Heavy and light chain homologs of ferritin are essential for blood-feeding and egg production of the ectoparasitic copepod *Lepeophtheirus salmonis*

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**A B S T R A C T**

The salmon louse, *Lepeophtheirus salmonis*, is a hematophagous ectoparasite of salmonid fish. Due to its blood-feeding activity, the louse is exposed to great amounts of iron, which is an essential, yet potentially toxic mineral. The major known iron storage protein is ferritin, which the salmon louse encodes four genes of (*LsFer1*-4). Two of the ferritins are predicted to be secreted. These are one of the heavy chain homologs (*LsFer1*) and the light chain homolog (*LsFer2*). Here, we perform functional studies and characterize the two secreted ferritins. Our results show that knocking down *LsFer1* and *LsFer2* both negatively affect the parasite's physiology, as it is not able to properly feed and reproduce. In a starvation experiment, the transcript levels of both *LsFer1* and *LsFer2* decrease during the starvation period. Combined, these results demonstrate the importance of these genes for the normal parasite biology, and they could thus potentially be targets for pest management.

1. Introduction

Iron is an essential element necessary for many cellular processes in all animals. Iron-containing proteins are present in several key biochemical pathways, such as DNA synthesis, energy metabolism and electron transfer reactions [1]. Free intracellular iron can rapidly be reduced to the ferrous (Fe2+) form, and may catalyze the generation of reactive oxygen species (ROS) which are highly cytotoxic reactive radicals that can cause harm to cellular compounds such as lipids, DNA and proteins [2]. In order to avoid cell harm, iron storage and distribution must be implemented by tightly regulated mechanisms, using designated proteins in the cell.

Ferritin is the major intracellular iron storage protein. Here, ferric iron (Fe3+) is sequestered and kept in a nontoxic, soluble and biologically non-reactive state [3]. In that state, the iron can be recruited to various proteins that require the metal as cofactor to perform their proper biological function. Typically, ferritin is composed of 24 subunits which can harbor up to 4000 Fe atoms [4]. Vertebrate ferritins are made up of the heavy (H) chain and the light (L) chain subunits, whereas arthropod ferritins are made up of homologs to these (heavy-chain homolog, HCH, and light-chain homolog, LCH) which are highly cytotoxic reactive radicals that can cause harm to cellular compounds such as lipids, DNA and proteins [2]. In order to avoid cell harm, iron storage and distribution must be implemented by tightly regulated mechanisms, using designated proteins in the cell.

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With the exception of intracellular iron within iron storage proteins, the majority of vertebrate iron is found in blood, complexed within the porphyrin ring of heme as a cofactor of hemoglobin used for gas transportation, but also in other forms such as bound to the iron transfer protein transferrin [7]. Blood-feeding parasites ingest high levels of iron and thus need to protect their cells from iron overload upon feeding and absorption. The salmon louse, *Lepeophtheirus salmonis*, is an obligate marine ectoparasite, feeding off the skin and blood of its salmonid fish host [8]. With an increasing resistance towards various chemical treatments [9], the salmon louse is considered to be an economically and ecologically important parasite to combat. The parasite’s life cycle is divided into eight stages which are as follows: the planktonic nauplius I and II, the infectious copepodid, the attached parasitic chalimus I and II, and the mobile parasitic preadult I and II and adult lice [10,11]. At 10°C, the development of *L. salmonis* from fertilization to mature adult lice is completed in approximately 40 (male) to 52 (female) days [12]. Once the female louse has reached the mature adult
stage, it will continuously produce pairwise egg strings. The hatching larvae have the potential to further infect hosts upon reaching the copepodid stage. In order to be able to successfully produce viable offspring, the louse needs to absorb several macro- and micronutrients from its blood meal. One of the many micronutrients required for oocyte development is the mineral iron, which the salmon louse requires proper storage for until it is needed in iron-proteins.

Here, we report the presence of four ferritin homologs (LsFer1-4) in the genome of *L. salmonis* and analyze the sequences. Further, we perform functional studies of the two ferritins with signal peptides (LsFer1 and LsFer2) by in situ hybridization, starvation as well as RNA interference (RNAi) to evaluate the importance of these genes in reproduction and blood-feeding of the adult female salmon louse. Results from the knockdown study illustrate the detrimental effects for the hemaphagous parasite lacking LsFer1 or LsFer2, as normal development and fecundity is negatively impacted upon RNAi mediated knockdown of the genes.

