Genome of the facultative scuticociliatosis pathogen *Pseudocohnilembus persalinus* provides insight into its virulence through horizontal gene transfer

Jie Xiong1, Guangying Wang1,2, Jun Cheng1,3, Miao Tian1,2, Xuming Pan4, Alan Warren5, Chuanqi Jiang1,2, Dongxia Yuan1 & Wei Miao1

Certain ciliates of the subclass Scuticociliatia (scuticociliates) are facultative parasites of fishes in which they cause a suite of diseases collectively termed scuticociliatosis. Hitherto, comparatively little was known about genetics and genomics of scuticociliates or the mechanism of scuticociliatosis. In this study, a laboratory culture of the facultatively pathogenic scuticociliate *Pseudocohnilembus persalinus* was established and its genome sequenced, giving the first genome of a marine ciliate. Genome-wide horizontal gene transfer (HGT) analysis showed *P. persalinus* has acquired many unique prokaryote-derived genes that potentially contribute to the virulence of this organism, including cell adhesion, hemolysis and heme utilization genes. These findings give new insights into our understanding of the pathology of scuticociliates.

Scuticociliatosis, caused by certain scuticociliates1, is one of the most important parasitological problems in marine aquaculture worldwide2. In recent years there have been many reports of fatal outbreaks of infection by several scuticociliates species including *Pseudocohnilembus persalinus*, *Uronema marinum*, and *Philasterides dicentrarchi* in east Asia (Korea, Japan, China), Europe (Spain, Portugal) and other regions of the world, which have led to serious economic losses because of high mortalities of many fish species, in particular olive flounder and turbot1-3,4. Most studies of scuticociliatosis have focused on species identification, histopathology and immunology with little attention paid to molecular mechanisms of pathogenicity, mainly due to the lack of basic research on topics such as the life cycle, genetics and genome. Unlike the hymenostome ciliate *Ichthyophthirius multifiliis*, an obligate parasite of freshwater fish with a typical parasitic life cycle and distinct polymorphism5,6, scuticociliates are generally free-living in limnetic and marine ecosystems, feeding on other microorganisms such as bacteria, microalgae, protozoa etc7. Under certain circumstances, however, these ciliates may become opportunistic histophagous parasites, actively feeding on cells and tissue residues of a host organism, living and reproducing within the host tissues without observable morphological change.

Scuticociliates belong to the Scuticociliatia, one of six subclasses of the class Oligohymenophorea, the others being Periculina, Hymenostomatia, Astomatia, Apostomatia, and Peritrichia8. Some
oligohymenophoreans are the best-known of all ciliates and are commonly used as model laboratory organisms. These include the hymenostome species *Tetrahymena thermophila*11,12 and *I. multifiliis* and the peniculian species *Paramecium tetraurelia*13,14. Genomes of these ciliates have been sequenced and comparative genomics analyses have provided a comprehensive understanding of the unique features of these free-living (*T. thermophila* and *P. tetraurelia*) and typical parasitic (*I. multifiliis*) ciliates. Thus, it is expected that the genomes of parasitic scuticociliates will provide new insights and better understanding of their mechanisms of pathogenicity.

The typical *Pseudocohnilembus* species, *P. persalinus* has been reported previously as a free-living marine ciliate15–17. Since Kim et al. (2004) isolated *P. persalinus* from diseased olive flounder in Korea, it has become recognized as an important facultative parasite causing scuticociliatosis in commercially important fishes such as rainbow trout18 and olive flounder19. *Pseudocohnilembus persalinus* can be cultured in the laboratory as a free-living form by feeding with bacteria, which enables sufficient DNA/RNA to be collected for genome/transcriptome sequencing. Here, we report the genome of *P. persalinus*, the first to be sequenced among scuticociliates and, following comparative genomics analyses with its close relatives, investigate the possibility that the acquisition of its virulence may be via horizontal gene transfer (HGT) from bacteria.

**Results and Discussion**

**Facultative parasitism of *P. persalinus* - a genomic view.** Like most other ciliates, *P. persalinus* has two types of nucleus, a macronucleus (MAC) and a micronucleus (MIC), the former controlling the physiological and biochemical functions of the cell and the latter as a germ-line reserve. The first ciliate genome to be sequenced was the MAC genome of *T. thermophila*22 and this was achieved by first physically separating the MAC from the MIC. However, such a method has not been established for *P. persalinus*. In most ciliates the MIC is either haploid or diploid whereas the MAC is polyploid. In *T. thermophila*, for example, the MAC has an average ploidy up to 45C11, and in *I. multifiliis*, the MAC has an estimated ploidy up to 12,000C20. Therefore, we anticipated that the MAC of *P. persalinus* also has a high ploidy level and utilized this natural enrichment of MAC in order to perform MAC genome sequencing as had previously been achieved in *I. multifiliis*20. By employing this simple sequencing strategy, a preliminary assembly of about 63.6 Megabase (Mb) sequences with 2403 scaffolds was constructed. The GC content distribution of the preliminary assembly showed that there are three peaks of 19%, 41% and 67% (Fig. 1, red), indicating the presence of contaminants. Compared to the second and third peaks, the first peak contained a majority of sequences (sequence length, about 87%, Fig. 1, red) with a small population of scaffolds with low GC content (scaffold number, Fig. 1, blue). Published oligohymenophorean ciliate genomes usually show a low GC content (22% in *T. thermophila*, 15% in *I. multifiliis* and 28% in *P. tetraurelia*, Table 1), therefore sequences represented by the first peak probably derive from *P. persalinus*. In order to verify this, BLASTX searches were performed for all preliminary assembled sequences against the NCBI non-redundant protein sequences database. The results support the suggestion that sequences in the first peak belong to *P. persalinus*, whereas sequences in the second and the third peaks probably derive from bacterial sources. In order to exclude bacterial sequences, two steps were applied: 1) based on their distribution like in *I. multifiliis*20, filtering all scaffolds with GC content higher than 25%; 2) filtering those scaffolds with more than 50% BLASTX hits in the remaining sequences after first step, these most likely being of prokaryote origin. In practice, almost all the filtered scaffolds were removed in the first step, with only two additional short scaffolds (1.5 and 1.6 Kb, respectively) being removed at the second step. This suggests that the GC content is a good index for discriminating ciliate from bacterial sequences. Finally, about 55.5 Mb *P. persalinus* genome sequences with 288 scaffolds were obtained (Table 1). The N50 of the *P. persalinus* genome is about 368 Kb which is comparable to the genome assemblies of *T. thermophila* (521 Kb) and *P. tetraurelia* (413 Kb) (Table 1).

