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Bacterial Vector-Borne Plant Diseases: Unanswered Questions and Future Directions

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ABSTRACT

Vector-borne plant diseases have significant ecological and economic impacts, affecting farm profitability and forest composition throughout the world. Bacterial vector-borne pathogens have evolved sophisticated strategies to interact with their hemipteran insect vectors and plant hosts. These pathogens reside in plant vascular tissue, and their study represents an excellent opportunity to uncover novel biological mechanisms regulating intracellular pathogenesis and to contribute to the control of some of the world’s most invasive emerging diseases. In this perspective, we highlight recent advances and major unanswered questions in the realm of bacterial vector-borne disease, focusing on liberibacters, phytoplasmas, spiroplasmas, and Xylella fastidiosa.

Key words: liberibacter, phytoplasma, spiroplasma, Xylella fastidiosa, vector-borne disease, effector, plant immunity

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INTRODUCTION

Bacterial vector-borne diseases cause some of the most serious crop losses worldwide. Examples of bacterial vector-borne pathogens include members of the genera Xylella, Candidatus Liberibacter, Spiroplasma, and Ca. Phytoplasma. Whereas X. fastidiosa proliferates in the xylem, liberibacters, spiroplasmas, and phytoplasmas are phloem limited and proliferate in sieve elements (Figure 1). These pathogens are transmitted by different groups of piercing-sucking insects in the order Hemiptera (Jiang et al., 2019) (Figure 2). Citrus, grape, and olive industries have been seriously affected by these bacterial vector-borne pathogens in recent decades. Huanglongbing disease (HLB) caused by Candidatus Liberibacter asiaticus (CLas) is the most damaging disease of citrus, resulting in estimated losses of over $7.8 billion in Florida since 2007 (Court et al., 2018), and the phytoplasma disease lime witches’ broom threatens the lime industry in the Middle East (Donkersley et al., 2018). Moreover, coconut phytoplasmas have destroyed millions of palms for centuries and are spreading locally (Gurr et al., 2016). Grape production has been significantly affected by Pierce’s disease of grapevine (X. fastidiosa), resulting in an annual cost of approximately $100 million in California alone (Tumber et al., 2014). Recently, European olive orchards have been devastated by the emerging disease olive quick decline syndrome caused by X. fastidiosa (Schneider et al., 2020). In the absence of disease control measures, the total economic loss to the olive industry is estimated to reach up to €5.2 billion in Italy and €16.86 billion in Spain over the next 50 years (Schneider et al., 2020).

Bacterial vector-borne disease research is challenging because of fastidious pathogen growth, the inability to culture multiple species, reliance on an insect vector for transmission, and the lack

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of model systems (Jiang et al., 2019). However, deciphering the interaction between pathogen, vector, and plants in diverse systems provides new opportunities to discover novel biology that regulates intracellular pathogens and cannot be addressed in more tractable systems. Transmission of bacteria directly into vascular tissues, such as the xylem or phloem, may be an adaptation that allows these pathogens to evade plant immune perception. Their virulence mechanisms are probably different from those of other well-characterized Gram-negative bacteria and provide unique opportunities to uncover novel biological insights (Toruno et al., 2016; Kruse et al., 2019). A greater understanding of pathogen host and vector manipulation, as well as plant responses, is critical for the effective control of vector-borne disease. Here, we highlight major advances and unanswered questions in bacterial vector-borne plant disease and provide a roadmap for future research directions.

WHAT ARE THE MINIMAL REQUIREMENTS FOR LIFE?

Bacterial Vector-Borne Pathogenicity Has Evolved at Least Four Times

Considering their small genome size, limited metabolic capacity, and life cycle that involves two hosts, vector-borne bacteria have an important role to play as models for study in research fields as varied as synthetic biology, evolution, phytopathology, and biochemistry. Genome reductions are predominantly caused by loss of metabolic genes, making these bacteria dependent on the acquisition of diverse metabolites from their hosts via various transporters, predominantly ABC transporters (Kube et al., 2012; Oshima et al., 2004).

Ca. Liberibacter and X. fastidiosa are both Gram-negative bacteria but belong to different orders, i.e., Rhizobiales (class Alphaproteobacteria) and Xanthomonadales (class Gammaproteobacteria), respectively (Figure 2). Spiroplasmas and phytoplasmas both belong to the class Mollicutes. However, spiroplasmas and phytoplasmas have different codon usages, and spiroplasmas share a common ancestry with mycoplasmas and phytoplasmas with the saprotrophic acholeplasmas (Gundersen et al., 1994; Gupta et al., 2019) (Figure 2). Spiroplasmas and phytoplasmas are both Gram-negative bacteria but belong to different orders, i.e., Rhizobiales (class Alphaproteobacteria) and Xanthomonadales (class Gammaproteobacteria), respectively (Figure 2). Spiroplasmas and phytoplasmas have different codon usages, and spiroplasmas share a common ancestry with mycoplasmas and phytoplasmas with the saprotrophic acholeplasmas (Gundersen et al., 1994; Gupta et al., 2019) (Figure 2).
insect transmission evolved independently multiple times. In all four cases, their evolution involved genome reductions and the retention and acquisition of virulence genes that enabled them to colonize animal and plant hosts.

