The extensive intergenerational molecular effects of ocean acidification on the olfactory epithelium transcriptome of a marine fish are associated with a better viral resistance

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Abstract

Background

Progressive climate-induced ocean acidification (OA) impacts marine life in ways that are difficult to predict but are likely to become exacerbated over generations. Although marine fishes can balance internal acid-base homeostasis efficiently, indirect ionic regulation effects that alter neurosensory systems can result in behavioural abnormalities. In marine invertebrates, OA can also affect immune system function, but whether this is the case in marine fishes of ecological and commercial importance is not yet understood. Farmed fish are highly susceptible to disease outbreak yet strategies for overcoming such threats in the wake of OA are wanting. Here, we exposed two generations of the European sea bass (*Dicentrarchus labrax*) to end-of-century predicted CO$_2$ levels (IPCC RCP8.5), with parents being exposed for four years and their offspring for two years. Our design included a transcriptomic analysis of the olfactory rosette (collected from the F1 offspring) and a viral challenge (exposing F1 offspring to betanodavirus) where we assessed survival rates.

Results

We discovered long-term intergenerational molecular trade-offs in both sensory and immune systems. Specifically, RNA-Seq analysis of the olfactory rosette, the peripheral olfactory organ, from two-year-old F1 offspring revealed extensive regulation in genes involved in ion transport and neuronal signalling, including GABAergic signalling. We also detected extensive OA-induced intergenerational up-regulation of genes associated with odour transduction, synaptic plasticity, neuron excitability and wiring and down-regulation of genes involved in energy metabolism. In addition, intergenerational exposure to OA induced up-regulation of genes involved in innate antiviral immunity (pathogen recognition receptors and interferon-stimulated genes) in combination with down-regulation of the protein biosynthetic machinery. Consistently, OA-exposed F1 fish challenged with betanodavirus, which causes damage to the nervous system of marine fish, had acquired improved resistance.

Conclusion

F1 exposed to OA-intergenerational acclimation showed superior viral resistance, though as their metabolic and odour transduction programs were altered, odour-mediated behaviours might be consequently altered. Our results reveal that trade-offs in adaptive plastic responses is a core feature of the olfactory epithelium transcriptome in OA-exposed fish, suggesting that intergenerational plasticity propagate with progressive exposure to OA and will have important consequences for how cultured and wild fish interacts with its environment.

Background
Atmospheric CO$_2$ concentration has increased from preindustrial levels of 280 ppm to the current value of 414 ppm (Dlugokencky and Tans 2021), and is expected to reach ~1 000 ppm by the end of this century (Caldeira and Wickett 2003; IPCC 2014; Bindoff et al. 2019). These changes in turn lead to ocean acidification (OA), including changes in the carbonate system equilibrium such as reduced carbonate ion concentrations and pH. According to the future warming scenario RCP 8.5, seawater pH will decrease by 0.3-0.4 units by 2100 in the North Atlantic Ocean (Caldeira and Wickett 2003; Orr et al. 2005; IPCC 2014).

OA alone or coupled with ocean warming poses global-scale threats to marine life ranging from the invisible scale of microbes to ecosystem levels (Danovaro et al. 2011; Hassenrück et al. 2016; Hutchins and Fu 2017; Cavicchioli et al. 2019). Given their efficient capacity for maintaining acid-base homeostasis, fishes have long been considered to be robust organisms capable of tolerating OA (Claiborne et al. 1999; Claiborne et al. 2002; Heuer and Grosell 2014; Esbaugh 2017). However, reported indirect impacts of OA include both inter- and intra-specific changes in sensory behaviours (Clements and Hunt 2015), which result from directly altered neurosensory systems (Munday et al. 2009; Dixson et al. 2010; Chung et al. 2014; Hamilton et al. 2014; Rossi et al. 2016; Ashur et al. 2017; Tresguerres and Hamilton 2017; Holmberg et al. 2019; Paula et al. 2019). Ultimately, these affect reproduction, fitness and mortality (Nilsson et al. 2012; Chivers et al. 2014; Faria et al. 2018; Servili et al. 2020). Although such effects are complex and vary with both biological traits and methods employed (Williamson et al. 2020), sensory behavioural alterations induced by acid-base regulation are generally explained as a GABAergic system dysfunction due to an inversion of the Cl$^-$/HCO$_3^-$ channels across neuronal membranes (Nilsson et al. 2012; Schunter et al. 2019). However, molecular and electrophysiological approaches also suggest that changes in calcium homeostasis and decrease in synaptic plasticity underlie OA impacts on synaptic activity in the olfactory bulb (Porteus et al. 2018). The capacity of the fish neurosensory system to cope with OA depends on the species considered and the parental phenotype (Lai et al. 2017; Tsang et al. 2020). In addition, OA-induced effects can vary over multiple generations (Miller et al. 2012; Munday 2014), as observed in Atlantic cod where parental acclimation to OA increased survival of offspring under elevated CO$_2$ (Stiasny et al. 2016). Whether such intergenerational acclimation is correlated with increased offspring viability in other economically exploited species is an open question.

Relative to the well-recognized influence of ocean warming on the emergence of pathogenic species and spread of viral diseases (Vezzulli et al. 2013; Cavicchioli et al. 2019), the influence of OA has received less attention, although it is known that virus abundance and virus-host interactions are altered by OA (Liu et al. 2010; Traving et al. 2014; Tangherlini et al. 2021). Even though the environment influences disease spread, it can also modulate the host immune responses to mitigate infection. Understanding how the environment, namely OA, influences the host immune system, and more specifically the sensitivity to pathogens, will become critical for predicting future changes in both population dynamics and aquaculture.

Earlier studies reported that OA can alter immune system parameters in marine invertebrates (Bibby et al. 2008; Mackenzie et al. 2014; Wang et al. 2016; Leite Figueiredo et al. 2016; Sun et al. 2017; Castillo et al. 2017; Migliaccio et al. 2019; Wang et al. 2021). Much less is known in marine fishes (Bresolin De Souza...
et al. 2014; Bresolin de Souza et al. 2016), and especially species of ecological and commercial importance, such as the European sea bass (*Dicentrarchus labrax*). Recently, the *cbln11* gene, which is involved in the pathogen defence, was reported to be up-regulated in the olfactory rosette of two successive generations of *D. labrax* exposed to OA, suggesting that OA can induce immune system changes (Mazurais et al. 2020). As the olfactory epithelium is in constant contact with seawater, it functions as a gateway for invading pathogens (Mori et al. 2005) and is capable of eliciting a strong antiviral response (Tacchi et al. 2014; Sepahi et al. 2019). However, although a transcriptomic analysis of the olfactory bulb of *D. labrax* after seven days of exposure to OA within a generation detected significant expression changes in genes involved in synaptic development, growth and ion transport, genes involved in the immune system were not significantly altered (Porteus et al. 2018).

