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Adaptation of genetically monomorphic bacteria: evolution of copper resistance through multiple horizontal gene transfers of complex and versatile mobile genetic elements

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Short title: Copper resistance in monomorphic bacteria

Abstract

Copper-based antimicrobial compounds are widely used to control plant bacterial pathogens. Pathogens have adapted in response to this selective pressure. Xanthomonas citri pv. citri, a major citrus pathogen causing Asiatic citrus canker, was first reported to carry plasmid-encoded copper resistance in Argentina. This phenotype was conferred by the copLAB gene system. The emergence of resistant strains has since been reported in Réunion and Martinique. Using microsatellite-based genotyping and copLAB PCR, we demonstrated that the genetic structure of the copper-resistant strains from these three regions was made up of two distant clusters and varied for the detection of copLAB amplicons. In order to investigate this pattern more closely, we sequenced six copper-resistant X. citri pv. citri strains from Argentina, Martinique and Réunion, together with reference copper-resistant Xanthomonas and Stenotrophomonas strains using long-read sequencing technology. Genes involved in copper resistance were found to be strain-dependent with the novel identification in X. citri pv. citri of copABCD and a cus heavy metal efflux resistance-nodulation-division system. The genes providing the adaptive trait were part of a mobile genetic element similar to Tn3-like transposons and included in a conjugative plasmid. This indicates the system’s great versatility. The mining of all available bacterial genomes suggested that, within the bacterial community, the spread of copper resistance associated to mobile elements and their plasmid environments was primarily restricted to the Xanthomonadaceae family.

The dynamics of genome evolution differ considerably across the bacterial diversity spectrum. Some bacteria display a level of polymorphism that makes it possible to distinguish strains from a few housekeeping genes. On the other hand, genetically monomorphic bacteria appear very similar and require full genome sequencing to uncover some diversity (Achtman 2008). Far from being anecdotic cases, genetically monomorphic bacteria include numerous successful species, including several human pathogens, such as Mycobacterium tuberculosis, Yersinia pestis or Escherichia coli O157:H7.

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(Achtman 2008), as well as plant pathogens, such as *Pseudomonas syringae pv. tomato* (Cai et al. 2011) or *Xanthomonas citri pv. citri* (Leduc et al. 2015).

Since the discovery of genetically monomorphic bacteria, they have been considered to be an evolutionary puzzle (Achtman 2008). Genetic variability drives adaptation, which would suggest that these bacteria have a low adaptive potential. However, many have caused worldwide epidemics, demonstrating an ability to thrive in contrasted environmental conditions. For years, a variety of antibiotics have been used to control genetically monomorphic bacteria that are pathogenic to humans. Yet, the bacteria have regularly succeeded in evolving antibiotic resistance through various molecular mechanisms (Davies & Davies 2010). Genome sequencing projects have led to the discovery of the few polymorphisms responsible for some of the bacterial adaptations. For instance, ciprofloxacin resistance in *Y. pestis* and rifampicin resistance in *M. tuberculosis* are due to the mutation of a single gene on their chromosome (Hurtle et al. 2003; Telenti et al. 1993). However, in many cases, the genes responsible for adaptation were actually horizontally acquired from the bacterial mobile gene pool (Behlau et al. 2013; Galimand et al. 1997; Nonaka et al. 2012).

Horizontal gene transfer (HGT, Bryant et al. (2012)) is the incorporation of novel DNA into the host genome. In many bacterial species, HGT is so frequent that the high number of alien genes compared to the number of core genes of a species has led to the development of the concept of an open pan-genome (Chaudhuri et al. 2010; Garcia-Vallve et al. 2000). HGT can occur through conjugation (i.e. DNA transfer via plasmids or integrative conjugative elements, ICE), transformation (i.e. the uptake of naked DNA from the environment), transduction (i.e. the incorporation of DNA through a bacteriophage) and various gene transfer agents (e.g. phage-like structures that are produced by a donor cell and released into the environment (Popa & Dagan 2011)). Although we do not yet fully understand the mechanism involved, nanotubes were recently highlighted because they allow DNA and protein transfer between bacterial cells (Dubey & Ben-Yehuda 2011). Using network theory-based analyses of shared gene content among bacterial species, plasmids were shown to be of prime importance in HGT (Halary et al. 2010). In genetically monomorphic bacteria, alien genes and, therefore, traces of HGT were shown to be largely restricted to bacteriophages and plasmids (Achtman 2012).

Plasmids are composed of backbone genes, essential to their evolutionary dynamics (i.e. genes involved in conjugation, replication control and stable inheritance). The presence of assisted or autonomous mobility genetic elements determines the plasmids’ ability to be mobilizable or conjugative (Smillie et al. 2010). Due to their relative stability in the plasmid genome, backbone genes may show traces of adaptation to bacterial hosts (Norberg et al. 2011). Besides backbone genes, plasmids typically carry accessory genes encoding traits that may be adaptive for the bacterial host (Pansegrau et al. 1994; Sen et al. 2013). Accessory genes are often located on mobile genetic elements themselves, either transposons (Haines et al. 2007; Trefault et al. 2004) or integrons (Tennstedt et al. 2005), defining hot spots of insertions within the plasmid genome. These elements are generally autonomous and can be transferred between genera, families and even kingdoms (Heinemann & Sprague 1989). Plasmid-encoded accessory genes enrich the species’ gene pool and provide new adaptive traits in response to environmental modifications, such as the use of a new antibiotic or the colonization of a new ecological niche (e.g. (Hobman & Crossman 2015; Ochman et al. 2000; Weinthal et al. 2007)). Therefore, it is now obvious that in genetically monomorphic pathogens as in others, in order to further our understanding of bacterial adaptation and ecology,
we need to understand plasmid evolutionary dynamics, i.e. deciphering the genomic structure of plasmids and the mechanisms involved in plasmid spread among bacteria. While several studies have already documented the history of specific accessory genes (Perry & Wright 2013), very few studies have documented the evolution of the whole plasmid genome for a given ecological function (but see Norberg et al. (2011) for a study of the backbone genome of IncP-1 plasmid family).

Among the traits typically coded by plasmid accessory genes, antibiotic and heavy metal resistance has had a tremendous impact on human, animal and plant health (Davies & Davies 2010; Seiler & Berendonk 2012). In agriculture worldwide, fungal and bacterial pathogens that cause damage to vegetable and fruit crops have been controlled chemically for over a century, with the intensive use of copper-based pesticides (Smith 1985). This has led to the development of copper resistance \( (\text{Cu}^8) \) in several plant pathogenic bacterial species, primarily in the \textit{Pseudomonas} and \textit{Xanthomonas} genera, two members of the Gammaproteobacteria class (Cooksey et al. 1990). For example, in most citrus growing areas, Asiatic citrus canker represents a significant risk that is both direct (yield decrease, alteration of fruit quality and partial defoliation) and indirect (ban of fresh fruit export because of the quarantine status of the pathogen) (Gottwald et al. 2007; Graham et al. 2004). The disease is caused by the genetically monomorphic bacterium \textit{X. citri pv. citri}, which is disseminated through contaminated water (splashing and wind-driven rains), tools and plant materials (particularly over long distances). Given its Asiatic origin, \textit{X. citri pv. citri} has geographically expanded and become established in many citrus-producing areas, including Florida (USA), South America and, recently, Africa (Leduc et al. 2015; Pruvost et al. 2014). In regions where \textit{X. citri pv. citri} has been established for a long time, control typically involves integrated pest management strategies that partly relies on repetitive applications of copper-based pesticides (Graham et al. 2004). \textit{X. citri pv. citri} copper resistance first emerged in 1994 in Argentina and it has persisted. There were no other reports of \textit{X. citri pv. citri} copper resistance until it was discovered in the French islands of Réunion (a French territory in the South West Indian Ocean) and Martinique (a French territory in the Eastern Caribbean Sea) in 2014, two decades later (Richard et al. 2016, Richard et al. 2017). The causative genes of Argentinian \textit{X. citri pv. citri} copper resistance, identified as \textit{copLAB}, were shown to be located on a large transmissible plasmid (Behlau et al. 2012; Behlau et al. 2011), suggesting that HGT has a role. Indeed, exactly the same gene set conferring copper resistance was identified in several other \textit{Xanthomonas} species and pathovars. In addition, the hypothesis of an HGT is endorsed by evidence of the major incongruence between the phylogeny of the concatenated \textit{copLAB} genes of \textit{Xanthomonas} species and the species’ phylogeny (Behlau et al. 2013).