**Bacterial Vector-Borne Pathogens Have Distinct Metabolic Requirements**

The four taxa of vector-borne plant pathogenic bacteria exhibit signatures of differential genome reduction rates, ranging from 2.5 Mb for *X. fastidiosa* to ±1.7–1.2 Mb for *Spiroplasma* and *Liberibacter* to <1 Mb for *Phytoplasma*. *X. fastidiosa* and *S. citri* are fastidious but can be cultured and genetically modified (Bové, 1997; Bové and Garnier, 2003; Perilla-Henao and Casteel, 2016). By contrast, *Ca. Liberibacter* and *Phytoplasma* have largely resisted transformation (Ha et al., 2019; Naranjo et al., 2019). Thus, it is likely that each pathogen relies on its own set of genes for growth, and culturability is not necessarily associated with genome size but rather with genes that were lost and are required for life in vitro. For example, mycoplasmas with genome sizes similar to those of phytoplasmas were cultured in rich artificial medium (Tully et al., 1977). Phytoplasmas lack different metabolic genes compared with other bacteria in the class Mollicutes (Chen et al., 2012). For instance, compared with mycoplasmas and *S. citri*, phytoplasmas lack a phosphotransferase system for the import of sugars and seem to import disaccharides by means of ABC transporter systems, import malate instead of lactate, and lack genes for generating their own ATP (which must be sequestered from insect and plant hosts) (Oshima et al., 2004; Bai et al., 2006; Kube et al., 2014).

**IMPORTANCE OF TISSUE TROPISM FOR PLANT COLONIZATION**

Tropism refers to the ability of a given organism to move and colonize a specific tissue. Research in vector-borne disease presents unique opportunities to investigate vascular patterning, vascular
defense, and novel components that regulate pathogen movement. Some insect vector-borne plant viruses encode an active strategy in the viral genome to maintain phloem localization, and this strategy may facilitate dispersal by phloem-feeding insect vectors and/or evasion of plant host immune responses (Peter et al., 2009).

Pathogen localization in a host is probably a combination of vascular flow, host genotype, and possibly bacterial preference for a tropic region (Figure 1). Accumulation in sink tissues, including roots and growing tissues, is often observed in phloem-limited pathogens; however, their distribution within the plant is variable (Figure 1). This distribution extends to individual sieve tubes in the phloem where, for example, CLas massively proliferates in some tubes while barely colonizing others (Achor et al., 2016; Hartung et al., 2010), and spiroplasmas accumulate near undifferentiated phloem cells (Bové et al., 2003). Recovery of apple trees from apple proliferation phytoplasma is associated with increased callose depositions that prevent recolonization of the apple tree crowns during the spring and with enhanced overall defense responses of recovered apple trees (Mussetti et al., 2010). Characterization of CLas and X. fastidiosa localization in citrus and related genotypes indicates that replication and establishment are correlated with the host’s ability to resist infection, leading to lower pathogen populations in genotypes that exhibit some resistance (Coletta-Filho et al., 2020; Ramadugu et al., 2016). X. fastidiosa exhibits differential tropism in the xylem of susceptible and resistant citrus genotypes; it is able to move to secondary xylem in susceptible genotypes but is confined to the primary xylem in resistant genotypes (Figure 1) (Niza et al., 2015). We lack a mechanistic understanding of how host genetic variability affects pathogen colonization and tropism. However, studies have correlated, for example, pit membrane carbohydrate composition with the ability of X. fastidiosa to colonize different grapevine varieties (Ingel et al., 2019). It is evident that individuals in a population respond differently to infection and that disease pressure and the environment play a role in mediating host responses. The composition of microbial communities can also influence disease. For example, the composition and diversity of citrus leaf and root microbiota were associated with HLB symptom severity (Blaustein et al., 2017).

To understand how the host regulates and responds to pathogen colonization, tools must be developed to investigate vascular biology; these may include inducible expression systems, cell-specific promoters, and robust silencing and gene-editing platforms. Recent advancements in phloem-limited viral vectors that enable expression of small genes and RNAs, such as Citrus tristeza viral vectors, hold promise for testing the importance of specific loci (Dawson et al., 2015). The development of advanced microscopy techniques will enable researchers to investigate bacterial movement during the course of infection and colonization across different pathogen strains and host genotypes (Newman et al., 2003). These tools will enable the identification of key loci important for vascular responses, including those that affect vascular patterning, limit pathogen distribution, and affect tropism. A greater understanding of how pathogens are distributed holds great promise for disease control by facilitating early detection of diseased plants, a task that is complicated due to uneven pathogen distribution. Early and efficient detection of bacterial pathogens that infect perennial woody plants, such as CLas, X. fastidiosa, S. citri, and Ca. Phytoplasma palmicola, will enable the removal of infected plants that serve as a reservoir for pathogen spread. A detailed understanding of vascular biology is also needed to develop delivery strategies for therapeutic molecules to control bacterial infection in plants.

