The pathogenicity of *Stenotrophomonas maltophilia* is regulated in part by its quorum sensing (QS) system. The main QS signaling molecule in *S. maltophilia* is known as diffusible signal factor (DSF), and the *rpf* gene cluster is responsible for its synthesis and perception. Two cluster variants have been previously described, *rpf*-1 and *rpf*-2, which differ basically in the conditions under which DSF is produced. Here, correlations between the *rpf* variant and antibiotic susceptibility, LPS electrophoretic profiles and virulence-related phenotypes were evaluated for a collection of 78 geographically and genetically diverse clinical strains of *S. maltophilia*. In general there were associations between previously established genogroups and the genetic variant of the *rpf* cluster. However, only few genotype-phenotype correlations could be observed. Resistance to the β-lactam antibiotics ceftazidime and ticarcillin was associated with strains carrying the *rpf*-1 variant, whereas strains of variant *rpf*-2, particularly those of genogroup C, showed higher resistance levels to colistin. Strains of variant *rpf*-2 were also significantly more virulent to *Galleria mellonella* larvae than those of *rpf*-1, most likely due to an increased ability of *rpf*-2 strains to form biofilms. A comparative genomic analysis revealed the presence of proteins unique to individual genogroups. In particular, the strains of genogroup C share an operon that encodes for a new virulence determinant in *S. maltophilia* related to the synthesis of an alternative Flp/Tad pilus. Overall, this study establishes a link between the DSF-based QS system and the virulence and resistance phenotypes in this species, and identifies potential high-risk clones circulating in European hospitals.

**Keywords:** quorum sensing (QS), diffusible signal factor (DSF), MLST (multilocus sequence typing), *rpf* cluster, biofilm, antibiotic resistance
INTRODUCTION

*Stenotrophomonas maltophilia* are metabolically and genetically diverse Gram-negative bacilli belonging to the γ-proteobacteria class that inhabit a wide range of environmental niches, mainly in association with plant rhizospheres (Ryan et al., 2009; Brooke, 2012). Although *S. maltophilia* are thought to be low-grade pathogens and a rather uncommon cause of bacterial disease in humans, they are increasingly recognized as significant opportunistic pathogens in healthcare settings worldwide, being one of the ten most common organisms found in respiratory tissue samples taken from intensive care unit (ICU) patients in Europe (Zuschneid et al., 2007; Gherardi et al., 2015). Several nosocomial outbreaks of *S. maltophilia* infection and/or colonization have been reported (Denton and Kerr, 1998), and the global prevalence and incidence of these bacteria as human pathogens have increased significantly during the last decade (Chang et al., 2015; Nayyar et al., 2017; Rutter et al., 2017), particularly in cystic fibrosis patients (Hatzigorou et al., 2019). The most common clinical manifestations of *S. maltophilia* infections include pneumonia, bacteremia, wound or urinary-tract infections (Looney et al., 2009; Gales et al., 2019), with crude mortality rates ranging from 14 to 69% in patients with bacteremia, being highest among patients receiving inappropriate initial antimicrobial therapy (Paez and Costa, 2008; Falagas et al., 2009; Kim et al., 2019). Most important, *S. maltophilia* show low susceptibility to many antibiotics, including those commonly used to treat infections by Gram-negative opportunistic pathogens (Sánchez, 2015). Not only have they a variety of intrinsic resistance mechanisms affecting almost all antibiotic classes but they are also able to acquire new resistances via horizontal gene transfer and mutations. The phenomenon of heteroresistance has also been described in *S. maltophilia* as a mechanism to withstand antibiotic treatment (Martínez-Servat et al., 2018). For these reasons, this species has been classified as one of the leading multidrug resistant (MDR) organisms in hospital settings (Brooke, 2014) and has now been included in the global priority list of the top 10 resistant microorganisms (TOTEM) isolated in ICUs (Rello et al., 2019).

The pathogenesis of *S. maltophilia* infections involves several virulence factors, including extracellular enzymes, bacterial motility and biofilm formation (Trifonova and Strateva, 2019). The ability to grow in matrix-enclosed biofilms is an important virulence-related trait of pathogenic bacteria. Among other functions, biofilms facilitate the development of bacterial persistence within the host and increase resistance against the host immune response (González et al., 2018) and antimicrobials (Stewart, 2015), including the last-resort antibiotic colistin (Mulcahy et al., 2008; Kim et al., 2015; Chua et al., 2016). Thus, biofilm formation is regarded as a form of phenotypic resistance (Olivares et al., 2013). *S. maltophilia* forms biofilms on a wide range of biotic and abiotic surfaces, such as indwelling medical devices (Di Bonaventura et al., 2007). Still, very little is known about the genetic mechanisms that control biofilm formation or virulence in these bacteria, and it is not clear whether there is an intrinsic difference in biofilm formation among genomically diverse environmental and clinical isolates (Pompilio et al., 2011). *S. maltophilia* produces a wide range of extracellular enzymes, including lipases, proteases, chitinases, esterases, DNases, and RNases, contributing to its colonization and virulence in different organisms (Ryan et al., 2009; Trifonova and Strateva, 2019). Furthermore, it is well known that *S. maltophilia* synthesizes diverse O-specific polysaccharide structures that constitute immunodominant antigens of the lipopolysaccharide (LPS) molecule and may contribute to the pathogenicity of the bacterium (Waters et al., 2007; Knirel, 2011). A diffusible signal factor (DSF)-mediated quorum sensing (QS) system, encoded by the *rpf* gene cluster, coordinates the regulation of a number of phenotypes in *S. maltophilia*, such as biofilm formation, secretion of proteases, virulence, and tolerance to a range of antibiotics (Fouhy et al., 2007; Ryan et al., 2009; Trifonova and Strateva, 2019). Two genetic variants of the *rpf* gene cluster (*rpf-1* and *rpf-2*) divide the strains into two phenotypically and genotypically distinct subpopulations (Huedo et al., 2014). Only *rpf-1* strains produce detectable DSF, which correlates with their ability to control biofilm formation, motility and virulence (Huedo et al., 2014, 2015). However, no clear link between the presence of a specific *rpf* variant and the isolation source of the strains has been found so far (Lira et al., 2017).

