Identification of a Cupin Protein Gene Responsible for Pathogenicity, Phage Susceptibility and LPS Synthesis of Acidovorax citrulli

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Bacteriophages infecting Acidovorax citrulli, the causal agent of bacterial fruit blotch, have been proven to be effective for the prevention and control of this disease. However, the occurrence of bacteriophage-resistant bacteria is one of hurdles in phage biocontrol and the understanding of phage resistance in this bacterium is an essential step. In this study, we aim to investigate possible phage resistance of A. citrulli and relationship between phage resistance and pathogenicity, and to isolate and characterize the genes involved in these phenomena. A phage-resistant and less-virulent mutant named as AC-17-G1 was isolated among 3,264 A. citrulli Tn5 mutants through serial spot assays and plaque assays followed by pathogenicity test using seed coating method. The mutant has the integrated Tn5 in the middle of a cupin protein gene. This mutant recovered its pathogenicity and phage sensitivity by complementation with corresponding wild-type gene. Site-directed mutation of this gene from wild-type by CRISPR/Cas9 system resulted in the loss of pathogenicity and acquisition of phage resistance. The growth of AC-17-G1 in King’s B medium was much less than the wild-type, but the growth turned into normal in the medium supplemented with D-mannose 6-phosphate or D-fructose 6-phosphate indicating the cupin protein functions as a phosphomannos isomerase. Sodium dodecyl sulfate analysis of lipopolysaccharide (LPS) extracted from the mutant was smaller than that from wild-type. All these data suggest that the cupin protein is a phosphomannos isomerase involved in LPS synthesis, and LPS is an important determinant of pathogenicity and phage susceptibility of A. citrulli.

Keywords: Acidovorax citrulli, cupin, pathogenicity, phage resistance

Bacteriophages, the most abundant entity in all microbial environments, were discovered by d’Herelle (1926), and were proposed for use as antibacterial agents. The use of phages to infect Vibrio cholera and other human pathogens were among the early studies of phages (Marks and Sharp, 2000). In addition to human pathogens, phages were investigated for the control of plant diseases in the pre-antibiotic era. For example, Coons and Kotila (1925) showed that phage application could prevent Pectobacterium atrosepticum from causing soft rot in potato tubers. However, most commercial use of phages was replaced with the introduction of antibiotics in the 1920s. Since then, awareness and consideration of phages has increased for three main reasons: widespread application of antibiotics, development of resistant bacteria, and the need for new antibacterial agents against antibiotic-resistant. Therefore, numerous studies have focused on the use of phages against pathogenic bacteria, and phages were recently reconsidered as an

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alternative solution for control (Park et al., 2018; Vu and Oh, 2020). However, similar to other biocontrol strategies, several factors limit bacteriophage biocontrol. Most of these factors are related to the nature of bacteriophages. For example, to successfully apply bacteriophages under physiological conditions, factors such as temperature, ultraviolet radiation, and pH must be optimized.

In addition to these limitations, which are generally related to bacteriophage structure, the occurrence of bacteriophage-resistant bacteria is a major issue in the development of bacteriophages as a control agent (Hassan et al., 2021). Coevolution of bacteriophages and bacteria occurs constantly through spontaneous mutations (Brockhurst et al., 2021) that allow bacteria to structurally change the composition of their cell wall, capsule, lipopolysaccharide (LPS), and flagella, which act as receptors for phages. For example, LPS alteration and OmpC surface protein silencing were found to be the two main factors associated with phage resistance in *Escherichia coli* (Pirhonen et al., 1988). Therefore, studies on phage resistance mechanisms and detailed analysis of the interactions between phages and bacteria are necessary to increase the chance of successful long-term application of phages.