2. Material and methods

2.1. Sequence analyses

Initial data for searching transcript and protein sequences and primer design were extracted from the *L. salmonis* genome annotation in Ensembl Metazoa (https://metazoa.ensembl.org/Lepeophtheirus_salmonis). All following analyses are based on the GenBank sequences: BT121711 (LsFer1), BT121232 (LsFer2), MK887318 (LsFer3), BT077723 (LsFer4a) and BT121164 (LsFer4b). Glycosylation prediction was performed using the NetNGlyc 1.0 Server (http://www.cbs.dtu.dk/services/NetNGlyc/) and the NetOGlyc 4.0 Server [13]. Protein BLAST searches were conducted against the Genbank and SwissProt databases [14]. Conserved domain search was conducted using InterProScan [15]. Signal peptides and cleavage sites in the aa sequences were predicted using Phobius [16]. Searches for iron-responsive elements (IREs) in the untranslated regions were performed using SIREs Web Server 2.0 [17].

2.2. Sequence alignment and phylogeny

Aa sequences of LsFer1-4 were used to obtain ferritins of various species from a BLASTp search [14] or from searching The Uniprot Consortium [18] for ferritins of model organisms. Alignment was performed in UGENE using MUSCLE alignment under default conditions [19,20]. Grey shading indicates percentage identity for alignment. For alignment, following sequences annotated with heavy chain were chosen: *Homo sapiens* (UniProt: P02794), *Salmo salar* (UniProt: P49946), *Xenopus laevis* (UniProt: P49948), *Caenorhabditis elegans* (NP_491198.1), *P. monodon* (AAQ54713.1), *H. sapiens* (BAN13552.1)*, Trichoplusia ni* (XP_026744300.1)*, *M. sexta* (AAK39636.1)*, Anopheles gambiae* (XP_124747.4)*. Light chain sequences chosen were: *H. sapiens* (UniProt: P02792), *X. laevis* (UniProt: Q7SKX5), *S. salar* (NP_001134896.1), *T. ni* (XP_026744299.1)*, M. sexta* (AAQ47171.1)*, *A. gambiae* (XP_001237468.3)*. Sequences with signal peptides are marked with *. For alignment however, the signal sequence was removed. From sequences without signal peptides, the start methionine was removed for alignment.

Sequences used for phylogenetic analysis were: *Caligus clemensi* (ACO15165.1 and ACO14799.1), *Caligus rogercresseyi* (ACO11534.1), *Drosophila melanogaster* (AAF07876.1, NP_524873.1 and NP_572854.1), *Aedes aegypti* (AAO41698.1 and AQY17412.1), *Ixodes ricinus* (ACJ70653.1), *Rhizophagus microplus* (UniProt: AA0334WXT0), *S. salar* (UniProt: P49947 and P49946, H. sapiens (UniProt: P02792 and P02794), *Mus musculus* (UniProt: P29391 and P09528), *C. elegans* (NP_504944.2 and NP_491198.1) and *H. longicornis* (BAN13552.1 and AAQ54713.1). The website http://www.phylogeny.fr was used to construct the phylogenetic tree with default settings [21].

2.3. Animals

*L. salmonis salmonis* was raised on Atlantic salmon (*S. salar*) under laboratory conditions in tanks with seawater (salinity 34.5 ppt and 10 °C) as described before [24]. Fish were daily handled commercial dry pellets and kept according to Norwegian animal welfare regulations. Experiments were approved by the governmental Norwegian Animal Research Authority (ID8589). Fish carrying lice from RNAi experiments were kept in single-fish tanks [25]. Fish were anesthetized with a mixture of methomidate (5 mg/l) and benzocaine (60 mg/l) upon samplings, or setup of salmon louse infections. Egg string pairs with larvae were incubated and hatched in single incubators in a seawater flow through system [24].

