group control I: calcium final concentration was 0 mmol/L (with EDTA treatment); Abiotic environment: bacteria only were cultured in plates containing RPMI 1640 medium; Cellular environment: bacteria were cultured in epithelial cell covered plates (80-90% cell confluence) containing RPMI 1640 medium. The recA gene was used as an internal reference. Relative changes in expression level of Ab-related genes between the experimental groups and control I group were calculated by the 2-\(\Delta\Delta Ct\) method.

Supplementary Files

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