



# Diluted Seawater and Ammonia-N Tolerance of Two Mangrove Crab Species. New Insights to Understand the Vulnerability of Pristine Islands Ecosystems Organisms

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Mégevand L, Theuerkauff D, L'Épine C, Hermet S, Corse E, L'Honoré T, Lignot J-H and Sucré E (2022) Diluted Seawater and Ammonia-N Tolerance of Two Mangrove Crab Species. New Insights to Understand the Vulnerability of Pristine Islands Ecosystems Organisms. Front. Ecol. Evol. 10:839160. doi: 10.3389/fevo.2022.839160 Mangrove ecosystems are the primary receptors of anthropogenic pollution in tropical areas. Assessing the vulnerability of these ecosystems can be expressed, among other indicators, by studying the health of 'ecosystem engineers'. In this study, mangrove forests facing opposing anthropogenic pressures were studied (i) in the uninhabited island of Europa (Mozambique Channel), considered as a pristine ecosystem, and, (ii) on the island of Mayotte, facing regular domestic wastewater discharges. Using an ecophysiological approach, the effects of diluted seawater (DSW) and increased ammonia-N were studied for two fiddler crab species: Gelasimus tetragonon (GT) on the island of Europa and Paraleptuca chlorophthalmus (PC) on the island of Mayotte. Osmoregulation curves and osmoregulatory capacity were determined along with O2 consumption rates after a 96 h exposure period. Histological analyses were also carried out on two important metabolic organs: the hepatopancreas and the posterior gills. Results indicate that both crab species are good hyper-hypo-osmoregulators but only PC can maintain its osmoregulatory capacity when exposed to ammonia-N. Oxygen consumption is increased in GT after 96 h of exposure to ammonia-N but this does not occur in PC. Finally, a thickening of the gill osmoregulatory epithelium was observed after 96 h in PC when exposed to ammonium but not in GT. Therefore, the two species do not have the same tolerance to DSW and increased ammonia-N. PC shows physiological acclimation capacities in order to better manage nitrogenous enrichments. GT did not show the same physiological plasticity when exposed to ammonia-N and could be more at risk by this kind of stress. These results along with those from other studies regarding the effects of domestic effluents on mangrove crabs are discussed. Therefore, the greater vulnerability of organisms occupying pristine ecosystems could induce major changes in mangrove functioning if crabs, that are engineer species of the ecosystem, are about to reduce their bioturbation activity or, even, disappear from the mangrove forests.

Keywords: Crustacea, pristine ecosystems, biomarkers, metabolism, ammonia-N, mangrove

#### Mangrove Crabs in Changing Environments

# INTRODUCTION

Ecological and ecosystem vulnerability assessment may represent a valuable tool in biodiversity risk management (De Lange et al., 2010). From the definition of Ippolito et al. (2010), ecosystem vulnerability may represent "the potential of an ecosystem to modulate its response to stressors over time and space, where that potential is determined by characteristics of an ecosystem vulnerability that include many levels of organization." Ecosystem vulnerability assessment may be expressed with a level of potential impact related to a certain stressor in a given environment: "the actual status of a polluted ecosystem or community represents the response of a (more or less) pristine ecosystem or community to a specific stressor or to multiple stressors" (De Lange et al., 2010). The authors, thus, state that combining the assessment of vulnerability of a pristine ecosystem or community with the actual status allows obtaining crucial information for risk management.

Among vulnerable ecosystems, mangrove forests that thrive intertidal areas are typically subjected to a variety of stressors, both natural and anthropogenic. These include high levels of pollutants and nutrients, severe hypoxia, high water turbidity, and fluctuations in temperature and salinity. Mangroves are significant ecosystems to study given their ecological importance in tropical areas. They accomplish a wide range of key ecosystem functions such as carbon storage, nutrient and global carbon cycling, provide breeding, recruitment, nursery and feeding areas both terrestrial and aquatic fauna (Nagelkerken et al., 2008; Amaral et al., 2009; Cannicci et al., 2009; Lugendo and Kimirei, 2021). Also, mangrove forests form a buffer zone protecting the shoreline against erosion and watershed (Zhang et al., 2012; Ouyang and Guo, 2016).

Located at the land-sea interface, mangroves are the receptacle of anthropogenic activities such as aquaculture, agriculture, urbanization, tourism or coastal pollution (Thomas et al., 1996; Alongi, 2002; Xiang et al., 2020; Lugendo and Kimirei, 2021).

Nitrogen inputs and the altered global cycle of N are one of the most important concerns about anthropogenic presence and activities (Vitousek et al., 1997). Human alterations of the nitrogen cycle can lead, amongst others, to changes in the composition and functioning of estuarine and nearshore systems, with accelerated losses of biological diversity (Nixon, 1995; Nixon et al., 1996; Vitousek et al., 1997). It has been predicted that tropical regions will receive the most dramatic increases in reactive N (Nr) inputs over the next few decades (Zhu et al., 2005; Galloway et al., 2008). In this context, our study aims at comparing some features of ecosystem vulnerability in both pristine and anthropized mangrove ecosystems.

Europa island  $(22^{\circ} 20' \text{ S}, 40^{\circ} 22' \text{ E})$  is the southernmost island of the Mozambique Channel. It is considered a pristine ecosystem as there is no permanent human settlement, making it a prime candidate for marine conservation (O'Donnell et al., 2017). Europa island is a low-lying (7 m elevation), coral island of 28 km<sup>2</sup> surrounded by sand dunes. A shallow lagoon, almost empty at low tide, is located on the north-western part of the island. It is fringed with 626 ha of mangrove forest (Boullet, 2014; Juhel et al., 2019).

Mayotte island (12°50′S 45°08′E) is a tropical island (374 km<sup>2</sup>) situated in the Comoros Archipelago, in the northern part of the Mozambique Channel. On this island, the mangrove forest covers about 700 ha of the territory (Jeanson et al., 2019). Large ecological gradients are encountered in Mayotte rivers and mangroves, one of the main anthropogenic pressures being the lack of connection between households and the sewage systems. Therefore, domestic wastewater is often discharged directly into rivers and mangroves (Herteman, 2010; Vasselon et al., 2017; Capdeville, 2018). Furthermore, the island of Mayotte possesses a fast growth of its human population, especially over the last two decades resulting in an exponential increase in anthropogenic pressures and degradations of its marine environment (Pusineri et al., 2014).

Studying ecosystem vulnerability in response to stressors must take into consideration indicators of ecosystem functioning, including early warning, diagnostic and retrospective indicators. Aquatic organisms including crustacean species, are frequently studied as early warning indicators of changes in environmental conditions (Burger, 2006; Ungherese and Ugolini, 2009). Some ecological functions provided by these species, such as ecosystem engineering, have also been identified (Brodie et al., 2018) as key elements to incorporate into conservation planning (see Coggan et al., 2018 for a review). Ecosystem engineers are species whose actions have "significant impacts" upon the physical structure of their habitat and the organisms that live in them (Jones et al., 1994). Amongst them, we find burrowing mangrove fiddler crabs, considered as ecosystem engineers through their bioturbation activity (Kristensen, 2008; Penha-Lopes et al., 2009; Kristensen et al., 2012). Their activity briefly consists in burying, macerating and ingesting litter as well as actively digging and maintaining burrows in the sediment (Giddins et al., 1986; Emmerson and McGwynne, 1992; Dittmann, 1996; Lee, 1997; Kristensen and Alongi, 2006).

