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Comparative analyses of transport proteins encoded within the genomes of Leptospira species

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A B S T R A C T

Select species of the bacterial genus Leptospira are causative agents of leptospirosis, an emerging global zoonosis affecting nearly one million people worldwide annually. We examined two Leptospira pathogens, Leptospira interrogans serovar Lai str. 56601 and Leptospira borgpetersenii serovar Hardjo-bovis str. L550, as well as the free-living leptospiral saprophyte, Leptospira biflexa serovar Patoc str. ‘Patoc 1 (Ames)’ The transport proteins of these leptospires were identified and compared using bioinformatics to gain an appreciation for which proteins may be related to pathogenesis and saprophytism. L. biflexa possesses a disproportionately high number of secondary carriers for metabolite uptake and environmental adaptability as well as an increased number of inorganic cation transporters providing ionic homeostasis and effective osmoregulation in a rapidly changing environment. L. interrogans and L. borgpetersenii possess far fewer transporters, but those that they all have are remarkably similar, with near-equivalent representation in most transporter families. These two Leptospira pathogens also possess intact sphingomyelinases, holins, and virulence-related outer membrane porins. These virulence-related factors, in conjunction with decreased transporter substrate versatility, indicate that pathogenicity arose in Leptospira correlating to progressively narrowing ecological niches and the emergence of a limited set of proteins responsible for host invasion. The variability of host tropism and mortality rates by infectious leptospires suggests that small differences in individual sets of proteins play important physiological and pathological roles.

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1. Introduction

Leptospirosis is an emerging zoonotic disease that affects about 900,000 people annually worldwide. It is caused by members of the bacterial genus Leptospira within the Spirochaeta phylum [1]. The disease poses a tremendous public health risk in tropical environments, especially as it is transmitted through contaminated water, infected tissues, and the urine of mammalian hosts [1]. Once infected, patients can potentially experience a variety of symptoms ranging from fever, myalgia, and fatigue to refractory shock, jaundice, renal failure, and pulmonary hemorrhage [1]. At-risk populations for this disease are primarily, but not exclusively, associated with tropical climates in developed and underdeveloped countries. Cases throughout the United States have also been reported, e.g. in Hawaii, Maryland, Louisiana, etc. [2–4]. Factors that increase risk include conditions of slum living, recreational water activities, and flooding [1]. Nonetheless, leptospirosis is found globally, and various animals such as rats, bats, and marsupials as well as domesticated animals such as dogs and cows can serve as reservoirs for its transmissions to humans [5–7]. Human to human transmission is rare, but it is believed that globalization and ecotourism contribute significantly to the emergence of this zoonosis [8].

The causative agents of leptospirosis, Leptospira spp., are spiral-shaped, thin, aerobic, gram-negative bacteria of highly divergent sprochetes, whose primary carbon sources are long-chain fatty acids [9]. Despite its threat to global public health, the Leptospira genus is not entirely filled with pathogenic species [10]. It is divided up into saprophytes (e.g. Leptospira biflexa, Leptospira wolbachii, Leptospira meyeri), pathogens (e.g. Leptospira interrogans, Leptospira borgpetersenii, Leptospira kmentyi), and intermediate pathogens (e.g. Leptospira licerasiae, Leptospira wolffii, Leptospira broomii) [10]. Among these characteristically distinct leptospires, practice among scientists has traditionally been to distinguish leptospires from each other by means of serotyping and antigenic similarity instead

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of genetic similarity [10]. Consequently, serotyping has resulted in the identification of over 230 different serovars among Leptospira with serovars crossing species lines [10]. The limited ability of sequencing technology at the advent of the identification and characterization of members of Leptospira was a driving force for this convention, but rapid sequencing technology such as qPCR has begun to enable scientists and clinicians to rapidly and effectively identify infecting leptospires and their phylogenetic relationships [11].

*L. interrogans* and *L. borgpetersenii* are two of many species that have been identified as pathogenic mediators of leptospirosis [9]. Their most common reservoir animals are rats and cows, respectively, although these two spirochetes are not exclusive to these two animals as noted above [12,13]. Working closely with these and other aforementioned animals, or being in proximity to them or their urine, gives increased likelihood to contracting the pathogen [14]. *L. biflexa*, on the other hand, is a free-living saprophyte isolated from stream water, the antigenic properties of which have been used as a basis for antigenic testing of pathogenic leptospires [15].

In the present study, the publicly available genomes of the following representative organisms were examined: *L. interrogans* serovar Lai strain 56601, *Leptospira borgpetersenii* serovar Hardjo-bovis strain L550, and *Leptospira biflexa* serovar Patoc strain ‘Patoc 1 (Ames)’. Table 1 presents the nomenclature for the organisms used in this study as well as additional information about them. *L. interrogans* and *L. borgpetersenii* allow comparison and identification of hallmarks of pathogenic leptospires, and *L. biflexa* enables comparison for identification of transport proteins and mechanisms unlikely to be related to pathogenicity.

In general, the mechanisms of pathogenesis in leptospirosis are relatively poorly understood. However, there are several suggested mechanisms [9]. The coiled shape of *Leptospira* is relevant to its corkscrew-like motility through viscous media, which provides an efficient mechanism of dissemination after entry into various organs such as the lungs, liver, kidneys, eyes, and brain [9]. Genes associated with motility and chemotaxis are known to play a role in virulence [16]. Among proposed factors that may facilitate virulence during migration through host tissues are hemolytic sphingomyelinases and phospholipases [16]. Additional components of leptospiral virulence promote adhesion and invasion of host cells, although intracellular pathogenicity has not been demonstrated [16].

Chromically infected animals (particularly rats, bats, and marsupials) are usually asymptomatic but show high levels of leptospiral excretion through the urine, supporting the hypothesis that renal colonization is important for *Leptospira* in reservoir selection and pathogenesis [9,17]. Kidney histological studies further support this hypothesis as kidneys show interstitial nephritis during infection, but no such damage in chronic carriers [18]. Kidney nephritis, along with damage to connective tissues, evident from hemorrhagic manifestations in lungs, supports virulence mechanisms involving invasion and damage to connective tissues [19]. Leptospiral lipopolysaccharide (LPS), known to be less toxic than the typical LPSs of other gram-negative bacteria, more strongly activates Toll-like receptor 2 (TLR2) than TLR4, conventionally achieved by gram-negative LPS in macrophages [20].

Ultimately, hemolytic sphingomyelinase and phospholipase activities, together with the identified motility and chemotaxis factors of *Leptospira*, damage host tissue and activate the inflammatory response of the host immune system to potentially cause significant damage, eventually resulting in death of the host [1]. Identification of transporters relevant to pathogenesis might reveal the presence of pore-forming toxins, transporters facilitating basic nutrient uptake, and the protein secretion systems necessary to release proteinaceous virulence factors. Furthermore, *Leptospira* pathogenic species are known to maintain poor viability in acidic urine relative to alkaline urine which suggests a preference of sodium cations (Na+) in some transport systems that utilize the proton motive force (pmf) or the sodium motive force (smf) [9].

