

Genomic Plasticity of the *rrn-nqrF* Intergenic Segment in the *Chlamydiaceae*[∇]

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In *Chlamydiaceae*, the nucleotide sequence between the 5S rRNA gene and the gene for subunit F of the Na⁺-translocating NADH-quinone reductase (*nqrF* or *dmpP*) has varied lengths and gene contents. We analyzed this site in 45 *Chlamydiaceae* strains having diverse geographical and pathological origins and including members of all nine species.

The complete genomes of six *Chlamydiaceae* species (1, 3, 12, 19–21, 23, 24) have revealed a high level of synteny, with large-scale rearrangements and niche-specific genes restricted to a “plasticity zone” located at the replication termination region. Analysis of the *rrn-nqrF* intergenic segments in these sequences reveals significant variation in length and gene content that is inconsistent with host range, tissue tropism, DNA-based phylogenies, and disease spectrum in humans and animals. Therefore, this segment may represent a new genomic plasticity zone in these bacteria. We were especially interested in a frameshifted invasin/intimin-like protein gene (*ilp*) at this location in *Chlamydia caviae* (20). Fragments of *ilp* in several *Chlamydia suis* strains were also identified by Dugan et al. (5). These observations led us to a comprehensive analysis of the *rrn-nqrF* intergenic sequences across the *Chlamydiaceae*. Strains are listed in Table 1. Data on PCR and sequencing primers are available upon request. Our findings are provided in a graphic summary (Fig. 1).

Species of the *Chlamydiaceae* can be separated into four groups based on the contents of their *rrn-nqrF* intergenic segments. Group 1 includes isolates from *Chlamydia pneumoniae*, *Chlamydia psittaci*, *Chlamydia abortus*, and *Chlamydia felis*, infectious to primate (respiratory tract and eyes), avian (respiratory and digestive tracts and eyes), ovine and bovine (digestive tract and placenta), and feline (respiratory tract and eyes) species, respectively. The *rrn-nqrF* segment is fully conserved in four *C. pneumoniae* strains (12, 19, 21) and is only 46 to 49% identical to segments in the three other species. The *rrn-nqrF* segment of the Japanese *C. felis* strain Fe/C-56 (1) is 99% identical to that of *C. psittaci* strain parakeet but only 63 and 61% identical to those of *C. psittaci* I-10 or 6BC and another *C. felis* strain, FEPN Baker, respectively. The last strain, however,

displays an *rrn-nqrF* intergenic sequence closely related to those of three *C. abortus* strains, S26 (24), B577, and LW508. Nucleotide substitutions, insertions, and deletions and hyper-variable segments including stretches of C's and T's (Fig. 2) are found in group 1 strains, causing various translational frames. A single open reading frame (ORF) at the *rrn-nqrF* site, identified as CF0129 in the *C. felis* strain Fe/C-56 genome (1), is also found in *C. psittaci* strains 6BC, parakeet, and I-10 and in *C. abortus* strain B577. However, mutations disrupt this ORF in *C. abortus* strains S26 and LW508, *C. felis* strain FEPN Baker, and four *C. pneumoniae* strains.

Group 2 includes isolates that carry at least partial *ilp* sequences at the *rrn-nqrF* site: *C. caviae* strain GPIC, *Chlamydia muridarum* strain MoPn, and eight *C. suis* strains, which infect guinea pigs (eyes and genital tract), mice (lungs), and pigs (digestive tract), respectively. Of those, only *C. suis* strain S45 (13) has intact *ilp*; all others have either fragments of the gene (*C. muridarum*) or a gene that is interrupted by mutation (*C. caviae* and Tet^c *C. suis*). The sequence similarity of *C. suis* Ilp to *Yersinia pseudotuberculosis* invasin (9) and enteropathogenic *Escherichia coli* intimin (11) is restricted to the transmembrane and immunoglobulin-like domains (D1). However, the general structure of the predicted Ilp polypeptide is similar to that of the invasin/intimin family and includes an extended rodlike structure consisting of several modular repeat domains (D2 to D7) and a distal domain (D8). Several conserved beta strands in D2 to D7 are consistent with the crystal structures of intimin D0 to D2 and invasin D1 to D3 (8, 16). Secondary-structure predictions, however, reveal no conserved salient features in D8 (<http://www.compbio.dundee.ac.uk/~www-jpred/>; domain definitions are per GenBank [accession number DQ076148]).

