ORIGINAL PAPER

A new intracellular bacterium, *Candidatus* Similichlamydia labri sp. nov. (Chlamydiaceae) producing epitheliocysts in ballan wrasse, *Labrus bergylta* (Pisces, Labridae)

Andreas Steigen · Egil Karlsbakk · Heidrun Plarre · Kuninori Watanabe · Aina-Cathrine Øvergård · Øyvind Brevik · Are Nylund

Received: 5 November 2014 / Accepted: 11 November 2014 / Published online: 22 November 2014 © The Author(s) 2014. This article is published with open access at Springerlink.com

Abstract Certain wrasse species (Labridae) are used as cleaner fish in salmon farms on the Norwegian coast, reducing salmon louse intensities. The pathogen repertoire of wrasse in Norway is poorly known, and the objective of the present study is to describe a novel intracellular bacterium detected in Norwegian Labrus bergylta. Histological examination of gill tissues from ballan wrasse, L. bergylta, revealed epitheliocysts occurring basally to the secondary lamellae in the interlamellar epithelium. Ultrastructurally, these had bacteria-filled inclusions with thickened membranes and radiating ray-like structures (actinae). 16S rRNA gene sequences from the gill bacteria showed the highest (97.1 %) similarity to Candidatus Similichlamydia latridicola from the gills of the latrid marine fish Latris lineata in Australia and 94.9 % similarity to Candidatus Actinochlamydia clariae, causing epitheliocystis in the freshwater catfish Clarias gariepinus in Uganda. A total of 47 gill samples from L. bergylta from Western Norway were screened by real time RT-PCR with an assay targeting Candidatus Actinochlamydiaceae 16S rRNA. Prevalence was 100 %. We propose the name Candidatus Similichlamydia labri sp. nov. for this new agent producing gill epitheliocysts in L. bergylta.

Communicated by Erko Stackebrandt.

A. Steigen (\boxtimes) · E. Karlsbakk · H. Plarre · K. Watanabe · Ø. Brevik · A. Nylund Department of Biology, University of Bergen, Bergen, Norway e-mail: andreas.steigen@bio.uib.no

E. Karlsbakk Institute of Marine Research, Bergen, Norway

A.-C. Øvergård Sea Lice Research Center (SLRC), Institute of Marine Research, 5817 Bergen, Norway **Keywords** Epitheliocystis · Chlamydiales · Actinochlamydiaceae · Similichlamydia labri · Labridae · *Labrus bergylta*

Introduction

The ectoparasitic salmon louse, Lepeophtheirus salmonis, is a serious problem to farming of Atlantic salmon, Salmo salar, in Norway (Heuch et al. 2005). Among other control strategies, salmon farmers in Western Norway use wrasse (Labridae) as cleaner fish to reduce lice levels and the harmful consequences due to L. salmonis. In Western Norway, salmon famers mainly stock four, wild-caught, wrasse species with their salmon: Labrus bergylta (ballan wrasse), Ctenolabrus rupestris (goldsinny), Symphodus melops (corkwing wrasse) and Centrolabrus exoletus (rock cook). The fish is transported over long distances on the Norwegian coast, raising concerns about the possible spread of pathogens to previously uninfected wrasse populations. To gain knowledge on naturally occurring pathogens in wrasse, a study of gill-associated bacteria and parasites was initiated in 2011. It was found that epitheliocysts were common in the gills of *L. bergylta* in Western Norway.

Epitheliocysts have been observed on the gills in many fish species. In most cases, these show a chlamydia-like intracellular development (cf. Nylund et al. 1998). The rRNA gene sequences also usually suggest membership within Chlamydiales (Draghi et al. 2004, 2007; Meijer et al. 2006; Nowak and LaPatra 2006; Karlsen et al. 2008; Horn 2008; Polkinghorne et al. 2010; Schmidt-Posthaus et al. 2011; Corsaro and Work 2012; Fehr et al. 2013; Steigen et al. 2013; Stride et al. 2013a, b). Exceptions include a betaproteobacterium, found to cause epitheliocystis in farmed Atlantic salmon (Toenshoff et al. 2012;



