The Intracellular Citrus Huanglongbing Bacterium, ‘Candidatus Liberibacter asiaticus’ Encodes Two Novel Autotransporters

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Abstract

Proteins secreted by the type V secretion system (T5SS), known as autotransporters, are large extracellular virulence proteins localized to the bacterial poles. In this study, we characterized two novel autotransporter proteins of ‘Candidatus Liberibacter asiaticus’ (Las), and redesignated them as LasA₁ and LasA₂ in lieu of the previous names Hyv₁ and Hyv₂. As a phloem-limited, intracellular bacterial pathogen, Las has a significantly reduced genome and causes huanglongbing (HLB), a devastating disease of citrus worldwide. Bioinformatic analyses revealed that LasA₁ and LasA₂ share the structural features of an autotransporter family containing large repeats of a passenger domain and a unique C-terminal translocator domain. When fused to the GFP gene and expressed in E. coli, the LasA₁ C-terminus and the full length LasA₂ were localized to the bacterial poles, similar to other members of autotransporter family. Despite the absence of a typical signal peptide, LasA₁ was found to localize at the cell surface by immuno-dot blot using a monoclonal antibody against the partial LasA₁ protein. Its surface localization was also confirmed by the removal of the LasA₁ antigen using a proteinase K treatment of the intact bacterial cells. When co-inoculated with a P19 gene silencing suppressor and transiently expressed in tobacco leaves, the GFP-LasA₁ translocator targeted to the mitochondria. This is the first report that Las encodes novel autotransporters that target to mitochondria when expressed in the plants. These findings may lead to a better understanding of the pathogenesis of this intracellular bacterium.

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Introduction

Autotransporters are large multi-domain virulence factors encoded by genomes of diverse gram-negative bacteria. A typical autotransporter consists of three functional domains: a Sec-dependent N-terminal signal peptide, a secreted passenger domain (α-domain) and a conserved C-terminal translocator domain (β-domain) [1]. The central passenger domain will ultimately be either attached to the cell surface or secreted. This type of self-transporting protein system is referred to as a type V secretion system (T5SS). Known virulence factors secreted by T5SS have been shown to be cytotoxic, contain protease activities, or functions such as adhesions. Based on structural features, autotransporters have recently been classified into three sub-types: classical autotransporters (T5aSS), two-partner secretion system (T5bSS) and trimeric autotransporters (T5cSS) [2]. The signal peptide directs export of the precursor protein across the inner membrane using the Sec machinery and then is cleaved by peptidase. Subsequently, the β-domain inserts into the outer membrane and forms a pore with 12 transmembrane β-strands through which the passenger domain is presumed to be exported [3]. Once the passenger domain is translocated to the cell surface, it is usually cleaved from the translocator domain and released extracellularly. In some cases the passenger domain is not cleaved and remains tightly associated with the cells [4]. The trimeric autotransporters (T5cSS) known as AT-2 are exemplified by the oligomeric coiled-coil adhesions from various pathogenic bacteria, such as YadA of Yersinia [5], Hia of Haemophilus [6], and Hap of Haemophilus [7]. Compared with the conventional translocator domain that typically contains about 300 amino acids, AT-2 contains a short translocator domain of about 70 amino acids that is sufficient for translocation of the passenger domains [8,9]. Deletion of...
the YadA translocator domain abolishes the ability to insert into the outer membrane [9]. Many AT-2 passenger domains contain large repeat units of about 70 residues. Phylogenetic clustering of these repeat units revealed that they share striking clustering patterns in which some of the repeats are almost identical in sequence [10]. It has been reported that autotransporters from a variety of rod-shaped pathogenic bacteria, including IcsA and SepA of Shigella flexneri, AIDA-1 of Escherichia coli, and BrkA of Bordetella pertussis, are localized to the bacterial poles [11]. Recently, it was demonstrated that the YadA translocator localized solely to the mitochondrial outer membrane when expressed in yeast and that four β-stands are sufficient for mitochondrial localization [12].

‘Candidatus Liberibacter asiaticus’ is a Gram-negative, fastidious alpha-Proteobacterium, causing huanglongbing (HLB), a devastating disease of citrus worldwide. HLB causes rapid decline and shortens the life span of infected trees [13]. Having a greatly reduced genome of approximately 1.23 Mb, Las bacteria reside in phloem sieve cells of infected citrus plants and are transmitted by the citrus psyllids, Diaphorina citri [14,15]. Intriguingly, even with such a small genome size, the Las psy62 genome contains multiple prophage-related regions, and two were identified as prophages/temperate phages, which occupy ca. one-sixteenth of the entire Las genome [15,16].

Within these prophage regions, two hypothetical hypervariable proteins (Hyv and Hyv II) were identified that contained multiple, nearly-identical, leucine-rich repeats (LRRs). The diversity and plasticity of these two genes may have implications for how these intracellular bacteria adapt to their host ecological niches [17].