The potential mechanisms through which bacteria become resistant to phages were previously found to be dependent on genotype (Barbu et al., 2016). Immunity of bacteria against bacteriophages may involve various mechanisms, such as mutations, restriction modification systems, and adaptive immunity via the CRISPR/Cas system. For example, the type I CRISPR/Cas system, which is abundant in bacteria, provides protection against the virulent phages φTE and φM1 in *P. atrosepticum* (Watson et al., 2019). Bacteriophage-resistant bacteria increase rapidly, as phage-susceptible genotypes are killed by phages in the same environment. A population of bacteria that has rapidly acquired resistance may have either positive or negative implications for the host. For example, diarrhea-causing *E. coli* showed decreased virulence after becoming resistant to bacteriophages. By contrast, *Enterococcus faecalis* resistant to bacteriophage φVPE25 was able to colonize the intestine, in addition to growing faster than the wild-type in the intestine (Chatterjee et al., 2020). Therefore, understanding bacteriophage-host interaction is essential prior to application of phages.

*Acidovorax citrulli* (previously known as *Acidovorax avenae* subsp. *citrulli*) is a gram-negative bacterium that causes bacterial fruit blotch (BFB) in *Cucurbitaceae*. *A. citrulli* strains have been divided into groups, with strains in group I showing virulence against plants in the *Cucurbitaceae* family other than watermelon, and group II strains primarily affecting watermelon. Like other plant pathogenic bacteria, *A. citrulli* disease relies on the type III secretion system that is responsible for secretion of effector proteins and mutation in the genes encoding type III secretion system and type IV secretion system affected virulence, motility, and biofilm formation of *A. citrulli* (Eckshtain-Levi et al., 2014; Tian et al., 2015). In addition to these secretion systems, several genes involved in pathogenicity of *A. citrulli* have been identified in the analysis of Tn5 mutants with significantly decreased the virulence but the detailed role of each gene remains unclear (Luo et al., 2015).

Despite the economic importance of *A. citrulli*, effective chemical control agents are limited and resistant cultivars are not available yet. Use of bacteriophages against *A. citrulli* has been proposed and studied since 2016, after the initial isolation of phage ACP17 (Rahimi-Midani et al., 2018), the issue of phage resistance has not been investigated yet. As phage-host interactions can provide insights into the bacterial genome and possibly pathogenic behavior, understanding such interactions is essential. Therefore, in this study, we aimed to identify the nature of bacteriophage-resistant in *A. citrulli* and any gene involved in this characteristic. Also, the relationship between phage resistance and pathogenicity was analyzed.

### Materials and Methods

**Preparation of bacterial mutants.** The *A. citrulli* mutant library of 3,264 strains was kindly provided by Professor Sang-Wook Han of Chung-Ang University, Republic of Korea. These mutants were constructed from the wild-type *A. citrulli* strain KACC17005 using the transposon mutation system EZ-Tn5 Transposome Kit (Lucigen, Madison, WI, USA) according to the manufacturer’s instructions.

**Screening of phage-resistant bacteria.** Bacteriophage ACPWH, isolated previously, was used to screen phage-resistant mutants (Rahimi-Midani et al., 2020). Bacteria from the mutant library were grown in King’s B (KB) medium containing 100 µg/ml each of kanamycin and rifampicin overnight at 37°C. Then, 300 µl of bacterial culture was mixed with 100 µl 0.7% agar in KB medium and poured into each well of a 48-well culture plate for spot assay. After hardening, 3 µl of bacteriophage ACPWH (titer, 10^6 pfu/ml) was spotted onto each well and the plates were incubated at 28°C for 24 h. Mutants resistant to bacteriophage ACPWH, which showed no clearing in the spot assay, were selected; the plaque assay was repeated three times to confirm resistance in the selected mutant.
Screening of avirulent *A. citrulli* mutants. In total, 39 strains of *A. citrulli* mutant resistant to phage ACPWH were used for pathogenicity test by seed coating of watermelon seeds as described previously (Rahimi-Midani et al., 2020). Briefly, commercial seeds were washed with 1% sodium hypochlorite for 20 min. After washing, the seeds were mixed with a suspension of bacterial mutants (10^5 cfu/ml, in phosphate-buffered saline). Nine seeds coated with each bacterium were sown in three plastic pots of 12cm diameter (3 seeds per pot) and maintained for 2 weeks, during which germination data were recorded. Screening was conducted in triplicate.