The genome of *P. persalinus* provides evidence of parasitism: 1) its genome size is similar to another ciliate fish pathogen, *I. multifiliis* (47.8 Mb), and is significantly smaller than those of free-living ciliates...
such as *T. thermophila* (103 Mb) and *P. tetraurelia* (72.1 Mb) (Table 1); 2) the *P. persalinus* genome harbors 13,186 predicted protein coding genes, which is about two-fold less than the free-living *T. thermophila* (Table 1); 3) protein domain analysis showed very similar domain composition between *P. persalinus* and *I. multifiliis*, not only in the types but also the number (Figure S1); 4) *Pseudocohnilembus persalinus* and *I. multifiliis* have a similar fraction of parasitic lifestyle-relevant gene families and a relatively high percentage of proteases compared to free-living ciliates (Table S1). Compared to *I. multifiliis*, *P. persalinus* has more proteases especially in the cysteine and serine classes (Table S1) which may be key to various functions of a parasitic lifestyle including immunoevasion, encystment/excystment, ex-sheathing, and cell and tissue invasion21,22. Besides the proteases, 106 *P. persalinus*-specific transporters (Table S2) were found when compared to *T. thermophila*, and these transporters were significantly enriched in the sodium ion, zinc ion, calcium ion, ammonium and phosphate transmembrane transport systems (Figure S2). Some of these transport systems may play important roles in unique aspects of parasite biology. Calcium, for example, is an important factor in the invasion of erythrocytes9, and has been shown to help the secretion of parasite proteins during the invasion process of *Toxoplasma gondii*23,24,25. Thus, calcium transporters may be involved in the invasion process in *P. persalinus*. Furthermore, we compared the gene compositions of *P. persalinus* to the well-known parasitic ciliate *I. multifiliis* and the free-living *T. thermophila*. The results showed that very few unique orthologs (74) shared in *I. multifiliis* and *P. persalinus* (two parasites) are absent in *T. thermophila* (Figure S3), suggesting *P. persalinus* has a distinct mode of parasitism compared to *I. multifiliis*. Scuticociliates infect aquatic organisms opportunistically and occurrences of scuticociliatosis seem to be influenced by environmental conditions, such as temperature and salinity, and weakening of the host due to bacterial infection26. The processes and mechanisms of the transition from free-living to parasitic lifestyle are unknown. By sequencing the genome of *P. persalinus*, this study revealed a number of features associated with parasitism.

**Horizontal gene transfer (HGT) genes may play an important role in the virulence of *P. persalinus***. HGT, the transfer of genetic material between species26, was discovered half of a century ago27, but it is the current wealth of genomic sequence data that is revealing its real impact on evolution. Genes acquired by HGT can sometimes be associated with important evolutionary adaptations, including parasitism and pathogenicity28. The first evidence for the role of HGT in the acquisition of virulence determinants was between pneumococci in infected mice29. Subsequently, a number of studies have reported horizontal transfer of virulence genes (i) between prokaryotic pathogens, (ii) from prokaryotic to eukaryotic pathogens, and (iii) between eukaryotic pathogens30-33. In the obligate ciliate parasite *I. multifiliis*, which harbors an endosymbiotic bacterium, relatively few (10) HGT genes were predicted30. Using a phylogenetic approach (see Materials and Methods section), 74, 5, and 54 putative HGT genes were identified in *T. thermophila*, *I. multifiliis* and *P. persalinus*, respectively. The 54 putative HGT genes in *P. persalinus* were dispersed among 42 different assembled scaffolds (Table 2 and Supplementary Information), and had similar GC contents to the rest of the genes (Fig. 2A). PCR analysis of DNA also verified that these genes are present in the ciliate genome (Figure S4). In addition, 52 of these putative HGT genes were predicted to contain introns (Fig. 2B) which is a main feature of eukaryotic genes. Analysis of transcriptome (RNA-Seq) data reveals that 85% (44) of the putative HGT genes contain at least one intron (Fig. 2C). It is unlikely that the acquisition of introns by these HGT genes was for generating a more complicated proteome through alternative splicing because no alternative splicing was found in the RNA-Seq data. It is possible that the intron gains were the result of adaptation of the transferred gene to its new host cell machinery34. The presence of introns strongly suggest the origin of the putative HGT genes is the *P. persalinus* genome itself rather than bacterial contaminants, and that the HGT events occurred long ago in evolutionary time.

| Species         | *P. persalinus* | *I. multifiliis* | *T. thermophila* | *P. tetraurelia* |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Genome size (Mb)| 55.5            | 47.8            | 103.0           | 72.1            |
| N50 (Kb)        | 368             | 66              | 521             | 413             |
| Scaffold number | 288             | 1375            | 1148            | 697             |
| Longest scaffold (Mb) | 2.0             | 0.4             | 2.2             | 1.0             |
| Sequencing method/platform | Illumina | Sanger/454 | Sanger | Sanger |
| Average GC content | 19%          | 16%             | 22%             | 28%             |
| Gene number     | 13186           | 8062            | 26460           | 39642           |
| Gene density (genes/Mb) | 238          | 169             | 256             | 548             |

**Table 1. Statistics of *Pseudocohnilembus persalinus* genome and comparison to three other sequenced oligohymenophorean ciliates.** *P. persalinus*: *Pseudocohnilembus persalinus*; *I. multifiliis*: *Ichnothyphthirius multifiliis*; *T. thermophila*: *Tetrahymena thermophila*; *P. tetraurelia*: *Paramecium tetraurelia*. 

