First Isolation of Virulent Tenacibaculum maritimum Isolates from Diseased Orbicular Batfish (Platax orbicularis) Farmed in Tahiti Island

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Abstract: The orbicular batfish (Platax orbicularis), also called ‘Paraha peue’ in Tahitian, is the most important marine fish species reared in French Polynesia. Sudden and widespread outbreaks of severe ‘white-patch disease’ have occurred since 2011 in batfish farms one to three weeks after the transfer of juveniles from bio-secured hatcheries to lagoon cages. With cumulative mortality ranging from 20 to 90%, the sustainability of aquaculture of this species is severely threatened. In this study, we report for the first time the isolation from diseased batfish of several isolates belonging to the species Tenacibaculum maritimum, a major pathogen of many marine fish species. Histopathological analysis, an experimental bath challenge and a field monitoring study showed that T. maritimum is associated with ‘white-patch disease’. Moreover, molecular and serological analyses performed on representative isolates revealed some degree of genetic diversity among the isolates, a finding of primary importance for epidemiological studies and the development of management and control strategies such as vaccination.

Keywords: tenacibaculosis; Tenacibaculum maritimum; Platax orbicularis; aquaculture; serotyping; 16S rRNA sequencing; multi-locus sequence analysis (MLSA); histology; qPCR diagnosis assay; experimental infection

1. Introduction

The orbicular batfish (Platax orbicularis, family Ephippidae) is a fish species inhabiting the top 30 metres depth over tropical reefs of the western Pacific [1]. It is widely distributed from the Indo-Pacific region to the Red Sea in East Africa, with a northern limit in south Japan and a southern limit in Australia and New Caledonia [2]. In French Polynesia, the batfish is highly appreciated as a food by the local population, but the wild stock is rapidly decreasing, probably as a result of high fishing pressure and climate change that severely disturb the coral communities [3]. In order to promote local, sustainable aquaculture, the governmental department of marine resources (DRM) decided to focus effort on captive breeding and rearing of orbicular batfish. Since 2011, the governmental hatchery VAIA (Vairao, Tahiti, French Polynesia) has produced eighty thousand fry annually, reared inland in bio-secure conditions (i.e., in filtered and UV treated seawater). Four production cycles...
are completed per year, allowing an annual production of 50–80 metric tons. One to two-month-old juvenile fish (mean weight, 1–10 g) are then transferred to different fish farms in lagoons. However, recurrent mortalities occur almost systematically during the first two months of growth in the lagoon cages, causing losses of 20–90% of production and threatening the sustainability of aquaculture based on this species. One to three weeks following their transfer to net cages in the lagoons, the fish show symptoms of disease, with loss of appetite, frayed fins, whitish patches on the tegument, followed by ulcers, necrosis and death. Little was known about the status of pathogens and diseases associated with this tropical fish species under local farming conditions, and, consequently, this disease was simply named ‘white-patch disease’, based on the clinical symptoms. Light microscopy examination of fragments of skin lesions revealed abundant rod-shaped and gliding bacteria potentially belonging to the genus *Tenacibaculum*.

The genus *Tenacibaculum* (family *Flavobacteriaceae*, phylum *Bacteroidetes*) currently comprises 31 validly named species (http://www.bacterio.net/tenacibaculum.html, accessed on 1 April 2021), all retrieved from marine environments [4]. Among these, *T. dicentrarchi*, *T. disolor*, *T. finnmarkense*, *T. gallaicum*, *T. maritimum*, *T. piscium* and *T. soleae* are responsible for ulcerative conditions that affect a large variety of cultured and wild marine fish species and are collectively known as tenacibaculosis [5]. Tenacibaculosis is generally associated with gross external lesions such as ulcerative and necrotic skin lesions, haemorrhagic mouth, frayed fins and tail rot [5]. The disease was originally described in 1977 in cultured red (*Pagrus major*) and black (*Acanthopagrus schlegeli*) seabream in Japan [6]. The causative agent was subsequently identified as *Tenacibaculum maritimum* (formerly *Flexibacter maritimus*) [7], a filamentous, Gram-negative, gliding bacterium. Since then, *T. maritimum* has been shown to be responsible for considerable losses in marine aquaculture worldwide, affecting a large variety of wild and cultured marine fish species. For example, *T. maritimum* was found to be associated with mortality events occurring in Atlantic salmon (*Salmo salar*) and rainbow trout (*Oncorhynchus mykiss*) in Australia [8], Chinook salmon (*Oncorhynchus tshawytscha*) in Canada [9], sole (*Solea senegalensis*) and turbot (*Scophthalmus maximus*) in Spain [10], sea bass (*Dicentrarchus labrax*) in Europe [11], Japanese flounder (*Paralichthys olivaceus*) in Japan [12], and black damselfish (*Neogyphioodon melas*) and Picasso triggerfish (*Rhinecanthus assasi*) in Egypt [13]. Although other *T. maritimum* isolates have also been retrieved from outbreaks in other countries and host fish species, pathogenicity has not been confirmed using experimental challenges in any of these examples [14,15].