Although it is tempting to conclude that the mechanistic basis of copper resistance has been elucidated in \textit{X. citri pv. citri}, a recent study in a close relative species has revealed that the genetic basis for copper resistance at the infraspecific level can be variable. In \textit{X. arboricola pv. juglandis}, copper resistance could be conferred by plasmid-borne \textit{copLAB} (Giovanardi et al. 2016), as well as by a \textit{copABCD} gene set located on the chromosome (Lee et al. 1994), which is sometimes associated with an ICE structure (Cesbron et al. 2015). The \textit{copABCD} system has been described in detail on a plasmid from other bacterial species, such as the tomato pathogen \textit{P. syringae pv. tomato} (Mellano & Cooksey 1988) or \textit{E. coli} (Brown et al. 1995). The diversity of gene sets involved in copper resistance as well as the diversity of the physical supports of these genes raise the question of the nature and origin of copper resistance in \textit{X. citri pv. citri} populations.
This paper aims to improve our understanding of the genetic basis for the adaptation to copper in *X. citri* pv. *citri*. To achieve this, we gathered an unprecedented collection of strains from the three known outbreaks of copper resistance to date: Argentina, Réunion and Martinique. By combining population genetics based on microsatellite and minisatellite markers, comparative genomics and network analysis, we address the following questions: how do copper-resistant strains from the three outbreaks relate to each other? Are the genes responsible for copper resistance similar for the different outbreaks of *X. citri* pv. *citri*? What is the genomic environment of these genes and how does it vary within pathovar *citri* and among xanthomonads? We discuss the evolutionary implications of our findings and propose future lines of research to increase our understanding of how adaptive genes spread in bacterial communities.

**Materials and methods**

**Bacterial strains**

The study included a total of 350 *X. citri* pv. *citri* Cu\(^R\) strains from the three known outbreaks of copper resistance (Argentina, n = 111; Réunion, n = 219; and Martinique, n = 20). Argentinian strains originated from a collection held at INTA and were collected from 1994 (the first outbreak) to 2015. Starting in early 2014, when copper resistance was discovered in Réunion, strains were collected in the field directly from active epidemics. In addition, a large collection of citrus strains from Réunion (held at the PVBMT laboratory and including strains collected since 1978) was tested for copper resistance. This revealed resistance in 219 Cu\(^R\) strains from all locations in Réunion, as well as the oldest copper-resistant strains found, dating from 2010. Martinique was a disease-free territory until the 2014 outbreak (Richard *et al.* 2016). Every strain characterized was Cu\(^R\) and 20 were collected from the first disease focus in Morne Rouge.

In addition to Cu\(^R\) haplotypes (n = 115 identified among 219 strains), all known haplotypes corresponding to copper-susceptible (Cu\(^S\)) *X. citri* pv. *citri* strains sampled in Réunion (n = 524) from all the citrus producing areas were also included in the study to assess the genetic relatedness between Cu\(^R\) and Cu\(^S\) strains on the island. This extensive dataset was built from strain collections sampled over two periods, 1978-1997 (historical strains) and 2009-2015 (contemporary epidemics).

Finally, as Cu\(^R\) xanthomonads were originally reported in strains causing bacterial spot of tomato and pepper and no genomic resources were available for such strains, reference Cu\(^R\) strains from the four species causing this disease were included in this study. They originated from Argentina, Mauritius, New Zealand and the USA (Table 1).

**Evaluation of copper resistance**

To assess copper resistance, we used the growth test on YPGA (yeast extract 7 g l\(^{-1}\), casein peptone 7 g l\(^{-1}\), glucose 7 g l\(^{-1}\), agar 18 g l\(^{-1}\), pH 7.2), supplemented with CuSO\(_4\).5H\(_2\)O (470 mg l\(^{-1}\)), and PCR using *copL* primers. We slightly modified the PCR protocol reported previously (Behlau *et al.* 2013) by using GoTaq Flexi polymerase (Promega), a lower primer concentration (5 pmol l\(^{-1}\)) and a higher annealing temperature (66°C). In the case of negative results using *copL* primers, *copA* and *copB* primers were also used, as reported previously (Behlau *et al.* 2013).
Minisatellite and microsatellite genotyping

To determine how the studied strains relate to the known diversity of *X. citri* pv. *citri* worldwide, a subset of available strains from Argentina (47 Cu\(^8\) and 45 Cu\(^5\)), Martinique (9 Cu\(^8\)) and Réunion (64 Cu\(^8\) and 103 Cu\(^5\)) were genotyped using 31 minisatellites (MLVA-31) developed for global epidemiology analyses and compared to the known worldwide diversity (http://www.biopred.net/MLVA/). Genotyping and discriminant analysis of principal components were performed as reported earlier (Pruvost et al. 2014). MLVA fragment sizes were transformed to tandem repeat numbers (rounding to the superior integer). Manhattan distances between strains were then calculated and Multidimensional Scaling (MDS) plots were computed using the BIOS2MDS R package (Pele et al. 2012). Categorical minimum spanning trees (MST) were built using the algorithm recommended for tandem repeat data combining global optimal eBURST (GOEBURST) and Euclidean distances in PHYLOVIZ v1.0 (Francisco et al. 2012).

The relatedness between strains was assessed at smaller spatio-temporal scales (i.e. at the country level) by examining the diversity of all *X. citri* pv. *citri* strains using a set of 14 microsatellites (MLVA-14), which represent the most discriminative *X. citri* pv. *citri* genotyping technique available for intrapathotype typing (Bui Thi Ngoc et al. 2009). Population genetics analyses were performed on two different datasets, as explained above. Moreover, allelic richness was computed using a rarefaction method with the HIERFSTAT R package (Goudet 2005). Population pairwise R\(_ST\) values were computed for estimating genetic differentiation among Cu\(^8\) populations at the country level. Significance was tested with 1000 permutations using ARLEQUIN 3.5.2.2 (Excoffier et al. 2005). The first dataset, which consisted of all Cu\(^8\) strains from the three resistance outbreaks, allowed us to decipher the genetic relationships between the three outbreaks. The second dataset, consisting of all Cu\(^8\) and Cu\(^5\) strains from Réunion (the only region where we have extensively sampled local populations), enabled us to examine the relationships between the Cu\(^8\) and Cu\(^5\) strains that were isolated both before and during the Cu\(^8\) *X. citri* pv. *citri* epidemics. When comparing Cu\(^8\) and Cu\(^5\) strains from Réunion, the optimal number of clusters in the dataset was assessed by computing silhouette scores from multiple K-means runs using the BIOS2MDS R package.

DNA sequencing and assembly

Based on strain diversity analyses and the Cu profiles (i.e. the combination of the growth test and PCR results), six *X. citri* pv. *citri* strains were selected for sequencing from Argentina, Martinique and Réunion (Table 1). In addition, six other reference strains of Cu\(^8\) xanthomonads causing bacterial spot disease on tomato and/or pepper and a Cu\(^8\) strain of *Stenotrophomonas* sp., a commensal bacterium collected from the citrus phyllosphere in Réunion (i.e. the same ecological niche as *X. citri* pv. *citri*), were also selected for genome sequencing. In total, 13 genomes were completely sequenced using the long-reads PacBio RSII technology, using one SMRT cell for each strain (Table 1). Assembly of the resulting raw reads was made using SMRT ANALYSIS HGAP v. 2.3 protocol with default parameters. An additional step of contig circularization was conducted using a combination of minimus assembler (Sommer et al. 2007) and the SMRT ANALYSIS resequencing v. 1 protocol.
Sequence annotations

Sequence data obtained from the 13 strains were screened for known copper-resistance systems using BLASTN and BLASTP algorithms. As for most strains, Cu<sup>+</sup> genes were found on a 230 kb plasmid with strong homology. We arbitrarily selected strain LH201 as a reference. Hereafter, this plasmid is referred to as pLH201.1. We carried out pLH201.1 CDS prediction and automatic gene annotation using the MaGe genome annotation platform (Vallenet et al. 2006) and then manual curation of the plasmid annotations using the extensive set of tools and databases available via the platform.