An additional important trait of *S. maltophilia* strains is their high genetic diversity (Gherardi et al., 2015) that facilitates adaptation to changing environmental conditions. Numerous genotypic studies of both clinical and environmental strains have revealed considerable variability among *S. maltophilia* isolates (Nicoletti et al., 2011; Pompilio et al., 2011, 2016; Ochoa-Sánchez and Vinuesa, 2017; Patil et al., 2018; Rizek et al., 2018; Steinmann et al., 2018; Bostanghadiri et al., 2019; Mojica et al., 2019). Due to these large genetic divergences, and because previously proposed species are recognized to be closely related to *S. maltophilia*, it is increasingly accepted to present it as the *S. maltophilia* complex (Svensson-Stadler et al., 2012; Patil et al., 2018; Vinuesa et al., 2018). PCR-based multilocus sequence typing (MLST) and whole-genome sequencing (WGS) have clearly shown the existence of several genomic or phylogenetic groups in the *S. maltophilia* complex, and have helped to clarify the taxonomic status of the species (Hauben et al., 1999; Kaiser et al., 2009; Ochoa-Sánchez and Vinuesa, 2017; Steinmann et al., 2018; Gröschel et al., 2020; Mercier-Darty et al., 2020). Even strains isolated from the same patient (Pompilio et al., 2016; Chung et al., 2017) or patients within the same hospital (Valdezate et al., 2004) can belong to distant phylogenetic groups with different phenotypes, probably due to the emergence of adaptive mutations as a result of selective pressure in the clinical environment or inside the host. In addition to interstrain variability, some isolates show cell-to-cell variation within a clonal population as a mechanism to survive environmental fluctuations (Abda et al., 2015). The complex genotypic and phenotypic heterogeneity between and within *S. maltophilia* isolates hampers not only the effectiveness of antimicrobial therapies but also investigations into the virulence and resistance determinants of this species.

To deepen the knowledge and understanding of the resistance and virulence phenotypes displayed by *S. maltophilia*, in a way
that is consistent with this high-diversity context, we have studied a panel of genetically diverse clinical S. maltophilia isolates with focus on the two genetic variants of the DSF-mediated QS system. Genetic characterization was performed using MLST and rpf typing, together with the analysis of shared and unique proteins. Phenotypic analyses included the determination of antibiotic resistance profiles, biofilm formation, protease secretion, cell motility and LPS banding patterns. Moreover, the virulence of the isolates was assessed using two different non-vertebrate model hosts. To our knowledge, a systematic analysis of the population structure based on the two genetic variants of the DSF-mediated QS system and its relationship with the resistance and virulence phenotypes of clinical S. maltophilia strains has not yet been carried out.

**MATERIALS AND METHODS**

**Bacterial Isolates and Growth Conditions**

The strain collection used in this study consisted of 78 clinical S. maltophilia isolates (Huedo et al., 2014) collected from point prevalence studies in ICUs of geographically distant European hospitals. The strain panel included isolates from sputum, blood and swabs from surgical wound, oropharynx, perineum, vascular ulcer, decubitus ulcer, or a bronchoaspirate (Supplementary Table S1). All clinical strains were isolated from different patients, except for five patients with two isolates each (Supplementary Table S1). Three strains of S. maltophilia, namely K279a, D457, and ATCC13637, were included as references. ATCC13637 was purchased from the American Type Culture Collection. Unless otherwise stated, cultures were routinely grown at 37°C in LB broth (10 g/L tryptone, 5 g/L yeast extract and 10 g/L NaCl) on a rotary shaker at 200 rpm. The species of all isolates was confirmed by matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry (data not shown).

**Genome Sequences and Comparative Genomics**

Annotated genome sequences of the unique isolates studied here were generated in the study by Gröschel et al. (2020) and are available from NCBI repository under the accession numbers RAX00000000-RAV00000000. Genome sequences from S. maltophilia strains K279a, D457, and ATCC13637 were obtained from the NCBI genome database. The encoded protein sequences were extracted from the genomes and compared to identify core and exclusive proteins. Orthologous protein sequences of the selected S. maltophilia strains were assessed by complete reciprocal best hit using UCSC blat (Kent, 2002) as comparison tool and accepting hits with a complete graph as previously reported (Callister et al., 2008) or the largest clique. This algorithm clusters the gene-derived proteins into core (hard and soft) and accessory proteins. Hard core proteins are those found in all genomes, while soft core proteins are found in 95–99% of genomes. Blastp and the HMM search program jackhammer (Johnson et al., 2010) were used to identify homologs of the protein sequences using several databases. Prediction of operons was done using the online software and database OperonDB (Pertea et al., 2009).

**Genotypic Characterization and Phylogenetic Analysis**

Genotypic characterization was basically performed on the basis of the MLST scheme for S. maltophilia (Kaiser et al., 2009). In addition, determination of the rpf variant was included in the genotyping as previously described (Huedo et al., 2014). The detailed MLST procedure was obtained from the S. maltophilia MLST database. The allele numbers for each locus and the sequence type (ST) were determined online at the S. maltophilia MLST database. A combination of the allelic sequences of the seven genes yielded the allelic profile for each isolate. Concatenated sequences of the seven housekeeping genes for each strain were aligned, and a phylogenetic tree was constructed using MEGA version 7.0.21 (Kumar et al., 2016) based on the neighbor-joining algorithm (Tamura–Nei model). Phylogenetic trees were visualized and annotated using the interactive web platform iTOL v3 (Letunic and Bork, 2016). To divide strains into clonal groups according to their MLST allelic profiles and to infer the population structure, BURST analysis (Feil et al., 2004) was used on the basis of five shared loci.

**Analysis of Antimicrobial Susceptibility**

The minimum inhibitory concentrations (MICs) of antibiotics were determined using Etest strips following the manufacturer’s instructions (bioMérieux, Madrid, Spain). Briefly, a 0.5-McFarland suspension was used to grow a confluent bacterial lawn on Mueller–Hinton (MH) agar plates (BioMérieux). Seven antibiotics with proven *in vitro* activity against S. maltophilia (Falagas et al., 2008; Wang et al., 2014; Chang et al., 2015) were used: ceftazidime, minocycline, levofloxacin, trimethoprim-sulfamethoxazole, amikacin, ticarcillin-clavulanate, and colistin. The MIC values were determined according to the Etest reading guide after 18 h of incubation at 37°C. The susceptibility test results were interpreted according to clinical susceptibility and resistance breakpoints (S) suggested by the Clinical and Laboratory Standards Institute (CLSI) (CLSI, 2019). For S. maltophilia, clinical breakpoints are now available for trimethoprim-sulfamethoxazole (≤2 μg/mL), ticarcillin-clavulanate (≤16 μg/mL), ceftazidime (≤8 μg/mL), minocycline (≤4 μg/mL), and levofloxacin (≤2 μg/mL).

For amikacin and colistin, CLSI breakpoints for *Pseudomonas aeruginosa* were used with susceptibility cutoffs of 16 and 2 μg/mL, respectively. Isolates with intermediate antibiogram results were considered resistant (non-susceptible). MDR was defined as non-susceptibility to at least one agent in three or more antimicrobial categories (Magiorakos et al., 2012). The MIC50 and MIC90 values represent the MIC value at which ≥50 and ≥90% of the strains within a test population are inhibited, respectively.

