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Evolutionary Dissection of the Dot/Icm System Based on Comparative Genomics of 58 Legionella Species

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Abstract

The Dot/Icm type IVB secretion system of Legionella pneumophila is essential for its pathogenesis by delivering >300 effector proteins into the host cell. However, their precise secretion mechanism and which components interact with the host cell is only partly understood. Here, we undertook evolutionary analyses of the Dot/Icm system of 58 Legionella species to identify those components that interact with the host and/or the substrates. We show that high recombination rates are acting on DotA, DotG, and IcmX, supporting exposure of these proteins to the host. Specific amino acids under positive selection on the periplasmic region of DotF, and the cytoplasmic domain of DotM, support a role of these regions in substrate binding. Diversifying selection acting on the signal peptide of DotC suggests its interaction with the host after cleavage. Positive selection acts on IcmR, IcmQ, and DotL revealing that these components are probably participating in effector recognition and/or translocation. Furthermore, our results predict the participation in host/effector interaction of DotV and IcmF. In contrast, DotB, DotO, most of the core subcomplex elements, and the chaperones IcmS-W show a high degree of conservation and not signs of recombination or positive selection suggesting that these proteins are under strong structural constraints and have an important role in maintaining the architecture/function of the system. Thus, our analyses of recombination and positive selection acting on the Dot/Icm secretion system predicted specific Dot/Icm components and regions implicated in host interaction and/or substrate recognition and translocation, which will guide further functional analyses.

Key words: Legionella, Dot/Icm system, positive-selection, negative-selection, diversifying-selection, evolution.

Introduction

Legionella are Gram-negative Gammaproteobacteria present in natural and man-made aquatic environments where they are replicating intracellularly within a wide range of amoeba species belonging to the genera Acanthamoeba, Hartmanella, Valkampfia, and Naegleria as well as inside ciliates such as Tetrahymena, Cyclidium, or Paramecium (Newton et al. 2010; Boamah et al. 2017). However, certain Legionella species are also well known human pathogens due to their capacity to replicate in mammalian cells like macrophages when accidentally reaching the human lung. In susceptible individuals, infection with Legionella can lead to an acute pneumonia known as legionellosis or Legionnaires’ disease (Newton et al. 2010).

The capacity of Legionella to infect eukaryotic cells relies on the Dot/Icm system, a type IV secretion machinery that is translocating proteins into the eukaryotic host cell. These effectors help this pathogen to subvert host functions to invade, replicate in, and escape from its hosts (Isberg et al. 2009; Hubber and Roy 2010). Based on sequence similarity, the Dot/Icm secretion machinery is classified as type IVB secretion system (T4BSSs) (Christie and Vogel 2000), in contrast to type IVA secretion systems (T4ASS) that are similar to the Agrobacterium tumefaciens VirB/VirD4 system. Recently, a classification based on the phylogenetic analysis of VirB4, the only ubiquitous protein in T4SSs, has been proposed (Guglielmini et al. 2013). Accordingly, T4SSs were classified in eight groups called MPF groups in reference to the
mating-pair formation proteins of the secretion systems. Based on this classification, T4ASSs belong to groups MPF (based on the T-DNA conjugation system of A. tumefaciens plasmid Ti) and MPF (based on plasmid F) whereas T4BSSs fall in the MPF group (based on the IncI plasmid R64). Both T4ASS and T4BSSs are bacterial multiprotein organelles specialized in the transfer of (nucleo)protein complexes across cell membranes (Christie et al. 2017).

The development of techniques to test experimentally whether a predicted Dot/Icm substrate is indeed secreted through this machinery (Zhu and Luo 2013) together with bioinformatics analyses has revealed that Legionella pneumophila is provisioned with a unprecedented number of over 330 Dot/Icm substrates (Burstein et al. 2009; Zhu et al. 2011; Lifshitz et al. 2013; Finsel and Hilbi 2015; Ensminger 2016; Escoll et al. 2016; Qiu and Luo 2017). Recent bioinformatics predictions of effectors in different Legionella species added many more proteins to this list (Burstein et al. 2016; Gomez-Valero et al. 2019). It is thought that this huge arsenal of effector proteins allows Legionella to adapt to its very large host spectrum of amoeba and ciliated protozoa. Interestingly, this effector repertoire is highly conserved among different L. pneumophila strains, but when comparing different Legionella species the scenario is very different, as most of the effectors lack orthologs in most or all Legionella species analyzed (Gomez-Valero et al. 2011; Gomez-Valero et al. 2014, 2019; Burstein et al. 2016). Indeed, of the over 330 substrates described in the species L. pneumophila, only 10 are present in all sequenced Legionella species, pointing to a very small core set of effectors (Burstein et al. 2016; Gomez-Valero et al. 2019). Nevertheless, bioinformatics predictions suggested that all species carry a large number of effectors. Therefore, this surprisingly small number of core substrates indicates that a very different effector set is present in different Legionella species (Gomez-Valero et al. 2019) probably because each Legionella species/strain faces a particular range of protozoan hosts in the environment. In contrast to the effector diversity, the Dot/Icm secretion system is highly conserved at interspecies level (Burstein et al. 2016; Gomez-Valero et al. 2019). This raises an intriguing question: which components of the Dot/Icm secretion system allow detection and translocation of such a broad spectrum of different proteins?