Table 1 Primers used for PCR and sequencing of the 16S rRNA gene and ITS from *Candidatus* Actinochlamydiaceae obtained from the gills of wrasse

| Primer | Sequence | Target gene | Publication |
|----------|-------------------------------|-------------|----------------------|
| Pic-F1 | AAG CAC TTT TGC CTG GGA GC | 16S | Present study |
| Chits-F1 | GGA ATT GCT AGT AAT GGC G | ITS | Present study |
| Chits-R1 | TGG TCT CCC CAG ATT CAG ACS | ITS | Present study |
| Chits-R2 | GTC TCC CCA GAT TCA GAC CG | ITS | Present study |
| 16sigF | CGG CGT GGA TGA GGC AT | 16S | Draghi et al. (2004) |
| 806R | GGA CTA CCA GGG TAT CTA AT | ITS | Draghi et al. (2004) |
| 16sB1 | TAC GGY TAC CTT GTT ACG ACT T | 16S | Draghi et al. (2004) |

Mitchell et al. 2013), and a gammaproteobacterium responsible for epitheliocystis in cobia *Rachycentrum canadum* larvae (Mendoza et al. 2013).

In this study, we describe and characterise a new species of chlamydia detected in the gills of ballan wrasse (*Labrus bergylta*) from Western Norway. This new species show unique ultrastructural characteristics, resembling those seen in epitheliocysts in the African sharptooth catfish, *Clarias gariepinus*, from freshwater in Uganda (Steigen et al. 2013).

Materials and methods

Fish

Specimens of ballan wrasse, *Labrus bergylta*, were collected during May–June 2012 and May 2013 in Raunefjorden close to Bergen, Western Norway. These fish were mature or had just spawned. An autumn sampling period was in October 2012. Farmed ballan wrasse (N=16) were also obtained from two different brood stock populations in Western Norway. The studied ballan wrasse caught in May 2013 measured 20–26 cm (N=15), in June 2012, 21–41 cm (N=27), and in October 2012, 27–37 cm (N=4). The fish were caught with fyke nets and transported live to the laboratory at the University of Bergen. Gill samples were taken from newly killed fish, and frozen at -20 °C (for DNA/RNA) or fixed (4 °C) for histology including transmission electron microscopy (TEM).

Histology and transmission electron microscopy

Gill tissue samples were immersed and fixed in a modified Karnovsky's fixative: distilled water replaced by Ringer's solution where sucrose was added (4 % w/v solution) (Nylund et al. 1998). The fixed tissues were used for histological studies and TEM, and were processed as described in Steigen et al. (2013).

PCR/RT-PCR

RNA was extracted from individual gill samples of all specimens from each species sampled (as in Steigen et al.

2013). The RNA was used for real time RT-PCR detection of chlamydiae belonging to the *Candidatus* family Actinochlamydiaceae using the ChV assay (Steigen et al. 2013). DNA was extracted from selected samples with low cycle threshold values (Ct), since these contain larger amounts of bacteria. The 16S rRNA gene, the ITS region, and the partial 23S of the *Candidatus* Actinochlamydiaceae present were sequenced from five ballan wrasse selected due to low Ct values obtained by the ChV assay. A second type of chlamydia (clade A) was detected in six ballan wrasse individuals by PCR. The ChV assay does not amplify the clade A chlamydia. The PCR was performed as described by Steigen et al. (2013). Primers used are presented in Table 1.

In situ hybridization

In situ hybridization was performed as described by Dalvin et al. (2013), with some modifications presented in Tröße et al. (2014). The digoxigenin labelled (DIGlabelled) sense and antisense RNA probes (776 nucleotides) were made with primers: 16SIGF and 806R (Draghi et al. 2004).

Phylogeny

The sequence data were preliminarily identified by GenBank searches done with BLAST (2.0). The Vector NTI Suite software package was used to obtain multiple alignments of sequences. Pairwise comparison of 16S sequences was done in GeneDoc (available at: www.psc.edu/biomed/genedoc). sequences Selected from other members of order Chlamydiales, available on the EMBL nucleotide database, were included in the comparisons. Phylogenetic analyses were performed using TREE-PUZZLE 5.2 (available at: http://www.treepuzzle.de), maximum likelihood (ML) (50,000 puzzling steps). The data were analysed using a GTR-I nucleotide evolution model, selected in jModelTest; with default settings and the Akaike information criterion option (Posada 2008). Phylogenetic trees were drawn using TreeView (Page 1996).



