Deletion of a Gene Encoding a Putative Peptidoglycan-Associated Lipoprotein Prevents Degradation of the Crystalline Region of Cellulose in *Cytophaga hutchinsonii*

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**INTRODUCTION**

Cell envelopes of Gram-negative bacteria consist of an outer membrane, an inner membrane, and a thin rigid layer of peptidoglycan located in the periplasm (Malanovic and Lohner, 2016). In order to maintain the integrity and stability of cell envelope, peptidoglycan interacts with many outer membrane and inner membrane proteins, one of which is the peptidoglycan-associated lipoprotein (Pal). Chapman et al. (1980) have shown that Pal is essential for the growth of *Cytophaga hutchinsonii* and its deletion results in the loss of the ability to degrade cellulose. The mechanism of cellulose degradation by *Cytophaga hutchinsonii* is still unclear. In this study, we deleted the gene *chu_0125*, encoding a homolog of Pal, and found that the deletion mutant prevented degradation of the crystalline region of cellulose.

**RESULTS**

**Growth and Crystallinity Analysis:**

Growth of the deletion mutant in cellulose medium was slower than that of the wild type. In addition, the degree of crystallinity of Avicel was increased in the deletion mutant compared to the wild type. These results indicate that *chu_0125* deletion mutant has a defect in growth on crystalline cellulose and affects the crystallinity of cellulose.

**Enzyme Activities:**

The endoglucanase and β-glucosidase activities on the cell surface were reduced by 60% and 30%, respectively, without the *chu_0125* gene. Moreover, compared to the wild type, the deletion mutant was more sensitive to some harmful compounds and released sixfold more outer membrane vesicles (OMVs) whose protein varieties were dramatically increased. These results indicated that *chu_0125* deletion affected the integrity of the outer membrane.

**Protein Stability:**

Western blotting revealed that CHU_3220, the only outer membrane protein necessary and specialized for degradation of the crystalline region of cellulose, was largely leaked from the outer membrane and packaged into OMVs. These results suggest that *chu_0125* deletion affected the integrity of the outer membrane and thus influenced the localization of some outer membrane proteins, including CHU_3220.

**DISCUSSION**

The deletion of *chu_0125* affects the integrity of the outer membrane and thus changes the localization of some outer membrane proteins. This might be the reason why deletion of *chu_0125* prevents degradation of the crystalline region of cellulose.

**Keywords:** *Cytophaga hutchinsonii*, peptidoglycan-associated lipoprotein, cellulose degradation, crystalline region, integrity of the outer membrane
lipoprotein (Pal) (Parsons et al., 2006; Godlewksa et al., 2009). Pal, widely distributed in Gram-negative bacteria, interacts with peptidoglycan through the C-terminal region and links to the outer membrane through the N-terminal lipid (Lazzaroni and Portalier, 1992). Pal is also an important component of the Tol-Pal complex consisting of an inner membrane subcomplex TolA-TolQ-TolR and an outer membrane associated subcomplex TolB-Pal (Lloubes et al., 2001). The Tol-Pal complex is considered to be involved in maintaining the proper structure and function of the cell envelope (Lloubes et al., 2001). Mutation in pal causes a defect in the integrity of the outer membrane, resulting in hypersensitivity to harmful compounds, leakage of outer membrane and periplasmic proteins, and formation of abundant outer membrane vesicles (OMVs) (Bernadac et al., 1998; Kowata et al., 2016). OMVs are released from the outer membrane into the extracellular milieu, carrying outer membrane proteins, periplasmic proteins, lipopolysaccharides, and phospholipids (Schwechheimer and Kuehn, 2015).

Cellulose, which is composed of the crystalline region and the amorphous region, is the most abundant biological resource on earth (Bayer and Lamed, 1992). To degrade cellulose, some aerobic bacteria secrete free cellulases to the extracellular milieu and some anaerobic bacteria produce cellulases (Lynd et al., 2002; Bayer et al., 2004). Cytophaga hutchinsonii is a widely distributed aerobic Gram-negative bacterium from the phylum Bacteroidetes, which exhibits gliding motility over surfaces and digests cellulose rapidly (Stanier, 1942; Xie et al., 2007). The strategy for C. hutchinsonii to digest cellulose is different from the free-cellulase and the cellulosome mechanisms (Zhu and McBride, 2017). C. hutchinsonii appears to use a contact-dependent digestion strategy and most of the cellulase activities are cell surface associated, indicating the importance of outer membrane in degrading cellulose (Wilson, 2009; Ji et al., 2013). However, the mechanism of cellulose degradation by C. hutchinsonii is still poorly understood. Several outer membrane proteins, including CHU_1276, CHU_1277, and CHU_0170, were demonstrated to be indispensable to cellulose utilization (Ji et al., 2014; Zhu and McBride, 2014; Zhou et al., 2016). Recently, Wang et al. (2017) reported that the outer membrane protein CHU_3220 was necessary for the degradation of the crystalline region but not for the amorphous region of cellulose. These studies further indicated the importance of outer membrane proteins in cellulose utilization. However, other factors that may be related to cellulose degradation need to be explored.