Structural and functional analyses of several components of the Dot/Icm system in the past few years have helped to better understand how the Dot/Icm machinery may work, but a detailed knowledge is still missing (Raychaudhury et al. 2009; Ghosal et al. 2017; Kwak et al. 2017; Chetrit et al. 2018; Meir et al. 2018). These studies have defined two major subcomplexes: the core transmembrane subcomplex and the coupling protein subcomplex (fig. 1). The first one is composed of the proteins DotC, DotD, DotF (IcmG), DotG (IcmE), and DotH (IcmK) and crosses the inner and outer membranes of

Legionella pneumophila (Vincent, Buscher, et al. 2006). In this complex, the DotH protein forms the outer membrane pore, localized properly thanks to the lipoproteins DotC and DotD, and receives energy from DotG (Sutherland et al. 2013). Additionally, DotF interacts with DotG and regulates energy transducing activity of DotG. The second complex recruits substrates and delivers them to the secretion channel. While only one protein, VirD4 mediates coupling between substrate recruitment and delivery to the secretion channel in type T4ASSs, a coupling complex is needed in T4BSSs. The coupling complex of the Dot/Icm system comprises the inner membrane AAA+ ATPase DotL (IcmO), which is a homolog of VirD4, two inner membrane/cytoplasmic components, DotM (IcmP) and DotN (Icm), a complex of two cytoplasmic chaperones, IcmS and IcmW (IcmSW) and the protein LvgA (Kwak et al. 2017). LvgA was originally described as a Legionella virulence factor (Edelstein et al. 2003), but later it has been shown that this protein is also part of the Dot/Icm secretion machinery (Vincent and Vogel 2006). Finally, to be functionally complete, the Dot/Icm secretion apparatus contains 16 other proteins in addition to the above-mentioned subcomplexes (table 1).

Given its function and localization, the Dot/Icm system is submitted to strong and competing evolutionary pressures: it needs to (i) avoid host defences, (ii) constantly adapt to new hosts/niches, and (iii) preserve interactions with other components of the Dot/Icm apparatus. Accordingly, some Dot/Icm elements have to remain unaltered to maintain the stability of the system and are thus expected to be highly conserved even among different Legionella species. Given that, most of the amino acid changes occurring within these proteins will be deleterious and thus removed by natural selection, the evolutionary pressure acting on these components/regions is negative/purifying selection. In contrast, other components/regions, in particular the ones directly exposed to the host system and/or the substrates, have to adapt constantly to different or new hosts/niches and are therefore evolving fast and show high variability. These components are usually under a positive selective pressure as the amino acid changes have a high probability to provide a selective advantage and thus to be fixed. Therefore, the study of the evolutionary pressures acting on the Dot/Icm system allows predicting which elements/regions are key for effector/host interaction and which remain unaltered, probably for maintaining the integrity of the system.

Such evolutionary forces can be better dissected among different species that diverged sufficiently long ago and live in different environments. Therefore, interspecies analysis provides a unique opportunity to identify elements under adaptive evolution. The genus Legionella is a privileged model for such analyses as the quasi entire genus has been sequenced. Although intracellular replication has been shown experimentally only for 35 of the 58 Legionella species included here
(summarized in Gomez-Valero et al. 2019) it can be assumed that all species replicate in eukaryotic hosts. This idea is also substantiated by a recent phylogenetic analysis of T4BSSs in Gram-negative bacteria that proposes that the acquisition of the T4BSS on the chromosome might be related to the alteration of the life style as intracellular bacterium. Furthermore, they suggest that the genes found only in the Dot/Icm systems of Legionella and related bacteria may encode components that are important for life as intracellular pathogens (Nagai and Kubori 2011). Thus, we used here the sequence of 80 Legionella strains belonging to 58 species (Gomez-Valero et al. 2019) and applied an evolutionary approach to identify the different functions of the Dot/Icm components.

Materials and Methods

The sequence of the 27 genes that constitute the Dot/Icm secretion system were extracted from the genomes of 80 Legionella strains belonging to 58 species, previously sequenced in our lab (Gomez-Valero et al. 2019) (supplementary table S1, Supplementary Material online). BLASTp of each Dot/Icm protein encoded by the L. pneumophila Paris genome against the orthologous proteins in each of the other Legionella species/strains was used to calculate the average amino acid identity of the Dot/Icm proteins (table 1) (Gomez-Valero et al. 2019). Each gene was translated and the corresponding proteins were aligned using the program PRANK v.170427 (Loytynoja 2014). PRANK uses a phylogenetic tree of the strain/species included in the alignment. Thus, a phylogenetic tree based on the core genome previously published (Gomez-Valero et al. 2019) of these strains was included in the study (supplementary fig. S1, Supplementary Material online). The resulting amino acid alignments were used as a guide for the alignment of the corresponding nucleotide sequences using the software DAMBE v.6.4.40 (Xia 2017). The obtained alignments were cleaned with Gblocks v.0.91b (Castresana 2000) with the less astringent conditions (allowing: smaller final blocks, gap positions within the final
blocks, and less strict flanking positions). Alignment properties were calculated using the tool infoalign from the EMBOSS package v. 6.6.0.0 (Rice et al. 2000) and taking L. pneumophila Paris as reference genome. Additionally, Shannon entropy calculation was done for some selected protein alignments using the Protein variability server (García-Boronat et al. 2008), to visualize variability along the protein sequence.

Intragenic recombination detection was carried out using the software 3seq v1.7 (Lam et al. 2018) according to recommendations by Martin et al. (2011). The program was also run on concatenated genes belonging to the same cluster. To confirm identified recombination events, we analyzed the congruence of each Dot/icm protein alignment with the phylogeny obtained from all alignments and the phylogeny derived from the core genome alignment. The congruence was evaluated applying the Shimodaira–Hasegawa (SH) test (Shimodaira and Hasegawa 1999) implemented in TreePuzzle (Schmidt et al. 2002). Tree-Puzzle was also used to evaluate the phylogenetic information contained in each Dot/icm protein alignment, using likelihood mapping (Strimmer and von Haeseler 1997) (supplementary table S2, Supplementary Material online).