*P. persalinus* genome itself rather than bacterial contaminants, and that the HGT events occurred long ago in evolutionary time.
| Gene ID       | Best hit in NCBI                                      | E-value  | Best hit species                        | Species category          | Function category by annotation |
|--------------|------------------------------------------------------|----------|-----------------------------------------|---------------------------|---------------------------------|
| PPERSA_00056740 | Ig family protein                                    | 6.00E-31 | Pectobacterium wasabia                  | Proteobacteria             | Cell adhesion                   |
| PPERSA_00029910 | Ig family protein                                    | 8.00E-23 | Dickeya zeae                            | Proteobacteria             | Cell adhesion                   |
| PPERSA_00047700 | Lysophospholipase L1                                 | 8.00E-34 | Amycolatopsis azaura                   | Actinobacteria             | Hemolysis related protein       |
| PPERSA_00098980 | phosphatidylinositol-specific phospholipase C1 like protein | 3.00E-06 | Flavobacterium sp.                     | Bacteroidetes             | Hemolysis related protein       |
| PPERSA_00020800 | phosphatidylinositol-specific phospholipase C1 like protein | 8.00E-10 | Streptomyces thermolacticus            | Actinobacteria             | Hemolysis related protein       |
| PPERSA_00035610 | hemolysin III family channel protein                | 1.00E-14 | Gordonia alkaniivorans                 | Actinobacteria             | Hemolysis related protein       |
| PPERSA_00117390 | hemopexin repeat-containing protein                  | 4.00E-39 | Flavobacterium beibenuense             | Bacteroidetes             | Heme related protein            |
| PPERSA_00079580 | hemopexin repeat-containing protein                  | 5.00E-45 | Flavobacterium beibenuense             | Bacteroidetes             | Heme related protein            |
| PPERSA_00031570 | hemin receptor                                       | 1.00E-06 | Mycobacterium chabuense                | Actinobacteria             | Heme related protein            |
| PPERSA_00130810 | 2OG-Fe(II) oxygenase                                 | 3.00E-143| Aeromonas hydrophila                   | Proteobacteria             | Oxidoreductase                  |
| PPERSA_00076120 | 2OG-Fe(II) oxygenase                                 | 8.00E-47 | Kordia algicida                        | Bacteroidetes             | Oxidoreductase                  |
| PPERSA_00055830 | D-amino acid dehydrogenase small subunit DadA       | 4.00E-23 | Flavimonobacterium sp.                 | Proteobacteria             | Oxidoreductase                  |
| PPERSA_00113410 | FAD-dependent pyridine nucleotide-disulfide oxidoreductase | 0.00E+00 | Methanothermococcus auratus            | Firmicutes                | Oxidoreductase                  |
| PPERSA_00109590 | amine oxidase                                        | 6.00E-37 | Microcystis aeruginosa                 | Cyanobacteria              | Oxidoreductase                  |
| PPERSA_00073400 | NAD-dependent epimerase/dehydratase                 | 3.00E-111 | Nitrosomonas sp.                       | Proteobacteria             | Oxidoreductase                  |
| PPERSA_00069080 | NAD-dependent epimerase/dehydratase                 | 1.00E-69 | Geobacter unamireducens                | Proteobacteria             | Oxidoreductase                  |
| PPERSA_00041150 | arsenate reductase                                   | 2.00E-18 | Pasteurella multocida                  | Proteobacteria             | Oxidoreductase                  |
| PPERSA_00069770 | glutathione S-transferase                            | 7.00E-21 | Lelio oryzae                          | Proteobacteria             | /                               |
| PPERSA_00097300 | major facilitator superfamily MFS_1                 | 2.00E-34 | Clostridium carboxidivorans            | Firmicutes                | /                               |
| PPERSA_00059750 | major facilitator superfamily MFS_1                 | 1.00E-14 | Clostridium drakei                     | Firmicutes                | /                               |
| PPERSA_00084980 | magnesium transporter                                | 2.00E-10 | Fischerella miccule                    | Cyanobacteria              | /                               |
| PPERSA_00107980 | MFS-type transporter                                 | 6.00E-45 | Bacillus massiliisenegalenis           | Firmicutes                | /                               |
| PPERSA_00125230 | Beta-lactamase                                       | 6.00E-37 | Paenibacillus sp.                      | Firmicutes                | /                               |
| PPERSA_00036040 | glyoxalase/bleomycin resistance protein/dioxygenase | 3.00E-36 | Spingopyxis sp.                        | Proteobacteria             | /                               |
| PPERSA_00034640 | thioesterase                                         | 2.00E-14 | Desulfovirile tiedei                  | Proteobacteria             | /                               |
| PPERSA_00036470 | thioesterase                                         | 3.00E-15 | Alcanivorax hungolagensis              | Proteobacteria             | /                               |
| PPERSA_00021250 | beta-lactamase                                       | 5.00E-50 | Cyanothece sp.                        | Cyanobacteria              | /                               |
| PPERSA_00131510 | 3-oxoacyl-ACP synthase                               | 4.00E-13 | Chlorobacillus sp.                     | Proteobacteria             | /                               |
| PPERSA_00117970 | DNA polymerase III subunit epsilon                   | 4.00E-18 | Gammamproteobacteria bacterium SCGC AAA003-E02 | Proteobacteria             | /                               |
| PPERSA_00073390 | 2-amin-3-ketobutyrate CoA ligase                     | 0.00E+00 | Candidatus Cloacamonomas               | Cloacimonetes              | /                               |
| PPERSA_00011350 | formyl transferase domain protein                    | 3.00E-16 | Streptomyces natalensis                | Actinobacteria             | /                               |
| PPERSA_00089970 | inosine/uridine-prefering nucleoside hydrolase       | 7.00E-37 | Legionella wadsworthii                 | Proteobacteria             | /                               |
| PPERSA_00125930 | bifunctional GMP synthase/glutamine amidotransferase protein | 0.00E+00 | Lentisphaera araneosa                  | Chlamydiae                | /                               |
| PPERSA_00035440 | Rhodanese domain protein                             | 5.00E-08 | Flavobacterium sp.                     | Bacteroidetes             | /                               |
| PPERSA_00043810 | rhodanese-related sulfurtransferase                  | 1.00E-117| Endozoicomonas elyscola                | Proteobacteria             | /                               |
| PPERSA_00057430 | SH3 protein, type 3                                  | 7.00E-58 | Gemmata obscuriglobus                  | Planctomycetes             | /                               |
| PPERSA_00050710 | hypothetical protein                                 | 6.00E-12 | Microcilla marina                      | Bacteroidetes             | /                               |
| PPERSA_00036150 | putative phosphatase                                 | 1.00E-12 | Photobacterium halotolerans           | Proteobacteria             | /                               |
| PPERSA_00010290 | PF08002 family protein                               | 3.00E-13 | Bacteroides bacterium                  | Bacteroidetes             | /                               |
| PPERSA_00059290 | photopyrone synthase                                 | 5.00E-14 | Photobacterium luminosus              | Proteobacteria             | /                               |
| PPERSA_00042620 | primase                                              | 5.00E-07 | Methanosarcina acetivorans            | Methanomicrobiota         | /                               |
| PPERSA_0006310 | 3-oxoacyl-ACP synthase                               | 7.00E-11 | Cupriavidus sp.                       | Proteobacteria             | /                               |
| PPERSA_0006300 | 3-oxoacyl-ACP synthase                               | 2.00E-10 | Pseudanabaena sp.                     | Cyanobacteria             | /                               |
| PPERSA_00045220 | 3-oxoacyl-ACP synthase                               | 2.00E-14 | Gramiscella mollensis                  | Fibrobacteres             | /                               |
| PPERSA_00054440 | acid phosphatase                                     | 1.00E-06 | Flavobacterium sp.                     | Bacteroidetes             | /                               |