In many cases, including some from the recent scientific literature [16], bacterial isolation and proper identification are usually not carried out, and tenacibaculosis diagnosis is often only based on clinical symptoms and microscopic observations. Indeed, the proper identification of the species involved in tenacibaculosis has long been hampered by a combination of factors, including the lack of selective culture medium to isolate *Tenacibaculum* strains responsible for outbreaks or to track/retrieve them from the different compartments of fish farms. Indeed, *Tenacibaculum* species are rather slow-growing and often only retrieved from external lesions (systemic infections seem rare). Consequently, bacterial cultures are almost systematically contaminated by fast-growing bacteria (e.g., *Vibrio* species), and obtaining pure cultures of *Tenacibaculum* is a challenge [17]. In addition, for over 3 decades (1977–2008), the only recognised pathogenic species in the genus was *T. maritimum*, and the recent discovery of an unforeseen diversity of pathogenic species in the genus was not followed by the development of the corresponding diagnostic tools [18].

The aims of the present study were to investigate the recurrent and acute outbreaks occurring in *Platax orbicularis* farms in French Polynesia, to characterise the causative agent using bacteriological, histological and molecular analyses, and then to conduct experimental challenges to confirm this causality, thus fulfilling Koch’s postulates.
2. Results

2.1. Microscopic Examination and Isolation of Bacteria

Two severe white-patch disease outbreaks, occurring within the first 2 months following transfer to net cages in a lagoon, were recorded in 2013 and 2016. These outbreaks occurred in two geographically distinct fish farming areas of Tahiti island (Tautira lagoon: 17°47′50″ S, 149°07′14″ W, and Vairao lagoon: 17°48′22″ S, 149°17′36″ W) with cumulative mortality reaching 80% and 62%, respectively. The main clinical signs were loss of appetite, erratic swimming and ulcerative skin lesions (Figure 1A). Wet mount examination of the skin mucus of diseased batfish revealed a significant amount of long (6.3 ± 0.6 µm) and rod-shaped bacteria (Figure 1B). In contrast, no alterations or clinical signs were observed in the internal organs.

Histopathological examination of skin lesions from moribund batfish revealed that the epidermis and dermis were severely damaged, with clusters of filamentous, Tenacibaculum-like bacteria and scattered inflammatory cells (Figure 2). In contrast, no evidence of histopathological changes was noticed in the internal organs.

Ten bacterial isolates were retrieved from samples of 10 moribund batfish exhibiting typical signs of ‘white-patch disease’ using Flavobacteriaceae-selective marine agar (Table 1). Two different colony morphotypes were observed after 48 h of incubation at 27 °C: the first morphotype consisted of pale, translucent colonies with uneven edges, extremely adherent to the agar (Figure 3A), while the second morphotype consisted of orange, opaque, diffuse and strongly iridescent colonies (Figure 3B,C).
Figure 2. Two representative cross-sections (H-E staining) of *Platax orbicularis* fingerlings affected by ‘white-patch disease’. (A,C): Severe necrosis affecting the hypodermis and dermis layers with invasion of *Tenacibaculum*-like cells visible at higher magnification (B,D) (arrows) and presence of inflammatory cells in damaged areas (asterisks).

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2.2. Genomic and Serological Characterisation

Analysis of nearly complete 16S rRNA sequences revealed that all isolates belonged to the genus *Tenacibaculum*. Seven isolates (TFA4, Aq 16–83, Aq 16–84, Aq 16–85, Aq 16–87, Aq 16–88 and Aq 16–89) shared 99.65% to 99.79% sequence identity with the *T. maritimum* type strain NBRC 15946T, with at least 98.9% coverage. The three remaining isolates (Aq 9–66, Aq 9–67 and Aq 16–91) displayed 99.37 to 99.93% sequence identity with the *T. mesophilum* type strain DSM 13764T, with at least 98.9% coverage. A tentative phylogenetic tree was drawn using MAFFT (Figure S1). The seven isolates displaying the first morphotype clustered with the *T. maritimum* type strain, while the three isolates belonging to the second morphotype clustered with the *T. mesophilum* type strain; bootstrap values were 100% and 91%, respectively.