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1. https://www.ncbi.nlm.nih.gov/genome/
2. https://blast.ncbi.nlm.nih.gov/Blast.cgi
3. http://pubmlst.org/smaltophilia/
In addition to the Etest method, colistin MICs were determined by the broth microdilution (BMD) method in cation-adjusted MH broth (CAMHB) in accordance with CLSI and the European Committee on Antibiotic Susceptibility Testing (EUCAST) joint recommendations (CLSI, 2019; EUCAST, 2019). Briefly, MICs were determined in sterile 96-well plates by twofold serial dilutions of colistin sulfate (Apollo Scientific Ltd., Batch AS405305) in 100 μL of CAMHB. Bacteria were first grown overnight in CAMHB using CLSI-recommended incubation conditions. After that, 100 μL of the bacterial suspensions with a final optical density at 550 nm (OD550) of 0.05 were added to the wells containing the antibiotic dilutions, and the MIC plates were read after 18 h of incubation at 37°C. The MIC was defined as the lowest antibiotic concentration that inhibited 80% of growth (based on OD measurements) in comparison to the control (CLSI, 2015).

**SDS–PAGE and Immunoblotting of Lipopolysaccharides**

For identification of identical ladder-like LPS patterns, LPS preparations from proteinase K-digested whole-cell lysates of *S. maltophilia* isolates were separated on 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS–PAGE) gels and stained with silver nitrate according to the method of Hitchcock and Brown (1983). The lysates were prepared from the biomass of overnight cultures grown on agar plates. For immunoblots, a rabbit polyclonal antibody against heat-inactivated *S. maltophilia* K279a cells was used. In order to detect serological cross-reactivity between O-specific polysaccharides, the antiserum was adsorbed with heat-killed cells of a K279a ΔrmtlBACD mutant as previously described (Steinmann et al., 2018). The LPS samples were electrotransferred from SDS–PAGE gels onto polyvinylidene difluoride (PVDF) membranes (Millipore, Merck, Darmstadt, Germany), followed by incubation of the blots with adsorbed anti-K279a antibody. The immunoblots were then treated with alkaline phosphatase-conjugated AffiniPure goat anti-rabbit immunoglobulin (Ig)G (H + L) (Dianova, Hamburg, Germany) and developed in the presence of nitroblue tetrazolium and 5-bromo-4-chloro-3-indolylphosphate substrate (Promega).

**Production of Extracellular Protease**

Extracellular proteolytic activity was assessed by means of casein hydrolysis on LB agar supplemented with 1% (w/v) skimmed milk. The bacterial suspensions (5 μL each with an OD550 of 0.5) were spotted onto the agar, and the plates were incubated at 30°C for 24 h. Proteolytic capacity was evaluated by measuring the diameter of the translucent halos.

**Twitching Motility**

Twitching was assessed using subagar stab inoculations (stab assay) from fresh overnight LB agar plates as previously described (Rashid and Kornberg, 2000). To prepare twitching plates, LB medium containing 1% (w/v) Noble agar (DifcoTM) was used. After incubation at 30°C for 24 h, the agar was removed from the plates, and the twitching zones were visualized by staining with 1% crystal violet (w/v) to measure their diameters. All experiments were conducted in triplicate.

**Biofilm Formation Assays**

To determine biofilm formation, overnight cultures of each isolate were grown aerobically (200 rpm) in LB at 30°C, followed by dilution of the cultures into fresh LB to an OD550 of 0.1. Sterile untreated 96-well microtiter plates (BrandTech 781662) were inoculated with 200 μL of the bacterial suspensions and incubated at 30°C for 24 h. Prior to biofilm quantification, cell growth was estimated in each well by measuring the optical density at 620 nm (OD620) using a microplate reader (Multilabel Plater Reader VICTOR3). Quantification of the amount of biofilm was performed by crystal violet (CV) staining as described previously (Huego et al., 2014). Briefly, wells containing adhered cells were washed three times with water, fixed at 60°C for 1 h and stained for 15 min with 200 μL of a 0.1% CV solution. The stained biofilms were rinsed with distilled water, allowed to dry at 37°C for 30 min and then extracted with 200 μL of 95% ethanol. The amount of biofilm was quantified by measuring the OD550 of dissolved CV using a microplate reader. Biofilm formation (OD550 of CV) was normalized to cell growth (OD620) and reported as relative biofilm formation. For this quantitative assay, we have used six to eight replicate wells for each isolate. Based on the ratio (OD550 of CV)/(OD620 of cell growth), the isolates were grouped into three categories to classify their ability to form biofilm: weak (0 to 0.85), moderate (0.85 to 1.5), or strong (> 1.5) biofilm formers (Zhuo et al., 2014).

**Virulence in Caenorhabditis elegans**

Determination of the virulence of *S. maltophilia* strains in the *C. elegans* CF512 infection model was based on the “slow killing” method (Tan et al., 1999). *C. elegans* CF512 [her-15(b26)II; fem-1(hc17)JV], a strain showing temperature-dependent sterility, was provided by the Caenorhabditis Genetics Center (CGC) and routinely maintained on NGM plates (1.7% agar, 50 mM NaCl, 0.25% peptone, 1 mM CaCl2, 5 μg/mL cholesterol, 25 mM KH2PO4, 1 mM MgSO4) seeded with *Escherichia coli* OP50 (provided by CGC) at 16°C. Bacterial strains were grown in Brain Heart Infusion (BHI) broth (BD Difco Bacto) overnight at 37°C, and 100 μL of each culture was spread onto a 5.5-cm-diameter NGM agar plate and incubated at 37°C for 24 h. Before usage, plates were cooled down to room temperature, and 15–20 adult hermaprodite CF512 worms were placed onto plates seeded with the bacterium of interest. Plates were incubated at 25°C (at which worms are sterile) and scored for live worms every 24 h. *E. coli* OP50 was used as a negative control. A worm was considered dead when it no longer responded to touch. Three replicates per strain were prepared.

**Virulence in Galleria mellonella**

Larvae of *Galleria mellonella* were obtained from our own hatchery, which was established in collaboration with Professor Fernando García del Pino from the Zoology Department at the Universitat Autònoma de Barcelona. To prepare bacterial inocula, *S. maltophilia* isolates were grown overnight at 37°C in 10 mL of BHI in a rotary shaker at 200 rpm. Then, cells were...
sedimented by centrifugation, washed in PBS and adjusted to contain approximately $5 \times 10^5$ CFU in a single dose of 5 µl. Bacterial burden of the doses was confirmed by plating on BHI medium. This inoculum size was selected based on previous studies (Betts et al., 2014; García et al., 2015) that have determined the optimal dose of S. maltophilia required to kill G. mellonella over a 24–96 h period. Ten larvae per group were infected through the left proleg using a 50-µl Hamilton® Microliter™ syringe with the aforementioned inoculum and incubated in the dark at 30°C in empty petri dishes. Mortality was determined every 24 h. Larvae were considered dead when they no longer responded to touch, which correlates with total melanization (black colored larvae).