2.4. Ontogeny

The genomic location for the validated transcript sequences (RACE-PCR) were determined using the software GMAP [26] with the Atlantic salmon louse genome (*LsalAtl2s*, https://metazoa.ensembl.org/Lepeophtheirus_salmonis) and the raw reads of existing RNA-sequencing data were recounted by featureCounts [27] at these regions. Available RNA-sequencing data are used as found in LiceBase (http://licebase.org, Dondrup et al., in prep) and from data of a time-series previously published [28]. Data from LiceBase are bulk samples of different developmental stages (nauplius I, nauplius II, planktonic copepods, attached copepodids at three different sampling time points, chalimus I and II, female and male of preadult I and II and adult lice, as well as eggs (unfertilized extracted from the genital complex of the adult female, immature fertilized eggs (0–24 hours after extrusion) and more mature fertilized eggs (2–7 days after extrusion)). In addition, samples of different tissues (intestine, ovaries, testis feet and antenns) are shown. The data from the time series [28] are RNA-sequencing samples of attached parasitic stages (chalimus I and II and preadult I), which were divided into different instar ages (called young (directly after previous molt), middle and old (directly before next molt)) as well as planktonic copepodids. In addition, unpublished RNA-sequencing data (Eichner et al., unpublished) for the transcript regions from nauplius I and II, taken at three different time points (directly after hatching or molting respectively, in the middle of the stage, and before the next molt) as well as attached copepodids, sampled one, three and five days after infection, are included. RNA-sequencing and data processing for the nauplius and attached copepodid samples was done using the same protocols as in Eichner et al. [28]. Six biological replicates were sequenced for each time point. Normalization was done by edgeR [29] after combining all data sets using the same protocol as in Eichner et al. [28].

2.5. Isolating RNA, synthesis of cDNA, qRT-PCR and RACE

Tissues for RNA isolation were stored in RNAlater (Ambion), at 4 °C overnight and then at −20 °C until usage. Total RNA extraction from adult female lice was executed using TRI Reagent (Sigma-Aldrich) as instructed by the manufacturer. Samples were homogenized using 5 mm stainless steel beads and a TissueLyser II (Qiagen) for 1 min at 30 Hz. Genomic DNA digestion was done by DNase I after manufacturer’s instructions (Invitrogen). Quantification of RNA and purity check were done using NanoDrop ND-1000 UV–vis Spectrophotometer (NanoDrop Technologies). DNase-treated RNA was reverse transcribed using the AffinityScript QPCR cDNA Synthesis Kit (Stratagene). qRT-PCR was performed on a QuantStudio 3 qPCR machine using PowerUp SYBR Green Master Mix (Applied Biosytems) on duplicate samples.
Table 1

Primer names used for RACE, in situ hybridization (ISH), RNA interference (RNAi) and qPCR. T7 promoter extension is underlined. Primer efficiency (E%) is given for qPCR assays.

| Primer name | Sequence (5’→3’) | Application/E% |
|-------------|------------------|----------------|
| LsFer1 5′ RACE | TCCAGTCAACAGAGCTCTAG | 5′ RACE |
| LsFer1 3′ RACE | GCAAGATTATACGTCTCCATTGGG | 3′ RACE |
| LsFer2 5′ RACE | TTATGGTTGGCGGCTACTGAGTGAATGGG | 5′ RACE |
| LsFer2 3′ RACE | GGTGCCACCATGATGACATGGTACAGT | 3′ RACE |
| LsFer3 5′ RACE | CAGCGGTCAGCCAGAATACTCG | 5′ RACE |
| LsFer3 3′ RACE | AAACATTGCTGAACCTAAATCGG | 3′ RACE |
| LsFer4 5′ RACE | AGTATTTGGATCTTCAGTTGGG | 5′ RACE |
| LsFer4 3′ RACE | ATGATGGCCAAATCGGCT | 3′ RACE |
| LsFer1 T7 fwd | TAAATAGCTGTTATGCTCTGATG | qPCR |
| LsFer1 T7 rev | TAATAGCTGCTATAGGGTGTGTATGGG | qPCR |
| LsFer2 T7 fwd | GGTTGCCACCATGATGATTGG | qPCR |
| LsFer2 T7 rev | GGTTGCCACCATGATGATTGG | qPCR |
| LsFer1 rev | TTATGGTTGGCGGCTACTGAGTGAATGGG | qPCR |
| LsFer2 rev | GGTGCCACCATGATGATTGG | qPCR |
| Cod_CP185 T7 fwd | TAAATAGCTGTTATGCTCTGATG | qPCR |
| Cod_CP185 T7 rev | TAATAGCTGCTATAGGGTGTGTATGGG | qPCR |
| SYBR LsFer1 fwd | GTGTTCCACAGGCGTACTGTT | qPCR-93% |
| SYBR LsFer1 rev | GGGGCTGGTGGTATGGT | qPCR |
| SYBR LsFer2 fwd | GCGATTTTCTCTCTGAGGATGGG | qPCR-99% |
| SYBR LsFer2 rev | GCGATTTTCTCTCTGAGGATGGG | qPCR |