Continued
The number of HGT genes in *P. persalinus* is similar to that in the free-living species *T. thermophila*, and far higher than the obligate parasite *I. multifiliis*. Therefore, the HGT genes in *P. persalinus* and *T. thermophila* were compared. In *T. thermophila*, 15 HGT genes are homologs of chemotaxis proteins (Table S3) which are related to the movement of an organism in response to a chemical stimulus such as the presence of food\(^\text{37}\). Fourteen genes are tetratricopeptide (TPR) repeat family homologs (Table S3) which have a variety of functions. Six are protein kinases (Table S3), the kinase families being extensively expanded in *Tetrahymena* compared to other organisms\(^\text{12}\). In *P. persalinus*, a gene ontology enrichment

| Gene ID          | Best hit in NCBI                  | E-value   | Best hit species          | Species category | Function category by annotation |
|------------------|-----------------------------------|-----------|---------------------------|------------------|---------------------------------|
| PPERSA_00098990  | acid phosphatase                  | 2.00E-06  | *Flavobacterium* sp.      | Bacteroidetes    | /                               |
| PPERSA_00121720  | rRNA adenine methyltransferase    | 1.00E-42  | *Planktothrix*             | Cyanobacteria    | /                               |
| PPERSA_00076020  | radical SAM domain protein        | 8.00E-81  | *Zavarzinella formosa*    | Planctomycetes   | /                               |
| PPERSA_00125500  | cytidine deaminase                | 8.00E-14  | *Methanoculleus* sp.      | Methanomicrobia  | /                               |
| PPERSA_00032590  | 2-hydroxy-6-oxo-6-phenylhexa-2,4-dienoate hydrolase | 6.00E-09 | *Bordetella avium*        | Proteobacteria   | /                               |
| PPERSA_00117680  | membrane-associated protein in eicosanoid and glutathione metabolism (mapeg) | 1.00E-16 | *Lutimonas meophitis*     | Proteobacteria   | /                               |
| PPERSA_00009640  | TPR repeat                         | 4.00E-12  | *Microscilla marina*      | Bacteroidetes    | /                               |
| PPERSA_00037920  | DTW domain protein                 | 1.00E-30  | *Photobacterium damselae* | Proteobacteria   | /                               |
| PPERSA_00083530  | 2-nitropropane dioxygenase        | 5.00E-63  | *Kyrpidia tusciae*        | Firmicutes       | /                               |

Table 2. 54 HGT genes in *Pseudocohnilembus persalinus* genome.

![A](image1.png)  ![B](image2.png)  ![C](image3.png)

Figure 2. GC content and intron number distribution of the 54 HGT genes. (A) the GC content distribution, similar to the GC content distribution of the assembled scaffolds; (B) the distribution of the predicted intron numbers, only two genes lack introns. (C) a RNA-Seq supported intron-containing HGT gene (PPERSA_00047700). These results suggest that the 54 HGTs belong to the *P. persalinus* genome rather than to bacterial contaminants.

The number of HGT genes in *P. persalinus* is similar to that in the free-living species *T. thermophila*, and far higher than the obligate parasite *I. multifiliis*. Therefore, the HGT genes in *P. persalinus* and *T. thermophila* were compared. In *T. thermophila*, 15 HGT genes are homologs of chemotaxis proteins (Table S3) which are related to the movement of an organism in response to a chemical stimulus such as the presence of food\(^\text{37}\). Fourteen genes are tetratricopeptide (TPR) repeat family homologs (Table S3) which have a variety of functions. Six are protein kinases (Table S3), the kinase families being extensively expanded in *Tetrahymena* compared to other organisms\(^\text{12}\). In *P. persalinus*, a gene ontology enrichment...
analysis suggested that the HGT genes are significantly enriched in functions such as oxidoreductase activity and iron ion binding, which clearly differ from the HGT genes in *T. thermophila* (Figure S5). Therefore, we carefully checked the functional annotations of HGT genes in *P. persalinus* and found a number of HGT genes (approximately 20%) may play important role in its virulence.

**Cell adhesion proteins.** Two of the Ig family of proteins were found in the HGT genes of *P. persalinus*. Domain analysis of these genes showed that both contain cadherin-like domains (Figure S6). Cadherins are a family of transmembrane proteins that play important roles in cell adhesion, forming adherens junctions to bind together cells within tissues. They are dependent on calcium ions (Ca²⁺) to function hence their name. Cadherins have been shown as important adhesins and invasins of pathogenic bacteria. Comparing the transporters between the facultatively parasitic *P. persalinus* and the free-living *T. thermophila* showed that the *P. persalinus*-specific transporter is significantly enriched in calcium ion transport (Figure S2 and Table S2). One of the two Ig proteins HGT genes in *P. persalinus* contain a He_pig domain (PF05345) which contains a conserved core region of about 90 residue repeats found in several haemagglutinins and other cell-surface proteins (http://www.ncbi.nlm.nih.gov/Structure/cdd/cddsrv.cgi?uid=pfam05345), indicating that this gene may contribute to cell (e.g. erythrocyte cell) adhesion (Fig. 3).

**Hemolysis related proteins.** It is well known that many bacterial pathogens induce hemolysis of host erythrocytes. These bacteria could produce proteins, usually called hemolysin, that cause lysis of erythrocytes by destroying their cell membrane. Many proteins have been identified as hemolysins including phospholipase and hemolysin III. Interestingly, three HGT phospholipase-related genes and a hemolysin III homolog were found in the *P. persalinus* genome (Table S2).

Bacterial phospholipases are a large group of enzymes that have a wide range of effects on host cells from minor alterations of cell membrane composition to increased vascular permeability and lethality, even at low concentrations. In the scuticociliate *Uronema marinum*, it has been suggested that phospholipase C could act as virulence factor that serves to actively disrupt host defense mechanisms. Among the three phospholipases in *P. persalinus*, two (PPERSA_00098980 and PPERSA_00002080) were identified as phosphatidylinositol-specific phospholipase C (PI-PLC) and the other one (PPERSA_00047700) was identified as lysophospholipase (Table S2). Phospholipase has been reported to function in erythrocyte membrane modification and hemolysis, for example the PI-PLC, which could release acetylcholinesterase linked to phosphatidylinositol. Thus, phospholipases may help *P. persalinus* utilize the host erythrocytes by disrupting their cell membrane (Fig. 3).