As an important component in maintaining the outer membrane integrity, peptidoglycan-associated lipoprotein might influence the location and function of some outer membrane proteins, so it might play a role in cellulose degradation by C. hutchinsonii. In this study, chu_0125, encoding a putative peptidoglycan-associated lipoprotein, was deleted. The outer membrane integrity of C. hutchinsonii was evaluated by examining its sensitivity to some harmful compounds, the quantification of OMVs and the OMV proteins. The cellulose utilization ability of the Δ0125 mutant was studied and the results showed that deletion of chu_0125 prevented degradation of the crystalline region of cellulose.

**Materials and Methods**

**Bacterial Strains, Plasmids, and Growth Conditions**

Cytophaga hutchinsonii ATCC 33406 was grown in PY6K medium (6 g/L peptone, 0.5 g/L yeast extract, 1 g/L KNO₃, 4 g/L glucose, pH 7.3), modified from PY6 medium (6 g/L peptone, 0.5 g/L yeast extract, 4 g/L glucose, pH 7.3) (Xu et al., 2012). To analyze cellulase activity, C. hutchinsonii was cultured in Stainer medium (1 g/L KNO₃, 1 g/L K₃HPO₄, 0.2 g/L MgSO₄·7H₂O, 0.02 g/L FeCl₃·6H₂O, 0.1 g/L CaCl₂, pH 7.3) (Stanier, 1942) plus 2 g/L glucose. Escherichia coli strains were grown in LB medium (10 g/L tryptone, 5 g/L yeast extract, 10 g/L NaCl, pH 7.0) at 37°C with shaking at 170 rpm. Antibiotics were used at the following concentrations: ampicillin (Ap), 100 µg/mL; erythromycin (Em), 30 µg/mL; chloramphenicol (Cm), 15 µg/mL. The strains and plasmids used in this study are listed in Table 1. Primers used in this study are listed in Supplementary Table S1.

**Construction of the chu_0125 Deletion Mutant**

Deletion of chu_0125 (yielding the Δ0125 deletion mutant) was performed as previously reported (Wang et al., 2014). Briefly, a 2.4-kbp fragment spanning the three flanking genes (chu_0122, chu_0123, and chu_0124) and the first 175 bp of chu_0125 was amplified with primers 0125H1F and 0125H1R. The fragment referred as H1 was digested with BamH1 and Kpn1 and cloned into pTSK digested with the same enzymes. A 2.0-kbp fragment spanning the two flanking genes (chu_0126 and chu_0127) and the last 198 bp of chu_0125 was amplified with primers 0125H2F and 0125H2R. The fragment referred to as H2 was digested with SalI and SacI and ligated into corresponding sites of pTSK, which flanked the ermF-FRT cassette opposite H1. The gene-targeting cassette was amplified with primers H1F and H2R, purified with a Cycle Pure kit (Omega, GA, United States) and transformed into 100 µL of competent cells of C. hutchinsonii by electroporation. The transformants were verified by PCR with two sets of primers, 0125UF/0125DR and 0125UF/0125UR. Then the correct transformants were used as parent strain for transforming the helper plasmid pCHF by electroporation to get the unmarked mutant. After incubation for about 15 days, the transformants were cultured without antibiotics to lose the pCHF. Then the cells were verified by PCR with primers 0125UF and 0125DR and the PCR products were sequenced to verify the scar sequence.