Specific plasmid features were characterized in more detail. We conducted a search of plasmid toxin-antitoxin systems using the on-line databases TADB (Shao et al. 2011) and RASTA (Sevin & Barloy-Hubler 2007). The conjugative apparatus of pLH201.1 (encoded by 16 tra genes dispatched in three clusters) was blasted against amino-acid sequences of (i) the plasmid database of NCBI and (ii) the ICEberg database (Wozniak & Waldor 2010). We only retained hits that displayed homology with at least one gene (on at least 70% of its length) from each of the three gene clusters involved in conjugation. Finally, ISfinder (Siguier et al. 2006) was used to detect known insertion sequences.

We used CIRCOS (Krzywinski et al. 2009) to plot a graphical representation of the plasmid. We computed GC and GC skew \([(G+C)/(G−C)]\) using a custom perl script available at https://github.com/DamienFr/GC-content-in-sliding-window.

The search for homologous sequences

In our search for sequences that are homologous to pLH201.1 among all the available sequences, we queried the Whole Genome Shotgun (WGS) and Non-Redundant (NR) public nucleotide databases. Results were automatically retrieved using BioPerl modules (SEARCHIO, SEQIO and EUTILITIES).

We first used CYTOSCAPE v. 3.3.0 to build a network of the gene sequences that were homologous with pLH201.1. We retained all the sequences that had a hit with at least one of the pLH201.1 genes (with 95% nucleotide identity over 95% of gene length). All the hits were blasted against each other to obtain the full network structure. The network was then plotted using an edge-weighted spring-embedded layout, which positions the nodes closer if the number of genes they share is higher.

We adopted a clustering strategy based on Average Nucleotide Identity (ANI) to determine the diversity of sequences related to pLH201.1 and eliminate the redundancy of the hit dataset. Nucleotide sequences that shared more than 10 genes with pLH201.1 (with 70% nucleotide identity over 70% of gene length) were clustered using a modified ANI algorithm (available at https://github.com/DamienFr/Clustering_withANI) that takes into account the identity of the homologous region in relation to the size of the complete sequence. For each sequence that was representative of a cluster, CDS prediction was performed using PRODIGAL (Hyatt et al. 2010). Sequences were then compared using a combination of MAUVE (Darling et al. 2004), MAFFT (Katoh & Standley 2013) and the APE (Popescu et al. 2012) and GENOPLOT R (Guy et al. 2010) R packages.

Results

In this study, we combined PCR screening, micro- and minisatellite typing and comparative genomics to place the strains that caused the outbreak into the known diversity and to characterize copper resistance genetic and genomic support.
Preliminary characterization of Cu<sup>+</sup> outbreak strains and placement within the pathovar diversity

Most Cu<sup>+</sup> X. citri pv. citri strains (98%) produced the expected amplicon when assayed by PCR with copL primers. Notably, seven strains from Argentina grew on YPGA supplemented with CuSO<sub>4</sub> although they were PCR negative when assayed with copL, copA and copB primers.

Strains sampled from the three outbreaks were assessed by MLVA-31 minisatellites and were all assigned to the same genetic cluster (formerly called DAPC1 in Pruvost et al. (2014)).

At the spatio-temporal evolutionary scale defined by these markers, strains from the three origins formed a single clonal complex (CC), i.e. a network of haplotypes composed of single-locus variants. A total of 34 haplotypes were delineated with no clear structure in terms of geographical origin or copper phenotype (Fig. S1, Supporting information; information on haplotypes available at http://www.biopred.net/MLVA/). Strains from Martinique were assigned to haplotypes also including strains from Réunion. Some MLVA-31 haplotypes were shared by Cu<sup>+</sup> strains from Martinique and Réunion. Most Cu<sup>+</sup> strains from Réunion were assigned to two frequent haplotypes (#173 and 175), which also contained Cu<sup>-</sup> strains of the same origin. Similarly, haplotype #51, the most frequent haplotype among Argentinian strains, included strains that differed in terms of the copper phenotype and/or cop PCR.

Relationships between outbreaks

Using microsatellite data (MLVA-14), we first characterized the genetic diversity of Cu<sup>+</sup> X. citri pv. citri strains originating from the three resistance outbreaks. A total of 83, eight and 115 haplotypes were identified among the 111, 20 and 219 Cu<sup>+</sup> strains from Argentina, Martinique and Réunion, respectively. Using a rarefaction method (n = 20), strains from Argentina, Martinique and Réunion showed a mean allelic richness of 3.96, 1.71 and 2.76, respectively. Two clusters were identified according to their distribution on the first axis of the MDS plot (68.0% of the total variance). Their identification was endorsed by the silhouette score derived from multiple K-means runs (Fig. 1A). Axes 2 to 5 contributed much less to total variance, ranging from 6.2 to 2.2%. Cluster 1 included all strains from Martinique and Réunion, while cluster 2 only had Argentinian strains. Cu<sup>+</sup> strains that were PCR negative for copLAB also grouped in cluster 2 without any apparent substructure. Within cluster 1, strains from Martinique and Réunion were moderately but significantly differentiated (R<sub>ST</sub> = 0.22; P < 0.001). In contrast, strains from Argentina (i.e. cluster 2) were strongly differentiated (R<sub>ST</sub> = 0.85; P < 0.001) from strains collected in Martinique and Réunion (i.e. cluster 1).

Relationships between copper-resistant and susceptible strains

Microsatellite data (MLVA-14) were also used to decipher the genetic relatedness of the Cu<sup>+</sup> and Cu<sup>-</sup> strains from Réunion. Silhouette scores derived from multiple K-means runs suggested a lack of strong structure in the dataset. For K > 2, strain assignation probabilities to clusters were often low (i.e. < 0.5). We decided to retain K = 2 because minimal strain assignation probabilities to clusters were > 0.8. All but two Cu<sup>+</sup> strains were assigned to cluster A, together with Cu<sup>-</sup> strains. In contrast, only two Cu<sup>+</sup> strains were assigned to cluster B, including almost exclusively Cu<sup>-</sup> strains (Fig. 1B). In a few cases, Cu<sup>+</sup> and Cu<sup>-</sup> strains shared the same haplotype. Most Cu<sup>+</sup> strains were genetically closely
related. They were structured as five CCs and 27 singletons, i.e. haplotypes sharing no single-locus variation with others (Fig. S2, Supporting information). Among these, a major CC (CC1) comprised 64 haplotypes, representing 152 strains (69%). An additional set of 48 strains (37 haplotypes), corresponding to smaller CC and singletons, consisted of double-locus variants of CC1. These strains did not join the main CC, which is probably due to incomplete sampling. Both clusters included historical (1978-1997) and contemporary strains (≥ 2009). The Cu² D07 strain (sampled from satsuma mandarin in 1989) was the closest historical strain to the main group of Cu² strains (i.e. a double-locus variant).

**Sequencing**

After PacBio reads assembly of the 13 fully sequenced strains, we obtained from one to 18 contigs per strain (see Table S1, Supporting information for details). It is important to note that all the chromosome sequences were successfully circularized and 34 of the remaining 70 contigs were circularized into plasmids. No plasmid was detected in the copper-resistant commensal strain of *Stenotrophomonas* sp. isolated from citrus in Réunion and one to four plasmids were detected in xanthomonads, depending on the strain.