Here, we reasoned that longer-term exposure to OA would serve as a proxy for *D. labrax* adaptations to OA, and that a transcriptome analysis of the olfactory epithelium from F1 generation offspring would help us understand, at a molecular level, how fish adapt over several years and across generations. We also asked how such OA-induced adaptations impacted on pathogen susceptibility by conducting an experimental betanodavirus challenge. Betanodavirus infections in *D. labrax* occur with high frequency at every life stage in aquaculture facilities (Le Breton et al. 1997; Munday et al. 2002) and causes viral nervous necrosis (VNN) by infecting the brain and retina, resulting in mass mortality and a range of neurological and behavioural abnormalities (Breuil et al. 1991; Le Breton et al. 1997; Munday et al. 2002; Costa and Thompson 2016). As betanodavirus can infect more than 120 species of marine and freshwater fishes as well as invertebrates (Bandín and Souto 2020), an outbreak represents a significant risk for breeding, rearing, and harvesting of fish in all types of water environments.

**Results**

Here, we designed a long-term experimental paradigm in which parental *Dicentrarchus labrax* (the F0 generation), were reared in the same pH conditions as their offspring (the F1 generation) (control treatment pH8.0, pCO$_2$ = 590 µatm or OA treatment pH7.6, pCO$_2$ = 1520 µatm, as predicted by 2100 in the IPCC RCP8.5 scenario). To generate the F1 generation, and as previously described, we collected gametes from 4-year-old F0 fish that had been reared from larval stage to adulthood in control or OA conditions (Crespel et al., 2017; Cominassi et al., 2019, 2020; Howald et al., 2019; Mazurais et al., 2020). For each pH treatment, we collected and pooled sperm and eggs from 20 males and 6 females (see methodological summary in figure 1). We then performed *in vitro* fertilization, after which the resulting F1 generation was returned to OA or control conditions (Crespel et al. 2017; Cominassi et al. 2019; Howald et al. 2019; Cominassi et al. 2020; Mazurais et al. 2020) (figure 1). After 2 years, we split our experimental F1 cohorts into two experimental arms; one in which we performed gene expression profiling of olfactory rosette tissue from F1 juveniles (7 each from control and OA rearing conditions, figure 1), and a second arm in which we conducted a betanodavirus challenge, as previously described, to investigate how long-term OA conditions impact on viral infection parameters in *D. labrax* (figure 1).
**Viral challenge**

In non-infected fish, and as expected, we did not detect any mortality, clinical symptoms or lesions. However, in infected fish, symptoms of VNN disease i.e. darkening of the body, whirling swimming and hyperactivity, were evident after five days in the control treatment (dpi 5, pH8.0) but appeared three days later in fish exposed to the OA treatment (dpi 8, pH7.6). Similarly, mortality on-set was staggered such that fish started dying at 7 dpi in the control treatment and at 9 dpi in the OA treatment), but after 25 days, no additional mortality was observed (figure 2). When we performed a Kruskal–Wallis test together with a Dunn's post hoc test to determine the Mean Day to Death (MDD), we found this to be 10.7 dpi in control and 12.8 dpi in OA sea water conditions. At the peak of mortality, the presence of the virus in the pool of organs was indistinguishable between treatments. At the end of the challenge, more fish survived infection in OA relative to control conditions, as reflected by a significantly higher survival rate (Kruskal-Wallis, H = 3.84, p-value = 0.05 in OA treatment (68%) than in the control (38%) (figure 2).

**Global gene expression profiling**

After a 24h fasting period, we dissected out the olfactory rosette from F1 juveniles (7 each, reared for 2 years in either OA and control conditions), extracted RNA and verified RNA integrity as described in Materials and Methods. cDNA library construction for the 14 samples and RNA-sequencing was performed by the GenomiX sequencing platform (MGX, Montpellier, France). Across the 14 libraries, Illumina sequencing generated a total of 772,449,468 reads with an average of 55,450,000 reads per library. More than 90% of the sequencing reads for each library (except for one sample that was 72%) could be mapped to *D. labrax* reference genome. Of the total 26,721 transcripts, 9,112 were differentially expressed (with an adjusted p-value ≤ 0.01), with 4,515 (49.6%) up-regulated and 4,598 (50.4%) down-regulated in the OA-exposed cohort (listed in additional file 2). To validate the RNA-Seq results, we performed qPCR analysis of the *samhd1*, *gvinp1*, *tlr3*, *cxcl14* and *nfat5* genes through which we obtained robust coefficient of correlation metrics (*r*$_{samhd1}$:0.90, *r*$_{gvinp1}$:0.74, *r*$_{tlr3}$:0.91, *r*$_{cxcl14}$:0.91 and *r*$_{nfat5}$:0.89).

**Regulation of pH, bicarbonate transport and chloride homeostasis**

Overall, we found that genes involved in regulation of pH, bicarbonate transport and chloride homeostasis were significantly differentially expressed (Table 1, additional file 3). Among the genes involved in pH regulation, members of the slc9a family genes were either up-regulated (*slc9a1*, *slc9a7*) or down-regulated (*slc9a2*, *slc9a9*) in OA-reared F1 juveniles whereas genes involved in bicarbonate transport, all slc4a family members, were up-regulated. 27 chloride transport-associated gene transcripts were up-regulated by OA exposure, of which the majority (22 transcripts) included Slc12a family genes (K-Cl cotransporters) and GABA receptors.
Table 1

| GO term                     | Regulation by OA | Number of transcripts | Regulated genes                                      |
|-----------------------------|------------------|-----------------------|-------------------------------------------------------|
| Regulation of pH            | Up               | 3                     | slc4a10, slc9a1, slc9a7                               |
|                             | Down             | 4                     | cahz, edn1, slc9a2, slc9a9                           |
| Bicarbonate transport       | Up               | 3                     | slc4a7, slc4a8, slc4a10                              |
| Chloride homeostasis        | Up               | 22                    | ano1, ano3, ano5, ano8, cftr, clcn2, clcn6, clcn7, clic4, clic5, gabra2, gabra4, gabrg2, gabrn2, girb, slc4a10, slc12a2, slc12a4, slc12a5, slc12a7, slc12a10 |
|                             | Down             | 5                     | abcc4, clcn5, fam131a, icln                          |