We chose to study two ecologically, morphologically and phylogenetically proximate species of Ocypodids fiddler crabs: Paraleptuca chlorophthalmus (PC) from the island of Mayotte and *Gelasimus tetragonon* (GT) from the island of Europa (Figure 1). As fiddler crabs, these two species are mainly characterized by a sexual dimorphism where female possess two small feeding claws when males possess one enlarged claw (up to 50% of total bodyweight). PC distribution range is restricted to the East African province of the Indian Ocean Subrealm of which it is endemic, whereas GT is the fiddler crab with the largest distribution range. It is widespread in the Indian Ocean and also in the Pacific Ocean (Figure 1) (Rosenberg, 2020). Only one observation of GT has been reported in Mayotte Island (Bouchard et al., 2013). PC has also been observed once in Europa Island (Poupin et al., 2012). We did not confirm this observation and the DNA sequencing analysis performed on the crabs collected on Europa Island did not reveal any representative of this species (see Supplementary Material). These two species are the most abundant crab species living in semi-open canopy habitats situated near tidal rivulets of the two studied mangroves. In these two areas, a high density of burrows (that originates from bioturbation activity) is observed. These habitats are the first receptors of anthropogenic spillages in the island of Mayotte.

Salinity variations throughout freshwater inputs and high levels of nutrients are two key stressors in such context. On the island of Europa, rainfall is the only source of freshwater, with a mean annual rainfall of 540 mm/year and a long dry period of 7 months in a semi-arid climate (Boullet, 2014; Lambs et al., 2016). Therefore, GT lives under strong marine influence, mostly in seawater and oligotrophic conditions. On the island of Mayotte, 24 permanent rivers reaching the sea have been listed (Lapègue, 1999), with a watershed area ranging from 2 to 23 km<sup>2</sup> (Lagarde et al., 2021). Therefore, PC is frequently experiencing salinity variations and osmotic changes under mesotrophic or eutrophic conditions, especially in anthropized areas.

In habitats with frequently changing environmental conditions (such as mangrove forests), organisms are exposed to multiple stressors that can neutrally, synergistically or antagonistically interact (Piggott et al., 2015; Renault et al., 2018). In order to maintain functional homeostasis, phenotypic adjustments may be expressed during the life history of an individual, ranging from physiological adjustments (acclimation) (Teets and Denlinger, 2013) to phenological changes and range shifts (Chuine, 2010; Briscoe et al., 2012; Renault et al., 2018). In some cases, these exogenous factors can generate physiological evolutionary changes because of genetic modifications (adaptation). This study does not aim to determine any potential environmental adaptation in the event that different sensitivities are observed between crab populations, and this should remain a working hypothesis only.

Considering these elements, we chose to study physiological endpoints on different levels of biological organization to assess the potential effects of salinity variations and short-term exposures to ammonia-N on the two crab species. Physiological biomarkers such as osmoregulatory capacity (the difference between hemolymph osmotic pressure and that of the external medium, OC), oxygen consumption and histological changes have been widely studied and proved to be informative proxies of crustacean responses to environmental changes such as salinity variations and ammonia-N inputs (Lignot et al., 2000; Gillikin et al., 2003; Ortega et al., 2017; Theuerkauff et al., 2018a,b; Ros et al., 2021). Resting metabolic rate is also considered as physiologically relevant (Ikeda, 2016; Borges et al., 2018) to assess the potential effects of an osmotic chock or ammonia-N exposure on crabs, independently from a potential locomotor or escape behavior.

Therefore, could GT and PC show different physiological tolerances to basic anthropogenic pressures, here: salinity reduction and high ammonia-N inputs? The working hypothesis is that vulnerability to diluted seawater and ammonia-N inputs must be different for the two species. Specifically: (1) PC could have a good tolerance capacity to reduced salinity and increased ammonium input within a short-term acclimation period; (2) GT, that is mainly recorded in stable environments, i.e., rarely exposed to eutrophic water and rarely to diluted seawater (Crane, 1975; Weis and Weis, 2004), could be vulnerable to anthropogenic discharges inducing salinity decrease; (3) there should be deleterious effects of Ammonia-N on osmoregulatory processes (aggravating factor) for both species since ammonia-N is known to disrupt physiological processes

such as osmoregulation, immunology, acid/base balance and gas exchange in Decapod crustaceans (Weihrauch et al., 2004; Romano and Zeng, 2013).

In the present study, crabs were collected from Europa (GT) and Mayotte (PC) mangroves and were artificially exposed to diluted seawater (5 ppt) and Ammonia-N (10 mg.L<sup>-1</sup>), a proxy of anthropogenic presence through an environmental concentration. Several biomarkers were considered in order to: (i) define the salinity tolerance range of the two species (osmolality curves), (ii) understand whether eutrophication can have an impact on the crab ability to maintain its hydromineral balance (osmoregulatory capacity), and, (iii) determine whether short-term acclimation (oxygen consumption and histological analyses) can occur to cope with eutrophication. If GT was found to be vulnerable to freshwater and Ammonia-N, what would be the consequences in terms of vulnerability to anthropogenic pressure for the pristine mangroves of the island of Europa? What could the consequences be in terms of risk assessment?

# MATERIALS AND METHODS

#### Study Sites Europa Island

The sampling site is mainly constituted of shrubby *Rhizophora mucronata* on coral mud with a sandy-silty texture, in an almost open canopy (**Figure 2**), near tidal channels and rivulets. The absence of freshwater inflows induces no transfer of nutrient or ions from inland to the mangrove (Lambs et al., 2016). High concentrations of nitrogen and phosphorus can be found in this Island on seabirds nesting sites due to amounts of guano, but the chosen sites in this study were not concerned (Zubia et al., 2016).

#### Mayotte Island

The study site is located in Boueni bay  $(12^{\circ}55'1''S, 45^{\circ}9'23''E)$ in the mangrove of Malamani, extending along both sides of a freshwater stream. On this site, vegetation structure is mainly represented by *Ceriops tagal* and *R. mucronata facies*, along with *A. marina* and *Bruguiera gymnorhiza* species. This open habitat presents no canopy cover, a large temperature gradient and regular water immersion/emersion, depending on the season, tides and time of the day. Malamani is part of the Chirongui area in the bay of Boueni, which had a human population density of 310 hab./km<sup>2</sup> in 2017.

A total of 30 crabs (15 from Europa and 15 from Mayotte) used during experiments were randomly chosen for DNA sequencing to confirm their taxonomical identification through a classical barcoding process (see **Supplementary Material**).

