Genomic Resources for Sea Lice: Analysis of ESTs and Mitochondrial Genomes

Motoshige Yasuike · Jong Leong · Stuart G. Jantzen · Kristian R. von Schalburg · Frank Nilsen · Simon R. M. Jones · Ben F. Koop

Abstract Sea lice are common parasites of both farmed and wild salmon. Salmon farming constitutes an important economic market in North America, South America, and Northern Europe. Infections with sea lice can result in significant production losses. A compilation of genomic information on different genera of sea lice is an important resource for understanding their biology as well as for the study of population genetics and control strategies. We report on over 150,000 expressed sequence tags (ESTs) from five different species (Pacific Lepeophtheirus salmonis (49,672 new ESTs in addition to 14,994 previously reported ESTs), Atlantic L. salmonis (57,349 ESTs), Caligus clemensi (14,821 ESTs), Caligus rogercresseyi (32,135 ESTs), and Lernaeocera branchialis (16,441 ESTs)). For each species, ESTs were assembled into complete or partial genes and annotated by comparisons to known proteins in public databases. In addition, whole mitochondrial (mt) genome sequences of C. clemensi (13,440 bp) and C. rogercresseyi (13,468 bp) were determined and compared to L. salmonis. Both nuclear and mtDNA genes show very high levels of sequence divergence between these ectoparasitic copepods suggesting that the different species of sea lice have been in existence for 37–113 million years and that parasitic association with salmonids is also quite ancient. Our ESTs and mtDNA data provide a novel resource for the study of sea louse biology, population genetics, and control strategies. This genomic information provides the material basis for the development of a 38K sea louse microarray that can be used in conjunction with our existing 44K salmon microarray to study host–parasite interactions at the molecular level. This report represents the largest genomic resource for any copepod species to date.

Keywords Lepeophtheirus salmonis · Caligus clemensi · C. rogercresseyi · Lernaeocera branchialis · Expressed sequence tags (ESTs) · Mitochondrial genome

Introduction

Copepods (Copepoda) are a group of small crustaceans found in various aquatic environments and they are described as the most abundant metazoans on earth (Humes 1994). The subclass Copepoda consists of over 250 described families, 2,600 genera, and 21,000 described species classified into ten orders (Walter and Boxshall 2008). Their life histories are diverse; planktonic and benthic copepods are an important ecological link in the
aquatic food chain (Gee 1987; Ohman and Hirche 2001),
but approximately one third of marine copepod species live
as associates, commensals, or parasites on invertebrates and
fishes (Humes 1994).
Parasitic copepods are commonly found both on farmed and
wild marine finfish (Johnson and Fast 2004). They feed
on host mucus, epidermal cells, tissues, and blood, the
result of which causes physiological stress, immune
dysfunction, impairment of swimming ability, and possibly
death (Boxaspen 2006; Costello 2006; Johnson and Fast
2004; Tully and Nolan 2002). Members of the family
Caligidae, especially the genera Caligus and Lepeophthei-
rus, are commonly referred to as sea lice (Costello 2006;
Johnson et al. 2004; Pike and Wadsworth 1999). They are
the most economically important parasites of the world
salmon farming industry and may cause direct and indirect
economic losses in the industry of €300 million (US
$480 million) annually (Costello 2009). In addition, there
is concern that salmon farms elevate the risk of sea lice
infections on wild salmon beyond that which naturally
occurs and lead to a depression in the abundance of wild
salmon stocks (Costello 2006; Heuch et al. 2005; Krkošek et al. 2007a; Krkošek et al. 2007b; Todd et al. 2006).
In the North Atlantic Ocean, Lepeophtheirus salmonis and
Caligus elongatus account for the most serious infestations
of cultured and wild salmonids (Johnson et al. 2004; Pike
and Wadsworth 1999). In the eastern north Pacific Ocean, L. salmonis and Caligus clemensi have been found on farmed
Atlantic salmon (Salmo salar) and wild Pacific salmon
(Oncorhynchus spp.; Beamish et al. 2009; Beamish et al.
2005; Saksida et al. 2007). While L. salmonis is prevalent in
both Atlantic and Pacific coasts, earlier studies suggested
that the Pacific and Atlantic populations of L. salmonis are
genetically distinct (Tjensvoll et al. 2006; Todd et al. 2004).
More recent genomic studies strongly suggest that distinct
species of L. salmonis exist in the Pacific and Atlantic
Oceans following a separation that occurred from 2.5 to
11 million years ago (Boulding et al. 2009; Yazawa et al.
2008). These parasites are referred to herein as the Pacific
and Atlantic forms of L. salmonis, respectively. In the
southern hemisphere, Caligus rogercresseyi is the dominant
species affecting salmonid aquaculture in Chile where the
parasites were found on farmed salmon in 99% of the
established cultured cages (Boxshall and Bravo 2000;
Carvajal et al. 1998).
Lepeophtheirus and Caligus species are distinguished
from each other based on morphological characters (Kabata
1979). The life cycle in L. salmonis has a total of ten
developmental stages, while C. elongatus and C. roger-
cresseyi are similar but appear to lack pre-adult stages
(Piasecki and MacKinnon 1995; González and Carvajal
2003). The host range of L. salmonis mainly includes
salmonids but the parasite has also been reported from non-
salmonid hosts, including sticklebacks, that co-occur with
salmon (Jones et al. 2006). In contrast, some Caligus
species have a broad host range of salmonids and non-
salmonids (Costello 2006; Johnson et al. 2004). Among its
salmonid hosts, L. salmonis displays clear preferences, with
heaviest infestations and greatest impacts occurring on
Atlantic salmon (S. salar) and sea trout (Oncorhynchus
truttu) followed by rainbow trout (Oncorhynchus mykiss),
chinook (Oncorhynchus tschawytscha), and coho salmon
(Oncorhynchus kisutch; Dawson et al. 1997; Fast et al.
2002; Johnson and Albright 1992). In contrast, C. roger-
cresseyi occurs in higher numbers on caged rainbow trout
compared with Atlantic or coho salmon (González et al.
2000). Thus, while L. salmonis and Caligus species exhibit
similar parasitic life history strategies, they display consid-
erable differences in morphology, life cycle, and host range.
Another parasite, Lernaeocera branchialis belongs to the
copepod family Pennellidae and is distantly related to
the caligid copepods, and this species is commonly found
on gadoids, particularly Atlantic cod (Gadus morhua)
and haddock (Melanogrammus aeglefinus) in the North Atlantic
Ocean and North Sea (Bricknell et al. 2006; Smith et al.
2007). This parasite has a negative impact on wild gadoids
and is a potentially serious pathogen of farmed Atlantic cod
(Smith et al. 2007). A compilation of genomic information
on parasitic copepods is an important tool for understanding
their biology as well as for the study of population genetics
and control strategies.
In this study, we report on over 150,000 expressed
sequence tags (ESTs) obtained from Pacific L. salmonis
(49,672 new ESTs in addition to 14,994 previously reported
ESTs), Atlantic L. salmonis (57,349 ESTs), C. clemensi
(14,821 ESTs), C. rogercresseyi (32,135 ESTs), and L.
branchialis (16,441 ESTs). These ESTs were assembled
into complete or partial genes and annotated by compar-
isons to known proteins in public databases. In addition,
whole mitochondrial (mt) genome sequences of two
Caligus species, C. clemensi and C. rogercresseyi, were
determined and compared to each other and to L. salmonis.
These studies show high levels of sequence divergence in
nuclear and mtDNA genes. This report describes the
production and characteristics of the largest genomic
resource for copepods.