For selection analyses, recombinant sequences from each alignment were removed, as these may interfere with the positive selection analysis. To ensure that differences among the compared sequences represented real fixation events along independent lineages, only one strain from each species was used in this analysis as artifacts may appear when comparing closely related strains (Kryazhimskiy and Plotkin 2008). Finally, for positive selection detection, we applied several methods implemented in the Hyphy package v. 2.3

### Table 1

| Protein Name | Protein Label for L. pneumophila strain Paris | No. of Sequences | Amino Acid Identity (%) | Gene Length | Alignment Length | Average Identical Nucleotides | % Change | Recombind |
|--------------|---------------------------------------------|------------------|-------------------------|------------|-----------------|-------------------------------|----------|-----------|
| IcmT         | Lpp0507                                     | 80               | 78–100 (87%)            | 261        | 249             | 196.4                         | 20.5     | No        |
| IcmS         | Lpp0508                                     | 80               | 74–100 (91%)            | 345        | 333             | 263.6                         | 20.4     | No        |
| IcmR         | Lpp0509                                     | 16               | 48–99 (94%)*            | 363        | 357             | 337.0                         | 6.6      | No        |
| IcmQ         | Lpp0510                                     | 80               | 49–99 (70%)             | 576        | 504             | 364.7                         | 27.1     | No        |
| IcmP / DotM  | Lpp0511                                     | 80               | 64–99 (79%)             | 1,131      | 1,098           | 834.8                         | 23.5     | Yes (1)  |
| IcmO / DotL  | Lpp0512                                     | 80               | 77–100 (89%)            | 2,352      | 2,325           | 1,854.6                       | 19.9     | Yes (10) |
| IcmN / DotK  | Lpp0513                                     | 80               | 50–99 (69%)             | 570        | 486             | 341.1                         | 29.0     | No        |
| IcmM / DotJ  | Lpp0514                                     | 80               | 33–99 (63%)             | 285        | 291             | 188.6                         | 35.0     | No        |
| IcmL / DotI  | Lpp0515                                     | 80               | 69–100 (90%)            | 639        | 633             | 504.7                         | 19.7     | No        |
| IcmK / DotH  | Lpp0516                                     | 80               | 65–99 (79%)             | 1,083      | 867             | 672.6                         | 21.9     | No        |
| IcmE / DotG  | Lpp0517                                     | 80               | 49–100 (71%)            | 3,147      | 2,706           | 1,904.1                       | 27.8     | Yes (43) |
| IcmG / DotF  | Lpp0518                                     | 80               | 37–99 (57%)             | 810        | 504             | 362.1                         | 27.4     | No        |
| IcmC / DotE  | Lpp0519                                     | 80               | 57–100 (76%)            | 585        | 525             | 384.7                         | 25.5     | No        |
| IcmD / DotP  | Lpp0520                                     | 80               | 57–99 (78%)             | 399        | 315             | 242.0                         | 22.5     | No        |
| IcmJ / DotN  | Lpp0521                                     | 80               | 72–100 (87%)            | 627        | 603             | 482.8                         | 19.5     | No        |
| IcmB / DotO  | Lpp0522                                     | 80               | 80–100 (88%)            | 3,030      | 3,006           | 2,366.4                       | 20.6     | Yes (1)  |
| IcmF         | Lpp0524                                     | 80               | 34–100 (70%)            | 2,922      | 2,811           | 1,985.7                       | 28.2     | No        |
| IcmH / DotU  | Lpp0525                                     | 80               | 36–100 (71%)            | 786        | 747             | 537.1                         | 27.5     | No        |
| DotV         | Lpp0537                                     | 79               | 44–100 (69%)            | 543        | 462             | 332.1                         | 28.1     | Yes (6)  |
| LvgA         | Lpp0590                                     | 80               | 42–100 (69%)            | 627        | 528             | 380.5                         | 27.3     | Yes (1)  |
| DotD         | Lpp2728                                     | 80               | 69–100 (84%)            | 492        | 465             | 364.2                         | 21.0     | No        |
| DotC         | Lpp2729                                     | 80               | 69–100 (81%)            | 912        | 810             | 634.4                         | 20.9     | No        |
| DotB         | Lpp2730                                     | 80               | 86–100 (94%)            | 1,134      | 1,101           | 878.1                         | 19.7     | No        |
| DotA         | Lpp2740                                     | 80               | 47–99 (60%)             | 3,108      | 2,025           | 1,419.0                       | 28.5     | Yes (20) |
| IcmV         | Lpp2741                                     | 80               | 49–99 (67%)             | 456        | 429             | 300.6                         | 29.4     | No        |
| IcmW         | Lpp2742                                     | 80               | 71–100 (89%)            | 456        | 453             | 361.0                         | 19.8     | Yes (1)  |
| IcmX         | Lpp2743                                     | 78               | 36–89 (51%)             | 1,419      | 711             | 464.6                         | 34.1     | Yes (4)  |

*Notes:* Amino acid (aa) identity represents the minimum, maximum, and average values (average between parentheses) of aa identity between each Dot/icm protein from L. pneumophila Paris against the corresponding orthologous in other species calculated by BLASTp. The parameters: alignment length, average of identical nucleotides, and the % of changes are calculated from the multiple alignment of each protein after cleaning with Gblocks and taking as a reference the genome of L. pneumophila Paris for calculating identity and % of change.

*a* L. pneumophila strain Paris.

*b* With respect to L. pneumophila Paris sequence.

*c* After cleaning with Gblocks.

*d* Number of recombinant sequences.

eIdentity value based on the comparison of 16 strains (15 belonging to the same species L. pneumophila and only one to a different species L. norrlandica).
(Pond et al. 2005): MEME (Murrell et al. 2012), FUBAR (Murrell et al. 2013), FEL (Pond et al. 2005), and aBSREL (Smith et al. 2015). FEL and FUBAR allow detecting sites under pervasive diversifying selection. Whereas both methods assume that selection pressure for each site is constant along the entire phylogeny, they differ in the way to calculate non-synonymous (dN) and synonymous (dS) substitutions. FEL uses a maximum-likelihood approach whereas FUBAR uses a Bayesian approach. Moreover, FUBAR seems to have more power than FEL, in particular when positive selection is present but relatively weak (Murrell et al. 2013). Additionally, we applied MEME to test the hypothesis that individual sites have been subject to episodic diversifying selection. Finally, aBSREL was applied to identify lineages, which have experienced episodic purifying selection independently of the sites affected by this selective pressure. We choose the P value according to the recommendations in the HYPHY package; 0.1 for FEL and MEME. For aBSREL, we choose a P values <0.01. For FUBAR, we do not have P values but posterior probabilities.