# **Animal Collection and Acclimation**

Male and female GT and PC (the same number for each sex) were collected in April 2019 and March 2020, respectively, in the mangroves of Europa and Mayotte (Malamani area) (see **Figure 2** for the sampling site). All animals were hand collected at low tide when crabs are active and out of their burrows. They were placed into individual bags or boxes in order to minimize stress and fights during transport to the laboratory



(either the field laboratory on Europa Island, or the CUFR marine laboratory at the University Center of Mayotte). Animals (~130 for PC, ~100 for GT) were placed for 3 days for acclimation in individual boxes containing natural filtered seawater (~33‰ salinity; 1,050 mOsm.kg<sup>-1</sup>), under a natural photoperiod (12 h light: 12 h dark) and without feeding prior to experimentation.

For the osmoregulation experiment,  $\sim 100$  PC were studied (N = 10 per salinity) in Mayotte and  $\sim 70$  GT were studied (N = 4-8 per salinity) in Europa Island. Sampling size was limited here due to the pristine nature of the ecosystem and legal environmental protection requirements. For the respirometry and osmoregulatory capacity, 24 crabs were studied for each species. No mortality was observed during acclimation and experimental processes for PC (Mayotte). No mortality was observed during acclimation for GT (Europa), but one female died after 72 h of ammonia-N exposure.

# Ammonia-N Exposure and Experimental Conditions

Three different experimental conditions were performed: some crabs were maintained in seawater ( $\sim$ 33 ppt), others were transferred to diluted seawater ( $\sim$ 5 ppt) or to 10 mg.l<sup>-1</sup> ammonia-N solution (hereafter noted, respectively, SW, DSW and N exposure). This solution was prepared by adding 1 mMol of ammonium chloride (Sigma, United States) to diluted seawater

(~5 ppt). Ammonia-N was chosen as experimental treatment because the ionized form of ammonia-N (NH4<sup>+</sup>) is the major component of domestic WW (Capdeville et al., 2018; Mégevand et al., 2021, submitted). This condition refers to total ammonia (ammonia-N). It represents the sum of unionized ammonia NH3 and ionized ammonia NH4<sup>+</sup> (Haywood, 1983; Lemarié et al., 2004; Bermudes and Ritar, 2008). The ammonia-N concentration was chosen on the basis of concentrations that may be released by treatment plants in Mayotte (Herteman, 2010). It corresponds to a sub-lethal concentration for a fiddler crab species such as *Uromastyx princeps*, adults sharing similar size and morphology with *P. chlorophthalmus* (Azpeitia et al., 2013). Ammonia-N exposure solutions were replaced every day. Salinity of SW and diluted seawater (DSW) were tested every day and readjusted if needed.

## **Osmolality Curves**

To determine osmolality curves, crabs were exposed at different salinities with 4–8 crabs per salinity for GT, and 8–10 crabs per salinity for PC. Salinity ranged from DSW (~4.5 ppt, ~135 mOsm.kg<sup>-1</sup>) to concentrated SW (~51 ppt, ~1,510 mOsm.kg<sup>-1</sup>). Animals were maintained in isolated boxes for a minimum of 72 h, which is sufficient time for osmoregulation to be achieved in such estuarine crustacean species (Lovett et al., 2001; Rivera-Ingraham et al., 2016).



After this period, a hemolymph sample was taken using a 0.1–0.5 ml hypodermic syringe with the needle inserted between the cephalothorax and the third and fourth periopod. Osmotic pressure (OP) of the hemolymph samples was immediately quantified in duplicate by freezing point depression osmometry (Model 3320, Advanced Instruments, Inc., Norwood, MA, United States).

At the end of this experiment, animals were put back in individual boxes in aerated SW before being released into the mangrove.

## **Oxygen Consumption Rate**

The experimental design consisted in a static, intermittent flow-through respirometry system based on Clark et al. (2013). Crabs were individually placed into 125 ml chambers allowing them to make sporadic movements considering their size, while ensuring accurate measurements of  $O_2$ consumption. Following procedures described in Killen (2014),  $O_2$  measurements were performed using a 4-channel fiberoptic system with contactless  $O_2$  sensor spots (FireSting  $O_2$ , PyroScience, GmbH, Aachen, Germany) where water oxygen content was quantified once every 5 s.

Water-mixing within the chambers was achieved with magnetic stirrers located under  $1 \text{ mm}^2$  mesh grid to avoid too much disturbance (Rivera-Ingraham et al., 2016). Sensors

were calibrated to 100 and 0% air saturation using air-saturated water and 80 mM Na<sub>2</sub>SO<sub>3</sub> solution, respectively. Chambers were filled with a tubing system providing control SW, DSW or contaminated water from aerated, filtered and temperature-controlled tanks ( $\sim$ 25°C). Crabs were gently introduced in the chambers and left to acclimate for 1 h in aerated seawater prior experiment.

Following acclimation, the system was filled with the exposure solution. A flush pump was switched on for 20 min to ensure proper mixing and was cut off during 40 min. During that time, the chambers were sealed. The decrease in oxygen content could be analyzed to indicate the rate of oxygen uptake. After the 40-min cycle, the pump was turned on to flush the metabolic chambers with aerated seawater (or DSW or DSW enriched with ammonia-N) during 20 min. Due to the presence of aerobic and anaerobic microorganisms capable of degrading organic compounds and consuming O<sub>2</sub> (Shchegolkova et al., 2016), a 20 min flushing was run to ensure 99% air saturation. The 40 min measurements allowed to make accurate O2 measurements without falling under the threshold of 70% air saturation in order to maintain aerobic metabolism and to avoid hypoxic stress (Ombres et al., 2011; Rodgers et al., 2016). O2 measurements were taken at T-1h (SW), T1h, T24h, and T96h after exposure (SW as control treatment, DWS or DSW with ammonia-N) for all individuals (N = 8 per condition). This 96 h exposure time

was chosen in order to better compare our results with data from the literature focused on the effects of pollutants and particularly ammonia-N on crustaceans. Many studies are based on the 96h LC50 with environmental concentrations of ammonia-N and regularly include histological analyses (De Freitas Rebelo et al., 2000; Wang et al., 2003; Barbieri et al., 2016; Weihrauch and O'Donnell, 2017). Only one female *GT* exposed to ammonia-N died after 72 h of exposure.

# **Osmoregulatory Capacity**

At the end of the 96 h respirometry exposure, a hemolymph sample from each animal was taken (N = 7-8 per condition) and OP was assessed as described above.

### **Sampling and Dissections**

Following the 96 h respirometry experiment, crabs were anesthetized and euthanized on ice for tissue sampling of the hepatopancreas, anterior and posterior gills in order to perform the histology measurements. Samples were fixed in Bouin's fixative solution for 48 h for histological analyses. Individual molting stage was verified through examination of an epipodite under a dissecting microscope after dissection (to avoid handlingassociated stress on the biological markers considered).