Materials and Methods

EST Analysis

Specimens belonging to the Pacific (British Columbia,
Canada (BC)) and Atlantic forms of L. salmonis (Norway
and New Brunswick, Canada), C. clemensi (BC), C.
rogercresseyi (Chile), and L. branchialis (Scotland, UK)
were collected and stored at −80°C or in RNAlater (Invitrogen) until RNA extraction. Total RNA was extracted from whole bodies (from various life stages and both sexes) using TRIZol reagent (Invitrogen) and spin-column purified using RNaseasy Mini kits (Qiagen). The purified RNAs were then quantified and quality checked by spectrophotometer (NanoDrop Technologies) and agarose gel, respectively. Approximately 1.0–3.0 μg of total RNA was converted into cDNA and normalized and was directionally cloned into pAL 17.3 vector (Evrogen Co.).

Clones from each library were robotically arrayed in 384-well microtiter plates as detailed previously (Koop et al. 2008). Plasmid DNAs were extracted and sequenced on an ABI 3730 DNA analyzer (Applied Biosystems) with M13 forward and M13 reverse primers (L. salmonis and C. rogercresseyi) or with M13 forward and SP6 primers (C. clemensi and L. branchialis). These sequence primers are shown in supplemental Table 1. The resulting ESTs were assembled with CAP3 (Huang and Madan 1999) with default parameters. The assembled total contigs (clusters + singletons) were annotated using RPS-BLAST and BLASTX comparisons with the Conserved Domain Database (CDD) and SwissProt (Bairoch and Apweiler 1996), respectively. The best BLAST match (E value threshold of 1 E −10) was used to identify contigs. Contigs that did not meet this threshold were annotated as “unknown.”

Table 1  Sea lice EST project summary

|                                      | L. salmonis (P) \(^a\) | L. salmonis (A) \(^b\) | C. clemensi | C. rogercresseyi | L. branchialis |
|--------------------------------------|----------------------|----------------------|-------------|-----------------|---------------|
| Number of clones \(^c\)              | 38,880 \(^e\)        | 51,607               | 7,680       | 19,200          | 8,448         |
| Number of sequences \(^d\)           | 64,666 \(^e\)        | 5734.9 \(^i\)       | 14,821      | 32,135          | 16,441        |
| Average trimmed EST length (bp) \(^f\) | 756                  | 644                  | 790         | 730             | 749           |
| Number of contigs \(^g\)             | 11,922               | 9,113                | 4,392       | 8,251           | 4,239         |
| Number of singletons                 | 4,186                | 5,145                | 1,662       | 3,106           | 2,199         |
| Number of putative transcripts       | 16,108               | 14,466               | 6,054       | 11,357          | 6,438         |
| Maximum contig size (no. of ESTs)    | 554                  | 1482                 | 15          | 34              | 21            |
| Average contig size (no. of ESTs)    | 4.0                  | 4.0                  | 2.5         | 2.8             | 2.6           |
| Number of transcripts with BLAST hits \(^h\) | 7,157 | 6,726                   | 3,775       | 5,830           | 3,951         |
| Percent with significant BLAST hits  | 44.4%                | 46.5%                | 62.4%       | 51.3%           | 61.4%         |

