VNTR Typing of the Bacterial Rice Pathogen Burkholderia glumae Reveals the Coexistence of Several Diverging Lineages in a Single Field in Colombia

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Vietnamese banana is responsible for the panicle blight disease of rice. This disease is present worldwide and can result in significant drop in yields. To estimate the genetic diversity of the bacterial strains present in a rice paddy field in Colombia, we sampled 109 strains from infected panicles. To detect fine genetic relationships among related haplotypes, and to overcome a very low nucleotide diversity detected in previous studies, we designed primers to amplify and sequence several highly variable minisatellite loci, or variable number tandem repeats (VNTRs), as well as part of the Toxoflavin toxA gene in all strains. Results show that the toxA nucleotide diversity defined four lineages and was similar to that detected in several fields in Japan; data suggest that B. glumae has spread from Asia to America without major loss of genetic diversity, and that five VNTR loci discriminated the strains within the field revealing single and multi-infections of the rice panicles with a wide distribution of the haplotypes among the different plots. Even though disease levels vary considerably from year to year, the bacterial genetic diversity is maintained within a field. We do not detect any geographical structuring within the field, nor any effect of the rice cultivar on the observed diversity. The consequences on the origin and evolution of the bacteria are discussed.

**Keywords:** Burkholderia glumae, Colombia, population structure, rice, VNTR

Rice is the staple food for >3.5 billion people worldwide, around half of the world’s population. Its cultivation is under threat from many diseases that decrease yields significantly and consequently have dramatic effects on the lives of human beings. Among these diseases, some have been studied for many years, such as rice blast (caused by **Magnaporthe oryzae**; Gladieux et al. 2018) or bacterial blight (caused by **Xanthomonas oryzae** pv. *oryzae*; Jiang et al. 2020). Although less studied, panicle bacterial blight caused by **Burkholderia glumae** appears to be one of the most worrying emerging diseases. **B. glumae** was first described in Japan at the end of the 1950s as causing symptoms of rotting grains and leaf sheaths in rice (Goto and Ohata 1956). This bacterial disease has now been described in many countries in Southeast Asia, and in Latin and North America (Mondal et al. 2015; Nandakumar et al. 2009; Riera-Ruiz et al. 2014; Zhu et al. 2008). The reduction in yield from this disease can reach up to 75% for the most affected plots (Trung et al. 1993), resulting from reduction of the weight of grains or from the sterility of the inflorescences. The determinants of re-emergence of the disease remain unclear. Prolonged periods of warm and wet conditions at the rice flowering time appear to be of primary importance (Lee et al. 2015). The correlation between high prevalence and high night-time temperatures, such as those reported in 1995 and 1998 in Louisiana, supports this hypothesis (Nandakumar et al. 2009). In today’s context of global warming, such high temperature conditions during flowering are likely to become more frequent in the coming years. This has led authors to suggest that **B. glumae** could become in the future the “next major pathogen of rice” (Ham et al. 2011).

Data on the epidemiology of this disease remain very fragmented. The bacterium is considered by some authors to be seed-borne, wherein it could be located in the epidermis or the parenchyma (Agarwal and Sinclair 1980). However, while **B. glumae** has been detected in seeds (Cottyn et al. 2009; Sayler et al. 2006), seed transmission and the significance of seed-borne initial inoculum for the development of bacterial blight has not been formerly assessed. The bacterium might also be present in the leaf sheaths, forming the source of inoculum for development in the panicles, from which the spread of the disease would be in close proximity to other plants, therefore producing an aggregated profile. The spatial distribution within a field would be a function of the most affected panicles, from which the spread of the disease would be by proximity (Trung et al. 1993). The available data do not go much further, with only microscopy results that monitored tagged bacteria in planta (Li et al. 2016). Moreover, detection of the bacterium in the sheath, leaf blade, boot, or spikelet remains challenging, with drastic variations of population size from sheath to leaf (Tsushima 2011), leading to apparent “disappearance” and “reemergence” between consecutive stages of development (Echeverri et al. 2021). Finally, this bacterium has been shown to interact with other pathogens such as fungi, which might also play a role in the epidemiology of the bacteria (Jung et al. 2018).