The results of the MLSA analysis (Figure 4) performed on *T. maritimum* isolates TFA4, Aq 16–85, Aq 16–88 and Aq 16–89 showed that none matched exactly with any of the sequence types (ST) already described in the pubMLST database. They were therefore treated as belonging to new STs: ST168, which was attributed to isolate TFA4, and ST167, which was attributed to isolates Aq 16–85, Aq 16–88 and Aq 16–89. Analysis of the number of locus variants revealed that these novel STs only share three allele types (AT), corresponding to loci gyrB, infB and rlmN, which reveal genetic heterogeneity among these two groups of isolates. The single and double locus variant analyses (SLV and DLV) were fairly congruent with the phylogenetic tree based on the concatenated nucleotide sequences of the seven housekeeping genes (Figure 4). SLV analysis showed that ST168 (TFA4) shares 6/7 loci with ST2, which up to now only included strain ACC13.1 (referenced as 002 in the pubMLST database), isolated from a diseased Senegalese sole (*Solea senegalensis*) in Portugal. In the DLV analysis, TFA4 shared 5/7 loci with ST3, 4, 10, 35, 36 and 130, which essentially comprise isolates from the south of Europe (except for strain 4646, isolated in Australia). Interestingly, isolates Aq 16–85, Aq 16–88 and Aq 16–89 (ST167) formed a singleton, meaning that they displayed at least three different ATs compared with all the strains included in the pubMLST database. These results revealed the
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Serological analysis revealed two different serogroups among the *T. maritimum* isolates. Slide agglutination tests showed that strain TFA4 specifically reacted with the anti-O3 antiserum, while strains Aq 16–85, Aq 16–88 and Aq 16–89 specifically reacted with the anti-O1 antiserum.

2.3. Pathogenicity Assays Using an Immersion Challenge

All batfish that were experimentally infected with *T. maritimum* TFA4 using the immersion challenge (IF group) exhibited typical clinical signs of ‘white-patch’ disease starting from 24 h post-infection (PI), although no mortality was recorded at this time point (Figure 5). At 30 h PI, fish in this group underwent significant (chi-squared-test with simulated \(p\)-value correction, \(p = 0.017\)) mortality, with a survival probability of 94.9% compared with 100% (no death events) in the non-infected fish group (NIF). A sudden mortality event was observed in the IF group between 30 and 48 h PI, with 94.9% and 25% survival probabilities, respectively. From 82 h until the end of the monitoring period (120 h PI), no increase in mortality was observed (7.1% survival probability for the IF group at 72 h PI to 120 h PI) even though all batfish displayed typical clinical signs of the disease. Nevertheless, the intensity of ulcerative skin lesions (i.e., the number and area of whitish patches) in the IF group from 72 h to 120 h PI was lower than that observed before 72 h PI. In fish from which mucus was partially removed prior to the bath challenge (IM-IF), batfish not only experienced the highest mortality rate (all fish died), giving a survival probability of 0% from 30 h PI, but they also died significantly earlier than those of the IF group with intact skin mucus (log-rank test comparing the survival curves from 0 h PI to 30 h PI, \(p = 0.02\)). No mortality was recorded in the NIF control group during the entire trial.
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Figure 4. Neighbour-joining tree based on the concatenated nucleotide sequences of 7 housekeeping genes (3894 bp). Isolates isolated in this study are shown in red. Genomic and background information on all *T. maritimum* isolates in the pubMLST database is included. Other information: strain number, strain name, country of isolation, fish host species, year of isolation, sequence type and allelic combination.
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All skin samples that were collected from diseased fish in the IF group before the onset of mortality were found positive by real-time PCR assay, with an average load of $7.8 \times 10^8 \pm 1.4 \times 10^8$ T. maritimum bacteria per μg of total extracted DNA. In contrast, no T. maritimum was detected in any of the four analysed skin samples in the NIF group. The virulence potential of two (Aq 16–84 and Aq 16–87) and three (Aq 9–65, Aq 9–66 and Aq 9–67) isolates belonging to the species T. maritimum and T. mesophilum, respectively, was also evaluated using the same immersion challenge protocol. The results of this separate trial showed that the two T. maritimum isolates exhibited levels of virulence similar to that of isolate TFA4, whereas the three T. mesophilum isolates were avirulent at a similar infection dose ($6.1 \times 10^4$ CFU mL$^{-1}$), with no mortality recorded in groups IF and NIF during a seven-day post-infection survey (data not shown).

