Two functionally distinct heme/iron transport systems are virulence determinants of the fish pathogen *Flavobacterium psychrophilum*

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**ABSTRACT**

Bacterial pathogens have a critical impact on aquaculture, a sector that accounts for half of the human fish consumption. *Flavobacterium psychrophilum* (phylum *Bacteroidetes*) is responsible for bacterial cold-water disease in salmonids worldwide. The molecular factors involved in host invasion, colonization and haemorrhagic septicaemia are mostly unknown. In this study, we identified two new TonB-dependent receptors, HfpR and BfpR, that are required for adaptation to iron conditions encountered during infection and for virulence in rainbow trout. Transcriptional analyses revealed that their expression is tightly controlled and upregulated under specific iron sources and concentrations. Characterization of deletion mutants showed that they act without redundancy: BfpR is required for optimal growth in the presence of high haemoglobin level, while HfpR confers the capacity to acquire nutrient iron from haem or haemoglobin under iron scarcity. The gene *hfpY*, co-transcribed with *hfpR*, encodes a protein related to the HmuY family. We demonstrated that HfpY binds haem and contributes significantly to host colonization and disease severity. Overall, these results are consistent with a model in which both BfpR and Hfp systems promote haem uptake and respond to distinct signals to adapt iron acquisition to the different stages of pathogenesis. Our findings give insight into the molecular basis of pathogenicity of a serious pathogen belonging to the understudied family *Flavobacteriaceae* and point to the newly identified haem receptors as promising targets for antibacterial development.

**Introduction**

Aquaculture is a rapidly growing sector which now produces more than half of the human fish consumption in the world [1]. However, its sustainable development faces the major challenge of controlling infectious diseases caused by poorly known aquatic pathogens. These include several Gram-negative bacteria belonging to the family *Flavobacteriaceae* (phylum *Bacteroidetes*), of which a majority of members are commensals and saprophytes commonly isolated from environmental sources such as soil, sediments and water [2]. Among them, *Flavobacterium psychrophilum* infects salmonids in freshwater environments and is responsible for diseases known as rainbow trout fry syndrome and bacterial cold-water disease which constitute a major health issue for the fish farming industry worldwide [3–8]. *F. psychrophilum* outbreaks cause complex rearing issues and substantial economic losses, impact animal welfare and due to repeated use of antibiotics the environment. A better understanding of the pathogenesis and biology of this pathogen is needed to enhance the development of alternative control strategies, such as efficient vaccines, bacteriophages therapy and selection of resistant fish lines [9–12].

Juvenile rainbow trout (*Oncorhynchus mykiss*) are particularly susceptible. In fry, the disease results in septicaemia leading to mortalities rising up to 70% without appropriate antibiotic treatment. Infected fish present signs of tissue erosion and necrotic lesions, typically in the jaw, the caudal peduncle or the caudal fin [13]. The bacterium is mainly found in the damaged skin and muscular tissues surrounding the lesions, and in the lymphoid organs, particularly in the spleen, which exhibits haemorrhages and hypertrophy and harbours a large number of bacteria [14–16]. *F. psychrophilum* cells are highly proteolytic, cytotoxic and adhesive [17–20]. The Type IX secretion system (T9SS) is required for pathogenicity and secretes several adhesins, peptidases and other uncharacterized proteins that likely function in adhesion to host cells, tissue degradation and pathogen spreading [21–23].

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Within the host, *F. psychrophilum* cells face a strong inflammatory response [24,25]. Host colonization relies on the resistance to stresses and the ability to efficiently retrieve nutrients such as amino acids, vitamins and essential metals. In vertebrates, iron is mainly sequestered in iron- or haem-binding proteins preventing toxic effects of free iron and limiting accessibility for bacterial growth. Pathogens have developed sophisticated mechanisms to fulfil their iron needs from various sources [26]. In Gram-negative bacteria, many of the iron acquisition systems rely on TonB-dependent receptors (TBDR) that mediate the active uptake of specific substrates across the outer membrane. TBDRs are β-barrel outer membrane proteins forming pores that are occluded by a globular periplasmic domain (plug) in the absence of substrate. Substrate transport is energized by the inner membrane TonB-ExbBD complex that transduces energy derived from the proton motive force [27]. A large variety of iron uptake TBDRs have been described, involving either the direct binding of host iron- or haem-binding proteins by the receptor, or the indirect substrate capture by high-affinity siderophores or hemophores. Expression of these genes is typically tightly regulated in response to iron availability in order to meet bacterial needs for iron while avoiding toxicity of excessive intracellular iron concentrations [28].

Despite the importance of iron homeostasis, the genes involved in iron uptake remain uncharacterized in *F. psychrophilum*. In vitro, growth of the bacterium is strongly inhibited by addition of an iron chelator, but can be restored by the supply of haemoglobin or transferrin, suggesting that the bacterium is able to uptake iron from these compounds [29]. Production of siderophores, haemolysis activity and haemoglobin consumption have been reported [18,21,29]. In addition, the inactivation of *exbD2*, a subunit of the TonB system, results in impaired growth under iron restricted conditions [30], pointing out the probable existence of TonB-dependent iron uptake systems in *F. psychrophilum*.

Here, we took advantage of a large *in vitro* condition-dependent transcriptome dataset [31] to identify iron-regulated TBDR encoding genes. By combining reverse genetics, *in vitro* phenotyping of deletion mutants and experimental infection in rainbow trout, we demonstrate the importance of two haem/haemoglobin receptors for the virulence of *F. psychrophilum*. The first receptor, that we named BfpR, acts under high-level haemoglobin conditions. The second receptor, HfpR, acts under iron-restricted conditions in concert with its cognate hemophore HfpY, a new member of the HmuY family widespread in the phylum *Bacteroidetes*.

**Materials and methods**

**Bacterial strains and growth conditions**

The strains used in this study are listed in Table 1 in Supporting Information. This study was performed using THCO2-90, a model strain for *F. psychrophilum* genetic manipulation [17,32]. *F. psychrophilum* was routinely grown aerobically in tryptone yeast extract salts (TYE) broth [0.4% (w/v) tryptone, 0.04% yeast extract, 0.05% (w/v) MgSO4 7 H2O, 0.02% (w/v) CaCl2 2 H2O, 0.05% (w/v) D-glucose, pH 7.2] or on TYES agar. When specified, media was supplemented with 5% foetal calf serum (FCS) for optimal growth. Growth experiments in liquid culture were carried out at 200 rpm and 18°C, as follows: strains were streaked on TYES FCS agar from –80°C freezer stock and incubated 4 days at 18°C, a pre-culture was then prepared by inoculation of one colony into 5 ml of TYE broth and incubation during 36 h. Cultures were performed by inoculation of 10 ml of broth medium at an OD600 of 0.02 with the pre-culture and growth was evaluated by measuring OD600 using a Biochrom WPA CO8000 spectrophotometer. Iron-depleted cultures were performed in 10 ml of TYE broth supplemented with 20 µM ethylenediamine-N,N’-bis (2-hydroxyphenylacetic acid) (EDDHA) and 1.25 µM haemoglobin, 5 µM haemin or 5 µM FeCl3 when specified. Stock solutions were prepared as follows: 2.5 mM lyophilized haemoglobin, 5 µM haemin or 5 µM FeCl3 when specified. Stock solutions were prepared as follows: 2.5 mM lyophilized haemoglobin (Sigma #H2500) in 0.1 M Tris 0.14 M NaCl, 10 mM haemin chloride (Fe-PPIX) (Frontier Scientific) in 50 mM NaOH, 50 mM FeCl3 (Merck) in water and 25 mM EDDHA (Sigma) in 50 mM NaOH. Haemoglobin-enriched cultures were performed by inoculation of the pre-culture at an OD600 of 0.005 into 5 ml of TYE broth supplemented with 1% haemoglobin (w/v) prepared from a fresh 10% haemoglobin (Sigma #08449) stock solution. Growth curves were determined by enumeration of colony-forming units (CFU) by plating serial dilutions of the cultures on TYES FCS agar plates. The elastin degradation assay was performed using TYE agar supplemented with 0.75% (w/v) elastin from bovine neck ligament (Sigma, E1625) as previously described [33]. *E. coli* strains were grown at 37°C in Luria Bertani broth or supplemented with 15 g/l of agar.