## **Histological and SEM Analysis**

For histological analyses, posterior gills (pair 7 or 8) and small samples of the hepatopancreas were rinsed in 70% ethanol following immersion in Bouin's fixative solution for 48 h. They were then dehydrated in a series of graded alcohols and embedded in paraffin. Tissue sections (4  $\mu$ m) were cut with a Microtome 2125RT Leica, then placed on Glycerin albumin precoated glass slides and deparaffinated. They were stained following Masson's Trichrome Staining Protocol (Martoja and Martoja-Pierson, 1967) using Haematoxylin Groat, Fuschine Ponceau and Aniline Blue staining solutions. Samples were examined using Leica DM6 microscope equipped with a Leica DMC 2900 and the associated software LAS X.

For the hepatopancreas analyses, pictures of the hepatopancreas for each individual were taken, each of them depicting 1-4 tubules. Images were scaled and analyzed using ImageJ software (version 1.51r). Tubules of the hepatopancreas were randomly selected for each individual. Four parameters were measured on each of those hepatopancreas tubules and, then, averaged for each individual (N = 3-8): the area of the tubules, the number of vacuoles per tubule, the area of the B-cell vacuoles, the percentage of B-cell vacuolization area per tubule. Posterior gill epithelial thicknesses were measured from areas where nuclei were visible using a  $40 \times$  objective (Figure 6C). Along this single-layered epithelium, cell heights were systematically measured from the base of the cuticle to the basal side of the cells facing the hemolymph lacuna. For statistical purposes, gills were divided into three sections along longitudinal (basal, central, and apical) and transversal axes (proximal, middle, and distal). For each of these areas, photographs were taken at the same magnification  $(60 \times)$ using the CapturePro software. Measurements of epithelial gill thickness were then performed using Image as well.

# STATISTICAL ANALYSIS

Statistical analyses were performed in R version 3.5.2 with Rstudio version 0.99.491 (Rstudio, Inc.), with an  $\alpha$ -level of 0.05 and 95% confidence intervals used to determine statistical significance in all tests. For osmoregulatory capacity and gills epithelial thickness, residuals were evaluated using Shapiro-Wilk and Levene's test, respectively, in order to test the data normal distribution behavior and variance homogeneity. Oneway ANOVAs were performed separately for posterior gills at each longitudinal level (apical, central, and basal) to assess potential differences of epithelial thickness between treatments (SW, DSW, and N) after 96 h exposure. To test potential effects of treatment on hepatopancreas parameters, we used generalized mixed-effects modeling with treatment as random effects (Manning, 2007; Zuur et al., 2013; Johnson et al., 2015) using the package glmmTMB (Brooks et al., 2017). The number of vacuoles per tubule, the mean area of the B-cell vacuoles per tubule, the percentage of B-cell vacuolization area per tubule in hepatopancreas were modeled in function of treatment (three-levels fixed factor) assuming a Gaussian distribution error with log link. Individual differences and repeated measurements among crabs and tubules were taken into account through a random intercept of tubule ID nested into crab ID. Then, type II ANOVAs (Wald chi-square test) were applied on the models using Anova function in CAR package (Fox and Weisberg, 2019). When significant effects of treatment were detected, they were investigated through pairwise comparisons using lsmeans package (Lenth, 2016) with Tukey correction. For O2 oxygen consumption, a generalized linear mixed model approach was taken assuming Gaussian distribution error with log link. Time (T-1h, T0h, T24h, and T96h) and treatment group (SW, DSW, and N) were considered as fixed factors. Crab ID was set as random factor nested within experimental tank (two tanks of exposure solution per treatment, N = 4 per tank), which are nested within treatment group (SW, DSW, and N). Interaction terms were examined using the approach using type II ANOVA and multiple pairwise comparisons followed by Bonferroni correction. All values are represented as average  $\pm$  SEM.

# RESULTS

## Tolerance to Salinity Variations and Osmoregulatory Capacity Under Osmotic and Ammonia-N Stress

Both GT and PC are strong osmoregulators across the tested salinities since they can maintain a gradient of greater than 600 mOsm.kg<sup>-1</sup> above the ambient medium when exposed to salinities close to FW (5–10 mOsm.kg<sup>-1</sup>, **Figure 3A**). Both species hyper-regulated at lower salinities (hemolymph osmolality actively maintained at levels above that of DSW) and hypo-regulated at higher salinity (hemolymph osmolality maintained below that of SW). The difference between external medium and hemolymph osmolality significantly increased as salinity decreased when both species were exposed to DSW



during 96 h (Figures 3B,C). GT can maintain osmoregulatory capacity values of ~50 mOsm.kg<sup>-1</sup> in SW and ~600 mOsm.kg<sup>-1</sup> in DSW after 96 h, and these values are significantly different [Figure 3B, one-way ANOVA F(2,20) = 89.44, p < 0.001]. The same results are observed in PC with osmoregulatory values of  $\sim 100 \text{ mOsm.kg}^{-1}$  in SW and  $\sim 600 \text{ mOsm.kg}^{-1}$  in DSW [Figure 3C, one-way ANOVA, F(2,20) = 1913, p < 0.001]. However, a break in the curve is detected at low salinities, when it became increasingly difficult for the animals to osmoregulate. For GT, this break occurs at a higher salinity ( $\sim 260 \text{ mOsm.kg}^{-1}$ ) than for PC (~118 mOsm.kg<sup>-1</sup>). Moreover, PC was able to maintain the same osmoregulatory capacity in DSW with ammonia-N as in DSW (Figure 3B), whereas a significant decrease in osmoregulatory capacity is observed in GT exposed to ammonia-N compared with DSW (Figure 3C): crabs cannot maintain the haemolymph-seawater interval as in DSW.

# Respiration Rate Across 96 h Diluted Seawater and Ammonia-N Exposures

Respiration rate measurements are depicted in **Figure 4A** for GT and **Figure 4B** for PC. They include temporal responses for each treatment at T-1h (SW), T0h, T24h, and T96h. In GT, a significant effect (p < 0.001) of exposure time on O<sub>2</sub> consumption ( $\mu$ mol O<sub>2</sub>·m<sup>-1</sup> g<sup>-1</sup>) was observed. The interaction between treatment and exposure time was significant [ $\chi^2(6, N = 31) = 16.831$ , p < 0.01]. *Post hoc* multiple comparisons on the interaction terms showed that crabs exposed to ammonia-N significantly increased

their respiration rate after 96 h of exposure compared to the other time steps and with DSW (**Figure 4A**). No effects of the osmotic or pollutant shock was observed immediately after exposure (T0h compared to T-1h).

In PC, no significant effect of treatment or exposure time was found, but their interaction was significant [ $\chi^2(6, N = 32) = 44.5581, p < 0.01$ ]. *Post hoc* multiple comparisons on the interaction terms showed that crabs exposed to SW significantly decreased their respiration rate after 96 h exposure compared with the other time steps (**Figure 4B**). Crabs exposed to DSW progressively increased their respiration rate until 96 h (significant difference between T-1h and T96h and between T0h and T96h). At 96 h of exposure, a significant difference was observed between SW and DSW only.