2.4. Kinetics of Tenacibaculum maritimum Infection during a Field Episode of Tenacibaculosis

In order to gain insight into T. maritimum pathogenesis under natural field conditions, a batch of batfish was monitored from its production under bio-secured conditions at the VAIA hatchery to its rearing in a net cage on a private farm in Tautira lagoon. Soon after the transfer to the net cage in the lagoon, a severe outbreak of ‘white-patch disease’ was observed, with the first typical signs appearing from D1 post-transfer and mortality from D3 (Figure 6).

Two peaks of mortality occurred, at D3–D6 and D13–D31. The second peak was higher, with cumulative mortalities increasing significantly from 11.2% at D13 to 74.5% at D31. No subsequent mortality occurred among the surviving batfish until the end of the study period (D36). Six days before the transfer to the lagoon cage (D-6), all sampled fish were qPCR negative for T. maritimum (Figure 7).

In contrast, the mucus of all batfish sampled just one day (D1) after the transfer to the lagoon net cages was found positive for T. maritimum by qPCR, although very different bacterial loads (mean values, $9.98 \times 10^6 \pm 1.57 \times 10^7$ cells per μg DNA) were observed. At D9, during the first stationary phase of mortality (D6–D13), asymptomatic batfish showed significantly lower bacterial loads in their mucus compared with D1 (Kruskal Wallis test; $p = 0.020$). Nevertheless, the amount of T. maritimum at D9 was significantly higher ($p = 0.047$) in batfish exhibiting clinical signs of tenacibaculosis compared with asymptomatic fish. During the second peak of mortality, the discrepancy between asymptomatic and symptomatic batfish was even more pronounced ($p = 0.016$). At the end of the mortality events (D36), T. maritimum DNA was absent from most surviving fish sampled.
(8/10). Indeed, only two asymptomatic batfish among the 10 tested were found positive by qPCR but at very low levels ($8.36 \times 10^2 \pm 6.02 \times 10^2$ cells per µg DNA), signalling the end of the outbreak.

Figure 6. Cumulative mortalities of a batch of batfish during a natural outbreak following their transfer to a net cage in Tautira lagoon. Fish were 10 g (mean weight) and reared at an initial density (D0) of 167 fish/m³.

Figure 7. Kinetics of *Tenacibaculum maritimum* bacterial cells quantified by qPCR in the mucus of asymptomatic (white boxes) and symptomatic (grey boxes) batfish from D-6 (at VAIA hatchery) to D36 post-transfer to net cage in the lagoon. Quantification results are expressed in numbers of *T. maritimum* cells per µg of total extracted gDNA. Because zero values (no detection) cannot be represented on a logarithmic scale, an arbitrary value of 1.1 was assigned to these negative results. Each box-plot shows mean (white circle), the 25th to 75th percentile (rectangular box), the minimum and maximum values (dots at the extremities), as well as individual quantification (black dots) from 5 to 10 batfish per sampling time and group of batfish.
qPCR results on the liver of the same batfish sampled for their mucus revealed that only one fish among the 50 tested, sampled at D1, was positive for *T. maritimum* at a low level (1.19 × 10^4 cells per µg gDNA). Similar results were obtained with the posterior intestine: only seven batfish were found qPCR positive, with the low value of 1.11 × 10^4 ± 1.37 × 10^4 cells per µg gDNA.

3. Discussion

Orbicular batfish aquaculture in French Polynesia started in 2004, and the first symptoms of ‘white-patch’ disease were observed in 2006. In this study, we showed that the ‘white-patch disease’ decimating farmed *Platax orbicularis* is associated with *T. maritimum* infection. To our knowledge, this is the first time that this bacterium has been isolated in French Polynesia and also the first time it has been retrieved from batfish. *T. maritimum* has been associated with a large variety of marine fish species: 38 according to Nowlan et al. (2020) [5], including the orbicular batfish (this study). Surprisingly, *Platax orbicularis* is the only tropical fish in which this pathogen has been reported so far. Nevertheless, the range of susceptible hosts for this bacterium is probably underestimated. In French Polynesia, tenacibaculosis has dramatic consequences for batfish farms, which often suffer mortality levels over 50%. In addition, the disease may also be of serious concern regarding the diversification of aquaculture programs launched by local authorities.