Pathogen Movement and Host Colonization

Phloem-limited bacteria must rely on intracellular communication to move and colonize different sections of the phloem. The late systemic spread in sieve elements colonized by CLas and phytoplasmas coupled with the slow seasonal migration of Ca. Phytoplasma mali (apple proliferation phytoplasma) from roots to new apple flush suggest that these pathogens can also move against the phloem flow, albeit at a slow rate (Hilf and Luo, 2020; Schaper and Seemüller, 1984; Wei et al., 2004). Phytoplasmas lack genes coding for cytoskeletal elements or flagella (Music et al., 2019; Oshima et al., 2004). CLas flagellar genes are repressed in planta, and flagellin expression can be detected only in the vector (Andrade et al., 2020; Yan et al., 2013). On one hand, motility mutants of S. citri that display only twitching movement are not impaired in pathogenicity. On the other hand, loss of helicity and motility seems to occur naturally at a very low rate in this pathogen (Duret et al., 2003; Fletcher et al., 2006). In spiroplasmas and other mollicutes, specialized tip structures consisting of exposed surface proteins are important pathogenicity determinants required for host membrane attachment and pathogen movement (Ammar et al., 2004). Similar surface proteins were also found in phytoplasmas and may play an analogous role (Kakizawa et al., 2006a, 2006b). Liberibacters and mollicutes use their pleomorphic shape to cross the tight space between sieve plates, even when they are fortified by callose (Figure 1) (Achor et al., 2020) (Waters and Hunt, 1980). Phytoplasmas have been shown to adhere to the inner surface of the sieve tube membranes (Marcone, 2009), and liberibacters often aggregate near sieve pores (Achor et al., 2020), a feature that may facilitate movement (Figure 1).

The ability of the Xanthomonadaceae family to colonize the xylem has been linked to the presence of CbsA, a cell-wall-degrading cellobiohydrolase (Gluck-Thaler et al., 2020). Deletion of CbsA in non-vector-borne vascular Xanthomonas oryzae pv. oryzae increased apoplastic colonization and symptom development. By contrast, X. fastidiosa cbsA mutants were severely reduced in vascular colonization and symptom development (Gluck-Thaler et al., 2020). CbsA expression correlates with the planktonic lifestyle and cell proliferation of X. fastidiosa, and it is also highly expressed in biofilm-deficient mutants (Gouran et al., 2016). Tracking studies of fluorescent-tagged cell-degrading enzymes, including CbsA, permit the study of movement in degradative enzyme mutants and will be required to understand the role of these enzymes in bacterial movement from primary to secondary xylem and between xylem vessels.

Our inability to culture Ca. Liberibacters and phytoplasmas present a significant challenge to the identification of important loci required for host or vector tropism. Other than Liberibacter crescentis BT-1, no liberibacters have been obtained in pure culture despite extensive efforts (Ha et al., 2019; Naranjo et al., 2019). Spiroplasmas represents an excellent model for the investigation of phloem-limited bacteria because of its culturability, broad host range, and potential for genetic manipulation (Renaudin
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et al., 2014). The development of high throughput genetic screens such as those that use TNseq will quickly advance our understanding of general mechanisms of host colonization in phloem-limited bacteria. The structure of the plant phloem may play a role in CLas acquisition and transmission by the psyllid vector by acting as a determinant for the establishment of plant feeding sites (George et al., 2017). Major unanswered questions also include the influence of genetic variation on pathogen tropism. Do tropism patterns observed under laboratory conditions with defined bacterial isolates reflect field conditions? What are the roles of pathogen and host genetic diversity in the colonization of particular vascular niches?

PLANT DEFENSE RESPONSE TO VECTOR-BORNE DISEASES

Plants possess sophisticated immune systems capable of perceiving conserved pathogen, damage, and insect features using surface-localized receptors and intracellular nucleotide-binding leucine-rich repeat (NLR) immune receptors that recognize secreted pathogen and insect proteins (Boissot et al., 2016; Boutrot and Zipfel, 2017; Lolle et al., 2020). Immediate plant defenses are triggered by vector feeding. Hemipteran insects secrete watery saliva as they probe into the apoplast and different cell types to acquire cell contents until the stylet reaches the phloem (Jaouannet et al., 2014). Insect effectors secreted during cell penetration can be perceived by NLR immune receptors (Boissot et al., 2016). Watery saliva contains insect features (herbivore-associated molecular patterns [HAMPs]) as well as microbial features from bacterial pathogens or symbionts (microbial-associated molecular patterns [MAMPs]) (Chaudhary et al., 2014; Prince et al., 2014). Honeydew is also known to contain HAMPs and MAMPs and to activate immunity (Schwartzberg and Tumlinson, 2014; Wari et al., 2019). These features are perceived by surface-localized pattern recognition receptors, resulting in the induction of plant immune responses (Chaudhary et al., 2014; Jaouannet et al., 2014; Steinbrenner et al., 2019).

The styles of hemipteran insects induce defense responses and Ca2+ bursts during probing (Vincent et al., 2017), and watery saliva also contains hemipteran effectors that can suppress immune responses, illustrating the importance of plant immunity in vector-borne disease (Mugford et al., 2016). HAMP and MAMP release during initial vector probing can activate immune responses that affect the phloem environment. The phloem itself is an important highway for systemic signaling in response to wounding and immune perception, which result in rapid waves of Ca2+ (Toyota et al., 2018). Many questions remain concerning innate immunity in the phloem. Do canonical plant immune receptors function in phloem tissues? Can bacterial effector proteins that move cell-to-cell via plasmodesmata be recognized by intracellular plant immune receptors?