**Neuronal plasticity and activity**

As predicted from an expression analysis of olfactory epithelium, GO analysis identified enrichment of up-regulated genes related to neuronal cell structures in OA-treated tissue (e.g. “neuron projection”, “synapse”, “postsynaptic density”, “voltage-gated sodium channel complex”, “neuronal cell body”) (additional file 4). For instance, within the 90 genes related to “neuron projection”, we found axonal function associated genes such as kinesin heavy chain isoform 5c-like (kif5c), semaphorins, ephrin type-b receptor 2 (ephb2), cyclin-dependent kinase 5 (cdk5), amyloid beta a4 precursor protein-binding family b member 1-like (apbb1) and growth associated protein 43 (gap43) (Table 2, additional file 5). Based on biological process GO enrichment, we found that genes associated with “signal transduction” (516 transcripts), including “regulation of small GTPase mediated signal transduction” (95 transcripts) or “cell surface receptor signalling pathway” (160 transcripts), were over-represented with OA-induced up-regulated genes (figure 3A, additional file 6). Among the 187 genes involved in “intracellular signal transduction”, we found known neuronal plasticity pathways such as the Neurotrophin TRK receptor signalling pathway (19 transcripts), the Glutamate signalling pathway (13 transcripts) and the Ephrin signalling pathway (9 transcripts) (Table 2, additional file 7). Genes related to “regulation of synaptic plasticity” (22 transcripts) were also enriched within OA-induced up-regulated genes (figure 3A, additional file 6), and genes coding for calcium calmodulin-dependent protein kinase type ii subunit beta-like (camk2b), synaptotogyrin-1 (syngr1) and neuroplastin-like (nptn) proteins exhibited significantly elevated gene expression levels (additional file 8). In addition, biological processes related to “regulation of action potential”, were also over-represented in OA-treated tissue (figure 3A, additional file 6) included genes involved in potassium, sodium and calcium ion transport (e.g. scn2b, scn1a, cacna1d, kcn2, kcn5) (Table 2, additional file 9). Consistently, “regulation of neurotransmitter levels” genes were enriched within genes up-regulated by OA, including genes involved in the GABAergic signalling pathway (up-regulated: cacna1a, gabra2, nf1, slc6a13) (Table 2, additional file 10).
Table 2
Examples of enriched gene ontologies and associated genes involved in neural plasticity and activity that were up-regulated (p≤0.01) by ocean acidification in the olfactory rosette of F1 European Sea Bass (Dicentrarchus labrax) juveniles. Several transcripts may originate from the same gene. This list is given an example, with no attempt to be exhaustive. See additional files 5-10 for an enlarged version that include gene description and associated GO terms. GO: Gene Ontology.

| GO term                                      | Subcategory                   | Number of transcripts | Examples of regulated genes                                      |
|----------------------------------------------|--------------------------------|-----------------------|------------------------------------------------------------------|
| Neuron projection                            | Axon guidance                 | 15                    | apbb1, cdk5, epha5, epha8, ephb2, kif5c, gap43, sptbn4, tgbf2     |
|                                              | Axon extension                | 3                     | cdk5, map1b, sema3a                                             |
| Intracellular signalling                      | Neurotrophin TRK receptor sig.| 19                    | adcy6, arhgef7, calm1, erbb4, gab1, kras, ngef, shc1, vav2       |
| transduction neuroplasticity                 | pathway                       |                        |                                                                  |
|                                              | Glutamate signaling pathway   | 13                    | gnaq, glur2b, gria1a, gria3b, gria4k, grik2, grik5, grin3a, grind2d |
|                                              | Ephrin signaling pathway      | 9                     | chn1, efna5, ek1, epha5, epha7, ephb1, ephb2, etk2              |
|                                              | Regulation of synaptic plasticity | 22                  | camk2b, cplx2, grik2, kcnn2, nf1, nptn, shank3, syngr1          |
| Regulation of action potential               |                               | 21                    | ank2, ank3, cacna1d, knip1, kcnc2, ryr2, scn1a, scn2b           |
| Reg. of the neurotransmitter levels          |                               | 34                    | cacna1a, gabra2, napb, nf1, rims1, slc6a13, snap25, stx1a, trpm7 |

Metabolism

GO processes enriched within down-regulated genes were primarily related to metabolism (figure 3B, additional file 11). A large number of these are implicated in biosynthetic processes (465 transcripts); peptide biosynthetic process (105 transcripts), regulation of translation (43 transcripts), carbohydrate derivative biosynthetic process (89 transcripts) and ATP biosynthetic process (20 transcripts) (Tables 3, additional file 12 and 13). Other down-regulated biological processes included oxidation-reduction process (266 transcripts), catabolic process (184 transcripts), cell division (62 transcripts) and viral process (16 transcripts) (figure 3B, additional file 14). Consistently, down-regulated gene programs included the ribonucleoprotein complex (166 transcripts), the mitochondria (217 transcripts) - including the mitochondrial respiratory chain complex I (12 transcripts), the mitochondrial matrix (29 transcripts), the catalytic complex (228 transcripts) and the oxidoreductase complex (29 transcripts) (additional file 15).
Table 3
Examples of enriched gene ontologies and associated genes related to metabolism, viral process and cell division that were down-regulated (p-value ≤ 0.01) by ocean acidification in the olfactory rosette of F1 European Sea Bass (Dicentrarchus labrax) juveniles. Several transcripts may originate from the same gene. This list is given as an example, with no attempt to be exhaustive. See additional files 12-14 for an enlarged version that include gene description and associated GO terms. GO: Gene Ontology.

| GO term                          | Subcategory                      | Number of transcripts | Examples of regulated genes                                      |
|---------------------------------|----------------------------------|-----------------------|------------------------------------------------------------------|
| Peptide biosynthetic process    |                                  | 105                   | gtf2h2, mrpl35_37, mrps10-12, mrps21, noa1, ssr1, rs17, rpl18    |
| Carbohydrate derivative         | Tricarboxylic acid cycle         | 22                    | aco1, aco2, cs, fh, idh1-3, mdh1, ogdh, pdhb, sdha-c, sucla2     |
| biosynthetic process            |                                  |                       |                                                                  |
|                                 | Gluconeogenesis                  | 9                     | aldoc, eno2, pck2, fbp1, pgk1, slc25a11, slc25a12, smek2, tpi1a, |
|                                 | ATP biosynthetic process         | 20                    | atp5i, atp5b, atp5g3, atp5a1, atpi, atp5f1, atp5g1, atp5o       |
|                                 |                                  | 266                   | aldh6a1, cyb5r3, cyb5r4, g6pd, gapdh, ndufs1-7, prdx1-6         |
| Oxidation-reduction process     |                                  |                       |                                                                  |
| Catabolic process               | Organic substance catabolic      | 169                   | afg3l1p, capn3, paox, pcks9, psma6, tpp1, usp14, usp2a          |
|                                 | process                          |                       |                                                                  |
|                                 | mRNA catabolic process           | 37                    | cnot11, ncbp1, ncbp2, pan2, pan3, pat1, smg5, smg8, uba52      |
| Viral process                   | Viral transcription              | 15                    | rl23a, rl37a, rpl19, rpl23, rpl38-39, rps25, rps27, rps29       |
| Cell division                   |                                  | 62                    | ccna2, ccnb1-2, ccne1-2, cdc2, cdc20, cdc34, cdc40, cdk2, fzr1  |

Finally, genes associated with AMP-activated protein kinase (AMPK) and mTOR signalling pathways, key energy-regulating pathways, were also differentially expressed between the two treatments. Specifically, mTOR signalling pathway genes (mtor, lamtor1, lamtor3, mlst8) were down-regulated while two AMPK subunit genes (prkag1-2) were found up-regulated by OA (additional file 2).