### **Histological Analyses**

The histology of the three main cell types of the pancreatic tubules of GT and PC is presented in **Figure 5I**: (1) R-cells (Restzellen, resorptive cells), the most abundant cell type in the hepatopancreas; (2) F-cells (Fibrillenzellen, fibrillar cells), which have a columnar shape and basophilic cytoplasm with an intense staining; (3) mature B-cells (Blasenzellen, blister-like cells), which are the largest cells showing large subapical vacuoles.

In GT, no effect of treatment was observed for all parameters measured in the hepatopancreas (One-way ANOVAS, **Table 1**, **Figures 5A–D**). In PC, a significant effect of treatment (p < 0.01) was reported regarding the percentage of B-cells vacuolization

in hepatopancreas tubules. Tukey *post hoc* showed an increased vacuolization process in DSW and ammonia-N compared to SW (**Figure 5H**).

No effect of treatment was observed in the gills of GT across the three levels of longitudinal axis (apical, central, and basal) (one-way ANOVAs, **Table 2** and **Figure 6A**). However, in PC, a significant effect of treatment (p < 0.0001) was reported at each level of the longitudinal axis (one-way ANOVAs, **Table 2**). Tukey *post hoc* tests revealed that the gill epithelium of crabs exposed to ammonium for 96 h was significantly thicker than those exposed to SW or DSW.

## DISCUSSION

# Osmoregulation and Ammonia-N Excretion

Osmoregulation is one of the key regulatory functions in aquatic organisms and represents an excellent indicator of the physiological status of crustaceans (Lignot et al., 2000). Crustaceans can be classified as strong, moderate, or weak osmoregulators depending on their capacity to tolerate low salinity and the magnitude of the hemolymph-seawater osmotic difference (Henry et al., 2012). Results indicate that both PC and GT are strong osmoregulators, but the breaking point in the osmolality curve (Figure 2) occurs at a higher salinity in GT compared to PC. Therefore, the physiological osmoregulatory limit toward low salinity for GT is reached before PC. Also, they do not share the same isosmotic point. It is higher in GT (~1,000 mOsm.kg<sup>-1</sup>) than in PC (~830 mOsm.kg<sup>-1</sup>). Similar results have been found in sympatric fiddler crabs living under different salinity conditions in the Gulf of Mexico (Thurman, 2004). Hypersaline species such as Leptuca subcylindrica and Minuca rapax are isosmotic for media between 840 and 940 mOsm.kg<sup>-1</sup> (26-30‰) whereas, freshwater/brackish water species such as Minuca longisignalis are isosmotic at 732 mOsm.kg<sup>-1</sup> (23‰). The ability to maintain hemolymph osmolality at different environmental salinities is well correlated with the habitat of the species (Lin et al., 2002), but may also depends on the species physiology itself. Consequently, a species living near estuaries or river mouths must be highly tolerant to salinity variation (hypothesis 1). Ecological knowledge about habitat preferences and distribution for both studied species tend to confirm this hypothesis. GT usually inhabits exposed, lower intertidal sandy or mud flat shores fringed with mangrove forest, coral rubble or rock (Dawn and Frith, 1977; Koga et al., 2000). This species is also considered as very marine, often living on shore and never near muddy river mouths (Crane, 1975). It has the largest longitudinal range of fiddler crabs, spanning the entire breadth of the Indian Ocean and Western Pacific subrealms, as described by Rosenberg (2020). PC does not live close to the open sea but near high-tide levels on the muddy banks and flats of mangrove estuaries, close to the mouth of streams or rivulets (Crane, 1975). It is one of the dominant species living in the study site of Mayotte and is one of the three endemic species from the East Africa Province (as defined by Rosenberg, 2020). This province also comprises three other

species extended into other provinces, including GT. Thus, both species can be found in Europa and Mayotte islands even if they do not share the same habitat level and salinity conditions.

Since benthic crustaceans can frequently experience elevated levels of localized ammonia-N when they bury in sediments with high organic contents (Weihrauch et al., 1999), it is believed that this necessitates an adaptive mechanism to remove excessive ammonia-N build-up within their hemolymph (Weihrauch et al., 2004; Romano and Zeng, 2012). Ammonia excretion has been widely studied in aquatic crustaceans and it is established that this function is intimately linked to the gills and their osmoregulatory function (see Weihrauch et al., 2004 for a complete description of the model).

The mechanisms for ammonia-N excretion in bimodal crabs are very diverse and complex (for a review, see Weihrauch and O'Donnell, 2017), as many membrane transporters are involved mainly in the anterior and posterior gills as well as in the antennal gland. The sodium pump Na<sup>+</sup>/K<sup>+</sup>-ATPase (NKA), K<sup>+</sup>-channels and Na<sup>+</sup>/H<sup>+</sup> exchangers (NHE) are among the key membrane proteins involved although more transporters could be at play (Masui et al., 2002, 2009; Lucu and Towle, 2003; Gonçalves et al., 2006; Freire et al., 2008; Weihrauch and O'Donnell, 2017). As already described, a number of these transporters playing a crucial role in trans-epithelial ammonia excretion are also involved in the ionoregulatory machinery (Weihrauch and O'Donnell, 2015, 2017). The best example for this is NKA, one of the key transporters responsible for energizing active NaCl transport processes in invertebrates and vertebrates alike (Larsen et al., 2014). The  $NH_4^+$  pump (often referred to NH4<sup>+</sup>/K<sup>+</sup>-ATPase) is equally important. However, this process reaches its limits when crustaceans are exposed to excessively high amounts of ammonia-N, generally resulting in reduced hemolymph Na<sup>+</sup> levels (Romano and Zeng, 2012). This phenomenon has been observed in several species living under various salinity conditions: the American clawed lobster, Homarus americanus (Young-Lai et al., 1991), the Kuruma shrimp, Marsupenaeus japonicus (Chen and Chen, 1996), the burrowing crab Neohelice granulata (Rebelo et al., 1999), the freshwater crayfish Pacifastacus leniusculus (Harris et al., 2001) and the mud crab Scylla serrata (Romano and Zeng, 2007a), amongst others. We can relate these data to our results on osmoregulatory capacity: increased ammonia-N excretion could induce a leak in Na<sup>+</sup> ions followed by a decrease in OC for GT, but not for PC. This may suggest that PC could set-up physiological mechanisms that allow a tolerance capacity to high ammonia-N levels in DSW. However, GT could not (rapidly) mobilize such tolerance. In most Crustaceans (i.e., H. americanus and *P. japonicus*), a decrease in hemolymph Na<sup>+</sup> levels is directly linked with reduced hemolymph osmolality and/or OC, but this is not observed in the euryhaline mud crab S. serrata (Romano and Zeng, 2007a). These authors propose that this crab species relies on other mechanisms to maintain osmotic pressure even in case of Na<sup>+</sup> reduced levels, as for example, by increasing the organic osmolytes such as hemolymph free amino acids (FAA). One can hypothesis that the difference observed in OC between GT and PC may result in different abilities to excrete ammonia-N (through NKA, NH4<sup>+</sup>/Na<sup>+</sup> exchanger, and/or other



transporters). With ammonia-N in DSW, GT may exhibit lower NKA activity, as reported in the blue swimmer crab *Portunus pelagicus* exposed to ammonia-N at low Na<sup>+</sup>/K<sup>+</sup> ratio (Romano and Zeng, 2011), leading to lower Na<sup>+</sup> levels in the hemolymph compared to PC. A possible explanation could be that PC may better cope with high levels of ammonia-N in its hemolymph than GT, by maintaining the activity level of its membrane transporters. Measuring ammonia-N, ion concentration and pH levels in the hemolymph would help to confirm this hypothesis.