**Bioinformatic analysis**

Comparison of the gene content between strains was performed on a selection of 22 genomes using the web
interface MicroScope [34]. The TBDR-encoding genes were identified using Interpro predictions of the TonB-dependent receptor family (IPR000531) or the highly conserved structural beta-barrel (IPR036942) and plug (IPR037066) domains. Orthologs were identified using BlastP Bidirectional Best Hit and the MicroScope default parameters (i.e. >80% protein identity, >80% coverage). The presence of truncated TBDR genes was verified by PCR and Sanger sequencing. The structure-assisted multiple sequence alignments of HfpR (THC0290_1814) and BfpR (THC0290_1679) with other heme/haemoglobin receptors and of HfpY (THC0290_1812) with other HmuY-like proteins were constructed with PROMALS3D [35] webserver.

**Plasmid construction and conjugative transfer into F. psychrophilum**

Plasmids (T1 in Supporting Information) were engineered by the Gibson assembly method using the NEBuilder® HiFi DNA Assembly Master Mix (New England Biolabs) with appropriate primers (T2 in Supporting Information). Plasmids were constructed in E. coli DH5α Z1, verified by PCR and DNA sequencing, transferred to E. coli MFDPir and subsequently introduced into F. psychrophilum OSU THCO2-90 by conjugation as previously described [31]. For selective growth of E. coli strains carrying plasmids, transformants were selected with 100 µg/ml of ampicillin. Cultures of E. coli MFDPir were supplemented with 0.3 mM dianinopimelic acid (Sigma-Aldrich Co.). Selection of F. psychrophilum transconjugants was carried out with 50 µg/ml of gentamycin and 10 µg/ml of erythromycin for the pCPGm1− and pYT313-derivative plasmids, respectively.

**Construction of transcriptional reporter plasmids and promoter activity monitoring**

pCPGm1−P<sub>les</sub>-mCh derivative plasmids carrying the promoter region upstream of hfpR (240 bp) or bfpR (various length: 306, 609 and 950 bp) were constructed as previously described [31]. Briefly, the vector fragment was amplified using primers TRO460/TRO461 with pCPGm1−P<sub>les</sub>-mCh as matrix and the promoter region using strain OSU THCO2-90 chromosomal DNA and either primers TRO462/TRO463 resulting in P<sub>hfpR</sub>-mCh fusion (pGi48), TRO464/TRO465 resulting in P<sub>bfpR,1</sub>-mCh (306 bp, pGi49), TRO789/TRO790 resulting in P<sub>bfpR,2</sub>-mCh (609 bp, pGi74) or TRO824/ TRO790 resulting in P<sub>bfpR,3</sub>-mCh (950 bp, pGi80) fusions. For promoter activity monitoring of P<sub>bfpR</sub>-mCh fusion, log-phase cultures (OD600 of 0.4) of the strain carrying the pGi47 plasmid performed in TYES broth supplemented with 20 µg/l gentamycin were distributed into 96-well microtitre plate, cultures were then supplemented with 20 µM of EDDHA and various iron sources (62.5 µM haemoglobin, 250 µM haemin, 5–250 µM FeCl₃) and incubated with continuous shaking at 18°C during 5 h. Cultures of strain carrying the pCPGm1−P<sub>les</sub>-mCh plasmid were used as negative control. Promoter activity monitoring of P<sub>bfp</sub>-mCh fusions was performed using bacterial colonies grown during 48 h on TYES agar supplemented with 1% haemoglobin or not. Colonies were scraped from the plates, then resuspended in TYES broth at a final OD600 of 1 for fluorescence measurement. Expression of the mcherry gene was monitored using whole-cell fluorescence with a Tectan Microplate Reader (Infinite 200 PRO). Excitation and emission wavelengths were set at 535 nm and 610 nm, respectively. Promoter activity (in arbitrary units) was estimated by dividing fluorescence intensity by the OD600.

**Construction of F. psychrophilum mutants**

Three suicide vectors pGi27, pGi29 and pGi38 allowing, respectively, hfpR, hfpY and bfpR deletion were constructed using the pYT313 plasmid [36] as previously described (M1 in Supporting Information) [31]. Each resulting plasmid was introduced by conjugation in strain OSU THCO2-90 and colonies selected for double crossing-over were screened by PCR using specific primers (T2 in Supporting Information) in order to identify those carrying the deletion genotype. The THC0290_1813 transposition mutant was produced as part of a Tn4351 mutant library as previously described [23,37]. Briefly, the pEP4351 suicide plasmid carrying Tn4351 was introduced by conjugation in strain OSU THCO2-90 and clones were selected on TYES agar supplemented with 10 µg/ml erythromycin. Identification of the Tn4351 interrupted locus was performed by digestion of genomic DNA by HindIII, ligation, inverse PCR of the resulting circular molecules using primers targeting Tn4351 and subsequent sequencing of the ampiclons. Sequences were used to locate the transposon insertion site into the genomic sequence.

**Construction of the F. psychrophilum chromosomal expression platform**

Several suicide plasmids allowing chromosomal recombination and gene insertion at a neutral intergenic region (between genomic positions 1,121,879 and 1,124,880 for strain OSU THCO2-90) were constructed
using pYT313 as backbone (T1 in Supporting Information). pGi39 was designed to carry the two DNA fragments upstream and downstream of the insertion site flanking an expression platform composed of a constitutive promoter PrpsL, a multiple cloning site and an intrinsic terminator, and pGi42 to carry the elastase encoding gene FP0506 under the control of PrpsL (M2 in Supporting Information).

**Complementation of the deletion mutants**

The chromosomal ectopic platform was used for reintroduction of deleted genes into the mutant strains. Several pGi39-derivative plasmids (pGi44, pGi45, pGi46 and pGi55; T1 in Supporting Information) were constructed and introduced by conjugation in the respective mutant strains, resulting in the chromosomal insertion of a single copy gene under the control of native promoter at the neutral intergenic region (M2 and F2 in Supporting Information).

**Recombinant MBP-HfpY purification and haem binding assays**

Recombinant HfpY was purified as a MBP-HfpY fusion protein from the pMBP-HfpY plasmid in E. coli Top10. pMBP-HfpY was constructed by gene synthesis (Proteogex). MBP-HfpY was purified by affinity chromatography on amylose resin as reported previously [38]. Briefly, bacteria were grown in 1L LB broth to OD600 = 0.6 and expression was induced with 0.5 mM isopropyl 1-thio-β-D-galactopyranoside (IPTG) overnight at RT. Cells were pelleted at 3500 g for 10 min, resuspended in 20 mM Hapes pH 7.5, 300 mM NaCl, containing 1 mM EDTA (binding buffer), and disrupted with glass beads (FastPrep, MP Biomedicals). Cell debris were removed by centrifugation at 18,000 g for 15 min at 4°C. MBP-tagged proteins contained in the supernatant were purified by amylose affinity chromatography (New England Biolabs) following manufacturer’s recommendations: the soluble fraction was mixed with amylose resin and incubated on a spinning wheel at 4°C for 1 h. The resin was then centrifuged, and washed 3 times with binding buffer. Purified proteins were eluted in binding buffer containing 10 mM maltose, and dialysed against 20 mM Hapes pH 7.5, 300 mM NaCl. Quantity and purity of the eluted fraction were determined with the Lowery Method (Biorad) and by SDS-PAGE followed by Coomassie staining, respectively. Haem binding was studied by UV-visible spectroscopy of MBP-HfpY in the presence of haemin in a microplate spectrophotometer (Spark, Tecan).

**Titration of MBP-HfpY with haemin**

Haem binding affinity and stoichiometry of MBP-HfpY were determined by adding 0.5 to 1 µl increments of a 200 µM haemin solution to cuvettes containing 100 µl test samples of 20 µM MBP-HfpY in 20 mM Hepes pH 7.5, 300 mM NaCl, or reference sample without protein. Spectra were measured from 230 nm to 750 nm in a UV-visible spectrophotometer Tecan (Spark). OD_{398} nm was plotted against haemin concentration and data were fitted to a one-binding site model (with calculated extinction coefficient of bound haemin of ε398 = 40 mM⁻¹. cm⁻¹) as described [38].