The understanding of the origin and emergence of this plant bacteriosis also remains very incomplete. The disease and its causative agent were described and isolated for the first time in Asia (Kurita and Tabei 1967). However, as of this writing, there is no evidence that Asia is actually the center of origin for the disease. Furthermore, at local scale, data on the fine genetic diversity of the bacteria is almost nonexistent. One study has looked at the genetic diversity in one country. Maeda et al. (2006) sampled and genetically typed 69 strains of **B. glumae** isolated from rice at 25 different sites in Japan. Characterization based on the sequencing of three loci was, however, not very informative on the genetic structure of the diversity of this species at this geographic scale because of the very low level of nucleotide diversity detected. With 25 isolates sampled in Arkansas in 2003, Sayler et al. (2006) detected 22 unique but closely related DNA fingerprint profiles. Cluster analysis revealed two major groups and
their genetic diversity was relatively low as revealed by 16S-23S rDNA internal transcribed spacer sequence analysis. Finally, little genomic data are available. Indeed, at the time this project was initiated, only two full genomes were available from GenBank (including the type strain LMG2196\(^T\); https://www.ncbi.nlm.nih.gov/genbank/; Lim et al. 2009), five others being available as draft genomes. However, these strains do not correspond to any logical sampling (in an epidemiological or evolutionary sense), but rather are specific strains from different laboratories. Two studies have compared genomes (Fory et al. 2014; Francis et al. 2013) and suggested a considerable amount of plasticity, largely based on accessory genes and genome islands, including multiple insertion/deletion (indel) events.

The goal in this study was to get the first image of the genetic diversity of \textit{B. glumae} isolates sampled from a single rice field in Colombia, a country where the disease has been present for several years with high incidence rates but with a high variation between years. Considering the lack of variable genotyping markers available for this species, we developed several \textit{B. glumae} variable-number-of-tandem-repeat (VNTR; Bakhtiari et al. 2018) markers to obtain a first snapshot of within field diversity, and thus circumvent the limits of the multi-locus sequence analysis approach. We detected a large genetic diversity, with several independent genetic lineages, raising the possibility of either several introductions of the bacterium into Colombia or considering this country as an alternative diversification zone.

### Materials and Methods

#### Sampling of isolates.

We sampled panicles from two rice varieties growing in a single field at the Fedearroz station in Saldaña, Tolima, Colombia (3\(^\circ\)54’50.80’’N; 74\(^\circ\)59’6.76’’W). In late summer in 2017, a trial included eight different varieties that were sown for agronomic and resistance to pathogens. Each variety was grown in four replicates, each in a 50 m\(^2\) plot (Fig. 1). For each of the two varieties known to be either susceptible (F2000) or moderately resistant (FNAL004-1) to panicle blight, we selected in each plot three panicles from three different plants at maturing stage with, whenever possible, two plants with clear symptoms of panicle blight and one plant with no or low level of symptoms. Each panicle was stored in an individual paper bag.

Back at the lab, we isolated strains from seeds. Briefly, from five to 10 seeds were put in a 2-ml Eppendorf tube (Hamburg, Germany) and ground with a small sterile pestle. One ml of sterile water was added, and tubes were left at room temperature for 2 h. Serial dilutions of maceration, from 1/10 to 1/1,000, were plated (100 \(\mu\)l) on King’s B agar (King et al. 1954) and incubated at 28\(^\circ\)C during 24 h. Each plate was checked under a stereoscope and several colonies were picked up according to their morphology that were similar to \textit{B. glumae} (pictures available upon request to G. Bena). All colonies collected were first streaked on King’s B plates to verify their purity, and, starting from a single isolated colony, all were grown in liquid Yeast Mannitol Broth. DNA for each sample was extracted with a classical proteinase K protocol. All strains were stored at \(-80\)^\circ C in 20\% glycerol.

#### Molecular characterization of isolates.

Toxollavin, the main phytotoxin produced by both \textit{B. glumae} and \textit{B. gladioli}, is one of the main enzymes involved in the pathogenicity of \textit{B. glumae} (Jeong et al. 2003) and, as of this writing, has never been detected in other species. In a first step, we tested for a positive amplification for all DNA samples targeting the methyltransferase \textit{toxA} gene, which is part of an operon of five genes that codes for the synthesis of the phytotoxin toxollavin, with primers designed in this study (Table 1). These primers were used both for confirmation of the taxonomic status of the isolate (positive or negative result for PCR amplification) and detection of genetic diversity among our isolates (by sequencing). All \textit{toxA} and subsequent VNTR PCR products were sent to GenoScreen (Lille, France) for purification and forward strand sequencing.