Common phloem defense responses include the production of Ca2+, callose deposition, and activation of phloem-specific forisomes and P proteins (Figure 1) (Jiang et al., 2019). Callose deposition in sieve pores is a common response to infection by phloem-limited pathogens. CLas infection induces the expression of multiple citrus callose synthases, including CalS3, CalS7, CalS8, CalS9, CalS11, and CalS12 (Granato et al., 2019). Callose hydrogel complexes have distinct biophysical properties that contribute to elasticity, which may account for the inability of callose to impair mass flow (Hunter et al., 2019; Abou-Saleh et al., 2018). Activation of Ca2+-dependent sieve proteins, including Sieve Element Occluding Relatives (SEOR) P proteins and forisomes, occurs during vector feeding (Figure 1) (Knoblauch et al., 2014). SEOR proteins form highly organized aggregates that disperse upon Ca2+ activation, surround the site of vector feeding, and block phloem acquisition to play an important role in resistance against phloem-feeding insects (Garzo et al., 2018; Peng and Walker, 2020). Surprisingly, Ca. Phytoplasma asteris exhibits reduced growth in Arabidopsis seor1 mutants, indicating that AtSEOR1 may also be a susceptibility gene (Pagliari et al., 2017). Development of genetic tools for investigating phloem responses will be necessary to provide clear genetic evidence linking phloem-specific defenses with disease resistance.

A common xylem defense response is the formation of tyloses, which are extensions of living parenchyma cells. Despite being a common and visible defense response, tylose formation occurs late in infection and fails to prevent bacterial movement in the X. fastidiosa–grape interaction; instead, it exacerbates disease symptoms (Stevenson et al., 2004). Both intracellular and surface-localized immune receptors that recognize bacterial and fungal xylem-colonizing pathogens such as Fusarium oxysporum and Ralstonia solanacearum have been cloned (Narasaki et al., 2009; Takken and Rep, 2010). Both Fusarium and Ralstonia must invade the root before reaching the xylem. There is evidence that immune perception is important in the activation of plant responses to X. fastidiosa infection, but perception is delayed by the presence of cell surface lipopolysaccharides (Rapicavoli et al., 2018).

CRITICAL PATHOGENICITY DETERMINANTS

Bacterial Vector-Borne Pathogens Have Different Retained and Acquired Virulence Capacities

Bacterial vector-borne pathogens appear to have differentially retained or acquired virulence genes. X. fastidiosa colonizes the plant xylem and the foregut of its insect vectors, which are xylem sap-feeding leafhoppers (sharpshooters) and spittlebugs,
through biofilm formation on the cuticular surface of the foregut lumen (Figure 2) (Killiny et al., 2013; Ionescu et al., 2016). Virulence factors involved in X. fastidiosa migration, rather than in biofilm formation in the xylem and the insect, are homologs of proteins from related free-living and plant pathogenic Xanthomonads (Almeida et al., 2012; Caserta et al., 2017). The latter possess type III secretion systems (TTSSs) for the penetration of plant cell membranes and the deposition in cells of an arsenal of protein effectors that modulate plant processes, but such systems are lacking in the X. fastidiosa genome (Lu et al., 2008; Toroño et al., 2016). Presumably, being xylem limited, X. fastidiosa does not require TTSSs and associated effectors, although it possesses other systems for the secretion of virulence factors that may unload from the xylem and may differ from those of plant pathogenic Xanthomonads (Lu et al., 2008).

In contrast to X. fastidiosa, Ca. Libriferi species, phytoplasmas, and spiroplasmas plant pathogens all colonize the phloem and, moreover, are invasive colonizers of their insect vectors. Ca. Libriferi are transmitted by phloem-feeding psyllids, spiroplasmas by phloem-feeding leafhoppers, and phytoplasmas by phloem-feeding leaffoppers, planthoppers, and psyllids (Figure 2). All three bacteria can be pathogenic to their insect vectors as well as to their plant hosts (Bövé, 1997; Marina Mann et al., 2018; Koimura et al., 2020). S. citri takes advantage of uncommon post-translational mechanisms to increase the antigenic diversity at its surface, which may play a role in insect colonization by this mollicute (Dubrana et al., 2017). Moreover, the virulence genes of phytoplasmas lie within large pathogenicity islands of ± 20 kb that are comprised of composite transposon-like elements and named potential mobile units (PMUs) (Bai et al., 2008). These PMU-encoded virulence genes are differentially expressed between phytoplasmas in the plant and vector (Toroño et al., 2010). The pathogenicity islands are prone to frequent mutations and deletions, recombine with each other, and can form extrachromosomal units, making phytoplasma genomes highly unstable but probably also giving rise to more opportunities for adaptation (Bai et al., 2006; Arashida et al., 2008; Music et al., 2018). PMU genes include effectors that modulate plant development (discussed below). There is evidence of horizontal DNA transfer, including that of PMU-like elements, among distantly related phytoplasmas and between spiroplasmas and phytoplasmas (Bai et al., 2004; Arashida et al., 2008; Music et al., 2019). Spiroplasmas genomes are also highly repeat rich; the repeats include phage-like elements and multiple plasmids with repeat-rich adhesin-like genes that are involved in virulence (Beven et al., 2015). Taken together, these findings suggest that there are ongoing acquisition and recombination of genomic virulence clusters that have integrated into the minimal genomes of vector-borne plant pathogens.