**Total RNA extraction**

Bacterial cells from 2 mL cultures were collected by centrifugation for 3 min after addition of ½ volume of frozen killing buffer (20 mM Tris-HCl pH 7.5, 5 mM MgCl₂, 20 mM NaCl) to the culture sample. The cell pellets were frozen in liquid nitrogen and stored at −80°C. For experiments performed in iron-depleted conditions, RNAs were extracted from pellets harvested at an OD600 of 0.8. For experiments performed in 1% haemoglobin-enriched conditions, RNAs were extracted from pellets harvested when bacterial concentration reached 10⁷ CFU/ml. Experiments were carried out from three independent cultures per biological condition. The ZymoBIOMICS RNA Miniprep Kit (Zymo Research) was used for total RNAs extraction, then 5 µg of RNA extracts were treated using DNase I (Qiagen) to remove residual genomic DNA and purified using the ZymoBIOMICS RNA Miniprep Kit. The RNA concentration was measured using a NanoDrop 1000 Spectrophotometer (NanoDrop Technologies, Inc.) and quality was assessed by electrophoresis on denaturing formaldehyde agarose gel.

**Reverse transcription PCR**

Transcriptional structure of the bfpR locus was determined by overlapping RT-PCR experiments. Total RNA was extracted from bacterial pellets harvested from cultures performed in TYES 1% haemoglobin. cDNA synthesis was carried out on 500 ng of DNase-treated RNA using the SuperScript™ II Reverse Transcriptase (Thermo Fisher) with random primers. Negative control reaction was run under similar conditions except that the reverse transcriptase was omitted (no RT). Polycistronic structure was detected by PCR amplification using cDNA as matrix and several primer sets designed to overlap adjacent genes (TRO677/TRO676 between the upstream region of bfpR and its
coding sequence, TRO674/TRO675 for the junction between THC0290_1678 and bfpR, and TRO672/TRO673 for the junction asd-THC0290_1678; T2 in Supporting Information). Negative and positive controls were run using no RT cDNA reaction and genomic DNA as matrix, respectively, and results were visualized by electrophoresis of PCR products on 1% agarose gel.

**Real-time quantitative PCR**

cDNA was generated from 500 ng of total RNA in a thermal cycler (Eppendorf) using the iScript™ Advanced cDNA Synthesis Kit for RT-qPCR (BIO-RAD) as recommended by suppliers. cDNA was diluted 1:25 in water and mixed with iTaq Universal SYBR Green Supermix (BIO-RAD) and forward and reverse primers with final concentration at 0.5 µM each. qPCR was performed using a Realplex Mastercycler (Eppendorf) or CFX system (BIO-RAD) with initial denaturation at 95°C for 3 min followed by 40 cycles of amplification as following: denaturation at 95°C for 5 s, annealing and extension at 60°C for 30 s. For each biological replicate, mean Ct values were calculated based on technical triplicate reactions and then normalized using the geomean of Ct values of two reference genes (rpsA and frr). Relative quantification of mRNA was expressed as $2^{-\Delta\Delta Ct}$ using TYES as a reference sample.

**Circular rapid amplification of cDNA ends (RACE)**

RACE was carried out using a general four-step protocol which enables the simultaneous cloning of 5' and 3' ends of RNA, as previously described [39]. Two RNA samples extracted from culture of the wild-type strain performed in TYES broth with 20 µM EDDHA or 1% haemoglobin were used respectively for hfp or bfpR mRNAs RACE experiments. Briefly, 8 µg of DNase-treated RNA samples were treated by 80 units of RppH (New England Biolabs) at 37°C for 1 hour to transform 5'-triphosphate groups carried by primary transcripts into 5'-monophosphate groups, then purified using the ZymoBIOMICS RNA Miniprep Kit. Circular ligation was performed by adding 80 units of T4 RNA ligase 1 (New England Biolabs) to 4 µg of the RppH-treated sample, the mix was incubated at 16°C overnight then heat inactivated at 70°C for 15 min. A control sample was prepared following the same protocol except that RppH was omitted in the reaction. First strand cDNA synthesis was performed using the SuperScript™ II Reverse Transcriptase (Thermo Fisher) with 400 ng of the ligation product as the matrix and a specific primer for each 3'-5' cDNA. 5'-3' end junctions were then amplified by PCR using cDNA as the matrix and specific primers (T2 and F4 in Supporting Information). PCR products were cloned into the pJET1.2 vector using the CloneJET PCR Cloning Kit (Thermo Fisher) then introduced in E. coli. Colonies were screened by PCR prior to Sanger sequencing of the insert. Three 3'-5' cDNAs carrying hfpR, hfpY or bfpR were produced using the gene-specific primers TRO806, TRO811 or TRO814, respectively. Detection of mRNA boundaries was performed as follows: for the hfpR cDNA using primers TRO806/TRO807 amplifying hfpR-THC0290_1813-hfpY mRNAs, for the hfpY cDNA using primers TRO812/TRO807 amplifying all hfpY mRNAs and for the bfpR cDNA using three sets of primers amplifying either all bfpR mRNAs (TRO815/TRO816), THC0290_1678-bfpR mRNAs (TRO815/TRO825) or asd-THC0290_1678-bfpR mRNAs (TRO815/TRO827). For bfpR RACE experiments, the first PCRs were followed by a nested PCR using TRO817/TRO818, TRO817/TRO826 or TRO817/TRO828, respectively, and the resulting PCR products were cloned into the pJET1.2 vector for Sanger sequencing.

**Rainbow trout experimental infections**

Three experimental infection models were used in this study differing in the infection route (intramuscular injection versus immersion), the rainbow trout lines or the size of fish (fingerling versus juvenile fish). Fish were reared at 10°C in recirculating aquaculture system (RAS) in 30-litre tanks with dechlorinated water, then transferred to BSL2 zone in 15-litre tanks with flow water (1 renewal per hour) for infection experiments. Commercial feed (@Le Gouessant) was delivered twice per day. Fish were fasted for 24 h or 48 h prior to infection by injection or immersion, respectively. Water quality was maintained by a constant renewal and daily cleaning. Bacterial cultures were performed in TYES broth at 18°C and used at the end of log phase (OD600 of 1, equivalent to 10^9 CFU/mL). Bacterial concentration was determined by colony counting on TYES agar after 4 days of incubation at 18°C. For the intramuscular injection model at the fingerling stage, fish (average body weight of 5 g) of the rainbow trout standard line (Sy^Aut) selected by INRAE were used. Six groups of 10 fish were anesthetized (MS222, Sigma) and challenged with the wild-type strain at different doses, as follows: 6 suspensions were prepared by ten-fold serial dilutions (10^{-1} to 10^{-6}) of a bacterial culture in TYES, then 50 µl of each suspension was injected per fish. The median lethal dose (LD50) of the wild-type was estimated using the
moving averages and interpolation method described by Thompson W.R [40] with three dose-groups used to calculate each moving average. This procedure was performed twice for the wild-type strain. A similar procedure was used for the eight deletion mutants and complemented strains except that only three doses (10^{-1} to 10^{-3}) were tested once. Survival curves were compared between strains at equivalent infectious doses using Kaplan–Meier method. F. psychrophilum bacterial counts were determined by plating serial dilutions of bacterial suspensions on TYES agar supplemented with FCS. After injection, fish were maintained in flow-through water at 10°C and mortalities were recorded twice a day for two weeks.