Ethics statement

All fish were treated according to the Norwegian Animal Welfare Act (01. 01. 2010) following regulations from the Norwegian Food Safety Authority.

Results

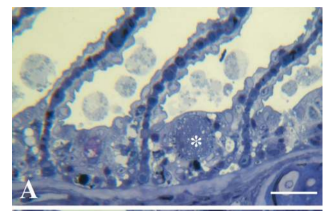
Occurrence of chlamydiae in Labrus bergylta

Screening (16S rRNA sequences) of gill tissues were carried out on 47 *L. bergylta*. PCR showed a 100 % prevalence of *Candidatus* Actinochlamydiaceae infections with Ct values always below 30. In May, samples average Ct was 18.5 (range 15.2–21.5), in June 21.4 (14.4–29.3), and in October 16.8 (15.0–18.1). Sequencing of the 16S rRNA gene from five samples provided identical or nearly identical sequences (identity 99.9–100.0, 2,102 nt compared) (type-A; accession nos: KC469556-8, KC469562, KC469564), with the highest affinity to *Candidatus* Similichlamydia latridicola.

Ballan wrasse was also found infected with a second chlamydia species (clade A), detected through PCR with general primers for Chlamydiaceae. The clade A (accession nos: KC469554-5, KC469559-61, KC469563) sequences showed the highest affinity to *Candidatus* Parilichlamydia carangidicola. Fish found infected with the clade A type of chlamydia were not used in the histological studies; the sequences, however, were included in the phylogeny.

Bacteria and inclusion morphology

Epitheliocysts in infected L. bergylta had a distinctive morphology (Fig. 1). The cysts occurred basal to the secondary lamellae, in cells resembling chloride cells (mitochondria and ER rich cells). The inclusions had numerous actinae radiating from the inclusion membrane (IM). Cysts in contact with the gill surface, reached 30 µm in diameter, but most inclusions measured <25 µm. Ultrastructurally, the IM was thickened and electron dense with regularly spaced actinae radiating from it (Fig. 2). The thickness of the IM was 50-100 nm, being thickest close to the actinae. The actinae consisted of the same electron dense material as the IM and showed longitudinal ridges on the surface producing a characteristic star-like pattern in transverse sections (Fig. 3). There was a central evagination of the inclusion vacuole into each actina, but the actinae did not appear to be hollow throughout. Near the IM, the actinae reached c. 650 nm in diameter, with up to 17 ridges, but some were smaller with only 6-7 ridges. The diameter and number of ridges tended to decrease with increasing distance from the inclusion. The actinae might reach to and apparently penetrate neighbouring



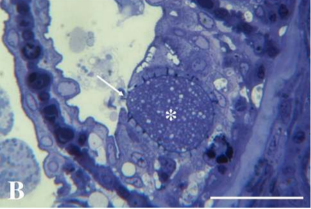


Fig. 1 *Candidatus* Similichlamydia labri sp. nov. from the gills of *Labrus bergylta*, semi-thin sections stained with toluidine *blue*. **a** Epitheliocyst (*asterisk*) with typical site in an interlamellar crypt. Actinae can be seen radiating from the inclusion membrane. $Bar = 25.0 \, \mu m$. **b** An epitheliocyst with radiating actinae, opening (*arrow*) on the surface of the gills basal to the secondary lamellae. $Bar = 30.0 \, \mu m$ (colour figure online)

cells; in some cases, the plasma membrane of the neighbouring cell disappeared in the contact region (Fig. 3a). The length of the actinae was typically $1.3{\text -}1.8~\mu\text{m}$, but could reach $2.1~\mu\text{m}$. The cytoplasm of the host cell was filled with mitochondria and membrane structures (endoplasmic reticulum), which occurred among the actinae and close to the IM.