However, most interesting for pathologists are the minimal requirements for pathogenesis. The minimal requirement for pathogenesis appears to be defined by genomic areas different from those involved in metabolism, although mechanisms that detect the processing of specific metabolites may be part of a larger machinery that controls the regulation of virulence gene expression. A genomic area that encodes virulence factors, such as a phytoplasma PMU, may be isolated and integrated into another culturable bacterium to assess mechanisms that enable the colonization of plants versus insect vectors.

**Effectors Biology**

The ability to secrete proteins, called effectors, is required for diverse organisms to colonize a host and/or cause disease (Toroño et al., 2016). Among bacterial vector-borne pathogens, phytoplasma effectors are well characterized. Although phytoplasmas are restricted to the phloem, effectors secreted into the cytoplasm of sieve cells can be unloaded from the phloem and move systemically, possibly via plasmodesmata (Bai et al., 2009; Hoshi et al., 2009; MacLean et al., 2011a). Phytoplasma effectors have dramatic impacts on plant development and induce several distinct symptoms characteristic of diseases. For example, SAP11 effectors from diverse phytoplasmas induce a range of phenotypes when stably expressed in plants, including crinkled leaves and silique, increased stem proliferation, altered root architecture, and abnormal glandular trichome development (Sugio et al., 2011b; Lu et al., 2014; Tan et al., 2016; Chang et al., 2018; Wang et al., 2018; Pecher et al., 2019). The role of these diverse plant phenotypes induced by phytoplasma effectors remains an important subject of future research.

Effectors targeting of transcription factors is a strategy for targeting multiple plant processes simultaneously. SAP11 proteins differentially interact with and destabilize members of the plant TEOSINE BRANCHED 1, CYCLOIDEA, PCF1 (TCP) transcription factor family, which regulates diverse plant developmental processes. Another phytoplasma effector, SAP54/phylogen, is responsible for the conversion of plant flowers into leaf-like tissues, a phenomenon known as phyllody that is commonly seen in infected plants (MacLean et al., 2011a, 2014; Maejima et al., 2014). SAP54/phylogen targets plant homoeotic MADS-box transcription factors that control floral development for destabilization through the host 26S proteasome (MacLean et al., 2014). A third phytoplasma effector, TENGU, induces dwarfism, stem proliferation, and sterility in plants by unknown mechanisms that suppress auxin and jasmonic acid signaling (Hoshi et al., 2009; Minato et al., 2014). Phytoplasma effectors appear to manipulate the entire host to their advantage. Not only do they maintain newly infected tissues in a juvenile or vegetative state that facilitates nutrient acquisition, but they also render plant tissues more attractive or less hostile to vectors that feed on infected plants (Sugio et al., 2011c; Orlovskis and Hogenhout, 2016; Tomkins et al., 2018).

In contrast to phytoplasmas, Ca. Libriferi effectors are underexplored, and Spiroplasma effector functions remain completely elusive. Effectors from Ca. Libriferi can inhibit immune-related protease activity (SDE1), suppress host cell death (HPE1), and suppress ROS production through peroxidase activity (Jain et al., 2015; Clark et al., 2018; Levy et al., 2020). It is likely that the immune-suppressing activity of libriferi effectors is just the tip of the iceberg. For example, the Ca. Libriferi effector SDE1 is multifunctional, localizes to diverse subcellular compartments, induces cell death in the model plant Nicotiana, and targets diverse host genes (e.g., DEAD-box RNA helicase DDX3 and immune proteases) to manipulate different plant responses (Pitino et al., 2016; Clark et al., 2018; Zhou et al., 2020). The moonlighting properties of SDE1 are reminiscent of viral proteins, which are capable of manipulating multiple cellular processes simultaneously through interactions with diverse host targets (DeBlasio et al., 2016).
Effectors from vector-borne pathogens exhibit variation in expression across host and vector and are influenced by plant genotype. Liberibacters and phytoplasmas deploy specific suites of effectors to differentially manipulate each organism and enable the switch between vector and host (Thapa et al., 2019; Shi et al., 2019). Filamentous pathogens spatially and temporally regulate suites of effectors during different infection stages (Torun˜o et al., 2016). Liberibacters, phytoplasmas, and spiroplasmas can infect perennial plants, including trees, residing in their hosts for years. Changes in effector expression over the course of long-term infection is an unexplored area. Is effector expression influenced by host, pathogen, and vector genotype? To what extent do bacterial effector proteins regulate tissue tropism during infection?

Functional characterization of effector candidates is key to understanding the tripartite interaction between plants, bacteria, and insects (Figure 3). Robust effector prediction pipelines will enable more accurate identification of effector candidates. Effectors from different vector-borne bacteria probably have divergent roles compared with those from apoplastic colonizing pathogens. The first challenge in characterizing effector function is to determine where these proteins function inside the host. For instance, both SAP11 and TENGU were demonstrated to accumulate in different cell types beyond the phloem using immunohistochemical analysis (Bai et al., 2009; Hoshi et al., 2009). The CLas SDE1 effector can also be detected in plant tissues by immunoblot when bacteria are undetectable by PCR (Pagliaccia et al., 2017). Depending on their localization, effectors can directly or indirectly affect bacterial growth and colonization. Because most bacterial effectors are small, it remains difficult to visualize effector localization and movement in vivo. Large tags, such as fluorescent proteins, may interfere with effector localization, movement, or function. In this regard, the development of new fluorescent-labeling methods with smaller tags for visualizing effector proteins in the hosts is highly desirable (Figure 3).