Prophages in many bacterial genomes are associated with bacterial pathogenicity and biofilm formation [18]. In the present study, we discovered that these two hypervariable proteins encoded by Las prophages are novel autotransporters (redesignated as LasA I and LasA II). We determined that LasA I and LasA II are polar and surface localized in bacteria in addition to being targeted to the mitochondria when expressed in plant cells. Previously, we demonstrated that the Las bacterium may act as an “energy parasite” by encoding a functional ATP translocase for direct ATP/ADP importation from their host cells [19]. Together these findings may lead us to understand how these intracellular bacteria modulate their host energy biosyntheses during their pathogenesis.

Results

Characteristics of unique autotransporters, lasA I and lasA II, in Las

The lasA I and lasA II, previously reported as hyv I and hyv II, are located in two prophage regions in the Las Psy62 genome [17]. The 2760 bp lasA I encodes a 919 amino-acid hypothetical protein with a predicted molecular mass of 38.9 kDa. LasA I has only one partial LRR. The translocator domains of LasA I and LasA II share 80% identity at the amino acid level while the passenger domains share 50% identity at the amino acid level. Using the SignalP signal peptide prediction software, no signal sequence was identified in LasA I or LasA II, and little information about the function of LasA I or LasA II was obtained from a BLAST search of the NCBI protein database. The LasA I and LasA II passenger domains share low level (about 25%) amino acid sequence similarities with the LRR protein of Colwellia psychrerythraea 34H (GenBank accession number: AAZ26055) and the cell wall associated biofilm protein of Staphylococcus epidermidis (ZP_06614153). Surprisingly, the passenger domains of LasA I and LasA II share similar LRR repeat structures with the Toll-like receptors (TLRs) that function as sentinels of the innate immune system by binding a variety of ligands, including lipopolysaccharide, flagellin and dsRNA, through a LRR ligand-binding domain [20]. LasA I and LasA II translocator domains were predicted to contain ten and twelve β-stranded secondary structures respectively by the YASPIN Secondary Structure Prediction program. However, the 3D structure predicted by the I-TASSER program did not form the typical β-barrel structure, which was reported for the translocator domain of the autotransporter Nap from Neisseria meningitidis [3,21]. Despite the absence of typical signal peptides and no significant sequence homology with other autotransporters at the amino acid level, sequence analyses predicted that LasA I and LasA II possess architectural features of the autotransporter family, including passenger domains with large repeated sequences that form coiled-coils and translocator domains containing β-stranded structures.

LasA I is an outer membrane protein and non-cleaved from cells

The full length gene lasA I was cloned into the pET102D-TOPO vector and protein expression was induced in E. coli BL21 (DE) cells (Invitrogen, Carlsbad, CA). A protein of the expected size for LasA I was shown on SDS-PAGE and confirmed by Western blot (Figure 1). LasA I was purified under hybrid conditions, and the elution fractions contained two bands detected by SDS-PAGE. The 120 kDa protein band including the 16 kDa fusion tag was verified by Western blot with an antibody against LasA I (N terminus, one full repeat and amino acids from part of the translocator domain). No signal was detected for the 40 kDa protein on the same Western blot (Figure 1B).

It has been shown that autotransporter passenger domains are transported to the cell surface and most of them are processed, thus releasing the passenger domain into the culture’s supernatant. To determine the subcellular localization of LasA I and its passenger domain, outer membrane proteins, surface-associated proteins and secreted proteins in culture supernatant were isolated. As shown in Figure 1, a 120 kDa protein was detected in outer membrane fraction by SDS-PAGE gel and confirmed by western blot, which suggests LasA I containing both passenger domain and translocator domain in these fractions. In contrast no LasA I protein was
detected from the culture supernatant using the anti-LasA\textsubscript{I} antibody (Figure S1A), which suggests that even though the LasA\textsubscript{I} protein contains the signal information required for cell pole localization in \textit{E. coli}, its passenger domain was not cleaved and released into the culture's supernatant in \textit{E. coli}.

The surface-associated protein was isolated from cell pellets and a Western blot was performed using an anti-LasA\textsubscript{I} antibody. No specific binding signal was observed (Figure S1A). Taken together, the results indicate that the LasA\textsubscript{I} is an outer membrane protein and its passenger domain was not cleaved and was still tightly associated with the translocator domain in \textit{E. coli}.