A hemolysin III family protein (Table 2, PPERSA_00035610) was identified among the HGT genes of *P. persalinus*. This gene is located in an assembled scaffold with 1.2 Mb in length and contains a RNA-Seq data-supported intron (Figure S7A). The coded protein contains a HlyIII domain (PF03006) and six
transmembrane helices (Figure S7B). It has been shown that hemolysin III produced by the bacteria *Bacillus cereus* and *Vibrio vulnificus* is capable of hemolytic activity. The *P. salinarum* hemolysin III gene closely resembles its *B. cereus* homolog (Figure S7C), especially in the functional domain region, suggesting that the protein for which this gene codes may play a role in the lysis of host erythrocytes (Fig. 3).

**Heme related proteins.** Hemolysis is the rupturing of the erythrocyte cell membrane and the release of its cytoplasm into surrounding tissue. By this process pathogens can acquire the erythrocyte cell contents and utilize it for their own metabolism. Erythrocyte cytoplasm is rich in hemoglobin which includes the iron-containing heme whose function is to bind and transport oxygen. Hemoglobin can be digested by a series of protease enzymes, releasing its amino acids and heme. Amino acids can be directly intercepted by the pathogen, whereas free heme generates oxidative stress known to damage cells if not utilized or transformed. In malaria, merozoites of *Plasmodium* invade erythrocytes, ingest host hemoglobin enclosing it in an acidic food vacuole and digest it using proteases. The released heme is then incorporated into haemoglobin. Due to its abundance in the host, heme is a valuable source of iron for invading micro-organisms during hemolysis, and makes the host dramatically more susceptible to infections and their complications. *Pseudocohniilembs salinarum* has two hemopexin repeat protein homologs (PPERSA_00117390 and PPERSA_00079580) that appear to be of bacterial origin acquired by HGT (Table 2). These proteins are reported to have a high binding affinity for heme and are probably heme carriers. The existence of these two HGT genes suggests that *P. salinarum* can uptake and utilize heme as a way to acquire and use iron (Fig. 3).

For many ciliates, e.g. *Tetrahymena*, the inclusion of inorganic iron salts in a culture medium produces a dramatic acceleration of growth and marked alterations in metabolism. Iron supplementation could be used in heme synthesis and lead to an increased concentration or activity of certain heme enzymes, particularly in the electron transport chain which plays an important role in ATP synthesis thus stimulating cell growth. In *P. salinarum*, the same heme de novo synthesis pathway was found as in *Tetrahymena* (Figure S8), suggesting that *P. salinarum* may also synthesize heme from iron in order to enhance growth. Therefore, it is reasonable to speculate that direct uptake of heme from the host could stimulate the reproduction of *P. salinarum* during infection (Fig. 3).

Hemin is the Fe(III) oxidation product of heme. An excess of heme can interact with the cell membrane resulting in formation of reactive oxygen species (oxidative stress) and causing cellular injury. For the host, the generation of hemin is a double-edged sword since it not only lyses pathogens, but also induces hemolysis of erythrocytes. Bacteria such as *Yersinia enterocolitica* have evolved heme utilization systems that enable them to acquire iron by intercepting hemin using heme receptor proteins. A heme receptor gene of bacterial origin (PPERSA_00031570) has been acquired by *P. salinarum* (Table 2), indicating that the ciliate has the ability to use heme. In addition to being a source of iron, the binding of free hemin by the bacterial receptor protein may help to reduce the oxidative stress for the ciliate.

Heme, or its Fe(III) oxidation product hemin, is catalytically broken down by heme oxygenase to carbon monoxide, bilirubin and iron. The iron can then be used by non-heme iron enzymes or participate in the de novo heme synthesis pathway. However, no homolog of heme oxygenase was found in *P. salinarum*, suggesting the presence of an alternative heme utilization system. In the bacterium *Ralstonia metallidurans* CH34, for example, some heme-related proteins share a transcription factor binding site (potential operon) with 2OG-Fe(II) oxygenase (http://regprecise.lbl.gov/RegPrecise/regulon.jsp?regulon_id=9937), indicating that 2OG-Fe(II) oxygenase may be involved in the heme utilization process, although its function was not determined. In *P. salinarum*, two 2OG-Fe(II) oxygenase (PPERSA_00130810 and PPERSA_00076120) were found as HGT genes (Table 2), raising the possibility that 2OG-Fe(II) oxygenase could function as heme oxygenase, cleaving the hemin ring to release the iron (Fig. 3). Thus, it appears that *P. salinarum* acquired by HGT almost the whole pathway for hemolysis and the utilization of heme.

One of the most salient clinical manifestations of scuticociliatosis is haemorrhagic lesions. Histopathological observations typically show many scuticociliates in the blood vessels, gills, fins, skin muscle, brain and lamina propria of the digestive tract, accompanied by necrosis and haemorrhages. Like *Uronema marinum* which destroys host tissue by proteases, *P. salinarum* may also utilize proteases to break the skin–blood barrier and gain entry to internal organs. Virulence HGT genes may thus contribute to the subsequent destruction of red blood cells and the acquisition of host-derived nutrients and energy for ciliate cell proliferation.

It has been reported that scuticociliatosis often accompanies bacterial disease (e.g. vibriosis), and the increased bacterial load probably helps ciliates to thrive and proliferate during the initial phase of infection. Therefore it can be speculated that the synergistic invasion by pathogenic bacteria and scuticociliates, and the presence of both in a shared environment, may provide the opportunity for the transfer of genetic materials from the former to the latter. Alternatively, many ciliates harbor bacterial symbionts which may also provide the conditions for HGT. Evidence for the presence of endosymbiotic bacteria in *P. salinarum* comes from the preliminary genome assembly, the second and third peaks with 41% and 67% GC (Fig. 1) representing two bacterial species. The homology search results showed these could be species closely related to *Pseudoalteromonas* and *Halomonas*, respectively. They are very likely bacterial symbionts harbored in *P. salinarum* because sequences of *Escherichia coli* DH5 alpha, the only food organism supplied to *P. salinarum* cultured in laboratory, were not found. The 54 HGT genes in
P. persalinus were not, however, included in these two peaks. Although homology searches showed that
these 54 HGT genes do not have an enriched bacterial source, many of the best homologs occur in two
large bacterial classes: Actinobacteria and Gammaproteobacteria (Table 2). Therefore, it is more likely
that the HGT genes in P. persalinus have multiple independent origins.