Statistical Analysis
Statistical comparisons between groups were performed using Welch’s $t$-test, i.e., assuming non-equal variance. The associations between two categorical phenotype and genotype variables were investigated using the z-test or Fisher’s exact test as appropriate (Fisher’s exact test works better with small sample sizes). The odds ratio (OR) with a corresponding 95% confidence interval (95% CI) for each variable was calculated. In all cases a $p$-value < 0.05 was considered statistically significant, and all tests were two-tailed. Data were analyzed using SPSS 15.0 for Windows (SPSS Inc., Chicago, IL, United States) or GraphPad Prism 5 (GraphPad Software, San Diego, CA, United States). For infection experiments, the time at which 50% of animals were scored as dead (LT50), was calculated using a non-linear regression based on the Hill equation (GraphPad Prism 5).

RESULTS
Genetic Diversity of the S. maltophilia Strains
To evaluate the genetic diversity and possible relations between the S. maltophilia isolates involved in this study, their MLST profiles were first analyzed. The 78 strains, isolated from patients of different European countries, mapped to 37 STs (Figure 1 and Supplementary Table S1). At the date of submission to the S. maltophilia MLST database, 30 of these STs were new and designated ST-76 to ST-120 to ST-143. Fifteen isolates (19.2%) belonged to seven previously described STs, whereas 63 (80.8%) belonged to the 30 novel STs, corroborating previous reports on the genotypic heterogeneity of S. maltophilia. Although no predominant clonal group was identified by BURST analysis, the three most frequent MLST genotypes were ST-77 (8 isolates), ST-132 (8 isolates), and ST-133 (7 isolates). The remaining STs occurred each with a frequency lower than 5.0%. When the entire S. maltophilia MLST database (522 isolates, 367 STs) was analyzed with the BURST algorithm, some of our isolates were part of discrete clonal groups, sharing six (clonal complexes, the most exclusive group definition) to five identical alleles (Supplementary Figure S1). For instance, four strains isolated from three different countries could be grouped into one of the few clonal groups (ST-8) detected so far in the global population of S. maltophilia (Supplementary Table S1 and Supplementary Figure S1).
Supplementary Figure S1). Likewise, our predominant ST (ST-77) comprises strains from different European countries and is part of a genetically related group of the clonal complex ST-103 (Supplementary Figure S1). In addition to MLST, rpf genotyping (Huedo et al., 2014) revealed that the rpf-1 variant was present in a total of 47 strains (60%), while rpf-2 was identified in 31 strains (40%) (Supplementary Table S1).

Among the 78 strains studied here, five pairs of isolates were confirmed to be epidemiologically linked. They were either cultured from two sources of the same patient on the same day (four pairs) or originated from samples taken from the same patient at different dates (pair OS91 and OS87) (Supplementary Table S1). In most cases, the paired isolates share the same ST, but isolates PU101 and PU102 are genetically distinct, with only three alleles in common. Also, based on the MLST typing and data from the source of isolation, among the 35 strains collected from Germany, and coming from a single hospital, there seems to be genetically related groups (Supplementary Table S1). In order to avoid interference of clonal strains with correlation tests, those isolates displaying identical MLST type and isolation data (i.e., same source, patient, and date) were considered as the same clone, so that only one representative isolate was selected for further investigations (Supplementary Table S1, highlighted in bold). However, those clonal strains that showed different antibiotic susceptibility profiles (see below) were not excluded from further analyses. Thus, the analysis of correlations between the phenotypic and genotypic parameters studied here was performed on 58 isolates. In addition, we have included the strains K279a, D457, and ATCC13637 as references in all experiments, yielding a total of 61 “unique strains” (36 rpf-1 and 25 rpf-2).

Classification of S. maltophilia Isolates Into Genomic Groups

For cluster analysis, taxonomic status verification and assignment of the S. maltophilia isolates to genomic groups, phylogenetic analyses were carried out based primarily on concatenation of the seven MLST genes (Figure 1 and Supplementary Figure S2). A neighbor-joining tree of the 522 isolates in the MLST database (Supplementary Figure S2) allowed us to cluster all these strains according to the 20 genogroups (#1 to #10 and A to J) as previously established by Hauben et al. (1999), Kaiser et al. (2009), and Mercier-Darty et al. (2020). This phylogenetic analysis showed a lack of clustering by isolation source or geographical origin (Supplementary Figure S2), which stresses again the heterogeneity of S. maltophilia isolates, in this case at a worldwide level. Even though most of the strains in the MLST database were of clinical origin, it is noteworthy that almost all environmental strains were clustered in groups #8, #9, and #10. The 78 strains in our panel could be clustered into 12 genogroups (Supplementary Table S1), including the new group G (4 strains) by Mercier-Darty et al. (2020). Therefore, our strain collection provides a good representation of the global diversity of S. maltophilia. Nineteen (31%) of the 61 unique isolates were clustered in genogroup #6, including the clonal complex ST-8 (Figure 1). Genomic group #6 comprises the so far largest number of strains in the MLST database (Supplementary Figure S2) and includes the reference strain ATCC13637 and the reference genome strain K279a. The other genogroups with a significant number of isolates in our collection were #2 (19%), #3 (13%) and C (15%), which is in agreement with the frequency of isolates belonging to these groups in the worldwide population. In agreement with previous studies (Corlouer et al., 2017; Mercier-Darty et al., 2020), genogroup #2 represents the most genetically diverse group in our collection. As proposed by Gröschel et al. (2020) and shown in Supplementary Table S1, the strains could be further classified into the monophyletic lineages named Sm1–Sm18 (termed S. maltophilia sensu lato) or the more distantly related lineages Sgn1–Sgn4 according to Vinuesa et al. (2018). All our strains, except the phylogenetically distant strain OU111 (genogroup A), belong to the S. maltophilia sensu lato cluster. None of them clustered into lineages in which there are isolates mislabeled as S. maltophilia or with an unclear taxonomic status, for example the genomic groups #8 and #10 (Ochoa-Sánchez and Vinuesa, 2017). In this respect, our results therefore could provide support for the multispecies concept to explain diversity among isolates of S. maltophilia.

Interestingly, there were statistically significant associations between the major MLST-based genomic groups and the rpf variants. In the genomic group #6, most of the strains contained the variant rpf-1 (P < 0.001), whereas the variant rpf-2 was predominant in genogroups C (P < 0.001) and #3 (P < 0.05).