\textbf{Polar localization of LasA\textsubscript{I} and LasA\textsubscript{II}}

Several autotransporters from a variety of rod-shaped pathogenic bacteria are polar-localized in the bacterial [11]. We examined the localization of LasA\textsubscript{I} and LasA\textsubscript{II} by constructing GFP fusion proteins. The expression of GFP and GFP fusion proteins was detected by Western blot with an anti-GFP antibody (Figure S1B). When GFP was fused with the translocator domain of lasA\textsubscript{I} (pET102-gfp-lasA\textsubscript{I}-TD), or the full length lasA\textsubscript{II} gene (pET102-gfp-lasA\textsubscript{II}), the expression of GFP was observed at the cell poles of \textit{E. coli} by confocal laser scanning microscopy (CLSM) (Figure 2D–I). In the control panel, the expression of GFP (pET102-gfp) without the fusion partner was observed in the whole cell, which indicates GFP itself is not directed to bacterial cell poles (Figure 2A–C).

\textbf{Exportation of the LasA\textsubscript{I} and LasA\textsubscript{II} passenger domains by the translocator domains}

Although the typical N-terminal signal sequence found in most autotransporters was not identified in LasA\textsubscript{I}, our results showed that the passenger domain of the LasA\textsubscript{I} protein is localized at the \textit{E. coli} cell surface. Immuno-dot blot results showed strong signals indicating LasA\textsubscript{I} passenger domain is transported out of the bacterial cells when expressed in \textit{E. coli}, and no signal was observed in the control strain of \textit{E. coli} (Figure 3). Proteinase K-treated \textit{E. coli} containing the LasA\textsubscript{I} constructs did not bind the LasA\textsubscript{I} antibody, indicating that the passenger domain of LasA\textsubscript{I} was degraded on the surface of the bacterial cells. LasA\textsubscript{I} degradation by proteinase K was confirmed by SDS-PAGE and Western blot. Proteinase K-treated \textit{E. coli} cells expressing LasA\textsubscript{I} contained no signal while the untreated control cells contained the full 120 kDa protein (Figure 1). Proteinase K has no ability to cross the bacterial membrane and only digest the surface protein of intact
bacteria. This confirmed that the passenger domain of LasA was digested on the surface of the bacterial cells.

The LasA translocator domain not only exports its native passenger domain but also the LasA-GFP fusion protein to the cell surface. When GFP alone was expressed, the GFP protein stayed inside the cells and no GFP binding signal was detected with an anti-GFP antibody. However, when GFP was fused with the translocator domain of LasA, or the full length lasA gene, the GFP proteins were detected on intact bacterial surfaces with an anti-GFP antibody (Figure 4). After proteinase K digestion, immuno-dot blot results showed no GFP signal (data not shown). However, when the whole-cell lysate was treated with proteinase K and analyzed by SDS-PAGE, a slight reduction in the intensity of the fusion proteins was observed with Coomassie blue staining (data not shown). This indicated that not all fusion proteins expressed in E. coli are exposed on the surface of the bacteria and that the translocation of the GFP fusion protein is not as efficient as with the native LasA passenger domain.

To further confirm the surface localization of LasA, an immunofluorescence assay (IFA) was performed. The E. coli cells expressing the LasA protein were not labeled by anti-LasA antibody even though the isolated LasA protein and the E. coli cells expressing the LasA protein can be detected by Western blot and immuno-dot blot. The absence of surface labeling indicates that the LasA protein produced in E. coli may not be secreted and folded properly.

LasA targeting to mitochondria

The autotransporter YadA translocator domain was expressed in yeast and imported into the mitochondria, which did not interfere with mitochondrial function [12]. To investigate the potential function and cellular localization of LasA and LasAII in plant cells and the role of the LasA and LasAII translocator domains, full-length LasA, full-length LasAII, and the translocator domain of LasA were cloned into the pGDY vector and transformed into Agrobacterium tumefaciens strain GV 2660. Transient expression results showed no detectable GFP in tobacco plants inoculated with these constructs, except with the pGDY vector alone. Co-inoculation of tobacco leaves with a P19 gene silencing suppressor and pGDY-lasAII-TD construct facilitated GFP expression (data not shown). Only a few cells had detectable GFP in the infiltrated zone when the full length lasA or lasAII constructs were co-inoculated with the gene silencing suppressor. CLSM and propidium staining results showed that the expression of pGDY-lasAII-TD appeared to localize in the mitochondria. MitoTracker labeling and CLSM confirmed that GFP-lasAII-TD targeted to the mitochondria in tobacco leaves. As shown in Figure 5G and G1, the appearance of yellow mitochondria confirmed the localization of the pGDY-lasAII-TD fusion protein. The autofluorescence of chloroplasts was not observed by MitoTracker detection using a 560nm low-pass filter as shown by differential interference contrast (DIC) (Figure 5D and H). In contrast, GFP alone in whole cells expressed mainly in the nucleus and did not show yellow mitochondria (Figure 5C). Our results demonstrated that the translocator domain of LasA contains sufficient structural information for targeting mitochondria. No obvious cell death response was observed in the infiltrated leaf zone; however, mitochondria aggregation was observed when infiltrated with full length lasA and lasAII constructs (Figure 6A2 and A3). In infiltrated leaves, enlarged mitochondria and morphology change in chloroplast were observed and both of them are detached from cell wall (Figure 6B2). In addition, aggregation and changes in mitochondrial morphology were observed in infected periwinkle (Figure 6C2). Collectively, these results suggest that lasA and lasAII may affect mitochondria and chloroplast function and manipulate energy production during Las infection.