**Innate immunity and anti-viral response**

Intergenerational exposure to OA induced up-regulation of a substantial number of genes involved in innate antiviral immunity, partially listed in Table 4 and more exhaustively in table additional file 16. In particular, these belong to four classes of the germline-encoded Pattern Recognition Receptors (PRRs), which include nucleotide oligomerization and binding domain (NOD)-like receptors (NLRs), C-type lectin receptors (CLRs), toll-like receptors (TLR) and a retinoic acid inducible gene-I-like receptor (RLRs). Indeed, NLRs were the most represented PRRs with 109 transcripts, including three encoding protein nlr3-like variants for which expression levels were elevated more than 30-fold (in additional file 16 and 17, see transcript IDs: DLAgN_00244700, DLAgN_00265820 and DLAgN_00234590). Other up-regulated genes
included the melanoma differentiation-associated gene 5 (*mda5* or *ifih1*) and Interferon-Stimulated Genes (ISG), which included GTPase-Immune-Associated Nucleotide-Binding Proteins (36 genes including *gimap4*, *gimap7* and *gimap8*), interferon-induced very large GTPase 1 (*gvinp1*) and GTP binding protein Mx (*mx*). Other up-regulated ISG transcripts included interferon-induced proteins with tetratricopeptide repeats (*ifit2*), MHC class I proteins, members of the TRIM protein family (*trim25*), an IFN-induced double-stranded RNA-activated protein kinase (*pkr1, prkrir*), the grass carp haemorrhagic virus (GCHV)-induced gene -1 (*gig1*), the nuclear autoantigen sp-100 (*sp100*), the sterile alpha motif and histidine/aspartic acid domain-containing protein 1 (*samhd1*), two apolipoproteins (*apo1,3*), a damage regulated autophagy modulator protein (*dram1*) and three caspases (*casp 1, casp 6* and *casp 7*). In addition, a small number of genes within the ISG category were also significantly down-regulated in the OA condition such as *gig2* or *irig*, *trim39* and *trim8*. While some transcripts associated with the interferon JAK-STAT signalling pathway were also up-regulated (*jak2, stat5.1, stat6*), several key genes in this pathway were down-regulated, such as mitochondrial antiviral signalling (*mavs*), myeloid differentiation factor 88 (*myd88*), tnf receptor-associated factor 3 (*traf3*) and several interferon regulatory factors (*irf4, irf5, irf6, irf8*).
Table 4
Categories and number of transcripts related to genes involved in innate immunity and/or antiviral defence up and down-regulated (p-value ≤ 0.01) by ocean acidification in the olfactory rosette of F1 European Sea Bass (*Dicentrarchus labrax*) juveniles. PRR: germline-encoded Pattern Recognition Receptors; ISG: Interferon (INF)-stimulated Genes. Several transcripts may originate from the same gene. This list is given as an example, with no attempt to be exhaustive. See Additional files 15-17 for an enlarged version that include gene description and associated GO terms. Nd: Not defined.

| Category and Subcategory | Number of transcripts | Examples of regulated genes | Function |
|-------------------------|-----------------------|-----------------------------|----------|
| **UP-regulated**        |                       |                             |          |
| PRRs                    |                       |                             |          |
| NLRs: nucleotide         | 109                   | *nlrc3; nlrc5;             | pathogen recognition receptor; intracellular signal transduction (Li et al. 2018; Zhang et al. 2019) |
| oligomerization and      |                       | *nlrp1b; nlrp4e;           |          |
| binding domain (NOD)-like receptors |           | *trpm2; nod2;             |          |
|                         |                       | *nwd1; spry2;             |          |
|                         |                       | *spry3*                    |          |
| CLRs: C-type lectin      | 8                     | *illr3; cd209;             | pathogen recognition receptor/sensing RNA viruses (Yang et al. 2015) |
| receptors               |                       | *c209d*                    |          |
| TLRs: toll-like receptors| 6                     | *tlr1l; tlr2-1; tlr21;    | pathogen recognition receptor/sensing RNA viruses (Matsuo et al. 2008; Langevin et al. 2013; Quiniou et al. 2013) |
|                         |                       | *tlr22; tlr3; tlr9*       |          |
| RLR: retinoic acid       | 1                     | *ifi1*                     | pathogen recognition receptor/sensing RNA viruses (Ohtani et al. 2011) |
| inducible gene-I-like    |                       | family                     |          |
| receptor                 |                       | **ISG**                    |          |
| GTPase-related family    | 36                    | *gimap7; gimap4; gimap8*   | secretion of cytokines; anti-apoptosis of lymphocytes (Kim et al. 2018) |
| GTPase imap family       |                       |                             |          |
| very large GTPase        | 9                     | *gvinp1*                   | Host resistance (Pilla-Moffett et al. 2016; Pavlovich et al. 2020) |
| GTP binding protein Mx   | 1                     | *mx*                       | antiviral effector; inhibition of viral replication (Jensen and Robertsen 2002; Haller et al. 2007; Su et al. 2009; Wu et al. 2010) |
| IFN-induced prot. with   | 1                     | *ifit2*                    | inhibition of viral translation and replication (Fensterl et al. 2012; Fensterl and Sen 2015) |
| tetratricopeptide        |                       |                             |          |
| repeats                 |                       |                             |          |
| MHC class I              | 13                    | Nd                         | immunomodulator; antigen presentation (Koppang et al. 1999; Hansen and La Patra 2002; Robertsen 2006) |
| Category and Subcategory | Number of transcripts | Examples of regulated genes | Function |
|-------------------------|-----------------------|----------------------------|----------|
| TRIM protein family     | 30                    | trim2,9,25,32,33; bty; btr01,06; mid2 | Sensors, restriction factors of viruses, regulators of IFN response (Short and Cox 2006; Ozato et al. 2008) |
| IFN-induced, d.s. RNA-activated protein kinase | 12 | pkr1; prkrir; eif2ak2; thap4, 7; pak3,4 | inhibition of viral translation and replication (Valero et al. 2015) |
| sterile alpha motif and histidine/aspartic acid domain-containing protein 1 | 4 | samhd1 | Inhibition of viral replication (Riess et al. 2017) |
| Nuclear autoantigen sp-100 | 1 | sp100 | inhibition of viral replication (Valero et al. 2015) |
| Apolipoprotein | 4 | apo1; apol3 | Degradation of the virus (Valero et al. 2015; Levraud et al. 2019a; Collados Rodríguez 2020) |
| damage regulated autophagy modulator protein | 1 | dram1 | Degradation of the virus (Valero et al. 2015; Cadwell 2016) |
| caspases | 2 | casp1; casp7 | apoptosis effector (Valero et al. 2015) |
| ISG signalling pathway | 3 | stat5.1; stat6; Jak2 | regulation of ISG expression (Wang et al. 2017; Levraud et al. 2019a) |
| Cytokines Interleukins & interleukins receptor | 22 | il17a/f-1; il20; il34; il1r1; il20ra; il6st | inflammation, protection against infection (Ogura et al. 2006; Magnadottir et al. 2011; Secombes et al. 2011) |
| Tumour Necrosis factors related genes including C1q-TNF | 15 | c1ql2; c1ql4l; c1gtnf1; tnfsf15; cd40 | inflammation (Kishore et al. 2004) |
| DOWN-regulated | | | |
| ISG signalling pathway | 7 | mavs; myd88; traf3; irf4; irf5; irf6; irf8 | regulation of ISG expression (Wang et al. 2013; Okamura et al. 2020) |
| Ribosomal proteins and translation initiation factors | 101 | rps2, 12, 14; wdr12; rpf2; nip7; eif4; eif4ebp3l; eif4a3; eif4ea; eif4b | inflammation, viral translation (Wang et al. 2013; Okamura et al. 2020) |
Finally, a large swath of cytokine-encoding transcripts were up-regulated, including 22 interleukin-associated genes (e.g. il34; il20; il17a/f-1; il1rl2; il6st; il20ra; il18r1; il7; il4r) and 15 tumour necrosis factor-associated genes (tnip2; c1ql2; c1ql4l; c1qtnf1; tnfshf15; cd40) (Table 4, additional file 5). In contrast, almost 100 transcripts related to genes typically involved in viral protein translation during infection (e.g. rps2, 12, 14; wdr12; nip7; eif4; eif4ebp3l; eif4a3; eif4ea; eif4b) were down-regulated (Table 4, additional file 5).