Identification of Core and Unique Proteins Within S. maltophilia Strains and Genogroups

In addition to the genotypic characterization, the genome-derived protein sequences of 60 unique strains of S. maltophilia were compared to identify core and exclusive proteins (assembled genome sequence of PR59 could not be determined). According to our ortholog identification method, the S. maltophilia strains included in this comparison share a core genome of 2,585 proteins. If we consider the soft core genome (proteins present in 95–99% of genomes), 349 additional proteins could be added to this set of shared proteins. We also identified the number of exclusive proteins shared by all members of each genogroup (Figure 2A and Supplementary Table S2). The analysis showed that the number of unique proteins to each genogroup varies considerably from genogroup to genogroup. Genogroups #5, #7, D and A clearly stand out having the highest number of unique proteins; this is likely due to these groups being represented by a single strain. It is worth mentioning that all members of genogroup C share 17 unique proteins, 12 of which are part of an operon that encodes a protein translocation system for the assembly of Flp or Tad (tight adherence) pilus (Figure 2B). Moreover, members of genogroup #4 share a unique operon for exopolsaccharide synthesis, and isolates in genogroup G share 41 proteins including peptidases and a transcriptional regulator, in addition to other unique enzymes and several hypothetical proteins (Supplementary Table S2). On the other hand, the comparative analysis did not identify proteins shared exclusively
by strains of the rpf-1 or rpf-2 variants. This result highlights the genetic heterogeneity among clinical S. maltophilia isolates and confirms, in addition to their distribution along genotypes, that the origin of the two rpf variants in S. maltophilia can be explained by horizontal gene transfer.

Diversity of S. maltophilia Isolates Based on Their LPS Profiles

The enormous structural diversity of surface-exposed O-specific polysaccharides of smooth-type LPS is assumed to result from adaptive mechanisms that enable the bacteria to withstand specific conditions in various environmental niches, which in turn provides the possibility of using O-polysaccharides as an epidemiological marker to discriminate between related and unrelated strains within a given species (Rimler, 1990; Aucken and Pitt, 1993; Santamaria et al., 1998). We therefore compared the LPS profiles of our S. maltophilia isolates as an additional marker to cluster the strains on the basis of identical LPS patterns in SDS-PAGE, the ladder-like appearance of which is caused by varying numbers of oligosaccharide repeating units in the O-specific polysaccharide (Supplementary Figure S3). As expected, we could observe significant heterogeneity of the LPS banding patterns, whereby none of the isolates investigated here lacked the O-specific polysaccharide (Supplementary Figure S3). This was also valid for the strains B5565, H5726 and PU126, but unequivocal evidence for the presence of smooth-type LPS could only be obtained using increasing amounts of LPS in SDS-PAGE (Supplementary Figure S3, inset). The three strains were therefore excluded from further analysis of LPS profiles. Based on the electrophoretic mobility of higher molecular mass LPS species, we assigned 76 strains to a total of 31 LPS groups, including 20 isolates with apparently unique LPS patterns (Supplementary Table S1). There were at least two different LPS profiles in each of the genogroups with more than one member, where genogroups F and C with dominating LPS groups 21 and 15, respectively, were the most homogeneous. In summary, apart from some exceptions, we could uncover a good correlation between the LPS profile and the ST of the isolates, which was largely in accord with the phylogenetic tree shown in Figure 2. Furthermore, the LPS samples were screened with a polyclonal antiserum raised against the O-antigen of strain K279a, which has been recently proposed to share similar characteristics with the S. maltophilia O8-antigen of a branched tetra-saccharide with three rhamnopyranosyl residues in the main chain and 3-O-methylxylose as substituent (Steinmann et al., 2018) (Supplementary Figure S3). Our immunoblot analyses indicated serological cross-reactivity of the antiserum with the O-antigens of all strains of LPS profile group 2 (including S. maltophilia K279a), which accounts for 14.5% of all examined isolates. Interestingly, the LPS of all strains of genogroup #4 (3.9%) cross-reacted as well with the antibody, suggesting that these isolates, despite showing different LPS banding patterns, expressed an O-specific polysaccharide either structurally similar or even identical to that of the reference strain K279a. Of note, each of the ten separate groups comprising two to six clonal strains did not only display an identical MLST type, but could also be assigned to one and the same LPS group, which supported the conclusion of epidemiologically linked strains that were isolated at the same hospital in Germany (Supplementary Table S1). All strains that reacted with the O-specific K279a antiserum carried the rpf-1 cluster, except for strain OC323 (genetically close to K279a) which encoded the rpf-2 variant (Figure 1). This could be in fact another indication that the two rpf variants in

![Figure 2](image-url)
correlation between antibiotic resistance phenotypes and genotypes

The antibiotic resistance pattern of our isolates was evaluated with the β-lactam antibiotics ceftazidime and ticarcillin-clavulanate, and the non-β-lactam antimicrobials amikacin, trimethoprim-sulfamethoxazole, levofloxacin, minocycline and colistin (MIC values in Supplementary Table S1). The MIC values of all antimicrobials were determined by the Etest, with the exception of colistin, which was additionally tested using the BMD method (see section “Materials and Methods”). All strains were susceptible to minocycline, and only 6.5 and 19.7% of the unique strains were non-susceptible to trimethoprim-sulfamethoxazole and levofloxacin, respectively. Conversely, high resistance rates were detected against the antibiotics amikacin (54.1%), ticarcillin-clavulanate (62.3%), and ceftazidime (62.3%), all of them with MIC values reaching >256 µg/mL (Table 1).

For colistin susceptibility testing, as expected (Moskowitz et al., 2010), the results of the BMD and Etest methods were significantly different. For our strain collection, the Etest resulted in a colistin MIC50 value of 2 µg/mL corresponding to susceptible strains (susceptibility breakpoint ≤ 2 µg/mL), whereas the BMD method yielded a colistin MIC50 value above the susceptibility breakpoint (>2 µg/mL) (Table 1). Moreover, while the colistin MIC90 value determined by both methods was consistent with a resistant phenotype, it was eightfold lower by the Etest than by the BMD method. This is due to the phenomenon of heteroresistance to colistin displayed by most of the S. maltophilia isolates (Martinez-Servat et al., 2018). Finally, taking the recommended BMD method as a reference, 67.3% of the strains could be considered as non-susceptible to colistin.