To ensure that the positive selection signal, we detected was not due to misalignments, in particular, in the case of the most variable Dot/Icm components, we used an alternative alignment program, MUSCLE v.3.8, and rerun FUBAR/aBSREL to confirm the results. Using this control, the only case that was not corroborated was positive selection acting on several codons at the beginning of DotK when using PRANK alignments. Detailed visual inspection on this region revealed that in fact it could not be aligned with high accuracy. In consequence, we did not take this result into account. Signal IP v.4.1 (Petersen et al. 2011) was used to detect secretion signals. The MOTIF search tool of the Japanese GenomeNet service (Kanehisa 1997) was applied for detecting pentapetide repeats using the Pfam library (Finn et al. 2016). The program IBS v.1.0.3 (Liu et al. 2015) was used for schematic representations of protein domains and ITOL v.4.3 (Letunic and Bork 2016) for representing phyllogeny together with partial alignments.

Results and Discussion

Recombination Is Important in the Evolution of the Dot/Icm System

The Legionella Dot/Icm system is highly conserved at interspecies level as previously reported (Burstein et al. 2016; Gomez-Valero et al. 2019) with an average amino acid identity of the Dot/Icm proteins ranging from 51% (icmX) to 94% (DotB) (Table 1). Due to this high homology at protein level, it was possible to align the corresponding genes with high accuracy based on protein alignments. Alignments were curated to avoid spurious orthologous codon detection while keeping most of the gene sequence (Table 1).

To analyze the role of recombination and natural selection on shaping the interspecies genetic diversity of the Dot/Icm components, we followed a two-step strategy. First, recombination analyses were carried out using the nonparametric 3seq to detect mosaic structures in the sequences. This analysis identified intragenic recombination in 9 of the 27 dot/icm genes with particular high recombination rates in dotG, dotA, and icmX as further detailed below (Table 1). As a second approach, we undertook recombination detection on concatenated genes, which was confirming our results (supplementary table S3, Supplementary Material online). Furthermore, we undertook a phylogenetic incongruence test (SH), which compares the likelihood of each protein alignment versus the phylogenetic trees derived from them and the core phylogeny to confirm these results (supplementary table S4 and fig. S2, Supplementary Material online). Combined these methods disclosed that recombination events are important in the evolution of the Dot/Icm system.

We then aimed to discover the Dot/Icm components, and more specifically the codons in these proteins, that may have been subjected to positive selection. After removing the above-detected recombinant sequences to avoid false positive results, amino acids that have been subjected to pervasive positive selection were searched for with the methods FEL and FUBAR. Furthermore, the sites that have evolved under positive selection even when this selection has acted only on a proportion of the branches in the phylogenetic tree were analyzed by MEME. Finally, we applied the aBSREL method (Smith 1992) to detect branches in the phylogeny of Legionella that could have been under positive selection for specific Dot/Icm components regardless whether codons affected by this selective pressure had been detected. Among the 27 Dot/Icm proteins, specific sites under positive selection were identified in IcmQ, DotM, DotF, IcmF, DotV, and DotC. Moreover, four genes that evolved through diversifying selection on different nodes of the phylogeny were identified (dotL, dotG, dotA, and icmX). No significant signs of positive selection for the remaining 17 genes could be detected.

Analysis of Proteins of the Core Transmembrane Subcomplex

High Intragenic Recombination Rates Support Host Exposure of DotG and DotA

DotG (IcmE/Lpp0517) is an integral membrane protein that was proposed to form a central channel spanning inner and outer membranes (Fig. 1) (Kubori et al. 2014). Recent in situ cryo-electron tomography confirmed that DotG couples the outer membrane core complex with the cytoplasmic complex by forming the cylinder domain that constitutes the central channel portion (Chetrit et al. 2018). When analyzing the DotG protein sequence at the genus level, we found that recombination plays a major role in the evolution of this protein as recombination events were identified in 43 of the 80 analyzed DotG sequences (Table 1). Despite this result, the DotG protein phylogeny is compatible with the core
phylogeny (supplementary fig. S2 and table S5, Supplementary Material online) when the SH test is applied (P value 0.82). This result may be due to the fact that the DotG protein is a large protein (1,048 aa in L. pneumophila) and recombination events are affecting only the N-terminal and middle part of the protein but not the C-terminal region (supplementary fig. S3, Supplementary Material online). Consequently, the phylogenetic signal of the alignment is strong enough to recover a tree with a similar topology as the core phylogeny.

The C-terminus of DotG of L. pneumophila has been reported to be similar to the TrbI domain of VirB10 family proteins of T4ASSs (Nagai and Kubori 2011). Our sequence analysis shows that this TrbI domain is conserved in the 75 analyzed strains/species for which the complete DotG sequence was available. In contrast to the well-conserved C-terminal region, the middle region is highly variable and contains a variable number of 1–13 pentapeptide repeats (supplementary table S4, Supplementary Material online). For L. pneumophila, Coxella burnetti, and Rickettsiella, it has been reported that DotG proteins are significantly larger than other homologs due to these repeats (Segal et al. 1998; Nagai and Kubori 2011). Here, we show that the DotG proteins of 58 Legionella species vary between 1,000 (cluster of L. pneumophila) and >1,500 amino acids (cluster of Legionella sainthelenensis–Legionella longbeachae and Legionella santicrucis) (supplementary fig. S4, Supplementary Material online). Interestingly, even between strains from the same species a considerable size variation can be present, as seen for the two Legionella oakridginesis strains analyzed here, that have 1,011 and 1,161 amino acids, respectively. The Helicobacter pylori HP0527 (CagY) protein, a homolog of DotG, contains also a large number of repeat regions following the well conserved C-terminal part, that can be deleted or extended by intragenic recombination leading to variations between strains. This mechanism has been suggested to help the pathogen to avoid or modulate the host immune response as CagY decorates the bacterial surface (Rohde et al. 2003; Barrozo 2013). Although, in Legionella, there is to date no experimental evidence that the variable sequence regions of DotG are surface displayed, our results suggest that some segments are surface exposed, and that DotG function could be similar to its homolog CagY. In addition, although specific sites under positive selection were not detected, several branches on the evolutionary tree of DotG have been subjected to diversifying selection (aBSREL) (table 2). Such a faster evolution supports again an interaction of DotG with the host system and/or with the secreted substrates.