**Features of plasmids associated with copper resistance in xanthomonads**

*Cop* genes were plasmid-borne for all the *Xanthomonas* strains sequenced. Consistent with previous WGS data from *Stenotrophomonas maltophilia*, these genes were present on the chromosome of the strain of *Stenotrophomonas* sp. isolated from citrus phyllosphere (Crossman et al. 2008; Davenport et al. 2014; Pak et al. 2015). A MAUVE comparison suggested that, with the exception of *X. euvesicatoria* LMG930, all other plasmids bearing *cop* genes were genetically related (Fig. 2). Consequently, the plasmid from the *X. citri* pv. *citri* strain LH201 was arbitrarily selected as our reference. MaGe annotation of this plasmid revealed two tRNA and 258 CDS, 176 of which (68%) encoded proteins of unknown function (Fig. 3). We identified the plasmid replication initiator gene *trfA* near a GC-skew switch. This commonly indicates the origin of replication (*oriV*) (Grigoriev 1998), suggesting that the *oriV* locus is located nearby. We revealed an 87 amino acid long conserved domain of a putative HigB-like addiction module killer toxin (e-value < 10⁻¹⁷) (Schuessler et al. 2013). A 222 amino acid long putative transcriptional regulator of the xenobiotic response element family, which might serve as an antitoxin protein, was found downstream and antisense to the toxin (e-value < 10⁻⁸).

The pLH201.1 encodes for all the apparatus required for conjugation with 16 Tra proteins, located at two different regions of the plasmid (region 1: 69 675 – 74 152; region 2: 182 783 – 205 787) and organized in at least three operons (Fig. 3). Using the NCBI plasmid and the ICEberg ICE databases, we conducted a search at the amino acid level, keeping only sequences that matched at least one Tra protein from each of the three pLH201.1 conjugative operons.

These 16 pLH201.1 Tra proteins shared best amino-acid identity (AAI) (from 20 to 58%) and organization with some IncA/C plasmids and SXT/R391-related ICE conjugative apparatus (Carraro et al. 2014; Fricke et al. 2009) (Table S2, Supporting information). On the 52 plasmid hits (120 kb to 582 kb) and 16 ICE hits, not a single Tra homologue displayed a AAI superior to 60%. Twelve matched ICEs belonged to the SXT/R391 family, the remaining four were from the SPI-7 family (one sequence) or unclassified (three sequences). Most of the plasmid hits (n = 23) were annotated as IncA/C.

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plasmids, while the remainder were spread in groups H, P, T, J, F, multireplicon and unknown plasmid incompatibility groups. All the plasmids conferred multi-drug resistance and were found in five bacterial families (Pseudomonadaceae, Enterobacteriaceae, Aeromonadaceae, Burkholderiaceae and Vibrionaceae) with contrasted ecological niches and geographical origins.

As the protein sequences of conjugal relaxases are useful for plasmid classification, we assessed the AAI of the pLH201.1 relaxase (TraI_{pLH201.1}) to that of each of the six MOB groups defined in the literature (Garcillan-Barcia et al. 2009). TraI_{pLH201.1} solely matched the MOB_{H} type (e-value < 10^{-4}) with its typical amino acid signature ((HQ)-x2-PASE-x-HHH-x3-GG-x3-H-x-L and (LV)-x-HD-(AVL)-GK). MOB_{H} relaxases are scarce and have only been reported in large plasmids (> 60 kb) (Smillie et al. 2010) and found in the incompatibility groups IncH, IncJ, IncT, IncP7 and IncA/C (Garcillan-Barcia et al. 2009). Of all known MOB_{H} clades and subclades, the TraI_{pLH201.1} appeared to be most closely related to the MOB_{H_{121}} sub-clade of MOB_{H_{122}} (Alvarado et al. 2012). TraI_{pLH201.1} also displayed a conserved domain of the PFL_4751 family of ICE relaxases (e-value = 2.20 e^{-21}), required for transfer of the SXT and R391 ICE types (Daccord et al. 2010). Similarly, the conjugative coupling factor TraD was homologous to the SXT-TraD domain (e-value = 0) found in conjugal-transposon-like mobile genetic elements (Beaber et al. 2002) and various groups of plasmids including IncA/C (Fernandez-Alarcon et al. 2011). TraD_{pLH201.1} also shared good AAI with the TrwB coupling factor (e-value of 2.35 e^{-19}) from IncW conjugal plasmids (Gomis-Ruth et al. 2002).

Globally, the content and organization of pLH201.1 conjugal apparatus clearly shared similarities with that of IncA/C plasmids and SXT/R391 ICE. However, the low AAI levels indicate that it may be a new system that has not yet been described.

**Cop genes are part of a Tn3-like transposon**

In IncA/C plasmids, genes associated with adaptive traits are often found as part of complex transposons (Harmer & Hall 2014) and display a higher GC content (Zhang et al. 2014). Globally, pLH201.1 had a GC content of 59.2%, lower than the 64.8% of the chromosome, but displayed a higher local GC content (63.3%) in the ~ 108 - 152 kb region. GC-skew profiles presented several variations. These lines of evidence suggest that the plasmid shows a mosaic structure. On pLH201.1, this region (hereafter referred to as TnpLH201.1, located at 108 034 – 151 931 bp) was surrounded by two inverted repeats of 34 bp. It contained genes that are syntenic and similar to genes from the plasmid-encoded Xanthomonas TnXo19, a Tn3-like transposon (Niu et al. 2015). This includes a transposon related cointegrate protein tnpT, a cointegrate protein tnpS, a transposase tnpA, a DNA recombination protein and a DNA helicase, that all display nucleotide identity (NI) between 70% and 90% with their TnXo19 homologs, on 94%, 70%, 26%, 100% and 100% of TnPLH201.1’s gene length, respectively. tnpT, tnpS and tnpA were all shown to be involved in the transposition of some Tn3-family transposons (Tsuda & Iino 1988; Yano et al. 2013). Inside TnpLH201.1, we found an additional copy of the 34 bp repeat, which formed two direct copies separated by 43 897 bp at one extremity of TnpLH201.1 and one inverted copy at the other extremity. This pattern, typical of composite transposons, has also been reported on a Tn3-like transposon from Pseudomonas putida (Lauf et al. 1998).
Gene content of the Tn3-like transposons

The pLH201.1 contained two clusters of heavy metal resistance genes that were included in TnpLH201.1. Apart from LM199, all the strains displayed a TnpLH201.1 homolog with a globally conserved and syntenic gene content, however sometimes showing rearrangements (Fig. 2 and Fig. S3, Supporting information). The first cluster was delineated by two 34 bp direct repeats and encompassed several genes, including the previously reported copLABMGCDF genes involved in copper resistance in Argentinian X. citri pv. citri populations (Behlau et al. 2011) and cusAB/smmD present in S. maltophilia (Crossman et al. 2008). The copLABMGCDF region was identical (100%) between the five (out of six) sequenced X. citri pv. citri strains that possessed this system and with the 10 328 bp region, which was described in X. citri pv. citri strain A44 (i.e. LM180 in the present study) and known to be functionally involved in Cu\(^+\) (Behlau et al. 2011) (Fig. S3 and Fig. S4, Supporting information). Interestingly, the Cu\(^+\) X. citri pv. citri strain from Argentina LM199 failed to produce PCR amplicons using primer pairs specific to copL, copA and copB of the copLAB system. Its genome sequence displayed a plasmid backbone extremely similar to that of pLH201.1 but with a distinct copper transposon region (Fig. 2), hereafter called TnpLM199. Indeed, the annotation of TnpLM199 revealed the presence of the alternate copper-resistance system copABCD. TnpLM199 however displayed a transposition apparatus (tnpA, tnpT and TnpS) similar to that of TnpLH201.1 (NI above 90% on more than 80% of each gene’s length) and comprised almost identical 34 bp inverted repeats at its extremities. The nucleotide sequences of copA, copB, copC and copD from LM199 respectively showed a NI of 97, 98, 98 and 98% with those of the known chromosomal system of X. arboricola pv. juglandis Xaj417 isolated from walnut (Pereira et al. 2015). Copper-resistance gene nomenclature is quite ambiguous: whereas copA and copB from copABCD respectively share 98 and 63% NI (both on 74% of the gene length) with their copLAB homologs, CopC\(_{\text{copABCD}}\) and CopD\(_{\text{copABCD}}\) amino-acid sequences only display low AAI levels with their copLAB counterparts (34% on 97% of gene’s length) (see Fig. S4, Supporting information). PCR primers targeting the four genes were designed (Table S3, Supporting information). The seven Cu\(^+\) X. citri pv. citri strains from Argentina that were copLAB negative by PCR all produced amplicons of the expected size for copABCD. The system found in pLM199, as in X. arboricola pv. juglandis, did not encode for CopRS, unlike E. coli and P. syringae. However, a transcriptional regulator that belongs to the MerR family was found close to the copABCD cluster. This regulator family has been reported to respond to environmental stimuli, including heavy metals. It also controls the expression of copA in E. coli (Brown et al. 2003; Stoyanov et al. 2001).