**Complementation of the Δ0125 Mutant**

The complementation of the Δ0125 mutant was performed as described previously (Wang et al., 2014). Briefly, a 2.2-kbp fragment spanning chu_0125, 187 bp upstream of the start codon and 96 bp downstream of the stop codon, was amplified with primers 0125CF and 0125CR. The fragment was digested and ligated into the pCH plasmid to generate pCH0125. Then the plasmid was transformed into 100 µL of competent cells of the Δ0125 mutant by electroporation. C0125 refers to the complemented strain of the Δ0125 mutant with pCH0125.
Filter Paper Degradation Assay
Filter paper degradation assays were carried out as previously described (Ji et al., 2012; Wang et al., 2014). Briefly, equivalent amounts of middle exponential phase cells from Stainer medium were spotted on Whatman NO. 1 filter paper overlaid on Stanier medium supplemented with 10 g/L agar. The plates were incubated at 30°C to observe the degradation.

Growth Analysis With Different Carbon Sources
When glucose was used as the carbon source, the growth curves were monitored by Bioscreen C analyzer (Oy growth curves Ab Ltd, Finland) (Bai et al., 2017a). Briefly, C. hutchinsonii strains were grown in Stainer medium to middle exponential phase and inoculated into 200 µL of Stanier, PY6, and PY6K medium in a sample plate. The plate was incubated at 30°C with shaking at medium speed, and the growth curves were monitored by the optical density at 600 nm every 3 h. When Avicel PH101 and cultured at 30°C with shaking at 160 rpm. Samples were taken at set intervals (0.5, 24, and 48 h). All the samples were centrifuged, resuspended in 0.2 M NaOH, boiled for 10 min, and centrifuged at 13,500 × g for 10 min to remove the supernatant. The pellets were then washed twice with distilled water and dried at 60°C overnight. The XRD of samples was observed with a D8 ADVANCE System Diffractometer (Bruker, Germany). The XRD crystallinity index ($C_{R,D}$) was calculated as described by Wang et al.: $C_{R,D} (%) = (I_{002} - I_{am})/I_{002} × 100$. $I_{002}$ is the height of the crystalline peak at 22° and $I_{am}$ is the intensity of the peak at 18° (Wang et al., 2017).

Cellulase Activity Assay
Cellulase activity assays were carried out as previously described (Bai et al., 2017a; Wang et al., 2017). Briefly, cells were grown in Stanier medium and harvested at different growth phase. For intact cell samples, cell pellets were resuspended in Na2HPO4-KH2PO4 buffer (100 mM, pH 6.8). For cell-extract samples, cell pellets were resuspended in Na2HPO4-KH2PO4 buffer with 2% (vol/vol) Triton X-100, and then the cells were incubated at 4°C for 5 h. The protein concentration was quantified as described by Bradford (Bradford, 1976). 1% (wt/vol) sodium carboxymethyl cellulose (CMC-Na) and 2 mM β-Nitrophenyl β-D-glucopyranoside (pNPG) were used as substrates to measure carboxymethyl cellulase (CMCase) activity and β-glucosidase activity, respectively. Cellulase activity of the intact cells was the cellulase activity of the cell surface. Intracellular cellulase activity was equal to cellulase activity of the cell extracts minus that of the intact cells. All the measurements were carried out in triplicate.

Disk Diffusion Susceptibility Test
Disk diffusion susceptibility test was performed as described by Bai et al. (2017b). C. hutchinsonii was grown in PY6K medium.
to middle exponential phase and spread evenly over PY6K agar plates. A 6 mm paper disk was placed on the center of the plate and 3 µL of reagents was added, followed by incubation at 30°C for 3 days. The inhibition zone diameters were determined by the average of three plates. The reagents for the test were sodium dodecyl sulfate (10%), ampicillin (100 µg/mL), dithiothreitol (2 mM), H₂O₂ (2%), and crystal violet (1%).

Observation of OMVs by Scanning Electron Microscopy (SEM)
The samples of OMVs were prepared as previously described with some modification (Nur et al., 2013). The strains were grown in PY6K medium in a 24-well culture plate with coverslips on the bottom. After incubation for 4 days, the coverslips were removed and washed twice with distilled water, and then fixed with 2.5% glutaraldehyde at 4°C overnight. All the samples were dehydrated through a graded series of ethylalcohol and dried by a critical point dryer (EM CPD300, Leica, Germany) for SEM (Quanta 250 FEG, FEI, United States).