A second Dot/icm protein showing a high recombination rate is DotA (Lpp2740), an integral cytoplasmic membrane protein (Roy and Isberg 1997), that is essential for the functioning of the T4SS (fig. 1) (Berger et al. 1994). The presence of recombination and frequent nonsynonymous mutations in the dotA gene has been reported for different L. pneumophila strains (Ko et al. 2003; Costa et al. 2010). Our present analysis of DotA shows also a high degree of interspecies variability (only 60% of average amino acid identity) and a high recombination rate with 20 different Legionella species being affected. A SH test confirmed this result as the phylogeny derived from DotA is not compatible with the phylogeny derived from the core genome (P value 0.007). The test shows incongruent evolutionary histories between dotA and the core genome, a result in agreement with recombination events affecting this protein (supplementary fig. S2, Supplementary Material online). Like for DotG, no specific site under diversifying selection was detected, but many branches in the tree are under positive selection (aBSREL) (table 2 and supplementary fig. S4, Supplementary Material online). It is highly probably that this is due to the fact that sequences affected by recombination had to be removed prior to the analyses, which reduced the power of detection of sites under positive selection for all methods used except for one (aBSREL). Thus, recombination and positive selection have played an important role in the evolution of DotA at the genus level.

DotA is an inner membrane protein but it is also secreted after cleavage of the 19 amino acids long leader peptide (Nagai and Roy 2001). Our results that show fast evolution of DotA further support the idea that DotA directly interacts with the host forcing DotA in the different Legionella species to constantly adapt to different hosts.

Positive Selection Analyses Suggest a Role of the Periplasmic Region of DotF in Substrate Recognition

DotF (IcmG/Lpp0518) is an inner membrane protein that is part of the core transmembrane complex of the Dot/icm apparatus (fig. 1). Our analysis shows that, like DotA, this protein has high variability at interspecies level as the percentage of change is 27.4% (table 1). Moreover, the analysis of selective forces acting on DotF points to at least two residues under diversifying selection (table 2). These two residues reside in the periplasmic domain (fig. 2) adjacent to the transmembrane domain of DotF. This finding suggests a role of this periplasmic region in effector interaction as predicted by Sutherland et al. (2013). Indeed, it has been demonstrated previously that DotF interacts with several Dot/icm effectors (Luo and Isberg 2004) and that this interaction takes place through the transmembrane and/or periplasmic domain of DotF (Sutherland et al. 2013; Kubori et al. 2014). Additionally, the recent electron cryotomography results suggested that the walls of the Dot/icm channel crossing the bacterial membrane have openings. The opening between the beta and gamma rings is localized just below DotD and next to DotF, suggesting that DotF plays a role in translocating effectors initially secreted to the periplasm, to the secretion chamber (Ghosal et al. 2017). Interestingly, the C-terminal
domain of DotF that contacts this opening is similar to the
C-terminal domain of PilP/GspC (Ghosal et al. 2017),
which in type II and III secretion systems recruits effectors
from the periplasm and delivers them to the translocation
channel. DotF was therefore suggested to play a role in
effector interaction or alternatively a role in stabilizing the
apparatus or triggering conformational changes (Ghosal
et al. 2017). Our current results showing diversifying se-
lection acting on DotF support a role of DotF in effector
interaction. Moreover, the fact that amino acids of DotF
for which we detect positive selection are exposed in the
periplasm supports the function of the periplasmic do-
main of DotF in effector binding. Indeed, from the 168
codons of DotF, 159 were under negative selection pres-
sure and only two are under positive selection (table 2).
Thus, the diversifying evolution detected in the periplas-
mic domain of DotF is probably reflecting adaptation to
changes in effector repertories in the different Legionella

### Table 2

| Gene Name | Gene Label | No. of Sequences | Methods Used to Infer Selection |
|-----------|------------|------------------|-------------------------------|
| icmT      | lpp0507    | 58               | FEL, MEME, FUBAR, ABSREL       |
| icmS      | lpp0508    | 58               |                               |
| icmR      | lpp0509    | 2                |                               |
| icmQ      | lpp0510    | 58               |                               |
| icmP, IdotM | lpp0511    | 56               |                               |
| icmO, IdotL | lpp0512    | 56               |                               |
| icmN, IdotK | lpp0513    | 58               |                               |
| icmM, IdotT | lpp0514    | 57               |                               |
| icmL, IdotI | lpp0515    | 58               |                               |
| icmK, IdotH | lpp0516    | 58               |                               |
| icmE, IdotG | lpp0517    | 35               |                               |
| icmG, IdotF | lpp0518    | 58               |                               |
| icmC, IdotE | lpp0519    | 58               |                               |
| icmD, IdotP | lpp0520    | 58               |                               |
| icmJ, IdotN | lpp0521    | 58               |                               |
| icmB, IdotO | lpp0522    | 58               |                               |
| icmF      | lpp0524    | 58               |                               |
| icmH, IdotU | lpp0525    | 58               |                               |
| dotV      | lpp0537    | 52               |                               |
| lgvA      | lpp0590    | 57               |                               |
| dotD      | lpp2728    | 58               |                               |
| dotC      | lpp2729    | 58               |                               |
| dotB      | lpp2730    | 58               |                               |
| dotA      | lpp2740    | 48               |                               |
| icmV      | lpp2741    | 58               |                               |
| icmW      | lpp2742    | 57               |                               |
| icmX      | lpp2743    | 53               |                               |

**NOTE.**—Number in parenthesis indicates either P value or posterior probability (FUBAR method) associated to the codon/branch inferred to be subjected to positive selection. Gray areas highlight those genes where positive selection was found.
species and defines the region of DotF involved in effector binding.