The bacteria inside the inclusions were polymorphic and varied from irregular shapes with several nucleoids or with no distinct nucleoids [reticulate body (RB)-like morphology] to coccoid shapes with distinct electron dense nucleoids [intermediate body (IB)-like morphology]. The irregular and electron lucent RBs could exceed 2 µm in length. The coccoid cells had a diameter of about 200–360 nm with a nucleoid diameter of about 100 nm, some cells larger than 300 nm were seen dividing. The smallest coccoid cells had a more condensed cytoplasm, and they were not seen dividing and measured 200–250 nm in diameter [possibly an early stage of elementary bodies (EB)]. A cell wall and a cytoplasmic unit membrane surrounded all morphs of the



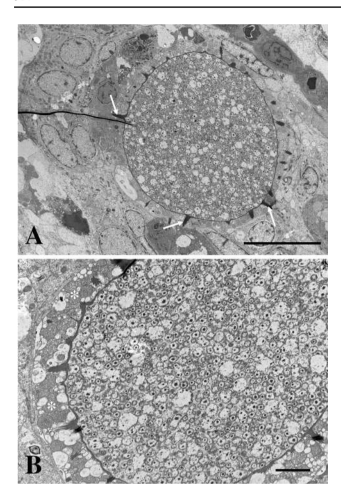


Fig. 2 Candidatus Similichlamydia labri sp. nov. from the gills of Labrus bergylta, TEM. **a** An epitheliocyst between two secondary lamellae showing a round inclusion containing bacteria and an inclusion membrane with radiating actinae (arrows). There is no significant reaction in the surrounding tissue. $Bar = 10 \mu m$. **b** Inclusion containing bacteria with different morphologies (RB-like and IB-like morphs). The actinae penetrate the host cell cytoplasm, which contains large amounts of mitochondria (asterisks). $Bar = 2.0 \mu m$

bacteria inside the inclusions, but the cell wall was more distinct in the coccoid cells.

There was no sign of a host response (inflammation, hyperplasia) associated with the epitheliocysts in ballan wrasse.

In situ hybridization

Inclusions in cells in the interlamellar epithelium of *Labrus bergylta* were specifically labelled with the 776 nt antisense riboprobe that was transcribed from 16S rRNA (accession no: KC469556) amplified from gills of the fish host. The ISH positive infected individual was among those studied by histology and TEM. Inclusions in adjacent histological sections that were incubated with the sense probe (776 nt) were not labelled (Fig. 4).

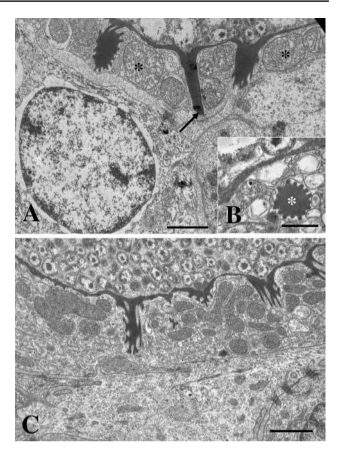


Fig. 3 *Candidatus* Similichlamydia labri sp. nov. from the gills of *Labrus bergylta*, TEM. **a** High magnification of the actinae penetrating the host cell cytoplasm and into the neighbouring cell membrane (*arrow*). Mitochondria indicated by asterisks. $Bar = 1.0 \, \mu m$. **b** Transverse section of an actinia with sprocket-like morphology (*asterisk*). $Bar = 0.5 \, \mu m$. **c** A large amount of mitochondria and endoplasmic reticulum is present in the host cell cytoplasm between the actinae. $Bar = 1.0 \, \mu m$

Phylogeny and sequence comparisons

Phylogenetic analyses showed that the 16S rRNA gene sequences of chlamydiae from ballan wrasse with epitheliocysts group with related chlamydiae from other Norwegian wrasse species into a major clade together with three previously described species: *Candidatus* Similichlamydia latridicola from striped trumpeter (*Latris lineata*) from a Tasmanian aquaculture facility, *Candidatus A. clariae* from *C. gariepinus* collected in fresh water in the vicinity of Lake Victoria, and *Candidatus* Parilichlamydia carangidicola from yellowtail kingfish (*Seriola lalandi*) in Australian waters (Figs. 5, 6). Two additional sequences obtained from the gills of *Oreochromis niloticus* (JQ480302, JQ480303) collected in Uganda also belong to this major clade (Fig. 5).