We reasoned that *ilp* of the *C. caviae* reference strain may have been inactivated through mutation during long-term in vitro culture. To address this question, the *rrn-nqrF* segment was amplified directly from early, archived specimens of *C. caviae* from the laboratory of Murray (18) (Table 1). Two specimens of the primary isolate that were maintained by passage through live animals, a 1962 specimen corresponding

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TABLE 1. Strains included in this study

Group, organism, and strain(s) or serovar(s)	Source	Accession no.	<i>rm-nqrF</i> intergenic segment length (bp)	Reference
Group 1				
<i>C. pneumoniae</i>				
AR39	Human	AE002161	2,047	19
J138	Human	NC002491	2,047	21
CWL029	Human	NC000922	2,047	12
TW183	Human	NC005043	2,047	This study
<i>C. psittaci</i>				
6BC	Parrot	DQ076132	2,334	This study
Parakeet	Parrot	DQ076133	2,218	This study
I-10	Pigeon	DQ076144	2,383	This study
I-6 and I-7	Parrots		2,200 ^d	This study
I-9 and I-11	Pigeons		2,200 ^d	This study
<i>C. abortus</i>				
S26/3	Sheep	CR848038	2,320	24
B577	Sheep	DQ076146	2,343	This study
LW508	Cow	DQ076147	2,346	This study
SV139 and I-17	Cattle		2,200 ^d	This study
I-1	Sheep		2,300 ^d	This study
I-5	Goat		2,300 ^d	This study
<i>C. felis</i>				
Fe/C-56	Cat	AP006861	2,229	1
FEPN sch562F4	Cat		2,200 ^d	This study
FEPN Baker	Cat	DQ076145	2,347	This study
Group 2				
<i>C. caviae</i>				
Yolk sac	Guinea pig	NC003361	4,463	20
Yolk sac 62H454 and 73H301	Guinea pig ^a		4,463	This study
<i>C. muridarum</i>				
MoPn	Mouse	AE002160	457	19
<i>C. suis</i>				
S45	Pig	DQ076148	4,184	This study
R19	Pig	AY428550	16,000 ^d	5
R27	Pig	AY428551	9,000 ^d	5
Group 3				
<i>C. trachomatis</i>				
A/HAR-13	Human	NC007429	519	3
D/UW-3/CX	Human	NC000117	518	23
E	Human	DQ851868	525	This study
G	Human ^b	DQ851869	518	This study
K	Human ^b	DQ851870	518	This study
D, H, and I	Humans ^b		518	This study
L2/LGV434 ^c	Human		519	
Group 4				
<i>C. pecorum</i>				
I-15	Cow	DQ076134	518	This study
I-12	Cow	DQ076135	469	This study
1710S	Pig	DQ076136	445	This study
I-19	Water buffalo	DQ076137	427	This study
LW679	Cow	DQ076138	419	This study
I-4	Goat	DQ076139	404	This study
L-17	Pig	DQ076140	388	This study
1708	Pig	DQ076141	380	This study
JP-1-751	Cow	DQ076142	356	This study
I-13	Cow	DQ076143	252	This study

^a The original isolate was from a guinea pig; no other *C. caviae* strains were available for examination.^b Clinical isolate(s) from the genital tract.^c The sequence for this serovar was retrieved from the Sanger Institute website at <http://www.sanger.ac.uk>.^d The length of the *rm-nqrF* segment is estimated by PCR and partial sequencing.

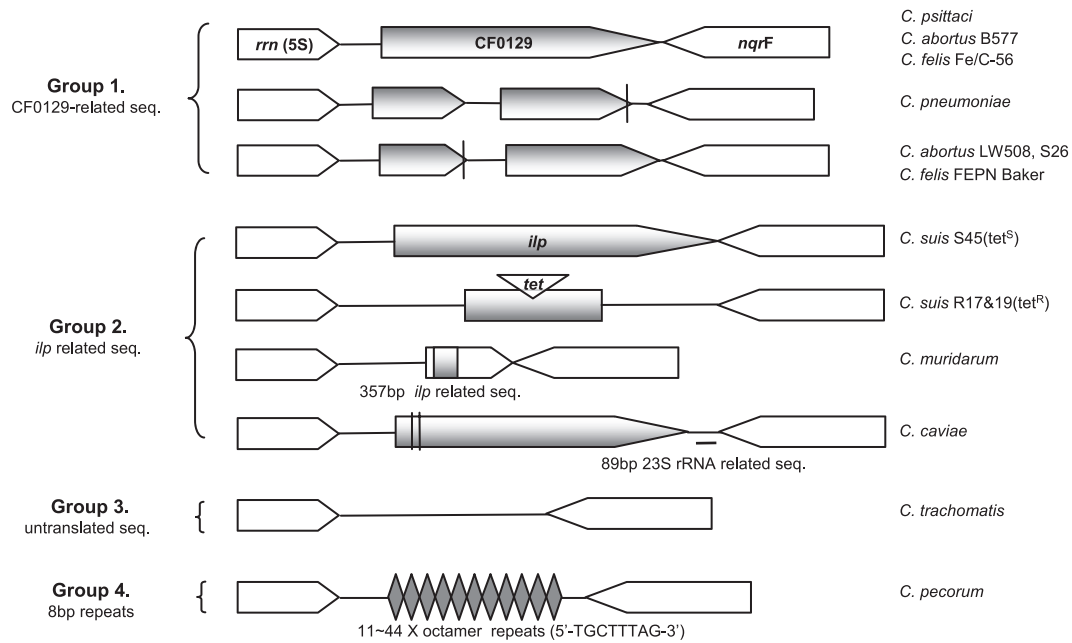


FIG. 1. Genetic diversity at the 5S *rrn-nqrF* site. The schematic diagram presents the genetic organization of the *rrn-nqrF* intergenic segment. *Chlamydia* species are separated into four groups based on genetic relatedness at the site. Vertical bars indicate mutations that cause frameshifts or translation stops. Homologous ORFs are indicated graphically as shaded arrows. The diagram is drawn approximately to scale. seq., sequence(s).