Quantification of OMVs
The quantification of OMVs was measured as previously described (Stewart, 1980; Wessel et al., 2013; Gugrati et al., 2014). Briefly, cells were grown in PY6K medium to middle exponential phase and harvested by centrifugation at 5,100 × g for 10 min at 4°C. Then the supernatants were filtered through a 0.45 μm-pore-size PVDF filter (Sangon Biotech, Shanghai, China) to remove all the cells. Cell-free supernatants were then centrifuged at 200,000 × g for 1 h to get vesicle pellets, followed by washing twice with PIPES (50 mM). Cell-free supernatants from which OMVs had been removed were labeled as OMV-free supernatants. For quantification, vesicle pellets were resuspended in MV buffer (50 mM Tris, 5 mM NaCl, 1 mM MgSO₄·7H₂O, pH 7.4) and extracted 1:1 with chloroform. The organic layers were removed into a new tube, combined with an equal volume of ammonium ferrothiocyanate and phospholipids complex at pH 7.4) and extracted 1:1 with chloroform. The absorbance of ammonium ferrothiocyanate and phospholipids complex at 470 nm was analyzed.

Preparation of Outer Membrane Proteins
Outer membrane proteins were prepared as described by Ji et al. (2014) and Wang et al. (2017). Briefly, the strains were grown in PY6K medium to middle exponential phase and harvested by centrifugation at 5,100 × g for 10 min at 4°C. Then the pellets were washed with PIPES (50 mM), resuspended in PIPES (50 mM) with 0.5 M NaCl, and incubated at 4°C for 20 min with shaking at 150 rpm. The cells were removed by centrifugation at 13,500 × g for 20 min at 4°C, and the supernatant containing the buffer-washed proteins was ultracentrifuged at 100,000 × g for 30 min at 4°C. Then the pellets were resuspended in PIPES (50 mM) as the outer membrane proteins, and the outer membrane proteins were subjected to SDS-PAGE. The weakened and missing protein bands in the profile of the ΔO125 mutant compared with that of the wild type were cut and analyzed by matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry.

Detection and Localization of CHU_3220
Outer membrane proteins were prepared as described above. OMVs were isolated as described above and resuspended in PIPES (50 mM) to obtain the OMV proteins. Extracellular proteins were prepared as described previously (Nandakumar et al., 2006). Briefly, cells were removed by centrifuging at 5,100 × g for 10 min. The supernatants were collected and filtered through a 0.22 μm-pore-size PVDF filter (Sangon Biotech, Shanghai, China). Cell-free supernatants and OMV-free supernatants were precipitated by treating with 10% (vol/vol) TCA (trichloracetic acid) in ice for 30 min. Then the precipitates were collected by centrifugation at 13,500 × g for 10 min at 4°C and washed three times with pre-cooled acetone to remove traces of TCA. Finally, the precipitated proteins were solubilized in PIPES (50 mM). The localization of CHU_3220 was detected by Western blotting analysis using the CHU_3220 antibody as described by Wang et al. (2017).

Bioinformatics
The genome sequence of C. hutchinsonii was obtained from the NCBI database1. CLUSTALW2 was used for the multiple alignment of related Pals. T-COFFEE3 and MUSCLE4 were applied to verify the CLUSTALW results. BoxShade5 was a tool to shade multiple alignment files.

RESULTS
Bioinformatic Analysis of CHU_0125
CHU_0125 was annotated as an outer membrane peptidoglycan-associated lipoprotein (Pal) in NCBI. Generally, Pal homologs, such as in E. coli, Haemophilus influenza, Actinobacillus pleuropneumoniae, Klebsiella pneumoniae, and Alkalomonas amylolytica only have an OmpA_C-like domain which plays a role in associating with peptidoglycan (Figure 1A) (Chen and Henning, 1987; Zlotnick et al., 1988; Frey et al., 1996; Hsieh et al., 2013; Zhai et al., 2014). The full-length of chu_0125 (GenBank accession number: ABG57418) is 1971 bp and encodes a protein of 656 amino acids. NCBI shows that in addition to the OmpA_C-like domain, CHU_0125 contains two other domains: TPR (tetratricopeptide repeats) and PD40. TPR serves as an interaction module and multiprotein complex mediator (Zeytuni and Zarivach, 2012). PD40 is a WD40-like domain responsible for regulating cellular function such as cell division, cell-fate determination, and mRNA modification (Neer et al., 1994). Sequence alignment revealed that OmpA_C-like domain

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1 https://www.ncbi.nlm.nih.gov/
2 http://www.ebi.ac.uk/Tools/msa/clustalw/
3 http://tcoffee.vital-it.ch/apps/tcoffee/index.html
4 http://www.ebi.ac.uk/Tools/msa/muscle/
5 http://www.ch.embnet.org/software/BOX_form.html
of CHU_0125 had 40% identity value with Pal from E. coli (EcPal) (Figure 1B), but CHU_0125 only had 15% coverage value with EcPal because of the TPR and PD40 domains (Figure 1A).