2.8. In situ hybridization

The localizations of LsFer1 and LsFer2 mRNAs were detected in the adult female salmon louse by using in situ hybridization (ISH) as earlier described [35, 36] with the DIG RNA Labelling Kit (Roche) on two sections per probe. RNA antisense probes of 568 bp (LsFer1) and 576 bp (LsFer2) were made from a target specific cDNA template (see Table 1 for primer sequences). Sense probes acted as negative control for transcript localizations.

2.9. Starvation

Adult female lice were removed from their host, the Atlantic salmon, and either sampled directly (zero days starved) in RNAlater or placed in individual incubators with running sea water [24]. The lice in the incubators were sampled one, two, four and eight days after being separated from their food source as described before [37]. RNA isolation and cDNA synthesis were performed as described in Section 2.5 on five replicates of each sampling time point. The transcript levels of LsFer1 and LsFer2 were finally assessed by qPCR.

2.10. Statistics

Statistics were performed by using IBM SPSS Statistics 23 for Windows. Data sets were first tested by Shapiro-Wilk’s test of normality and Levene’s test for equality of variances. An independent t-test was used to evaluate the mean between two groups. For non-normal data sets, a Mann-Whitney U test was conducted. A p-value ≤0.05 was considered statistically significant. Data are presented as mean values ± standard deviations (SD) or as individual values.

3. Results

3.1. Sequence analyses

cDNA sequences of ferritins LsFer1-4 in the salmon louse were obtained by 5′ and 3′ RACE using sequence specific primers (Table 1). Main results of sequence analyses are summarized in Table 2. No IREs were predicted for any of the genes. LsFer1 consists of an open reading frame (ORF) of 639 bp. A protein BLAST search of the L. salmonis full
length LsFer1 protein showed 78% identity with a Ferritin subunit precursor from *Caligus rogercresseyi* (Genbank accession: AC011534.1), 54% identity with Ferritin, lower subunit-like from *Eurytemora affinis* (Genbank accession: XP_023345228.1), and 40% identity with Ferritin 1-like protein *A from Daphnia pulex* (Genbank accession: ABK91577.1). LsFer2 consists of an ORF of 672 bp. A protein BLAST search of the *L. salmonis* full length LsFer2 protein showed 69% identity with a Ferritin light chain, oocyte isoform from *Caligus elegens* (Genbank accession: AC015165.1), and 29% identity with 32 kDa ferritin subunit from *Galleria mellonella* (Genbank accession: AA47694.1). The ORFs of LsFer1 and LsFer2 were analyzed in InterPro-Scan, and both proteins were characterized as containing a ferritin-like domain (PF00210). In LsFer1 there is a predicted ferroxidase site, a ferrihydrite nucleation center and an iron ion channel, which are not predicted in LsFer2. LsFer2 has a predicted dinuclear metal binding motif. A scan in Phobius shows that LsFer1 has a signal peptide from aa 1 → 21 with a cleavage site from aa 17 → 21, and a non-cytoplasmic region from aa 22 → 213. A Phobius scan for LsFer2 shows that the protein has a predicted signal peptide from aa 1 → 24 with a cleavage site from aa 17 → 24, and a non-cytoplasmic region from aa 25 → 224. LsFer1 has two predicted O-linked glycosylation sites at aa 29 and 36 and no predicted N-linked glycosylation sites. LsFer2 has no predicted O-linked glycosylation site, and one predicted N-linked glycosylation site at aa 55.