To further characterize the resistance patterns, the criteria suggested by Magiorakos et al. (2012) for related non-fermenting Gram-negative bacteria were applied. Among the 61 unique isolates, 37 (60.7%) were found to be MDR strains, whereas 24 (39.3%) were not. On the other hand, 22 different antibiotic resistance profiles were observed among the 61 strains (Supplementary Table S1 and Figure 1), the most common being those that combine resistance to ceftazidime, amikacin, ticarcillin-clavulanate, and colistin. Notably, in three out of five strain pairs isolated from the same patient, the members of the same pair showed different resistance profiles (Supplementary Table S1). The two genetically different strains isolated from the oropharinx and the perineum of the same patient, PU101 and PU102, displayed the same resistance profile. These strains were also assigned to different LPS groups. Strain OL11 was the most resistant isolate in our panel (six out of seven antibiotics). It had been isolated with PL12 from different anatomical sites of the same patient. The two strains are genetically identical, possess the same LPS profile (Supplementary Table S1) and are resistant to almost all antibiotics, except that PL12 showed susceptibility to colistin. On the other end, strains OU152, OU343, and OL16, all belonging to different and distant genogroups (Figure 1), were susceptible to all antibiotics tested (Supplementary Table S1). Overall, no significant correlations were observed between the resistance profiles (antibiotic resistance combinations) and the MLST genotypes. However, statistically significant associations could be observed between specific genomic groups and the susceptibility of the isolates to certain antibiotics (Supplementary Table S3). For instance, almost all strains of the genogroup #6 were susceptible to ticarcillin-clavulanate, whereas strains of genogroups #2 and F were mostly susceptible to colistin but ceftazidime resistant. The genetically related strains of genogroups #2 and F showed similar resistance profiles and were therefore considered as a whole in association studies. In addition, we found high-level resistance to levofloxacin in all strains of the new genogroup G. Almost all strains of the genogroup C were susceptible to the β-lactams ceftazidime and ticarcillin-clavulanate, but were quantitatively more resistant to colistin (Supplementary Table S3 and Figure 1). Interestingly, the variant of the rpf cluster showed some correlation with the resistance profile of the isolates. Most of the rpf-1 strains turned out to be resistant to ceftazidime (P = 0.014), and, from a quantitative point of view, the rpf-2 resistant strains inclined toward higher MIC values for colistin compared to the rpf-1 variant, although without statistical significance (Figure 3A). This analysis also showed higher resistance levels to ceftazidime and ticarcillin-clavulanate among resistant strains of the rpf-1 variant.

Correlation Between Genotypes and Virulence-Related Phenotypes

Virulence-associated phenotypes, including biofilm formation, production of extracellular enzymes and motility, were finally assayed for our strain collection. All isolates were protease and biofilm producers (Supplementary Figure S4). However, the amount of biofilm formed in vitro varied considerably among S. maltophilia isolates. The rpf-2 strain OU110 exhibited the highest biofilm-forming ability. Most of the isolates (44.3%, 27/61) produced moderate biofilms; with a mean of 1.14 and standard deviation of ±0.43) was observed in 24.6% (15/61) of the isolates, whereas the remaining 19 strains (31.1%) showed weak biofilm formation capacity (mean: 0.38, SD: ±0.24). With respect to genotypes, we found statistically significant associations (P < 0.005) between the isolates of genomic group C and their ability to form biofilm (Supplementary Table S4). On the contrary, 90.9% of strains of genomic groups #6, F and #2 were moderate to weak biofilm producers, though without significant associations. Once again, despite variations in phenotypic traits, the biofilm-forming ability of the strains could be related to the rpf variant. Quantitatively, the biofilm formation capacity of rpf-2 strains (relative biofilm mean 1.38) was significantly higher (P < 0.05) than that of rpf-1 strains (relative biofilm mean 0.92) (Figure 3B). With respect to the number of isolates, strains of variant rpf-1 are mostly moderate to weak biofilm formers (P < 0.001).
### TABLE 1 | In vitro activity (µg/mL) of seven antimicrobial agents against the 61 “unique” strains of S. maltophilia studied here.

| Method | Agent Class or sub-class | MIC50<sup>a</sup> | MIC90<sup>a</sup> | Range | % of susceptible isolates<sup>b</sup> |
|--------|--------------------------|------------------|-----------------|-------|-----------------------------|
| Etest  | Ceftazidime (CAZ)        | Cephalosporins   | 16              | >256  | 1–>256                      | 37.7 |
|        | Minocycline (MIN)        | Tetracyclines    | 0.25            | 0.5   | 0.032–2                     | 100  |
|        | Levofloxacin (LEV)       | Fluoroquinolones | 0.5             | 16    | 0.125–>32                   | 80.3 |
|        | Trimethoprim-sulfamethoxazole (SXT) | Folate pathway inhibitors | 0.25 | 0.5 | 0.125–>32 | 93.5 |
|        | Amikacin (AMK)           | Aminoglycosides  | 32              | 256   | 2–>256                      | 45.9 |
|        | Ticarcillin-clavulanate (TTC) | Penicillins     | 64              | >256  | 0.5–>256                    | 37.7 |
|        | Colistin (COL)           | Polymyxins      | 2               | 8     | 0.064–32                    | 75.4 |
| BMD<sup>c</sup> | Colistin                | Polymyxins      | 8               | 64    | <0.25–>256                  | 32.7 |

<sup>a</sup>MIC90 and MIC50 values were defined as the lowest concentration of the antibiotic on a two-fold scale at which 90 and 50% of the isolates were inhibited, respectively.

<sup>b</sup>Susceptibility breakpoints for S. maltophilia established by the Clinical and Laboratory Standards Institute (CLSI, 2019). For amikacin and colistin, CLSI breakpoints for P. aeruginosa were used. Intermediate and resistant strains were grouped together into a non-susceptible category.

<sup>c</sup>Broth microdilution.

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**FIGURE 3** | Phenotypic resistance and virulence-related phenotypes within each rpf-variant subpopulation of S. maltophilia. (A) Box and whisker plots (range bars indicate maximum and minimum values) showing log2-fold change increase from susceptibility cut-off for each antibiotic: CAZ, ceftazidime; LEV, levofloxacin; AMK, amikacin; TTC, ticarcillin-clavulanate; COL, colistin. Only resistant isolates were included. (B) Profiles for virulence traits of isolates from different rpf-variant subpopulations (*p < 0.05; unpaired t-test with Welch’s correction).

( Supplementary Table S5). On the other hand, there was no statistically significant correlation of protease secretion and twitching motility with the phylogenetic clustering, except for strains belonging to genogroup C or those that are of the rpf-1 variant, which are less casein exoprotease producers (Supplementary Figure S4 and Supplementary Tables S4, S5).

The virulence capacity of the strains was investigated by infecting both the nematode C. elegans and larvae of the insect G. mellonella. The results were compared and correlated with the assayed virulence-related phenotypes. The S. maltophilia strains evaluated in this study were differentially virulent to C. elegans in the slow killing assay, with no correlation found with genomic groups (Supplementary Figure S4 and Supplementary Table S4). The strains required 4–10 ten days to kill 50% of the worms (mean LT50 = 6.9 days). Complete killing by E. coli OP50, used as a negative control, was achieved after 15 days. No significant correlation was found between the assayed virulence-related phenotypes and the virulence of the strains in the C. elegans model. The two least virulent strains in this model, PC274 (LT50 = 10.5 days)
and the type strain ATCC13637 (LT50 = 10.0 days), showed low motility and biofilm formation capacity and produced lower amounts of extracellular proteases. Nevertheless, the most virulent strain EV5520 (LT50 = 3.8 days) displayed also weak virulence-related phenotypes, indicating that virulence factors other than those examined here contribute to the infection process in the nematodes. In general the type strain ATCC13637 showed obvious signs of attenuation in all assays (Supplementary Figure S4), as previously reported using a zebrafish infection model (Ferrer-Navarro et al., 2013), most likely due to numerous in vitro passages since it was isolated in 1960.