to the first passage and a 1973 specimen, were analyzed. In both cases, two frameshift mutations identical to those obtained from genomic analysis are present.

Group 3 includes only strains of the human pathogen *Chlamydia trachomatis*. PCR amplicons from six genitotropic strains, including clinical isolates of serovars D, G, H, I, and K and one serovar E reference strain, were compared. Sequences obtained from serovars D, G, and K are identical to the published sequence of serovar D strain UW-3/Cx (23). However, the serovar E sequence contains a 7-nucleotide insertion and 11 additional mutations, 7 of them located within the putative *rrn* transcription terminator (not shown). The *rrn-nqrF* intergenic sequence of lymphogranuloma venereum strain L2/434 (12) is identical to that of ocular strain A/HAR-13 (3) but less identical (96 to 99%) to that of urogenital strains (Table 1). The *rrn-nqrF* intergenic segments of all *C. trachomatis* strains and serovars contain no recognizable ORF.

Group 4 includes 10 isolates of *Chlamydia pecorum* obtained from mammalian species on two continents. Amplicons of the segments from these isolates range between 400 and 700 bp (Fig. 3A). Sequence analysis reveals multiple copies (between 11 and 44) of the simple sequence repeat (SSR) 5'-TGCTTT

AG-3' (Fig. 3B). BLAST searches reveal no matches, suggesting that this SSR is unique to *C. pecorum*.

Taken together, our results indicate an unusual degree of genetic plasticity at the *rrn-nqrF* locus in the family *Chlamydiaceae* that transcends phylogenetic boundaries. For instance, the *ilp* sequence, albeit degenerate or mutated, is common to *C. suis*, *C. muridarum*, and *C. caviae*, i.e., three species representing three different hosts and two phylogenetic lineages of the *Chlamydiaceae* (7). The apparent genetic plasticity of this site stands in sharp contrast with that of the neighboring rRNA operon, whose functional constancy and genetic stability are a basis of current phylogenetic analysis. It is possible that the plasticity of the *rrn-nqrF* segment stems from its being frequently exposed during chlamydial development. There are only one or two copies of *rrn* in the *Chlamydia* genomes. Elevated transcription activity of the *rrn* operon would require frequent unwinding of supercoiled DNA, which might promote physical damage (15), consistent with the site's being a preferred target for recombination. The presence of an 89-bp sequence nearly identical to a chlamydial 23S rRNA domain 1 sequence, flanking *ilp* in *C. caviae*, may indeed be a vestige of a previous horizontal *ilp* transfer event (Fig. 1). The identifi-

<i>C. felis</i> FEPN Baker	535 AACCTGCTGGAGAAGGTACCCCCCCCCCCCC~GCAACTGCTACCTTA----880	GGATCTTCATTTTTTTTTTTAGAGCTCTC
<i>C. abortus</i> S26	535 AACCTGCTGGAGAAGGTACCCCCCCCCCCCC~GCAACTGCTACCTTA----879	GGATCTTCATTTTTTTTT~AGAAGCTCTC
<i>C. abortus</i> LW508	535 AACCTGCTGGAGAAGGTACCCCCCCCCCCCCCGCAACTGCTACCTTA----881	GGATCTTCATTTTTTTTT~AGAAGCTCTC
<i>C. abortus</i> B577	535 AACCTGCTGGAGAAGGTACCCCCCCCCCCCC~GCAACTGCTACCTTA----877	GGATCTTCATTTTTTTTT~AGAAGCTCTC
<i>C. psittaci</i> I-10	566 AACATACTATA~~~GGTACCCCCCCCCCCCC~GCAACGAATACCTTA----906	GGATCTTCATTTTTCTTT~AGAAGATCTC
<i>C. psittaci</i> 6BC	527 AACCTACTATA~~~GTTACCCCCCCCCCCCC~GAAACGCATACCTTA----868	GGATCTTCATTTTTCTTT~AGAAGATCTC

poly(C) tract

poly(T) tract

FIG. 2. Multiple alignment of the poly(C) and poly(T) tracts and flanking sequences at the *rrn-nqrF* sites in three *Chlamydia* species. Base numbering is from the 3' end of *rrn*.

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