Discussion

We previously reported the genetic diversity and characteristics of two hypervariable proteins (HyvI and HyvII) from the Psy62 Las genome and global Las isolates that
contain up to 12 nearly identical tandem repeats [17]. In the present study, we discovered that LasA<sub>I</sub> and LasA<sub>II</sub> are two novel autotransporters. Most known autotransporters are virulence proteins in animal and human pathogens [2].

Typically autotransporters contain an N-terminal signal peptide, a passenger domain and a C-terminal translocator domain. However, no typical signal peptide was predicted in LasA<sub>I</sub> or LasA<sub>II</sub>, and while the amino acid sequences of LasA<sub>I</sub> and LasA<sub>II</sub> translocator domains contained predicted β-stranded structures, they shared no homology with translocator domains from other autotransporters. We propose that these proteins are new members of the autotransporter family because the translocator domains of LasA<sub>I</sub> and LasA<sub>II</sub> not only deliver their native passenger domains, but also exported GFP fusion proteins (GFP-LasA<sub>I</sub>-TD and GFP-LasA<sub>II</sub>-TD) onto the bacterial cell surface. These findings reveal that LasA<sub>I</sub> and LasA<sub>II</sub> are unique autotransporters and the T5SS may play an important role in Las pathogenesis. Furthermore, using transient gene expression in tobacco leaves, we demonstrated that LasA<sub>I</sub> contains sufficient structural information for targeting the host mitochondria as do other members of the autotransporter family [12].

Secreted proteins play a central role in the interactions of bacteria and their hosts. Gram-negative bacteria have evolved several specialized secretion systems to deliver effectors into their hosts, such as the Type I secretion system (T1SS) and the Type IV secretion system (T4SS). As an intracellular bacterium, Las does not have a T3SS or a T4SS, but may use the Sec secretion system [15,22]. Among the known bacterial secretion systems, the autotransporter or T5SS is the simplest pathway. Since the first report in the 1980s, the autotransporter family has been continuously expanding. Most of the characterized T5SS secreted proteins contribute to the virulence of animal or human pathogens [2]. The relatively few autotransporters reported from plant pathogens include the adhesins HecA/HecB of Erwinia chrysanthemi [23] and the XatA of Xylella fastidiosa which is important for virulence and is also associated with bacterial autoaggregation and biofilm formation [24]. In addition, the EstA autotransporter was reported as a member of the esterase family from the rice root colonizing and beneficial bacterium, Pseudomonas stutzeri A15 [25].

The amino acid sequences of autotransporters are highly divergent except for the conserved translocator domain. LasA<sub>I</sub> and LasA<sub>II</sub> proteins are unique autotransporters because they share no homology with any other members of the autotransporter family. Only coiled-coil domains were predicted and no signal peptides or anchor domains were identified in the LasA<sub>I</sub> and LasA<sub>II</sub> proteins when using the Trimeric Autotransporter Adhesins (TAAs) domain annotation tool [26]. Since the translocator domains of LasA<sub>I</sub> and LasA<sub>II</sub> have the ability to export their native passenger domains and GFP fusion proteins to the E. coli cell surface, the C-terminal translocator domains of LasA<sub>I</sub> and LasA<sub>II</sub> may form β-barrel structures through which the passenger domain can pass [2]. However, using the I-TASSER program to predict the 3D structures of LasA<sub>I</sub> and LasA<sub>II</sub>, the β-barrel structures of these translocator domains are atypical [21]. In other translocator

Figure 5. Mitochondrial localization of the Las autotransporter LasA<i><sub>I</sub></i>. A-G1: confocal laser scanning micrographs. GFP expression and MitoTracker labeling were detected in tobacco leaves infiltrated with pGDY and pGDY-lasA<sub>I</sub>-TD plasmids, respectively. A, E: GFP detection with 505-530 nm BP filter; B, F: MitoTracker detection with 560 nm LP filter; C, G: merged scans; D, H: differential interference contrast (DIC) micrographs of tobacco cells with chloroplasts (red arrows). E1, F1, G1: magnifications of yellow boxes in panels E, F and G. Mitochondria (yellow arrows). doi: 10.1371/journal.pone.0068921.g005
domains, such as that of NalP from Neisseria meningitidis, the crystal structure contains a 12-stranded β-barrel with a hydrophilic pore filled by an N-terminal α-helix [3]. It is not surprising that the predicted structures of the LasAII and LasAII translocator domains are different from other translocators as there are no conserved amino acids between the Las translocator domains and the known translocator domains. The crystallized structures of LasAII and LasAII will be important for understanding the passenger domain export mechanism.

