Multilocus sequence typing methods for the emerging *Campylobacter* species *C. hyointestinalis, C. lanienae, C. sputorum, C. concisus,* and *C. curvus*

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Multilocus sequence typing (MLST) systems have been reported previously for multiple food- and food animal-associated Campylobacter species (e.g., C. jejuni, C. coli, C. lari, and C. fetus) to both differentiate strains and identify clonal lineages. These MLST methods focused primarily on campylobacters of human clinical (e.g., C. jejuni) or veterinary (e.g., C. fetus) relevance. However, other, emerging, Campylobacter species have been isolated increasingly from environmental, food animal, or human clinical samples. We describe herein four MLST methods for five emerging Campylobacter species: C. hyointestinalis, C. lanienae, C. sputorum, C. concisus, and C. curvus. The concisus/curvus method uses the loci aspA, atpA, glnA, gltA, glyA, ilvD, and pgm, whereas the other methods use the seven loci defined for C. jejuni (i.e., aspA, atpA, glnA, gltA, glyA, pgm, and tkt). Multiple food animal and human clinical C. hypintestinalis (n = 48), C. lanienae (n = 34), and C. sputorum (n = 24) isolates were typed, along with 86 human clinical C. concisus and C. curvus isolates. A large number of sequence types were identified using all four MLST methods. Additionally, these methods speciated unequivocally isolates that had been typed ambiguously using other molecular-based speciation methods, such as 16S rDNA sequencing. Finally, the design of degenerate primer pairs for some methods permitted the typing of related species; for example, the C. hyointestinalis primer pairs could be used to type C. fetus strains. Therefore, these novel Campylobacter MLST methods will prove useful in differentiating strains of multiple, emerging Campylobacter species.

Keywords: MLST, emerging, Campylobacter hyointestinalis, Campylobacter lanienae, Campylobacter concisus, Campylobacter curvus, Campylobacter sputorum

INTRODUCTION

Campylobacters are a major cause of human bacterial gastrointestinal illness in the industrialized world (Mølbak and Havelaar, 2008; Olson et al., 2008); campylobacterioses (12.68 cases per 100,000) were second only to Salmonella infections (16.2/100,000) in the United States in 2008 (Anonymous, 2009). The majority of *Campylobacter* strains isolated from human clinical samples have been identified as C. jejuni subsp. jejuni or, to a lesser extent, C. coli (Lastovica and Allos, 2008). Recently, pathogenic campylobacters outside of the C. jejuni/C. coli group, termed here as emerging Campylobacter species, have been isolated more frequently from food and/or food animals. Recovery of these more fastidious, emerging Campylobacter species from food has not been reported often; isolation of such strains is likely limited by the culture conditions employed, conditions that favor *Campylobacter* species such as C. jejuni and C. coli. However, Lynch et al. (2011) using novel culture conditions, reported the isolation of multiple emerging Campylobacter spp., e.g., C. concisus, C. curvus, and C. sputorum, from chicken, beef, and pork samples. Emerging campylobacters isolated from food animals are often strains of species associated typically with livestock, such as *C. hyointestinalis* in sheep, cattle, and swine (Hakkinen et al., 2007; Salihu et al., 2009; Oporto and Hurtado, 2011), *C. lanienae* in cattle and swine (Sasaki et al., 2003; Inglis et al., 2004; Oporto and Hurtado, 2011), and *C. sputorum* in cattle and sheep (Terzolo, 1988; On et al., 1998).

The clinical relevance of the emerging *Campylobacter* spp. is as yet undetermined. Many of the emerging campylobacters are isolated infrequently from human clinical samples, although, as with isolation from food, recovery of these strains from clinical samples is probably limited by the isolation methods and media used. Nevertheless, emerging *Campylobacter* species are isolated from human clinical samples (Edmonds et al., 1987; Gorkiewicz et al., 2002; Lastovica and Allos, 2008; Bullman et al., 2011). Although the frequency of human illness associated with emerging *Campylobacter* spp. might be quite low, especially when compared to *C. jejuni*-associated gastroenteritis, it is possible that some emerging species could be associated with more severe illness. One such example is *C. concisus*, for which a strong association with Crohn's disease and ulcerative colitis, has been reported recently (Man et al., 2010; Mahendran et al., 2011; Mukhopadhya et al., 2011).

Although molecular detection methods exist for many of the emerging campylobacters, population analyses, epidemiology, and source tracking of these organisms are limited by the strain typing methods available for these taxa. Molecular typing methods such as amplified fragment length polymorphism (AFLP) analysis and pulsed field gel electrophoresis (PFGE) methods have been employed on emerging Campylobacter strains (reviewed in On et al., 2008); however, sequence-based typing methods are not available for many species. One such sequence-based typing method is multilocus sequence typing (MLST). MLST methods amplify and sequence defined regions of moderately conserved housekeeping loci. At each locus, regions with distinct sequences receive arbitrary but unique allele numbers; similarly, each different allelic profile is assigned a unique sequence type (ST). The first Campylobacter MLST method was developed for C. jejuni (Dingle et al., 2001). This method sequences portions of seven genes: aspA, atpA (uncA), glnA, gltA, glyA, pgm (glmM), and tkt. The C. jejuni MLST method has been used in multiple typing studies and has been used successfully for strain typing and characterization, identification of clonal complexes and lineages, epidemiology, and investigation of host/source-associations (reviewed in Maiden and Dingle, 2008). Since the description of the C. jejuni MLST method, other Campylobacter MLST methods have been constructed that type C. coli (Dingle et al., 2005; Miller et al., 2005), C. lari (Miller et al., 2005), C. upsaliensis (Miller et al., 2005), C. helveticus (Miller et al., 2005), C. fetus (van Bergen et al., 2005), and C. insulaenigrae (Stoddard et al., 2007). Besides the primary use of Campylobacter MLST data for strain typing, MLST data for multiple taxa within Campylobacter are a valuable resource for studies on lateral gene transfer and evolution. MLST data can be used also to identify putative and perhaps clinically relevant taxonomic subdivisions within a species (Miller et al., 2005); additionally, MLST can provide genotypic information for novel species that are diverse phenotypically (Stoddard et al., 2007), especially those for which the only molecular speciation method is 16S rDNA sequencing.