In this study, we described a reproducible bath challenge protocol that demonstrated that *T. maritimum* is able to infect orbicular batfish by immersion, thus fulfilling Koch’s postulates. Experimental infection procedures using immersion challenges have been broadly used with fish pathogens in recent years because they are likely to mimic the natural infection process more accurately than injection challenges. In particular, immersion does not bypass the first line of fish defence (i.e., the skin mucus barrier), unlike the more common subcutaneous, intraperitoneal and intramuscular injection routes. It was also reported that, compared with immersion, some injection challenges fail to induce tenacibaculosis [11,19–21] or can lead to high mortality rates in negative controls due to stress and local lesions caused by the injection [22]. However, comparative analyses of challenge protocols are rather difficult to perform due to the many factors reported influencing pathogenicities, such as the bacterial strain [9,20,21], culture conditions (i.e., growth medium and temperature), infection dose [12,19], duration of immersion [23], physical [21] and chemical characteristics of seawater, zootechnical practices (e.g., fish density and animal feed), the host fish species [8], physiological status (e.g., age [21]) and genetic background (e.g., susceptible, resistant). In this study, no significant difference in mortality rates was observed when batfish were infected with isolates TFA4, Aq 16–87, or Aq 16–84, although these isolates differed in some genetic traits. However, due to the high virulence of these isolates in our immersion challenge model, further studies using a lower dose or shorter immersion time might reveal virulence differences between these strains. Although physical alteration of the fish skin was not necessarily noticed before morbidity and mortality, batfish with impaired mucus developed clinical signs more rapidly and experienced higher mortality rates (100% mortality at 23 h post infection) than those where mucus was intact. These results are in agreement with similar studies performed on other fish species [11,23]. Indeed, mucus has been largely documented as an important component of the fish innate immune system and a physical and chemical barrier against pathogens [24].

Some of the isolates from diseased batfish were shown to belong to another *Tenacibaculum* species, *T. mesophilum*, a bacterium initially retrieved from a marine sponge. In addition, *T. mesophilum* strain HMG1 was shown to degrade malachite green, an antimicrobial that has long been used in aquaculture but is now banned in many countries [25]. Immersion challenges performed with isolates Aq 9–66 and Aq 9–67 showed that both strains were totally avirulent. Further studies would be needed, however, to determine whether *T. mesophilum* isolates can play a role in the pathogenesis of tenacibaculosis, primarily caused by *T. maritimum*, by acting as secondary colonisers of the lesions.
Although this study was conducted with only 10 isolates, an unexpected diversity of *T. maritimum* isolates was found. Our results demonstrate the presence of two distinct groups: isolates Aq 16–85, Aq 16–88 and Aq 16–89, belonging to serotype O1 and to sequence type ST167; and isolate TFA4, belonging to serotype O3 and to sequence type ST168. Such diversity among *T. maritimum* isolates was also noticed among Australian isolates (Figure 4). In agreement with Van Gelderen et al. (2010) [26], no correlation between serotype and geographic distribution was observed in the present study. In contrast with some authors [27,28] who reported a correlation between the host and the serotype of *T. maritimum*, our results are more consistent with other studies [9,26,29] that revealed that several different serotypes could be found associated with the same fish species.

**Table 1.** List of bacterial isolates retrieved from farmed *Platax orbicularis* affected by ‘white-patch’ disease, with their sources and characteristics. Virulence was evaluated by experimental bath challenge on *T. maritimum*-free batfish. Significant differences (*p* < 0.05) in mortality rate (see paragraph 3-3) between non-infected and infected fish are indicated by ‘yes’ or ‘no’. Isolate identification was performed using the EzBioCloud software [30], based on >99% identity of their 16S rRNA sequences with the closest type strain. ST refers to the MLST sequence type. n/a: not analysed.