High Conservation and Positive Selection Characterize the Outer Part of the Core Complex

Whereas DotG and DotF contact the inner membrane proteins, three other proteins: DotH (Lpp0516), DotC (Lpp2729), and DotD (Lpp2728) constitute the outer part of the core transmembrane complex (fig. 1) (Nakano et al. 2010). Moreover, it was reported that DotH is associated with a fibrous structure that covers the entire bacterial surface under certain conditions enhancing bacteria internalization (Watarai et al. 2001). If exposed on the bacterial surface, we would expect that this protein interacts with the host system and is subjected to an evolutionary arm race. However, except an intrinsically disordered N-terminal region of 50 amino acids (15% of the protein) DotH shows a high degree of interspecies conservation (table 1). Moreover, when removing this region, no signatures of recombination or of positive selection are detectable (table 2). Thus, although we cannot discard that the disordered N-terminal region contains amino acids...
under positive selection our results suggest that DotH is not directly exposed to the host system as previously described as no sign of diversifying selection acting on this protein was detected.

DotC and DotD, like DotH show a high degree of conservation (table 1) and contain signal peptides responsible for their secretion through the outer membrane. Our analyses identified three codons in the DotC N-terminal region within the signal peptide that are under positive selection (table 2 and fig. 2). The DotC signal peptide is well conserved as it was detected in 63 of the 80 sequenced DotC proteins (supplementary table S6, Supplementary Material online). In addition to targeting preprotein secretion, it has been shown that the cleaved signal peptides can play additional roles as hormones, neurotransmitters, or self-antigens (Hegde and Bernstein 2006). It is thus tempting to suggest that the released signal peptide of Legionella DotC is detected by, for example, the host immune system explaining why positive selection is acting on this sequence. However, alternative explanations such as relaxed selection due for example to alternative start codons cannot be discarded. Additionally, we also detected positive selection acting on amino acid 250 of DotC (fig. 2), however, only MEME is supporting this and very few species are affected by this amino acid change, thus this result needs to be taken with care. In contrast, DotD sequence conservation is high in all species and DotD shows no signs of recombination or of positive selection acting on it. The crystal structure of this protein showed that it has striking structural similarity to the N-terminal subdomain of secretins and N0 domain proteins (Nakano et al. 2010). They found that the DotD/N0/T3S domain is present in outer membrane components of many even distantly related secretion systems indicating that negative selection is acting on it which is in line with our results.

In summary, although highly conserved at the sequence level, the outer membrane side of the core complex shows positive selection pressure acting on the signal peptide sequence of DotC suggesting an additional role of this protein in the host cell after cleavage.

**DotB and DotO are Subjected to Strong Negative Selection**

DotB (Lpp2730) is a protein that forms stable homohexameric rings and hydrolyses ATP (Sexton, Miller, et al. 2004). The corresponding mutant is defective for growth in macrophages, but Sexton et al. (2005) obtained some dotB alleles with partial activity. Two of these were unable to export a subset of T4SS substrates indicating a possible role of DotB in substrate selection. Our comparative analysis shows that DotB is the DotIcm protein with the highest degree of conservation (93% average amino acid identity among different species; table 1) and even DotB from C. burnetti can complement a L. pneumophila dotB mutant (Zamboni et al. 2003). Furthermore, no recombination or sites under positive selection were identified in the DotB proteins. A similar result was found for the ATPase DotO (Lpp0522) where conservation at the sequence level is high and neither branches nor sites under positive selection were detected. Recently, it has been reported that DotB is a dynamic entity as it can be free in the cytosol or associated with the DotIcm system through the DotO ATPase (Chetrit et al. 2018) where it constitutes the disc at the base of the cytoplasmic complex of the DotIcm system. Our results demonstrate that these proteins are mostly under negative selection and suggest that the basal structure composed by DotB-DotO is highly structurally constrained to allow the passage of all effectors across the DotB-DotO energy complex.

**Analysis of Proteins of the Coupling Subcomplex**

**Recombination and Diversifying Selection Shape DotL and DotM Evolution**

The DotIcm system-coupling subcomplex (T4CP) contains integral inner-membrane proteins that play a dual role of recruiting substrates and escorting them to the secretion conduit (Gomis-Ruth et al. 2004). DotL and DotM are part of this subcomplex (fig. 1). DotL (IcmO/Lpp0512) is an inner membrane protein related to Escherichia coli VirD4 and TrwB (Buscher et al. 2005), both structural prototypes of coupling subcomplexes in T4SS. However, compared with TrwB, DotL contains an additional 200-residue segment at the C-terminus of unknown function that is found also in Coxiella, Yersinia, and Pseudomonas species (Kwak et al. 2017).

Our analysis shows a high degree of interspecies conservation of DotL. This result is in line with the knowledge that DotL is involved in multiple interactions with DotN, IcmS, IcmW, and DotM and therefore under high structural constraints to maintain the architecture of the coupling subcomplex. Moreover, DotL plays an important role in intracellular replication since the corresponding mutants are defective in replication in a variety of host cells (Sutherland et al. 2012). Despite its high conservation several branches are affected by diversifying selection (aBSREL) but we did not detect any site under positive selection. Furthermore, we detected intragenic recombination events affecting mostly L. pneumophila strains and the species Legionella dumoffi and Legionella worsellensis. The regions involved in recombination were always located in the P-loop Ntpase domain (supplementary fig. S5A, Supplementary Material online), but were not affecting DotL regions that interact with IcmS-W, DotN, and DotM (Vincent et al. 2012). Taken together, most of the DotL sequence is highly conserved whereas the variability accumulates mainly in the DotM-interacting domain of the transmembrane region and in particular in the segment of the C-terminus (supplementary figs. S5B and S6, Supplementary Material online).