The ability of bacterial strains to colonize rainbow trout at the fry stage (average body weight of 2 g) was compared using the bath infection model on the rainbow trout homozygous line A36, which is highly susceptible to F. psychrophilum infection [10,23,41]. Groups of 40 fish in duplicates were used for each strain. Bacterial cultures were prepared as follows: two independent cultures of the wild-type, deletion and complemented mutant strains were grown in TYES broth until late-exponential phase (OD600 of 1). The bacterial cultures were directly diluted (200-fold) into 101 of aquarium water. Bacteria were maintained in contact with fish for 24 h by stopping the water flow and were subsequently removed by restoring the flow. During the experiment, water was maintained at 10°C under continuous aeration and physical parameters (NH_4+, NO_2-) were monitored. Sterile TYES broth was used for the control group. F. psychrophilum bacterial counts were determined at the beginning and at the end of the immersion challenge by plating serial dilutions of water sample on TYES FCS agar. Five fish from each tank (n = 10 per strain) were randomly chosen and euthanatized (tricaine, 300 mg/l) at 6 h post-infection to sample gills and spleen. Organs were mechanically disrupted in tubes containing 400 µl of peptone (10 g/l water) and 1 mm ceramic beads (Mineralex SAS, Lyon, France) using a FastPrep-24” 5 G instrument at 6.0 m s^{-1} for two cycles of 20 s (MP Biomedicals), and bacterial loads were determined by plating serial dilutions of lysates on TYES FCS agar.

Virulence of the bfpR deletion mutant and the wild-type strain was evaluated by intramuscular injection at the juvenile stage (average body weight of 65 ± 9 g) using the rainbow trout line A36. Bacterial suspension was prepared by 10-fold dilution in sterile TYES of a culture at an OD600 of 1. For each strain, 22 fish were anesthetized (MS222, Sigma) and inoculated with 100 µl of bacterial suspension then randomly distributed into two compartments of a 300-litre tank parted using a perforated polycarbonate divider and maintained in flow-through water (1 renewal per hour) at 10°C: the first subset (n = 10) was used for survival estimation by recording mortalities during 2 weeks and the second subset (n = 12) for sampling of spleen, head kidney and blood at 4 days post-infection and bacterial loads determination. Following anaesthesia, blood (150 µl per fish) was collected from the caudal vein using syringes containing heparin (500 units/mL) and the bacteraemia was determined by plating fresh blood and serial dilutions on TYES FCS agar plates. Infectious doses and bacterial loads in organs were determined as described above.

Statistical differences of bacterial loads between groups were analysed using the Mann–Whitney test, and survival curves were determined using the Kaplan–Meier method and compared using the Mantel-Cox log-rank test with GraphPad Prism 8.2.0 (GraphPad Software, San Diego, CA, USA).

**Ethics statements**

The infection challenges were conducted at the INRAE-IERP Fish facilities of Jouy-en-Josas (building agreement n°C78-720), performed in accordance with the European Directive 2010/63/UE regarding animal experiments, and approved by the institutional review ethics committee, COMETHEA, of the INRAE Center in Jouy-en-Josas, France. Authorizations were approved by the Direction of the Veterinary Services of Versailles (authorization number # 19,210-2015100215242446).

**Results**

**Identification of two TonB-dependent receptors with potential function in haem utilization**

The distribution of TBDRs was analysed in a collection of 22 genomes of F. psychrophilum isolates representative of a variety of host species, geographical origins and clonal complexes [42]. A total of 33 TBDR-encoding genes were retrieved. With an average number of 25 per genome, most (21) of these predicted TBDRs were conserved in all isolates, though 3 were pseudogenes in one isolate (Figure 1). Only a few TBDRs have predicted substrates, among which are the cobalamin transporter BtuB and 2 putative receptors homologous to the ferric hydroxamate receptor FhuA and the ferric citrate transporter FecA [43]. Among TBDR genes found only in a subset of the isolates, 3 may be associated to a particular host fish species. The first one, THC0290_1129 in strain OSU
THCO2-90, is only present in the genomes of CC-ST9 isolates infecting coho salmon. The 2 others, FP2456 and FP1882 in strain JIP 02/86, are present in all isolates belonging to the clonal complexes associated with rainbow trout (CC-ST10 and CC-ST90), but truncated and absent in the other genomes, respectively.

In order to further identify, among these TBDRs, those that are involved in iron acquisition, we searched for genes whose expression profile indicated specific regulation under blood supplementation, fish plasma exposure or conditions of iron limitation in the condition-dependent transcriptome dataset previously established for strain OSU THCO2-90 [31]. Two TBDR genes were up-regulated in the presence of blood: (i) THC0290_0369 without predicted substrates, and (ii) THC0290_1679, a member of the haem/haemoglobin receptor family (TIGR01785), hereafter named bpfr (for blood-induced Flavobacterium psychrophilum receptor). bpfr was also overexpressed in the presence of fish plasma but was not modulated by iron limitation. Four TBDR-encoding genes were up-regulated in cells submitted to low-iron conditions, but down-regulated when blood was supplied (Figure 1 and F1 in Supporting Information): (i–ii) FhaA and FecA siderophore receptors homologs; (iii) THC0290_0681 with unknown substrate; and (iv) THC0290_1814, the most highly deregulated gene, hereafter named hfpR (for haem F. psychrophilum receptor). hfpR is conserved in synteny with a gene encoding a HmuY-like hemophore protein (hereafter named hfpY), suggesting functional similarity with the haem utilization system Hmu characterized in other Bacteroidetes [44–47].

Here, we further characterized the Bpfr and Hfp systems predicted to transport haem. Expression profiles suggested a distinct regulation of the corresponding genes (F1 in Supporting Information), so we examined their transcriptional structure and regulation (see below). To analyse their respective role in bacterial growth, we also constructed the single and double deletion mutants Δbpfr, ΔhfpR, ΔhfpY and ΔhfpRΔbpfr, by allelic replacement in F. psychrophilum OSU THCO2-90 (T1 in Supporting Information). The ectopic complementation of these mutants was performed by inserting a single gene copy under the control of the native promoter at a different chromosomal position than the endogenous deleted gene copy. For ΔhfpRΔbpfr, the 2 genes were inserted arranged in tandem. For this purpose, a new plasmid was designed allowing chromosomal gene insertion at a neutral site conserved within the F. psychrophilum species and the efficacy of this expression platform was validated using reporter genes (F2

**Figure 1.** Conservation and gene expression of the 24 TonB-dependent receptors encoded in the genome of strain OSU THCO2-90. (a) *in silico* prediction of TBDR encoding genes. The list of TBDR encoding genes was retrieved by *in silico* prediction of the plug and β-barrel domains. Conservation in a selection of 22 isolates: gene present (blue), truncated (grey), absent (white). (b) MLST and genomic data are from Duchaud et al. [42]. CC-ST: clonal complex-sequence type. (b) fish host: Oncorhynchus mykiss, Salmo trutta, Tinca tinca, Gasterosteus aculeatus, Plecoglossus altivelis, Oncorhynchus kisutch, Salmo salar. (b) TBDR-encoding genes significantly upregulated (green) and down-regulated (red) in strain OSU THCO2-90 in iron-limited condition (TYES + 25 µM 2,2’-dipyridyl), under rainbow trout plasma exposure or in the presence of blood (TYES agar supplemented with 10% horse blood). Differential expression is expressed as log2-fold change values retrieved from Guérin et al. [31] (https://fpeb.Migale.inrae.fr).
and F3 in Supporting Information). The growth of the mutants was analysed in vitro under different conditions of iron and in vivo in the natural host.

**High levels of haemoglobin promote the accumulation of monocistronic bfpR mRNA**

BfpR is encoded downstream of the aspartate-semialdehyde dehydrogenase (asad) and a hypothetical protein (THC0290_1678; Figure 2(a)). Structure of the bfpR transcript is uncertain: expression level across biological conditions differs between bfpR, asd and THC0290_1678, and the increase of mRNA level in colonies grown on blood-supplemented TYES agar occurs upstream bfpR, suggesting that bfpR is transcribed as a monocistronic mRNA. However, no transcription start site (TSS) was detected in the corresponding region (Figure 2(a)) [31].