\(^a\) L. salmonis Pacific form

\(^b\) L. salmonis Atlantic (Canada, Norway) form

\(^c\) Number of clones which from at least one sequence (5' or 3') was obtained

\(^d\) Number of 5' and 3' EST sequences obtained

\(^e\) Twenty-eight thousand thirty-two clones and 49,672 sequences were obtained from this study, while 5,760 clones and 14,994 sequences were previously reported (Yazawa et al. 2008)

\(^f\) Vector, low quality, and contaminating bacterial sequences are trimmed

\(^g\) A contig (contiguous sequence) contains two or more ESTs

\(^h\) Number of transcripts that have a RPS-BLAST or BLASTX hit of less than 1 E −10 to the Conserved Domain Database (CDD) or SwissProt databases

Reference full-length cDNAs (FLcDNAs) were identified as detailed previously (Leong et al. 2010). A single clone containing an entire coding sequence (CDS) for a gene product is considered a reference FLcDNA.

Complete Mitochondrial Genome Sequences of C. clemensi and C. rogercresseyi

The total genomic DNAs were extracted from an adult male C. clemensi and C. rogercresseyi as previously described (Yazawa et al. 2008). A sample placed in 5% Chelex-100 resin (Sigma) solution (5% Chelex-100 resin, 0.2% SDS in TE, with proteinase K (100 μg/ml)) was incubated for 30 min at 55°C, and the proteinase K was then inactivated for 10 min at 90°C. The sequence determination of the complete C. rogercresseyi mt genome was carried out as previously described (Yazawa et al. 2008). The PCR primer sets that were used were designed for 15 fragments (Supplemental Table 1) based on the EST sequences encoding mtDNA. PCR amplification was performed using 1.0 μl of extracted total genomic DNA of C. rogercresseyi with an initial denaturation step of 2 min at 95°C and then 30 cycles as follows: 30 s of denaturation at 95°C, 30 s of annealing at 55°C, and 3 min of extension at 72°C. PCR products were cloned into pCR2.1 vector (TA Cloning Kit, Invitrogen) with the manufacturer's protocol, and each positive PCR product was sequenced as described above.
The entire mt genome for *C. clemensi* was amplified by a long PCR method for three long fragments (5.4, 5.0, and 3.0 kb) and by PCR as described above for one short fragment (0.8 kb). The three PCR fragments were amplified using the PCR primer sets shown in Supplemental Table 1 and by using Long PCR Enzyme mix (Fermentas) following the manufacturer’s protocol. The long PCR amplification was performed using 100 ng of extracted total genomic DNA of *C. clemensi* with an initial denaturation step of 2 min at 94°C and then a two-step PCR procedure (40 cycles of 95°C for 10 s and 68°C for 2 min at 94°C and then a two-step denaturation step of 2 min at 94°C and then a two-step extension. The three long PCR products were cloned into pCR-XL-TOPO vector (Invitrogen) with an initial step of 2 min at 94°C and then a two-step PCR procedure (40 cycles of 95°C for 10 s and 68°C for 7 min), and 10 min of final extension. The three long PCR products were sequenced by primer walking (supplemental Table 1). The one short fragment was cloned into pCR2.1 vector and sequenced as described above.

Protein-coding and rRNA genes of *C. clemensi* and *C. rogercresseyi* were identified by alignment with the Pacific *L. salmonis* mt gene sequences (GenBank: EU288200). The majority of the tRNA genes was identified using tRNAscan-SE 1.21 (Lowe and Eddy 1997), using the same parameters as described by Tjensvoll et al. (2005). The remaining tRNA genes were identified based on the sequence homology with *L. salmonis* tRNA sequences.

Pair-wise Kimura two-parameter (K2P) distances (Kimura 1980) of 16S rRNA and *cox1* sequence homology with *C. rogercresseyi*, *C. clemensi*, and *L. branchialis* were calculated in MEGA5 (Tamura et al. 2007), with default settings.

**Results and Discussion**

**EST Analysis and Comparison of the Nuclear Genes**

Normalized cDNA libraries were constructed for Pacific *L. salmonis*, Atlantic *L. salmonis*, *C. clemensi*, *C. rogercresseyi*, and *L. branchialis*. The 114,967 clones obtained from these cDNA libraries (28,032 Pacific *L. salmonis*, 51,607 Atlantic *L. salmonis*, 7,680 *C. clemensi*, 19,200 *C. rogercresseyi*, and 8,448 *L. branchialis*) were sequenced with M13 forward and M13 reverse (*L. salmonis* and *C. rogercresseyi*) or with M13 forward and SP6 primers (*C. clemensi* and *L. branchialis*). A summary of the EST project is shown in Table 1. From these clones, 153,977 high-quality ESTs were obtained from Pacific *L. salmonis* (49,672 ESTs), Atlantic *L. salmonis* (57,349 ESTs), *C. clemensi* (14,821 ESTs), *C. rogercresseyi* (32,135 ESTs), and *L. branchialis* (16,441 ESTs). The average trimmed length of these ESTs was 734 bp. These EST sequences are available in GenBank.