Development of several Campylobacter MLST methods was assisted by the availability of draft genome sequences (Miller et al., 2005). Development of the novel Campylobacter MLST methods described in this study utilized recent draft genomes of C. hyointestinalis, C. lanienae, and C. sputorum (Miller et al., unpublished data), in addition to the closed C. concisus and C. curvus genomes available in the NCBI Microbial Genomes database. We anticipated that the draft genomes would contain some sequencing errors; however, enough reliable sequencing data was available to design MLST primers that could be used to type these five Campylobacter species. Therefore, in this study we describe four novel MLST methods that can be used to type: (1) C. concisus and C. curvus; (2) both subspecies of C. hyointestinalis (subspp. hyointestinalis and lawsonii) and C. fetus (subspp. fetus and venerealis); (3) C. lanienae; and (4) all three biovars of C. sputorum (bvs. fecalis, paraureolyticus, and sputorum). All four MLST gene sets are identical to the *C. jejuni* gene set [i.e., *aspA*, *atpA*(*uncA*), *glnA*, *gltA*, *glyA*, *pgm*, and *tkt*], with the exception of the *C. concisus/C. curvus* MLST method in which *ilvD* replaces *tkt*. A sample set of 213 isolates of diverse geographic origin and source was typed in this study. For all four methods, a total of 163 STs and 729 alleles were identified, indicating that these new MLST methods provide resolution similar to the previous MLST methods described.

MATERIALS AND METHODS GROWTH CONDITIONS AND CHEMICALS

All *Campylobacter* strains were cultured routinely under microaerobic conditions (1.5% O₂, 10% H₂, 10% CO₂, and 78.5% N₂) at 37°C on Brain Heart Infusion agar (Becton Dickinson, Sparks, MD, USA) or Anaerobe Basal Agar (ABA; Oxoid, Lenexa, KS, USA) supplemented with 5% (v/v) laked horse blood (Hema Resource and Supply, Aurora, OR, USA). PCR enzymes and reagents were purchased from New England Biolabs (Beverly, MA, USA) or Epicentre (Madison, WI, USA). All chemicals were purchased from Sigma-Aldrich Chemicals (St. Louis, MO, USA) or Fisher Scientific (Pittsburgh, PA, USA). DNA sequencing chemicals and capillaries were purchased from Applied Biosystems (Foster City, CA, USA). PCR and sequencing oligonucleotides were purchased from Eurofins MWG Operon (Huntsville, AL, USA).

ISOLATION OF CAMPYLOBACTER FROM FERAL SWINE AND CATTLE

Cattle feces were inoculated into wells of a six-well microtiter plate containing 6 ml 1× Anaerobe Basal Broth (Oxoid) amended with Preston supplement (Oxoid), using a sterile cotton swab. Plates were placed inside plastic ZipLoc bags and incubated under microaerobic conditions (as above) for 24 h at 37°C, while shaking at 40 rpm. After incubation, a 10-µl loop of each enrichment culture was plated onto ABA amended with 5% laked horse blood and CAT supplement (Oxoid). Feral swine feces were plated directly, using a sterile cotton swab, onto ABA amended with 5% laked horse blood and CAT supplement. All plates were then incubated under microaerobic conditions at 37°C for 24 h. Bacterial cultures were than filtered through 0.2 µm mixed cellulose ester filters onto ABA plates and incubated at 37°C under microaerobic conditions. After 24 h, single colonies were streaked onto new ABA plates and incubated 24–48 h at 37°C for purification.

CAMPYLOBACTER SPECIATION

Campylobacter strains isolated from the feces of California feral swine or cattle were speciated initially by 16S rDNA sequencing, using the primer pairs 27F (5' AGA GTT TGA TCC TGG CTC AG 3') and 1392R (5' GAC GGG CGG TGT GTA C 3'; Lane, 1991). However, the 27F/1392R primers were not able to type *C. hyointestinalis* strains past the species level and several strains could not be typed unequivocally. To improve speciation, the atpAF/atpAR primer pairs from Miller et al. (2005) were used. These primers can amplify all *Campylobacter* taxa described currently, with the exception of *C. avium* (data not shown). Using these *atpA* primers, campylobacters of uncertain type were amplified and sequenced; the sequences were then compared by alignments and phylogenetic analyses to strains of known species/subspecies identification, especially those whose genomes had been sequenced. Unlike the 16S primers, the *atpA* primers could speciate unambiguously all

of the strains isolated in this study and could identify clearly both subspecies of *C. hyointestinalis*. The *atpA* primer pairs, however, could not differentiate the three known biovars of *C. sputorum* (i.e., *sputorum, fecalis*, and *paraureolyticus*; On et al., 1998). The MLST results provided further confirmatory speciation data; *atpA* speciation agreed completely with subsequent MLST speciation.

DETECTION OF UREASE AND CATALASE ACTIVITY

The biovar *paraureolyticus* can be distinguished from the other two biovars of *C. sputorum* by the production of urease (On et al., 1998). Therefore, to identify putative bv. *paraureolyticus* strains, *C. sputorum* isolates were assayed for urease activity, as follows: a 10µl loop of an overnight *C. sputorum* culture was resuspended in 2 ml urease reagent (3 mM NaH₂PO₄, 110 mM urea, 7 µg/ml phenol red, pH 6.8) and incubated for 1 h at RT. *C. sputorum* cultures were typed as bv. *paraureolyticus* based on a positive reaction (solution turning from yellow/orange to magenta). Genome-sequenced strains of biovars *sputorum* (strain RM3237) and *paraureolyticus* [strain RM4120 (LMG 11764)] were used as negative and positive controls, respectively. All tests were performed independently at least twice.

The *C. sputorum* biovars are distinguished also by the production of catalase: bv. *fecalis* is catalase-positive while the other two biovars are catalase negative (On et al., 1998). To test for catalase activity, a 10- μ l loop of an overnight *C. sputorum* culture was resuspended in 200 μ l 3% H₂O₂ on a glass slide. Presence of bubbles indicated a positive reaction. Genome-sequenced strains of biovars *sputorum* (strain RM3237) and *fecalis* [strain RM4121 (CCUG 20703)] were used as negative and positive controls, respectively. All tests were performed independently at least twice.