Several methods were combined to define the structure of the bfpR transcript. First, a putative polycistronic mRNA encompassing asd, THC0290_1678 and bfpR was predicted by RT-PCR using overlapping primers. Circular 3'-5' RACE experiments were then run in an attempt to identify the most abundant bfpR transcript and to distinguish between primary and processed RNAs (Figure 2(a); F4 in Supporting Information). bfpR amplification from a primer within asd was unsuccessful, attesting of the very low quantity of the polycistronic mRNA predicted by overlapping RT-PCR. In contrast, cDNA was successfully amplified using primers within bfpR. The majority of transcripts ended at +83 bp relative to the bfpR stop codon, while no discrete 5'-end position was identified. Instead, most transcripts started within a 50 bp-long region spanning from −399 to −347 bp relative to the bfpR start codon, which coincides with the upshift of expression signal detected by microarrays in blood supply conditions. By using a primer within THC0290_1678, transcripts were detected by nested PCR only and started mainly at −673 (±1) bp relative to bfpR start codon. This 5' end is 22 bp downstream the start position of

![Figure 2. Transcriptional organization of the bfpR locus and its regulation by haemoglobin concentration.](image-url)
THC0290_1678. However, the location of translation initiation is difficult to predict in *Bacteroidetes* [48] and another ATG codon located 38 bp downstream the 5’ end may be the correct start. Various fragments (306, 609 and 950 bp) of the upstream region of *bfpR* were cloned into a transcriptional reporter plasmid and a significant promoter activity was detected only for the one carrying the longest fragment when colonies were grown in TYES agar supplemented with 1% haemoglobin (Figure 2(b)). These results indicate that the majority of *bfpR* mRNAs are monocistronic and may originate from processing of a longer transcript starting 673 bp upstream of *bfpR*.

We therefore compared the effect of haemoglobin supply on *bfpR* mRNA level by RT-qPCR assays (Figure 2(c)). Addition of large amounts of haemoglobin in TYES broth led to 10-fold increase of *bfpR* mRNA level. This increase was not observed for the upstream gene THC0290_1678. Altogether, we concluded that a primary transcript encompassing THC0290_1678 and *bfpR* is synthetized then probably subject to maturation, resulting in the *bfpR* mRNA that accumulates in presence of large amounts of haemoglobin.

**BfpR contributes to growth in the presence of high amounts of haemoglobin**

On the basis of its expression profile, we analysed the role of BfpR on the growth of *F. psychrophilum* under high-haemoglobin condition. The wild-type strain exhibited a slight delay in the presence of 1% haemoglobin (equivalent of 620 µM haem), but final viable cell counts were higher than in TYES broth (F5 in Supporting Information), indicating that haemoglobin availability can enhance bacterial growth. In these conditions, the growth rate of Δ*bfpR* was reduced compared to that of the wild-type and partially restored in the complemented Δ*bfpR* strain (Figure 3). RT-qPCR showed that ectopic expression of *bfpR* in this complemented Δ*bfpR* strain resulted in reduced mRNA level compared to the wild-type (F6 in Supporting Information), indicating that adequate regulation of *bfpR* expression is important for growth in these conditions. Growth curves of Δ*hfpR*Δ*bfpR* and Δ*bfpR* were similar. These results showed that among the two TBDRs, only BfpR is required for optimal growth in the presence of high amounts of haemoglobin.

**The hfp operon is transcribed in response to iron scarcity**

The Hmu systems characterized so far in *Porphyromonas gingivalis* and other *Bacteroidetes* are transcribed in a polycistronic transcript encoding at least 1 HmuY-like hemophore, 1 TBDR and 4 uncharacterized proteins (a putative CobN/magnesium chelatase and 3 putative membrane proteins) [46,47]. In *F. psychrophilum*, only the genes encoding a TBDR (*hfpK*) and the HmuY-like protein (*hfpY*) are present. They are located within a 19 kb region containing other iron-responsive transcripts encoding proteins that have a putative role in iron assimilation or transmembrane electron transfer (T3 in Supporting Information) [31,49]. The upregulated polycistronic mRNA deduced from transcriptomic data is bound by a putative TSS and a predicted intrinsic terminator and encodes, from 5’ to 3’: (i) HfpR; (ii) THC0290_1813, a predicted 18 kDa inner membrane protein of unknown function belonging to the DoxX family (IPR032808); and (iii) HfpY (Figure 4(a)). Expression signal measured by high-density microarrays was higher in the *hfpY* part of the operon for all biological conditions suggesting that a monocistronic *hfpY* mRNA may also be produced (F7 in Supporting Information). We confirmed this transcriptional structure by 3’-5’ RACE experiments: two 5’-ends and one unique 3’-end were identified (Figure 4(a); F4B in Supporting Information). RppH enzymatic treatment increased cDNA amplification for the polycistronic

![Figure 3. *bfpR* is required for *in vitro* growth under high haemoglobin concentration. Growth under high haemoglobin concentration determined by viable cell measurements (CFU/ml) of cultures in TYES broth supplemented with 1% haemoglobin. Plain lines: wild-type (black) and deletion mutants (Δ*bfpR*: green; Δ*hfpR*Δ*bfpR*: brown), dashed lines: ectopic complemented mutants. Values represent the mean and standard deviation of four independent experiments.](image-url)
Figure 4. Transcriptional organization of the hfp locus and its regulation by iron availability. (a) genomic view of the hfpR-THC0290 _1813-hfpY locus, microarray expression profiles and putative TSSs from Guérin et al. [31] are shown (see legend of Figure 2). For iron limitation, cells were grown in TYES broth supplemented with 25 µM 2,2'-dipyridyl (red) or not (black); for blood supplementation, cells were from colonies grown on TYESA covered by 10% defibrinated horse blood (violet) or not (black). 3'-5’ RACE: schematic representation of 5’ and 3’ ends identified by circular 3'-5' RACE experiments. Positions of the two 5’ ends and of the 3’ end are relative respectively to start positions of hfpR, hfpY, and stop position of hfpY. (b) transcriptional regulation of the hfpR promoter by iron availability. Promoter activity was measured using whole-cell fluorescence of the wild-type strain carrying the reporter plasmid with 240 bp DNA upstream of hfpR (P_less). Bacteria were grown in iron depleted TYES broth (E20, 20 µM EDDHA) supplemented with a range of FeCl₃ (upper panel) or with different iron sources (lower panel) as indicated. Cultures were inoculated at OD600 0.4 and fluorescence was measured during 5 h as described in materials and methods. Strain carrying the empty plasmid (P_less) was used as control. Values represent the mean and standard deviation of three independent experiments. (C) hfpR, 1813 and hfpY mRNA levels correlate with iron availability. RT-qPCR measurements were performed under iron depletion (upper panel) or high concentration of hemoglobin (lower panel) as described in Figure 2 using the wild-type strain grown in TYES broth, iron-depleted TYES (E20) supplemented with 1.25 µM haemoglobin (E20 Hb) or in TYES supplemented with 1% haemoglobin (Hb 1%).
form only indicating a 5'-triphosphorylated RNA. These results confirmed the TSS position and the existence of 2 mRNAs, while suggesting that the hfpY mRNA originates from a post-transcriptional maturation event.

Expression of the hfp locus was then monitored under iron-depleted and -replete conditions using a transcriptional reporter (P_hfpR-mCh) and RT-qPCR assays. Addition of the iron-chelator ethylenediamine-N,N'-bis (2-hydroxyphenylacetic acid) (EDDHA) in TYES broth resulted in a strong increase of promoter activity, which was countered by FeCl₃ supply in a dose-dependent manner (Figure 4(b)). Promoter activity was also limited by haem or haemoglobin supply, although with a lower efficiency than when FeCl₃ was used (Figure 4(b)). RT-qPCR confirmed up-regulation of the three genes composing the hfp operon under iron limitation and the effect of haemoglobin supply (Figure 4(c)). Moreover, a fivefold down-regulation was measured in the presence of high level of haemoglobin in iron-replete TYES broth (Figure 4(c)), which is consistent with the down-regulation previously observed by microarrays in the presence of blood [31].

We concluded from these results that two mRNAs are produced at the hfp locus: a primary transcript encompassing hfpR, THC0290_1813 and hfpY, and a matured mRNA carrying hfpY only, resulting in more abundant mRNA for hfpY than for hfpR. The expression level of these 2 RNAs is tightly controlled in response to iron availability through the regulation of the hfpR promoter, which is activated by iron scarcity and downregulated in presence of iron sources.

**HfpR and HfpY are required for haem utilization under iron scarcity**

The effect of iron limitation on growth was first analysed on the wild-type strain. Addition of EDDHA, a chelator of ferric iron Fe(III), to TYES broth resulted in a strong growth inhibition, confirming that iron is an essential element for *F. psychrophilum*. Growth was restored by the supply of haemoglobin, haemin or FeCl₃, indicating that the bacterium is able to scavenge these different iron sources from the environment (Figure 5).