| Isolate | Source and Date of Isolation | GPS Location | 16S rDNA GenBank Accession Number | Bacterial Species | Virulence | ST |
|---------|-----------------------------|--------------|----------------------------------|-------------------|-----------|----|
| TFA4    | Skin lesions, Tautira lagoon, Tahiti, 2013 | 17°47′50″ S, 149°07′14″ W | MW690171 | *T. maritimum* | yes | ST168 |
| Aq 9–66 | Skin lesions, Tautira lagoon, Tahiti, 2013 | 17°47′50″ S, 149°07′14″ W | MW690177 | *T. mesophilum* | no | |
| Aq 9–67 | Skin lesions, Tautira lagoon, Tahiti, 2013 | 17°47′50″ S, 149°07′14″ W | MW690178 | *T. mesophilum* | no | |
| Aq 16–83 | Skin lesions, Vairao lagoon, Tahiti, 2016 | 17°48′22″ S, 149°17′36″ W | MW690172 | *T. maritimum* | n/a | n/a |
| Aq 16–84 | Skin lesions, Vairao lagoon, Tahiti, 2016 | 17°48′22″ S, 149°17′36″ W | MW690173 | *T. maritimum* | yes | n/a |
| Aq 16–85 | Skin lesions, Vairao lagoon, Tahiti, 2016 | 17°48′22″ S, 149°17′36″ W | MW690174 | *T. maritimum* | n/a | ST167 |
| Aq 16–87 | Skin lesions, Vairao lagoon, Tahiti, 2016 | 17°48′22″ S, 149°17′36″ W | MW690175 | *T. maritimum* | yes | n/a |
| Aq 16–88 | Skin lesions, Vairao lagoon, Tahiti, 2016 | 17°48′22″ S, 149°17′36″ W | MW690176 | *T. maritimum* | n/a | ST167 |
| Aq 16–89 | Skin lesions, Vairao lagoon, Tahiti, 2016 | 17°48′22″ S, 149°17′36″ W | MW690180 | *T. maritimum* | n/a | ST167 |
| Aq 16–91 | Skin lesions, Vairao lagoon, Tahiti, 2016 | 17°48′22″ S, 149°17′36″ W | MW690179 | *T. mesophilum* | n/a | |

Additional work is needed for an in-depth exploration of the genetic diversity of *Tenacibaculum* isolates associated with batfish in French Polynesia in order to evaluate their virulence potential and develop management and disease control strategies. Because the natural ecology of *T. maritimum* is still unknown, more in-depth epidemiological studies will also be necessary to decipher the mode of transmission and the natural route of infection of this pathogen.
4. Materials and Methods

4.1. Sampling of Diseased Fish

Ten symptomatic orbicular batfish (mean weight $5.1 \pm 2.3$ g, crop density of 170 fish/m$^3$) were collected from two different farms located in Tahiti island. Fish were transported alive in the microbiology laboratory, using buckets filled with seawater and protected from direct sunlight for further examination. They were sampled during two severe outbreaks in 2013 and 2016 that had caused $>50\%$ cumulative mortality just 2 weeks after the fish had been transferred from the VAIA hatchery to net cages in the Tahiti lagoon. The sea temperature was $26.5\, ^\circ\text{C}$ in 2013 and $27\, ^\circ\text{C}$ in 2016, and water salinity was consistently $3.3\%$. All fish showed erosion and ulceration of the skin surface. After being euthanised with an overdose of Benzocaine ($150 \text{ mg L}^{-1}$), they were examined by microscopy and microbiological and histological techniques. This method of euthanasia, reproducible and safe to the operator, induces a depression of the central nervous system activity, rapid unconsciousness and death of P. orbicularis, without compromising further diagnostic techniques.

4.2. Direct Microscopic Examination and Isolation of Bacteria

Skin lesion scrapings from moribund batfish were performed using sterile surgical scalpels. Wet mount preparations of these samples were then examined under a light microscope (Leica DM 1000 LED). For bacterial isolation, a sterile swab cotton-tipped applicator (COPAN) was used. Smears of skin samples were deposited onto plates of Flavobacteriaceae-selective marine agar (FSMA) developed by an accredited veterinary diagnostic laboratory (Labofarm, Loudéac, France). A total of ten dominant bacterial isolates were isolated after sub-culture (Table 1). Bacterial isolates were cryo-conserved at $-80\, ^\circ\text{C}$ in Marine Broth medium supplemented with 15% of glycerol.

4.3. Histopathological Examination

Skin fragments of approximately 1 cm$^2$, as well as internal organs (anterior kidney, spleen, liver), were collected from four fish showing typical skin lesions of ‘white-patch’ disease using a sterile surgical scalpel. Samples were fixed in Davidson’s solution (25% formaldehyde, 37.5% ethanol, 12.5% acetic acid and 25% water) for 48 h at room temperature, then washed and kept in 70% ethanol [31]. Skin samples were progressively dehydrated in an ascending series of alcohol (70 to 100% ethanol) followed by a xylene bath, using dehydration automate (Leica, ASP 3005), then embedded in paraffin, cut into 3 $\mu$m sections using a rotary microtome (Microm HM 340E, Thermo Fisher Scientific, Waltham, MA, USA) and stained with haematoxylin–eosin (H-E) using a fully-automated integrated stainer (Leica, CV5030 autostainer XL). Several sections were analysed to ensure reproducibility, using a Leica DM 1000 LED microscope equipped with a Dino-Lite camera (AnMo Electronics).