BLAST search and sequence comparisons revealed that sequence similarity of the chlamydiae to *Candidatus* S. latridicola is 97.1 % (1,395 nt compared), to



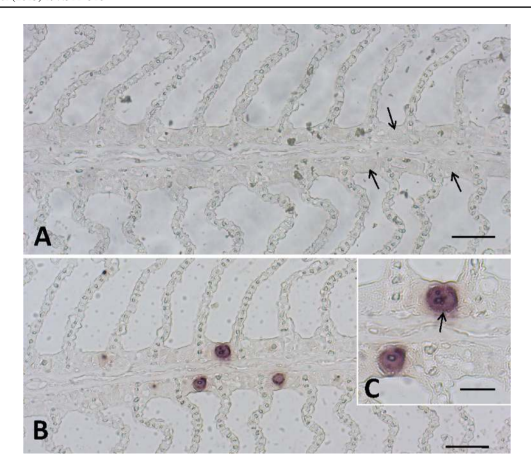


Fig. 4 Sections of gill tissue from *Labrus bergylta* processed for in situ hybridization. **a** Primary gill filament stained with a sense probe, showing the absence of staining of the inclusions (*arrows*). $Bar = 50.0 \mu m$. **b** Primary filament showing dark-blue stained Candidatus Similachla-

mydia labri inclusions, stained with antisense DIG-labelled RNA probe against 16S rRNA of the bacterium. $Bar = 50.0 \, \mu m$. c Magnification of the IHC stained inclusion in figure B with discernible actiniae (arrow). $Bar 20.0 = \mu m$ (colour figure online)

Candidatus A. clariae 94.9 % (1,464 nt compared), and to Candidatus P. carangidicola 93.3 % (1,103 nt compared). The clade A sequences from ballan wrasse showed 93.8 % similarity with the chlamydia sequences (1,464 nt compared) from ballan wrasse with the described epitheliocysts. Phylogenetically, the two types grouped into two subclades (Fig. 5). One type grouped with Candidatus S. latridicola and Candidatus A, clariae, while the second type (clade A) grouped with Candidatus P. carangidicola (Fig. 6).

Discussion

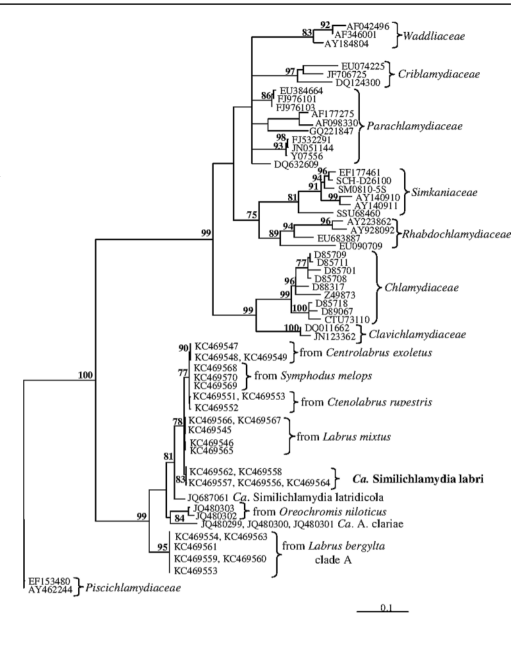
None of the Chlamydiales associated with epitheliocystis in fish have been cultured. New species are proposed as *Candidatus* species, based on the uniqueness of the 16S rRNA gene sequences, ultrastructure, and host specificity. A 16S rRNA gene sequence divergence of 1.0–1.3 % may

separate different species (Stackebrandt and Ebers 2006; Kim et al. 2014).

The cysts and inclusions in infected L. bergylta showed similarities to those found in Candidatus Actinochlamydia clariae described from the gills of the African sharptooth catfish Clarias gariepinus in Uganda (Steigen et al. 2013). Ultrastructurally, they share a thickened electron dense inclusion membrane and the presence of actinae. The inclusions and the epitheliocysts reach a slightly larger size in ballan wrasse than in Candidatus A. clariae in catfish (30 vs. 23 µm in diameter). However, Candidatus Similichlamydia latridicola is more closely related to the L. bergylta chlamydiae than Candidatus A. clariae according to the 16S rRNA sequences. This species produce large cysts (>50 µm) in the secondary lamellae in the gills of Latris lineata. However, actinae have not been reported from Candidatus S. latridicola and are not evident from images in Stride et al. (2013b), and the ultrastructure of that candidate species is unknown.