LsFer3 has an ORF of 540 bp. A protein blast shows a 59% identity with a ferritin of *Calanus sinicus* (Genbank accession: APG62655.1) and 58% identity with *Pseudodiaptomus annandalei* (Genbank accession: AGT28487.1). LsFer4a and b encode similar ORFs with only slight variations in their C-terminals affecting 3 aa only as well as 2 additional aa in LsFer4b. LsFer4a has an ORF of 513 bp whereas LsFer4b has an ORF of 519 bp. Both LsFer4a and 4b have 60% identity with a ferritin of *Calanus sinicus* (Genbank accession: APG62655.1) and a 58% identity with a ferritin heavy chain A-like of *Eurytemora affinis* (Genbank accession: XP_023344839.1). According to Phobius scans, LsFer3-4 lack signal peptides. InterPro-Scan predicts that LsFer3-4 have a ferritin-like domain (PF00210) and have ferroxidase activity, a ferrihydrite nucleation center and an iron ion channel.

### 3.2. Sequence alignment and phylogeny

Sequence alignment (Fig. 1) shows that LsFer1, 3, 4a, and 4b contain all conserved aa of the ferroxidase site. In addition, the conserved aspartic acid of the iron entry pore is found in all these *L. salmonis* ferritins, while the conserved glutamic acid is not found in LsFer1. Like the insect sequences shown in the alignment, it is replaced by threonine. There are only minor differences in the aa sequence between LsFer4a and LsFer4b, as the nucleotide sequence at the C-terminal end shows differences. LsFer2 contains only two of the conserved aa making up the functional ferritin motif. However, all of the conserved cysteins found in the light chain of insects responsible for disulfide bridges are also found in LsFer2. Two cysteine residues common in disulfide bridges are also conserved in LsFer1.

To investigate the relationship between LsFer1-4 and other vertebrate and invertebrate ferritins, a maximum likelihood phylogenetic analysis was performed (Fig. 2). The analysis shows that LsFer1 clusters together with secretory invertebrate HCH ferritins. LsFer2 clusters together with secretory invertebrate LCH ferritins. Both LsFer1 and LsFer2 have their closest relation to ferritin sequences of the parasitic copepod *C. clemensi*. LsFer3-4 cluster together in a separate clade, close with intracellular ferritins (HC and LC) of both vertebrates and invertebrates.

### 3.3. Ontogeny

To determine spatial and temporal expression patterns of the ferritin subunits, RNA-sequencing data were used to count the transcripts (Fig. 3). The two secreted ferritins LsFer1 (Fig. 3a) and LsFer2 (Fig. 3b) have similar absolute expression counts and pattern, and they are highly expressed in the intestine. LsFer3 (Fig. 3c) is nearly exclusively found in male lice and transcripts are highly expressed in testis. LsFer4a and LsFer4b are differing at the 3′ end only, and the total count of LsFer4 is shown (Fig. 3d). LsFer4 (Fig. 3d) is expressed in the intestine, but is also found in all other tissues investigated, it is highly expressed already in immature eggs.

### 3.4. In situ hybridization

To localize and confirm where the secretory ferritin transcripts (LsFer1 and LsFer2) are expressed, we utilized *in situ* hybridization. Both transcripts were found to be expressed in cells lining the intestinal tract of the adult female louse (Fig. 4).

### 3.5. Starvation

To evaluate the effect starvation has on the expression of LsFer1 and LsFer2, transcript levels were measured by qPCR in a time series after removing the lice from their host. With increased time absent from the food source, decreased levels of both transcripts were found in the adult female *L. salmonis* (Fig. 5). More specifically, the expression of LsFer1 is significantly down-regulated from day 2 and throughout the starvation period (t-test: day 1: \( p = 0.132 \), day 2: \( p = 0.001 \), day 4: \( p = 0.005 \), day 8: \( p = 0.0000213, n = 5 \)). At eight days of starvation, the expression of LsFer1 is decreased with on average 67% (Fig. 5a). LsFer2 shows a similar profile, but shows a significant decrease already from day 1 of starvation (t-test: day 1: \( p = 0.007, \) day 2: \( p = 0.0000219, \) day 4: \( p = 0.0000493, \) day 8: \( p = 0.00000669, n = 5 \)) and at day 8 of starvation, LsFer2 has decreased with on average 82% (Fig. 5b).