Identification of mutated genes. Gene identification was carried out using the rescue cloning method, as described in the manual of the EZ-Tn5 Tnp Transposome Kit. Briefly, bacteria were grown in 100 ml KB broth medium containing kanamycin (100 μg/ml). Bacterial DNA was extracted using the AccuPrep Genomic DNA Extraction Kit (Bioneer, Daejeon, Korea). One microgram of total extracted DNA was treated with EcoRI enzyme and incubated for 1 h at 37°C. Digested DNA was ligated with the EcoRI adaptor (GGATCCGCGGCC) and PCR was conducted using the EcoRI adaptor primer (GGATCCGCGGCCNN) under the following conditions: 95°C for 5 min; 32 cycles of 94°C for 20 s, annealing for 20 s at 52°C, and extension at 72°C for 20 s; followed by an extension phase at 72°C for 3 min. The PCR product was used for direct sequencing without cloning.

Complementation. The gene encoding a cupin protein was amplified from wild-type *A. citrulli* using cupin protein gene promoter forward and reverse (F and R, respectively) primers, which are listed in Table 1. PCR was performed using Taq plus (Epsisi-Biotech, Daejeon, Korea), as follows: 95°C for 5 min; 32 cycles of 94°C for 20 s, annealing for 20 s at 60°C, and extension at 72°C for 20 s; followed by an extension phase at 72°C for 3 min. DNA was purified using the ExpiN PCR SV kit (GeneAll Biotechnology, Seoul, Korea). The 1.3-kb PCR product was cloned into Takara T-Vector pMD9 (Takara Korea Biomedical Inc., Seoul, Korea) and used for complementation.

A phage-resistant and avirulent mutant strain was used for complementation with the cloned cupin protein gene. One hundred milliliters of exponential-phase bacterial culture (OD600 of ~0.6) was centrifuged at 2,800 × g for 30 min. The pellet was resuspended in chilled 10% glycerol and centrifuged at 2,800 × g for 30 min. Glycerol washing and centrifugation were repeated three times. The electrocompetent cells were resuspended in 10% glycerol and stored at −80°C. Plasmids were electroporated into the electrocompetent cells under the conditions of 2.0 kV, 200 ohms, and 25 μF using Gene Pulser II electroporator (Bio-Rad, Hercules, CA, USA), and recovered in super optimal broth with catabolite repression medium at 37°C with shaking for 1 h. Electroporated cells were plated onto KB agar plates containing 100 μg/ml of ampicillin. Transformation was confirmed through PCR with the M13 forward and reverse primers. Positive strains were used for the plaque assay and seed coating experiment. Thirty surface-sterilized watermelon seeds were coated with the avirulent mutant or the complemented strain. The seeds were sown in six pots.

| Primers | Sequence | Annealing temperature (°C) |
|---------|----------|---------------------------|
| Transposon left-end reverse (TLR) | CTACCCCTGTGGAACACCTACATCT | 59 |
| Transposon right-end forward (TRF) | ACCTACAAACGCTCTCATCAACC | 60 |
| Transposon left-end forward (TLF) | AGATGTAGGTGTTCCACAGGGTAG | 60 |
| Transposon right-end reverse (TRR) | GGTGTGATGAGGCTTTGTTGAGG | 59 |
| Cupin protein gene promoter forward (CPF) | GGACCAACTCGTCGTAGCCC | 60 |
| Cupin protein gene downstream reverse (CDR) | GTTGGAGACGGTCCGC | 59 |
| EcoRI adaptor sequence F | GGATCCGCGGCCNN | 52 |
| SDM-Cupin F | TACTAAGGAGGTTGACGCGGAACGTGGTGTTTCGAGCTAGA | 50 |
| SDM-Cupin R | TTCGTTCTACGTCGTTACGAGCTTAC | 48 |
| GRNA-Check F | TAAACCGGTGCGTGTTGACT | 58 |
| GRNA-Check R | GACTCCTGCCTCTTTC | 57 |

Bold sequences indicate the single guide RNA.
measuring 15 cm × 8 cm × 12 cm per treatment, which were maintained for 20 days to record the germination rate.