The 49,672 Pacific *L. salmonis* ESTs obtained in this study along with 14,994 Pacific *L. salmonis* ESTs from our previous study (Yazawa et al. 2008) were assembled into 11,922 contigs and 4,186 singletons (16,108 putative transcripts). There is a total of 14,466 putative transcripts for Atlantic *L. salmonis*, 6,054 for *C. clemensi*, 11,357 for *C. rogercresseyi*, and 6,438 for *L. branchialis*. These putative transcripts were annotated using RPS-BLAST and BLASTX comparisons with the CDD and SwissProt (Bairoch and Apweiler 1996), respectively. The best match (E value threshold of 1 E−10) was used to identify putative transcripts. Of the 16,108 Pacific *L. salmonis* putative transcripts, 7,157 (44.4%) matched at least one entry in the databases while the others remain unidentified. Similarly, 6,726 (46.5%) Atlantic *L. salmonis*, 3,775 (62.4%) *C. clemensi*, 5,830 (51.3%) *C. rogercresseyi*, and 3,951 (61.4%) *L. branchialis* putative transcripts have significant BLAST hits (Table 1).

A collection of reference FLcDNA clones is an important resource for identifying genes, determining their structural features and for experimental analysis of gene functions. Possible reference FLcDNAs were defined as having an entire open reading frame (ORF) corresponding to a full-length protein and were identified as described previously (Leong et al. 2010). Using an E value filter of E ≤10−5, the top ten SwissProt high-scoring segment pairs (HSPs) from BLASTX for each putative transcript were analyzed in succession to identify the correct ORF. Of the 16,108 Pacific *L. salmonis* putative transcripts, 1,435 transcripts were identified as possible FLcDNAs. There are 1,086 Atlantic *L. salmonis* FLcDNAs, 1,223 *C. clemensi* FLcDNAs, and 1,574 *C. rogercresseyi* FLcDNAs. These reference FLcDNAs were submitted to NCBI’s FLIC database.

A relational database with an intuitive web interface was developed to process and display the large quantities of EST data, their assemblies, and their associated annotation information (Fig. 1). This interface provides the ability to search using sequence data, identifiers, accession numbers, and descriptive keywords. The BLAST search allows users to perform homology searches with sequences of interest, identifying potential transcripts names, and then visualizing these sequences and EST alignments. These EST contigs have predicted ORFs and BLASTX HSPs displayed in a single view. This database contributes to the identification and analysis of proteins and to the development of microarrays for gene expression analyses.

![Fig. 1](https://example.com/fig1.png)
Sequence similarities and putative transcripts were compared among the nuclear genes of the five copepods (Pacific *L. salmonis*, Atlantic *L. salmonis*, *C. clemensi*, *C. rogercresseyi*, and *L. branchialis*) by BLASTN for nucleotide (nt) sequences and tBLASTX for amino acid (aa) sequences (Table 2). We previously reported that a total of 155 nuclear genes from Pacific and Atlantic *L. salmonis* showed an average of 96.8% nt identity over an average of 756 bp (Yazawa et al. 2008). In this study, a total of 8,121 nucleotide and 8,827 translated aa sequences matched between the Pacific and Atlantic *L. salmonis* putative transcripts. These sequences showed an average of 96% identity at the nt level over an average of 626 bp and 88% at the aa level over an average of 187 aa (Table 2). Nuclear gene sequences were quite different not only between the genera *Caligus* and *Lepeophtheirus* (81–82% nt, 70–72% aa identities), but also between the two *Caligus* species (83% nt, 71% aa identities; Table 2). The range of nuclear gene sequence divergence was quite similar among these species (17–19% nt and 28–30% aa sequence divergences). As expected, nucleotide sequences of *L. branchialis*, the only species examined from the family Pennellidae, were very different from the caligid sequences: only 4–6% of the total queries (254–405 sequences) matched the nuclear genes of the four other copepods. We speculate that the matched genes are conserved among copepods and therefore we could not determine the divergence between nt sequences of *L. branchialis* and the four caligid copepods. However, the 2,634–3,375 translated aa sequences of *L. branchialis* (44–52% of query sequences) did show significant matches with sequences of the four other copepods. These translated aa sequences showed 59–62% identities over averages of 121–132 aa (Table 2). Although these comparisons provide only a very rough estimate of overall sequence similarity, they clearly indicate a high level of sequence divergence among these copepods nuclear genes.