A second challenge lies in the identification of effector targets. Because the host range of some pathogens can be quite specific, the tools and resources for identifying effector targets may not be as readily available as in model organisms such as Arabidopsis. One potential solution is the establishment of more pathosystems or surrogate systems for studying vector-borne bacterial diseases. For example, Nicotiana has been used to screen for CLas effectors that induce cell death and could therefore be useful in the identification of plant receptors for engineering resistance in citrus (Pitino et al., 2016) (Figure 3). The next challenge is to understand how effectors interfere with host processes and facilitate pathogen spread. As mentioned above, the reprogramming of plant development by phytoplasma effectors SAP11 and SAP54 also modulates plant–insect interactions and promotes insect vector attraction and fecundity (Sugio et al., 2011a; MacLean et al., 2014; Orlovskis and Hogenhout, 2016). However, whether bacterial effectors modulate plant–insect interaction more generally requires further exploration. Moreover, current studies on vector-borne bacteria largely focus on the function of an effector within one organism, either a plant or insect. It remains to be seen how effector-triggered changes in one organism may interfere with processes in another, resulting in effective disease spread.

**VECTOR COMPATIBILITY AND VECTOR IMMUNITY**

**Does the Vector Actively Limit Pathogen Colonization and Tropism?**

The most important hemipteran vectors that transmit bacterial pathogens are members of Sternorrhyncha, including psyllids (Psylloidea) and Auchenorrhyncha, including leafhoppers (Membracoidea), froghoppers/spittlebugs (Cercopoidea), and planthoppers (Fulgoroidea) (Figure 2) (Bove and Garnier, 2003; Perilla-Henao and Casteel, 2016; Sicard et al., 2018). Vector-
borne bacterial pathogens depend on their insect vectors for spread and transmission to plants, and insect-bacteria interactions are as intricate as those between plant and bacteria. Some pathogens provide fitness benefits to their vectors; however, neutral or positive effects are not always observed. Similar to the plant host, the tissue tropism of the pathogen within the vector defines the mode of insect transmission.

Transmission occurs in distinct phases: acquisition of the bacterium from infected plants, translocation through the body tissues (which includes bacterial propagation; X. fastidiosa has a different mode of insect colonization), and inoculation of a new recipient host plant (Ammar et al., 2011; Heck and Brault, 2018; Koinuma et al., 2020). The anterior midgut of the alimentary canal and type III cells of salivary glands were identified as the major sites of onion yellows phytoplasma infection in a leafhopper vector (Koinuma et al., 2020). For CLas acquisition, the insect developmental stage is critical, but how this relates to tissue tropism is not well understood. CLas must be acquired during the nymphal stage for adults to transmit efficiently (Inoue et al., 2009; Ammar et al., 2016; George et al., 2018). Moreover, young citrus tissue (called flush) where the nymphs hatch and develop becomes infectious just 15 days after inoculation with CLas (Lee et al., 2015). Thus, asymptomatic spread coincides with the vector’s reproductive cycle. This pattern of spread underscores the need for research on early detection and removal of infected plant material to prevent disease outbreaks.

Molecular methods, such as in situ hybridization, confocal microscopy, electron microscopy, and qPCR, have been used to localize bacteria to specific insect tissues (Ammar et al., 2011; Ghanim et al., 2016, 2017; Kruse et al., 2017; Mann et al., 2018) and have identified apoptosis as involved in the developmental regulation of CLas acquisition. The gut is a physical barrier to acquisition and the first site of proliferation for propagative pathogens (Figure 2). In adult D. citri, CLas induces extreme apoptosis and nuclear disruption in the midgut and malpighian tubules, whereas nuclei in foreguts and hindguts appear normal (Ghanim et al., 2016; Mann et al., 2018). Comparative transcriptome analyses of CLas infected and uninfected D. citri midguts revealed extensive transcriptional changes, including multiple differentially expressed genes involved in the ubiquitin-proteasome and immune system (Yu et al., 2020). Collectively, these data highlight extensive cellular and transcriptional changes in adult D. citri midguts in response to CLas acquisition.

Whereas adult D. citri midguts are severely affected by CLas, nymphal midgut nuclei are not affected (Mann et al., 2018). Evidence for midgut cell apoptosis as a barrier to CLas acquisition by adult D. citri comes from comparative studies on Bactericera cockerelli, the potato psyllid. In this psyllid, Ca. Liberibacter solanacearum (CLso) does not cause apoptotic nuclear disruption in any cells of the gut of adult psyllids, and adult psyllids are able to acquire and transmit CLso (Tang and Tamborinedeguy, 2019). Development of strategies that induce apoptosis in the psyllid midgut, by manipulation of caspase genes, for example, may be a novel means to control acquisition and transmission of liberibacters by psyllids. For example, the legume lectin concanavalin A has been reported to induce apoptosis in the potato psyllid midgut when ingested from an artificial diet (Tang et al., 2020).