MULTILOCUS SEQUENCE TYPING

Each MLST amplification mixture contained: 1× MasterAmp PCR buffer (Epicentre, Madison, WI, USA), 1× MasterAmp PCR enhancer (Epicentre), 2.5 mM MgCl₂, 250 µM (each) dNTPs, 50 pmol each primer, and 1 U Taq polymerase (New England Biolabs). For strains where genomic DNA was extracted using kits or standard isolation protocols, 50 ng purified genomic DNA was added to each reaction tube. Otherwise, 2 µl of a boilate was added. Boilates were prepared by resuspending a 1-µl loop of a pure culture or a single Microbank bacterial storage bead (Pro-Lab, Austin, TX, USA) in 100 µl TE and heating at 80°C for 5 min, then 100°C for 20 min, and cooling to 4°C. MLST amplifications were performed on a Tetrad thermocycler (Bio-Rad, Hercules, CA, USA) with the following settings: 94°C for 30 s, 53°C for 30 s, and 72°C for 2 min (30 cycles). Amplicons were purified on a BioRobot 8000 workstation (Qiagen, Valencia, CA, USA). Cycle sequencing reactions were performed on a Tetrad thermocycler, using the ABI PRISM BigDye terminator cycle sequencing kit (version 3.1; Applied Biosystems) and standard protocols. Cycle sequencing extension products were purified using BigDye XTerminator (Applied Biosystems). DNA sequencing was performed on an ABI PRISM 3730 DNA Analyzer (Applied Biosystems), using POP-7 polymer and ABI PRISM Genetic Analyzer Data Collection and ABI PRISM Genetic Analyzer Sequencing Analysis software. Sequences were trimmed, assembled, and analyzed in SeqMan (v 9.1; DNASTAR, Madison, WI, USA).

ALLELE NUMBER/SEQUENCE TYPE ASSIGNMENT

The Perl program MLSTparser3 (Miller et al., 2009) was modified to include the novel MLST methods for *C. concisus*, *C. curvus*, *C. hyointestinalis*, *C. lanienae*, and *C. sputorum*. The expanded MLSTparser3 was used to identify the MLST alleles and ST of each *Campylobacter* strain typed in this study. New *Campylobacter* MLST databases were created¹; allele and ST data generated in this study were deposited in this database and are available online. The allelic profiles for all 213 strains are listed in Table S1 in Supplementary Material.

PHYLOGENETIC ANALYSES

A dendrogram of unique *Campylobacter* STs was constructed by concatenating the allele sequences comprising each ST. Allele sequences for each strain were concatenated in the order *aspA-atpA-glnA-gltA-glyA-pgm-tkt* with the exception of *C. concisus* and *C. curvus* allele sequences, that were concatenated in the order *aspA-atpA-glnA-gltA-glyA-ilvD-pgm*. Composite concatenate lengths were 3345 bp (*C. concisus/C. curvus*), 3312 bp (*C. fetus, C. hyointestinalis,* and *C. lanienae*), or 3321 bp (*C. sputorum*). Sequence alignments were performed using CLUSTALX (ver. 2.1)², and dendrograms were constructed using the neighbor-joining method with the Kimura two-parameter distance estimation method (Kimura, 1980). Phylogenetic analyses were performed using MEGA version 5.1 (Tamura et al., 2011). Polymorphic sites and d_n/d_s ratios were calculated using START2 (Jolley et al., 2001)

RESULTS AND DISCUSSION

DESIGN OF THE NOVEL CAMPYLOBACTER MLST METHODS

Construction of the novel *Campylobacter* MLST methods was facilitated by the availability of genome sequences for all of the taxa typed in this study. The genome-sequenced strains were: the completed genomes of *C. concisus* strain 13826 (NC_009802.1), *C. curvus* strain 525.92 (NC_009715.1), and *C. fetus* subsp. *fetus* strain 82–40 (NC_008599.1), and the draft genomes of *C. hyointestinalis* subsp. *hyointestinalis* (*Chh*) strain RM4092 (LMG 9260), *C. hyointestinalis* subsp. *lawsonii* (*Chl*) strain RM4096 (CCUG 27631), *C. lanienae* strain RM3663 (NCTC 13004), *C. sputorum* bv. *sputorum* strain RM3237, *C. sputorum* bv. *fecalis* strain RM4121 (CCUG 20703), *C. sputorum* bv. *paraureolyticus* strain RM4120 (LMG 11764), and strain RM6914, exemplar of a novel *C. concisus*-like clade (Mandrell et al., manuscript in preparation).

Primer design based on a sequence from a single strain might not lead to a successful MLST method if the sequence variation within that taxon prevents the design of primer pairs that efficiently amplify all strains. Therefore, MLST gene sequences from related species would be aligned. Based on this alignment, primers would be designed to bind to regions, 100–200 bp upstream and downstream of the allelic endpoints, that demonstrate a high degree of conservation among the aligned taxa. One to four degenerate bases would be incorporated into the MLST primers, if necessary, to optimize primer binding. This approach was used previously to construct successfully other *Campylobacter* MLST

¹http://pubmlst.org/campylobacter/

²http://www.clustal.org/

methods (Miller et al., 2005). Therefore, the full *aspA*, *atpA*, *glnA*, *gltA*, *glyA*, *pgm*, and *tkt* gene sequences were extracted from the completed and draft genomes and various alignments were performed. Based on sequence similarity between the various *Campy-lobacter* taxa, we developed four novel MLST methods to type the strains in this study: Method 1 for typing both *C. concisus* and *C. curvus*; Method 2 for typing *C. fetus* and both subspecies of *C. hyointestinalis*; Method 3 for typing *C. lanienae* strains; and Method 4 for typing all three biovars of *C. sputorum*.