**Knockout of the cupin gene by CRISPR/Cas9.** In order to confirm the function of the cupin protein gene in pathogenicity and phage susceptibility, this gene was knocked out from the wild-type *A. citrulli* by the CRISPR/Cas9 system. A single guide RNA (sgRNA) for the gene was designed using online tools (Broad Institute), which are listed in Table 2. This sgRNA was then ligated into the all-in-one vector (Addgene plasmid no. 78098, http://n2t.net/addgene:78098) using the site-directed mutagenesis (SDM) method, as described by Liu and Naismith (2008) (Supplementary Fig. 1) with the SDM primers listed in Table 1. The vector was then electrically transformed into wild-type *A. citrulli* as described above. The transformed bacteria were grown on KB medium containing ampicillin (100 μg) and selected colonies were grown on KB ampicillin medium, and used for spot assays, as described previously (Khan Mirzaei and Nilsson, 2015).

**Growth patterns of the wild-type strain and mutants.** As one major group of bacterial cupin protein is the phosphomannose isomerase (PMI) that catalyzes the interconversion of D-fructose-6-phosphate (F6P) and D-mannose-6-phosphate (M6P), the growth of wild-type *A. citrulli* and the phage-resistant and avirulent mutant AC-17-G1 were compared in KB medium containing 5 mM each of M6P and F6P. The wild-type and mutant strains were cultured overnight in KB medium and KB medium containing kanamycin (40 μg/ml), respectively and were then inoculated into KB medium, KB medium containing kanamycin, and KB medium containing 5 mM each of M6P or F6P and incubated at 26°C. The optical density was recorded hourly. Plaque assay was conducted with phage ACPWH and mutant AC-17-G1 cultured in these mediums.

**Comparison of LPS in wild-type and mutant.** LPS was extracted from the wild-type and the mutant AC-17-G1 as described previously (Szzermer-Olearnik and Boratyński, 2015). Briefly, bacteria were grown overnight in KB medium and centrifuged at 2,800 × g for 30 min. The pellet was resuspended in 55% hot water-phenol solution. The resulting mixture was incubated at 60°C for 1 h and centrifuged at 21,206 × g for 10 min. The upper phase was removed and treated with DNase (100 μg/ml), RNase (50 μg/ml, Takara Korea), and proteinase K (100 μg/ml, Bioneer). The treated sample was subjected to precipitation in 100% ethanol, as described previously (Darveau and Hancock, 1983), and the extracted LPS was analyzed via sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE).

### Results

**Screening of mutants with phage-resistant and reduced virulence.** A total of 3,264 transposon mutant strains were used for the selection of mutants resistant to phage ACPWH, which can lyse the wild-type host *A. citrulli* KACC17005. Thirty-nine mutants showing no lysis in the spot assay were obtained and further screened for pathogenicity. Seeds coated with 37 mutants had no germination or germinated with severe symptoms resulting in death within a few days as the seeds coated with wild-type *A. citrulli*. Seeds coated with one mutant designated as AC-5-E1 showed a germination rate of 55% and less severe symptoms of BFB. By contrast, seeds coated with the mutant AC-17-G1 showed 88.8% germination with reduced or no BFB symptoms (Fig. 1A). Therefore, this mutant was used for the identification of genes involved in both phage susceptibility and pathogenicity (Fig. 1B).

**Identification of the mutated gene in mutant strain AC-17-G1.** DNA extracted from the phage-resistant less-virulent mutant AC-17-G1 was digested with EcoRI, which cut one site in the transposon gene and an unknown site near the integration site. An EcoRI linker was ligated to the digested DNA and amplified with a primer specific to the adaptor. A PCR product of almost 1 kb was obtained, and sequence analysis of this product showed that Tn5 was integrated into the cupin protein gene of *A. citrulli* KACC17005, as annotated to the genome sequence (GenBank accession no. NZ_CP023687.1). This finding was

### Table 2. Guide RNAs used for CRISPR/Cas9 in this study

| Name  | Sequence | Size (bp) | PAM  |
|-------|----------|-----------|------|
| Sgrna1| AGCCGAACGTGTGGTTCGAGCCCGGGGATC | 30 | GGG |
| Sgrna2| GATGATCAGCTACGCCACCGACGGCGGGGG | 30 | GGG |
| Sgrna3| CGCGCGCCGTACATCATCGCGTGCGGCGG | 30 | CGG |
| Sgrna4| AGCCGAACGTGTGGTTCGAGCCCGGGGATCG | 31 | CGG |