Pathogenic species of *Leptospira* must encode the proteins that mediate virulence. Both *L. interrogans* and *L. borgpetersenii* might be expected to show similar pathogenesis-related transporters, but this postulate had not been examined. *L. biflexa* would be expected to have few, or incomplete sets of these proteins [21]. Studies on *L. biflexa*, *L. interrogans*, and *L. borgpetersenii* have suggested that *L. biflexa* is most closely related to the common *Leptospira* ancestor, and that pathogenicity was an acquired feature [21]. Consequently, the saprophytic and free-living nature of *L. biflexa* suggests that its genome enables it to live with high versatility in a range of environments [21,22]. The suggested flexibility of *L. interrogans* to live within a host and also in the external environment combined with the greater dependence of *L. borgpetersenii* for survival in the host in spite of its relatively small genome, due to insertion sequence (IS)-mediated reduction [22], suggests that increased pathogenicity and host tropism favor decreased versatility and reduced genetic diversity. The encoded transport proteins should reflect these characteristics.

### 2. Materials and methods

The spirochete genomes analyzed were the most complete and up to date versions for each organism at the time these studies were initiated. The FASTA formatted protein coding sequences of *Leptospira interrogans* serovar Lai strain 56601, *Leptospira biflexa* serovar Patoc strain ‘Patoc 1 (Ames)’, and *Leptospira borgpetersenii* serovar Hardjo-bovis strain L550 were used [21–23]. Each protein sequence from the respective proteomes was queried and blasted against the Transporter Classification Database (TCDB; www.tcdbr.org [24]) using the program GBLast [25]. GBLast retrieves the TC top hit sequence, TC number, protein size in number of amino acyl

<table>
<thead>
<tr>
<th>Species name</th>
<th>Leptospira interrogans serovar Lai strain 56601</th>
<th>Leptospira borgpetersenii serovar Hardjo-bovis strain L550</th>
<th>Leptospira biflexa serovar Patoc strain ‘Patoc 1 (Ames)’</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chromosome I RefSeq</td>
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<td>NC_008508.1</td>
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<td>9.36</td>
</tr>
<tr>
<td>Pathogenic?</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>
residues, the number of predicted TMSs using the HMMTop 2.0 Program, the E-value for the query and hit proteins, regions of sequence similarity, and regions of TMS overlap. The low complexity filter was not used as it is normally of value only for larger datasets including proteins with multiple repeat elements. The Web-based Hydrophathy, Amphipathicity and Topology (WHAT) program [26,27] was used with a window size of 19 residues and an angle of 100° (as is appropriate for alpha helices) to display the hydropathy plot for individual proteins in conjunction with TOPCONS for consensus prediction (TOPCONS, www.topcons.net) in order to resolve the differences in the number of TMSs between the proteins retrieved and their TCDB homologs. The plot generated by WHAT allows the user to judge if a program such as HMMTOP has missed a TMS or has predicted a TMS inappropriately. A cut-off E-value of 0.001 was used with the G Blast program to eliminate most false positives and proteins with unreliable degrees of sequence divergence.

Proteins with no predicted TMSs were eliminated so that only integral membrane proteins, primarily multispanning membrane proteins, were retrieved. Proteins with only an N-terminal signal sequence are numerous because these proteins include almost all secreted proteins that are exported via the general secretory (Sec) pathway or twin arginine translocase (TAT) [28,29]. The topological prediction programs too often miss these TMSs, recording the proteins to have zero TMSs. Consequently, the number of zero or one TMS proteins retrieved were not reliable and were therefore not always recorded. Furthermore, TMSs detected by G Blast are those only of the α-helix-type transporters and not the β-sheet-type porins. Transporters known to have β-sheet, porins found primarily in TC subclasses 1.B and partly in 1.C, were further analyzed for β-strands using PRED-TMBB with all three decoding methods [30].

Transport proteins thus obtained from query Leptospira sequences were tabulated, and unusual characteristics were identified based in part on topologies that differed from corresponding family members in TCDB as well as E-values obtained with G Blast. Unusual properties can result from events such as genetic deletion or fusion, sometimes resulting in the gain or loss of extra domains or the generation of multifunctional proteins. Such results can be reflective of the protein sequence, but they can be artifacts due to sequencing errors or incorrect initiation codon assignment. In the latter cases, but not the former, the protein sequences were either corrected when possible or eliminated from our study.

Candidate proteins were examined in greater detail to estimate their probable substrate specificities on the basis of their predicted structures and numbers and degrees of sequence similarity with entries of known function in TCDB. Homology was examined using the Protocol 1 and Protocol 2 programs as well as GSAT [25]. Transport proteins were also classified into families and subfamilies of homologous transporters according to the classification system presented in TCDB [24,31,32]. Regions of sequence similarity were examined to ensure that homology was in the transmembrane region(s) and not merely in hydrophilic domains [33]. The substrate specificities of particular homologs identified in the sequenced genomes were initially predicted based on homology to functionally characterized proteins. Assignment to a family or subfamily within the TC system often allows prediction of substrate type with confidence [34–36].

3. Results

Three Leptospira genomes were analyzed for the occurrence of transport proteins using the Transporter Classification Database (TCDB) [24] and the G Blast program [25]. The results are summarized according to TC subclass in Table S1. Examining the total number of transport proteins present in these three genomes, we see that L. borgpetersenii has the fewest at 260. The most found in any genome is 337 for L. biflexa, 77 more than in L. borgpetersenii. However, the two pathogenic species (L. interrogans and L. borgpetersenii) combined contained 110 unique transport proteins lacking in L. biflexa, the saprophyte.

3.1. Transport protein subclasses

TC subclass 1.A in TCDB includes all α-type channels except for holins, which are found in the 1.E subclass. 17, 21, and 26 of these α-type channel proteins were identified in L. interrogans, L. borgpetersenii, and L. biflexa, respectively. L. biflexa possesses the greatest number of unique families in subclass 1.A, many of which are cationic channels including two mercuric ion channels, suggesting greater versatility in the saprophyte than in the pathogens. Surprisingly, L. interrogans possesses eight proteins belonging to the 1.A.30 Outer Membrane Transporter Energizer (Mot-Exb) superfamily compared to the fifteen found in both L. biflexa and L. borgpetersenii.

TC subclass 1.B includes outer membrane β-type porins. 38 were identified in L. interrogans, 32 in L. borgpetersenii, and 39 in L. biflexa. The distribution of these porins does not suggest a major contribution to pathogenicity. Since these proteins localize to the outer membrane via β-strands instead of α-helices, those containing zero or one predicted α-helical TMSs were included in our study.

TC subclass 1.C includes pore-forming toxins. L. interrogans encodes ten putative toxins showing sequence similarity to established toxins belonging to four families, whereas L. borgpetersenii contains eight and L. biflexa contains seven. The 1.C.67 family of SphH Hemolysins is notably absent in L. biflexa but present in L. interrogans and L. borgpetersenii with four and two members, respectively. Although hemolysins have not been unequivocally shown to be essential for leptospiral pathogenesis, their presence in pathogens is likely to be of significance [16]. Hemolysins have been shown to strongly induce proinflammatory cytokines [20]. The three other families represented in subclass 1.C contain similar representation in the three leptospires examined. It should be noted that toxins with zero or one predicted transmembrane α-helices were included in this study as many secreted toxins can exist in both soluble and membrane integrated forms, and many are known to be pore-forming β-type toxins (see TCDB).