**Discussion**

Because the olfactory epithelium in fish is in direct contact with water, its cells are particularly exposed to environmental changes. However, in the context of global climate change and its longer-term impact on marine life, it's difficult to predict how fish and other species will adapt to an increasingly acidic marine environment. Indeed, impacts of olfactory systems in the context of OA were reported to impact on olfactory behaviour in broadly related work (Munday 2014; Porteus et al. 2018; Jiahuan et al. 2018; Williams et al. 2019; Porteus et al. 2021). For instance, *Dicentrarchus labrax*, when exposed to predicted end-of-century OA (around 1 000 µatm) for up to two weeks, experienced declined capacity for detecting food sources and avoiding predator-associated scents (Porteus et al. 2021). However, these relatively short exposures resulted in marginal effects on gene expression of the olfactory bulb (Porteus et al. 2018; Williams et al. 2019).

In this study, we experimentally investigated intergenerational long-term consequences of OA on the olfactory epithelium transcriptome in F1 *D. labrax*. Specifically, we sought to understand how predicted end-of-century OA alters the olfactory system in a way that reflects long-term adaptations in F1 offspring, and how such changes might impact on overall robustness, such as survival after infection, a common problem in aquaculture. Cumulatively, we identified ~9,000 OA-induced differentially expressed transcripts, representing roughly a third of the total number of transcripts (34%), that revealed a profound modification of the *D. labrax*’olfactory epithelium transcriptomic profile.

Overall, among differently expressed genes, we identified genes broadly related to olfactory function. These included genes involved in ion balance, GABA signalling, olfactory neuronal signal transduction and neuron excitability, and their regulation reflected significant adaptations to synaptic plasticity (figure 4). We also observed transcriptome changes indicative of a metabolic depression in the olfactory epithelium, as exemplified by down-regulation of genes involved in ATP synthesis and the mitochondrial electron transport system [cytochrome c oxidase (cox) genes] in combination with lower expression of genes involved in energetically costly processes (e.g. macromolecules biosynthesis). At a general level, we also detected higher expression of genes related to the AMPK signalling pathway, consistent with related work in the brain of fish exposed to OA as well as in other marine species (Schunter et al. 2018; Cao et al. 2018). Extending from these findings, and unexpectedly, we also discovered that *D. labrax*, upon intergenerational exposure to OA, up-regulated gene programs conferring elevated resistance to virus infection. Consistent with these molecular changes, we found in our betanodavirus infection study that
the OA-exposed F1 cohort succumbed to infection later and that fewer fish died than control F1 fish (figure 4).

Across different organs in OA-exposed and control fish, virus presence and concentrations were similar, indicating that the viral infection in and of itself was similar, and that the difference in disease resistance was due to a superior defence relative to fish not exposed to OA. In related work, Bresolin de Souza and collaborators found that a three-month exposure to OA in Atlantic juvenile halibuts was associated with increased activity of complement component C3, lysozyme and fibrinogen, known factors in innate and complement systems though antiviral and inflammatory activities (Bresolin De Souza et al. 2014; Bresolin de Souza et al. 2016). Apart from this work, OA effects linked to immune function has been reported for shellfish (e.g. Leite Figueiredo et al., 2016; Castillo et al., 2017; Sun et al., 2017), but our insight into long-term adaptations for marine fish, and their potential ramifications for aquaculture remain largely unknown (Baag and Mandal 2022). As a progressively acidic marine environment will inevitably become a reality in the future, such knowledge is urgently needed. Here, we reveal for the first time that fish may acquire an improved capacity to resist a viral infection upon years of intergenerational exposure to OA. In our transcriptomic characterization of the olfactory epithelium of fish exposed to intergenerational OA, we detected up-regulation of gene programs linked to innate antiviral immunity; pathogen receptors genes, Interferon-Stimulated Genes (ISG) and ribosome-related genes.

Detection of an invading pathogen is a critical first step in the initiation of a robust immune response (uncapped viral RNA/DNA and/or cell wall components) through Pattern recognition receptors (PRRs) (Janeway and Medzhitov 2002). In the OA-exposed F1 cohort, we detected a robust up-regulation of four out of the five types of PRRs relative to control F1 fish, suggesting an enriched extent of pathogen detection as a first contingency response upon viral invasion. Of note, these are germline-encoded genes, suggesting that their up-regulation can partly result from trade-offs in parents carried over to offspring, although our analysis does not allow us to distinguish this possibility from solely F1 generation adaptations. Of the up-regulated PRR classes, nucleotide oligomerization and binding domain (NOD)-like receptors (NLRs) NLRs were the most abundant. NLRs are barely present in mammals, but through evolution expanded into very large families in fish and other species lacking adaptive immunity (Howe et al. 2016). NLR can act as sterile and intracellular pathogen stress signal receptors and are therefore poised to play a central role in the inflammatory response (Laing et al. 2008; Levraud et al. 2019a) and homeostasis of microglia, the only central nervous system immune cells present in fish (Shiau et al. 2013; Wang et al. 2019).