# Implications of Osmoregulation Strategies in the Whole Metabolism of Both Species

Increased energy expenditure for osmoregulation and ammonia-N excretion may be reflected by increased metabolic indicators such as oxygen consumption (Gomez-Jimenez et al., 2004).

Our results showed no effects of DSW in respiration rate of GT across the 96 h of exposition compared to the controls in

DSW. In PC, respiration rate progressively increases in DSW, whereas O<sub>2</sub> consumption in SW progressively decreased until 96 h of exposure. Euryhaline crabs are able to compensate for diffusion loss of ions in dilute media by active uptake of gill epithelium (Genovese et al., 2004). The most energy-demanding ion transport mechanism is the active transport driven by the NKA (Lucu and Towle, 2003), with active ion pumping in specialized epithelial cells that are also mitochondria rich (Freire et al., 2008; McNamara and Faria, 2012). Ammonia-N excretion and hyper-osmoregulation mainly rely on these mechanisms (Weihrauch et al., 2004) which generally increases when estuarine/marine crustaceans are exposed to diluted seawater (reviewed in Romano and Zeng, 2012). Based on the large loss of salt in diluted media, these crustaceans must invest considerable amounts of energy in active transport/uptake of salt in order to compensate for the large rates of diffusive salt loss (Henry et al., 2012). When osmotic stress is coupled with ammonia-N stress, an increase in O2 consumption is observed in GT after 96 h exposure, but this was not observed in PC.



**FIGURE 5** | Histological analyses from transverse sections of different tubules of GT (A–D) and PC (E–H) hepatopancreas exposed to seawater (SW), diluted seawater (DSW) and 10 mg.L<sup>-1</sup> ammonia-N (N). Measurements and counts were carried out on randomly distributed tubules per individual. Data were averaged per individual. (A,E) Areas of the tubules expressed as mean per individual; (B,F) B-cell vacuole areas expressed as mean per tubule per individual; (C,G) number of B-cell vacuoles in the tubule; (D,H) percentage of B-cell vacuolization area per tubule; (I) light microscopy illustration of a tubule cross section (GT). The photograph shows a tubule epithelium containing absorbing R-cells (R), enzyme synthetizing F-cells (F), and tubule-clearing B-cells (Bv). L, lumen; N, nucleus. Values are means  $\pm$  SEM (N = 3-8 per treatment). Different letters represent statistically significant differences between treatments at p < 0.05 after Wald Chi-square test and multiple comparisons (Tukey).

This could be directly linked to the OC decrease observed in this species. An increased energy demand is likely to occur in case of ammonia-N prolonged exposure. For instance, GT could rely on different ionic transport mechanisms without necessarily modifying its gill epithelial thickness (as discussed in Section "Histological Changes to Explain Paraleptuca chlorophthalmus and Gelasimus tetragonon Differential Ammonia-N Tolerance"). Even with an increased uptake of O2, it is possible that the maintenance of GT hydromineral balance and ammonia-N excretion function is less efficient when facing these stressors than in PC. This could highlight a physiological trade-off because of the lack of other adaptive mechanisms, but also due to any physiological differences between the two species. In PC maintained in DSW, the increase in O<sub>2</sub> uptake could be due to an increased energy demand for osmoregulation (among others). However, a lower oxygen consumption in DSW with ammonia-N could suggest an increased anaerobic metabolism or a different compromise between gas exchange and osmoregulation at the gill surface (aka the osmo-respiratory compromise) (Robertson et al., 2015). A clear functional specialization has been reported for several crab species, posterior gills being the main effectors of the "compensatory" process, while anterior gills are mostly involved in gas exchange (Taylor and Taylor, 1992; Pequeux, 1995; Genovese et al., 2004). Anterior gills are also known to be actively involved in ammonia excretion (Weihrauch et al., 1998, 1999) and in some hyper-regulating species such as green shore crab, active ammonia excretion rates being lower in posterior osmoregulatory gills compared to

anterior gills (Weihrauch et al., 1998). These results indicate that osmoregulatory processes are not directly linked to ammonia excretion and that excretion of toxic ammonia could proceed independently of other physiological processes (Weihrauch and O'Donnell, 2017). Large areas in the lamellae of the posterior gill epithelium reveals thick ion-transporting cells with welldeveloped basolateral membranes associated with numerous mitochondria in several species of euryhaline crabs such as *Neohelice granulata* (previously *Chasmagnathus granulatus*) (Genovese et al., 2004). Most of the NKA activity is generally restricted to specific areas of posterior gill lamellae lined by this kind of epithelium (Towle and Kays, 1986).

# Histological Changes to Explain *Paraleptuca chlorophthalmus* and *Gelasimus tetragonon* Differential Ammonia-N Tolerance

In DSW, no changes were observed in the thickness of the gill epithelium for the two species (**Figures 6A,B**). However, when exposed to ammonia-N for 96 h, branchial epithelium thickness of PC significantly increases along the longitudinal axis, whereas this does not occur for GT. An increase in osmoregulatory epithelial patches surface of posterior gills in crabs exposed to either dilute or hyper-saline water (i.e., when crabs osmoregulate) has already been observed (Genovese et al., 2000; Lovett et al., 2006). The authors suggest that these gills could be involved both in active ion uptake and active excretion of ions depending on

A	Hepatopa	ancreas tubu	le areas (µm	2)						
Factor	DF	χ <sup>2</sup>		Р						
	GT-PC	GT	PC	GT	PC					
Treatment	2	0.9819	1.4586	0.6121	0.4822					
В	Number of vacuoles/tubules									
Factor	DF	χ <sup>2</sup>		Р						
	GT-PC	GT	PC	GT	PC					
Treatment	2	1.4951	1.3823	0.4735	0.501					
с	B-cells vacuole areas (μm <sup>2</sup> )									
Factor	DF	Х	2	Р						
	GT-PC	GT	PC	GT	PC					
Treatment	2	1.0779	4.4643	0.5834	0.1073					
D	% of vacuolization/tubule									
Factor	DF	χ <sup>2</sup>		Р						
	GT-PC	GT	PC	GT	PC					
Treatment	2	3.1161	9.5381	0.2105	0.008489					

**TABLE 1** | Results of analysis of deviance for histological measurements from transverse sections of different hepatopancreas tubules of *G. tetragonon* and *P. chlorophthalmus* as a function of treatment.