All S. maltophilia isolates were virulent to G. mellonella larvae in killing assays. The selected inoculum size of $5 \times 10^5$ CFU resulted in 20–60% death at 24 h post-inoculation with most strains, while complete killing was observed after 96 h. Some strains were considered to be highly virulent as they were able to kill approximately 60% of the larvae after 24 h of inoculation and reached 100% after 48 h. These highly virulent strains belong to almost all genogroups and rpf variants. Nevertheless, on average, rpf-2 strains were more virulent than rpf-1 strains ($P = 0.05$) on the basis of LT50 values (Figure 3B). The association results showed again no significant correlation between virulence-related phenotypes and the capacity of the strains to kill the larvae.

**DISCUSSION**

In recent years S. maltophilia has received considerable attention. First, the frequency of MDR phenotypes found among isolates is particularly high, as demonstrated by the panel studied here. Second, S. maltophilia is capable of surviving in a wide variety of environmental niches, partly due to its capacity to form biofilms on diverse biotic and abiotic surfaces, including medical devices. Finally, S. maltophilia is characterized by a high rate of genomic re-arrangements and hypermutator activity, which allows rapid adaptation to new niches (Berg and Martinez, 2015). With this presentation card, it does not come as a surprise that, despite not being a particularly pathogenic or virulent species, S. maltophilia has found its way as an emerging nosocomial pathogen (Brooke, 2012; Adegoke et al., 2017).

Although 60.7% of the clinical strains investigated in this work were MDR, most isolates were susceptible to trimethoprim-sulphamethoxazole and all to minocycline. Monotherapy with trimethoprim-sulphamethoxazole, fluoroquinolones or tetracyclines such as tigecycline, minocycline, and doxycycline remains the most effective antimicrobial treatment against this organism (Chang et al., 2015; Hand et al., 2016). However, resistance to levofloxacin was detected in almost one fifth of the strains evaluated in this study. Even though trimethoprim-sulphamethoxazole is indeed the first choice antibiotic against S. maltophilia, resistance to this drug has been observed in 6.5% of the strains included in this work, as it has also been reported in other studies (Valdezate et al., 2001). Minocycline emerges as the best alternative to treat S. maltophilia, but more studies are needed to introduce this antibiotic into clinical practice (Hand et al., 2016). On the other hand, colistin is usually the last resort antibiotic for treatment of infections caused by MDR strains, despite its proven toxicity (Petrosillo et al., 2008; Wood et al., 2010). In the present study, in addition to a high proportion of colistin-resistant isolates, we have shown that S. maltophilia was capable of developing colistin-resistance heterogeneity, deduced from the observed discrepancies between the two commonly used susceptibility testing methods (Etest vs. BMD) (Lo-Ten-Foe et al., 2007). This is in agreement with our previous, more detailed studies on heteroresistance with four colistin resistant S. maltophilia strains (Martinez-Servat et al., 2018). Based on the strain characterization presented here, this phenotypic heterogeneity seems to be widely spread among clinical isolates regardless of their genotypes and resistance profiles. Due to the emergence of resistances to almost all antibiotic classes and the observed heteroresistance phenomenon, combination therapies and novel agents may soon be needed to treat infections caused by MDR S. maltophilia strains.

An important finding of this study is that resistance to some antibiotics appears to be influenced by the rpf variant. For instance, while the rpf-1 subpopulation displayed higher levels of resistance to the β-lactam antibiotics ceftazidime and ticarcillin, strains of the rpf-2 group showed higher resistance levels to colistin (Figure 3A). The differential resistance profiles found for the rpf-1 and rpf-2 subpopulations may be attributable to inherent DSF production and its lack in rpf-1 and rpf-2 subpopulations, respectively (Hudeau et al., 2014). While rpf-1 strains synthesize DSF under standard conditions activating the QS, rpf-2 strains require detection of exogenous DSF to trigger DSF production (Hudeau et al., 2015). In this line, it has been reported that DSF signals stimulate the production and release of outer membrane vesicles (OMVs), which may contain β-lactamases (Devos et al., 2015). Furthermore, some antibiotic resistance mechanisms seem to be controlled by DSF in the plant-growth-promoting model strain S. maltophilia R551-3 (rpf-1), including the biosynthesis of β-lactamases and multidrug efflux pumps (Alavi et al., 2013). On the other hand, in time-kill curve experiments, S. maltophilia's DSF seemed to influence polymyxin tolerance in P. aeruginosa, with addition of DSF leading to an enhanced tolerance to polymyxin B and colistin (Ryan et al., 2008). We have recently shown that naturally occurring DSF signals and synthetic derivatives potentiate the activity of the polymyxin antibiotic colistin against S. maltophilia and some members of the Burkholderia cepacia complex (Hudeau et al., 2019). Accumulation of DSF on the surface may slightly increase the permeabilization of the bacterial membrane due to its lipophilic nature, potentiating colistin activity against rpf-1 strains. Worth mentioning, there appear to be associations between MLST genotypes and antibiotic resistance regardless of the rpf cluster variant. Strains of genogroup C were mostly susceptible to ticarcillin-clavulanate in addition to ceftazidime. Moreover, all strains of genogroups #4 (rpf-1) and G (rpf-2) were resistant to levofloxacin, and most strains of genogroups #2 and F (mostly rpf-1) were susceptible to amikacin. A significant proportion of β-lactam-resistant isolates arise in S. maltophilia through altered expression...
of β-lactamases (Sánchez, 2015) or amino acid substitutions in efflux pumps (Blanco et al., 2019). Both intrinsic and acquired resistance mechanisms are also involved in resistance to aminoglycosides and quinolones in S. maltophilia, such as expression of multidrug-resistant efflux pumps, presence of antibiotic inactivating enzymes or mutations in the genes coding for antibiotic targets (García-León et al., 2014; Sánchez, 2015). The comparative genomic analysis found no unique proteins shared by members of a genomic group that would be involved, in a straightforward, direct way, in antibiotic resistance (Supplementary Table S2). Taken together, these findings confirm that mutations, together with horizontal gene transfer, affect the resistance patterns in S. maltophilia isolates beyond genomic diversity.