### 3.6. RNAi phenotype, histology and knockdown effect of LsFer1 and LsFer2

In order to study the functions of LsFer1 and LsFer2 in vivo, two independent RNAi knockdown experiments were conducted. In both experiments, the knockdown animals’ recovery rates (lice sampled at the end of the experiment/lice injected at the start of the experiment) are similar to the control animals’, ranging from 17%–59% (Table 3). However, the fecundity of the parasite is affected by knocking down LsFer1 or LsFer2. In both experiments, larvae from control animals hatched from all egg strings. However, knockdown of LsFer1 resulted in one hatched out of ten egg string pairs (exp. 1), and when repeated, no egg strings (out of 6 pairs) hatched (exp. 2). Knockdown of LsFer2 resulted in egg strings present in 10 of 23 animals, and only 5 pairs of these hatched (exp. 1). In experiment 2 of LsFer2 knockdown, all animals (7 out of 7) had egg strings upon termination, but none of these hatched.

In all control adult female salmon lice, a salmon blood-filled intestinal tract is seen as a red line stretching from the mouth part in the cephalothorax, through the genital segment with maturing oocytes, and finally through the abdomen ending in the rectum (Fig. 6a). When knocking down LsFer1 (Fig. 6b) and LsFer2 (Fig. 6c), an alteration in phenotype is seen. In both knockdown groups, the animals have little or no blood in their intestines, as seen by visual inspection (Table 3 and Fig. 6). While all lice in the control groups had a high amount of blood
in their intestine, this was not the case in the LsFer1 and LsFer2 knockdown animals (Table 3). In order to more closely examine the knockdown effects, sections were made for histology (Fig. 6). In the control louse (Fig. 6a), oocytes are stacked and well-structured with lipid-droplets (seen as white dots) and with chorions that define boundaries between individual oocytes. In the LsFer1 knockdown animal, oocytes appear as a homogenous mash and without visible chorions subdividing individual oocytes (Fig. 6b). Histology of the LsFer2 knockdown louse reveals a genital segment with little and highly under-developed oocytes present (Fig. 6c). Lice shown in Fig. 6 are representative of both experiments. Photographs of all lice from exp. 2 are shown in Supplementary Fig. S1.

The efficacies of the knockdowns of LsFer1 and LsFer2 were assessed by qPCR. Knockdowns were successful for both treatments in both experiments, and experiment 2 is presented with individuals’ levels of transcripts (Fig. 7). In experiment 1, LsFer1 is on average 85% down-regulated (Mann-Whitney U: p = 0.008, n = 5), and LsFer2 is on average 78% down-regulated (t-test: p = 0.005, n = 5). In experiment 2, knockdown of LsFer1 resulted in an on average 93% down-regulation (t-test: p = 0.000171, n = 5) (Fig. 7a). Knockdown of LsFer2 resulted in a down-regulation of 60–86% for four out of five animals tested, whereas up-regulation of 41% was found in the last one (Fig. 7b). This individual had a similar phenotype and morphology as the rest of the knockdown group, and is therefore not omitted from the dataset or statistical analysis (Mann-Whitney U: p = 0.076, n = 5).

4. Discussion

In this study, we present the characterization and first functional studies of ferritin in the obligate hematophagous ectoparasitic salmon louse, Lepeophtheirus salmonis. We report the presence of four ferritin homologs (LsFer1-4) in the genome of the salmon louse, and find that two of these (LsFer1-2) have predicted signal peptides in the N-terminus of their aa sequences. Functional studies show that LsFer1-2 are expressed in the midgut, and both are essential for the blood-feeding and reproduction of the adult female salmon louse.