The second PRR class, C-type lectin receptors (CLRs), mediate bacteria- and fungi-associated responses, but following viral recognition, can induce both protective and detrimental effects depending on the pathogen (Hoving et al. 2014). Members of the mannose receptor, the asialoglycoprotein receptor and c-type lectin domain families, were represented among the up-regulated CLR genes (additional file 2). Interestingly, the melanoma differentiation-associated protein 5 (mda5 or ifih1) gene, which encodes for a member of RLRs and several members of the TLRs gene family, both considered as key virus sensors
were also up-regulated (Takeuchi and Akira 2009; Ranjan et al. 2009). Stimulation of MDA5 or TLRs by RNA viruses is known to result in type I interferon response activation (Langevin et al. 2013).

Interferons are signalling cytokines secreted by every type of cell in order to interfere with viral progression by regulating the expression of more than 1000 genes at the transcriptional level (Klamp et al. 2003; Paul et al. 2007; Li et al. 2009; Kuenzel et al. 2010). These genes are called Interferon-induced genes (ISG) and exhibit a wide array of antiviral properties (de Veer et al. 2001; Wang et al. 2017). A large proportion of orthologous ISGs have been identified in fish and humans and very few fish ISG have no human ortholog (e.g. gig1, gig2, vig-B319 (Levraud et al. 2019b)). The mechanisms of antiviral activity of fish-specific ISGs remain poorly understood (Schneider et al. 2014; Poynter and DeWitte-Orr 2016). Here, we found that OA-exposed fish expressed several up-regulated ISGs whose functions may partially explain their enhanced resistance to viral infection. These include the IFN-induced double-stranded RNA-activated protein kinases (pkr1, prkrir), a particular ISG subset that functions as PRR, antiviral effector, and inhibitors of virus translation and replication combined (Clemens and Elia 1997; Schneider et al. 2014; Poynter and DeWitte-Orr 2016); genes from the GTPase imap family (gimap8, 4, 7), known to mediate cell-autonomous resistance against pathogens (Haller and Kochs 2002; Klamp et al. 2003; Haller et al. 2007); and genes from the TRIM family (trim16, 39) that play pivotal role in viral restriction, modulation of immune signalling, autophagy and formation of cellular structures (Langevin et al. 2019). The regulation of ISG transcription can take place via the classical interferon mediated JAK-STAT signalling pathway (Darnell et al. 1994; Wang et al. 2017) and/or through a variety of non-canonical pathways independent of interferon induction (Mostafavi et al. 2016; Wang et al. 2017). Here, we found that genes involved in the JAK-STAT pathway were both up- and down- regulated in the OA-exposed fish cohort. To better understand ISG signalling pathway regulation in the context of viral infection in fish, analyses at the post transcriptional levels (e.g. phosphorylation status) for JAK-STAT and non-canonical pathways would be necessary.

Finally, we found that intergenerational exposure to OA triggered a down-regulation of genes coding for ribosome biogenesis, ribosomal proteins and eukaryotic translation initiation factors, indicative of reduced ribosome abundance and malfunction. This is confirmed by the GO analyses that show strong down-regulation of both translation and macromolecule biosynthetic process. During an infection, the virus hijacks the host ribosomes to produce new viral particles along with host cellular factors to initiate viral translation (Lee et al. 2013; Li 2019). We hypothesised that as OA induced a malfunction of the host's ribosomes, viruses may lack the biosynthetic machinery to translate and transcribe their nucleic acids. This, together with the up-regulation of PRRs and ISGs might explain the observed superior resistance during the viral challenge.

VNN outbreaks are considered one of the most relevant infectious constraint for the culture of a variety of fish species. Hence, the new insights we provide into how a marine teleost of economic interest undergoes intergenerational acclimation could be of precious concern to aquaculture. Although juvenile fish exposed to intergenerational acclimation were more resistant to VNN, based on the changed transcriptome, their metabolic and odour transduction programs were altered, which may in turn alter the
olfactory perception of a wide array of chemical cues that may consequently impact odour-mediated behaviours including feeding, homing and other inter- or intra-specific interactions such as sociability, mating, competition and predator detection (figure 4). Understanding how the interplay between acidification and warming over generations modulate olfactory behaviour and viral resistance could be useful to develop new strategies for maintaining the health and production in aquaculture facilities for either commercial or scientific purposes.

Conclusions

We report here that intergenerational exposure to OA induces a deep modification of the transcriptomic profile in the olfactory epithelium of the *D. labrax* that include plastic responses related to ion balance and transport, neuronal activity and plasticity, energy metabolism and innate immunity (figure 4). This intergenerational plasticity may be considered as an acclimation (adaptive plasticity) to prevent more severe OA-induced physiological disruption at the whole organism level (Bailey et al. 2017; Schunter et al. 2018). It is noteworthy that immune system factors such as cytokines, interferons and interleukins also play a role in central nervous system and brain development, and can induce changes in neural network activity, supporting the intricately interplay between the immune and the nervous system (Mousa and Bakheit 2013; Clarkson et al. 2017). Additional experimentations based on electrophysiology and behavioural tests would help determine whether the regulation that we observed in the neuronal plasticity and activity gene programs are associated with a perturbation of the olfactory function, as suggested by the regulation of key processes associated with energy metabolism. Likewise, further studies will be necessary to more globally characterize OA-induced effects on immune status in both cultured and wild fish and its capacity to resist the most fish pathogens in a changing ocean.

Methods

Animal experiments in this study were conducted following the European Commission recommendation 2007/526/EC and Directive 2010/63/EU for the accommodation and care of animals used for experimental and other scientific purposes. The OA conditioning was performed at the French Research Institute for Exploitation of the Sea (IFREMER) in Plouzané, within the facilities of the Laboratory of Adaptation, Reproduction and Nutrition of fishes (ARN) (Agreement number: B29-212-05). The viral challenge was performed at the French Agency for Food, Environmental and Occupational Health & Safety (ANSES) in Plouzané (Agreement number: D29-212-3). Each procedure was the subject of a specific authorization issued by a French Ethics Committee for animal testing (APAFIS #2018032209421223 for OA conditioning, and APAFIS #202001161613768 for viral challenge).