In TnpLH201.1, genes homologous to cusAB/smmD, a heavy metal efflux resistance-nodulation-division (HME-RND) (> 95% AAI) of S. maltophilia were identified. These sequences corresponded to that of known copper/silver efflux pumps. On all strains from Réunion and Martinique (five X. citri pv. citri, one X. gardneri and one Stenotrophomonas sp.) and in one Argentinian strain (LM180), this cluster was surrounded by two almost perfect 907 bp-long direct repeats. We only found one copy on other Cu\(^+\)-bearing DNA molecules (plasmids for all strains apart from Stenotrophomonas sp. LM091) for all the other strains and none in LH3. Interestingly, on TnpLH201.1, the two copies of 907 bp comprised an Ile tRNA (UAU anticodon), which is absent on the chromosome of X. citri pv. citri LH201. Codon usage revealed a bias in the use of the ATA codon between chromosomal genes (1.4% of the Ile-encoding codons) and plasmid-borne genes (6.5%) (data not shown).
The second cluster contained genes also involved in heavy metal resistance: \textit{czcA, czcB, czcC} and \textit{czcD} (which encode a cobalt/zinc/cadmium efflux system), and three genes putatively involved in arsenic resistance, an arsC gene encoding an arsC reductase, a NADPH-dependent FMN reductase (\textit{arsH}) and an arsB gene encoding an arsB transporter. Again, \textit{czc} and \textit{ars} genes were found to be highly (> 95%) and moderately (30 to 67%) identical, respectively, to those of \textit{S. maltophilia}. The latter genes were located downstream of the \textit{arsR} transcriptional regulator, which is induced by arsenite and antimonite (Wang et al. 2004). This cluster also encoded a copper-dependent transcriptional regulator \textit{hmrR} from the MerR family, which is highly identical (97%) to that of \textit{S. maltophilia}.

The region between the two clusters was surrounded by 66 bp long direct repeats and encoded a metal chelator protein (also described in TnXo19 from \textit{X. oryzae pv. oryzae}, (Niu et al. 2015)), a heavy metal efflux protein, a copper-sensing transcriptional repressor (which binds to a gene promoter and to copper with a higher affinity to copper) and a metal binding exoribonuclease (which might help degrade the transcriptional repressor in the presence of copper).

Slight variations were observed between the TnplH201.1 homologues. For example, pLL074-4 displayed a 6,945 bp insertion at position 154,316. This insertion contained 40 bp inverted repeats at both ends. We found an identical transposon (100% NI) in a \textit{X. citri pv. citri} strain jx-6 plasmid pXAC33, which encoded a transposase, a resolvase and the twitching motility protein PilT. In addition, pLMG930 displayed an inserted gene (99% NI with \textit{S. maltophilia} iron permease), a deletion of the \textit{ars} cluster and several gene rearrangements in the region between the \textit{copLAB} and the \textit{czcABCD} clusters (Fig. S3, Supporting information).

Therefore, TnplH201.1 appeared to be a hotspot of insertions, deletions and rearrangements, consistent with previous data on \textit{Xanthomonas} Tn3-like transposons (Ferreira et al. 2015; Niu et al. 2015). Two different Tn3-like transposons encoded for copper resistance. Other genes putatively involved in heavy metal resistance were conserved in the sequenced plasmids, except for a single strain (\textit{X. euvesicatoria} LMG930).

\textbf{TnplH201.1 is found in various genomic environments}

Homologues of the transposon TnplH201.1 were found in diverse genomic environments within the other sequenced strains (see Fig. 2, Fig S5 and Table S4, Supporting information for values of nucleotide divergence between the blocks of nucleotide identity defined in Fig. 2). First, we found that TnplH201.1 was integrated in plasmids that were highly homologous and syntenic to the pLH201.1. These conserved plasmids were present in \textit{X. citri pv. citri} strains LM180, LH276, LJ207-7 and LL74-4 (from the three regions studied) and from other \textit{Xanthomonas} species pathogenic to solanaceous species: \textit{X. gardneri} JS749-3 (Réunion) and \textit{X. vesicatoria} LM159 (Argentina). Then we found that highly homologous copies of the transposon were also integrated in rearranged pLH201.1 homologues present in strains of other \textit{Xanthomonas} species pathogenic to solanaceous species: \textit{X. 'perforans' LH3} (synonym \textit{X. euvesicatoria}; Mauritius), \textit{X. gardneri} ICMP7383 (New Zealand) and \textit{X. vesicatoria} LMG911 (New Zealand) (Table 1). We also observed a highly similar transposon homologue that was integrated in a markedly different plasmid environment (\textit{X. euvesicatoria} LMG930, USA). The conjugative apparatus of pLMG930 displayed homology to that of pBVIE04 from \textit{Burkholderia vietnemensis} G4, an ecologically versatile rice root-associated nitrogen-fixing betaproteobacterium (Chiarini et al. 2006). Indeed, genes of pBVIE04 involved in conjugation are located in three separate genomic regions, each of which are very well conserved in pLMG930,
sharing 93%, 90% and 92% NI, on 7 068, 5 037 and 7 792 bp, respectively. Lastly, TnpLH201.1 was integrated in the chromosome of the citrus-associated strain of *Stenotrophomonas* sp. LM091 (Réunion) with a NI of 97.7% on 43 276 bp.

**Networks of gene sharing**

As the clues indicated a mosaic structure for the Cu\(^8\) plasmid, we searched for pLH201.1 homologues in the public NCBI databases NR and WGS. Networks of gene sharing (Fig. 4) revealed that pLH201.1 homologues present in *X. citri* pv. *citri* could only be identified from *X. gardneri* and *X. vesicatoria*, consistent with data produced in this paper. In contrast, genes homologous to TnpLH201.1 were detected from 14 species included in five genera (*Xanthomonas, Stenotrophomonas Pseudoxanthomonas, Pelomonas* and *Pseudomonas*). Globally, Cu\(^8\) gene homologues were found further apart in the taxonomy than plasmid backbone homologues.

30% of the genomes represented on the network only shared one or two genes with pLH201.1, 86% of which (26% of the total) only matched known insertion sequences.