Mitochondrial Genome Sequences of *L. salmonis*, *C. clemensi*, and *C. rogercresseyi*

Metazoan mt genomes typically range between 15 and 20 kb in size, containing 37 genes: 13 protein-encoding genes (PCGs), 22 transfer RNA (tRNA) genes, two ribosomal RNA (rRNA) genes and a major non-coding region (NCR; Boore 1999). In this study, whole mt genome sequences of two *Caligus* species, *C. clemensi* and *C. rogercresseyi*, were determined. The sizes of the entire mt genomes were 13,440 bp for *C. clemensi* [Genbank: HQ157566] and 13,468 bp for *C. rogercresseyi* [Genbank: HQ157565], and thus, these mt genomes are the shortest among 57 crustacean mt genomes (average length: 15,785 bp) reported so far (Genbank: November 2010). There are two reasons for the small size of these mt genomes. First, the major NCRs of the *C. clemensi* (104 bp) and *C. rogercresseyi* (129 bp) mt genomes were much shorter than that of *L. salmonis* (Pacific form, 1,441 bp; Atlantic form, 2,146 bp) and that of other crustaceans (average length, 875 bp), except for that of the amphipod *Metacrangonyx longipes* (76 bp; Bauzà-Ribot et al. 2009). Second, while both *Caligus* mt genomes contained the typical set of 12 protein-encoding, 21 tRNA and two rRNA genes found in other animal mt genomes, both mt genomes lacked the PCG, *nad4L*, and a tRNA gene, *trnL* (CUN).

Interestingly, the *C. clemensi* mt genome is adenine and thymine (A + T)-rich (PCG, 74.5%; whole genome, 75.6%) compared to *C. rogercresseyi* and *L. salmonis* (PCG, 63.6–64.9%; whole genome, 65.2–66.5%; Supplemental Table 2). In crustaceans, the mt genomic A–T content values range from 60.9% for *Ligia oceanica* (Isopoda; Kilpert and Podsiadlowski 2006) to 77.8% for *Argulus americanus* (Branchiura; Lavrov et al. 2004). The reason for the variability in A–T richness within the mitochondrial genome among taxa is not clear.

Like the nuclear genes, the mtDNA gene sequences also exhibited large divergence, not only between *L. salmonis* and the two *Caligus* species (66.7–68.8% nt and 64.2–65.4% aa identities), but also between the two *Caligus* species (68.8% nt and 63.6% aa identities). The range of mtDNA sequence divergence was quite similar among the three caligid copepods. The percent nt and aa identities among the *L. salmonis*, *C. clemensi*, and *C. rogercresseyi* sequences are 63.6–68.8% (Table 3). The *cox1* gene is the most conserved PCG among the three mt genomes (79.1–82.6% nt and 91.2–94.1% aa identities), while *nad2*, *nad4*, *nad5*, and *nad6* exhibit a large sequence divergence (56.1–62.2% nt and 40.0–51.9% aa identities; Table 3).