Based on infection experiments, some researchers have concluded that the scuticociliates P. persalinus,
P. hargisi and U. marín are not primary pathogens of oysters, rather they are non-pathogenic,
free-living, bacteriophagous and/or saprophagous ciliates that opportunistically attached to lesions on the
host that are originally produced by bacterial infection or some other cause61. However, the identification
here of virulence HGT genes in P. persalinus provides evidence that there is a molecular basis for its
pathogenicity. Recent attempts to develop vaccines targeting antigens such as those responsible for host
cell immobilization, proton-translocating inorganic pyrophosphatases, cathepsin L-like cysteine protease,
etc. have met with limited success because of different levels of virulence and serotype-specific protection
among species/strains of pathogens64–67. It is anticipated that the virulence HGT genes identified here will
help us to gain a better understanding of the pathology of scuticociliatosis and provide potential antigen
candidates for the development of anti-scuticociliatosis vaccines.

Conclusions
In summary, we report the genome of scuticociliate Pseudocohnilembus persalinus, the first marine cil-
iate genome as far as we know. The genome of P. persalinus genome is 55.5 Mb, i.e., about half the
size of the model free-living ciliate Tetrahymena thermophila. The P. persalinus genome harbors many
prokaryote-derived horizontally transferred genes; function analysis showed that many of these HGT
genesis are potential virulence factors. These findings help to increase our knowledge and understanding
of the mechanism of the common fish disease, scuticociliatosis.

Methods
Pseudocohnilembus persalinus culture, total DNA and total RNA extraction. Pseudocohnilembus
persalinus was isolated from water in a shrimp-farming pond (36°08′N,120°43′E; water tem-
perature 27°C; salinity 20%; pH 7.5) in Qingdao, China68. The species was identified by its morphol-
ogy, morphogenesis69 and 18S rDNA marker (Figure S9). In order to obtain sufficient DNA and RNA
material for sequencing, P. persalinus cells were cultured in the laboratory using sterilized sea water with
Escherichia coli DH5 alpha as a food source. Contamination by bacteria was prevented by treating the
cell cultures with lysozyme (200 μg/ml for 2 hours at 28 °C) before the DNA and RNA extraction. The
total DNA was extracted using the method described for Tetrahymena69, and the total RNA was extracted
using the RNeasy Protect Cell Mini Kit (Qiagen, Valencia, CA) according to protocol in TetraFGD70.

Pseudocohnilembus persalinus genome and transcriptome sequencing. The P. persalinus
genome was sequenced using the Illumina platform. Paired-end (about 190bp insert size) and mate-pair
(about 2 Kb insert size) libraries were constructed and sequenced using the standard protocol of Illumina
(https://icom.illumina.com/). Briefly, genomic DNA was fragmented and fragments of appropriate size
(see above) were selected. For mate-pair library construction, fragment ends were biotinylated and circu-
larized, and the fragments were then enriched using biotin. Fragment ends were then repaired, A-tailed
and ligated with sequencing adaptors. Adapter-ligated fragments were PCR amplified using Phusion
polymerase, denatured with sodium hydroxide and diluted in hybridization buffer. The prepared libraries
were loaded onto the flowcell and sequenced.

For transcriptome sequencing, Poly-A mRNAs were isolated using Dynal magnetic beads (Invitrogen)
and fragmented by heating to 94°C. First strand cDNAs were synthesized with reverse transcriptase
and random hexamer primers, and then the second strands were synthesized with DNA polymerase
and random hexamer primers. Double strand cDNAs were end-repaired and a single adenosine moi-
ety was added. Illumina adapters were ligated and gel-electrophoresis was used to select DNA frag-
ments about 200bp size. Libraries were PCR-amplified using Phusion polymerase. Cluster formation,
primer hybridization and pair-end sequencing were performed using proprietary reagents according to
manufacturer-recommended protocols (https://icom.illumina.com/).

Genome assembly and bacterial contamination exclusion. The paired-end reads and mate-paired
reads of the Illumina sequencing were assembled using SOAPdenovo software71, which uses the de Bruijn
graph data structure to construct contigs. A series of K-mer values (from 33 to 79) were used to assem-
bly the P. persalinus genome, and the assembly with the longest N50 length was chosen by deleting
scaffolds shorter than 1 Kb. Bacterial contaminants were excluded in two stages, sample preparation (see
above) and bioinformatics analysis. In the bioinformatics analysis stage, bacterial contaminants were first
excluded using the GC content, any scaffolds with a GC content more than 25% being discarded. The
remaining scaffolds were then BLAST searched against the NCBI non-redundant protein database; any
scaffolds with more than 50% hits belonging to the prokaryotes were excluded. The remaining scaffolds
were regarded as the P. persalinus genome assembly sequences.
**Gene prediction and annotation.** Using the RNA-Seq data, transcripts were both de novo assembled using Trinity, and reference-guided assembled using the TopHat and Cufflinks pipeline. A combination of de novo and reference-guided assembled transcripts were validated by aligning the putative transcripts onto the assembled genome using PASA. A set of the so-called best models generated by PASA was used to train the gene prediction software Augustus and GlimmerHMM. The training parameters were then used by the two programs to de novo predict the gene models. The Augustus software could accept the cDNA or protein evidence, therefore the assembled transcripts were also used as the cDNA evidence for Augustus. Finally, an integrated set of gene models was created using Evidence Modeler by merging all of the predicted gene models.

Homologs of *P. persalinus* genes were BLAST searched against the NCBI non-redundant protein database. The KEGG pathway information was annotated using the KAAS server. Protein domain information was annotated using the Pfam database. Gene ontology (GO) information was annotated using the Goanna server. For each gene set, GO enrichment analysis was also carried out using BinGO. FDR correction was used to control the false positive rate. If a GO term in a test gene set showed a corrected p value less than 0.05 compared with the reference set (all the GO annotated genes), the GO term (function) was determined to be significantly overrepresented in the test gene set. To annotate the proteases, a batch BLAST against the MEROPS database using all predicted genes. Genes with BLAST hit E-values less than 1e-10 were regarded as proteases, and the best hit in the MEROPS database was used to assign the protease class. To annotate the membrane transporters, all the genes were BLAST searched against the TCDB transporters database, and the transmembrane helices were predicted using SCAMP, Toppred and TMHMM. A gene was regarded as a membrane transporter if it had a BLAST hit in the TCDB transporters database with an E-value less than 1e-03 and showed at least one transmembrane helix in all three programs.