In a second step, we typed all strains of \textit{B. glumae} determined according to their \textit{toxA} sequences. As previous works had shown the low genetic diversity detected among genomes available in GenBank or among isolates sampled all around Japan (Maeda et al. 2006), we used highly variable VNTR loci to characterize different genomes. We used the online bioinformatics tool Polloc-V (http://bioinfo-web.mpl.ird.fr/xantho/utils) to screen for VNTR loci in five \textit{B. glumae} genomes from the NCBI data bank (LMG2196\(^T\), BGR1, AU3208, 3252-8, and 336gr-1). Polloc-V identifies groups of loci based on the similarity rate of their flanking regions. In a first analysis, we used the following selection criteria:

- full loci size from 30 to 1,000 bp;
- pattern size from 5 to 9 bp;
- number of repetitions set to six or above;
- stringency set to maximal values 2 (match)–3 (mismatch)–5 (indel);
- minimum percentage of similarity between repeat units = 80%.

In the first row of analysis, only a few primers could be selected, either because they could not be detected in one or several genomes (from indels or draft status) or because of their lack of diversity. We then selected 14 of the 366 total loci from each of the five genomes to type the isolates. These 14 loci were then used to type all strains of \textit{B. glumae} collected during the study period, which could be grouped in 19 distinct clusters (Fig. 2).

#### Fig. 1. Map of the trial established during September 2017 in the Experimental Center las Lagunas - Fedearroz in Saldaña, Colombia. Each plot (50 m\(^2\) each) is numbered from 1 to 8, with four replicates (10 to 40). The whole field was flooded. Rice genotypes shown in each plot are in italics. Plots in gray shading were sampled. The blue cross in a circle is where the global positioning system track was taken (3\(^\circ\)54’50.80’’N; 74\(^\circ\)59’6.76’’W).
perform a second analysis with the same parameters but with the pattern size extended from 10 to 90 bp (i.e., corresponding to minisatellites rather than microsatellites). Similarly, we only kept sequences that were recovered from the five genomes and for which diversity could be detected. For all loci detected that fulfilled these constraints, we designed a set of primers in the flanking regions of the tandem repeat sequence using the web interface tool, Primer3Plus (https://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi; Untergasser et al. 2007). All pairs of primers designed were tested in silico using the suite of command-line tools NCBI-BLAST+ (https://blast.ncbi.nlm.nih.gov/Blast.cgi?PAGE_TYPE=BlastDocs&DOC_TYPE=Download; Camacho et al. 2009) against B. glumae genomes to confirm their specificity for the genomic region containing the locus. Whenever possible, melting temperatures were set around 60°C (Table 1). Moreover, because the genome of B. glumae consists of two chromosomes and four plasmids (Lim et al. 2009), we tried as much as possible to have loci on each of the two chromosomes.

All primer pairs designed have been tested for positive amplification with a subsample of 24 randomly selected isolates from our derived sample. To obtain the highest level of diversity, including both the number of repeats and single nucleotide polymorphisms (SNPs), we sequenced these loci for all individuals. We retained for further analysis only those primers for which we obtained amplification for at least 95% of the individuals and for which sequencing revealed variability, both in terms of number of repeats and in terms of SNPs, both within the repeats and in the flanking regions of the locus. All sequences obtained were verified and corrected if necessary using the software ChromasPro v.1.34 (Technelysium Pty Ltd, South Brisbane, Queensland, Australia). The different haplotypes of each locus were detected using the tool Mothur (Schloss et al. 2009). Each different haplotype sequence has been deposited in GenBank. We finally included in the final dataset other B. glumae genomes (full or draft) that were released after the typing of all our isolates.

**Data analyses.** At a first step, using the tool BLASTn (https://blast.ncbi.nlm.nih.gov/Blast.cgi?PAGE_TYPE=BlaltDocs&DOC_TYPE=Download), we recovered all the B. glumae toxA sequences of at least 473 bp available in GenBank and we reconstructed a Maximum Likelihood phylogenetic tree using the program MEGA6 (https://www.megasoftware.net; Tamura et al. 2013). The best model of molecular evolution was chosen according to the “Akaike’s information criterion” function implemented in MEGA6. The toxA sequences of the sister species B. gladioli were only included when we searched for traces of selection along the branch joining the two species (see next paragraph). None of the data extracted from GenBank suggests toxA transfer between the two species, which confirms the species determination based on the toxA sequence (data not shown). As a second step, to simplify the phylogeny obtained, we used the software package Mothur (https://mothur.org). We kept only one or two representatives of each haplotype detected, retaining isolates from different countries when their sequences were identical to reflect the geographical distribution of diversity.