After clustering 180 NCBI sequences sharing more than 10 genes with pLH201.1 (with 70% NI over 70% of gene length), we obtained 62 clusters (Fig. S6, Supporting information). Three patterns of homology emerged. The first pattern (two clusters) comprised homologues to the complete pLH201.1. The second pattern was found with clusters whose backbone region displayed NI with the entire pLH201.1 backbone but differed from TnpLH201.1 (three clusters). Within this group, all clusters displayed a highly similar backbone region and a conserved gene content, which suggests that they are closely related. However, their accessory gene regions were different with the lack of *copLABMGCDF* (*X. perforans*, contig accession JZUY01000051, that can be circularized with a 21bp perfect-match overlap), or the incorporation of other gene clusters coding traits such as cobalt efflux or ion transport (*X. perforans*, accession JZVH01000033). In addition, this cluster comprised a contig of a *X. euvesicatoria* pv. *alli* strain from Réunion matching the whole backbone region of pLH201.1 (accession JOJQ01000000) that we were unable to circularize. Finally, the third pattern consisted of multiple clusters that only shared NI with TnpLH201.1 or parts of it. However, the distance between them was sufficient to form distinct clusters. This confirmed that highly similar TnpLH201.1 homologues insert in diverse genomic environments.

**Discussion**

In response to the use of copper-based antimicrobial compounds to control plant bacterial pathogens, copper-resistant strains have emerged repeatedly in different parts of the world. Determinants of copper resistance have often been reported to be plasmid-borne, as in the case of *X. citri* pv. *citri*, the causal agent of Asiatic citrus canker (Behlau et al. 2012). Until now, the understanding of the ecology of these resistance determinants and the evolution of the associated plasmids has been limited by the lack of genomic data. In the present study, we provide a comparative genomic analysis of plasmids associated with Cu\(^8\) in several *Xanthomonas* species, including *X. citri* pv. *citri*. 

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Multiple acquisitions of copper resistance in X. citri pv. citri

An unprecedented collection of X. citri pv. citri Cu\textsuperscript{R} strains from all the geographical areas where its emergence has been reported (Argentina, Réunion and Martinique) was genotyped using two complementary sets of markers (minisatellites and microsatellites). Cu\textsuperscript{R} strains were genetically related, so much so that they were assigned to a single lineage (DAPC1) and formed a single clonal complex based on minisatellite data. This lineage corresponded to the wide host range pathotype A, known to be responsible for the worldwide emergence of Asiatic citrus canker in the 20\textsuperscript{th} century from Asia (i.e. its native origin) (Pruvost et al. 2014). Nevertheless, within this clonal complex, microsatellite analysis revealed a substantial differentiation into two genetic clusters likely with no epidemiological link. One cluster included Argentinian strains, while the other encompassed all strains from Réunion and Martinique. Genomic data confirmed the differentiation between French and Argentinian X. citri pv. citri strains (with intergroup genetic divergence tenfold higher than intragroup one, see Table S4, Supporting information for details). In Réunion, the genetic relatedness between Cu\textsuperscript{R} contemporary outbreak strains and Cu\textsuperscript{S} 'historical' strains isolated two to four decades ago suggested that the establishment of Cu\textsuperscript{R} strains on the island was unlikely to be the result of the recent introduction of genetically distinct strains. Instead, our data support the hypothesis that strains, which were genetically similar to the Cu\textsuperscript{S} populations characterized in the early years of the disease in Réunion, acquired a pLH201.1-like plasmid from a presently unknown source (i.e. Asiatic canker was reported for the first time in Réunion in 1968 - Brun (1971)). On the contrary, the weak genetic divergence between strains from Réunion and Martinique show a possible epidemiological link between them (Fig. S5 and Table S4, Supporting information).

Contrasting with their epidemiological structure and genetic divergence, the Argentinian LM180 strain and the French ones displayed extremely similar plasmids (Fig. S5 and Table S4, Supporting information), suggesting independent copper-resistance acquisition by these two groups of strains, and showing the mobility of plasmid-encoded adaptive traits at very large geographical scales.

This mobility and the scenario of independent acquisition was further supported by the fact that in Réunion a Cu\textsuperscript{S} X. gardneri strain (a pathogen of tomato and pepper) was found to carry a copy of pLH201.1 (average of 0.05 different nucleotides per kb) and a X. euvesicatoria pv. allii contig (Gagnevin et al. 2014), which corresponded to the pLH201.1 backbone. The latter could not be circularized and, therefore, we were unable to confirm that it was a plasmid which lacked TnpLH201.1, despite the fact that the strain was PCR negative for copLAB and had a Cu\textsuperscript{S} phenotype. In Argentina, some Cu\textsuperscript{R} strains varying in copLAB PCR amplification were not genetically differentiated based on microsatellite data. This suggests the independent acquisition of two distinct copper-resistance systems within Argentinian lineages.

Several putative copper resistance systems in X. citri pv. citri

For the Argentinian X. citri pv. citri A44 (LM180), Cu\textsuperscript{R} genes primarily shown to be experimentally functional comprise a transcriptional regulator (copL) and two copper-binding proteins (copAB) (Behlau et al. 2011). PCR tests provided evidence that most known Cu\textsuperscript{R} X. citri pv. citri strains possess this copLAB system. Using transposon mutagenesis, Behlau et al. (2011) demonstrated that the disruption of the copLAB genes was sufficient to lower the copper-resistance level to that of Cu\textsuperscript{S} strains. However, when inserted in a Cu\textsuperscript{S} strain of a closely related species, Xanthomonas 'perforans', the copLAB system alone did not confer the level of copper resistance of wild-type Cu\textsuperscript{R} strains. The
authors suggested that this was due to the fact that their recipient strain had a different genetic background (Behlau et al. 2011). In 12 out of 13 Cu$^+$ sequenced strains, we identified an additional gene cluster that could be involved in copper resistance and may explain the partial phenotype restoration observed by Behlau et al. (2011). The HME-RND system, which is also present in S. maltophilia (Crossman et al. 2008) and forms a channel through the periplasm, is composed of an inner membrane pump (here CusA), a periplasmic protein (CusB) and an outer membrane protein (SmmD) (Routh et al. 2011). The cusAB/smmD is not widespread among xanthomonads. It is not present in the draft genome of the strain complemented with copLAB by Behlau et al. for their functional analysis (data not shown) and, to date, has solely been detected from the draft genome of the Cu$^+$ X. vesicatoria ATCC 35937 and mentioned in the NR database in S. maltophilia and S. acidaminiphila. Here, we added X. citri pv. citri, X. gardneri, X. euvesicatoria and Stenotrophomonas sp. to the list of species with both the copLAB and the cusAB/smmD systems.

In contrast to the majority of strains that have been studied, some X. citri pv. citri strains from Argentina were Cu$^+$ and PCR positive for copABCD but not copLAB. One of the strains (LM199 from Argentina) was sequenced and revealed that its plasmid hosts a different Tn3-like transposon (i.e. containing copABCD) in a genetically related backbone. Both the copLAB and the copABCD systems were reported from distinct strains of the walnut pathogen X. arboricola pv. juglandis on plasmid and chromosome, respectively (Behlau et al. 2013; Lee et al. 1994). Hence, we were able to establish that at least two distinct cop systems were associated with copper resistance in Argentinian X. citri pv. citri. Why polymorphism exists in copper-resistance systems is intriguing. Currently, we lack the necessary elements to test whether it is adaptive, i.e. occurs in response to environmental variations in copper concentration or fortuitous and driven by bioavailability.

The importance of HGT for the adaptation of genetically monomorphic bacteria

For all the studied X. citri pv. citri strains, copper-resistance systems were found on closely related plasmids of approximately 230 kb in size. Extensive annotation of the Réunion X. citri pv. citri plasmid pLH201.1 revealed that it bears all the genetic elements required for conjugation, confirming in vitro tests (data not shown) and previous data on strain A44 (LM180) from Argentina (Behlau et al. 2012). The pH201.1 showed no strong homology to plasmids described previously. However, its relaxase is such that it belongs to the MOB$_{h12}$ plasmid family. The content and organization of its conjugative apparatus clearly have similarities with IncA/C plasmids and SXT/R391 ICE. IncA/C plasmids have a very broad host range and are found in very diverse environments and geographical areas. Recently, an unknown MOB$_{h12}$ plasmid from a marine environment, which also has similarities with IncA/C plasmids and SXT/R391, was reported (Nonaka et al. 2012). This suggests that the MOB$_{h12}$ plasmid family could be wider than previously thought.