The final MLST primer sets are listed in **Table 1**. Methods 2, 3, and 4 use the same seven loci and allelic endpoints of the *C. jejuni* MLST method, i.e., *aspA*, *atpA*, *glnA*, *gltA*, *glyA*, *pgm*, and *tkt*. However, for the *C. concisus/C. curvus* MLST method, the sequence diversity at the *tkt* locus was too great for the construction of suitable primers. Therefore, *tkt* was replaced by *ilvD* in Method 1; *ilvD* was used in a *C. jejuni* MLST method described previously (Manning et al., 2003). The Method 1 *aspA*, *atpA*, *glnA*, *gltA*, *glyA*, and *pgm* alleles also use the same endpoints of their *C. jejuni* counterparts; the *ilvD* allelic endpoints are unique to this study.

CHARACTERIZATION OF THE FOUR NOVEL *CAMPYLOBACTER* MLST METHODS

A total of 213 strains were typed in this study. Complete descriptions of each strain, including isolation source, date, and location (if known), allelic profiles and STs are listed in Table S1 in Supplementary Material. Strains typed in this study were isolated over a 30-year period (1981–2010) and were also geographically diverse: strains from each species were isolated on two to three continents (Table S1 in Supplementary Material). All of the *C. concisus* and *C. curvus* strains typed were isolated from human clinical samples, whereas nearly all of the *Chl* and *C. lanienae* strains typed were isolated from pigs and feral swine; strains from the other three taxa were a mixture of isolates from humans, cattle, and swine (**Table 2**).

With a few exceptions, the primary MLST primers listed in **Table 1** were able to amplify successfully all seven loci and provide high quality sequence data for all 213 strains to generate unambiguous ST. However, despite our best efforts to overcome sequence variation in the initial primer design, in a few instances (\sim 1–2%), the main primer pairs did not provide sequence quality high enough for an unambiguous ST. Thus, alternate primer pairs (annotated as "A" in **Table 1**) were used to amplify and sequence these alleles. No strain was excluded from the final strain list because a ST could not be obtained.

GENETIC DIVERSITY

Phylogenetic analysis of the MLST STs validated the taxonomic relationships observed previously (Debruyne et al., 2008) for the six *Campylobacter* species typed in this study. Although the use of different gene sets comprising Method 1 and Methods 2, 3, and 4 prevented the simultaneous analysis of all six species, a clear delineation between *C. concisus* and *C. curvus* strains was observed (**Figure 1**), as well as unambiguous segregation of the other four species (**Figure 2**), that included the related *C. hyointestinalis*, *C. fetus*, and *C. lanienae* taxa. Moreover, the two *C. hyointestinalis* subspecies, *Chh* and *Chl*, formed distinct clusters (**Figure 2**).

However, the two *C. fetus* subspecies could not be discriminated by phylogenetic analysis, consistent with previous observations (van Bergen et al., 2005). Divergent STs (*C. lanienae* STs 1 and 4; *C. sputorum* STs 8, 13, 14, and 15) were identified within some strains (see below and **Figure 2**). In *C. sputorum*, these divergent STs formed a cluster (termed Csp₂) distinct from a cluster (termed Csp₁) containing the other *C. sputorum* strains.

Many MLST STs and alleles were identified in this study (**Table 3**). With the exception of *C. fetus*, for which a previous study identified also a relatively small number of highly clonal STs (van Bergen et al., 2005), the majority of strains within each of the remaining five species possess unique STs. Phylogenetic analysis of the STs of these five species indicated that the least amount of variation exists within *Chh* and each of the two *C. sputorum* clades, whereas the greatest amount of variation detected here resides clearly within the *C. concisus* and *C. curvus* strain sets (**Figures 1–3**). Even with the inclusion of the divergent *C. lanienae* and *C. sputorum* STs, the average number of base substitutions per site was less in each case (0.0330, *C. lanienae*; 0.0265, *C. sputorum*; **Figure 3**) than that calculated for the *C. concisus* STs (0.0641; **Figure 3**).

The high degree of variation across the C. concisus STs is reflected by the large number of alleles and polymorphic sites identified within this strain set: for the 70 C. concisus strains, the number of alleles detected at any locus ranged from 55 (atpA) to 64 (*ilvD*; Table 3). This high density of alleles translated into the large number of polymorphic sites identified at each of the seven C. concisus loci (Table 4A). Over 100 polymorphic sites were present within the alleles of each of the C. concisus MLST loci (Table 4A), a large number when compared to the relatively few polymorphic sites detected within the Chh alleles, even when the relative sizes of the strain sets were factored into the comparison. For some species, the numbers of polymorphic sites were inflated by the presence of divergent alleles or strains within the strain set. When these alleles and strains were removed from the appropriate strain sets, the number of polymorphic sites decreased substantially. For example, removal of the divergent C. sputorum strains present within C. sputorum clade Csp₂ (Figure 3) eliminated 23 of 24 polymorphic sites at the glnA locus (Table 4A).

While MLST is often used as a strain typing tool, it can be used also to investigate the population structure of an organism, to identify lineages, for example, that demonstrate a higher association with disease or a particular host. Genes used for MLST methods, typically core housekeeping genes, are usually under purifying or neutral selection. Positively selected genes would be influenced by external/environmental pressures and would evolve more through recombination rather than through the accumulation of point mutations; thus, such genes are not generally used in MLST methods, since they may not provide an accurate representation of the clonal structure of a bacterial population (Maiden, 2006; Perez-Losada et al., 2011). One method of determining the level of selective pressure on a gene is by calculating the ratio between non-synonymous (d_n) and synonymous (d_s) base substitutions. The rate of synonymous base substitution in genes should equal the neutral substitution rate, in the absence of codon usage bias. Non-synonymous base substitutions (that result in an amino acid change) would be the result of positive