To explore a possible selection that would have shaped and constrained the nucleotide diversity of toxA sequences, we estimated the number of synonymous and non-synonymous mutations in the dataset using the tool DnaSP 6 (http://www.uh.edu/dnasp; Librado and Rozas 2009). We tested for selection along the gene by estimating the dN/dS between all pairs of sequences (i.e., the ratio of the number of non-synonymous substitutions per non-synonymous site to the number of synonymous substitutions per synonymous site). We also tested for a significant departure from a neutral evolution of this ratio along the branch joining the B. glumae and the B. gladioli clades using the online tool datamonkey (https://www.datamonkey.org/; Weaver et al. 2018) with the “aBSREL” test function (https://www.datamonkey.org/absrel; Smith et al. 2015).

**VNTR diversity and network reconstruction.** The different alleles of each VNTR locus were detected on the basis of the total length (and thus the number of repetitions) but also considered point mutations (same length but different sequences). The discriminatory power of each locus was compared with the Hunter–Gaston discriminatory index. This index is based on the probability that two unrelated strains randomly sampled in our population will be placed into different typing groups.

To visualize the different clonal complexes and genetic relationships among haplotypes, we applied a two-step procedure. We considered separately the nucleotide diversity detected in the toxA sequences and in the flanking region of each VNTR locus and the variations in the number of repeats for each VNTR locus. To begin, we retrieved all the nucleotide diversity from toxA and flanking regions, for which the rate of point mutations must be considerably lower than the rate of mutations within the repeat region. We concatenated these sequences for each isolate. With this data matrix, a neighbor-net split graph was built, using the software SplitsTree4 (http://www.sofsea.com/review/SplitsTree4.html; Huson and Bryant 2006). This graph allowed us to distinguish between the different groups of sequence type (ST) but also potentially illustrate recombination effects. Then, for the different groups of haplotypes obtained on the graph, we applied a network reconstruction using the program PHYLOVIZ v.2 (https://www.phylolviz.net/; Nascimento et al. 2017), with the algorithm of global optimal (“goeBURST” function) distances. It was not a straightforward task, as one highly polymorphic VNTR locus (BG02_2) revealed, after sequencing either differences in length involving several indel events or alleles with a different nucleotide sequence (involving puntual mutations within repeats) but the same total number of repeats (i.e., homoplasy). We thus reconstructed the network based on the haplotypes coding schemes rather than on the VNTR length of each loci, and considered in the network all single and double locus variants.

**Structure of the genetic diversity in the field.** We tested whether there was a significant effect of the variety or plot on the bacterial genetic diversity recovered from each plant. Initially, we simply mapped on the haplotype network the origin of the isolates, whether it was the variety or the plot. We then tested whether the distribution of the different haplotypes between varieties and between plots deviated from a random distribution, using the program Genepop (https://cran.r-project.org; Rousset 2008), with the “population differentiation” option and an unbiased estimate of the P value performed on a contingency table. Subsequently, to disentangle plot and variety effects, we tested for a significant differentiation “between variety” and “among plot within each variety” in an analysis of molecular variance function

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**Table 1.** List of the primers used, all of which were designed in this study

<table>
<thead>
<tr>
<th>Locus</th>
<th>Primer’s name</th>
<th>Primer’s sequences (5’ to 3’)</th>
<th>Tm°/elongation time</th>
<th>Location (chromosome/gene)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ToxA</strong></td>
<td>ToxA1F</td>
<td>TTTCGCGGCGTGAATCTATC</td>
<td>56°C</td>
<td>Chr2; toxoflavin methyltransferase</td>
</tr>
<tr>
<td></td>
<td>ToxA1R</td>
<td>GTCTGAGCAGCAGTTGACGT</td>
<td>30 s</td>
<td>Chr1; hypothetical protein</td>
</tr>
<tr>
<td><strong>BG09</strong></td>
<td>BG09F</td>
<td>AAACGGTCCCGATTCTTTC</td>
<td>57°C (deleting)</td>
<td>Chr1; hypothetical protein</td>
</tr>
<tr>
<td></td>
<td>BG09R</td>
<td>TGTTGTCGACAACGGCAGC</td>
<td>1 min</td>
<td>Chr2; arginine metabolism</td>
</tr>
<tr>
<td><strong>BGM03</strong></td>
<td>BGM03F</td>
<td>ATCATGACCGCTTCTTCC</td>
<td>57°C</td>
<td>Chr2; intergenic sequence</td>
</tr>
<tr>
<td></td>
<td>BGM03R</td>
<td>TCCGCGTCTAACTTCTTACGA</td>
<td>1 min</td>
<td>Chr2; iron regulon</td>
</tr>
<tr>
<td><strong>BG12</strong></td>
<td>BG12F</td>
<td>CCGCCGATCGTTCCTTAAAC</td>
<td>57°C</td>
<td>Chr2; iron regulon</td>
</tr>
<tr>
<td></td>
<td>BG12R</td>
<td>TGTAGTGACCGCGGTTGAAA</td>
<td>1 min</td>
<td>Chr2; iron regulon</td>
</tr>
<tr>
<td><strong>BG02</strong></td>
<td>BG02F</td>
<td>GTCCGCGTATGTGCAATGC</td>
<td>57°C</td>
<td>Chr2; lipoprotein, bacterial collagen</td>
</tr>
<tr>
<td></td>
<td>BG02R</td>
<td>GTAGAAAACCTGGCCGAGT</td>
<td>1 min 30 s</td>
<td>Chr2; lipoprotein, bacterial collagen</td>
</tr>
</tbody>
</table>