4.4. Molecular and Serological Studies

The 16S rDNA sequences were PCR-amplified using the universal 27F and 1492R primers (Table S1) on 5 $\mu$L of a boiled bacterial colony. Briefly, a bacterial colony was resuspended in 100 $\mu$L of molecular-free DNA water, heated 10 minutes at 98 $^\circ\text{C}$ and used immediately as a DNA template. The purity and length of the amplicons were verified by agarose gel electrophoresis. Amplicons were Sanger sequenced by GATC-biotech (https://eurofinsgenomics.eu/, accessed 20 January 2021) using the six universal sequencing primers listed in Table S1.

For each isolate, the six sequences were visualised and aligned to create a consensus sequence (with $\geq 2\times$ coverage over 80% of the sequences) using the Benchling software (2020). For primary taxonomic assignation, the 16S rRNA consensus sequences were searched against the EzBioCloud database [21] (accession numbers are given in Table S2). In addition, a tentative phylogenetic tree was constructed using the MAFFT online service [32]. The evolutionary distance was calculated using 1000 bootstrap replicates (Figure S1).
In order to characterise the genetic diversity of presumptive *T. maritimum* isolates in greater depth, multi-locus sequence analysis (MLSA) was performed on four selected isolates (TFA4, Aq 16–85, Aq 16–88, Aq 16–89) using sequences retrieved from their draft genomes [18]. These isolates were selected according to their background information: Aq 16–85, Aq 16–88 and Aq 16–89 were sampled from three different infected fish during an outbreak at the Vairao fish farm in 2016, while isolate TFA4 was isolated from a symptomatic fish at the Tautira fish farm in 2013 (Table 1). The MLSA profile defined by Habib et al. (2014) [33] consists of the sequences of seven housekeeping genes (*atpA, gyrB, dnaK, glyA, infB, rlmN* and *tgt*). The profiles of the new allele and sequence types (ATs and STs, respectively) were generated and analysed using the *Tenacibaculum* pubMLST database (https://pubmlst.org/tenacibaculum/, accessed on 10 December 2021) [34]. Results were visualised using the incremented Interactive Tree of Life (iTOL) v3 tool [35].

In order to characterise the isolates identified as *T. maritimum* more precisely, their serotype was determined as described by Avendaño-Herrera et al. (2004) [36]. This method uses a slide agglutination test and a dot blot assay on both whole-cell preparations and heat-stable O antigens of each strain. Antisera against serovars O1 (PC503.1), O2 (PC424.1) and O3 (ACC13.1) were used in all assays.

4.5. Experimental Infection by Immersion and Quantification of *Tenacibaculum maritimum* in Mucus Samples by Real-Time qPCR

A batch of *Platax orbicularis* fingerlings (mean weight 9.7 ± 2.6 g) reared at the VAIA bio-secured hatchery was transferred for acclimatisation to a 1 m³ fibreglass tank containing seawater (salinity, 32 PPT; water temperature, 26–27 °C) for 10 days. Prior to infection, 494 fish were randomly selected and transferred to ten 150 L tanks (49 to 50 fish per tank) filled with 5 µm filtered seawater. Three groups were tested in three or four replicates according to the condition: (i) non-infected fish (NIF), (ii) fish infected with *T. maritimum* isolate TFA4 (IF) and (iii) fish with impaired mucus (IM) infected with *T. maritimum* isolate TFA4 (IM-IF). The mucus of the latter fish was partially removed by gently wiping one side of the fish with a sponge soaked in filtered seawater. Fish were challenged with a pure culture of isolate TFA4 obtained by incubation at 27 °C for 48 h (stationary phase as determined by preliminary growth monitoring assays using ten-fold serial dilutions) in autoclaved nutrient broth composed of 4 g L⁻¹ peptone and 1 g L⁻¹ yeast extract (Becton, Dickinson and Co., Sparks, MD, USA) in 5 µm filtered seawater under orbital shaking at 200 rpm. Bath challenges were performed for two hours with isolate TFA4 at a final concentration of 5.3 × 10⁴ CFU mL⁻¹ for groups IF and IM-IF or with nutrient broth in the case of the mock-treated NIF control group. The infected fish were then rinsed twice with filtered seawater to remove all non-adherent bacteria, and fish from the NIF control group were manipulated in the same way. Mortality was monitored twice daily from day 0 (D0) until day 5 (D5), at which point the fish were euthanised using 150 mg L⁻¹ Benzocaine. Any fish that died or were found moribund over the experimental period (D0–D5) were promptly removed from the tanks during the monitoring. The non-parametric Kaplan–Meier method (R package survival) was used to test for differential survival performances among groups at the same date or within groups throughout the sampling period. Differences were considered significant at *p* < 0.05.