LasAII and LasAII localize at bacterial poles as do other reported autotransporter members, including IcsA and SepA of Shigella flexneri, AldA-I of diffusely adherent E. coli and BrkA of Bordetella pertussis [11]. It has been shown that NalP from spherically shaped N. meningitides and BrkA from B. pertussis localize at the pole of E. coli, suggesting that autotransporters contain information required for polar localization [11]. It is interesting to note that in the IcsA protein of S. flexneri two regions within the passenger domain were involved in pole targeting [27]. In contrast, the LasAII translocator domain alone has the ability to target the bacterial poles. Further investigation should identify the region(s) essential for LasAII and LasAII to localize to the bacterial poles.

Typically T5aSS autotransporters exposed at the cell surface are proteolytically cleaved at the junction of the passenger domain and the outer-membrane embedded translocation domain [2]. Although LasAII and LasAII were surface-localized autotransporters, LasAII was present in greater amounts in whole-cell lysates and outer membrane but undetectable in the culture supernatant, indicating either that its inefficient cleavage in E. coli or that the cleaved passenger domain remains tightly associated with the translocator domain. The passenger domains of T5cSS, such as Hia from H. influenzae, are usually not cleaved and stay tightly associated with the cells [4]. In E. coli, BrkA is proteolytically cleaved at the bacterial surface, and the extracellular domain, though cleaved, remains tightly associated with the translocator domain [11]. Further efforts to confirm the LasAII and LasAII surface localization failed by IFA, although production of LasAII and GFP fusion proteins in E. coli was confirmed by immuno-dot blot and Western blot. The exported native passenger domains could not bind the primary antibody, indicating that they may not fold properly on the cell surface of E. coli. This was also observed with the IcsA of S. flexneri, which can be labeled at the surface of wild type S. flexneri but cannot be labeled in the E. coli cells expressing IcsA [28]. Once a pure Las culture is obtained, it will be possible to confirm the LasAII and LasAII surface localization by IFA and determine whether passenger domains are cleaved.

To understand the function of these novel autotransporters, lasAII and lasAII were constructed for Agrobacterium-mediated transient expression. Because no GFP expression was detected with the infiltrations containing our constructs, we used the gene silencing suppressor p19 to enhance the ectopic expression in plant leaves since post-transcriptional gene silencing (PTGS) is reported as a general feature in Agrobacterium-mediated transient expression [29–31]. As expected, strong GFP expression was observed when the lasAII translocater domain (pGDY-lasAII-TD) was co-infiltrated with the p19 construct; while no GFP was detected when infiltrated with the lasAII translocater domain (pGDY-lasAII-TD) alone. However, only a few GFP expressing cells were detected from the leaves co-inoculated with p19 and the full gene constructs of either lasAII or lasAII (pGDY-lasAII and pGDY-lasAII, respectively). This result may be due to GFP-LasAII and GFP-LasAII fusion proteins improperly folded in the plant cells. Surprisingly, we observed the mitochondria aggregation in these infiltrated cells even though there was no detectable GFP expression. We speculate that LasAII and LasAII may have the ability to self-cleave and produce functional subunits targeted to the mitochondria similar to the autotransporter VacA from Helicobacter pylori [32]. Further investigations are underway to confirm this hypothesis. Collectively, these results suggest lasAII and lasAII from a Las prophage/phage (bacterial virus) can act as inducers of PTGS in plant cells. To the best of our knowledge, this is the first evidence of bacterial prophage/phage gene inducing PTGS in plants. Further characterization of the PTGS conferred by LasAII and LasAII may shed light on the evolution and adaptation of the Las bacterium.

Figure 6. Mitochondria aggregation and morphology in plant cells. A1-A3: confocal laser scanning micrographs. MitoTracker labeling was detected in tobacco leaves infiltrated with pGDY (A1), pGDY-lasAII (A2) and pGDY-lasAII (A3). Arrows indicate normal and aggregated mitochondria respectively. B1-B4: transmission electron microscopy micrographs. B1: normal mitochondria (red arrow), chloroplast (blue arrow) and cell wall (green arrow) from infiltrated pGDY. B2: enlarged mitochondria (red arrow), abnormal chloroplast (blue arrow) and detached cell wall (green arrow) from infiltrated pGDY-lasAII. B3: normal mitochondria (arrow) from healthy periwinkle. B4: aggregated abnormal mitochondria (arrow) from Las-infected periwinkle.