Hebert et al. (2003) reported that *cox1* divergences among the 13,320 species in the animal kingdom ranged from a low of 0.0% to a high of 53.7% and the mean divergence value of 11.3%. The *cox1* divergences in the Crustacea showed the mean species divergence value of 15.4% (Hebert et al. 2003). Interestingly, our present study showed that the *cox1* divergences among the three caligid copepods were higher than the mean divergence value of Crustacea. The *cox1* interspecific divergence between *C. clemensi* and *C. rogercresseyi* is 20.2% and between the genera *Caligus* and *Lepeophtheirus* 26.0%. Øines and Schram (2008) compared among the *cox* fragment (a total 504 aligned base pairs) of 18 caligid copepods and the 16S rRNA fragment (a total of 438 aligned base pairs) of 11 caligid copepods. They found that an average K2P distance of *cox1* were 0.218 and those of 16S rRNA were 0.221 (Øines and Schram 2008). In the present study, the K2P distance of *cox1* (a total of 1,539 aligned base pairs) among the *L. salmonis*, *C. clemensi*, and *C. rogercresseyi* is 0.202–...
| Type                  | No. of queries | No. of matches | Percentage with match<sup>a</sup> | Average | Maximum | Standard deviation for length | Average identities | Maximum identities | Minimum identities | Standard deviation for% identities | Average positive AAs |
|----------------------|----------------|----------------|----------------------------------|---------|---------|--------------------------------|--------------------|-------------------|-------------------|-------------------------------|--------------------|
| Atlantic form L. salmonis vs Pacific form L. salmonis blastn | 14,466         | 8,121          | 56%                              | 626 bp  | 2,891 bp | 365.25                          | 96%                | 100%              | 78%                | 4.41                          | –                  |
| tblastx              | 14,466         | 8,827          | 61%                              | 187 aa  | 820 aa   | 110.46                          | 88%                | 100%              | 22%                | 18.15                         | 91%                |
| C. clemensi vs Pacific form L. salmonis blastn              | 6,054          | 1,598          | 26%                              | 327 bp  | 1,316 bp | 208.89                          | 81%                | 98%               | 76%                | 2.83                          | –                  |
| tblastx              | 6,054          | 3,852          | 64%                              | 151 aa  | 569 aa   | 79.38                           | 72%                | 100%              | 12%                | 17.09                         | 83%                |
| C. clemensi vs Atlantic form L. salmonis blastn              | 6,054          | 1,595          | 26%                              | 318 bp  | 1,316 bp | 197.67                          | 81%                | 98%               | 76%                | 2.98                          | –                  |
| tblastx              | 6,054          | 4,906          | 68%                              | 145 aa  | 539 aa   | 75.68                           | 72%                | 100%              | 16%                | 16.65                         | 83%                |
| C. clemensi vs C. rogercressyi blastn               | 6,054          | 1,893          | 31%                              | 338 bp  | 1,305 bp | 202.55                          | 83%                | 100%              | 77%                | 3.11                          | –                  |
| tblastx              | 6,054          | 3,715          | 61%                              | 142 aa  | 456 aa   | 71.72                           | 71%                | 100%              | 21%                | 17.09                         | 82%                |
| C. clemensi vs L. branchialis blastn               | 6,054          | 257           | 4%                               | 278 bp  | 1,226 bp | 186.83                          | 81%                | 97%               | 77%                | 3.01                          | –                  |
| tblastx              | 6,054          | 2,634          | 44%                              | 125 aa  | 436 aa   | 63.27                           | 59%                | 100%              | 21%                | 16.47                         | 74%                |
| C. rogercressyi vs Pacific form L. salmonis blastn           | 11,357         | 1,931          | 17%                              | 301 bp  | 1,309 bp | 178.46                          | 82%                | 99%               | 76%                | 2.86                          | –                  |
| tblastx              | 11,357         | 5,937          | 52%                              | 139 aa  | 557 aa   | 72.40                           | 70%                | 100%              | 18%                | 17.45                         | 81%                |
| C. rogercressyi vs Atlantic form L. salmonis blastn           | 11,357         | 1,973          | 17%                              | 292 bp  | 1,498 bp | 175.48                          | 82%                | 100%              | 76%                | 3.40                          | –                  |
| tblastx              | 11,357         | 6,383          | 56%                              | 133 aa  | 521 aa   | 68.41                           | 70%                | 100%              | 21%                | 16.81                         | 82%                |
| L. branchialis vs Pacific form L. salmonis blastn             | 6,438          | 417           | 6%                               | 264 bp  | 1,079 bp | 168.51                          | 81%                | 99%               | 76%                | 2.99                          | –                  |
| tblastx              | 6,438          | 3,284          | 51%                              | 132 aa  | 466 aa   | 69.24                           | 62%                | 100%              | 22%                | 16.22                         | 77%                |
| L. branchialis vs Atlantic form L. salmonis blastn             | 6,438          | 405           | 6%                               | 260 bp  | 1,079 bp | 163.04                          | 81%                | 98%               | 77%                | 2.75                          | –                  |
| tblastx              | 6,438          | 3,375          | 52%                              | 126 aa  | 466 aa   | 64.90                           | 62%                | 100%              | 22%                | 16.31                         | 76%                |
| L. branchialis vs C. rogercressyi blastn                      | 6,438          | 254           | 4%                               | 250 bp  | 1,211 bp | 156.59                          | 81%                | 99%               | 76%                | 2.73                          | –                  |
| tblastx              | 6,438          | 3,021          | 47%                              | 121 aa  | 465 aa   | 59.81                           | 59%                | 100%              | 17%                | 16.38                         | 74%                |

<sup>a</sup>The number of queries that had BLASTN hit with an E value <1 E<sup>−10</sup> and 100 bp minimum of alignment length or that had tBLASTX with an E value <1 E<sup>−10</sup> and 50 aa minimum of alignment length

<sup>b</sup>First match that conformed to parameters was taken from the top five hits of blast output. If no suitable match was found in the top five hits, it was not included in the results
Table 3 Comparison of the *L. salmonis*, *C. clemensi*, and *C. rogercressyi* mtDNA genes

| Genes        | In nucleic sequence (%) | In deduced amino acid sequence (%) |
|--------------|-------------------------|-----------------------------------|
|              | Pacific form *L. salmonis* vs *C. clemensi* | Pacific form *L. salmonis* vs *C. rogercressyi* | Pacific form *L. salmonis* vs *C. clemensi* | Pacific form *L. salmonis* vs *C. rogercressyi* | Atlantic form *L. salmonis* vs Pacific form *L. salmonis* |
| rmS          | 77.2                    | 76.4                               | 74.9                               | 98.8                                      |
| rmL          | 68.3                    | 67.2                               | 71.2                               | 96.9                                      |
| atp6a        | 72.0                    | 67.7                               | 72.0                               | 96.8                                      |
| atp6         | 63.9                    | 65.4                               | 65.3                               | 91.9                                      |
| cob          | 71.0                    | 70.8                               | 71.3                               | 93.7                                      |
| cox2         | 76.6                    | 75.7                               | 78.5                               | 93.5                                      |
| cox3         | 73.2                    | 71.7                               | 72.4                               | 92.0                                      |
| nadi         | 72.2                    | 71.6                               | 70.5                               | 92.8                                      |
| nad2         | 57.9                    | 57.8                               | 59.3                               | 90.9                                      |
| nad3         | 68.5                    | 57.8                               | 65.0                               | 91.6                                      |
| nad4         | 61.9                    | 58.8                               | 58.7                               | 91.2                                      |
| nad4Lb       | N.A.                    | N.A.                               | N.A.                               | 94.3                                      |
| nad5         | 62.2                    | 58.8                               | 61.8                               | 90.7                                      |
| nad6         | 59.6                    | 56.1                               | 59.1                               | 93.8                                      |
| Average      | 68.8                    | 66.7                               | 68.8                               | 93.5                                      |