Fig. 5 Phylogenetic tree showing the relationship between *Candidatus* Similichlamydia labri sp. nov. from *L. bergylta* and selected members of other families within order Chlamydiales. The analysis is based on a 1,210 nt long edited alignment of 16S rDNA sequences. The *scale bar* shows the number of nucleotide substitutions as a proportion of branch lengths. *Ca. = Candidatus*



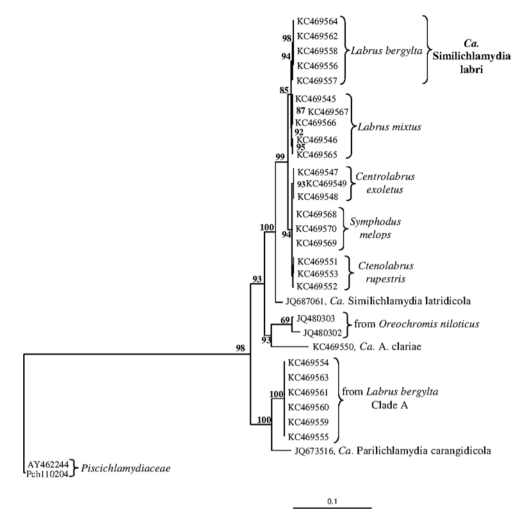
Another difference between the epitheliocysts from infected ballan wrasse and *Candidatus* A. clariae is the appearance of the host cell cytoplasm. The inclusions of both chlamydiae were surrounded by a limited volume of host cell cytoplasm containing mitochondria and endoplasmic reticulum (cf. Steigen et al. 2013). However, a layer of vacuoles often surrounded the inclusions of *Candidatus* A. clariae setting them apart from the rest of the host cell (Steigen et al. 2013). Such vacuoles were not common around the inclusions from ballan wrasse.

We did not observe bacterial stages in the cysts from ballan wrasse similar to the elementary bodies (EB) of *Candidatus* A. clariae. The EBs from the latter contains a 'polar cap area' penetrated by rod-like structures arranged in a hexagonal pattern and protruding from the surface of the cap, and an eccentric nucleoid (Steigen et al. 2013). The lack of mature EB's in the epitheliocysts from the studied ballan wrasse prevents a thorough comparison with the developmental stages of *Candidatus* A. clariae. However, further sampling should solve this issue.

The 16S rRNA gene sequences of the *L. bergylta* chlamydiae show 94.9 % similarity to those from *Candidatus* A. clariae. This suggests that they belong to the same family, but to a separate species in a different genus (cf. Everett 2000; Bush and Everett 2001; Stackebrandt and Ebers 2006). Based on 16S RNA gene sequence similarity (97.1 %), the *L. bergylta* chlamydia is congeneric with *Candidatus* S. latridicola. The clade A sequences obtained from *L. bergylta* from Western Norway (accession no: KC469554-5, KC469559-61, KC469563) and



Fig. 6 Phylogenetic tree showing the relationships between possible members of the family *Candidatus* Actinochlamydiaceae. Sequences of 16S rRNA gene sequences from *Candidatus* Piscichlamydia salmonis have been used as outgroup. The analysis is based on a 1,103 nucleotides long alignment of 16S rRNA gene sequences. The *scale bar* shows the number of nucleotide substitutions as a proportion of branch lengths. *Ca. = Candidatus*



Candidatus P. carangidicola (accession no: JQ673516) from Australian waters are the nearest relatives to the Ca. Actinochlamydiaceae, grouping in a separate clade (Fig. 5). The morphology of the clade A chlamydiae is unknown.

The epitheliocyst producing agent found and described in the present study shows a close relationship to species found in both fresh- and seawater, in cold and in warm regions. The wide geographical distribution is remarkable, with related species off Australia and Norway, and in Lake Victoria. A high global diversity therefore seems likely for these parasitic bacteria, meriting further investigations. Material now being processed in our laboratory will expand this distribution further and add new *Candidatus* species.