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With the aid of the gene silencing suppressor p19, we revealed that the LasA<sub>I</sub> translocator domain-GFP fusion protein targeted to the mitochondria of tobacco leaf cells. Using the YASPIN secondary structure program, LasA<sub>I</sub> and LasA<sub>II</sub> were predicted to contain at least ten β-stranded structures in the translocator domain. It is worth noting that four β-strands in the YadA autotransporter of *Yersinia* are sufficient for its mitochondrial localization in yeast [12]. Furthermore, several bacterial proteins without typical N-terminal signal sequences also target mitochondria [33], indicating that the lack of signal peptides in LasA<sub>I</sub> and LasA<sub>II</sub> is not exceptional.

Proteins containing tandem repeats are associated with diverse functions, and the variable numbers of tandem repeats affect the pathogenicity or antigenicity of several human and animal pathogens [34]. Deletions or insertions of these repeats within the *lasA*<sub>I</sub> and *lasA*<sub>II</sub> genes were reported in samples of distinct geographical origins and a single origin [17]. It is interesting that the tandem repeats of the LasA<sub>I</sub> and LasA<sub>II</sub> passenger domains contain characteristics of the LRR family of proteins. The LRR proteins are important for immune responses, adhesion, invasion, signal transduction, and DNA/RNA processing [35]. The LRR motif of these proteins forms a "horseshoe-shaped" molecule that provides a versatile scaffold for protein–protein interactions [36]. Several LRR proteins have been shown to be located on the cell surface and play a role in surface adherence and aggregation [37]. Compared to the tobacco cells expressing the GFP-LasA<sub>I</sub> translocator domain, which did not affect mitochondrial morphology, we observed mitochondrial aggregation in cells infiltrated with full length *lasA*<sub>I</sub> or *lasA*<sub>II</sub>. This phenomenon could be explained if the translocator domain was integrated into the mitochondrial outer membrane with the LRR passenger domain facing the cytosol, thus causing mitochondrial aggregation. By transmission electron microscopy, mitochondrial aggregation was observed in Las infected periwinkle, which agrees with our observation that mitochondrial aggregation is caused by LasA<sub>I</sub> and LasA<sub>II</sub> expression in tobacco. Most of the reported LRR proteins contain an N-terminal signal peptide for secretion across the bacterial membrane and a C-terminal membrane attachment region followed by a hydrophobic transmembrane [37]. In contrast, LasA<sub>I</sub> and LasA<sub>II</sub> lack the classical signal sequence but the translocator domain can export the LRR passenger domain across bacterial membranes. AdpC, a LRR protein lacking a signal peptide, was also reported to be located on the outer membrane surface when it was expressed in a heterologous *E. coli* host [38]. Further investigation into whether LasA<sub>I</sub> and LasA<sub>II</sub> passenger domains target mitochondria are critical to understanding the functions of these proteins.

In conclusion, ‘Ca. Liberibacter asiaticus’ is an obligate, intracellular bacterium with a significantly reduced genome. We are the first to demonstrate that Las encodes two novel autotransporters (LasA<sub>I</sub> and LasA<sub>II</sub>) that target mitochondria when expressed in plant cells. Although the functions of these effectors remain to be elucidated, we hypothesize that Las encodes these autotransporters to modulate energy biosynthesis since Las may directly import ATP/ADP from the cytosol of host cell for its energy and biosynthesis [19]. On the other hand, these proteins may serve as suppressors for plant immune responses since Las encodes a functional flagellin that induces PAMP-triggered immunity in tobacco leaves [39]. Future work will focus on the functional elucidation of LasA<sub>I</sub> and LasA<sub>II</sub>, including an investigation into whether LasA<sub>I</sub> and LasA<sub>II</sub> passenger domains target mitochondria, identification of the eukaryotic binding partners, and characterization of protein structures. These studies will lead to a better understanding of Las pathogenesis, and thereby yield a better control strategy for HLB.

### Materials and Methods

#### Bacterial strains, plants and cultivation

Strains and plasmids used in this study are listed in Table 1. *Escherichia coli* Top10 (Invitrogen, Carlsbad, CA) was used as a host for plasmid construction and *E. coli* BL21 (DE3) cells (Invitrogen, Carlsbad, CA) for recombinant protein expression. *E. coli* was grown in Luria-Bertani (LB) medium at 37°C. *Agrobacterium tumefaciens* strain GV2660 was cultured at 7°C in LB and used to mediate transient expression in the plant leaves of *Nicotiana benthamiana*. Antibiotics were used at the following concentrations: carbencilin, 50 μg/mL; kanamycin, 50 μg/mL.

*N. benthamiana* seeds were stored at 4°C for 2 days prior to germination. Subsequently the seeds were germinated in chambers programmed for cycles of 16 h light and 8 h dark at 26°C. The seedlings were then transferred into FaFard 4P mix soil in plastic containers and grown in controlled greenhouse conditions.