*Comparisons of amino acid sequences of atp8 genes were not conducted because these sequences are very short in size (31 aa)

* nad4L genes are absent in the two Caligus species

0.270 (Supplemental Table 3), which is similar to an average K2P distance found by Øines and Schram (2008). However, the 16S rRNA among the three copepods showed a very high genetic distance. The K2P distance of the 16S rRNA (a total of 1,085 aligned base pairs) were 0.333 between *C. clemensi* and *C. rogercressyi* and 0.422 (Supplemental Table 3). These molecular distance values support an ancient separation between *C. clemensi* and *C. rogercressyi* as well as between *Lepeoptheirus* and *Caligus*.

In our previous study, a molecular clock based on 16S rRNA and calibrated by copepod data suggested that the forms of *L. salmonis* existing in the Pacific and Atlantic Oceans evolved from a common ancestor following a separation that occurred from 4.6–11 million years ago (Yazawa et al. 2008). In this study, the molecular estimates of the age of divergence between the *L. salmonis* (Pacific) and the two *Caligus* species were calculated based on the 16S rRNA gene using the same method as previously reported (Yazawa et al. 2008). The results suggest that the separation between the *L. salmonis* (Pacific) and the two *Caligus* species occurred approximately 45–113 million years ago (Table 4). In addition, the separation between the two *Caligus* species was estimated to have occurred 37–87 million years ago (Table 4). Salmonids are believed to have evolved from an ancestor in which a whole genome duplication event occurred 25–100 million years ago (Ohno 1970). Thus, our present results suggest that the *L. salmonis* and *C. clemensi* have been in existence for 45–106 million years and that parasitic association with salmonids is likely also quite ancient (Table 4).

The order of the genes in the two *Caligus* mt genomes is identical despite extensive sequence divergence. In contrast, the order of genes in the two *Caligus* mt genomes is quite different from that in the *L. salmonis* mt genome. The gene arrangement in the region between nad4 and trnL1 (UUR; approximately 10 kb) is well conserved between *L. salmonis* and the *Caligus* species. However, the gene arrangements adjacent to their control regions (CRs) are very distinct, and the *Caligus* mt genomes show a novel gene arrangement (Fig. 2). The region around the CR is more prone to gene rearrangement in both vertebrate (Macey et al. 1997) and invertebrate (Dowton and Austin 1999) mt genomes. In the *L. salmonis* mt genomes, this region is comprised of six rRNA and atp6 genes) is in a row (Tjensvoll et al. 2005; Yazawa et al. 2008). However, in the *Caligus* mt genomes, this region is separated by rrs-nad6-trnA-trnK2-trnQ-trnT-cytb-CR, and divided into trnK2-trnN-trnG-trnV and atp6-trnY-trnR (trnY also had a position change; Fig. 2). As mentioned above, the nad4L and trnL2 (CUN) genes are absent in the *Caligus* mt genomes. These two genes normally reside in this region.
and have probably been lost due to rearrangement. It is likely that this rearrangement event also has led to the trimming of their CRs in the two *Caligus* mt genomes.

In the mt genomes of most animals, nad4L and atp8 are located together with nad4 and atp6, respectively (nad4L-nad4 and atp8-atp6), and nad4L-nad4 and atp8-atp6 are translated from a single mRNA (Amalric et al. 1978; Berthier et al. 1986). In contrast, several genes separate nad4 and nad4L in the mt genomes of *L. salmonis* and in the mt genomes of all copepods characterized so far: *Tigriopus japonicas* (Machida et al. 2002), *Tigriopus californicus* (Burton et al. 2007), *Paracyclopina nana* (Ki et al. 2009), and the partially sequenced mt genomes of *Eucalanus bungii* and *Neocalanus cristatus* (Machida et al. 2004). The atp6 and atp8 are also separated in the two *Caligus* species and in *L. salmonis* (Fig. 2). In addition, it has been reported that atp8 is absent in the mt genome of *P. nana* (Ki et al. 2009). Thus, it is most likely that these separations of nad4-nad4L and atp6-atp8 occurred during copepod evolution and led to the loss of nad4L in the two *Caligus* species and to the loss of atp8 in the *P. nana*.

In summary, the mtDNA genes of the two *Caligus* species showed high levels of sequence divergence (Table 3). The A+T content is also quite different between

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**Table 4** Ranges of 16S rRNA gene divergence based on Kimura two-parameter distance and crustacean molecular clock calibrations

| Distance (K2P) | Divergence Range (Myr) |
|---------------|------------------------|
|               | Ano | Fid | Gra (low) | Gra (high) |
| Pacific form *L. salmonis* vs. *C. clemensi* | 0.405 | 106.2 | 45.0 | 62.3 | 46.0 |
| Pacific form *L. salmonis* vs. *C. rogercressyi* | 0.431 | 113.0 | 47.8 | 66.2 | 48.9 |
| *C. clemensi* vs. *C. rogercressyi* | 0.333 | 87.4 | 37.0 | 51.2 | 37.8 |

The values for “Distance” are the Kimura two-parameter (K2P) distance between the species. Rates of molecular evolution used for the 16S rRNA gene include 0.38% K2P/million year (Myr) for anomurans (Ano; Cunningham et al. 1992), 0.90% K2P/Myr for fiddler crabs (Fid; Sturmbauer et al. 1996), and 0.65 (low)–0.88% (high) K2P/Myr obtained from grapsid crabs (Gra; Schubart et al. 1998).