PAM, protospacer adjacent motif.
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Further confirmed through PCR amplification with primers specific to Tn5 and the cupin protein gene, as shown in Fig. 2. Transposon is inserted between position 672 and 673 of the cupin protein open reading frame. The adenine residue of the AUG start codon and the last guanine residue of the TAG stop codon correspond to the nucleotide number 3094447 and 3095583 of NZ_CP023687.1, respectively. PCR with the cupin protein gene promoter F primer and transposon left-end R primer resulted in a DNA fragment of 1,155 bp (Fig. 2, lane F), while PCR with the transposon right-end F primer and cupin protein gene downstream R primer resulted in a DNA fragment of 874 bp (Fig. 2B, lane R). Sequence analysis of these PCR products confirmed that Tn5 is integrated 672 bp from the AUG codon of the cupin protein gene.
Complementation of mutant AC-17-G1. The phage-resistant and less-virulent mutant AC-17-G1 was complemented with the cupin protein gene from the wild-type. Transformed cells were selected on KB medium containing ampicillin, to which the wild-type is sensitive. The presence of introduced DNA was confirmed through PCR with the primers M13 F and m13 R, and a DNA fragment of the expected size was detected. The presence of the wild-type cupin protein gene was further confirmed through PCR with cupin-specific primers.

The selected complemented line was then tested for sensitivity through a plaque assay. As shown in Fig. 3, the complemented mutant showed sensitivity to phage ACP-WH, in contrast to mutant ACP-17-G1, which was resistant to this phage.

Pathogenicity test of the complemented mutant. The mutant AC-17-G1 that had been complemented with the cupin protein gene from wild-type A. citrulli was further investigated for the recovery of pathogenicity. As shown in Fig. 4, watermelon seeds coated with the mutant AC-17-G1 showed 90% germination with no BFB symptoms. However, the mutant complemented with the wild-type cupin protein gene led to virulence as severe as in wild-type A. citrulli.

Fig. 3. Recovery of mutant AC-17-G1 for phage sensitivity by complementation with wild-type gene. Plaque assay with phage ACPWH was conducted with mutant AC-17-G1 (A) and the same mutant complemented with wild-type cupin protein gene (B).

Fig. 4. Pathogenicity test of the complemented mutant. (A) Germination of watermelon seeds coated with wild-type, mutant AC-17-G1 and mutant AC-17-G1 complemented with wild-type gene. (B) Representative pots sown with watermelon seeds coated with designated bacterial strains.
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Confirmation of gene mutation using CRISPR/Cas9. SDM was performed on the cupin protein gene of wild-type *A. citrulli* using the CRISPR/Cas9 system. We designed four sgRNA for the all-in-one vector and introduced them into wild-type *A. citrulli*. Among colonies selected on ampicillin plates, those recovered from the transformant with sgRNA4 showed resistance against ACPWH (Fig. 5A). The cupin protein gene from this mutant was amplified through PCR with the cupin protein gene promoter F and R primers, and its sequence was analyzed. As shown in Fig. 5B, the mutant with sgRNA showed the deletion of four nucleotides in the middle of the cupin protein gene.

**Fig. 5.** Confirmation of site-directed mutagenesis of the cupin protein gene using the CRISPR/Cas9 system. (A) Spot assay with phage ACPWH against wild-type *Acidovorax citrulli* (W), mutant AC-17-G1 (G), and a cupin protein gene deleted mutant generated by CRISPR/Cas9 (C). (B) Sequence analysis of the mutant produced by CRISPR/Cas9 system. Four nucleotides were deleted in the middle of the cupin protein gene in the mutant created using sgRNA4. The deleted four nucleotides are shown in bold. The single-guide RNA sequence is underlined and the PAM sequence is indicated with dashes.