We suggest that the described chlamydia from *Labrus bergylta* is given status as a new species, *Candidatus* Similichlamydia labri n.sp. The 16S RNA gene sequences, site, host, and geographic location clearly distinguish it from *Candidatus* Similichlamydia latridicola.

Taxonomy

Candidatus Similichlamydia labri [labr'i M.L. gen. sing. of Labrus, generic name for the fish host Labrus bergylta; of (living in) members of the genus Labrus].

Type host: *Labrus bergylta* Ascanius (Labridae)

Site: intracellular inclusions in chloride cells in the gills Type locality: Raunefjorden, near Bergen, Hordaland County, Norway (60°16′N, 05°13′E)

16S and ITS rDNA Sequence: GenBank Acession No. KC469556

The provisional taxon *Candidatus* Similichlamydia labri represents an intracellular bacterium within membrane-bound vacuoles (inclusions) in the cytoplasm of chloride cells in the gills of *Labrus bergylta*. Inclusion membranes are thickened, ridged, and give rise to ridged actinae. The developmental stages include RBs of varying size, shape and number of nucleoids, and smaller coccoid cells,



<300 nm in diameter. The bacterium cell wall is thin; typical EBs have not been observed.

Acknowledgments The authors wish to thank the skilled and dedicated staff at the University of Bergen's Marine Biological Station at Espegrend for providing fish for our investigations.

Open Access This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

References

- Bush RM, Everett KDE (2001) Molecular evolution of the Chlamydiaceae. Int J Syst Evol Microbiol 51:203–220
- Corsaro D, Work TM (2012) Candidatus Renichlamydia lutjani, a gram-negative bacterium in internal organs of blue-striped snapper Lutjanus kasmira from Hawaii. Dis Aquat Org 98:249–254
- Dalvin S, Nilsen F, Skern-Mauritzen R (2013) Localization and transcription patterns of LsVasa, a molecular marker of germ cells in Lepeophtheirus salmonis (Krøyer). J Nat Hist 47(5–12):889–900
- Draghi A, Vsevolod LP, Kahl MM, Stanton JB, Brown CC, Tsongalis G, West AB, Frasca S (2004) Characterization of "Candidatus Piscichlamydia salmonis" (order Chlamydiales), a Chlamydialike bacterium associated with epitheliocystis in farmed Atlantic salmon (Salmo salar). J Clin Microbiol 42:5286–5297
- Draghi A, Bebak J, Popov VL, Noble AC, Geary SJ, West AB, Byrne P, Frasca S (2007) Characterization of a *Neochlamydia*-like bacterium associated with epitheliocystis in cultured Arctic charr Salvelinus alpinus. Dis Aquat Org 76:27–28
- Everett KD (2000) *Chlamydia* and *Chlamydiales*: more than meets the eye. Vet Microbiol 75:109–126
- Fehr A, Walther E, Schmidt-Posthaus H, Nufer L, Wilson A, Svercel M, Richter D, Segner H, Pospischil A, Vaughan L (2013) Candidatus Syngnamydia venezia, a novel member of the phylum Chlamydiae from broad nosed pipefish, Syngnathus typhle. PLoS One 8(8):e70853. doi:10.1371/journal.phone.0070853
- Heuch PA, Bjørn PA, Finstad B, Holst JC, Asplin L, Nilsen F (2005) A review of the Norwegian 'National action plan against salmon lice on salmonids': the effect on wild salmonids. Aquaculture 246:79–92
- Horn M (2008) Chlamydiae as symbionts in eukaryotes. Ann Rev Microbiol 62:113–131
- Karlsen M, Nylund A, Watanabe K, Helvik JV, Nylund S, Plarre H (2008) Characterization of *Candidatus* Clavochlamydia salmonicola: an intracellular bacterium infecting salmonid fish. Environ Microbiol 10:208–218
- Kim M, Oh H-S, Park S-C, Chun J (2014) Towards a taxonomic coherence between average nucleotide identity and 16S rRNA gene sequence similarity for species demarcation of prokaryotes. Intern J Syst Evol Microbiol 64:346–351
- Meijer A, Roholl PJM, Ossewaarde JM, Jones B, Nowak BF (2006) Molecular evidence for association of *Chlamydiales* bacteria with epitheliocystis in leafy seadragon (*Phycodurus eques*), silver perch (*Bidyanus bidyanus*), and barramundi (*Lates calcarifer*). Appl Environ Microbiol 72:284–290