Growth Properties of the Δ0125 Mutant With Glucose as the Carbon Source
To study the function of chu_0125 in C. hutchinsonii, we deleted chu_0125 and obtained the Δ0125 mutant (Supplementary Figure S1). Then we complemented the Δ0125 mutant with pCH0125 and got the C0125. The growth properties of the wild type, Δ0125 mutant, and C0125 in PY6 and Stainer medium were tested. In Stainer medium, the growth curve of the Δ0125 mutant was similar to that of the wild type and the C0125 (Figure 2A). However, in PY6 medium, it was found that the Δ0125 mutant exhibited a long lag period and a reduced biomass at the stationary phase compared with the wild type and the C0125 (Figure 2B). Then we cultured the strains in PY6 medium supplemented with one kind of the inorganic salts of Stainer medium and found that when PY6 medium was supplemented with 1 g/L KNO₃, the Δ0125 mutant could grow as well as the wild type (Figure 2C), while the other four kinds of inorganic salts of Stainer had almost no effect on the growth of the Δ0125 mutant (Supplementary Figure S2). To further determine whether K⁺ or NO₃⁻ played a role in improving the growth of Δ0125 mutant, KCl and NaNO₃ were used as the providers of K⁺ and NO₃⁻, respectively. The growth curves showed that KCl had no effect on the growth of the Δ0125 mutant (Figure 2E) while NaNO₃ was beneficial to the growth of the Δ0125 mutant (Figure 2D), suggesting that NO₃⁻ played a role in improving the growth of the Δ0125 mutant. To eliminate the effect of Na⁺, the growth curves of strains in PY6 medium supplemented with 1 g/L NaCl were monitored, indicating that Na⁺ had no effect on the growth of C. hutchinsonii (Figure 2F). All these results suggested that the Δ0125 mutant could not grow as well as the wild type in PY6 medium and additional NO₃⁻ was important for the growth of the Δ0125 mutant. In the following tests, we used PY6 medium supplemented with 1 g/L KNO₃ called PY6K medium to replace the PY6 medium. Compared with PY6 medium, in Stainer and PY6K medium the strains did not have a long stationary phase, which was observed before in our laboratory. However, the reason for this unusual behavior was still unknown.

Deletion of chu_0125 Prevents Degradation of the Crystalline Region of Cellulose
In order to study the effect of chu_0125 on cellulose degradation, the ability of C. hutchinsonii to digest different kinds of cellulose was monitored. As shown in Figure 3A, the wild type and the C0125 could degrade filter paper while the Δ0125 mutant was deficient in filter paper degradation. We also measured the growth curves of the wild type, Δ0125 mutant, and C0125 when Avicel or RAC was used as the sole carbon source. As shown in Figure 3B, the growth rate of the Δ0125 mutant was lower than that of wild type and C0125 with Avicel as the sole carbon source while the growth rate of the Δ0125 mutant was similar to that of wild type and C0125 with RAC as the sole carbon source (Figure 3C). When C. hutchinsonii was cultured in Avicel, quantification of the residual Avicel showed that the wild type utilized 81% of the Avicel after incubation for 51 h while the Δ0125 mutant utilized only 12% of the Avicel in the same time (Figure 3D). These results suggested that the Δ0125 mutant was able to digest RAC as well as the wild type but it was defective in the degradation of cellulose.

In addition, the degree of crystallinity of Avicel which was incubated with C. hutchinsonii was tested by XRD. After incubation for 48 h, the crystallinity of Avicel was reduced from 65.1 to 57.4% and 58.5% by the wild type and C0125 respectively (Figures 4A,C), while that of the Δ0125 mutant increased to 70.7% (Figure 4B). The result of the increase in crystallinity of Avicel by the Δ0125 mutant suggested that the Δ0125 mutant mainly utilized the amorphous region of cellulose when cultured in Avicel. Furthermore, we observed the arrangement of cells on filter paper by scanning electron microscopy (SEM). As shown Supplementary Figure S3A, cells of the wild type grew and arranged on the surface of the fibers, while that of the Δ0125 mutant grew and gathered in the gully of fibers (Supplementary Figure S3B). Surface morphology of Avicel was also investigated by SEM. The surface of Avicel was smooth and flat after incubated with the wild type (Supplementary Figure S3C) while that of the Δ0125 mutant was gully shaped (Supplementary Figure S3D). Given our previous finding that the Δ0125 mutant could mainly degrade the amorphous region of cellulose and was defect in degradation of crystalline region of cellulose, we deduced that the gully shaped surface might be caused by the selective degradation of cellulose by the Δ0125 mutant.