The Relationship between Vector-Borne Pathogens and Their Vectors

Studies on the coevolution of two stunting mollicutes, the Dalbulus leaffopper spp., and maize-related grasses eloquently highlight how host associations in nature have favored positive relationships between plant pathogenic bacteria and their insect vectors. Maize bushy stunt phytoplasma (MBSP) and corn stunt spiroplasma (S. kunkelli) are two major maize stunting pathogens primarily transmitted by two maize specialists, Dalbulus maidis and D. elimatus. These two species belong to a larger genus that also includes D. gelbus and D. quinquenotatus, which feed on teosinte and gamagrass. Although these insects can be force-fed on maize, in nature they are rarely found in maize. When D. gelbus and D. quinquenotatus are infected by MBSP or S. kunkelli, the insects often die before or at the time of pathogen transmission to maize. By contrast, D. maidis and D. elimatus seem to be less affected by bacterial colonization and in some cases even benefit from the infection (Madden et al., 1984; Nault et al., 1984; Moya-Raygoza and Nault, 1998). Therefore, Dalbulus spp. that have been exposed to MBSP and S. kunkelli for a longer time are likely to develop better tolerance than those that have had little exposure to these bacteria.

A bacterium’s association with specific vector(s) probably determines not only the tissue where it becomes established in vector and host, but also its host range. Liberibacter species associate exclusively with psyllid vectors, and both vector and bacteria associate with a limited number of hosts (Figure 2). By contrast, phytoplasmas infect nearly 200 plant species and can be transmitted by at least 30 polyphagous species of phloem-feeding insects, including psyllids, leaffoppers, and planthoppers (Weintraub and Beanland, 2006; Ku et al., 2013) (Figures 1 and 2). The host range of spiroplasmas is more restricted than that of phytoplasmas, and spiroplasmas are transmitted by several species of leaffoppers (Figures 1 and 2). The case of X. fastidiosa is substantially different. There is no evidence of X. fastidiosa–vector specificity, meaning that insects in the abovementioned groups should be considered vectors until proven otherwise (Sicard et al., 2018). Most xylem sap-sucking insects are generalists, and this may be linked to the ability of X. fastidiosa to colonize hundreds of host plant species, usually without causing disease (Sicard et al., 2018).

GOING BEYOND MINIMAL SYSTEMS: HOW DOES THE GENETIC DIVERSITY OF THE PATHOGEN, VECTOR, AND HOST INFLUENCE DISEASE?

Technical limitations, including culturability, insectary infrastructure, and a lack of studies on biology under field settings, have led to important gaps in our understanding of bacterial vector-borne disease. How diverse are bacterial and vector populations in the field and natural ecosystems? How does this diversity affect epidemiology and capacity to cause plant disease? Can laboratory findings be translated to the field?

Vector Diversity

Natural variation in the ability of insect vectors to transmit plant and animal pathogens has been demonstrated extensively in many
species, including the psyllid *D. citri*, which transmits CLas. This natural variation has been used as a powerful tool to dissect the genetic and molecular mechanisms of vector competence and vector–pathogen interactions (Guo et al., 1996; Habekuss et al., 1999; Terradot et al., 1999; Bencherki et al., 2000; Lucio-Zavaleta et al., 2001; Burrows et al., 2006, 2007; Gray et al., 2007; Yang et al., 2008; Cilia et al., 2011; Ogada et al., 2016; Ammar et al., 2018). Variability in transmission efficiency has also been found for phytoplasmas and spiroplastas (Gonella et al., 2019). For phytoplasmas, the interaction between phytoplasma membrane proteins and insect factors determines vector specificity (Suzuki et al., 2006). It is well known that pathogens encounter barriers to transmission within the insect vector, and movement of pathogens across the gut and salivary tissues is also variable across populations and is genetically regulated (Ammar et al., 2018). Future research should leverage the natural variation in *D. citri* populations for CLas acquisition and transmission (Coy and Stelinski, 2015; Ammar et al., 2018; Hall, 2018), genomic resources for *D. citri* (Hosmani et al., 2019), and genome-wide association studies to identify psyllid genes that regulate CLas acquisition and transmission. By understanding individual-level diversity instead of generalizing across populations, we can also begin to understand why lab-generated management strategies may not always work in the field.

Although vector diversity is likely to significantly impact the vector transmission efficiency of phloem-limited bacterial pathogens, the role of within-species vector diversity in *X. fastidiosa* transmission is unknown and, if existing, likely to be linked to vector behavior and ecology. This may be explained by *X. fastidiosa* colonization of the cuticular surface of the foregut, which is a conserved substrate among these insect groups. *X. fastidiosa* does not interact with live insect host cells, such as the phloem-limited bacteria discussed here (Sicard et al., 2018). Evidence has shown that an array of factors—including host tissue preferences in relation to pathogen distribution within plants (Daugherty et al., 2011)—influence transmission rates and among-species differences in vector transmission of *X. fastidiosa* (Sicard et al., 2018), although evidence of within-species differences is so far absent or limited (Almeida and Purcell, 2003; Lopes et al., 2009; Krugner et al., 2012).