Although we did not detect specific sites under selection in DotL, the high variability localized in the C-terminal region is a sign of a fast evolution rate of this part of the protein, which is
Another component of this coupling complex is DotM (ICmP/Lpp0511), a protein that possesses a cytoplasmic domain that has just been crystallized (Meir et al. 2018) and that is thought to interact with DotL through their transmembrane domains (fig. 1) (Vincent et al. 2012). Indeed, we observed that the transmembrane region of DotM is highly variable like the DotM-interacting domain of DotL as mentioned above and that positive selection is acting on codon 22 localized at the end of the first transmembrane helix domain of DotM. These results further suggest coevolution of both proteins. The analyses of the crystal structure of the DotM cytoplasmic domain revealed that it contains large patches of basic residues suggesting that it might form a recruiting platform for domain revealed that it contains large patches of basic residues which is in agreement with our results that identify a weak positive selection signal on codon 362 of the cytoplasmic domain (table 2 and supplementary fig. S7, Supplementary Material online). Moreover, this codon under positive selection has undergone amino acid replacement alternating between polar and neutrally charged residues suggesting that this impacts DotM-effector interactions.

Taken together, we show that despite a general high degree of sequence conservation of both DotM and DotL, specific regions of these two proteins are under fast evolution pointing to their potential role in the interaction with substrates. In contrast to DotM and DotL, the proteins DotN and IcmT that were also suggested to be involved in effector recruitment (Meir et al. 2018) show a high conservation and no recombination nor sites and/or branches under positive selection.

Evolution of Cytoplasmic Proteins
Extremely Fast Evolution is Acting on the IcmRQ Complex
IcmQ and IcmR are essential for growth of L. pneumophila in macrophages (Coers et al. 2000). These proteins interact in vivo in L. pneumophila (Dumenil and Isberg 2001) through the middle region of IcmR and the N-terminal region of IcmQ (Raychaudhury et al. 2009). When not bound to IcmR, IcmQ can insert into the lipid membrane forming pores through its N-terminal part (Dumenil et al. 2004). Our analysis of IcmQ (Lpp0510) shows a moderate conservation (70% average amino acidic identity), no recombination but at least one amino acid under positive selection, localized at the beginning of the protein (table 2 and fig. 2). Curiously, this amino acid is located in the part of the protein that has been defined as the interacting with IcmR (amino acids 1–57 in L. pneumophila Paris). The alignment shows that most of the hydrophobic residues previously defined to be involved in this interaction (Raychaudhury et al. 2009) are also conserved in all Legionella species analyzed here (supplementary fig. S5, Supplementary Material online). The structure of full-length IcmQ in complex with IcmR revealed that the C-terminal domain of IcmQ contains a NAD+ binding domain (Farelli et al. 2013). An alignment of IcmR from different bacteria was used to define the essential residues of this IcmR NAD+ domain (Farelli et al. 2013). Here, our interspecies alignment revealed which of these residues have a higher conservation among species and are therefore potentially essential for IcmR function (supplementary fig. S8, Supplementary Material online). The presence of this module has allowed to suggest that IcmRQ binds to membranes, where it may interact with, or perhaps modify, a protein in the T4SS when NAD(+) is bound (Farelli et al. 2013).
IcmR (Lpp0509) is the Dot/Icm protein with the highest rate of evolution of all Dot/Icm proteins. An icmR gene similar to \textit{L. pneumophila} icmR was identified only in the species \textit{Legionella norrlandica}, the phylogenetically closest species to \textit{L. pneumophila} (Gomez-Valero et al. 2019). Thus, IcmR from \textit{L. pneumophila} strain Paris can only be aligned with homologous proteins belonging either to strains from the same species or to the closely related \textit{L. norrlandica}, which explains the high amino acid identity values obtained (table 1) despite the high evolutionary rate of IcmR. Indeed, in all other \textit{Legionella} species analyzed, one or two nonhomologous genes replace this gene in the same position where icmR is present in \textit{L. pneumophila} (Gomez-Valero et al. 2019).

Feldman et al. (2005) had shown that the genes that replace icmR in \textit{Legionella} hackeliae and \textit{Legionella micdadei} are functional homologs of \textit{L. pneumophila} icmR (designated FIR proteins). Despite the lack of sequence homology, two conserved structural regions were predicted in the FIR proteins containing nonidentical, hydrophobic side chains that may contribute to the binding between IcmR and IcmQ (Raychaudhury et al. 2009). We have shown that these two regions in FIR proteins are also conserved in 58 different \textit{Legionella} species (Gomez-Valero et al. 2019). However, the absence of homology at the sequence level for IcmR in the different \textit{Legionella} sp. constitutes a limiting factor for the analysis of diversifying selection. Homology among five or more strains was present only in two subgroups, one containing the species \textit{Legionella gratiana}, \textit{L. santhelensi}, \textit{L. longbeachae}, \textit{Legionella cincinnatiensis}, and \textit{L. santicrucis} and the other one containing \textit{L. pneumophila} and \textit{L. norrlandica} strains. Therefore, we used these two groups to search for positive selection within IcmR. Among \textit{L. pneumophila} and \textit{L. norrlandica} strains (16 sequences), codons 39 and 90 were identified as being under positive selection. In contrast, within the \textit{L. longbeachae} cluster, codon 10 was under selection (data not shown). Thus, positive selection seems to act on specific amino acids of IcmR. The reason why this gene is so extremely divergent is not known. Originally, it was suggested that the FIR–IcmQ complex is secreted upon contact with a protozoan host cell what would explain the positive selection acting on it (Dumenil and Isberg 2001). Later, the crystal structures of the N-terminal domain of IcmQ with the interacting region of IcmR suggested that IcmQ is associated with the inner bacterial membrane (Raychaudhury et al. 2009) and consequently not exposed to the host system. Therefore, the diversifying selection acting on these proteins and more specifically the high evolutionary rate of IcmR is probably linked to the large variety of Dot/Icm effectors secreted in the different \textit{Legionella} species by this system.