TC subclass 1.E consists of Holins. Both L. interrogans and L. borgpetersenii encode a protein that hits the Mycobacterial 4 TMS Phage Holin (MP4 Holin) (TC#1.E.40.3.6). Holins have a variety of proposed functions in prokaryotes and may play a role in cell lysis and biofilm formation [37]. The presence of these holins in L. interrogans and L. borgpetersenii and their aforementioned functions may promote pathogenicity in these leptospires.

The largest number of transporters for all three species is found in TC subclass 2.A, secondary carriers. L. interrogans encodes 75, L. borgpetersenii 70, and L. biflexa 110. Given the substantially greater number of secondary carriers found in L. biflexa, the relative presence (or lack thereof) of a variety of transporters may help to distinguish between the two Leptospira pathogens and free living bacteria. L. biflexa appears to have much greater metabolic flexibility than its two pathogenic cousins.

TC subclass 2.C includes energizers for motility and outer-membrane transport. All three species encode a single TonB protein that is necessary to energize the stable construction of the outer membrane. The MotAB protein pair energizes motility while the ExbBD protein pair forms an H’ or Na’ channel that energizes transport across the outer membrane with the periplasmic accumulation of solutes, dependent on TonB and outer membrane receptors (OMRs; TC#1.B.14). Further analyses of these energizers will be reported below.
TC subclass 3.A are pyrophosphate hydrolysis-driven primary active transporters, usually multi-component systems. With 56, 49, and 67 integral membrane transport proteins of this subclass found in L. interrogans, L. borgpetersenii, and L. biflexa, respectively, these proteins make up a significant proportion of the total transport proteins found in these organisms. The variety and wealth of transporters found in this subclass clearly suggests that they play an important role in spirochetes. The 3D TC subclass of ion-pumping electron carriers are represented in L. interrogans, L. borgpetersenii, and L. biflexa with 25, 30, and 32 proteins, respectively.

TC class 4 includes group translators that are believed to modify their substrates in processes coupled to transport. The TC 4.B subclass includes members of the nicotinamide ribonucleoside (NR) group translocating uptake permease (PnuC) family. Only L. biflexa encodes such a protein.

TC subclass 4.C includes fatty-acyl-coenzyme A ligases that activate fatty acids for lipid biosynthesis and may function in transport via group translocation. All three species each contains one member of this family. GBLAST revealed more fatty-acyl-coenzyme A ligases in all three leptospires, but the absence of transmembrane segments in addition to an unproven transport function in leptospires warranted their exclusion from our analyses.

All three Leptospira species have proteins belonging to TC subclass 4.D, probable group translocating glycosyl transferases. L. interrogans and L. biflexa encode three, whereas L. borgpetersenii encodes four. Proteins in this family have demonstrated exopolysaccharide synthesis activities thought to be coupled to polysaccharide secretion [38]. As exopolysaccharides can contribute to biofilm formation, all three leptospires likely benefit from the presence of these proteins, for both free-living and host colonization purposes.

Subclass 5.A includes electron-carriers that transfer electron pairs from one side of the membrane to the other, thereby influencing cellular energetics. L. interrogans was found to have two, whereas L. borgpetersenii and L. biflexa have three. Among these are disulfide bond oxidoreductases and prokaryotic molybdopterin-containing oxidoreductases. These proteins might play a role in establishing the proton motive force, but they probably do not contribute to pathogenicity.

Subclass 5.B in TCDB consists of one electron transmembrane carriers. None of the leptospires was found to contain integral membrane carriers in this subclass. However, all three contain multiple copies of cytochrome c peroxidases (TC#5.B.3.1.1) for extracellular reduction of Fe2O3 (unpublished results).

Subclass 8.A represents auxiliary proteins with one in L. interrogans and L. borgpetersenii, and three in L. biflexa. All three encode a stomatin-like protein which may help with localization and insertion of proteins into the outer membrane. L. biflexa is the only one to encode a membrane fusion protein (TC#8.A.3.2.1), which probably functions with an ABC exporter.


TC Subclass 9.B includes a variety of proteins that are putatively classified as transporters. Further study of a given 9.B protein might either confirm its involvement in transport, or warrant its removal from the TC classification system if a transport function is disproven. L. interrogans and L. borgpetersenii have 34 while L. biflexa has 38 of these proteins.

3.2. Transporter superfamilies and families

Fig. 1 details protein families found in some, but not all three leptospiral species. 21 families are shown to be unique to L. biflexa, dwarfing the three families each unique to L. interrogans and L. borgpetersenii. This is reflective of the large differences in the numbers of total transporters (Table S1) and the disproportionately high number of 2.A carriers (Fig. 1). Analyses of these families may reveal key features of free-living saprophages.

Unique families of transporters found in L. biflexa, presented in Fig. 1, reveals ten families in the 2.A subclass. Families of unique transporters with notable substrates are TC#1.A.72, 2.A.51, 2.A.56, 2.A.59, 2.A.102, and 4.B.1 that transport mercury, chromate, dicarboxylates, arsenic, sulfites, and nicotinamide mononucleotide and related compounds, respectively. All of these substrates are transported only by L. biflexa, suggesting increased versatility over L. borgpetersenii and L. interrogans.

The only families of transporters unique to L. interrogans (Fig. 1) include TC#1.A.43 and TC#2.A.121, members of which transport fluoride and sulfate, respectively. L. borgpetersenii does not appear to have unique transporter families. Families belonging to both pathogens are TC#1.A.23, 1.C.67, 1.E.40, 2.A.114, 2.A.115 and 9.B.125, encoding small mechanosensitive ion channels, hemolysins, holins, peptide transporters, multidrug exporters, and unknown substrates, respectively. The families unique to L. interrogans may confer increased environmental versatility over L. borgpetersenii, but the shared transport families may play roles in pathogenesis.

3.3. Interesting facets of channel proteins

A limited number of channel protein families is represented in the three Leptospira examined. Most channel proteins are involved in ionic and water homeostasis, but some also serve functions in stress responses. These will be described below.

Only a single member of the Voltage-gated Ion Channel Superfamily (TC#1.A.1) was identified, and this protein was found only in L. biflexa. It proved to be a 6 TMS cyclic nucleotide-dependent channel, almost certainly a potassium channel like those characterized in cyanobacteria and other spirochetes [39].

A single member of the MIP Family (TC#1.A.8) of aquaporins and glycerol facilitators was found in each of the three leptospires. These three proteins are probably aquaporins capable of transporting three-carbon compounds such as glycerol and dihydroxyacetone [40]. The high scores, all matching the same TCDB entry, suggest that these three proteins are orthologous.

Ammonium channels are prevalent in leptospires but in variable numbers; thus, L. interrogans has two dissimilar paralogs, and L. biflexa has three, but L. borgpetersenii has only one. Interestingly, it appears that one of these proteins in each organism hits the homolog from Azospirillum brasilense (TC#1.A.11.1.4) with excellent comparable scores. These three proteins are undoubtedly orthologs. It is worth noting that the A. brasilense protein is subject to multiple mechanisms of regulation which may be applicable to the spirochete proteins as well [41].