In addition to antibiotic resistance, the genetic variant of the DSF QS system has an impact on pathogenicity of S. maltophilia as well, and many virulence determinants are involved in this process. Biofilm formation is associated with an increase in virulence of S. maltophilia. Although the biofilm forming capacity varied considerably among the strains studied here, consistent with previous observations from our group (Huedo et al., 2014), we have shown that strains carrying the rpf-2 variant are stronger biofilm producers than strains carrying the rpf-1 version (Figure 3B). In a previous study, we showed that, although the protein variant RpfF-2 is functional in DSF synthesis, its permanent repression by RpfC-2 results in the absence of DSF production under the conditions tested. In contrast, the rpf-1 subpopulation produces larger amounts of DSF and, in that case, DSF seems to contribute to biofilm dispersion (Huedo et al., 2014). Similar results were observed for the closely related bacterium Xanthomonas campestris (encoding a very close RpfF-1 variant), in which DSF also appears to disperse biofilms (Dow et al., 2003). Likewise, in P. aeruginosa, a related DSF signal, formerly termed cis-DA, displayed a strong dispersion capacity of mature biofilms of a large number of bacterial pathogens (Davies and Marques, 2009). Overall, it seems likely that the absence of DSF production confers on rpf-2 members a stronger ability to form biofilms, and this may partially explain why these strains are more virulent in larvae of G. mellonella. Pompilio et al. (2016) observed that S. maltophilia biofilm formation is directly associated with mortality rate in G. mellonella. In general, DSF signaling in S. maltophilia regulates factors contributing to virulence, such as motility, biofilm formation and colonization (Alavi et al., 2013; An and Tang, 2018). However, no correlation could be detected when comparing virulence and biofilm formation with casein proteolytic activity or twitching motility, suggesting that the latter attributes are not major virulence traits of S. maltophilia, at least in the two infection models tested. On the other hand, no significant association between antibiotic resistance and the amount of biofilm formed could be observed. Thus, our results are similar to those reported by Pompilio et al. (2016) but diverge from those reported by Liaw et al. (2010), who concluded that MDR isolates show higher levels of biofilm formation in comparison to non-MDR strains.

In S. maltophilia, the high intra-specific genetic variability, together with a low diversity of the known resistance and virulence factors (Berg and Martínez, 2015) significantly impedes the establishment of relationships with epidemiological relevance. Although we could not uncover significant correlations between all phylogenetic groups and virulence-related phenotypes, we have identified groups of genetically related strains that are scattered throughout European countries and could represent high-risk clones. For instance, although the number of strains tested is small, it can be observed that genogroup C comprises stronger biofilm-forming strains that are almost all resistant to the last-resort antibiotic colistin. However, all these strains are susceptible to other antibiotics tested herein and, thus, there remain good treatment options against infections caused by them. Nevertheless, strains of this group associated with human infections (Supplementary Table S1) have also proven to be highly virulent in different infection models (Ferrer-Navarro et al., 2013; Huedo et al., 2014), and can quickly acquire a resistant phenotype (Alonso and Martínez, 1997; Martínez-Servat et al., 2018). Interestingly, all strains of group C seem to express an additional type of the Flp/Tad pilus encoded by a unique cluster of genes (operon SMD_0948-SMD_0960 in D457). This operon seems to differ from the TadE-like pil gene cluster located at Smnt2867-Smnt2875 in the K279a genome (Crossman et al., 2008), although some components are similar. In strain D457, the closest ortholog to the Flp/Tad pilus in K279a is the system encoded by operon SMD_2505-SMD_2513. The Flp/Tad pilus has been categorized as a subtype of the type II secretion system (Tomich et al., 2007), and the encoding cluster is present in a wide variety of bacterial species, for which it functions as an important virulence factor. It is considered a target of horizontal gene transfer and is essential for biofilm formation (Tomich et al., 2007; Nykyri et al., 2013). To our knowledge, it is the first time that this alternative structure has been found encoded in the genome of strains of S. maltophilia, which could contribute to the virulence of these strains. Besides, the genogroup #6 includes the largest subgroup of our strain collection. Of note, the predominance of clinical isolates in genogroup #6 has also been shown by others, mostly in cystic fibrosis patients (Kaiser et al., 2009; Vasilieuska-Schulz et al., 2011; Corlouer et al., 2017; Ochoa-Sánchez and Vinuesa, 2017; Steinmann et al., 2018). Three of a total of four trimethoprim-sulfamethoxazole resistant isolates of our collection belong to genogroup #6, this antibiotic being a first-line therapeutic choice in S. maltophilia infections.

Phylogenomic analyses linked with comparative LPS profiling and O-serotyping have recently revealed correlations between genomic classification of a number of S. maltophilia isolates and electrophoretic banding patterns of their LPS in silver-stained polyacrylamide gels and immunoblots with an antibody raised against the O-antigen of the reference strain S. maltophilia K279a (Steinmann et al., 2018). The results obtained here confirm the usefulness of LPS profile typing in combination with immunoblotting as a supportive tool for identification of epidemiologically related S. maltophilia strains. The LPS-based approach was not only suited to supplement the genotypic characterization of our diverse strain panel, but
could also corroborate previous data on the high variability of O-polysaccharide structures among *S. maltophilia* strains.

**CONCLUSION**

Overall, this study shows that the QS system plays a pivotal role in pathogenicity and persistence in *S. maltophilia*. Furthermore, since there are significant correlations between the *rpf* variant and the phylogenetic groups of the isolates based on their MLST profiles (Supplementary Table S4), this genotyping method could be valid as an epidemiological tool. Genome sequencing studies also revealed genes that are exclusive to certain groups of isolates and could contribute to the observed phenotypes, showing that other factors besides DSF synthesis contribute to the virulence and resistance of *S. maltophilia*. More in-depth genomic analysis (e.g., single nucleotide polymorphism detection), together with phenotypic characterization of a larger collection of representative clinical isolates of *S. maltophilia*, will give a much deeper understanding of genetic diversity, phylogeny, population structure and epidemiology of this pathogen. For instance, this will allow the discovery of point mutations involved in antibiotic resistance, for example, in clonal strains isolated from the same patient. Altogether, this could have significant implications in understanding the biology of this species and, even more important, in the management of *S. maltophilia* infections.

**DATA AVAILABILITY STATEMENT**

The datasets generated for this study can be found in the https://pubmlst.org/smaltophilia/.

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**AUTHOR CONTRIBUTIONS**

DY, PH, SM-S, OC-S, UM, XC, FL, and IR performed the experiments presented in the work. OC-S and DY did the comparative genomics analyses. DY, PH, SM-S, OC-S, and UM participated in the writing of the manuscript. All authors contributed to the design and interpretation of the results, revised the manuscript, and approved it for publication.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020.01160/full#supplementary-material
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