#### Table 1. Strains and plasmids used in this study.

| Strains or plasmids | Description | Source |
|---------------------|-------------|--------|
| **Escherichia coli** |             |        |
| TOPO10              | Chemically competent cells for cloning | Invitrogen |
| BL20(DE3)           | Chemically competent cells for protein expression | Invitrogen |
| **Plasmids in Escherichia coli** |             |        |
| pCR 2.1             | Cloning vector, Amp<sup>r</sup>, Km<sup>r</sup> | Invitrogen |
| pET102D-TOPO        | Expression vector, Cm<sup>r</sup> | Invitrogen |
| pET102-gfp          | pET102D carrying gfp, Cm<sup>r</sup> | This study |
| pET102-lasA<sub>I</sub> | pET102D carrying lasA<sub>I</sub>, Cm<sup>r</sup> | This study |
| pET102-gfp-lasA<sub>I</sub>-TD | pET102D carrying gfp and lasA<sub>I</sub> translocator domain, Cm<sup>r</sup> | This study |
| pET102-gfp-lasA<sub>II</sub> | pET102D carrying gfp and lasA<sub>II</sub>, Cm<sup>r</sup> | This study |
| **Agrobacterium tumefaciens** |             |        |
| GV2660              | Strain for transient expression in the plant | 40 |
| **Plasmids in Agrobacterium tumefaciens** |             |        |
| pGDY                 | GFP transient expression vector, Km<sup>r</sup> | 40 |
| pGDY-lasA<sub>I</sub> | pGDY carrying lasA<sub>I</sub>, Km<sup>r</sup> | This study |
| pGDY-lasA<sub>II</sub>-TD | pGDY carrying lasA<sub>II</sub> translocator domain, Km<sup>r</sup> | This study |
| pGDY-lasA<sub>II</sub> | pGDY carrying lasA<sub>II</sub>, Km<sup>r</sup> | This study |
| p19                 | Gene suppressor, Km<sup>r</sup> | 31 |
Plasmid construction

For *Agrobacterium-mediated* transient expression, the full length *lasA* gene, *lasA* gene and translocator domain (TD) of *lasA*, were amplified using genomic DNA from infected plants using the primers listed in Table 2. The respective PCR products were cloned into a binary vector, pGDY, in which gene expression was under control of a CaMV 35S promoter [40], generating pGDY-*lasA*, pGDY-*lasA*, and pGDY-*lasA*-TD. The recombinant plasmids were verified by sequencing and then transformed into *A. tumefaciens* GV2660 by electroporation.

For protein expression, the GFP-pET-F and GFP-pET-R primers were used for PCR amplification of either GFP alone, the N-terminal GFP fusion with the translocator domain of *lasA*, or the N-terminal GFP fusion with the full length *lasA* from pGDY, pGDY-*lasA*-TD and pGDY-*lasA* plasmids, respectively (Table 2). Full length *lasA* was amplified from infected plant DNA with primers lasA-pET-F and lasA-pET-R (Table 2). PCR products for each gene were ligated into the pET102/D-TOPO vector and transformed into *E. coli* TOP10 cells according to the manufacturer’s instructions (Invitrogen, Carlsbad, CA). Selected clones were cultivated at 37°C in LB broth for plasmid isolation and sequence verification. The clones were designated as pET102-gfp, pET102-*lasA*, pET102-gfp-*lasA*-TD and pET102-gfp-*lasA*.

Purification of LasA

The pET102-*lasA* consensus clone was transformed into *E. coli* BL21 (DE3) expression cells (Invitrogen, Carlsbad, CA). LasA expression was induced and the protein purified using ProBond™ purification system under hybrid conditions according to the manufacturer’s instructions (Invitrogen, Carlsbad, CA). The purified protein was separated using one-dimensional sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and stained with Coomassie blue. Membrane transfer was performed by iBlot according to the manufacturer’s instructions (Invitrogen, Carlsbad, CA). Western blotting was performed with a primary antibody against purified partial LasA (N-terminus, one repeat and part of the translocator domain) from an immunized mouse (ProMab Biotechnologies, Richmond, CA). Goat anti-mouse HRP-conjugated antibody was used as the secondary antibody and detected by chemiluminescence following the manufacturer’s instructions (Life Technologies, Carlsbad, CA).