The three Leptospira species examined possess either one or two homologs of Epithelial Chloride Channels (TC#1.A.13), characterized only in animals. Although bacterial homologs have been identified, in none of them is the function known. Our results reveal that these spirochete proteins exhibit the same topology as the mammalian proteins, suggesting a similar function. We suggest that these proteins will prove to exhibit chloride channel activities comparable to those found in eukaryotes.

The three leptospires display either zero or one small mechanosensitive ion channel (MscS; TC#1.A.23), and interestingly, L. biflexa is the one that lacks such a protein. All three organisms lack an MscL channel (TC#9.A.22). Proteins of both of these families are known to function in osmotic adaptation [42].

All three spirochetes possess a member of the MgtE family
The three Leptospira species examined possess multiple paralogs of the H⁺ or Na⁺-translocating MotAB/ExbBD/TolQR channel-forming constituents (TC#1.A.30). While MotAB proteins function to energize motility [43,44], ExbBD channels energize transport across the outer membrane [45], and TolQR channels are believed to energize assembly of the outer membrane, promoting stability of this structure [46]. All three spirochetes possess two MotAB energizers which presumably function in motility, possibly one utilizing the proton motive force and the other utilizing the sodium motive force [47]. On the other hand, the occurrence of ExbBD/TolQR energizers are variable in these three species with two in L. interrogans, four in L. borgpetersenii, and five in L. biflexa. These results suggest that L. interrogans, like Escherichia coli, possesses the equivalent of one ExbBD complex and one TolQR complex [48–50]. However, the other two leptospires have an increased number of these H⁺ or Na⁺ channel proteins. The functions of these proteins will be interesting targets of future investigations.

Remaining families of channel proteins are present only in select Leptospira species. The CorA Metal Ion Transporter Family (TC#1.A.35) is only represented in L. interrogans and L. biflexa. L. interrogans possesses a member of the Camphor Resistance Family (TC#1.A.43). These proteins have recently been shown to be fluoride export channels which protect the bacterium against the toxic effects of fluoride [51,52]. For both the Homotrimeric Cation Channel Family (TC#1.A.62) [53] and the Mer Superfamily (TC#1.A.72) [54,55] only L. biflexa has constituent channels. While the former proteins have not been characterized in bacteria, the latter function in the uptake of mercuric ions for the purpose of reduction to metallic mercury by a cytoplasmic mercuric reductase, a detoxification reaction [56].

### 3.4. Interesting facets of β-type porins

β-type porins represent a significant portion of the channel proteins found in the Leptospira examined. The leptospiral outer membrane is of particular interest as it contains cell surface antigens that can be used for vaccine production, and they can also serve as potential drug targets [57]. Members of sixteen different families of outer membrane porins were identified in at least one of the three Leptospira species examined, and interestingly, fourteen of these families are represented in all three species. Just two of the families (POP; 1.B.5 and SAP; 1.B.16) were found only in L. biflexa, not in the two pathogenic species. While the POP Family is concerned with anion transport, the SAP Family mediates urea and short-chain amide transport.

Some of the families represented in all three organisms have only a single protein per organism, and these may be orthologs of each other as all three proteins hit the same TC entry (see for example 1.B.4, 1.B.6, 1.B.9, and 1.B.13). However, striking differences occur in some of the other families, for example, the Outer Membrane Receptor (OMR) Family (TC#1.B.14), where each organism exhibits different sets of these pore-forming receptors. This fact can be explained by the different specificities of these receptors as illustrated in Table S1. Similar observations were made for the Outer Membrane Factor (OMF) Family (TC#1.B.17), where again, the
different specificities of these porins provide an explanation. It seems likely that the complement of OMRs and OMFs reflect the specific environments in which these organisms are found.

The remaining families in this subclass consist of macromolecular transporters for protein secretion (TC#1.B.22; TC#1.B.48), outer membrane protein insertion (TC#1.B.33), lipid export (TC#1.B.42), outer membrane lipid insertion (TC#1.B.46), and polysaccharide export for protection and biofilm formation (TC#1.B.55). All leptospires possess members of these families, which represent core components of the Leptospira outer membrane proteome.

3.5. Interesting facets of secondary carriers (TC subclass 2.A)

In most organisms, the Major Facilitator Superfamily (MFS; TC#1.A.1) is the largest superfamily of secondary carriers. However, in the pathogenic leptospires, the MFS is poorly represented. Both species have only six MFS members and zero members of the related GPH Family (TC#2.A.2). By contrast, L. biflexa has fifteen MFS porters and two GPH porters. Thus, its MFS representation is almost three times that of the pathogens. Of particular note is the presence of multiple multidrug efflux pumps of MFS subfamily 21, nitrate/nitrite transporters, and several members of unknown function. Also, while the pathogens possess only a single member of the ABC amino acid transporter family, L. biflexa has three such members, each derived from a separate subfamily.

Divalent cation transporters can mediate either uptake or efflux of these essential but potentially toxic substances. While the CDF Family (TC#2.A.4) catalyzes heavy metal export and has equal numbers of its members in all three leptospires, the ZIP Family (TC#2.A.5) and the NRAMP Family (TC#2.A.55) catalyze heavy metal uptake and are found only in L. biflexa. Interestingly, the Chromate Resistance (CHR) Family (TC#2.A.51) and the Arsenical Resistance-3 (ACR3) Family (TC#2.A.59) are also restricted to L. biflexa.

The RND Superfamily is by far the largest superfamily of secondary carriers present in these spirochetes with sixteen members in L. interrogans, fourteen in L. borgpetersenii, and twenty in L. biflexa. These proteins are divided about equally between heavy metal efflux pumps and multidrug resistance pumps. Only the SecDF and SecF proteins, present in single copy in all three organisms, fall outside of these two groups. These two proteins function together as a single RND pump to facilitate proton-driven protein secretion via the General Secretory Pathway (Sec; TC#3.A.5) [58]. Finally, the two pathogens, but not L. biflexa, possess a single member of the poorly characterized putative Hydrophobe/Ampiphile Efflux-3 (HAE3) subfamily (TC#2.A.6.7) within the RND superfamily.

The Drug/Metabolite Transporter (DMT) Superfamily is the third largest superfamily in these spirochetes. L. interrogans has four such members, L. borgpetersenii has five, and L. biflexa has nine. Most of the top hits in TCDB have not been functionally characterized, so specific substrates cannot be assigned. However, all known members of this superfamily function in the transport of small metabolites and drugs.

Interestingly, all three leptospires have a single homolog of the Sweet family of putative sugar transporters (TC#2.A.123). The homologs identified have a 3 TMS subunit structure. Seven 7 TMS Sweet family members have been shown to transport sugars such as glucose and fructose [59]. Presently, two 3D structures of a 3 TMS Sweet glucose transporter from L. biflexa have been solved [60]. Based on these structures, transport mediated by this protein appears to be that of a secondary carrier [60]. Table 51 reveals the presence of secondary carriers belonging to many other families, and almost all of these are well-represented in all three leptospires. These families will not be further discussed here.