Results from the sequence analysis tools used here give support to the hypothesis that LsFer1-4 are homologs of ferritin as there are significant sequence similarities with ferritins of other species found by BLAST searches. Since LsFer1,3 and 4 have the highly conserved motifs for the ferroxidase site, whereas LsFer2 does not, LsFer1,3 and 4 are likely heavy chain homologs (HCH), whereas LsFer2 is likely a light chain homolog (LCH). This is further supported by the phylogenetic analysis, which shows that LsFer1 mainly clusters together with other HCHs and LsFer2 mainly with other LCHs (Fig. 2). LsFer3-4 clusters phylogenetically closer together with ferritin subunits without a signal peptide. Classically secreted ferritins are commonly found in arthropods where they are reported to be involved in detoxification, storage and distribution of iron. As a well-studied phenomenon in insects, ferritin is loaded with absorbed iron in the endoplasmic reticulum (ER), and is then secreted to the hemolymph, a tissue fluid analogous to...
blood in vertebrates, and subsequently used as a vehicle for iron transportation to various tissues [38]. In the current study, we find that both LsFer1-2 ferritin subunits are predicted to have signal peptides, and are thus likely destined for the secretory pathway. Since the salmon louse lacks the iron transfer protein transferrin (unpublished, http://licebase.org), it may utilize secreted ferritin to shuttle iron to recipient tissues and still avoiding iron-related cytotoxicity. The localizations of the precursor and mature LsFer1-2 proteins are however not determined, as only transcript localization is assessed here. Further work is required to determine if ferritin acts as an iron transporter in the salmon louse. Also, whether LsFer1 and LsFer2 are subunits of the same protein, or if they make up two different proteins remains an open question.

Fig. 2. Unrooted phylogenetic tree of vertebrate and invertebrate ferritin subunits. The phylogenetic analysis was performed on the Phylogeny.fr platform. Secreted ferritins are marked with the green box and intracellular ferritins with the blue box. Salmon louse ferritins (LsFer1-4) are written in red letters. LsFer1 clusters together with secreted invertebrate HCH ferritins, and LsFer2 clusters together with secreted invertebrate LCH ferritins. LsFer3-4 cluster together and are close to branches of intracellular heavy and light chain ferritins of both vertebrate and invertebrate origin. Numbers at branches represent support values (approximate likelihood-ratio). HC(H) = heavy chain (homolog), LC(H) = light chain (homolog). Scale bar: 0.3 amino acid substitutions per site.

Fig. 3. Ontogeny. RNA-sequencing data (left and middle column: LiceBase, right column: Eichner et al., 2018 [28] with additional data from nauplius and attached copepodids (unpublished data)) for LsFer1 (a), LsFer2 (b), LsFer3 (c) and LsFer4 (d). Available reads were recounted for the transcribed regions of the ferritin genes found by RACE-PCR. See Section 2.4 “Ontogeny” for explanations of the abbreviations used in the figure.
unanswered question. Since LsFer2 lacks conserved aa residues for ferroxidase catalytic activity, it is highly unlikely that the LCH subunit may constitute an iron storage molecule on its own. Also, LsFer1 and LsFer2 contain cysteine residues reported to form intersubunit disulfide bonds in other secretory ferritins [6].

Characterization of the LsFer1-4 genes by RNA-Seq showed that LsFer1-2 have similar expression patterns. Both secreted ferritin subunits are highly expressed in the intestine of the salmon louse, consistent with having an iron storage and/or transportation function. Additionally, LsFer1-2 are expressed in the larval, non-feeding stages of the salmon louse, which could indicate extra functions of the ferritin subunits, or they could be necessary for detoxification of iron from the turnover of maternally derived iron-proteins. The similar expression patterns of LsFer1-2 are expressed in the larval, non-feeding stages of the salmon louse, which may constitute an iron storage molecule on its own. Also, LsFer1 and LsFer2 have a more uniform expression pattern, and probably has an intracellular iron storage function in several tissues of the salmon louse. Further analysis by in situ hybridization confirmed that the transcripts of LsFer1 and LsFer2 are both expressed in cells lining the intestinal tract of the adult female salmon louse. Epithelial cells in the gut are the entry site for iron absorbed from the blood of the parasite’s host, and is where iron storage proteins need to be ready and available. In the starvation experiment, the expression of LsFer1-2 is decreasing with the increased starvation time. A reason for this could be that the salmon louse is an ectoparasite that is either on or off its host. It does not experience a gradient in iron availability, other than remnants of blood in the gut after feeding (seen as a fainter red color with increased starvation time). Expressing a nutrient storage molecule when there is an absence of nutrients available is unnecessary and requiring energy. Similar to our results, an HCH ferritin in A. aegypti was found in the midgut of the mosquito and was regulated by blood-feeding [39]. Also, Wu et al., found that a ferritin homolog in abalone Haliotis discus hannai was up-regulated by dietary iron [40]. Walter-Nuno et al., also found that blood-feeding increased the expression level of several genes involved in heme and iron metabolism, including ferritin [41]. These findings indicate that ferritin gene expression is a consequence of the dietary iron status. A common way of controlling ferritin transcript translation in a cell is by iron regulatory proteins (IRPs) that inhibit ferritin translation during iron shortage by binding to an IRE upstream of the ferritin ORF [42]. Yet, no IREs were predicted in any of the ferritin sequences in the salmon louse, even though the louse does have an IRP-IRE complex [37]. We do not know if uptake of iron is controlling ferritin translation here as it does post-transcriptionally in vertebrates [43] and other various invertebrates [44,45]. Not all organisms depend on translational control by IRE however, e.g. plants regulate ferritin levels on a transcriptional level in response to iron and oxidants [46]. How ferritin expression is regulated in the salmon louse remains an unanswered question.