GT, Gelasimus tetragonon (Europa Island); PC, Paraleptuca chlorophthalmus (Mayotte). Wald Chi-square tests are performed for testing the significant differences of fixed categorical variables in GLMM (N = 7–8 per treatment for 3 treatment group). Statistically significant effects are indicated by asterisks: \*\*p < 0.01 and appear in bold.

the medium salinity, especially with the progressive increase of NKA activity during 6 days observed in crabs exposed to dilute seawater (Lovett et al., 2006). In PC, this response may be directly related to a strategy allowing increased ammonia-N excretion and osmoregulation may be through NKA increased activity, as discussed above. By increasing the thickness of the basolateral epithelium of posterior gill cells, PC may increase the number of ion transporters (mainly NKA) and mitochondria allowing, therefore, an increase in its osmoregulatory and ammonia-N excretion capacities. The increase in epithelial thickness does not occur in DSW although a higher metabolic rate is observed in this condition. Therefore, PC could rely on two different strategies depending on the nature and level of stress (increased metabolic rate in DSW and/or enhanced ion transport mechanisms when exposed to ammonium by the lengthening of the basolateral epithelium), whereas GT cannot exhibit such plasticity regarding ammonia-N exposure and increases its metabolic rate.

However, though not quantified, no major disruption of cellular structure has been observed in either species exposed to DSW or Ammonia-N. In Romano and Zeng (2007b), Ammonia-N exposure for 96 h lead to different dose-dependent disruptions of gills in *P. pelagicus*. At 10 mg.l<sup>-1</sup> ammonia-N (a similar concentration as in the present study), gill lamellae showed

**TABLE 2** Results of analysis of variance for gill epithelial thickness measured on different parts of the longitudinal axis of the gill one-way ANOVAs).

<b>F</b> <b>PC</b> 2 63	<b>GT</b> 0.8836	<b>PC</b> 20.635	GT	PC	GT	P PC				
2	0.8836		GT	PC	GT	PC				
		20.635				10				
63		20.000	0.551	15.36	0.579	3.68E-06	***			
	1.6029	1.343								
Gill epithelial thickness – central part										
F	MS		F		Р					
PC	GT	PC	GT	PC	GT	PC				
2	2.054	14.839	1.503	13.79	0.231	1.07E-05	***			
63	1.367	1.076								
pithe	lial thickne	ess – basa	al part							
F	MS		F		Р					
PC	GT	PC	GT	PC	GT	PC				
2	2.057	20.844	0.841	11.37	0.437	6.08E-05	***			
63	2.447	1.833								
	PC           2           63           apithe           F           PC           2           63	F         N           PC         GT           2         2.054           63         1.367           apithelial thickne         N           PC         GT           PC         GT           2         2.057           63         2.447	F         MS           PC         GT         PC           2         2.054         14.839           63         1.367         1.076           epithelial thickness – base         F         MS           PC         GT         PC           2         2.057         20.844           63         2.447         1.833	F         MS         GT         PC         GT         GT </td <td>F         MS         F           PC         GT         PC         GT         PC           2         2.054         14.839         1.503         13.79           63         1.367         1.076         1.076         1.079           printelial thickness – basal part           F         MS         F           PC         GT         PC         GT         PC           2         2.057         20.844         0.841         11.37           63         2.447         1.833         1.076         1.076</td> <td>F         MS         <math>F</math>         GT         PC         GT         PC         GT         PC         GT         I.303         13.79         0.231         3.379         3.379         3.379         3.379         3.379         3.379         3.379         3.379         3.379         3.379         &lt;</td> <td>F         MS         F         P           QC         QT         PC         QT         PC         QT         PC           2         2.054         14.839         1.503         13.79         0.231         1.07E-05           3         1.367         1.076         PC         P         P           appithelial thickness - basal part         P         P         P           PC         QT         PC         QT         P           QC         QT         PC         QT         P           QC         QT         PC         QT         PC           QT         PC         QT         PC         QT         PC           QT         QT         PC         QT         PC         QT         PC           QT         QT         PC         QT         PC         QT         PC           QT         QT         QT         QT         QT         QT         QT           QT         QT         QT         QT         QT         QT         QT           QT         QT         QT         QT         QT         QT         QT           QT         QT</td>	F         MS         F           PC         GT         PC         GT         PC           2         2.054         14.839         1.503         13.79           63         1.367         1.076         1.076         1.079           printelial thickness – basal part           F         MS         F           PC         GT         PC         GT         PC           2         2.057         20.844         0.841         11.37           63         2.447         1.833         1.076         1.076	F         MS $F$ GT         PC         GT         PC         GT         PC         GT         I.303         13.79         0.231         3.379         3.379         3.379         3.379         3.379         3.379         3.379         3.379         3.379         3.379         <	F         MS         F         P           QC         QT         PC         QT         PC         QT         PC           2         2.054         14.839         1.503         13.79         0.231         1.07E-05           3         1.367         1.076         PC         P         P           appithelial thickness - basal part         P         P         P           PC         QT         PC         QT         P           QC         QT         PC         QT         P           QC         QT         PC         QT         PC           QT         PC         QT         PC         QT         PC           QT         QT         PC         QT         PC         QT         PC           QT         QT         PC         QT         PC         QT         PC           QT         QT         QT         QT         QT         QT         QT           QT         QT         QT         QT         QT         QT         QT           QT         QT         QT         QT         QT         QT         QT           QT         QT			

GT, Gelasimus tetragonon (Europa Island); PC, Paraleptuca chlorophthalmus (Mayotte).

N = 7–8 per treatment for 3 treatment group. Statistically significant effects are indicated by asterisks: \*\*\*p < 0.001 and appear in bold.

local infiltrations of hemocytes with normal lamellae structures whereas 60 mg.l $^{-1}$  ammonia-N generated extensive hemocyte infiltrations, disrupted pillar cells and lamellae collapses.