3.6. Interesting facets of primary active transporters

TC subclass 3.A contains the largest superfamily of transporters found in all three Leptospira species, the ABC Superfamily (TC#3.A.1). While L. interrogans and L. borgpetersenii possess 28 and 27 of these proteins, respectively, L. biflexa possesses 41. The ABC Superfamily is represented in all domains of life and is known to transport a wide variety of substrates for both uptake and export. Of note, L. biflexa is the only leptospire to possess ABC transporters for putrescine/spermidine (polyamines), phosphate, thiamine, zinc (Zn$^{2+}$), iron siderophores, and fatty acyl-CoA. However, all three organisms possess good representation of oligopeptide transporters, suggesting that these substances are important for the nutrition of these organisms. All three organisms have ABC uptake systems for sulfate and lipids.

ABC efflux systems are present in numbers that are similar to those of the uptake systems in all three spirochetes. The primary transporters for these efflux systems possess 1) lipids and lipoproteins, 2) proteins and peptides, 3) exopolysaccharides, and 4) multiple drugs. Most of these transporters, except for those specific for lipids, are found in similar numbers in the three spirochetes examined. Only a few ABC export systems are specific to L. biflexa. One of these is a putative organoanion (fatty acid?) exporter (TC#3.A.1.203.8), and the others undoubtedly exhibit specificity for specific proteins (TC#3.A.1.109, 3.A.1.110, 3.A.1.111). Interestingly, a single member of the Membrane Fusion Protein (MFP) Family (TC#3.A.1.1) was found only in L. biflexa, correlating with the presence of these ABC protein exporters.

All three leptospires possess orthologous sets of the integral membrane components of the ATP synthases in the F-ATPase Superfamily (TC#3.A.2) for subunits a, b, and c. The reversibility of the enzyme for both the establishment of the proton motive force and ATP synthesis is a key characteristic of this system. Additionally, all three Leptospira have an H$^{+}$- or Na$^{+}$-translocating pyrophosphatase (TC#3.A.10). While the TC hit for L. biflexa (TC#3.A.10.1.1) is different from those for L. interrogans and L. borgpetersenii (TC#3.A.10.1.6), sequence comparisons of these entries showed that these proteins are probably orthologous.

Of note is the variance of P-type ATPases (TC#3.A.3) in Leptospira species. L. interrogans possesses six of these transporters with substrate specificities for magnesium (Mg$^{2+}$), copper (Cu$^{2+}$), and potassium (K$^{+}$). L. borgpetersenii, however, possesses only two, one specific for copper (Cu$^{2+}$) and the other for calcium (Ca$^{2+}$). While the Mg$^{2+}$ and K$^{+}$ systems catalyze uptake, the Cu$^{2+}$ and Ca$^{2+}$ systems probably catalyze efflux. L. biflexa has six including a putative Na$^{+}$/K$^{+}$ ATPase, two copper (Cu$^{2+}$) transporters, a calcium (Ca$^{2+}$) transporter, and a heavy metal (Cd$^{2+}$, Zn$^{2+}$, Cd$^{2+}$) transporter. The diversity of these transporters presumably reflects the types of stress that these organisms encounter. Thus, most prokaryotic P-type ATPases function in stress relief [61,62].

All three leptospires have proteins with sequence similarity to the three integral membrane components of the General Secretory Pathway (Sec) Family (TC#3.A.5), which transports most secreted proteins across the inner cytoplasmic membrane. The presence of SecDF (TC#2.A.6.4) as well as the associated YajC protein (TC#9.B.18) in all three species reveals the genus--wide presence of the integral membrane constituents of the general secretory pathway. In addition, we found constituents of the outer membrane protein secreting Main Terminal Branch (MTB) Family (TC#3.A.15). These proteins proved to be distantly related to the MTB constituents previously tabulated in TCDB, and consequently we have entered all constituents of this system from L. interrogans into TCDB under TC# 3.A.15.4.1. MTB family members are believed to export hundreds of proteins across the outer membranes of gram negative bacteria, initially secreted across the cytoplasmic membrane by the Sec system [63].
Not surprisingly, all motile leptospires possess the flagellar (Type III) secretion complex (TC#3.A.6). The constituents recorded in Table S1 include six integral membrane constituents of these systems in all three leptospires whose near-identical E-values for all components suggest orthology for the three complete systems in this genus. The striking similarity between Type III secretion systems and flagellar export systems gives rise to the possibility that these systems export virulence factors in addition to flagellar subunits as has been demonstrated for these systems in other bacteria [64,65].

As expected, all leptospires have the Septal DNA Translocase (TC#3.A.12), involved in DNA transfer across the completed septa of newly dividing cells. However, while these organisms lack a type IV protein secretion system involved in conjugation, they do possess components of Bacterial Competence-related DNA Transformation (DNA-T) systems (TC#3.A.11). Interestingly, while *L. interrogans* and *L. biflexa* appear to have all constituents of these systems, only two were found in *L. borgpetersenii*. Possibly this last organism has lost some of the constituents of these systems and therefore has lost competence. Surprisingly, nothing seems to have been published on competence for DNA uptake in *Leptospira* species.

None of the spirochetes examined appears to have a Na+–transporting carboxylic acid decarboxylase of the NaT-DC Family (TC#3.B.1). These organisms do have decarboxylases, but they lack the integral membrane protein which is required for Na+–extrusion. We therefore conclude that this mechanism for generating a sodium motive force is lacking in these organisms, in agreement with the conclusion that these ion pumps are largely restricted to anaerobes [66].

Constituents of most, but not all, of the primary proton pumping electron transfer complexes present in mitochondria and many aerobic bacteria were found in the leptospires. These include the proton-translocating NADH dehydrogenase (TC#3.D.1), proton-translocating transhydrogenase (TC#3.D.2), and proton-translocating cytochrome oxidase (TC#3.D.4), but not the proton-translocating quinol:cytochrome c reductase (TC#3.D.3). Additionally, leptospires possess prokaryotic succinate dehydrogenase (TC#3.D.10). These results are consistent with the conclusion that leptospires use electron transfer as a primary mechanism for generating a proton motive force, subsequently used for ATP synthesis. As expected, these aerobic bacteria possess members of the disulfide bond oxidoreductase D (DsbD) and Molybdopterin-containing Oxidoreductase (PMO) Families (TC#5.A.1 and 5.A.3, respectively), but surprisingly not the single electron transferring DsbB complex. In this regard, it is important to note that members of the DsbD family can serve any of a variety of biological functions (see TCDB).

### 3.7. Possible group translocators (TC class 4)

None of the leptospires possess a phosphoenolpyruvate-dependent sugar transporting phosphotransferase system (PTS) although proteins of such systems have been found in other spirochetes [67]. However, *L. biflexa* appears to have a nicotinamide ribonucleoside uptake permease (TC#4.B.8), thought to function by a group translocation mechanism [68]. Each spirochete also has a membrane–associated acyl-CoA ligase (TC#4.C) that could function in transport [69]. Finally, each leptospire possesses three or four polysaccharide synthase/exporters (TC#4.D), all of which give low scores to the proteins in TCDB. These putative enzyme/porters may catalyze vectorial glycosyl polymerization [38,70].