Animal husbandry and experimental setup

F1 juveniles (2 years old) of the European sea bass *Dicentrarchus labrax* used in the present study originated from an *in vitro* fertilisation of 4 years old F0 parents exposed from larval to adult stages to control conditions (~pH8.0) or to OA conditions (~pH7.6) (Crespel et al. 2017; Cominassi et al. 2019;
Howald et al. 2019; Cominassi et al. 2020; Mazurais et al. 2020). Sperm and eggs were collected and pooled from 20 males and 6 females of each pH-treatment (see methodological summary in figure 1). Luteinizing hormone releasing hormone (LHRH) was injected to stimulate synchrony in oocytes full maturation. Taking great care of maintaining parental pH conditions, eggs were hatched and the resultant F1 offspring were reared in water at the same pH as their parents. Rearing conditions during larval and juvenile stages were similar to those described in previous studies (Howald et al. 2019; Cominassi et al. 2020; Mazurais et al. 2020). For both treatments, seawater temperature and salinity followed seasonality of the Bay of Brest. 402 juveniles (201 per treatment) were distributed evenly in 6 culture tanks (400 L, three tanks = three replicas per treatment) that were part of an open-circuit system. To guarantee high quality, seawater pumped 500 m off the coastline at a depth of 20 m passed through a sand filter, a tungsten heater, a degassing column packed with plastic rings, a 2-µm filter membrane, and a UV lamp. Seawater for the control treatment was then poured into each of the three replicas tanks. Seawater for the OA treatment was injected with CO$_2$ (through manipulation of a flowmeter (Aalborg, USA) connected to a CO$_2$ bottle (Air Liquide, France)) in a header tank equipped with a degassing CO$_2$ column to favour mixing. Then, low pH seawater was poured into each of the 3 replicas tanks. pH in NIST scale and temperature in the six tanks were daily measured with a WTW 3110 pH meter (Xylem Analytics Germany, Weilheim, Germany; with electrode: WTW Sentix 41) calibrated daily with pH4.0 and pH7.0 buffers (WTW, Germany). Total alkalinity was measured once a week following the protocol of Strickland and Parsons (Strickland and Parsons 1972): a 50 ml sample of tank water was mixed with 15 ml HCl (0.01 M) and pH was measured immediately. Total alkalinity was then calculated with the following formula:

$$TA = \frac{V_{HCl} \times C_{HCl}}{V_{sample}} - \left( \frac{V_{HCl} + V_{sample}}{V_{sample}} \right) \times \frac{H^+}{H^+}, \left[ \text{mol l}^{-1} \right]$$

with, total alkalinity (TA, mol l$^{-1}$), volume (V, l) of HCl or of the sample, concentration (C, mol l$^{-1}$) of HCl, hydrogen activity (H$^+$, 10$^{-pH}$) and hydrogen activity coefficient (YH$^+$, here = 0.758). F1 juveniles were fed ad libitum with a diet that meets their nutritional requirements (Vitalis Cal, Skretting, Stavanger, Norway). No significant difference was observed in the mean weights (t test, t=0.02, df=64.52, p-value = 0.98) of the 2-year-old juveniles between the two treatments (n=35). F1 juveniles from the two treatments behaved and fed in a relatively similar way. Mortality events were not detected.

**RNA extraction**

Prior to sampling, fish were fasted for 24 h. Then, 2-year-old juveniles were first anesthetized (20 mg L$^{-1}$), and then euthanized with a lethal dose (200 mg L$^{-1}$) of tricaine methane sulfonate 222 (MS222, Pharmaq, Fordingbridge, Hampshire, UK). Sampling consisted in dissecting the olfactory rosette from 7 individuals per treatment that were quickly stored in RNA Stabilization Reagent (RNAlater, Qiagen, Hilden, Germany) following recommendations from the supplier. Total RNA was extracted using Extract-All reagent
(Eurobio, Courtaboeuf, Essonne, France) combined with Nucleospin RNA column that includes one step of DNase treatment (Macherey–Nagel, Düren, Germany) according to the manufacturer’s instructions. The concentration and purity of extracted RNA were verified (ratio>2) using an ND-1000 NanoDrop® spectrophotometer (Thermo Scientific Inc., Waltham, MA, USA). The integrity of RNA was checked by electrophoresis using an Agilent Bioanalyzer 2100 (Agilent Technologies Inc., Santa Clara, CA, USA). All samples showed an RNA integrity (RIN) score >9. RNA samples were stored at −80°C for further RNA sequencing and qPCR analyses.

**RNA-Sequencing analysis**

RNA samples extracted from the olfactory rosettes were sent to the sequencing platform GenomiX (MGX, Montpellier, France) for transcriptome analysis through RNA sequencing (RNA-Seq).

**cDNA Libraries:** Libraries of cDNA were constructed using TruSeq Stranded mRNA kit (Illumina, San Diego, CA, USA) according to the manufacturer’s instructions. Briefly, poly-A RNAs were purified using oligo-d(T) magnetic beads from 863ng of total RNA. Poly-A RNAs were fragmented and underwent a reverse transcription using random hexamers. During the second strand generation step, dUTP substituted dTTP in order to prevent the second strand to be used as a matrix during the final PCR amplification. Double stranded cDNAs were adenyalted at their 3’ ends and ligated to Illumina's indexes. Ligated cDNAs were amplified following 15 PCR cycles. PCR products were purified using AMPure XP Beads (Beckman Coulter Genomics, Brea, CA, USA). Libraries were validated using Standard Sensitivity NGS kit on Fragment Analyzer (Agilent Technologies, Santa Clara, CA, USA).

**Libraries sequencing:** Fourteen libraries were pooled in equimolar amounts. The balance between all samples of the pool was assessed by using KAPA Library quantification kit (Roche, Bâle, CHE). The pool was then sequenced on a Novaseq 6000 (Illumina, San Diego, CA, USA) on a SP flow cell in single-read 100 nt mode according to the manufacturer’s instructions. This sequencing produced between 40 and 50 million passed filter clusters per library.

**Sequencing quality control:** Image analyses and base calling were performed using the NovaSeq Control Software and Real-Time Analysis component (Illumina, San Diego, CA, USA). Demultiplexing and trimming were performed using Illumina’s conversion software (bcl2fastq 2.20). The quality of the raw data was assessed using FastQC from the Babraham Institute and the Illumina software SAV (Sequencing Analysis Viewer). FastqScreen was used to estimate the potential level of contamination.

**Transcriptomic and Gene Ontology (GO) analysis**

Raw reads mapping and gene quantification were done using STAR aligner (v2.7.2c) (Dobin et al. 2013) to the *D. labrax* reference genome guided by the reference gene annotation (Tine et al. 2014). Differential expression analysis was performed with the DESeq2 package (Bioconductor) (Love et al. 2014) using an adjusted p-value cut-off of 0.01. The SRR accession numbers for the raw sequence data are SRR15222852-65. *Cbln11* structural annotation was carried out manually in GenomeView (Abeel et al. 2012) based on BlastHit of *Cbln11* sequences available at NCBI and from mapped RNA-Seq reads. Gene
structure was exported in *gtf* format and added to the current annotation. The datasets supporting the conclusions of this article are available in the SRA repository under accession numbers: SRR15222859-65 (https://www.ncbi.nlm.nih.gov/sra/?term=SRR15222852) and SRR15222852-58 (https://www.ncbi.nlm.nih.gov/sra/?term=SRR15222859).

To extract biological meaningfulness and to visualize potentially affected pathways, a Gene ontology (GO) enrichment analyses was performed separately on significantly up- or down-regulated genes under OA. Raw reads from RNA-Seq were imported into Galaxy instance of Ifremer (Augusto et al. 2017). An obo GO file and the product annotation file for *D. labrax* (Tine et al. 2014) were used to analyse significantly (p<0.01) up or down-regulated genes. FDR was corrected with the Benjamini-Hochberg test. The corrected p-value to apply to the graph output was set to 0.01.