We then analysed the role of HfpR and HfpY by comparing the growth of deletion mutants and respective complemented strains under iron scarcity (Figure 5). Supplementation by FeCl₃ fully restored the growth of all strains (Figure 5). In contrast, growth inhibition was only partially restored by haemoglobin and haemin supplementation in mutant ΔhfpR which still exhibited a severe growth delay compared to complemented ΔhfpR and wild-type strains. ΔhfpY was by far less affected but its growth rate still significantly reduced compared to the wild-type in haemoglobin supplemented medium containing EDDHA, and this growth defect was partially restored in the complemented ΔhfpY strain. In contrast, no growth defect was observed in ΔhfpY when haemin was supplied instead of haemoglobin (Figure 5). Partial growth restoration in the ectopic complemented ΔhfpY strain is likely due to its reduced hfpY mRNA level compared to the wild-type as measured by RT-qPCR (F6 in Supporting Information).

We also evaluated the growth on haemoglobin of a transposition mutant inactivated in the unknown function gene lying between hfpR and hfpY (THC0290_1813); no difference was observed compared to the wild-type, indicating that this gene is dispensable for haem utilization under iron-restricted conditions (F8 in Supporting Information).

Finally, the behaviour of ΔhfpR was analysed. In contrast to ΔhfpR and ΔhfpY, no growth defect was observed under iron limitation when haemoglobin, haemin or FeCl₃ was supplied, and growth of ΔhfpRΔhfpR was indistinguishable from that of ΔhfpR (Figure 5). We conclude that HfpR and HfpY are important for utilization of haem compounds under iron scarcity whereas BfpR is dispensable, and that there is no functional compensation by other genes for the lack of HfpR under these conditions.

**F. psychrophilum HfpY is a haem-binding protein**

HfpY exhibits only low level of similarity with other HmuY-like hemophores (below 26% identity in amino-acids sequence; F9 in Supporting Information); we thus investigated the ability of HfpY to bind haem in *vitro*. Recombinant HfpY was purified as a fusion to the maltose-binding protein (MBP) from *E. coli*. Spectroscopic monitoring showed that while the maximum absorbance of free haemin was located between 361 and 384 nm, addition of purified MBP-HfpY resulted in a Soret peak at λ_max 398 nm typical of haem coordination to protein (Figure 6(a)). Q bands were also visible at 503, 530 and 623 nm in the difference spectra. Haem titration assays indicated a stoichiometry of 1:1 for the holoprotein which is similar to the other characterized HmuY-like hemophores (Figure 6(b)). Based on these results, we conclude that HfpY of *F. psychrophilum* is a haem-binding protein.
The two TBDRs are required for virulence in rainbow trout

Wild-type, deletion mutants and complemented strains were analysed for virulence in rainbow trout using three experimental infection models that differ by the infection route and fish size. The roles of hfpR, hfpY and bfpR were first evaluated by measuring bacterial loads in organs after immersion challenge in fry (2 g), a model that mimics the natural route of infection. At 6 h post-infection, *F. psychrophilum* was detected in the gills of fish from all infected groups at a similar concentration (average of $9 \times 10^3$ CFU/gill). In contrast, colonization of spleen was significantly lower for groups infected with ΔhfpR, ΔhfpY or ΔhfpRΔbfpR relative to the wild-type (Figure 7(a)). No significant difference was observed between bacterial loads of fish infected with ΔbfpR or the wild-type, though values were more scattered with ΔhfpR. Bacterial colonization of the spleen was fully restored by complementation of
Figure 6. HfpY is a haem-binding protein. (a) UV-visible absorption spectra of MBP-HfpY complexed with hemin. 40 μM MBP-HfpY was purified from E. coli and mixed with equimolar concentration of hemin. UV-visible spectra (in 200 μl) of the complex and same concentration of hemin were obtained in a microplate spectrophotometer (Spark Tecan). (Inset) 5X magnification of the 500- to 700 nm region. Results are representative of three independent experiments. (b) Titration of MBP-HfpY with hemin followed by absorbance at 398 nm. 20 μM of MBP-HfpY were mixed with increasing concentrations of hemin as indicated. For each spectrum, the absorbance of the same concentration of hemin was subtracted and the difference absorption spectra are shown, from light to dark blue color as a function of hemin concentration. The absorbance difference at 398 nm was plotted against hemin concentration (inset). The curve is representative of 3 independent experiments and was fitted using the nonlinear regression function of GraphPad Prism 4 software, which determined that the stoichiometry of the HfpY-hemin complex was 1:1.

The ΔhfpRΔbfpR double mutant compared to the wild-type, but only partially restored in the complemented ΔhfpY strain.

Survival curves were then compared in fingerling (mean weight, 5 g) after intramuscular injection to quantify the virulence. The median lethal dose (LD50) of the wild-type was determined as 1.9 (±0.3) × 10^5 CFU. Relative assessment of the virulence of mutants was achieved by comparing survival rates relative to wild-type. At an infection dose corresponding to 10 × LD50 of the wild-type strain, 100% of fish died rapidly in groups infected with the wild-type or complemented strains whereas mortality was delayed and survival was significantly increased for fish infected with either ΔbfpR or ΔhfpY (Figure 7(b)). A similar difference between strains was also observed using a lower infectious dose (F10 in Supporting Information). Using this infection model, the survival curves of fish inoculated with ΔbfpR or the wild-type were similar. In addition, the levels of mortality observed with ΔhfpRΔbfpR and ΔhfpR were similar (F10 in Supporting Information).

Since massive haemolysis may not be encountered in the fingerling infection model, we also investigated the role of BfpR in virulence on larger fish. Two groups of juvenile rainbow trout (mean weight, 65 g) were infected by intramuscular injection and the survival curves or bacterial loads in blood, spleen and kidney 4 days post-infection were determined. In this model, bacteraemia and bacterial loads in spleen and kidney were consistently significantly lower in the group infected with ΔbfpR relative to the wild-type (Figure 7(c)). The fish groups infected with ΔbfpR displayed a higher survival rate compared to those infected with the wild-type strain (Figure 7(d)).

Altogether, the results showed that both Hfp and BfpR systems are necessary for full virulence of F. psychrophilum in rainbow trout with no apparent functional redundancy.

Discussion

Here, we report the identification of two haem/hemoglobin outer membrane receptors conserved in the species and demonstrate that they act differently to support bacterial growth, under high haemoglobin condition for BfpR and under iron scarcity for HfpR. We also show by experimental infection in a natural host, the rainbow trout, that both receptors are required for full virulence (Figure 8). The findings have relevance to other important fish pathogens such as Flavobacterium columnare and Tenacibaculum maritimum that possess homologs of the two uptake systems reported here.

Expression of the hfp and bfpR loci are tightly regulated and indicate a role under distinct iron conditions