**Bacterial horizontal transferred genes identification.** *Pseudocohniembus persalinus* HGT genes were identified by two steps, similar to the strategy used in Ricard et al. Firstly, similarity searches were performed to screen the potential prokaryotic origin genes by using a BLASTP search against the NCBI non-redundant database (Figure S10). In this step, the *E*-value 1E-05 was used as the cutoff, and if a *P. persalinus* gene had a best hit belonging to the prokaryotes, it was regarded as a candidate gene. Phylogenetic approaches are widely used to identify HGT genes, so we also employed phylogenetic analyses to further identify the *P. persalinus* HGT genes based on the screened candidates. All candidate genes retrieved from the first step were BLASTP searched (E-value: 1E-05) against both the prokaryote and eukaryote protein databases (generated from the Refseq data: ftp://ftp.ncbi.nlm.nih.gov/refseq/release/) in order to retrieve homologs in both eukaryotes and prokaryotes. For a protein in *P. persalinus*, if there were homologs (E-value: 1E-05) in both domains, the sequences were retrieved in order to construct phylogenetic trees. Two methods (programs) were used, namely FASTTREE and PHYML. For FASTTREE, all the homologs with E-value less than 1E-05 were used to construct the tree; for PHYML, only the top ten homologs (if present) were used to construct the tree. Sequences alignments were performed using MUSCLE. A gene clustered in the prokaryotic clade which had a eukaryotic outgroup was accepted as an HGT gene, a technique now widely used to identify HGT genes. As shown in Figure S11, only gene in *P. persalinus* with this kind of tree topology was considered an HGT gene. If a *P. persalinus* gene only has homologs with E-values less than 1E-05 in prokaryotes, the phylogenetic analysis could not work. HGTs were determined if there were at least 5 prokaryotic hits and the E-value of the best hit in prokaryotes and eukaryotes differed by more than 5 orders of magnitude (i.e., the E-value of best hit in eukaryotes will be larger than 1 if the best prokaryotic hit E-value is 1E-05). Some HGT genes may have diverged significantly after the HGT event. For example, gene PPERSA_00031570 with an E-value 1E-06 to its best BLAST hit (prokaryotic protein), has a bacterial-like globin (Pfam domain: PF01152) and is therefore highly likely to be of prokaryotes origin. In such cases, an E-value 1E-05 was used if no homolog was found in the eukaryotes. In addition, to confirm the existence of HGT genes, PCR analysis was performed for 20 of 54 HGT genes identified by the bioinformatics (for primers, see Table S4), 100% of which were verified.

**Data access.** This Whole Genome Shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession LDAU00000000. The version described in this paper is version LDAU01000000. The raw genome sequences reads have been deposited in Sequence Read Archive (SRA) under accession SRX883501 and SRX883503. Transcriptome data has also been deposited in SRA under accession SRX849480.

**References**

1. Noga, E. J. *Fish disease: diagnosis and treatment* (John Wiley & Sons, 2010).
2. Cheung, P. I., Nigrelli, R. F. & Ruggieri, G. D. Studies on the Morphology of Uronema-Marinum Dujardin (Ciliata, Uronematidae) with a Description of the Histopathology of the Infection in Marine Fishes. *Journal of Fish Diseases* 3, 295–303 (1980).
3. Kim, S. M., Cho, J. B., Kim, S. K., Nam, Y. K. & Kim, K. H. Occurrence of scuticociliatosis in olive flounder Paralichthys olivaceus by Phialidium dicentrarchii (Ciliophora : Scuticociliatida). *Diseases of Aquatic Organisms* 62, 233–238 (2004).
4. Lee, B. Y., Kim, Y. C. & Park, M. S. Morphology and biology of parasite responsible for scuticociliatosis of cultured olive flounder Paralichthys olivaceus. *Diseases of Aquatic Organisms* 47, 49–55 (2001).
47. Toutant, J. P., Roberts, W. L., Murray, N. R. & Rosenberry, T. L. Conversion of human erythrocyte acetylcholinesterase from an amphiphilic to a hydrophobic form by phosphatidylinositol-specific phospholipase C and serum phospholipase D. *European Journal of Biochemistry* **180**, 503–508 (1989).

48. Chen, Y. C., Chang, M. C., Chuang, Y. C. & Jeang, C. L. Characterization and virulence of hemolysin III from Vibrio vulniﬁcus. *Current Microbiology* **49**, 175–179 (2004).

49. Hendrickx, W. A. Hemoglobin - Structure, Function, Evolution, and Pathology - Dickerson,Re, Geis,Scientific Reports **304**, 193–198 (1983).

50. Piccard, H., Van Den Steen, P. E. & Opdenakker, G. Hemopexin domains as multifunctional liganding modules in matrix metal loproteinas and other proteins. *Journal of Leukocyte Biology* **81**, 870–892 (2007).

51. Aikawa, M., Huff, C. G. & Sprinz, H. Comparative Feeding Mechanisms of Avian and Primate Malarial Parasites. *Military Medicine* **131**, 969–& (1966).

52. Banerjee, R. et al. Four plasmepsins are active in the Plasmodium falciparum food vacuole, including a protease with an active-site histidine. *Science* **308**, 1639–1642 (2005).

53. Egan, T. J. et al. Fate of haem iron in the malaria parasite Plasmodium falciparum. *Biochemical Journal* **365**, 343–347 (2002).

54. Chipman, A. D. et al. The First Myriapod Genome Sequence Reveals Conserved Arthropod Gene Content and Genome Organisation in the Centipede Strigamia maritima. *Plos Biology* **12** (2014).

55. Shug, A. L., Elson, C. & Shrago, E. Effect of iron on growth, cytochromes, glycogen and fatty acids of *Tetrahymena pyriformis*. *J Nutr* **99**, 579–86 (1969).

56. Balla, J. et al. Endothelial-Cell Heme Uptake from Heme-Proteins - Induction of Sensitization and Desensitization to Oxidant Damage. *Proceedings of the National Academy of Sciences of the United States of America* **90**, 9285–9289 (1993).

57. Orjih, A. U., Banyal, H. S., Chevli, R. & Fitch, C. D. Hemin Lyses Malaria Parasites. *Science* **214**, 667–669 (1981).

58. Li, S. D., Su, Y. D., Li, M. & Zou, C. G. Hemin-mediated hemolysis in erythrocytes: Effects of ascorbic acid and glutathione. *Acta Biochimica Et Biophysica Sinica* **38**, 63–69 (2006).

59. Soojikjovic, I. & Hanke, K. Hemin Uptake System of Yersinia-Enterocolitica - Similarities with Other Tonn-Dependent Systems in Gram-Negative Bacteria. *Embo Journal* **11**, 4359–4367 (1992).

60. Azad, I., Al-Marzouk, A., James, C., Almatar, S. & Al-Gharabally, H. Scuticociliatosis-associated mortalities and histopathology of natural infection in cultured silver pomfret (*Pampus argenteus* Euphrasen) in Kuwait. *Aquaculture* **262**, 202–210 (2007).