Cellulase Activity Determination
To study the influence of chu_0125 on cellulase activity, endoglucanase and β-glucosidase activities of C. hutchinsonii were examined. As shown in Figures 5B,D, intracellular cellulase activities of the Δ0125 mutant were almost as same as that of the wild type. But the endoglucanase activity of intact cells of the Δ0125 mutant was only 40% (the average of three growth phase) of that of the wild type (Figure 5A) and β-glucosidase activity of the Δ0125 mutant was 70% of that of the wild type (Figure 5C), indicating that deletion of chu_0125 affected both endoglucanase and β-glucosidase activities on the cell surface.

Defect in the Integrity of the Outer Membrane
Studies showed that deletion of pal impaired the integrity of outer membrane. To study the function of chu_0125 in C. hutchinsonii, susceptibility of the wild type, Δ0125 mutant, and C0125 to some harmful compounds, including sodium dodecyl sulfate, ampicillin, dithiothreitol, H₂O₂, and crystal violet, was tested. As shown in Table 2, the Δ0125 mutant was more sensitive to all these reagents than the wild type, suggesting that the outer membrane permeability of the Δ0125 mutant had been impaired.

The OMVs of the wild type and the Δ0125 mutant were also observed by SEM. As shown in Figure 6A, both the wild
type and the Δ0125 mutant were able to secret OMVs. But the surface of the Δ0125 mutant was rougher and more OMVs were observed compared with the wild type. Moreover, we quantified the productions of OMVs and found that the OMV production of the wild type was only about 15% of that of the Δ0125 mutant (Figure 6B). We also analyzed the cargo proteins of OMVs by SDS-PAGE. OMVs of the Δ0125 mutant contained far more kinds of proteins than those of the wild type (Figure 6C). Considering the difference of the outer membrane permeability, OMV productions, and OMV proteins between
wild type and Δ0125 mutant, we deduced that the outer membrane integrity of the Δ0125 mutant was impaired.

Analysis of Outer Membrane Proteins

Previous studies showed that defects in outer membrane integrity also resulted in release of outer membrane proteins (Bernadac et al., 1998; Llamas et al., 2000). To determine the changes of the Δ0125 mutant in outer membrane proteins, we prepared the outer membrane fractions and examined by SDS-PAGE. As shown in Figure 7, compared with the wild type, several bands of outer membrane proteins were weakened or missing in the protein profile of the Δ0125 mutant, which...
TABLE 2 | Inhibition zone diameters of wild type and Δ0125 mutant.

| Reagent                | Inhibition zone diameter (mm) | WT     | Δ0125  | C0125 |
|------------------------|------------------------------|--------|--------|-------|
| Sodium dodecyl sulfate | 24.6 ± 0.9                   | 30.4 ± 0.4* | 27.6 ± 0.5 |
| Ampicillin             | 44.2 ± 0.1                   | 53.7 ± 0.2* | 48.3 ± 0.5 |
| Crystal violet         | 25.0 ± 0.8                   | 30.0 ± 0.8* | 27.6 ± 0.9 |
| Dithiothreitol         | 25.6 ± 0.4                   | 32.0 ± 0.6* | 25.3 ± 1.2 |
| H2O2                   | 37.2 ± 0.8                   | 45.7 ± 0.8* | 37.7 ± 1.2 |

*The inhibition zone diameters were the average of three triplicates ± standard deviation. Statistical analysis was carried out using the Student’s t-test analysis. Significant differences (T-test *P < 0.05, **P < 0.01) were detected between WT and Δ0125 mutant.

were identified by MALDI-TOF mass spectrometry and listed in Table 3. One of the weakened outer membrane proteins was CHU_3220, which was reported to be necessary for the degradation of the crystalline region of cellulose (Wang et al., 2017). Zhu et al. (2013, 2016) reported that disruption of chu_1107 caused no effect on cellulose degradation. The other three genes, chu_1075, chu_3437, and chu_0522, were singly deleted and it was found that these mutants were able to degrade cellulose as well as the wild type (Supplementary Figure S4). Arrow 5 was identified as CHU_0125, suggesting that CHU_0125 was located on the outer membrane. The disappearance of the band of CHU_0125 in the Δ0125 mutant also confirmed that chu_0125 was successfully deleted in the Δ0125 mutant.