Locus	Allele size (bp)		Oligonucleo	tide primers				Meth	po		
						-		2		m	4
		Forward (5′–3′)		Reverse (5'-3')		ပိ	S	ರ	ъ	ច	Cs
asp	477	aspCCCFN1	GGHCAAAGCACAAATGAYRCTTATCC	aspCCCRN1	GCCWAGDACTGATTTTARGCAAGC	۵_	٩				
		aspCCF1	CAAAGCACAAATGACACTTATCCAA	aspCCR1	GRACTGATTTTAGGCAAGCCTCAGG	۷	∢				
		HFaspF	CTTTGAWAAAAAGCAGARGAGTTTAA	HFaspR	GCTGTAACGATACCGATTGAGTTATAA			٩	٩		
		HYOaspF	AAGAGTKGCTATTTATGAAAGACTTTGTG	HYOaspR	AATGCTGTAACGATACCGATTGAGT				Þ		
		LANaspF	TTTAGCCACAGCTATGGAGTATCTCAA	LANaspR	ATATGGGTTRAAWGCTGTAACRATACC					٩	
		HFLaspXF	AAYATGAAY GCAAACGAAGTTATAGC	LANaspR	ATATGGGTTRAAWGCTGTAACRATACC					∢	
		SPUTaspF	GAAACATTTGCAAGACGAACTAGAT	SPUTaspR	AAATGCTGTTACTATACCAACTGATCC						٩
atpA	489	atpCCCFN1	GATACYATCATYAAYCAAAAAGGTCAAGA	atpCCCRN2	GGTATTTHGCCTCKATATAWGGATAWAGC	٩	٩				
		atpCCF1	ACTATCATCAAYCAAAAAGGTCAAG	atpCCR2	GGATAWAGCTCMGCTTCAAATTTTGT	۷	۷				
		HFatpF	GTAT YAAAGCTATWGACGGTTTGGTTC	HFatpR	GAGYGGGCTATAAGGAGGTTG			٩	٩.		
		HYOatpF	ATGTVGCTATMGGTCAAAARCAATC	HYOatpR	TTTCTACWGGRAGYGGGCTATAAGG				۷		
		LANatpF	AACCAAAAAGGTCAAGATGTTATATGT	LANatpR	ATTTTCTACTGGAAGTGGGCTATAAGG					٩	
		HFLatpXF	CMAAAGGHATYATGGCTAGAAAAT	HFLatpXR	TTCRATATCTTTATCHAGWGCTTTTTT					۷	
		SPUTatpF	ACTATCATAAATCAAAAAGGCGGAG	SPUTatpR	TTCTCAATACCAAGRGGTGAATAAG						٩
gInA	477	gInCCCFN1	GSTTGGCAYAGCATAAGYTACAAC	gInCCCRN2	GTTYGTGCTWGGGTTTGTRAAGGC	٩	٩				
		gInCCF1	GSYTGGCACAGCATAAGCTACAAC	gInCCR2	ACCRTCTTTCCARACTGATTGATGC	∢	∢				
		HFgInF	GGCATCACGTATCKTATAATAAAAGC	HFgInR	ATGTACGTGCATACCGCTTCC			٩	٩		
		HYOgInF	TCTTATAATATAAAAGCCGTGAGCGAA	HYOgInR	CCRTCTTTCCATATACTTTGATGTACG				A		
		LANgInF	TGGCAYCAYGTATCWTATAATATAAAAGC	LANgInR	ATGGACRTGCATACCRCTWCCATTATC					٩	
		HFLgInXF	TTTYGAATWTTGTRAWGAAAATGAAGT	HFLgInXR	AGAGTAWGTWAGAATGCTTGGKGCTTC					∢	
		SPUTgInF	AGGAACTTGGCATCATGTTTCTTAT	SPUTgInR	CCATCTTTCCAAATAGATTGATGAA						٩
gltA	402	gltCCCFN1	GGGMTACACCTCRACKGCGATGTG	gltCCCRN2	CBCCRTGWGCCCAGCCCC	۵.	٩				
		gltCCF1	TACATCGACGGKCTAAARGGCGAG	gltCCR2	GATCTCYTCWAGCTGGCGGGATGAC	۷	۷				
		HFgltF	CTATAACRTTTATMGATGGWGAAAAGG	HFgltR	ATCAACYCTATCTGGAGTTCCTATCAT			٩	٩		
		HYOgItF	TATCACGTTTATAGATGGTGAAAAAGG	HYOgItR	YCTATCTGGAGTTCCTATCATYTCAAG				۷		
		LANgItF	ATGCATAGMGGMTATGATATAGCGTGG	LANgItR	CATCAACTCTATCTGGAGTWCCKATCA					٩	
		HFLgItXF	TACTGGTATGTTTACRTTTGATAGGGG	HFLgItXR	GAAATCMACATTTGGATATARATTTCT					∢	
		SPUTgltF	AAAAAGCATATTAAAACATCGTGG	SPUTgltR	TTATCCACACTTCCTATCATTTCTAGTT						٩