Complex regulatory mechanisms act in bacterial pathogens to maintain iron at intracellular levels needed for bacterial development and below its toxicity threshold
Figure 7. Evaluation of the virulence of wild-type and deletion mutant strains in rainbow trout. (a) bacterial loads in the spleen of rainbow trout fry after immersion challenge (A36 line, 2 g average weight). Each tank was inoculated at a bacterial concentration of 1 (±0.3) × 10^7 CFU/mL and fish were maintained in contact with bacteria for 24 h at 10°C under optimal oxygenation conditions. At the end of the challenge, bacterial concentration in water was 2 (±0.7) × 10^7 CFU/mL. Ten fish from each group were sacrificed at 6 h post-infection and bacterial loads in the spleen were determined. The colour of circles indicates strains: wild-type: black, ΔhfpR: red, ΔbfpR: green, ΔhfpRΔbfpR: brown, ΔhfpY: blue. Open circles are for corresponding ectopic complemented mutants. Horizontal dashed line indicates the detection threshold. Values are significantly lower in the spleen of groups infected with the ΔhfpR and ΔhfpY mutants compared to the wild-type and complemented strains (Mann–Whitney: *, 0.033; **, 0.002; ***, <0.001). (c) Kaplan–Meier survival curves of rainbow trout fingerling (Sy*Aut line, 5 g average weight) after intramuscular injection. Each group (n=10) was challenged at a dose corresponding to 10 x LD50 of the wild-type strain. Upper panel: 3 × 10^6 CFU for the wild-type (black), 2 × 10^6 CFU for ΔhfpR (red plain line) and ecto:hfpR (red dashed line); lower panel: 6 × 10^6 CFU for the wild-type, 7 × 10^6 CFU for ΔhfpY (blue plain line) and ecto:hfpY (blue dashed line). The survival curves for fish inoculated with either the wild-type or the ectopic complemented mutants were significantly different from those inoculated with the ΔhfpR or ΔhfpY mutants (p-value of the Mantel-Cox log-rank test is indicated). (c-d) experimental infection of juvenile rainbow trout (A36 line, 65 g average weight) after intramuscular injection. Fish were inoculated with 1 × 10^7 CFU for the wild-type (black) or ΔbfpR mutant (green) strains; a subset (n = 12) was used for bacterial loads determination in blood, spleen and head kidney at 4 days post-infection (c) and another subset (n = 10) for survival estimation (d). Significant differences were observed between both the Kaplan–Meier survival curves (Mantel-Cox log-rank test p-value) and the bacterial loads (Mann–Whitney test).
Here, we show that hfpR and hfpY are upregulated in an iron-dependent manner. The promoter of the operon was induced under low-iron condition, whereas availability of ferric iron led to a dose-dependent transcriptional repression, which was also observed in presence of a high level of haemoglobin. In many bacteria, such regulation is mediated by Fur family iron-responsive regulators. Typically, under iron-replete condition the Fe^{2+}-Fur holoprotein binds to the promoter of iron-regulated genes thus preventing transcription, while under low-iron apo-Fur is released and transcription occurs [51]. *F. psychrophilum* genome contains one member of the Fur family (THC0290_0812). Several attempts to construct a deletion mutant failed at the second recombination event, plasmid excision resulting only in wild-type genotype. This may be either due to the requirement of the Fur homolog for growth or to impaired expression of the downstream purine biosynthesis gene purA [31].

Several evidences indicate that post-transcriptional regulatory events occur in both hfpR and hfp mRNAs. In the wild-type strain, mRNA levels for hfpY were higher than for hfpR under all tested biological conditions (F7 in Supporting Information) [31]. In-depth analysis of the transcriptional structure of the hfp locus revealed the existence of two mRNAs (hfpR-1813-hfpY and hfpY) and only the polycistronic RNA was triphosphorylated at the 5' terminus, indicating a single primary transcript. Moreover, no binding site of sigma factor was found in the sequence surrounding the 5' end position of hfpY mRNA, which is consistent with a post-transcriptional regulation. We also showed that hfpY mRNA level was lower in the ΔhfpY strain carrying the chromosomal transcriptional fusion P_{hfpR-hfpY} than in the wild-type, suggesting that the natural 32-nucleotide 5' untranslated region of hfpY mRNA is important for its stability. This lower expression was associated to only partial restoration of *in vitro* and *in vivo* phenotypic defects of ΔhfpY, indicating that accurate hfpY mRNA quantity is needed for full activity of the Hfp system. Moreover, a putative low permissive transcriptional state of the ectopic chromosomal site is unlikely as hfpR mRNA level in complemented ΔhfpR carrying the P_{hfpR-hfpR} fusion was similar than in wild-type. It is noteworthy that several hmuY transcripts are also
produced in *P. gingivalis*, the monocistronic form being more abundant than the bicistronic *hmuy-hmuR* one, suggesting a similar hemophore upregulation in the two species despite distinct operonic structure [46]. In bacteria, the stability of RNAs released by processing of a polycistronic RNA may differ, which affects the relative expression levels of genes within an operon. Our hypothesis is that a stable *hfpY* transcript is released after maturation of *hfpR*-1813-*hfpY* mRNA, resulting in higher quantity of *hfpY* compared to *hfpR*, independently of iron level. A similar regulation was recently described in *Pseudomonas aeruginosa* for the *hasR-hasA* operon encoding the hemophore-dependent Has system [52]. Production of large amounts of hemophore likely provides a significant advantage to maintain effective concentration relative to its cognate receptor while diffusing into the surrounding environment.

In contrast to *hfp*, we showed that *bfpR* mRNA accumulated in the presence of haemoglobin under iron-replete condition. The upregulated RNA was monocistronic and likely originated from maturation of a longer RNA starting at –673 bp from the *bfpR* start codon and carrying a putative hypothetical gene (THC0290_1678). RNA level of the 5’ region of this long transcript was unchanged in the presence of haemoglobin. The underlying mechanism, presently not known, likely involves the condition-dependent modulation of 1678-*bfpR* mRNA cleavage by a ribonuclease. Typically, such regulation can be mediated by structural change of the target RNA or by binding of a small regulatory RNA, this interaction acting positively or negatively on the stability and translation of the target [53]. For instance, a sRNA-dependent mRNA cleavage of the 5’ UTR of *cola* mRNA, which encodes the collagenase in *Clostridium perfringens*, releases a more stable transcript allowing overexpression of the toxin [54]. Several *F. psychrophilum* sRNAs were shown to accumulate in the presence of blood [31] and are regulator candidates for the *bfpR* post-transcriptional regulation. It is also noteworthy that the regulatory event leading to *bfpR* accumulation occurs also under specific conditions without haemoglobin such as in biofilm [55] or in stationary phase [31].

**Role of the two haem/haemoglobin receptors in haem acquisition and host colonization**

For vertebrate pathogens, the most abundant form of iron is bound within haem, and mainly sequestered into haemoglobin in erythrocytes and into other host proteins. In rainbow trout, serum albumin, hemopexin-like proteins and haemoglobin are among the most abundant proteins in plasma [56].

In vitro, when ferric iron was deprived and haem or haemoglobin was the sole iron source, the growth of *F. psychrophilum* was strongly impaired in the absence of HfpR, but only slowed in the absence of HfpY. The importance of the hemophore was nevertheless obvious in *vivo*, the lack of HfpY resulting in reduced mortality and bacterial loads in spleen. These results indicate that the outer membrane receptor is able to independently capture and transport haem, although cooperation with its cognate hemophore is needed in *vivo*, likely to out-compete the host sequestration systems (Figure 8). The effectiveness of hemophore-based receptors to capture very low levels of haem was exemplified for several acquisition systems, such as the Has system found in *Serratia marcescens* and other Gram-negative bacteria: the affinity of HasR for the hemophore was 100-fold higher than for free haem [57,58].

Direct binding of various substrates to the HfpR receptor was not assayed here, but meaningful information can be deduced from phenotypic results. The Hfp system was able to provide nutrient iron by the uptake of haem from haemoglobin. Previously, it was established that *F. psychrophilum* cells are highly proteolytic and that a T9SS-deficient mutant is unable to consume haemoglobin [21]. This indicates that the Hfp system may require additional factors to efficiently retrieve haem from hemoproteins. We propose that haemoglobin cleavage by exopeptidases contributes to release haem molecules which are then captured by the hemophore HfpY for delivery to the outer membrane receptor. Other hemoproteins such as serum albumin or hemopexin may similarly serve as alternative haem source after their proteolysis by secreted peptidases.

The expression profile of *bfpR* and the phenotypic characterization of the corresponding deletion mutant suggested a role in adaptation to haemoglobin overload, a stress condition that the pathogen likely undergoes during its lifecycle (Figure 8). Indeed, infected fish develop haemorrhagic septicaemia [13]. *F. psychrophilum* cells are haemolytic and high bacterial load in blood could result in the release of large amounts of haemoglobin. Bacterial cells also accumulate at high level in spleen and kidney, two haematopoietic tissues in teleost fish. In particular, the spleen acts as a filter for blood-derived substances and plays a major role for pathogens clearing and turnover of erythrocytes through phagocytosis by melanomacrophages [59]. This infection niche thus accumulates large amount of haem. We assume that the haemoglobin concentration (1%) used for *in vitro* experiments properly mimics massive haemolysis, as total
haemoglobin levels of 6% to 12% (g/dl) in blood of healthy rainbow trout were reported [60]. The results showed that the presence of 1% haemoglobin did not affect the viability of *F. psychrophilum*, rather, the growth of wild-type was only slightly reduced and the final biomass was increased. In contrast, the lack of BfpR resulted in a greatly reduced growth rate. Under these conditions, the Hfp uptake system was strongly repressed, though free iron was available as nutrient iron source. We concluded that *F. psychrophilum* cells can tolerate high concentration of haemoglobin (equivalent to 620 μM haem) without toxicity and that BfpR is involved in this adaptation. Several Gram-negative pathogenic bacteria are reported to accumulate haem in their membrane with no apparent toxicity, whereas others are highly susceptible [61]. Additional experiments are ongoing in order to better characterize the function of BfpR in the physiology of *F. psychrophilum* under haemoglobin overload.