* Tm°, melting temperature.
* Location on chromosome and gene are given according to LMG21967® genome sequence and annotation (GCA_000609955.1); all PCR cycling parameters start with an initial denaturation step (95°C for 5 min) followed by 30 three-step cycles (94°C/30 s, Tm/30°, 72°C/elongation time).
performed with the program Arlequin v.3.0 (http://cmpg.unibe.ch/ software/arlequin3c; Excoffier et al. 2005). The linkage disequilibrium among all pairs of loci was tested with the “log likelihood ratio” statistic implemented in the program GenePop.

Results

Identification of B. glumae isolates and toxA diversity. We sampled 24 different panicles from eight plots and two varieties. Three panicles showed no symptoms (plots 103, 202, and 407), six showed mild symptoms, and the last 15 showed severe symptoms. We isolated strains based on their morphology from each of the panicles. The number of strains isolated per panicle varied from four to 14, and the number per plot varied from 24 to 31, for a total of 207 strains that were stored in glycerol at −80°C. The toxA amplification was positive for 119 samples. We sequenced all 119 of these PCR products, and sequences obtained clearly clustered 10 isolates within a B. gladioli clade with all other sequences of this species obtained from GenBank. Although known to be pathogenic on rice, we did not include these strains in the study because they represented <9% of the entire data sets and were not associated to a variety nor a specific plot. We continued the analyses with the remaining 109 isolates (Supplementary Table S1). These strains were distributed over the eight plots, with a variation from seven (plot 304) to 22 strains (106) per plot. For two panicles (one symptomless panicle from plot 103 and one with mild symptoms from plot 304) no strains for which toxA could be amplified were obtained. Conversely, we obtained six strains from a panicle in plot 202 that showed no symptoms (Supplementary Table S1).

We retrieved 88 toxA sequences from GenBank, originating from Japan, China, Korea, Vietnam, the U.S.A., Puerto Rico, and Colombia, corresponding to the main areas where the disease has been reported so far. We aligned these sequences with ours, which is 473 bp over the 738 bp of the entire toxA gene. Along the 473 bp we only detected five SNPs, two of which were specific to our dataset, giving seven different haplotypes, four of which were detected in our study. The alignment of 23 complete toxA sequences, extracted from GenBank with our four haplotypes, revealed only eight different mutations, four being singletons (i.e., limited to a single sequence). The phylogenetic reconstruction resulted in seven different clades, three including strains isolated from different continents (Fig. 2). Of the eight mutations detected, three resulted in amino acid substitution, while the five SNPs detected in our data are all silent. We did not detect any evidence of a significant departure from a neutral selection using the aBSREL test (https://www.datamonkey.org/absrel; Smith et al. 2015) implemented with the online tool datamonkey, either when testing the overall B. glumae phylogeny, or the branch joining the two B. glumae and B. gladioli species, or the branch joining the two Burkholderia species with two homologous sequences from Pseudomonas species (data not shown).

VNTR loci selection. We designed 14 pairs of primers that framed different VNTR loci. The quality and efficiency of these primers were first tested on DNA of the type strain LMG2196T. Among the 14 pairs of primers, one pair did not produce any amplicon and was discarded. The 13 others resulted in a single band of the expected size. We then tested these primers on a subsample of 24 B. glumae isolates randomly chosen from our study. The rate of success ranged from 42 to 100% for the primers, with only three primer pairs giving 100% amplifications. We retained these last three loci and a fourth primer pair, which failed to amplify two samples. Locus BG09 is located on chromosome 1 whereas the three others (BG02, BGm03, and BG12) are on chromosome 2.