Plasmids can confer a broad range of adaptive traits to their host, such as antibiotic resistance (Ochman et al. 2000), heavy metal resistance (Hobman & Crossman 2015), UV tolerance, hormone production, pathogenicity determinants and toxin production (Sundin 2007; Vivian et al. 2001). These adaptations can lead to the colonization of new ecological niches. They may even be responsible for major evolutionary events, such as the emergence of new pathogenic populations. For example, different allelic forms of the pPATH plasmid have transformed strains of the commensal bacterial species Pantoea agglomerans into gall-forming pathogens of gypsophila and beet (Weinthal et al. 2007). Our results strongly support the acquisition of a new adaptive
phenotype through plasmid incorporation by different X. citri pv. citri populations. However, the mechanisms of xanthomonad adaptation through HGT might not be restricted to plasmid acquisition. Indeed, chromosomally encoded resistance was reported on *Stenotrophomonas* and xanthomonads causing bacterial spot of tomato and pepper or bacterial blight of walnut. In addition, several genomic islands, including genes of plasmid origin, were detected on the chromosome of *X. citri pv. citri* (Gordon et al. 2015).

**Importance of plasmid-borne mobile genomic elements.**

In the present study, we provide evidence that copper-resistance gene clusters on pH201.1 were encoded on Tn3-like transposon (referred to here as TnpLH201.1). Tn3-like transposons have been reported for other plasmids in *Xanthomonas* (Ferreira et al. 2015; Niu et al. 2015), as well as other genera (Lauf et al. 1998). IncA/C plasmids often carry a complex transposon-based cluster of resistance genes involved in the spread of multi-drug resistance between bacteria (Harmer & Hall 2014).

This feature could mitigate the apparently limited host range of pH201.1, by providing a second layer of mobility. Indeed, three distinct species (*X. euvesicatoria*, *S. maltophilia* and *Stenotrophomonas* sp.) harbour a transposon almost identical to the one hosting the copLAB gene system in *X. citri pv. citri* (>99% NI) in a genomic environment that is markedly different from that of pH201.1 (i.e. a 179 kb plasmid for *X. euvesicatoria* and the chromosome for the two other species). This supports the hypothesis that the transposon is a source of mobility for the Cu⁸ gene cluster. Moreover, the copABCD system found on LM199 has 98% NI with that encoded on the chromosome of *X. arboricola pv. juglandis*, while encoded on a pH201.1-related plasmid (NI of 90% on 85% of pLM199 length, see Fig. 2 and Table S4, Supporting information). The pLM199 comprised all the 16 genes from the pH201.1 conjugative gene set. In this regard, the transposon TnpLH201.1 can be considered as an autonomous vehicle. Indeed, it encodes for Cu⁸ proteins, transcriptional regulators, a transposition apparatus and a single tRNA. The latter, which is required for the transcription of the genes encoded on the transposon, is absent on the chromosome.

To lower the fitness cost of plasmid carriage, chromosomal genome and plasmids co-evolve (Harrison & Brockhurst 2012). This process could limit the spread of entire alien plasmids and, instead, favour the incorporation of the transposon into plasmids that are already present within a restricted taxonomic group and therefore already adapted to their host.

**Barriers to HGT and importance of reservoir bacteria**

Optima of genome functioning leave strong imprints, such as GC content and codon usage. These differences tend to limit exchange of DNA between distantly-related bacteria (Popa & Dagan 2011). Indeed, within networks of shared DNA among bacterial genomes (with 95% NI), *Xanthomonas* tend to form an isolated cluster. Only plasmids with lower NI within the species connect *Xanthomonas* to some distantly-related bacterial genera (Halary et al. 2010). Our network approach with pH201.1 yielded similar results. We identified complete or nearly complete pH201.1 homologues, as well as genes involved in Cu⁸ or conjugation, primarily in the Xanthomonadaceae family.
The present study has provided evidence that distantly-related Xanthomonas species (e.g. the tomato pathogen X. gardneri and the citrus pathogen X. citri pv. citri) carry the same plasmid. Xanthomonas is a bacterial genus largely composed of plant pathogenic bacteria with a high degree of host specialization (Leyns et al. 1984). As Xanthomonas lineages that have a different host range colonize distinct ecological niches, they would not be expected to share DNA directly through conjugation because it requires cell-to-cell contact. However, in some agricultural contexts (intercropping, for example), the physical proximity of plant species contaminated with distinct bacterial pathogens could facilitate contact and HGT. In addition, extreme weather events that have been reported to spread bacterial cells over long distances (Irey et al. 2006) may contribute to the mixing of xanthomonad populations. A key factor for gene transfer between populations probably lies in the ability of Xanthomonas to survive transiently on plant surfaces, in natural plant openings or even on non-host plant species (Robinson & Callow 1986). In fact, xanthomonads were reported to form mixed-biofilm structures on plant surfaces (Cubero et al. 2011; Jacques et al. 2005), which have been recognized as highly favourable to HGT within the phyllosphere (Van Elsas et al. 2003).

Indirect transfer of copper resistance between xanthomonads may occur. Different reservoirs of bacteria resistant to antimicrobials can be found in different environmental compartments that interact and share interfaces (Nesme et al. 2014). Indeed, a resistome to environmental or industrial copper does exist and combines different genes associated with copper resistance (He et al. 2010). By tracking the dispersal and availability of this type of resistance in the natural environment or agro-ecosystem and linking it to other settings, we should be able to understand and predict how the ecosystem functions (Vieites et al. 2009).

Following sporadic reporting over several decades (Vauterin et al. 1996), commensal xanthomonads are now being more carefully characterized in terms of taxonomy or taxonomic placement (Triplett et al. 2015) or pathogenicity gene repertoires and mobile genetic elements (Cesbron et al. 2015). The extent to which these commensal Xanthomonas strains or commensal bacteria act as reservoirs or hubs for adaptive genes is still unknown. In the context of increased HGT frequencies between phylogenetically related species, the significance of Stenotrophomonas (and other genera in the Xanthomonadaceae family) as a major source of adaptive genes for xanthomonads in agricultural ecosystems has largely been underrated. At least two commensal Stenotrophomonas species displayed a highly identical copy of TnpLH201.1. Despite its relative individual insignificance as a pathogen, S. maltophilia is of major relevance in terms of plant, animal and human health because it constitutes a gene reservoir that is available for gene transfer within the community. Indeed, the panoply of resistance genes that it harbours could provide a source of antibacterial resistance determinants that are transferable to bacterial pathogens, such as the copper-resistance system presented here or other types of resistance relating to human health reported previously (Crossman et al. 2008). Our study highlights the importance of conducting further research on entire microbial communities in order to improve our understanding of the emergence of pathogenic bacteria.
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Data Accessibility

The MLVA-31 and MLVA-14 data generated in this study are available in the Xanthomonas citri genotyping database (http://www.biopred.net/MLVA/) and at https://agritrop.cirad.fr respectively. Sequences produced in this study are deposited in the GenBank database (Table 1).
Author Contributions