Deletion of chu_0125 Affected the Localization of CHU_3220

CHU_3220 is a large protein located on the outer membrane and it is the only reported protein to be necessary and specialized for the degradation of the crystalline region of cellulose (Wang et al., 2017). Given the reduced amount of CHU_3220 in the outer membrane of the Δ0125 mutant as described above, we tested the localization of CHU_3220 in the Δ0125 mutant. The outer membrane fractions and extracellular fractions were prepared for Western blotting. As shown in Figure 8, CHU_3220 could be detected both in the outer membrane fractions of the wild type and the Δ0125 mutant. However, the amount of CHU_3220 in the outer membrane of the Δ0125 mutant was about 25% of the amount of wild type as analyzed by Gel-Pro analyzer. In extracellular fractions, the band of CHU_3220 was obvious in the Δ0125 mutant but it was not observed in the wild type. The results suggested that deletion of chu_0125 led to CHU_3220 being largely leaked from outer membrane into extracellular milieu. To detect the exact location of CHU_3220 in the extracellular fractions, we prepared OMV fractions and OMV-free extracellular fractions.

FIGURE 6 | OMVs of wild type (WT) and Δ0125 mutant (Δ0125). Observation of OMVs by SEM (A). Quantification of OMVs (B). SDS-PAGE of OMV proteins (C).

The loading samples were normalized by calibration of cell biomass. Quantification of OMVs was carried out in triplicate, and error bars indicate standard deviations from these replicates. Significant differences (T-test *P < 0.05, **P < 0.01) were detected between WT and Δ0125 mutant.
In the Δ0125 mutant, CHU_3220 was detectable in OMV fractions and undetectable in OMV-free extracellular fractions which implied that CHU_3220 leaked from outer membrane and was packed into OMVs. Analysis of the OMV proteins by LC-MS/MS found that the unique peptides of CHU_3220 could be detected in the Δ0125 mutant while they could not be detected in that of the wild type, which was in accordance with the result of Western blotting. All these suggested that in the wild type, CHU_3220 was located on the outer membrane, while in the Δ0125 mutant a lot of CHU_3220 leaked from outer membrane and was packed into OMVs.

**DISCUSSION**

*Cytophaga hutchinsonii* is a widely distributed gliding cellulolytic bacterium, which belongs to the phylum Bacteroidetes of Gram-negative bacteria (Stanier, 1942). Proteins of the Pal family are ubiquitous in Gram-negative bacteria, interacting with peptidoglycan and forming the Tol-Pal complex to maintain the integrity of the cell envelop (Lloubes et al., 2001). Mutation of pal results in various changes, such as damage of the outer membrane permeability and formation of outer membrane vesicles (Bernadac et al., 1998; Kowata et al., 2016). Sequence alignment showed that the OmpA_C-like domain of CHU_0125 had a 40% identity value with EcPal. Our results showed that the Δ0125 mutant was hypersensitive to some harmful compounds. Meanwhile, deletion of chu_0125 resulted in the mutant producing about 6-fold more OMVs than the wild type (Figure 6B). Llamas et al. (2003) reported that the Tol-Pal complex was necessary for appropriate function of certain uptake system. We found that compared with the wild type, the Δ0125 mutant exhibited a long lag phase and a reduced

| Band | CHU no. | Description | MW (kDa) | pI |
|------|---------|-------------|----------|----|
| 1    | CHU_1075 | β-glycosidase-like protein | 274.2 | 5.09 |
| 2    | CHU_3437 | Hypothetical protein | 245.8 | 4.82 |
| 3    | CHU_3220 | Hypothetical protein | 198.9 | 4.97 |
| 4    | CHU_1107 | Endoglucanase | 135.2 | 5.39 |
| 5    | CHU_0125 | Outer membrane peptidoglycan-associated lipoprotein | 73.8 | 5.55 |
| 6    | CHU_0522 | Hypothetical protein | 29.1 | 7.09 |

* Band numbers listed here are corresponding to those denoted in Figure 7.
biomass at the stationary phase in organic medium (PY6 medium) but grew normally in inorganic medium (Stainer medium). When the PY6 medium was supplemented with 1 g/L NO\textsubscript{3}−, the growth of Δ0125 mutant recovered (Figure 2). We deduced that limited uptake of some nutrients might be responsible for the hypotrophy of the Δ0125 mutant in PY6 medium.