**Fig. 6.** Growth of *Acidovorax citrulli* mutant AC-17-G1 in King’s B (KB) medium supplemented with D-mannose-6-phosphate (M6P) and D-fructose-6-phosphate (F6P) and recovery of phage sensitivity. (A) The growth of wild-type and mutant strains in KB medium and KB medium supplemented with M6P and F6P. (B) Plaque assay with phage ACPWH and mutant AC-17-G1 grown in KB medium (K), KB medium supplemented with F6P (F), and M6P (M).
which resulted in a frame shift of the open reading frame (ORF) and premature termination of the protein.

**Growth of the mutant in the presence of F6P and M6P.** To determine whether the cupin protein is a PMI, we compared the growth of mutant AC-17-G1 in the presence of M6P and F6P. As shown in Fig. 6A, mutant AP-17-G1 exhibited slow growth in KB medium. However, when M6P or F6P was added to the medium, the mutant grew at the same rate as the wild-type in KB medium. In addition, plaque assay showed that mutant AC-17-G1 grown in media containing M6P or F6P was sensitive to phage ACPWH contrast to the bacterium grown in KB medium was resistant to this phage (Fig. 6B).

**LPS characterization.** LPS was extracted from wild-type and phage-resistant avirulent mutant AC-17-G1 and analyzed using SDS-PAGE. As shown in Fig. 7, LPS extracted from the mutant was approximately 10 kDa in size, whereas that from the wild-type was approximately 12 kDa.

**Discussion**

Recently, the use of bacteriophages for biocontrol has been revisited as an alternative to antibiotics. To develop this novel alternative strategy, detailed characterization of bacteriophages and their host interactions are essential. Bacteria can resist attachment of bacteriophages through various mechanisms to avoid infection in nature. Bacteria can also stop phage adsorption, degrade phage DNA, and initiate cell death upon infection. However, bacteria-phage coevolution not only increases the level of population diversity, but has also been shown to affect bacterial ecological behavior (Koskella and Meaden, 2013). For example, reports have suggested that bacterial strains resistant to lytic phages were more susceptible to other phages (Koskella and Brockhurst, 2014), had reduced carbon metabolism (Xu et al., 2019), and even showed structural changes (Oechslelin, 2018). The fitness cost for bacteria has been shown to depend on the timing of infection and exposure time to phages.

In this study, we used the T5 transposon mutation method for mutation of *Acidovorax citrulli* strains, resulting in 39 of 3,624 total mutants showing resistant to phage infection in spot assay. Our study focuses on phage-resistant bacteria with reduced virulence. Through seed coating assay for pathogenicity with these phage-resistant mutants, a mutant AC-17-G1 with severely reduced pathogenicity was selected (Fig. 1). Sequence analysis of the selected mutant AC-17-G1 with phage resistance and reduced virulence showed that Tn5 is inserted in a gene that encodes a protein with the cupin domain, which has two histidine motifs with variable length regions (Fig. 2).

Cupin superfamily proteins exhibit extremely diverse functions and contain a considerable number of proteins with unknown functions. Khuri et al. (2001) showed that cupin proteins are a diverse group of proteins present in all three kingdoms of life. Moreover, cupin protein are found in most of microorganism including bacteria with various functions.

The function of the cupin protein in phage resistance and virulence was confirmed by complementation of the mutated gene with the corresponding gene cloned from wild-type. The complemented bacteria recovered phage sensitivity as shown in Fig. 3 and showed the same pathogenicity as the wild-type (Fig. 4). These results indicated that the cupin protein gene of *A. citrulli* encodes a protein involved in phage sensitivity. These was further confirmed by generation of a site-directed mutant by using CRIPR/Cas9 system. A mutant showing no lysis in spot assay with phage ACPWH indicating the loss of phage sensitivity (Fig. 5) was selected and sequence analysis of the targeted area showed deletion of four nucleotides in the cupin protein gene. Pathogenicity test of this mutant showed similar virulence in seed coating assay (data not shown). Therefore, it was concluded that the cupin protein gene of *A. citrulli* encodes a protein that affect both of phage sensitivity and pathogenicity.