-ocus	Allele size (bp)		Oligonucleo	tide primers				Metl	poq		
						-		7		m	4
		Forward (5'–3')		Reverse (5'-3')		ပိ	S	പ്	ဗ	ច	Ö
glyA	507	glyCCCFN1	ACAAACAAATACGCMGAAGGCTA	glyCCCRN1	GATATCWGCRTCTTTWCCGCTAAA	۵.	۵.				
		glyCCF1	AARSAGCTTTTTGGMTGCGAA	glyCCR1	GATRTCWGCGTCTTTTCCGCTAA	۷	۷				
		HFglyF	GCAAAYGTTCAGCCAAATAGC	HFglyR	TITATTTACTGTTATTCCWGCRTTACC			٩	٩		
		HYOglyF	TTGCWAATGTTCAGCCAAATAGC	HYOglyR	ATTCCRGCATTRCCAAGAGCGATAT				۷		
		LANglyF	TGCWAATGTTCAGCCAAATAGCG	LANglyR	CAAGAGCGATATCRGCRTCTTTACC					٩	
		HFLglyXF	GATWGCTAGTGARAAYTTCACATATCC	HFLglyXR	GYCCTCTTTCATACCYCTTGC					∢	
		SPUTglyF	TGCAAATGTTCAACCAAACTCAG	SPUTglyR	GTAATTCCAGCATTTCCTAAAGCAT						٩
ivD	492	ilvCCCFN1	CGACTGGGCTAAAAGACGAGGA	ilvCCCRN1	TATGTGAGCGATRTTTYGGCTGAT	۵.	٩				
		ilvCCF1	CGACYGGGCTAAAGACGAGGAC	ilvCCR1	CKATGTGAGCGATGTTTTGGCTGAT	Þ	A				
шba	501	pgmCCCFN1	CARATMAAAAATTCHTTCCCAAAAGAG	pgmCCCRN1	CTTTTABGCATTTARGGCTTTTAYRAA	۵.	٩				
		pgmCCF1	ATCAAAAATTCMTTCCCAAAAGAGC	pgmCCR1	ATCAMRTTTTCRGTRCCAGAGTATCTAA	Þ	٩				
		HFpgmF	AAAAGGTTTRMGAGTTGTTTTGGACGT	HFpgmR	TAACGTTTTCWGTVCCWGAATATCTAAA			٩	٩		
		HYOpgmF	GCTTACCTTAAAAGGTTTRCGAGTTGT	HYOpgmR	TCATCACGTCCTTCAAGCAAAAG				Þ		
		LANpgmF	GCTTACYTTAAAAGGCCTRMGAGTTGT	LANpgmR	AAGAAGCAGYCTAATCAAATTYTCTGT					٩	
		HFLpgmXF	CAATMGCRTTTTTAACCGAAGATATG	HFLpgmXR	AATTITTCYACYTCTTCCATYTTTTA					∢	
		SPUTpgmF	TTCCAAAAGAGCTTACAATGTATGG	SPUTpgmR	TGTTCCTGAATATCTAAATAGTGAGCG						₽.
tkt	459, 468 (Cs)	HFtktF	TTTTRTGTGCVGATATGGTTCAAAA	HFtktR	TATGATWTCTTCRCCAAGMGGAGC			۵	٩		
		HYOtktF	CAATGGGACTTGCTGATTTAATGAG	HYOtktR	TCTTTGCMTCTTTTATGATATCTTCGC				Þ		
		HFLtktXF	AATAAGATTTTTRTGTGCVGATATGGT	HFLtktXR	AAGAGTGAATTTARMAGCTCTTTTTA					٩	
		LANtktF	CATCTAAAKCAYAATCCMAAAAATCC	LANtktR	ATCTCWKCGCCAAGMGGAGC					∢	
		SPUTtktF	TTGCGGATATTATGGTTGTTTTAGC	SPUTtktR	GATTAAATCCTGCGAGTTTTTTGC						٩

Co, C. concisus; Cv, C. curvus; Cf, C. fetus (both subspecies); Ch, C. hyointestinalis (both subspecies); Cl, C. lanienae; Cs, C. sputorum; P. primary MLST primer; A. alternate MLST primer.

Table 2	Source of the	Campylobacter	strains typed in	this study.
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Species	Subspecies	Strains	Human	Cow/cattle	Pig/feral swine	Other/unknown
concisus		70	70	0	0	0
curvus		16	16	0	0	0
fetus		21	6	4	8	3
hyointestinalis	hyointestinalis	39	14	16	9	0
hyointestinalis	lawsonii	9	0	1	8	0
lanienae		34	1	0	32	1
sputorum		24	2	9	8	5

selection. Thus, the ratio of non-synonymous to synonymous base substitutions (d_n/d_s) would be an indicator of potential positive selection: ratios > 1 would be evidence of possible positive selection, whereas ratios < 1 would be more indicative of purifying or stabilizing selection. The d_n/d_s values for previous *Campylobacter* methods were quite low: the highest d_n/d_s values for *C. jejuni* (0.093), *C. coli* (0.173), *C. lari* (0.047), *C. upsaliensis* (0.097), and *C. insulaenigrae* (0.110) were substantially <1 (Colles et al., 2003; Miller et al., 2005; Stoddard et al., 2007). Similar ratios (highest d_n/d_s values) calculated in this study for *C. concisus* (0.0295), *C. curvus* (0.0468), *Chh* (0.0516), *Chl* (0.0655), *C. lanienae* (0.0562), and *C. sputorum* (0.0426; **Table 4B**) are consistent with the previous methods, indicating that these MLST loci are also not subject to positive selection.

IDENTIFICATION OF PUTATIVE LATERAL GENE TRANSFER EVENTS AND NOVEL TAXA

Characterization of the Campylobacter MLST methods also identified putative lateral gene transfer events. An allele that was nearly identical to, and clustered phylogenetically with, alleles from another taxon was determined to represent a putative lateral gene transfer event. For example, strains RM14410 and RM14403 in the Chh strain set contain alleles pgm-14 and pgm-16, respectively (Table S1 in Supplementary Material), that are 97.8-99.6% similar at the nt level to pgm alleles from Chl but only 95.2–96.2% similar at the nt level to *pgm* alleles from *Chh* (data not shown); these two alleles are also clearly related to other Chl pgm alleles, based on phylogenetic analysis (Figure 4A). Strains RM14410 and RM14403 also contain the *atpA* alleles *atpA*-2 and *atpA*-13, respectively, that cluster with other Chh atpA alleles (Figure 4B). Alleles at the other five loci for these two strains cluster also with other Chh alleles (data not shown), indicating that RM14403 and RM14410 are Chh strains in which a putative lateral transfer event has occurred at the pgm locus. The extent of gene transfer in these two strains could not be determined by MLST and will require further genome sequence analyses. It is not surprising that Chl alleles were discovered in Chh strains: these two taxa are highly related, being subspecies of the same species, and are isolated often from the same food animals (i.e., cattle and swine). No putative lateral transfer events were observed within C. concisus or C. curvus and no alleles were identified that originated tentatively in another species.