The functional study by knocking down LsFer1 and LsFer2 in the salmon louse mediated by RNAi had negative consequences for the fecundity of the parasite. Fewer of the adult female lice had egg strings, and when present, they had an overall much lower hatching success in both knockdown groups compared with the control group. Histological analyses demonstrated that oocyte development was impaired, and this is probably why eggs hatching was highly unsuccessful. Knockdown of genes involved in iron metabolism is already known to limit reproductive success of other hematophagous parasites, such as the tick’s reproduction and development [44,47,48]. Furthermore, knockdown of a ferritin in the vector Rhodius prolixus resulted in drastically reducing (95%) viable eggs for the hematophagous insect [41]. As the salmon louse is semi-transparent, its red, blood-filled gut can easily be observed

unanswered question. Since LsFer2 lacks conserved aa residues for ferroxidase catalytic activity, it is highly unlikely that the LCH subunit may constitute an iron storage molecule on its own. Also, LsFer1 and LsFer2 contain cysteine residues reported to form intersubunit disulfide bonds in other secretory ferritins [6].

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by visual inspection. Animals in the knockdown group had less blood in
their intestines than animals in the control group, which were fully
engorged. This phenomenon has also been reported in ticks upon
knockdown of ferritin [48]. An explanation for this could be as free iron
generates ROS, having a reduced capacity of sequestering and thus
detoxifying iron could lead to detrimental effects for the parasite upon
blood-feeding. A solution for the salmon louse could be to cease the
blood-feeding behavior. Downstream effects of not ingesting blood
would include severe undernutrition and consequently cause egg pro-
duction to be halted.

In conclusion, this report on four ferritin homologs in the salmon
louse is the first where we have functionally characterized the two se-
cretory subunits (LsFer1-2). LsFer1 and LsFer2 were both found to be
vital for blood-feeding and reproduction in the hematophagous parasite

![Fig. 6. Phenotype and histology. Phenotype and toluidine blue stained histological sections of an untreated control adult female louse (a), a LsFer1 RNAi knockdown louse (b) and a LsFer2 RNAi knockdown louse (c). Of every histological section, a magnified photo of the genital segment is shown in the lower panel. All three lice are representative of their group. CT = cephalothorax, GS = genital segment, AB = abdomen, i = intestine, o = oocyte.](image)

![Fig. 7. Knockdown efficacy. RNAi mediated knockdown effect of (a) LsFer1 and (b) LsFer2 measured by qPCR from experiment 2. Results are displayed as individual animals' values, and the mean value for each group is indicated by a dotted line. ** = p ≤ 0.01. n = 5.](image)
in a knockdown study and starvation experiment. Our results confirm several previous findings on the importance of ferritin throughout arthropods. Further work on these genes and proteins could provide a deeper knowledge of the iron metabolism of *L. salmonis* and should be studied further. Considering the detrimental effects the knockdown of *LsFer1-2* had on the egg production in the salmon louse, these results could find their application as treatment targets in the aquaculture sector in order to combat the infamous parasite.

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**Declaration of Competing Interest**

None.

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**Appendix A. Supplementary data**

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