PL, VR, CV and OP conceived and designed the study. BIC provided essential bacterial strains and produced phenotypic data on Argentinian strains. DR, CB, KB, PG, SJ, MT, BIC, CV and OP carried out the experiments. OP, CV and VR conducted population biology analyses. DR and PL conducted genomic data analyses. BF, IR and AR helped to design and carry out the study. DR, VR, AR, BF, IR, AC, CV, OP and PL wrote the manuscript. All the authors have read and approved the final manuscript.
| Strain | Other numbers | Accession | Genus | Species | Pathovar | Copper resistance location | Country | Date | Host |
|--------|---------------|-----------|-------|---------|----------|---------------------------|---------|------|------|
| LMG930 |                | CP018463-CP018467 | Xanthomonas | euvesicatoria | plasmid | USA | 1969 | Pepper |
| LMG911 |                | CP018725-CP018727 | Xanthomonas | vesicatoria | plasmid | New Zealand | 1955 | Tomato |
| LM091  |                | CP017483 | Stenotrophomonas | sp. | chromosome | Réunion | 2015 | Tangor |
| LM199  | Xcc 15-4632 (INTA) | MSQV00000000 | Xanthomonas | citri | citri | plasmid | Argentina | 2015 | Orange |
| LM180  | Xcc 03-1638 (INTA); A44† | MSQW00000000 | Xanthomonas | citri | citri | plasmid | Argentina | 2003 | Pomelo |
| LM159  | Bv-5-4a (INTA) | CP018468-CP018471 | Xanthomonas | vesicatoria | plasmid | Argentina | 1987 | Pepper |
| LL074-4|                | CP018847-CP018849 | Xanthomonas | citri | citri | plasmid | Martinique | 2014 | Grapefruit |
| LJ207-7|                | CP018850-CP018853 | Xanthomonas | citri | citri | plasmid | Réunion | 2012 | Kaffir lime |
| LH3    |                | CP018472-CP018476 | Xanthomonas | 'perforans'‡ | plasmid | Maurice | 2010 | Tomato |
| LH276  |                | CP018854-CP018857 | Xanthomonas | citri | citri | plasmid | Réunion | 2010 | Kaffir lime |
| LH201  |                | CP018858-CP018860 | Xanthomonas | citri | citri | plasmid | Réunion | 2010 | Kaffir lime |
| JS749-3|                | CP018728- | Xanthomonas | gardneri | plasmid | Réunion | 1997 | Tomato |
| Accession | Characteristics of copper-resistant bacterial strains used for long-read sequencing |
|-----------|---------------------------------------------------------------------------------|
| CP018730  | Xanthomonas gardneri plasmid New Zealand 1980 Tomato |
| ICMP7383  | CP018731-CP018734 Xanthomonas gardneri plasmid New Zealand 1980 Tomato |

ICMP (International Collection of Microorganisms from Plants, Landcare Research, Auckland, New Zealand), BCCM/LMG (Belgian Coordinated Collections of Microorganisms, University of Ghent, Belgium), INTA (Instituto Nacional de Tecnología Agropecuaria)

† As designated by Behlau et al. (2011).
‡ X. 'perforans' was reclassified as X. euvesicatoria

**Tables**

**Table 1** Characteristics of copper-resistant bacterial strains used for long-read sequencing
Figure captions

Fig. 1 Genetic diversity derived from microsatellite data of Xanthomonas citri pv. citri strains differing in geographical origin and their susceptibility to copper. (A) Manhattan distances represented by multidimensional scaling (MDS) among copper-resistant haplotypes from Argentina, Martinique and Réunion. The MDS plot showing axes 1 and 2 represented 68.0% and 6.2% of the total variation, respectively. Blue and red dots indicate strains bearing the copLAB and copABCD system, respectively, as confirmed by PCR. (B) Manhattan distances represented by multidimensional scaling (MDS) among copper-resistant and copper-susceptible haplotypes from Réunion, sampled between 2009 and 2015. Historical strains sampled between 1978 and 1997 were included as supplementary (suppl.) individuals in the MDS analysis. The MDS plot showing axes 1 and 2 represented 27.5% and 14.8% of the total variation, respectively. The black arrow localizes strain D07, which was identified as the historical strain genetically closest to the main clonal complex of epidemic copper-resistant strains.

Fig. 2 Alignments between pLH201.1 and sequences carrying pLH201.1 homologues from strains sequenced in this study. All sequences are plasmids, unless specified otherwise. On pLH201.1, blocks represent curated gene prediction (Mage), whereas on other genomes the blocks represent uncurated Prodigal gene prediction. The pLH201.1 genes are categorized into several groups according to the caption. The comparison zone shows homologous nucleotide sequences. Colour varies according to genetic distance between homologous blocks defined by a MAUVE alignment.

Fig. 3 Circular representation of the plasmid pLH201.1. Circles, from the outer to the inner, represent: (1) predicted genes and their function according to the caption at the centre of the circle (putative genes are followed by *); (2) difference between mean GC content of the chromosome of LH201 and pLH201.1’s % of GC content in a 2 000 bp sliding window with a 200 bp step; and (3) GC-skew using a 4 000 bp sliding window with a 200 bp step.

Fig. 4 Network of all NCBI genomes sharing homologous genes with pLH201.1 (nucleotide identity > 95% on > 95% of pLH201.1 gene length). Edges appear closer if the number of genes they share is higher and diameter of the nodes is proportional to the number of genes shared with pLH201.1. (A) Nodes are coloured depending on the taxonomy of the organisms, (B) Nodes are coloured in green if at least one gene in the sequence is homologous with TnpLH201.1, otherwise they are red.

Fig. S1 Categorical minimum spanning tree of DAPC1 strains from Argentina, Martinique and Réunion, which differ in their susceptibility to copper (268 strains – 34 haplotypes), based on minisatellite data. These strains were organized as a single clonal complex (i.e. a network of haplotypes linked by single-locus variations). Dot diameter is representative of the number of strains per haplotype. Colour indicates the strain origin and copper phenotype: light green = copper-susceptible Argentina; dark green = copper-resistant Argentina; khaki = copper-resistant non-copLAB Argentina; red = copper-resistant Martinique; light blue = copper-susceptible Réunion; dark blue = copper-resistant Réunion.
**Fig. S2** Categorical minimum spanning tree of copper-resistant *Xanthomonas citri* pv. *citri* strains from Réunion based on microsatellite data. These strains were organized as five clonal complexes (i.e. networks of haplotypes linked by single-locus variations) and 28 singletons (i.e. haplotypes with no single-locus variants). Dot diameter represents the number of strains per haplotype. Single and double-locus variations were represented as solid and dotted lines joining haplotypes, respectively.

**Fig. S3** Alignments between TnpLH201.1 and sequences carrying TnpLH201.1 homologues from strains sequenced in this study and *S. maltophilia* K279a. All sequences are plasmids, unless specified otherwise. On TnpLH201.1, blocks represent curated gene prediction (Mage), whereas on other genomes the blocks represent uncurated Prodigal gene prediction. The pLH201.1 genes are categorized into several groups according to the caption. The comparison zone shows homologous nucleotide sequences. Colour varies according to genetic distance between homologous blocks defined by a MAUVE alignment.

**Fig. S4** Distance tree of the copper-resistance region of all the sequenced strains as well as some sequences coding for known copper-resistance systems extracted from public database along with and graphical representations of the copper-resistance genes organisation. The alignment used for the tree computation was obtained after the concatenation of alignments of each gene.

**Fig. S5** Heatmap representation of the genetic divergence between six sequenced strains. Divergence values (proportions of variable nucleotides between two sequences) were obtained from the comparison of all homologous regions. Whereas the lower triangle represents the divergence between the chromosomes, the upper triangle represents divergence between the plasmids.

**Fig. S6** Clusters of sequences with homologues to pLH201.1 genes. Each line represent a cluster of sequences obtained from GenBank that are homologous to at least 10 pLH201.1 genes. Squares represent homologous genes and are ordered as in the pLH201.1 sequence. Squares are coloured according to their level of nucleotide identity with pLH201.1, as indicated on the scale on the right. The names at the left of each line indicate the species of the parental sequence of the cluster, its GenBank identifier (gi) and the number of sequences in the cluster. Psp: *Pseudoxanthomonas* sp.; Sac: *Stenotrophomonas acidaminiphila*; Sma: *Stenotrophomonas maltophilia*; Sni: *Stenotrophomonas nitritireducens*; Ssp: *Streptomyces* sp.; Xar: *Xanthomonas arboricola*; Xax: *Xanthomonas axonopodis*; Xca: *Xanthomonas campestris*; Xci: *Xanthomonas citri*; Xeu: *Xanthomonas euvesicatoria*; Xga: *Xanthomonas gardneri*; Xor: *Xanthomonas oryzae*; Xpe: *Xanthomonas perforans*. 