We amplified and sequenced the four loci for all our isolates. Characteristics of each VNTR locus are given in Table 2. The length of each repeat (8 to 12 bp) classify them at the frontier between minisatellites and microsatellites. The BG02 locus is composed of two VNTRs in a row, the first with two to five repetitions of a 12-bp pattern, the second just afterward with 27 to 51 repetitions of a 9-bp pattern. In all subsequent analyses, we treated them separately, respectively as BG02_1 and BG02_2. Loci BG12 and BGm03 are nearly perfect, with only one SNP detected in each; BG02_1 is perfect, whereas BG09 and BG02_2 are compound VNTR loci, with, respectively, four and seven different repeats. However, in each case, the different repeats only differ from each other by a single nucleotide difference. The 109 typed strains produced 19 multi-locus haplotypes (Supplementary Table S2). The number of alleles, when considering both number of repeats and SNPs, ranged from six to 16 alleles (Table 2). When only considering number of repeats, the number of alleles ranged from four to 14. The HDGI scores ranged from 0.645 to 0.924. The combination of the five loci (toxA + the four VNTRs) has a discriminatory power of 0.938.

Within plot and within panicles diversity. Each sampled plot contains from two (plots 103 and 303) to six different haplotypes (plot 106). There is no correlation between genetic distance among haplotypes and their co-occurrence within a plot (Fig. 3). We can also note the low intrapanel diversity observed. Indeed, of the 21 panicles for which we isolated several strains of B. glumae, only six contained several haplotypes (Supplementary Table S2). Among these six panicles, only one (plot 106 panicle 2) suggests intrapanelular diversification (i.e., haplotype ST11). ST11, composed of only one isolate, is closely related to ST10 (n = 7 isolates) as a single locus variant, and further differs by a unique repeat. It probably emerged as a result of a mutation of ST10. The other panicles contain very divergent haplotypes differing by at least three VNTR loci, belonging to different genetic groups (i.e., groups including single- and double-locus variants) according to the network.

The four loci display different mutation step models. Sequencing suggested that each locus did not evolve by following the same mutation step model. Within the small clonal complexes, each allele differs from another only by a single repeat for VNTR loci BG12, BG09, BGm03, and BG02_01. Conversely, for BG02_02 we detected differences between alleles along the branches of clonal complexes ranging from one to 12 repeats. ST3 and ST12 each differ (by three repeats) at locus BG02_2 (38 and 41, respectively), and the alignment of the sequences shows that this is the result of a single three-repeat indel event. Conversely, ST1 and ST2 also differ by three repeats (34 and 31, respectively), but the alignment shows three different mutation events, involving three different indel events (Fig. 3). We mapped onto the network the different number of mutation events between haplotypes for BG02_2.

Network reconstruction and linkage disequilibrium. We reconstructed network haplotypes in a two-step analysis using the programs SplitsTree4 and Phylovis v.2 (Fig. 3). The concatenated data of VNTR flanking regions and toxA sequences resulted in a 1,629-bp matrix with 35 polymorphic sites, including three singletons. In a first step, using the Neighbor-Net method implemented in SplitsTree4, we reconstructed a split network with four clearly separated groups of isolates, showing little evidence of recombination among them. In a second step, for each group, we reconstructed the network considering different alleles of each haplotype rather than the length of each VNTR locus to avoid homoplastic pitfalls (Supplementary Table S3). The tree cut-off, which is the threshold of number of differences (i.e., different alleles at each locus) below which haplotypes are considered to belong to the same clonal complex, was set to three. When considering only VNTR loci that have 19 different haplotypes and 171 potential links between them, 21 display <3 different alleles. The average number of different alleles for the five VNTR loci between haplotypes is 3.78, reflecting the genetic divergence among haplotypes. The network resulted in five groups and two singletons. Three groups each clustered together two haplotypes that differ between them at a single locus. The two other groups are more divergent, with up to two different alleles among haplotypes.

Considering all isolates, we detected a highly significant linkage disequilibrium among all loci (P < 10−6 for each locus pair). The analysis of molecular variance showed a lack of variety effect when the plot effect was removed in the model (P = 0.26), reflecting the lack of specificity or preference of the various bacterial haplotypes by one of the two varieties.

Discussion

B. glumae is a bacterium that induces the rice panicle blight disease present in Asia and America and can occasionally cause extremely high yield losses. Our study explores for the first time in detail the diversity and genetic structure of bacterial populations in a cultivated rice field. The first step was to isolate bacteria from rice panicles.
Fig. 2. Maximum likelihood phylogeny of the toxoflavin toxA sequence. Tree is based on partial 473-bp sequences. The seven types detected are numbered from 1 to 7, while numbers in parentheses and italics indicate the number of sequences retrieved from GenBank that fall in each clade (thus excluding sequences from our study). Types 2, 3, and 5 clustered sequences are from different countries.