In some instances, phylogenetically divergent alleles within a strain set were indicative of either highly divergent strains or perhaps novel taxa. Here, as in other MLST studies (Miller et al., 2005, 2009), putative lateral gene transfer events were identified at only one of the seven MLST loci. For example, Chh STs containing pgm-14 or pgm-16 (ST-26_{hh} and ST-28_{hh}, respectively; Figure 2) were divergent at only the pgm locus. The alleles for each of the remaining six loci within each ST were of likely Chh origin, and phylogenetic analysis of the concatenated allele sequences clearly placed these two STs within the Chh clade. However, some STs (ST-1_{lan}, ST-4_{lan}, ST-8_{sp}, ST-13_{sp}, ST-14_{sp}, and ST-15_{sp}) contain three to seven variant alleles (Table S1 in Supplementary Material). The C. lanienae STs ST-11an and ST-41an differ substantially from the other typed C. lanienae strains: concatenated nucleotide sequences across all seven loci for ST-11an and ST-4_{lan} are on average only approx. 92 and 88% similar, respectively, to the concatenated sequences of the other C. lanienae STs (Figure 2), which display an average 98% cross-similarity (data not shown). Therefore, these two C. lanienae STs may be exemplars of novel C. lanienae-related taxa (for comparison, the concatenated Chh nucleotide sequences are approx. 94% similar to those of the other C. hyointestinalis subspecies and 87% similar to those of the related species C. fetus; Figure 2). Additionally, six phylogenetically related, urease-negative strains of C. sputorum, all isolated from cattle over a 19-month time period, may be members of a novel taxon. Within these six strains, four divergent ST were identified (ST-8_{sp}, ST-13_{sp}, ST-14_{sp}, and ST-15_{sp}). The concatenated allele sequences of these four STs are 95% similar to STs from the three established C. sputorum biovars, that display only 1% sequence divergence across the 3321-bp (Figure 2). Thus, it is possible that these six strains are members of a C. sputorum-like taxon, perhaps a novel C. sputorum subspecies or biovar. Nevertheless, for both the divergent C. lanienae and C. sputorum strains, additional biochemical and molecular tests will need to be performed to definitively establish their taxonomic position within Campylobacter.

SUBTYPING OF C. CONCISUS AND C. SPUTORUM STRAINS

Previous studies investigating the diversity of *C. concisus* organized strains from this species into two major genetically diverse clusters or genomospecies (GS), based on strain typing using 23S rRNA PCR (Engberg et al., 2005; Kalischuk and Inglis, 2011) or AFLP (Aabenhus et al., 2005; Kalischuk and Inglis, 2011). Included in the *C. concisus* strain set here were several strains characterized previously by AFLP (Aabenhus et al., 2005). In agreement with these previous studies, phylogenetic analysis of the *C. concisus* STs identified two clusters: each cluster contained almost exclusively GS1 or GS2 strains (**Figure 1**). Within *C. concisus*, two to eight



FIGURE 1 | Dendrogram of *C. concisus* and *C. curvus* **STs**. Allele sequences for each strain were concatenated in the order *aspA-atpA-glnA-gltA-glyA-ilvD-pgm* and aligned using CLUSTALX. The dendrogram was constructed using the neighbor-joining algorithm and the Kimura two-parameter distance estimation method (Kimura, 1980). Bootstrap values >75%, generated from 1000 replicates, are shown at the nodes. Scale bar represents substitutions per site. Genomospecies 1–4 designations, as assigned by Aabenhus et al. (2005), were placed to the right of STs representing strains from that study. Arrows indicate the STs of the *C. concisus* and *C. curvus* genome-sequenced strains. The dendrogram contains also the concatenated *aspA, atpA, glnA, gltA, glyA, ilvD*, and *pgm* allele sequences of *Campylobacter rectus, Campylobacter showae, Campylobacter mucosalis,* and *Campylobacter hominis.* These allele sequences were extracted from draft (*C. rectus, C. showae,* and *C. mucosalis*) and completed (*C. hominis*) genome sequences.



sputorum bv. fecalis; sp (p), *C. sputorum* bv. paraureolyticus. Arrows indicate the STs of the genome-sequenced strains for each taxon. *Urease-positive strains. Values in parentheses at the nodes represent the average %nt similarity of the STs split at each node, following pairwise comparisons of the concatenated allele sequences.

alleles at each of the seven MLST loci were identified in more than one ST (Table S1 in Supplementary Material). It is noteworthy perhaps that of these 33 "common" MLST alleles, only one

Species	Subspecies	Strains	STs		Alleles								
				aspA	atpA	gInA	gltA	glyA	ilvD	pgm	tkt		
concisus		70	66	60	55	62	59	61	64	59	N/A		
curvus		16	11	8	7	6	6	7	9	7	N/A		
fetus		21	5	1	4	2	2	2	N/A	1	2		
hyointestinalis	hyointestinalis	39	31	8	6	7	5	12	N/A	12	13		
hyointestinalis	lawsonii	9	8	7	8	7	6	7	N/A	7	7		
lanienae		34	26	16	9	13	11	13	N/A	16	12		
sputorum		24	16	6	6	3	6	9	N/A	6	7		

Table 3 Sequence types and alleles identified by the	novel MLST methods.
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taxa. Each value represents the average number of base substitutions per site between concatenated allele sequences. Analyses were conducted using the Kimura two-parameter model (Kimura, 1980). The analysis involved 86 nucleotide sequences. Codon positions included were first + second + third+ non-coding. All positions containing gaps and missing data were eliminated. There were a total of 3312 positions in the final dataset. Evolutionary analyses were conducted in MEGA5 (Tamura et al., 2011). Branch lengths used to calculate the averages here were used to construct the dendrograms of **Figures 1** and **2**. NT, not tested. The *C. lanienae* and *C. sputorum* strain sets contained divergent STs. The STs ST-1_{lan} and ST-4_{lan} were removed from the *C. lanienae* strain set Clan_{all} to create Clan₁. ST-8_{sp}, ST-13_{sp}, ST-14_{sp}, and ST-15_{sp} were removed from the *C. sputorum* strain set Csp_{all} to create Csp₂; the remaining STs formed Csp₁.

allele (glyA-31) was identified in both GS1 and GS2 strains, suggesting that minimal genetic exchange occurs between the two genomospecies.

Division of *C. concisus* strains into two primary genogroups is not merely an academic exercise. In a study analyzing *C. concisus* strains isolated from diarrheic and non-diarrheic individuals, Kalischuk and Inglis reported that GS1 strains were isolated predominantly from healthy individuals while the GS2 cluster contained isolates primarily from diarrheic individuals (Kalischuk and Inglis, 2011); this correlation between diarrheal disease and GS2 *C. concisus* species was also observed by Aabenhus et al. (2005). Moreover, GS2 strains were reported to exhibit higher levels of epithelial invasion (Kalischuk and Inglis, 2011). Therefore, GS1 or GS2 strains would be predicted to lead possibly to different clinical outcomes, and the proper placement of *C. concisus* isolates into these two genomospecies would be critical not only clinically but also for epidemiological studies. The MLST method described here provides another accurate tool for *C. concisus* genomotyping.