Our analysis combined with the crystal structures suggest that IcmR may have a central role in substrate interaction and thus needs constantly to adapt to the changing effector repertoire in the different species.

### Selection Analysis of Dot/Icm Components of Yet Unknown Function

IcmX (Lpp2743) is a 50-kDa periplasmic protein that is essential for \textit{L. pneumophila} pathogenesis (Sadosky et al. 1993; Edelstein et al. 1999; Matthews and Roy 2000) and required for pore formation in the membrane of the eukaryotic cell (Matthews and Roy 2000). Our analysis reveals that, although present in all analyzed \textit{Legionella} species, IcmX is one of the least conserved proteins of this system (table 1), especially in the N-terminal region. Consequently, many regions of the gene could not be included in our analysis due to uncertainty in their corresponding alignment. However, we still detected intragenic recombination in icmX in four \textit{Legionella} species (table 2) and one branch of the tree under positive selection (aBSREL). This diversifying selection in some lineages may reflect a role of IcmX as signal transmitter to the host cell. It has been reported that a truncated IcmX product is secreted into culture supernatants by \textit{L. pneumophila} (Matthews and Roy 2000), although its translocation across eukaryotic cell membranes has not been detected. Additionally, it has been shown that IcmX is a surface exposed protein (Khemiri et al. 2008). Together, these results strongly suggest that IcmX is a protein exposed to the host, and therefore, in different \textit{Legionella} species, it interacts with different protozoan hosts explaining its fast evolution.

The Dot/Icm system comprises also many membrane-associated proteins, such as DotK/IcmN (Lpp0513), IcmF (Lpp0524), DotU (Lpp0525), DotE (Lpp0519), DotV (Lpp0537), DotP (Lpp0520), Dotl (Lpp0515), DotU (Lpp0514), and IcmV (Lpp2741). Among those, IcmF and DotU proteins prevent DotH degradation, stabilize the \textit{L. pneumophila} T4SS (Sexton, Pinkner, et al. 2004) and recruit the DotCH complexes to the poles of the cell (Jeong et al. 2017). Our analysis revealed no recombination or diversifying selection acting on DotU, but IcmF contains one residue under positive selection (MEME) and several branches are affected by diversifying selection (supplementary fig. S9, Supplementary Material online). The amino acid under positive selection is located in a conserved segment among IcmF homologs and outside the C-terminal region predicted to contact DotCH (Ghosal et al. 2017). This is consistent with the fact that the IcmF segment predicted to interact with DotCH is likely subjected to structural constraints that prevent amino acid changes. Among DotK, DotE, DotP, Dotl, DotU, IcmV, and DotV, recombination and positive selection were detected only for DotV and positive selection affects the last codon of one of the predicted transmembrane helices (Nagai and Kubori 2011) (supplementary fig. S11, Supplementary Material online).

These results demonstrate that IcmX, IcmF, and DotV are subjected to diversifying selection suggesting that they play a role in host/effecter interaction and are thus interesting targets for future functional studies.
Concluding Remarks

It is well known that the constant arms race between pathogens and hosts selects for the maintenance of polymorphisms thereby allowing adaptations and counter-adaptations to occur. The Legionella Dot/Icm type IV secretion system or at least a part of it has to contact the host cell for the delivery of effectors. It is thus a target of the pathogen recognition systems and a hot spot of selection. Our study analyzing these evolutionary forces acting on it revealed high rates of recombination and/or positive selection for proteins DotA, DotG, and IcmX suggesting, in line with previous studies, that these proteins are directly interacting with the host system. Moreover, our analysis highlights DotC whose signal peptide is subjected to diversifying selection which may indicate that after cleavage it is released and plays a role in the host cell. In contrast, our results did not support the suggestion that DotH and DotO have a role in host interaction.

In contrast to the high conservation of the Dot/Icm secretion system at interspecific level, the effector repertoire is very variable (Burstein et al. 2016; Gomez-Valero et al. 2019) suggesting that proteins involved in effector binding have to evolve through diversifying selection to adapt to different effector sets in the different species. Indeed, we detected diversifying selection acting on DotL and DotM that had been suggested previously to be involved in effector recruitment. Moreover, we detected amino acids and regions under positive selection in DotF, the cytoplasmic domain of DotM and the C-terminal region of DotL pointing to protein segments probably involved in interacting with Dot/Icm substrates. In the case of IcmR, several amino acids were under positive selection, despite a limited analysis possibility due to lack of conservation of this protein at interspecific level. The positive selection acting on the protein together with its extremely high rate of evolution clearly points to a key role in substrate recruitment. Furthermore, IcmF, DotV, and DotK contain amino acids and/or branches under positive selection suggesting they are involved in effector/host interactions. In contrast, DotO and DotB that constitute the base of the cytoplasmic Dot/Icm complex are under strong negative selection as neither signs of recombination nor of positive selection and a high degree of interspecific conservation were detected suggesting that these proteins are essential for the maintenance of the architecture and or function of the T4SS.

In conclusion, our evolutionary studies of the Dot/Icm components allowed identifying those proteins and amino acids of this secretion system that may be functionally important for host/effector binding. Indeed, the detection of diversifying selection acting on pilius proteins of the type III secretion system of Pseudomonas syringae (Gutman et al. 2006) or on the type IV secretion system of Bartonella (Nystedt et al. 2008) suggested their role in host–pathogen interactions. For the Dot/Icm secretion system, we are far from a complete understanding of its structural and functional mechanism, but the availability of interspecies genome data allows new ways to analyze their components and thereby to predict their role. Functional analyses of the proteins predicted here through the analysis of evolutionary forces acting on them will be exciting to gain further insight into this important secretion system.

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

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Author Contributions

L.G.V., I.C., and A.C.O. did the data analyses and interpretation. The article was written by L.G.V. and C.B. with input from coauthors. The project was conceived, planned, and supervised by L.G.V. and C.B.

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