Table 2. Genetic diversity of the five loci retained in the typing scheme of *Burkholderia glumae*.  

<table>
<thead>
<tr>
<th>Locus</th>
<th>Number of alleles</th>
<th>Number of alleles based only on repeat numbers</th>
<th>Number of SNPs(^b) in flanking regions(^c)</th>
<th>Allelic range variation (size of the repeat)</th>
<th>Type of VNTR</th>
<th>PCR product size range (bp)</th>
<th>HGDI(^b) scores</th>
</tr>
</thead>
<tbody>
<tr>
<td>toxA</td>
<td>4</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>–</td>
<td>560</td>
<td>0.698</td>
</tr>
<tr>
<td>BG09</td>
<td>9</td>
<td>8</td>
<td>13</td>
<td>5–19 (9 bp)</td>
<td>C(^b) (4 types)</td>
<td>522–648</td>
<td>0.872</td>
</tr>
<tr>
<td>BGm03</td>
<td>6</td>
<td>4</td>
<td>6</td>
<td>2–5 (10 bp)</td>
<td>P(^b) (1)</td>
<td>404–445</td>
<td>0.745</td>
</tr>
<tr>
<td>BG12</td>
<td>7</td>
<td>5</td>
<td>7</td>
<td>2–11 (8 bp)</td>
<td>P (1)</td>
<td>460–532</td>
<td>0.795</td>
</tr>
<tr>
<td>BG02</td>
<td>16</td>
<td>14</td>
<td>4</td>
<td>2–5 (12 bp) and 27–51 (9 bp)(^d)</td>
<td>P and C (7)</td>
<td>312–528</td>
<td>0.924</td>
</tr>
</tbody>
</table>

\(^a\) The basic statistics were obtained from the Saldaña collection (\(n = 119\)).

\(^b\) SNP, single nucleotide polymorphism; VNTR, variable number of tandem repeats; HGDI, Hunter–Gaston discriminatory index; P, perfect (numbers of SNPs detected within VNTR); C, compound (number of types detected).

\(^c\) From 68 to 394 bp around the VNTR loci.

\(^d\) Locus BG02 includes two VNTR loci in a row.
expected, there is a strong correlation between the severity of the disease and our ability to recover isolates, even if we managed to recover a few isolates from asymptomatic panicles, as done in a previous study from China (Zhu et al. 2008). Bacteria invaded the panicles and the grain and increased in density during infection (Pedraza et al. 2018), which clearly explains our success in recovering more easily isolates from diseased panicles. We were, however, unable to detect any correlation between the genotype of the isolates and the health status of the panicle, although our sampling was not strictly adapted to detect such a link. Pathogenicity tests should be carried out to detect the variation in symptoms induced by each bacterial genotype and to see if some of them are more aggressive than others.