61. Song, J. Y. et al. Pathogenicity of *Miamisensis* avidis (syn. *Philasterides denticranchi*), *Pseudocohnilembus persalinus*, *Pseudocohnilembus bargisi* and *Uronema marina* (Ciliophora, Scuticociliatida). *Diseases of Aquatic Organisms* **83**, 133–43 (2009).

62. Lee, E. H., Kim, C. S., Cho, J. B., Ahn, K. J. & Kim, K. H. Measurement of protease activity of live *Uronema marina* (Ciliata: Scuticociliatida) by fluorescence polarization. *Dis Aquat Organ* **54**, 85–8 (2003).

63. Görtz, H.-D. Symbiotic associations between ciliates and prokaryotes. in *Nucleic Acids Research* **34**, 2878–2879 (2004).

64. Mallo, N., Lamas, J., Pizzonzo, C. & Leiro, J. M. Presence of a plant-like proton-translocating pyrophosphatase in a scuticociliate parasite and its role as a possible drug target. *Parasitology*: 1–14 (2014).

65. Shin, S. P. et al. Expression and characterization of cathepsin L-like cysteine protease from *Philasterides denticranchi*. *Parasitol Int* **63**, 559–65 (2014).

66. Leon-Rodriguez, L., Lutzardo-Alvarez, A., Blanco-Mendez, J., Lamas, J. & Leiro, J. Biodegradable microparticles covalently linked to surface antigens of the scuticociliate parasite *P. denticranchi* promote innate immune responses in vitro. *Fish Shellfish Immunol* **34**, 236–43 (2013).

67. Leon-Rodriguez, L., Lutzardo-Alvarez, A., Blanco-Mendez, J., Lamas, J. & Leiro, J. Expression and characterization of cathepsin L-like cysteine protease from Philasterides denticranchi. *Parasitol Int* **63**, 359–65 (2014).

68. Leon-Rodriguez, L., Lutzardo-Alvarez, A., Blanco-Mendez, J., Lamas, J. & Leiro, J. A vaccine based on biodegradable microparticles induces protective immunity against *scuticociliatosis* without producing side effects in turbot. *Fish Shellfish Immunol* **33**, 21–7 (2012).

69. Pan, X., Ma, H., Shao, C., Lin, X. & Hu, X. Stomatogenesis and morphological redescription of *Pseudocohnilembus persalinus* (Ciliophora: Scuticociliatida). *Acta Hydrobiologica Sinica* **36** (483–494) (2012).

70. Savilev, S. V. PCR-based detection of a rare linear DNA in cell culture. *Bio Protoc* **4**, 70–80 (2002).

71. Xiong, J. et al. Thrombopoiesis and thrombolysis in *Drosophila* model hemostatic system. *Proceedings of the National Academy of Sciences (Oxford)* **2013**, bat008 (2013).

72. Xie, Y. et al. SOAPdenovo-Trans: de novo transcriptome assembly with short RNA-Seq reads. *Bioinformatics* (2014).

73. Grabherr, M. G. et al. Full-length transcriptome assembly from RNA-Seq data without a reference genome. *Nature Biotechnology* **29**, 644–U130 (2011).

74. Trapnell, C., Pachter, L. & Salzberg, S. L. TopHat: discovering splice junctions with RNA-Seq. *Bioinformatics* **25**, 1105–1111 (2009).

75. Trapnell, C. et al. Differential gene and transcript expression analysis of RNA-seq experiments with TopHat and Cufflinks. *Nature Protocols* **7**, 562–578 (2012).

76. Haas, B. J. et al. Improving the Arabidopsis gene annotation using maximal transcript alignment assemblies. *Nucleic Acids Research* **31**, 5654–5666 (2003).

77. Stanke, M., Diekhans, M., Baertsch, R. & Haussler, D. Using native and syntenically mapped cDNA alignments to improve de novo gene finding. *Bioinformatics* **24**, 637–644 (2008).

78. Majaros, W. H., Pertea, M. & Salzberg, S. L. TigrScan and GlimmerHMM: two open source ab initio eukaryotic gene-finders. *Bioinformatics* **20**, 2878–2879 (2004).

79. Haas, B. J. et al. Automated eukaryotic gene structure annotation using EVidenceModeler and the program to assemble spliced alignments. *Genome Biology* **9** (2008).

80. Bateman, A. et al. The Piam protein families database. *Nucleic Acids Research* **28**, 263–6 (2000).

81. Orjih, A. U., Banyal, H. S., Chevli, R. & Fitch, C. D. Hemin Lyses Malaria Parasites. *Science* **214**, 667–669 (1981).

82. Bernsel, A. et al. Prediction of membrane-protein topology from first principles. *Proceedings of the National Academy of Sciences of the United States of America* **105**, 7177–7181 (2008).

83. Vonheijne, G. Membrane-Protein Structure Prediction - Hydrophobicity Analysis and the Positive-inside Rule. *Journal of Molecular Biology* **225**, 487–494 (1992).

84. Sonnhammer, E. L., Von Heijne, G. & Krogh, A. A hidden Markov model for predicting transmembrane helices in protein sequences. *Proc Intl Conf Intent Syst Mol Biol* **6**, 175–82 (1998).

85. Ricard, G. et al. Horizontal gene transfer from bacteria to rumen ciliates indicates adaptation to their anaerobic, carbohydrates-rich environment. *Bmc Genomics* **7** (2006).

86. Li, Z. W., Shen, Y. H., Xiang, Z. H. & Zhang, Z. Pathogen-origin horizontally transferred genes contribute to the evolution of Lepidopteran insects. *Bmc Evolutionary Biology* **11** (2011).
87. Stanhope, M. J. et al. Phylogenetic analyses do not support horizontal gene transfers from bacteria to vertebrates. Nature 411, 940–944 (2001).
88. Price, M. N., Dehal, P. S. & Arkin, A. P. FastTree 2—Approximately Maximum-Likelihood Trees for Large Alignments. Plos One 5 (2010).
89. Guindon, S., Dufayard, J. F., Hordijk, W., Lefort, V. & Gascuel, O. PhyML: Fast and Accurate Phylogeny Reconstruction by Maximum Likelihood. Infection Genetics and Evolution 9, 384–385 (2009).
90. Edgar, R. C. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Research 32, 1792–1797 (2004).
91. Zhao, H. et al. Host-to-Pathogen Gene Transfer Facilitated Infection of Insects by a Pathogenic Fungus. Plos Pathogens 10 (2014).

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Author Contributions
W.M. and J.X. designed the project. M.T., X.P. and J.X. collected samples and prepared DNA and RNA. J.X., G.W., J.C., C.J. and D.Y. performed the analyses. J.X., W.M. and A.W. wrote the manuscript. All authors read and approved the final manuscript.

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