Although we confirmed that the cupin protein of *A. citrulli* could be one of, if there any, complete determinant for phage resistance, this protein is not the only gene involved in pathogenicity. Mutant AC-17-G1 showed complete resistance to phage ACPWH but showed reduced behavior.
virulence and was not non-pathogenic. In the analysis of *A. citrulli* Tn5 mutants, Luo et al. (2015) identified insertion of Tn5 at eight genes including cation diffusion facilitator family transporter, alpha/beta hydrolase fold, and inosine monophosphate dehydrogenase which resulted in reduced pathogenicity. Zhang et al. (2018) showed that deletion mutation of the genes *hrpG* or *hrpX* that involved in the regulation of type 3 secretion system reduced virulence on watermelon seedlings as well as hypersensitive responses. In addition, mutation of the *acrR* gene of *A. citrulli* that involved in the motility and flagella formation resulted in virulence with disease index of 31.7 compare to 48.88 of the wild-type (Guan et al., 2020). Compare to the virulence of these mutants, mutant AC-17-G1 severe decrease in virulence, 0% vs. 88.8% germination in seeds coated with wild-type and mutant AC-17-G1, respectively. This suggested that the cupin protein is a major factor for virulence.

The pathogenicity of *A. citrulli* strains is related to multiple factors, including the type III and type IV secretion systems and flagella. However, genes in this bacterium are spread through the genome in structures called pathogenicity islands (PAIs). PAIs are responsible for various virulence factors, and the ORF of the cupin protein gene is located within a PAI in the genome of *A. citrulli*. Homology of cupin protein-coding sequences was identified in three *A. citrulli* strains (AA78-5, AA99-2, and ATCC19860), which all showed similar locations in PAIs and similar roles in pathogenicity. Due to differing annotation methods, homologues of cupin protein have generally been reported as hypothetical proteins, but this gene has been confirmed to play a role in pathogenicity in *A. citrulli*. Notably, the genes responsible for virulence in the type III secretation system have not been fully clarified; therefore, this result will help elucidate PAIs in *A. citrulli* and further characterization of gene involved in pathogenicity is required.

The importance of phage-resistant bacteria to the development of antibacterial agents has attracted researchers, who have identified genes related to phage resistance. Studies have divided the mechanisms of phage resistance in bacteria into two groups, LPS-related and non-LPS-related genes. For example, six receptors for bacteriophages in bacteria into two groups, LPS-related and non-LPS-receptor phage. This was further confirmed by SDS-PAGE analysis of LPS extracted from wild-type and mutant. LPS from mutant AC-17-G1 was smaller than that from wild-type (Fig. 7). Studies with mutant strains defective in LPS biosynthesis-related genes have indicated that LPS is required for bacterial virulence. For example, mutations in LPS biosynthesis-related genes of *Salmonella enterica* resulted in the loss of its virulence (Kong et al., 2011). Similarly, mutation of genes involved in the synthesis of the core oligosaccharide (OS) of LPS, resulted in clear reduction in the virulence of *Burkholderia glumae* that causes bacterial panicle blight and seedling rot in rice (Lee et al., 2019). Since PMI is involved in the production of intermediates used for the synthesis of the OS of LPS, our data coincide with these previous data.

It has been suggested that selection of host populations for the resistance to parasites can results in the alteration of other traits as trade-off. For example, mutants of *Pseudomonas syringae* that have evolved resistance via a single mutation showed significant reduction of the growth within plants tissue and reduction in pathogenicity (Meaden et al., 2015). Since the mutants were screened for phage resistance first and then tested for pathogenicity, mutant AC-17-G1 is not the case of natural selection for resistance against phage and loss of its pathogenicity as the cost. However, similar evolutive changes could happen especially considering the prevalence of phages infection *A. citrulli* as previously reported (Rahimi-Midani et al., 2020) and this could be an obstacle for the application of phages as the biocontrol agents of BFB. However, these results also support the hypothesis that phage resistance not only reduces pathogenicity, but can also increase the number
“good strains” of *A. citrulli* in the presence of phages.

**Conflicts of Interest**

No potential conflict of interest relevant to this article was reported.

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**Electronic Supplementary Material**

Supplementary materials are available at The Plant Pathology Journal website (http://www.ppjonline.org/).

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