### 3.8. Poorly characterized transporters (TC class 9)

TC subclass 9.A represents known transport systems that function by an unknown mechanism of action. Three such systems are found in all three spirochetes, and no other members of this subclass were identified. The first of these families is the FeoB family of ferrous iron uptake transporters (TC#9.A.8). The second family is a multicomponent protein secretion system characterized only in Bacteroidetes (TC#9.A.25). Homologs of only one constituent of this family were identified, suggesting that the complete system is not present. Members of the third family, (HlyC/CorC; TC#9.A.40) may function as divalent cation channels. TC subclass 9.B includes putative transporters, where even transport function is not established. These proteins are listed in Table S1 but will not be discussed.

### 3.9. Transporter substrates

The substrates of transporters found in the three leptospires likely reflect the physiological characteristics of these organisms. In Fig. 3, the distribution of substrates by category and subcategory is shown. Each chart in the top row represents the percentage of substrate types in each category for a given leptospire, and the chart below corresponds to the subclasses.

All three spirochetes have very similar percentages of the various substrate categories and subcategories. The most obvious difference between the three species is the relatively larger size of certain categories and subcategories in *L. biflexa*. Whereas *L. interrogans* and *L. borgpetersenii* have 87 and 89 inorganic substrate transporters, respectively, *L. biflexa* has 123, suggesting that this organism must be capable of maintaining intracellular ionic homeostasis under a much greater range of environmental conditions than for the two pathogens. Similarly, *L. biflexa* has a greater number of transporters for each category of substrate except nonselective transporters. The same can be seen in most subcategories where *L. biflexa* has more transporters than the other two spirochetes.

Correlating with its greater capacity for maintaining ionic homeostasis, *L. biflexa* has 98 proteins involved in cation transport whereas *L. interrogans* and *L. borgpetersenii* have 68 and 71, respectively. The prevalence of these transporters correlates with the disproportionately high number of *L. biflexa* secondary carriers that can utilize protons (H+) or sodium (Na+) for symport or antiport. This fact also indicates a reliance on transport that is energized by the proton (or sodium) motive force over other energy-coupling mechanisms such as those driven by ATP hydrolysis. In addition to ionic homeostasis, cation symport and antiport facilitate osmotic regulation [42] and heavy metal resistance [71]. Other inorganic monovalent and divalent cationic substrates pumped by these three organisms include potassium (K+), calcium (Ca2+), magnesium (Mg2+), and various cations of metals including copper, iron, zinc, cobalt, cadmium, mercury, and manganese.

Transporters specific for inorganic anions number 17 for *L. interrogans*, 15 for *L. borgpetersenii*, and 22 for *L. biflexa*. Anions, compared to cations, represent a much smaller proportion of the inorganic substrates transported by these leptospires. The latter play roles in redox processes, establishing, for example, the pmf or smf for energization. Anion transporters are found primarily in TC subclass 2.A, taking up or exporting bicarbonate, phosphate, arsenate, arsenite, telluride, chromate, chloride, and fluoride. Sulfate uptake, on the other hand, is mediated primarily by homologs of the CysPTW A ATP-dependent ABC system.

The three *Leptospira* species have only a small percentage of their transporters dedicated to organoanion transport. *L. biflexa* has four of these transporters, whereas *L. interrogans* has five and *L. borgpetersenii* has three. Fatty acids and other carboxylic acids, bile acids and their conjugates, taurine, and carnitine are the main substrates in this subclass of carbon sources. These spirochetes also
exhibit good representation of carbonate transporters with six in *L. biflexa*, two in *L. borgpetersenii*, and three in *L. interrogans*. Pyruvate, malate, succinate, acetate, hydroxybenzoate, citrate, and fumarate are all probably transported. These leptospires have outer membrane porins dedicated to fatty acid and hydrophobic compound uptake (TC#1.B.9.3.3). Fatty acid group translocation across the inner membrane may be catalyzed by transporters in the FAT Family (TC#4.C.1) [72–74].

Sugars and polyols taken up include glycerol, glycerol-3-phosphate, monosaccharides, and disaccharides. *L. biflexa* possesses eleven such proteins, whereas *L. interrogans* and *L. borgpetersenii* each possesses seven. As *Leptospira* utilize fatty acids as primary carbon sources, this subclass of carbon sources may play roles in osmoregulation, alternative metabolic pathways involving differentially expressed genes, and membrane construction [72–74].

*L. biflexa* has disproportionately high numbers of proteins involved in the transport of amines, amides, and polyamines (ten proteins compared to two each in *L. interrogans* and *L. borgpetersenii*). Among such transported substrates are putrescine, spermidine, ethanolamine, choline, and quaternary ammonium compounds. Transport of polyamines is associated with cellular growth and proliferation and may alleviate stress resulting from elevated external pH [75–77].

All three leptospires have transporters for amino acids and their conjugates, primarily members of the TC subclass 2.A. Representatives can be found in the APC (TC#2.A.3), DMT (TC#2.A.7), AGCS (TC#2.A.20), and ABC (TC#3.A.1) families. There appears to be substantial diversity in the types of amino acids transported. Of additional note, the three *Leptospira* species possess at least seven homologs to a putative alanyl teichoic acid synthesis protein DltB (TC#2.A.50). These homologs possess domains that strongly match with a DltB domain (e=48), and another weaker match for an O-acetyltransferase (e=24) (unpublished results). The strong match warrants inclusion of these proteins in this study as potential transporters of activated α-alanine, an amino acid conjugate.

The three spirochetes included in this study possess several proteins that function in the transport of peptides and their conjugates. These peptides can be di-/tripeptides, oligopeptides, and peptidoglycan fragments. These transporters additionally transport antibacterial agents, various nitrogen sources, and precursors of cell wall biosynthesis and primarily belong to the POT/PTR and ABC families.

Proteins can be secreted by *leptospires* using multiple systems. The General Secretory Pathway (TC#3.A.5) and the outer membrane secretory pathway (MTB) (TC#3.A.15) probably provide the primary pathways for protein secretion across the two membranes of the cell envelope. However, flagellar proteins and possibly some virulence proteins are secreted by the Type III Secretory Pathway (TC#3.A.6). Finally, the Outer Membrane Insertion BAM complex (TC#1.B.33) probably inserts most outer membrane proteins into this structure.

A key feature of *Leptospira* is its outer membrane, composed of lipids, porins, lipoproteins, and lipopolysaccharide (LPS); this last constituent consists primarily of lipid A and O-antigen. The Outer Membrane Lipopolysaccharide Export Porin (LPS-EP) Family (TC#1.B.42), the Outer Membrane LolAB Lipoprotein Insertion Apparatus Family (TC#1.B.48), the Multidrug/Oligosaccharidyl-lipid/Poly saccharide Flippase Family (TC#2.A.66), and members of the ABC Superfamily (TC#3.A.1) are the primary systems dedicated to the export of lipids and LPS precursors to the outer membrane. All three leptospires in this study possess proteins for the transport of drugs with sixteen in *L. borgpetersenii*, and twenty in both *L. interrogans* and *L. biflexa*. Multidrug resistance pumps are known to be prevalent in free living organisms which need to defend themselves against toxic substances produced by other microbes [78]. Further characterization of the members of the Drug/Metabolite Transporter Superfamily (TC#2.A.7) should provide a more accurate representation of the substrates transported by members of this superfamily. Drug exporters function to protect the cell from endogenously produced antibiotics, to remove exogenous and harmful substances produced by other microbes, and to export drug-like secondary metabolites such as siderophores, lipids, signaling peptides and periplasmic redox cofactors [78].