Experimental infection in rainbow trout fingerling using intramuscular injection or immersion models highlighted the major contribution of the Hfp system to virulence and host colonization. In contrast, the lack of BfpR had no significant impact in these experiments. It is noteworthy that experimental infection of rainbow trout at the early stage (1 to 5 g) by strain OSU THCO2-90 induced acute mortality 2- to 3-day post-infection, this tiny window making monitoring of the infectious process difficult. We thus infected juvenile rainbow trout (65 g) which immune system is expected to be more mature, allowing a better resistance against infection. Mortalities occurred between 4- and 8-days post infection with the wild-type, whereas the lack of BfpR resulted both in reduced bacterial loads in blood, spleen and kidney and in increased fish survival. This indicates that BfpR contributes to *in vivo* fitness in *F. psychrophilum*, although this conclusion needs to be strengthened by further replicate experiments. Overall, we propose that both BfpR and Hfp systems support haem uptake with non-redundant functions and act at distinct substrate concentration ranges to adapt the physiology of *F. psychrophilum* to iron sources available in the surrounding environment. Many pathogenic bacteria evolved to produce multiple haem receptors, each of them contributing to host colonization. For instance, *P. aeruginosa* produces three outer membrane haem receptors: the hemophore-dependent systems HxuA and Has act mostly as sensors of extracellular haem, while the hemophore-independent PhuR receptor ensures the uptake of high amounts of haem providing nutrient iron for growth [62,63].

**Sequence properties of the BfpR and HfpR receptors and the HfpY hemophore**

Distinct TBDRs typically display similar 3D-structural organization but low-sequence similarity (20%). Predicting the substrate of TBDRs is challenging as specificity resides more in a few residues located in external loops and internal surface of the channel than in the global sequence conservation of the beta-barrel domain. Despite the high diversity between sequences, conserved amino acid motifs have been identified in haem/haemoglobin receptors characterized in other Gram-negative bacteria [64–66]. Multiple sequence alignments showed that the FRAP/NXXL motifs and the histidine residue located within the external loop L7 in other haemoglobin/haem receptors were conserved in BfpR (F11A in Supporting Information), suggesting that it interacts directly with haemoglobin. In the case of HfpR, the loop L7 was extended (78 AA instead of 34 AA for *P. gingivalis* HmuR) and motif conservation was only partial. No obvious conservation of the FRAP/NXXL motifs was found for the two other TBDRs (THC0290_0681 and THC0290_0369) that are overexpressed by low-iron conditions and in the presence of blood, respectively (Figure 1, F11B in Supporting Information). These FRAP/NXXL motifs and histidine residue are thought to mediate haemoglobin binding and haem utilization, although mutagenesis studies of *P. gingivalis* HmuR and *Haemophilus ducreyi* HgbA showed a different behaviour and other residues can be involved in substrate interaction [64–66].

We noticed that, compared to other TonB-dependent haemoglobin receptors, BfpR possesses an additional domain called NTE for N-terminal extension (Carboxypeptidase D regulatory-like, IPR008969) which is frequently found in TBDRs (i.e. 16 out of 23 TBDRs in strain OSU THCO2-90). This domain of unknown function may contribute to substrate uptake as recently exemplified for the levan acquisition system in *Bacteroides thetaiotaomicron* [67].

The low sequence similarity of BfpR and HfpR with other characterized haem/haemoglobin receptors makes the transfer of knowledge hazardous and future investigations are needed to decipher the haem transport mediated by BfpR and HfpR at the molecular level. Moreover, the very low sequence conservation of HfpY with other members of the HmuY family makes it probable that the hemophore of *F. psychrophilum* functions using an original haem-binding mechanism. Indeed, despite a general conservation of haem binding ability in all five HmuY-like proteins described so far in strict anaerobes (*P. gingivalis, Tannerella forsythia,*...
Prevotella intermedia and Bacteroides vulgatus), in depth studies revealed important differences in their mode of interaction with haem; for instance, haem coordination involved His134/His166 in P. gingivalis HmuY but Met140/Met169 in T. forsythia Tfo [45,68].

Other iron acquisition systems are predicted in the genome of F. psychrophilum

The majority of TBDRs encoded in the F. psychrophilum genome have unknown functions. Based on the expression profiles in plasma or blood, several other candidates may also contribute to iron acquisition from haem or other iron sources. Occurrence analysis of TBDR encoding genes in F. psychrophilum genomes identified several receptors present only in genomes from specific lineages. Molecular epidemiology studies of F. psychrophilum outbreaks revealed an epidemic population structure with limited number of clonal complexes that are associated with host fish species [6,8,42]. For instance, strains isolated from coho salmon, including OSU THCO2-90 used in this molecular genetic study, belong to CC-ST9, whereas the most prevalent observed clonal complex, CC-ST10, contains the majority of isolates recovered from rainbow trout worldwide. Phenotypic diversity between isolates was reported and a few related genetic determinants identified so far by comparative genomics [33,69–74]. The two haem uptake systems characterised here using experimental infection in rainbow trout belong to the core genome and are unlikely responsible for host specificity. Further experimental infection studies performed in other fish species are necessary to decipher molecular mechanisms underlying this specificity. Here, we found such candidate genes: a gene cluster containing an uncharacterized TBDR-encoding gene (FP2456 in strain JIP 02/86) present only in genomes belonging to the two predominant clonal complexes infecting rainbow trout (i.e., CC-ST10 and CC-ST-90). The FP2456 protein belongs to the TonB-dependent siderophore receptor family (TIGR01783) and genes conserved in synteny encode two putative siderophore biosynthesis enzymes and a MFS-type efflux transporter. Further studies aiming to characterize the role of this predicted siderophore system in iron acquisition and in vivo fitness in rainbow trout are of great interest but are currently limited by the lack of efficient mutagenesis tools to manipulate these isolates. Along the same line, the importance for virulence of the uncharacterized TBDR-encoding gene (THC0290_1129 in OSU THCO2-90) present specifically in isolates belonging to CC-ST9 could be evaluated in a coho salmon infection model [75].

Conclusion

This study provides molecular knowledge on three novel molecular factors involved in the infection process in F. psychrophilum, a serious fish pathogen responsible for important losses in trout and salmon farming industries worldwide. The results will help to determine future directions of study for many aspects of haem acquisition. A brief list of interesting topics would be the characterization of regulatory events that coordinate expression of HfpR and BpR receptors in response to environmental changes and identification of additional proteins involved in the process such as exopeptidases releasing haem from hemoproteins, putative haem oxygenase degrading haem to free ferrous iron, or inner membrane transporter mediating transfer to the cytoplasm. F. psychrophilum likely evolved several acquisition strategies to take advantage of the diversity of iron sources encountered during its lifecycle. Mutagenesis of other putative iron acquisition systems may also unveil additional factors favouring host colonisation and disease development. F. psychrophilum infections are a major threat for salmonid farming worldwide and innovative control strategies are needed to limit their impact. This study shows that haem outer membrane receptors have a major contribution to bacterial fitness in the fish host and are promising targets for innovative antibacterial strategies [76].

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Data availability statement

The authors confirm that the data supporting the findings of this study are available within the article and its supplementary materials.

Supporting information

Supporting Information.pdf This file contains Supplementary Methods (M1 to M3), Supplementary Figures (F1 to F11) and Supplementary Tables (T1 to T3).

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