- Mendoza M, Guiza L, Martinez X, Caraballo X, Rojas J, Aranguren LF, Salazar M (2013) A novel agent (*Endozoicomonas elysicola*) responsible for epitheliocystis in cobia *Rachycentrum canadum* larvae. Dis Aquat Org 106:31–37
- Mitchell SO, Steinum TM, Toenshoff ER, Kvellestad A, Falk K, Horn M, Colquhoun DJ (2013) *Candidatus* Branchiomonas cysticola is a common agent of epitheliocystis in seawater-farmed Atlantic salmon in Norway and Ireland. Dis Aquat Org 103:35–43
- Nowak BF, LaPatra SE (2006) Epitheliocystis in fish. J Fish Dis 29:573-588
- Nylund A, Kvenseth AM, Isdal E (1998) A morphological study of the epitheliocystis agent in farmed Atlantic salmon. J Aquat Anim Health 10:43–55
- Page RD (1996) Tree view: an application to display phylogenetic trees on personal computers. Appl Biosci 12:357–358
- Polkinghorne A, Schmidt-Posthaus H, Meijer A, Lehner A, Vaughan L (2010) Novel *Chlamydiales* associated with epitheliocystis in a leopard shark *Triakis semifasciata*. Dis Aquat Org 91:75–81
- Posada D (2008) jModelTest: phylogenetic model averaging. Mol Biol Evol 25:1253–1256
- Schmidt-Posthaus H, Polkinghorne A, Nufer L, Schifferli A, Zimmernmann DR, Segner H, Steiner P, Vaughan L (2011) A natural freshwater origin for two chlamydial species, *Candidatus* Piscichlamydia salmonis and *Candidatus* Chlavochlamydia salmonicola, causing mixed infections in brown trout (*Salmo trutta*). Environ Microbiol 14:2048–2057
- Stackebrandt E, Ebers J (2006) Taxonomic parameters revisited: tarnished gold standards. Microbiol Today 33:152–155
- Steigen A, Nylund A, Akoll P, Fiksdal IU, Karlsbakk E, Nylund S, Odong R, Plarre H, Semyalo R, Skår C, Watanabe K (2013) 'Cand. Actinochlamydia clariae' gen. nov., sp. nov., a unique intracellular bacterium causing epitheliocystis in catfish (Clarias gariepinus) in Uganda. PLoS One 8(6):e66840. doi:10.1371/journal.pone.0066840
- Stride MC, Polkinghorne A, Miller TL, Groff JM, LaPatra SE, Nowak BF (2013a) Molecular characterization of "Candidatus Parilichlamydia carangidicola", a novel Chlamydia-like epitheliocystis agent in yellowtail kingfish, Seriola lalandi (Valenciennes), and the proposal of a new family, "Candidatus Parilichlamydiaceae" fam.nov. (Order Chlamydiales). Appl Environ Microbiol 79:1590–1597
- Stride MC, Polkinghorne A, Miller TL, Nowak BF (2013b) Molecular characterisation of 'Candidatus Similichlamydia latridicola gen. nov., sp. nov.' (Chlamydiales: Parilichlamydiaceae), a novel Chlamydia-like epitheliocystis agent in striped trumpeter, Latris lineata (Forster). Appl Environ Microbiol 79(16):4914. doi:10.1128/AEM.00746-13
- Toenshoff ER, Kvellestad A, Mitchell SO, Steinum T, Falk K, Colquhoun DJ, Horn M (2012) A novel betaproteobacterial agent of gill epitheliocystis in seawater farmed Atlantic salmon (*Salmo salar*). PLoS One 7(3):e32696. doi:10.1371/journal.pone.0032696
- Tröße C, Nilsen F, Dalvin S (2014) RNA interference mediated knockdown of the KDEL receptor and COPB2 inhibits digestion and reproduction in the parasitic copepod *Lepeophtheirus* salmonis. Comp Biochem Physiol Part B Biochem Mol Biol. doi:10.1016/j.cbpb.2013.12.006