The rest of the “Drugs, vitamins, siderophores & cofactors” category of substrates in these three organisms is comprised of thirteen transporters in *L. interrogans* and *L. biflexa*, with only eight in *L. borgpetersenii*. All three leptospires transport Vitamin B12 (cobalamin), while only *L. biflexa* has a transporter designated as a Vitamin B1 (thiamine) transporter, and it belongs to the ABC Superfamily (TC#3.A.1.17.8). Siderophore transporters, also of the ABC-type, are found in all three organisms, consistent with a need for iron in these aerobes.

Nonselective channels include α-type channels, β-barrel porins, and pore-forming toxins. The three leptospires examined have transporters in these subclasses with sixteen in *L. interrogans*, fifteen in *L. borgpetersenii*, and eleven in *L. biflexa*. These nonselective transporters can play receptor roles in the outer membrane, induce toxin-like effects in other bacteria, and regulate cellular osmolarity (see TCDB). The greater numbers of these proteins in these two pathogens suggests that they play important roles in virulence.

All three leptospires have a substantial proportion of transporters that are poorly defined, about 12%. This represents a diverse subset of all leptospiral proteins with no clearly demarcated function [21–23]. Nonetheless, these proteins which may transport unknown substrates via ill defined mechanisms of action are likely to serve important, possibly genus-specific roles in metabolism and pathogenicity.

4. Discussion

4.1. Comparative leptosomal proteome analyses

Members of the genus *Leptospira* cause leptospirosis, a zoonotic disease with global prevalence, affecting nearly one million people annually with mortality ranging 5–25%. Current treatment of its wide variety of symptoms relies heavily on symptom management and antibiotic administration. Antibiotics currently in use to treat leptospirosis include penicillin, doxycycline, and cephalosporins, the efficacies of which remains mixed and questionable [79–83]. Prophylaxis usually involves vaccines of typically heat-attenuated leptospires with limited results. Chemophylactic treatment involves extended administration of doxycycline, a procedure that has been shown to reduce the incidence of the disease [84–86]. Variation in the susceptibility of different strains of *Leptospira* to these treatments has been reported, but instances of asymptomatic carriers in humans complicate the issue [87,88]. The causes of the variable clinical manifestations of leptospirosis are poorly understood, and the entire *Leptospira* genus is poorly characterized.

Comparative analyses of transport proteins provide clues as to the metabolic, pathological, and drug resistance properties of these spirochetes. By comparing and contrasting two known pathogenic members, *Leptospira interrogans* and *Leptospira borgpetersenii*, with a free-living saprophyte, *Leptospira biflexa* [21–23,89], we have generated data that will help define the metabolic capabilities of these organisms, allow identification of transporters common to these leptospires, and distinguish transport systems required for pathogenic versus saprophytic life.

Bioinformatics is a powerful tool for analyzing and assessing
4.2. Distinguishing transporters of three leptospires

All identified transporters are compiled with their characteristics in Table S1, revealing many of the conclusions drawn in our study. We show that L. borgpetersenii possesses 260 transport proteins, L. interrogans possesses 270, and L. biflexa possesses 337, showing that the two pathogens have fewer transporters than the free-living L. biflexa. This difference arises because of decreases in the secondary carriers, primary active transporters, and channel proteins in the former two organisms as seen in Fig. 2. This fact reflects decreased transport capabilities and therefore metabolic diversity and potential for homeostatic control of the pathogens relative in the free-living saprophyte.

Fig. 3 shows the very significant difference in the inorganic cation transporters of L. biflexa (98) compared to L. interrogans (68) and L. borgpetersenii (71). Transporters associated with these substrates include pmf and/or smf generators, osmotic and ionic homeostatic stress response regulators, and heavy metal resistance proteins. These cation transporters confer upon L. biflexa the ability to survive in external environments through effective osmotic regulation, metabolic versatility, and by competing with other environmental microbes.

L. biflexa has increased numbers of transporters for carbon sources and amino acids relative to L. interrogans and L. borgpetersenii. Among these substrates are carboxylates, sugars, polyols, non-carboxylic organoanions, amines, amino acids & their conjugates, and peptides & their conjugates. With just two exceptions, L. biflexa has more transporters in each of these subcategories of substrates. This undoubtedly reflects its superior metabolic versatility, conferring the ability of this free-living organism to grow under a wide range of environmental conditions. Pathogenic species of Leptospira are known to lack proteins related to carbohydrate, nitrogen, and amino acid metabolism that correlate with their protracted growth in artificial media [92]. Many of the aforementioned transporters, as seen in Table S1, belong to the 2.A TC subclass, the carriers of which are known to demonstrate lower affinities, but greater efficiencies at lower energy cost, than ABC transporters.

The relatively high affinity ABC transporters are well represented in all three leptospires, but L. biflexa has significantly more (67) of these transport proteins than L. interrogans (56) or L. borgpetersenii (49). Uptake systems for peptides and sulfate are present, but L. biflexa possesses a system for putrescine/spermidine uptake, as well as ones for siderophore, zinc (Zn\(^{2+}\)), and vitamin acquisition. High affinity acquisition of putrescine, critical in cell survival, demonstrates a crucial component of L. biflexa saprophytism [93]. Similarly, uptake systems for iron siderophores, thiamine, and zinc (Zn\(^{2+}\)) serve to accumulate them to high concentrations within the cell for use as cofactors so pathogens have better access within a host. Macromolecular ABC export systems are similarly well represented in all three species, transporting proteins and polysaccharides. These spirochetes also exhibit differential abilities to export drugs including antibiotics. L. biflexa, however, uniquely has homologs of exporters for fatty acyl-CoA and putative adhesin proteins. Fatty acyl-CoA export may function to acylate outer membrane constituents including adhesin proteins, likely to play a role in biofilm formation.

Interspecies differences in transporter classes and substrate categories are limited in the Leptospira species analyzed here. Individual transport proteins serve as likely contributors to pathogenesis in L. interrogans and L. borgpetersenii. Both pathogens encode members of the SpH Hemolysin Family (TC#1.C.67) that are notably absent in L. biflexa [94–96]. SpH2 (Uniprot # P59116; Table S1), is a sphingomyelinase with all active site residues essential for catalysis in vitro. Sphingomyelinase C (Uniprot # Q04XS2) of L. borgpetersenii is closest in sequence to SpH2. Both pathogens, but not L. biflexa, possess proteins with strong sequence similarity to a member of the Mycobacterial 4 TLS Phage Holin Family (TC#1.E.40). The proposed roles of prokaryotic holins in cell lysis and biofilm formation indicate the potential role these proteins may play in pathogenesis [37]. Both L. interrogans and L. borgpetersenii possess a member of the Putative Peptide Transporter Carbon Starvation CstA Family (TC#2.A.114). Mutation of a homolog in Campylobacter jejuni displayed decreased host-pathogen interactions [97]. All three leptospires possess proteins exhibiting sequence similarity to a member (TC#1.B.6.1.20) of the OmpA-OmpF Porin family. The L. interrogans protein queried has been shown to be Loa22, a protein essential for leptospiral virulence [98]. The relative dissimilarity of the L. biflexa homolog could render it avirulent.