**qPCR validation**

To validate some results from the transcriptomic analysis, qPCRs were performed. iScript™ cDNA Synthesis kit (Bio-Rad Laboratories Inc., Hercules, CA, USA) was used to reverse transcribe in duplicate 500 ng of DNase-treated RNA samples into first strand cDNA by following conditions provided by the manufacturer. Negative RT consisting in RT reaction without retro-transcriptase enzyme were also performed for all samples. The relative quantification of mRNA exhibiting differential expression through the RNA-Seq approach [SAM and HD domain containing deoxynucleoside triphosphate triphosphohydrolase 1 (*samhd1*), Interferon-induced very large GTPase 1 (*gvinp1*), Toll-Like Receptor 3 (*tlr3*), Chemokine C-X-C motif ligand 14 (*cxcl14*), Nuclear factor of activated T-cells 5 (*nfat5*)] was performed by qPCR using primers designed for the assembled transcripts and blasted using the NCBI BLAST tool to verify specificity (additional file 1). Elongation factor 1-alpha (ef1α) and Ribosomal Protein L13a (rpl13a) were tested as housekeeping genes for normalization but only rpl13a was used since ef1α did not meet the acceptance criteria in terms of gene expression stability measure (M) and coefficient of variation (CV).

The efficiency of the qPCR reaction tested for each primer pair through standard curves were around 100% with R² >0.999. Transcript expression was quantified using the CFX96 Touch Real-Time PCR Detection system (Bio-Rad Laboratories Inc.) and the protocol previously described (Mazurais et al. 2020).

**Viral Challenge experiment**

Betanodavirus strain W80 isolated from diseased *D. labrax* displaying typical signs of VNN was used in this study. According to Castri et al. (Castri et al. 2001), a stock of virus was produced at 24°C on the SSN-1 (Striped Snakehead fish; *Ophicephalus striatus*) cell line (L15 medium, 10% FBS, pH7.6) and frozen. Cell debris was removed by centrifugation for 15 min at 2000 g; the virus was then aliquoted and stored at −80°C. Before the challenge, viral titration was carried out on one of the aliquots after a single freeze-thaw cycle based on the tissue culture infectious dose technique (TCID₅₀) described by (Dussauze...
The infectious titer of the viral production was calculated according to the method of Kärber (Kärber 1931), and was found to be $1 \times 10^8 \text{TCID}_{50}/\text{mL}$.

After 2-years rearing of the F1, 148 juveniles per treatment were divided into four flow through tanks of 400L (figure 1) and were challenged by immersion in a bath with W80 at 25°C to mimic the environmental conditions where the clinical signs of disease were reported. For the viral challenge, the water flow was interrupted, the oxygenation was increased and the water volume was reduced to 100 L. Three out of four tanks per treatment were exposed for 3 h with an infectious dose of $5 \times 10^4 \text{TCID}_{50}/\text{mL}^{-1}$ of W80. The fourth tank per treatment was exposed in the same conditions to SSN-1 cell supernatant free from the virus and used as a negative control without virus. After a 3-hours bath exposure, the water flow was restored to slowly dilute the virus titration. The system was maintained open and the water temperature ($25^\circ C \pm 2^\circ C$) was continually measured and recorded with a wireless probe (Cobalt, Oceasoft®) coupled to an acquisition system (ThermoClient 4.1.0.24). Juveniles were fed once a day with commercial pellets (Neo Start Coul 2 from Le Gouessant Aquaculture) except on the day of the viral infection.

Mortality was recorded twice a day during 45 days post-infection (dpi). Dead fish were stored at −20°C until viral examination. The presence and concentration of the virus was checked on a pool of organs (spleen, kidney, heart, brain). Virus concentrations were determined in six fish that died at the peak of mortality during the viral challenge (9 dpi in the control treatment; 11 dpi in OA treatment) and in five survivor fish per treatment by immunofluorescence assay on the SSN-1 and E11 cell lines according to the adapted protocol by Dussauze et al. (Dussauze et al. 2015).

Survival curves were estimated using the Kaplan–Meier method and were compared by a log rank analysis using the online platform BiostaTGV. Firstly, survival curves for each replica within each treatment were compared, and if no significant difference was observed, results from the 3 replicas per treatment were pooled. In case of significant difference, each replica was analysed separately and compared to the other treatment. In addition, a Kruskal–Wallis test together with a Dunn's post hoc test were performed (XLSTAT software 2020) to determine whether the Mean Day to Death (MDD) and the final survival rate varied with treatment.

**Declarations**

_Ethics approval and consent to participate:_ As indicated in the Methods section, our two experiments followed European Commission recommendations on fish ethics and well-being and were authorized by the French Ethics Committee for animal testing (APAFIS #2018032209421223 and APAFIS #202001161613768).

_Consent for publication:_ Not applicable.

_Availability of data and materials:_ The datasets generated and analysed in this article are available in the SRA repository under accession numbers: SRR15222859-65 and SRR15222852-58.
(https://www.ncbi.nlm.nih.gov/sra/?term=SRR15222852 and https://www.ncbi.nlm.nih.gov/sra/?term=SRR15222859).

**Competing interests:** The authors declare that they have no competing interests.

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**Authors’ contributions:** DM & FM conceived the scientific questions and experimental design. DM collected RNA samples and TV extracted the RNA and verified RNA quality and integrity. A-AG performed the RNA sequencing. MC-R, AC & CN performed the transcriptomic and Gene Ontology analysis. LM performed qPCR’s. MD & JC conducted the viral challenge experiment and treated and analysed the outcoming data. MB provided ideas concerning olfactory behaviour. MC-R & DM led the writing of the manuscript. All authors contributed significantly to the revision of the manuscript and approved it for publication.

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Figures

Figure 1

Methodological summary. It is shown the rearing times and procedures applied on *Dicentrarchus labrax* parental lineage (F0) and their offspring (F1) exposed to either the control (pH 8.0) or the acidified (pH 7.6) treatments; r: replica; nc: negative control.
Survival rate (%) of F1 *Dicentrarchus labrax* after viral challenge. Curves indicate the survival rate of non-infected juveniles (white circle) and infected juveniles (black circle) with W80 strain either from the control treatment (pH 8.0, continuous line) or the acidified (pH 7.6, broken line) treatment. Each challenge was performed in triplicate with 37 fish per replica tank. Survival was monitored during 45 days post-infection.

**Figure 3**

Gene ontology enrichments for biological processes. Enriched (q-value ≤ 0.01) biological processes related to the (A) up-regulated and (B) the down-regulated genes in F1 *Dicentrarchus labrax* exposed to ocean acidification.

**Figure 4**

Conceptual framework. The impacts of intergenerational ocean acidification (OA) on F1 *Dicentrarchus labrax* juveniles are shown for both the viral challenge and the ARN-Seq analysis.

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