At 24 h post-infection, before the onset of mortality, four fish displaying skin lesions were randomly sampled from the two infected groups (IF and IM-IF) in addition to four fish from the NIF group. These were used to quantify *T. maritimum* cells in fish mucus using TAQMAN real-time PCR (see primers and probe in suppl. Table 1) following the protocol developed by Fringuelli et al. (2012) [37] with minor modifications. Briefly, skin mucus samples were obtained from lesions on symptomatic fish using cotton swabs (COPAN), directly diluted in 1.5 mL microcentrifuge tubes containing 0.5 mL of lysis solution (100 mM NaCl, 10 mM Tris pH 8, 25 mM EDTA, 0.5% SDS and 200 µg ml⁻¹ proteinase K) and incubated overnight at 55 °C. DNA was extracted using the conventional phenol/chloroform/isoamyl alcohol (25/24/1) method. DNA quantity and purity were assessed using a NanoDrop ND
1000 spectrophotometer (Thermo Fisher Scientific). In order to obtain a standard curve, bacterial cells of isolate TFA4 from a stationary phase culture in nutrient broth were enumerated using a Malassez counting chamber (2.35 × 10⁸ bacteria ml⁻¹), and DNA from 1 ml of the bacterial suspension was extracted. The DNA was then spiked at a final concentration of 1.33 ng µL⁻¹ in salmon sperm gDNA (SSD, Thermo Fisher) at 10 ng µL⁻¹ in artificial seawater (ASW, Sigma), then serially diluted 10-fold in SSD at 10 ng µL⁻¹ in ASW. A linear range of values was obtained for PCR amplification on an Mx3000 Thermocycler (Agilent) using Brilliant III Ultra-Fast QPCR Master Mix (Agilent) following the supplier’s recommendations (5 µL DNA at 10 ng µL⁻¹ in a total reaction volume of 20 µL), with six successive sample 10-fold dilutions tested in triplicate. Cycle threshold (Ct) values ranged from 16.05 to 33.14, corresponding to 1.44 × 10⁵ to 2.06 × 10⁵ cells of isolate TFA4 per PCR well, while correlation (linear regression with r² coefficient) and qPCR reaction efficacy were 0.995 and 99.9%, respectively.

4.6. Detection and Quantification of *Tenacibaculum maritimum* during a Field Episode of Tenacibaculosis

Juvenile batfish (mean weight 10 ± 3.3 g) reared in the bio-secure facilities of the VAIA hatchery were carefully transferred (D0) to the Tahiti Fish Aquaculture farm in Tautira lagoon and kept in a single net cage of 1 m³ (167 fish/m³ density). Ten fish were collected at five sampling times: day 6 before transfer (D-6) (i.e., in the VAIA hatchery) and D1, D9, D17 and D36 post-transfer to Tautira lagoon. When gross signs of ‘white-patch disease’ were observed, five moribund fish (symptomatic) and five apparently healthy (asymptomatic) ones were sampled. These fish were euthanised as detailed above, and the liver, posterior intestine and some skin mucus (collected with a cotton swab in the lesion area in the case of symptomatic fish) were individually and aseptically sampled and preserved in 500 µL RNAlater (Ambion) at −80 °C. Approximately 100 mg of tissue were used to quantify *T. maritimum* by qPCR [37]. Throughout this survey, no curative treatments were given, mortality was monitored daily, and moribund animals (euthanatised with an overdose of benzocaine) or dead fish were removed and discarded.

**Supplementary Materials:** The following are available online at: https://www.mdpi.com/article/10.3390/pathogens11020131/s1, Figure S1: Phylogenetic relationships of 16S rDNA nucleotide sequences of the isolates recovered in this study and the type strains of all Tenacibaculum species (See Table S2 for accession number); Table S1: List of PCR primers and probe used in this study; Table S2: Accession number of the type strains used in Figure S1.

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