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**Fig. 2** Genomic organization of the *C. clemensi* (13.440 bp) and the *C. rogercressyi* (13.468 bp) mt genomes. The complete mt genomes of the Atlantic (15,445 bp) and Pacific (16,148 bp) *L. salmonis* were previously reported, and these mt genomes are identical in gene organization (Tjensvoll et al. 2005; Yazawa et al. 2008). Boxes represent mtDNA genes. tRNA genes are denoted by the single letter amino acid code, and an underline indicates tRNA genes located on negative strand. *rrnL* and *rrnS* refer to 16S and 12S rRNA; *cox1*, *cox2*, and *cox3* refer to cytochrome oxidase subunits I, II, and III; *cob* refers to cytochrome b; *nad1–6* and *nad4L* refer to NADH dehydrogenase subunits 1–6 and 4 L, *atp6* and *atp8* refer to ATP synthase subunits 6 and 8, respectively, and *CR* refers to control region. Transcription directions for the protein-coding and tRNA genes are shown by arrowheads.
the two Caligus mt genomes (Supplemental Table 2). In addition, the orders of the genes in the two Caligus mt genomes are identical to each other, but different from the order in the L. salmonis mt genome (Fig. 2).

Sea Lice as Ectoparasite Model System

Since parasites by definition depend on a live host for growth and survival, in vitro culture system is typically very difficult to establish. Although procedures for experimental infections are established for some parasitic species, manipulation of the parasites may still be very difficult since removing them from the host is lethal for the parasite in general. Sea lice have life cycle features that make them promising as a model system. The life cycle features, consisting of both free-living larval developmental stages and pre-adults and adult stages that can move unrestricted on host surface, enable manipulation of these parasites. For L. salmonis, recent advances in larval production systems and infection procedures (see Hamre et al. 2009) have been crucial for the establishment of defined laboratory strains of the salmon louse with different properties (e.g., drug-resistant strains, inbred strains). Stable and predictable production conditions further enables specific breeding to create various types of hybrids (e.g., susceptible and drug-resistant family groups). The improvement of rearing facilities has been a crucial facilitator for establishment of RNAi in L. salmonis (Dalvin et al. 2009).

Systemic RNAi is easily achieved in pre-adult or adult lice by injection of dsRNA in the animal. In addition, soaking free-living larval stages (e.g., copepodids) in dsRNA enables RNAi in copepodids (Campell et al. 2009). In addition, the genomes of both the Pacific and Atlantic variants of L. salmonis are currently being sequenced and together with the present cDNA resources this will open up for a new avenue in sea lice research. There is a wide diversity of arthropod parasites and good experimental parasite model systems are scarce, and we anticipate that experimental studies on salmon louse and other sea lice species will contribute to increase our knowledge about ectoparasites in general, particularly when more parasite genomes become available.

Conclusions

We sequenced over 150,000 ESTs from Pacific L. salmonis (49,672 new ESTs in addition to 14,994 previously reported ESTs), Atlantic L. salmonis (57,349 ESTs), C. clemensi (14,821 ESTs), C. rogercresseyi (32,135 ESTs), and L. branchialis (16,441 ESTs; Table 1). A relational database with an intuitive web interface was developed to process and display the large quantities of EST data, their assemblies and associated annotation information, as well as possible full-length gene information (Fig. 1). This database provides a novel resource for the study of sea louse biology, population genetics, and control strategies. This genomic resource represents the largest compilation of any copepod species and provides the material basis for the development of a 38K microarray that can be used in conjunction with our existing salmon 44K microarray to study host–parasite interactions at the molecular level.

The nuclear genes showed a high level of sequence divergence among the caligid copepods examined: L. salmonis, C. clemensi, C. rogercresseyi, and L. branchialis (Table 2). In addition, whole mt genome sequences of two Caligus species, C. clemensi (13,440 bp) and C. rogercresseyi (13,468 bp), were determined and compared. The L. salmonis, C. clemensi, and C. rogercresseyi mtDNA genes also exhibited extensive sequence divergence, ranging among these species from 66.7 to 68.8% nt and from 63.6% to 65.4% aa identities (Table 3). Both nuclear and mtDNA genes showed very high levels of sequence divergence between these ectoparasitic copepods which suggested that they have been in existence for 37–113 million years and that parasitic association with marine organisms is likely also quite ancient. However, while the order of the genes in the two Caligus mt genomes is the same, they are different from L. salmonis (Fig. 2). The large sequence divergence observed among these copepods may help to explain an extensive variety of morphology, life history, and host association in copepods.

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