**Pathogen and Host Diversity**

Molecular tools facilitate the investigation of bacterial diversity. However, much of this work has been limited in scope, biased toward agricultural crops, and reliant on a limited number of genetic markers. The widespread advent of next-generation sequencing tools has enabled whole-genome sequences to be generated. Because there are more sequences available for *X. fastidiosa* (~350 isolates at the time of writing), we focus on this pathogen to illustrate aspects of pathogen diversity that require investigation.

There are currently two species of *Xylella*, the widely studied *fastidiosa* and the less-known *taivawensis*. Because *X. fastidiosa* studies have focused on diseased crops, we do not understand what diversity may exist within *X. taivawensis* or *X. fastidiosa* in natural ecosystems or whether there are other species of *Xylella* (Almeida and Nunney, 2015). It is unreasonable to assume that the only *Xylella* not considered to be endemic to the Americas is found in an island in Asia; even within the Americas, there is evidence of unsampled clades of *X. fastidiosa* (Castillo et al., 2020). Similarly, in the Americas, and more recently in Europe, the vast majority of *X. fastidiosa* for which there are genetic data are linked to crops of economic importance. *X. fastidiosa* genetic diversity can be found within regions, orchards, trees, and even branches (Coletta-Filho et al., 2014). There is little knowledge of what this variation, particularly at the within-crop and local spatial levels, may mean biologically. Finally, when extrapolated to other vector-borne plant pathogenic bacteria, these knowledge gaps only become more extreme.

Available genomic data are now beginning to be used for an array of evolutionary and ecological questions that were previously impossible to address. Genome sequences provide sufficient genetic resolution to study recent pathogen dispersal pathways, with implications for a greater understanding of pathogen biology and clues for how to avoid future introductions (Landa et al., 2019). Similarly, genomic data have elucidated how endemic pathogens have adapted to an introduced insect vector, leading to a novel phytoplasma disease (Malembic-Maher et al., 2020). Plant pathogen population genomic studies, including one focused on *X. fastidiosa* that colonizes grapevines in California, are resulting in novel interpretations of the geographical structure of pathogens in the landscape, as well as possible adaptive changes associated with temperature gradients (Vanhove et al., 2020). We expect this emerging field to continue yielding novel and relevant insights into the biology of vector-borne plant pathogens, but the difficulty of obtaining large populations of clonal isolates for genome sequencing remains.

Variations in host responses are also an important component of vector-borne disease. For example, CLas can proliferate on multiple plant hosts within the subfamily Aurantioidae to which citrus belongs. While all cultivated citrus is susceptible to HLB, other members of the Aurantioidae subfamily exhibit a gradient of responses (Ramadugu et al., 2016). Some genotypes are not suitable hosts for *D. citri* (white sapote, *Casimiroa edulis*), others lack high CLas populations despite being preferred hosts for the psyllid (*curry leaf, Bergera koeniggi*; pink wampee, *Clausena excavata*), and some exhibit tolerance to HLB despite higher CLas populations (orange jasmine, *Murraya paniculata*) (Westbrook et al., 2011; Ramadugu et al., 2016). Similar variability in plant responses to *X. fastidiosa* infection exists. For example, although all sweet orange varieties are susceptible to *X. fastidiosa* infection, leading to citrus variegated chlorosis, varieties of lemon, mandarin, grapefruit, and other citrus fruits are tolerant or resistant (Coletta-Filho et al., 2020). European grapevine varieties (*Vitis vinifera*) express a range of susceptibility to *X. fastidiosa* (Rashed et al., 2013), but many American hybrids and species of *Vitis* are tolerant to this pathogen (Krivanek et al., 2005).

Insect–plant–pathogen interactions at the ecological level and their consequences for disease incidence and pathogen spread are poorly understood. The main challenges are associated with the need to extrapolate controlled experimental manipulations to the field or landscape scales. This is usually performed with mathematical modeling, yet experimental work on these interactions is rarely designed with models in mind. Thus, captured data are often inadequate for downstream inferences. Similarly, manipulations are usually performed at single time points and may not measure the role of vector choice on pathogen

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transmission. The impact of disease symptoms on vector behavior, for example, is a function of time after infection (Daugherty et al., 2017), and the incorporation of time into experiments is therefore essential for understanding disease incidence. Although there are data supporting the hypothesis that insect vectors of phloem-limited bacterial pathogens are attracted to infected plants, the opposite has been demonstrated for *X. fastidiosa* (Daugherty et al., 2011). In the case of *X. fastidiosa*, it is clear that time after infection is key to determining vector behavior, ultimately driving a decrease in plant transmission: plants become better sources of inoculum, but that is offset by vector discrimination against infected and symptomatic plants (Daugherty et al., 2017). In addition, experimental and modeling work has revealed distinct transmission dynamics in tolerant and susceptible plants (Zeilinger et al., 2018). Studying how time after plant infection with a pathogen affects vector behavior and the design of experiments with mathematical models in mind should provide a more holistic understanding of disease dynamics and translate results more effectively to field situations.

CONCLUDING REMARKS

Bacterial vector-borne diseases represent some of the most economically important and invasive agricultural diseases. Because of their unique biological attributes, including the colonization of both host vascular tissue and their insect vector, the study of these diseases is poised to uncover novel biological mechanisms. We hope the ideas presented in this perspective serve as an invitation for new generations of scientists to investigate many unanswered questions about vector-borne diseases and push the field forward.

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