We recovered four different \textit{toxA} haplotypes, involving four SNPs among the 109 isolates. Quite surprisingly, the study from Maeda et al. (2006) typed 67 strains of \textit{B. glumae} samples recovered from all over Japan, and in examining the same portion of the sequence as we did, only three SNPs and four haplotypes were found. The diversity of the partial \textit{toxA} gene found in a single field in Colombia is therefore similar, if not higher, than the diversity found throughout Japan. Moreover, by reconstructing the phylogeny (although again based on a limited number of mutations), it appears clear that there is no global geographical pattern of \textit{toxA} diversity (Fig. 2). For instance, \textit{toxA} type 3 is shared by isolates recovered from the U.S.A. (336gr1), Vietnam (NCPPB3923), Japan (LMG2196 and others), Korea (BGR1), and Colombia (this study). It is also the case for type 2 (Japan and Colombia) and type 5 (Japan, and a lung isolate taken from a hospital in the U.S.A.). This finding demonstrates that, so far, the toxoflavin diversity is widespread over the different regions from which \textit{B. glumae} isolates have been recovered. Because the majority of the mutants detected are silent (i.e., no change in the amino acid), we may reject the hypothesis that they have arisen independently by convergence and selection (i.e., are homoplastic). We can then suggest that if the disease really originated in Asia, the bacterium has spread from Asia to America, either once or repeatedly, without significant genetic bottleneck. Conversely, we can also suggest that Asia may not be the center of origin of the disease, which could be America instead. This strong hypothesis, however, requires a much larger sampling, both in Asia and America, to be tested. Finally, it can be noted that the majority of sequences recovered belong to haplotype 3 (59 sequences out of 87 in total), and that this haplotype is the one with the widest geographical distribution. Whether this predominance is the result of greater competitiveness in its ability to infest the host or to infest a wider host spectrum, to better survive in the environment (which could have led to a recent expansion), or is simply random should be tested by artificial inoculations and large sampling in other parts of the world. It is finally interesting to note that the overall diversity, even over the whole sequence of \textit{toxA}, remains extremely low. This low nucleotide diversity could be the result of high purifying selection because of its role in pathogenicity. We tried to detect evidence of such selection, but the test we applied was not significant; perhaps this was from the very low number of mutations along the sequences, which could restrict the power of this test. Whether these mutations have an effect or not, however, should be studied in depth in the future and their impact in the infection process should be assessed. At the beginning of this project, we faced one main pitfall to select primers: only two genomes (LMG2196 \textsuperscript{2} and BGR1) were completed at the time of this analysis; the three others remained at the draft level (from 96 to 2,457 scaffolds). The fragmentations of genome sequences
are often positioned within the repeated sequences, limiting the possibility of searching for sequences conserved on either side of the VNTR loci to design the primers. We did succeed in designing primers for 14 different loci, but many failed to amplify all of our subsample of isolates. Such a low level of success with our primers was quite disappointing. The primers were not simply designed on the basis of five genomes available in GenBank; we also verified that they were present in three other genomes added after our initial analyses. These eight genomes had been sampled from 1967 (for type strain LMG2196) to 2017 (Bp9029), taken from two continents and seven different countries. We were therefore expecting them to be representative of the species diversity, which was clearly not the case. As suggested earlier (Francis et al. 2013), one explanation for this failure could be that *B. glumae* has a high degree of plasticity in its genomic structure, with several genomic islands. These rearrangements and indels could be from an admixture of several independent lineages newly introduced into the field that will gradually recombine and homogenize over time and generations, or whether *B. glumae* is a species with a very low ability to recombine. One point that could partly resolve the issue is the geographical distribution of haplotypes in the field. After mapping the origin of each haplotype on the network, we found that there is no correlation between genetic proximity and the plot of origin. A plot may contain several divergent haplotypes, just as some haplotypes have been recovered from different plots (Fig. 3). Bacteria may disperse rapidly in the field (which is certainly possible). Alternatively, seeds used for the trials may have been heavily contaminated at the time of sowing. Indeed, previous studies have shown that seeds may contain *B. glumae* isolates (Cottyn et al. 2009; Sayler et al. 2006). However, because there is no effect of the rice variety on the bacterial diversity recovered, we had to hypothesize that the two lots of seeds have been produced under the same bacterial population contamination pressure. Finally, this distribution on the different plots could suggest that these different haplotypes have been present for a long time, and therefore would mean that the degree of recombination among strains must remain very low.

These results suggest key elements in the presence of the bacterium in Colombia. Colombia is a region of major diversity and diversification of *B. glumae* as a rice pathogen, with the genetic diversity observed being the result of migration events from Asia (the region assumed to be the center of origin of the pathogen) and also possibly from other regions where the disease is present, such as northern America. Indeed, the presence of *toxA* haplotypes shared between Asia and Colombia, coupled with the high conservation of sequences of this gene, supports the hypothesis of recurrent migration events between the two continents. This is not surprising, based on the high exchange of rice germplasm for breeding purposes. At the same time, the high number of multi-loci haplotypes and VNTR alleles reflect the diversification process that occurs in Colombia.

The assumption that the seed lots used were already heavily contaminated, the coexistence of several genetically highly divergent bacterial lineages within a field shows that these bacterial lineages are locally maintained and can reinfect a field year after year, either by wind or by irrigation water. Such a dynamic must play a great role in the epidemiology of the disease and its ability to maintain itself over years in a geographic area. It would be particularly relevant to study the extent to which such diversity may have a role in the dynamics of the disease and its adaptability to the different rice varieties that can be deployed in the field. We have not yet tested for differing virulence or aggressiveness in rice cultivars. However, it certainly might be the case, because previous studies have shown that different strains may have variable toxoflavin production, which affects their ability to colonize the plant, and their particular virulence (Karki et al. 2012). It has been also reported that *B. glumae* can act as a biocontrol agent (some strains are producers of antibiotics) and that may help for survival in the seeds and other tissues (Cottyn et al. 2009). Colombia experiences large year-to-year variations in disease incidence and severity. Our results suggest that this is not from annual variations in the diversity of strains present but more likely the result of environmental fluctuations; previous studies have shown that the disease is dependent on particular environmental conditions for its development.
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