Campylobacter concisus GS4 strains were isolated from severely immunodeficient patients and identified initially by limited phenotyping; comparative AFLP analysis and other DNA-based testing indicated their taxonomic position required clarification (Aabenhus et al., 2005). Given these data, and our MLST results, it is likely that these strains are *C. curvus* and not *C. concisus*. These species share many phenotypic traits (On et al., 1996) and are difficult to distinguish with limited phenotypic testing.

At present, C. sputorum biovars are identified by their catalase or urease phenotypes (On et al., 1998). No PCR/sequencing methods exist that differentiate the three described biovars, although wholegenome macro-restriction profiling has been used to suggest biovar clonality (On et al., 1999). However, putative biovar-associated alleles were observed at three of the C. sputorum MLST loci (Table S1 in Supplementary Material). For example, at the atpA locus, by. sputorum strains are *atpA*-1, by. *fecalis* strains are *atpA*-3, and bv. paraureolyticus strains are either atpA-2 or atpA-4; strains of the cow-associated C. sputorum clade described above are either atpA-5 or atpA-6. Similar associations exist at the glyA and pgm loci. Indeed, the three biovars segregate also when all seven loci are examined phylogenetically (Figure 2), although the differences are quite small. Obviously, the size of the C. sputorum strain set typed here is too small to reach any definitive conclusions, although the associations are intriguing.

CONCLUSION

The four MLST methods described in this study typed successfully all 213 *Campylobacter* strains, representing at least ten *Campylobacter* taxa that included both subspecies of *C. fetus* and *C. hyointestinalis* and all three biovars of *C. sputorum*. Each method identified multiple novel STs; the small number of STs present in some taxa were more likely due to the limited size of the sample sets for those taxa than a limitation of the method itself. These methods were successful despite the high degree of variation in some species, e.g., *C. concisus*. Also, the concatenated *C. concisus* and *C. curvus* ST sequences were only 81% similar at the nt level (**Figure 1**); likewise, the *C. hyointestinalis* and *C. fetus* ST sequences were 87% similar (**Figure 2**), yet MLST Methods 1 and 2 could type either set of strains unequivocally. Indeed, the ability to sequence such variable strains provided an unexpected bonus to these MLST methods. The methods described here identified putative novel *C*.

Species	Subspecies	Strains	aspA	atpA	gInA	gltA	glyA	ilvD	pgm	tkt
A. POLYMORP	HIC SITES									
concisus		70	131	105	120	102	129	140	128	N/A
curvus		16	90	47	47	51	40	63	41	N/A
hyointestinalis	hyointestinalis	39	9	9	10	27	15	N/A	30 (17)	24
hyointestinalis	lawsonii	9	23	36	10	11	64	N/A	19	21
lanienae		34	67 (29)	86 (51)	58 (19)	57 (26)	71 (26)	N/A	142 (31)	96 (38)
sputorum		24	58 (5)	33 (17)	24 (1)	28 (9)	39 (14)	N/A	18 (10)	20 (8)
B. RATIOS OF	NON-SYNONYM	OUS (d _n)T	O SYNONY	MOUS (d _s)	BASE SUBSTI	TUTIONS				
concisus		70	0.0257	0.0028	0.0052	0.0036	0.017	0.0295	0.0095	N/A
curvus		16	0.0417	0.0057	0.008	0.0468	0.0168	0.0149	0.0091	N/A
hyointestinalis	hyointestinalis	39	0	0	0	0.0262	0.0516	N/A	0.0249 (0.0110)	0.0381
hyointestinalis	lawsonii	9	0.0169	0.0276	0.0216	0	0.0113	N/A	0.0655	0.0251
lanienae		34	0.015	0.0204	0.0026 (0)	0.0502	0.0112	N/A	0.0562 (0.0748)	0.0421
			(0.0149)	(0.0235)		(0.0298)	(0.0257)			(0.0419)
sputorum		24	0.041	0 (0)	0 (0)	0 (0)	0.0102	N/A	0.0293 (0.0264)	0.0426
			(0.0715)				(0.0511)			(0.1672)

Table 4 | Diversity within the Campylobacter MLST loci.

Numbers in parentheses represent polymorphic sites and d_n/d_s ratios recalculated following removal of the divergent pgm-14 and pgm-16 alleles within the Chh profiles and removal of the divergent ST-1_{lan} and ST-4_{lan} (lanienae) and ST-8_{so}, ST-14_{so}, and ST-15_{so} (sputorum) sequence types.



lanienae- and *C. sputorum-*related taxa and it is likely that these methods could further characterize and type as-yet-undescribed *Campylobacter* species or subspecies. For example, MLST Method 2 has been used to type reptile-associated *C. fetus*-like organisms (data not shown).

For many campylobacters, sequence data is restricted currently to ribosomal rRNA loci. While these rDNA sequences can provide crucial speciation data for many taxa, some groups of campylobacters cannot be differentiated readily by 16S rDNA sequencing. One such example includes *C. hyointestinalis* and *C. lanienae* strains. Some of the strains in this study from these species could not be typed unequivocally by 16S rDNA sequencing; however, MLST could readily place all strains in their proper taxonomic positions. MLST has been shown also to be of value in identifying strains of species with multiple phenogroups, such as *C. insulaenigrae* (Stoddard et al., 2007). In this study, a *C. sputorum* clade was typed that, based on established phenotypic characterization, would likely have been classified as bv. *fecalis*. While additional tests need to be performed, MLST cast some doubt that these strains were *C. sputorum* bv. *fecalis*.

Eighteen of the thirty validly described *Campylobacter* taxa can now be typed by MLST. This number is likely an underestimate, as some of the *C. lari*-like species (e.g., *C. peloridis*) described recently can be typed also using the *C. lari* MLST method (data not shown). The ability of MLST to type and speciate campylobacters, as well as identify putative horizontal gene transfer, indicates that the multiple *Campylobacter* MLST methods now available provide a valuable tool in the epidemiology, typing, and evolution of emerging campylobacters.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at http://www.frontiersin.org/Cellular_and_Infection_Microbiology /10.3389/fcimb.2012.00045/abstract

Table S1 | Campylobacter strains typed in this study.

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