Protein fraction preparation and Western blot

Whole-cell protein lysates, culture supernatant protein (CS), outer membrane protein (OM) and surface associated protein (AD) were prepared from *E. coli* BL21 containing plasmid pET102-*lasA*, as previously described [11,41–43]. For the CS fraction, culture supernatants were filtered through 0.22 µm pore size filters and concentrated approximately 100-fold by passage through Amicon centrifuge tubes with a molecular mass limit of 50 KDa (Millipore, Billerica, MA). The final pellet was dissolved in SDS-PAGE buffer. The outer membrane protein was isolated on the basis of sarkosyl insolubility [43]. Briefly the cells were collected and broken down by sonication. Total membrane proteins were separated by ultracentrifugation at 28000 rpm for 1hr at 4°C. To obtain the outer membrane protein, the pellet was suspended in 20mM Tris buffer (pH7.4) containing 0.5% sarkosyl and centrifuged again at 28000 rpm for 1hr at 4°C. The final pellet was dissolved in SDS-PAGE buffer. To obtain the AD fraction, the protein secreted but remaining bound to the cell surface, and the cell pellets were suspended in PBS and N-hexadecane was added. The suspensions were vortexed and centrifuged, and the liquid phase was filtered. The proteins were precipitated by acetone and dissolved in SDS-PAGE buffer. Proteins were separated on SDS-PAGE and detected by Western blot as described above.

Immuno-dot blot and proteinase K treatment

Immuno-dot blotting was performed as described previously [39]. The protein expression from bacterial cells containing different constructs was induced as described above. Bacterial cultures were centrifuged at 2,000 g for 10 min at 4°C, washed three times with PBS and adjusted to a final concentration of 0.45 as determined by measurements of the optical density at 660 nm. Three microliters of each serial dilution (1:1, 1:5, 1:10 and 1:20) was spotted onto a nitrocellulose membrane in three replicates. The membrane was air dried and the Western blot procedure was performed as described above. The proteinase K treatment of intact cells was performed in PBS with 10 mM MgCl₂ [44] and then detected by immuno-dot blot. The proteins, either untreated or treated with proteinase K, were recovered for SDS-PAGE analysis and Western blotting.

**Table 2. Primers used in this study.**

| Primer | Sequence |
|--------|----------|
| **Primers for transient expression** | |
| lasA-F | 5’- GCCAGATCTAATTGAAAAATGACG-3’ |
| lasA-R | 5’- ATTCGCTAGTTAGCTACCAAATTAAC-3’ |
| lasA-F | 5’- ATGGATCTAGACT*CAGGACACTAGGAGG-3’ |
| lasA-TD-F | 5’- ATGGATCTAGAGCACTAGAAGG-3’ |
| **Primers for protein expression** | |
| GFP-pET-F | 5’- CACCATGGTGAGCAAGGGCGAGGA-3’ |
| GFP-pET-R | 5’- TTATCTAGATCCGCTGGATCC-3’ |
| lasA-pET-F | 5’- CACCATAGTACAAAGTAAACATGG-3’ |
| lasA-pET-R | 5’- ATAGCTACAAATTAACATCCTTC-3’ |

*Restriction enzyme sites are in italics and underlined*
the same way [31]. The final cell suspension with an OD$_{600}$ of 1.0 was mixed with the p19 suppressor and infiltrated into 4 week-old N. benthamiana leaves with a 1 mL needleless syringe. The experiments were performed with ten independent replicates.

After two days of infiltration, the infiltrated zone was excised and the epidermal layers were peeled for mitochondrial staining with MitoTracker Red CMXRos according to the manufacturer’s protocol (Invitrogen, Carlsbad, CA) and imaged using a confocal laser scanning microscope (CLSM), Zeiss LSM 510. GFP was detected with a 505-530 nm BP filter and MitoTracker Red CMXRos according to the manufacturer’s protocol. The experiments were performed with ten independent replicates. Images were taken with a 505 nm LP filter and 40X or 63X (oil) objectives.

Polar localization by confocal microscope

The E. coli BL21 (DE3) cells containing pET102-gfp, pET102-gfp-lasA$_1$-TD and pET102-gfp-lasA$_2$ constructs were cultured and induced as described above. The collected cells were mounted in antifade solution (Invitrogen, Carlsbad, CA) for CLSM. Images were taken with a 505 nm LP filter and 40X or 63X (oil) objectives.

Transmission electron microscopy (TEM)

Midribs were sampled from healthy and Las-infected pierwinkle. The infiltrated zone was sliced from tobacco. The samples were fixed and sectioned for TEM micrographs as described previously [44].

Supporting Information

Figure S1. LasA1 protein fractions and GFP fusion proteins. A: Western blot of E. coli containing the pET102-

lasA$_1$ construct. Lane 1: whole-cell lysate; lane 2: culture supernatant; lane 3: cell associated protein. B: Western blot of E. coli containing GFP and GFP fusion proteins. Lane 1: E. coli containing plasmid pET102-gfp; lane 2: E. coli containing plasmid pET102-gfp-lasA$_1$-TD; lane 3: E. coli containing plasmid pET102-gfp-lasA$_2$. (TIF)

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Author Contributions

Conceived and designed the experiments: GH YPD. Performed the experiments: GH LZ MB. Analyzed the data: GH. Contributed reagents/materials/analysis tools: GH MB. Wrote the manuscript: GH YPD.

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