We further analyzed the domain structure of CHU_0125, which was found to be different from the domain structures of reported Pals. The reported Pals only possessed an OmpA_C-like domain while CHU_0125 had two other domains, TPR and PD40 (Figure 1A). We also analyzed the putative Pals in other bacteria in the phylum Bacteroidetes, including Porphyromonas gingivalis, Flavobacterium johnsoniae, Flavobacterium columnare, Pontibacter indicus, Hymenobacter sedentarius, and Aquimarina megaterium and found that they all possessed other domains in addition to the OmpA_C-like domain, suggesting the putative Pals in the phylum Bacteroidetes might be different from other reported Pals. Moreover, Cascales et al. reported that the C-terminal SYGK/E motif, a TolA box, was required for the binding of Pal to TolA in E. coli (Cascales and Lloubes, 2004). The KNRR motif was also needed for Pal to interact with cell-envelope proteins in E. coli (Cascales and Lloubes, 2004). These two motifs were also conserved in some reported Pals (Figure 1B). But the two motifs were changed to GYGE and TNRR in CHU_0125 respectively, suggesting that CHU_0125 might interact with other proteins in a different way from Ecpal (Figure 1B). In addition to these differences in protein structures, some phenotypes of the Δ0125 mutant were also different from those of Δpal in E. coli. In E. coli, deletion of pal causes a motility defect (Santos et al., 2014). However, in C. hutchinsonii, cells of the Δ0125 mutant could spread as normally as the wild type on soft agar (Supplementary Figure S5). Gerding et al. (2007) revealed that the Tol-Pal complex was a part of the cell division machinery, but the Δ0125 mutant did not exhibit cell division deficit as observed by SEM. In addition, we deleted tola (chu_0135) and tolB (chu_1429) and found that the Δ0135 mutant and the Δ1429 mutant could digest filter paper and grew in PY6 medium as same as the wild type, indicating that the integrity of the Tol-Pal complex was not necessary for cellulose degradation in C. hutchinsonii (Supplementary Figure S6).

Previous study showed that C. hutchinsonii appeared to use a contact-dependent digestion strategy, indicating the importance of the outer membrane which directly contacted with cellulose in cellulose degradation. However, only several outer membrane proteins, including CHU_1276, CHU_1277, CHU_0170, and CHU_3220, were reported to be important for the digestion of cellulose. In this study, we found that an outer membrane protein CHU_0125 was necessary and specialized for the degradation of the crystalline region of cellulose by C. hutchinsonii. Deletion of chu_0125 impaired the integrity of outer membrane and this impairment of the outer membrane integrity resulted in the decrease of some outer membrane proteins in the Δ0125 mutant (Figure 7 and Table 3). Genes of these proteins listed in Table 3 were singly deleted, but the mutants were able to digest cellulose as same as the wild type except the Δ3220 mutant. Our previous study showed that the Δ3220 mutant could not digest crystalline region of cellulose, which was similar to the phenotype of the Δ0125 mutant in degradation of cellulose (Wang et al., 2017). In addition, Western blotting showed that amounts of CHU_3220 were leaked from outer membrane and packaged into OMVs in the Δ0125 mutant. These results indicated that deletion of chu_0125 influenced the localization of CHU_3220, which might affect the degradation of the crystalline region of cellulose by the Δ0125 mutant. But how the misplaced CHU_3220 affected the cellulose degradation of the Δ0125 mutant needed to be further studied.

CONCLUSION

This work revealed that deletion of chu_0125 which encoded a putative outer membrane peptidoglycan-associated lipoprotein prevented degradation of the crystalline region of cellulose, which improved our understanding of the function of Pal. This study also enhanced the importance of the integrity of outer membrane in the degradation of the crystalline region of cellulose by C. hutchinsonii.

AUTHOR CONTRIBUTIONS

XW, ZW, XB, and XL conceived and designed the experiments. XW, ZW, and XB performed the experiments. XW, XB, and WZ analyzed the data. XW, ZW, XB, YZ, WZ, and XL wrote the paper. XW, XB, and XL revised the manuscript. All authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2018.00632/full#supplementary-material
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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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