4.3. Transporter hallmarks of leptospira

Leptospira is a branch of a divergent phylum and represents a genetically isolated group of bacteria [21]. Transporters identified in the leptospires in this study potentially serve novel roles for these gram-negative aerobes. All three leptospires possess a significant number of z-type channels, most of which transport inorganic ions and small metabolites. The largest representative of these channel components is the MotAB/ExbBD/ToIQR channel-forming protein (TC#1.A.30) with roles in motility, energized outer membrane transport, and outer membrane stability, respectively. The presence of two MotAB energizers may permit one system to utilize the proton motive force and the other the sodium motive force [47]. L. interrogans is notably deficient in ExbBD/ToIQR energizers relative to L. borgpetersenii and L. biflexa, which raises questions about the role of these energizers in Leptospira [48–50].

Of strong clinical relevance is the leptospiral outer membrane proteome (surfaceome), constituted largely by a collection of β-type porins. It contains cell surface antigens for potential vaccine production and drug targets [57]. As mentioned above, all three leptospires possess Loa22 (or a homolog), a surface-exposed porin, necessary for leptospiral virulence. Aside from this virulence factor,
Leptospires possess outer membrane transporters for the transport of small molecules, larger molecules including siderophores, proteins and membrane constituents. The leptospiral outer membrane plays a role in transport of substrates from the extracellular space to the periplasm.

The largest TC subclass identified in *L. interrogans*, *L. borgpetersenii*, and *L. biflexa* is 2.A, carrier proteins catalyzing uniport, antiport, and symport. The diversity of substrates transported is due to members of about 40 families of carriers. This distribution of transporters suggests an important role of secondary carriers in nutrient acquisition and drug resistance over primary active transporters such as ABC systems. The distribution of secondary carriers and primary active transporters reveals the prioritization of the acquisition and export of various molecules. Metabolic flexibility should dictate utilization of low-affinity secondary carriers, whereas a specific metabolic need might

Fig. 2. Distribution of transporters based on TC (A) classes and (B) subclasses in *L. interrogans*, *L. borgpetersenii*, and *L. biflexa*. 
necessitate other higher-affinity systems.

P-type ATPase (TC#3.A.3) distribution varies between all three leptospires, but all three possess an exporter of copper (Cu^{2+}), indicating a critical need for strict intracellular copper regulation. Only *L. interrogans* possesses ATPases for the uptake of Mg^{2+} and K^{+}, suggesting these ions play critical roles for this organism to survive in the external environment and/or in the host. *L. biflexa* possess ATPases for Ca^{2+} export, heavy metal resistance (Co^{2+}, Zn^{2+}, Cd^{2+}), and Na^{+}/K^{+} exchange. These proteins, unique to *L. biflexa*, likely highlight its effective osmoregulation and capacity for membrane potential maintenance.

All three leptospires possess multiple systems for protein secretion. The primary pathways for protein secretion across the two membranes of the cell envelope are probably provided by the

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**Fig. 3.** Distribution of transporters based on substrate (A) category and (B) subcategory in *L. interrogans*, *L. borgpetersenii*, and *L. biflexa*. 
General Secretory Pathway (TC#3.A.5) and the outer membrane secreting Main Terminal Branch (TC#3.A.15). Flagellar proteins and potential virulence proteins are secreted by the Type III Secretory Pathway (TC#3.A.6). This particular pathway is critical for virulence, as inhibition of flagellar motility in leptospires has been shown to render them avirulent [64].

Common to all three leptospires are primary proton pumping electron transfer complexes inherently present in mitochondria and many aerobic bacteria. These systems include the proton-translocating NADH dehydrogenase (TC#3.D.1), proton-translocating transhydrogenase (TC#3.D.2), and proton-translocating cytochrome oxidase (TC#3.D.4). The presence of these proton-translocating systems is consistent with the conclusion that electron transfer is used as a primary mechanism to generate a proton motive force, subsequently to drive solute transport via secondary carriers and used for ATP synthesis. The prevalence of secondary carriers correlates with this primary mechanism of energy generation.

As leptospires are genetically divergent compared to most well studied bacteria, they are expected to share a strong core of proteins and possess unique systems for pathogenesis and life in the open environment [21–23]. A significant portion of the identified transporter proteome in these leptospires are immediate, such as transporters from TC subclasses 9.A and 9.B. Further identification and characterization of these proteins, in addition to the remaining encoded non-protein-transport proteins, should provide a more complete understanding of leptospirosis pathogenesis and saprophytism.

Key attributes of Leptospira are aligned with their motile and chemotactic abilities. The embedded flagelli permitting corkscrew-like motility favors these organisms in host dissemination and environmental survival [99]. Transport proteins identified in this study, including a flagellar export system, chemotaxis proteins, and flagellar motor energizers, play roles in survival and virulence. Chemotaxis toward specific molecules like glucose may facilitate tissue tropism of Leptospira pathogens [99]. L. biflexa can persist over long periods of time in distilled water by forming biofilms, and aggregation has been suggested to allow environmental survival and host colonization [74,100]. Transport proteins that excrete exopolysaccharides, signaling molecules, and adhesion proteins also promote biofilm formation for persistence [101–103].

Transport proteins represent a subset (about 10%) of the entire proteome of an organism. However, intracellular processes are dependent on what materials are available in the cell. By providing an overview of the molecules transported and how they are imported and exported, conclusions about the metabolism and physiology of an organism can be drawn. In the case of Leptospira, the overall transportome (transporters of the proteome), reveals key characteristics of saprophytism and pathogenesis. L. biflexa demonstrates high flexibility and versatility in its transportome with a relatively large subset of secondary carriers and transporter families not found in the pathogens. In addition, L. biflexa possesses high-affinity transporters for critical cofactor import and increased numbers of uptake systems for carbon and nitrogen sources. Meanwhile, the pathogens possess remarkably similar transport protein profiles, suggesting that host tropism and environmental survival for the two relies on similar transporters. Both possess factors that may be associated with pathogenesis, absent in L. biflexa, such as pore-forming toxins, holins, and virulence-related outer membrane porins. The increased versatility of L. biflexa as a free-living organism likely reflects the inverse as the decreased versatility in the pathogens forces them to realize progressively narrower ecological niches. As leptospirosis manifests with a variety of symptoms, small differences within individual proteins and the leptospiral proteome may play roles in determining virulence and mortality in humans [104]. The findings reported here on these leptospiral transporters should improve understanding of the pathology of leptospirosis and allow more specific experimentation with L. biflexa as a model system for the Leptospira genus.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.micpath.2016.06.013.

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