In Crustaceans, the hepatopancreas is a key metabolic organ, expressing a variety of transporters (Ahearn et al., 1992). Its numerous tubules are involved in food digestion, detoxification of xenobiotics and nutrient absorption (Ceccaldi, 1998; Zilli et al., 2003; Ortega et al., 2014). Histological analyses of these tubules revealed few changes for the B-cells in both GT and PC (Figures 5A-G). Even if their role is still discussed, B-cells can be considered to have two functions: elimination of waste material and intracellular digestion of nutrients (Vogt, 1994). The supranuclear vacuoles resulting from the coalescence of subapical vacuoles take the form of huge lysosomes (Vogt, 1993) and their content can be discharged into the lumen by holocrine secretions (Nott et al., 1985). Our results showed a higher vacuolization in PC exposed to Ammonia-N that could be linked to an increased intracellular digestion process through B-cells vacuoles, as it has been found in a Sesarmid crab species exposed to domestic wastewater (Mégevand et al., 2021). This increase in activity would potentially lead to an increased energy expenditure but in this case, our hypothesis is that ammonia-N could have a limited effect on the hepatopancreas functioning. As mentioned by Chu (1987), the hepatopancreas overall contribution to osmoregulatory ion uptake is believed to be minimal, whereas gills are the main site for respiration, osmoregulation, ammonia-N excretion, these gills being the main organs involved for the overall organismal adaptation to changing conditions. In Ocypodids such as PC and GT,



the study of the antennal gland would also be of interest in order to understand ammonia excretion mechanisms. As shown for several fiddler crabs such as Minuca pugnax or Leptuca pugilator, ammonia is known to be actively transported across the duct and excreted in the primary urine which contains a high concentration of ammonium (Green et al., 1959; De Vries et al., 1994). Other ecomorphological markers in addition to those used on the hepatopancreas and gills would be interesting to study as indicators of adaptation to changing environments. For example, Lim and Goh (2021) demonstrated that the setae present on the maxillipeds are great indicators of adaptation, especially for efficient food extraction regarding sediment quality. Fiddler crabs with greater diversity and density in setae could be able to adapt to more variable habitats since their ability to extract food from different sediment types would be better.

Coping with an aquatic environment with low salinity and elevated ammonia-N is highly species- and habitat-specific. Therefore, a fully euryhaline species such as PC, that is adapted to nutrient-rich environments (Gross, 1982; Henry et al., 2012) is likely to be less vulnerable to nutrient enrichment and may be able to develop other coping mechanisms. However, this study focuses on potential short-term acclimation capacities. It does not take into account any possible environmental determinism that could arise from micro-evolution after prolonged exposure to ammonia-N enriched conditions (in Mayotte) or differential abiotic factors such as eutrophic or oligotrophic environments.

A research program with multi-generational expositions of the two populations would be of high interest, especially for understanding ecosystem health and ecological risk implications on a longer timescale (although technically complicated to implement).

# Ecosystem and Risk Assessment Implications

Acclimatory and adaptive capacities to different environments induce species-specific responses among Decapod crustaceans that rely on various mechanisms to maintain their metabolism (Weihrauch et al., 1999; Henry et al., 2012; Romano and Zeng, 2013).

However, brackish and intertidal environments are among the most stressful aquatic biotopes and the establishment of Crustacean communities in such habitats suppose highly adapted physiological features (Gilles and Pequeux, 1983; Charmantier et al., 2002) to cope with low and varying salinities (Henry et al., 2012), as well as constantly changing environments. For example, it has been shown by Yong and Lim (2021) that ocypodid crabs such as *Ocypode gaudichaudii* possess a feeding plasticity when confronted to different habitat types, which is related to their burrowing and foraging strategies. In the case of our study, salinity variations and nitrogen enrichment phenomena could also induce different levels of adaptation in feeding and foraging habits, which would be interesting to study in both species in parallel with the individual physiology.

In the present study, PC appears to have a better physiological tolerance to decreased salinity compared to GT, even if GT is a relatively strong osmoregulator (hypothesis 1). Ammonia-N triggers short-term acclimation in PC that was not observed in GT suggesting that this species could be more vulnerable to anthropogenic discharges (hypothesis 2). GT copes with 96 h ammonia-N exposure through increased metabolic rate to excrete this product, while struggling to maintain its osmoregulatory balance. However, this response does not occur in DSW condition without ammonia-N. This suggests that some cumulative deleterious effects under Ammonia-N exposure and osmotic stress occur for this species (hypothesis 3).

Ammonia-N was used in this study as a first indicator of anthropogenic effluents, but it is more likely that crab communities are confronted with different contaminants with combined effects (e.g., complex and hypoxic wastewater), often consisting of repeated exposures. Worldwide, effects of domestic effluent discharges (controlled or not) on crab community assemblage have been observed in several different mangroves with various results (Wear and Tanner, 2007; Penha-Lopes et al., 2009; Wickramasinghe et al., 2009; Capdeville, 2018). The results of the present study point (at least) to a short-term physiological acclimation to salinity variation and pollution for species such as PC, acclimated to euryhaline and anthropized conditions (already proposed in Mégevand et al., 2021, submitted), but it has been demonstrated that individual abundance for this species decreased in chronically WW impacted areas (Capdeville et al., 2018). If GT is unable to develop short-term response to salinity and ammonia-N inputs, it is likely that this species would be even more sensitive to longer exposure to pollution, or to more complex contaminants such as hypoxic WW. If the population decline of engineering species can be dreadful for the functioning of freshwater-influenced mangroves (e.g., Mayotte), what about isolated and pristine ecosystems like Europa Island? What would be the acclimation potential of representative species like GT in the event of anthropogenic effluents? In view of the

results obtained in this study and those obtained during the various works mentioned above, it is possible to be pessimistic about the ecosystem risks incurred by a pristine mangrove such as that of Europa if it were to face waters discharges of anthropogenic origin.

# CONCLUSION

The two crab species studied here showed different physiological plasticity that led to different tolerance to salinity variations and ammonia-N exposure. In accordance with our first hypothesis, P. chlorophthalmus (Mayotte Island) has a better osmoregulatory capacity in diluted seawater than G. tetragonon (Europa Island), which remains, however, a strong hyper-hypo-osmoregulator. Exposure for 96 h to ammonia-N at a concentration of 10 mg.l<sup>-1</sup> in diluted seawater (5ppt) generates branchial morphological acclimation in PC most likely in order to increase the osmoregulatory and Ammonia-N excretion functions. These changes do not occur in GT, this species experiencing an increase in oxygen consumption over time as its osmoregulatory capacity decreases (hypothesis 2). The two species do not have the same tolerance to short-term nitrogen inputs (as a simulation of anthropogenic presence) (hypothesis 3). Burrowing crabs ensure crucial ecological functions associated with their bioturbation activity. A change in crab community composition will not ensure the maintenance of these functions as functional redundancy may not happen. Consequences in terms of mangrove functioning and vulnerability are difficult to predict and linked to many others parameters. However, our study emphasizes the importance of considering small/medium scale ecophysiological indicators of species vulnerability, especially when developed in engineer species, to answer large-scale questions of ecosystem risk assessment.

# DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/**Supplementary Material**, further inquiries can be directed to the corresponding author.

# **AUTHOR CONTRIBUTIONS**

LM: conceptualization, methodology, investigation, formal analysis, and writing – original draft. DT: conceptualization, methodology, investigation (osmoregulation curves and osmoregulatory capacity), formal analysis, and writing – review and editing. CLÉ: investigation (histological analyzes). SH: methodology (histological analyzes), investigation (histological analyzes), and writing – review and editing. EC: methodology, investigation (molecular analysis of rRNA sequences), and writing – review and editing. TL'H: investigation (histological experiments) and writing – review and editing. J-HL: supervision, conceptualization, investigation, and writing – review and editing. ES: project administration, funding acquisition, supervision, conceptualization, investigation, and writing – review and editing. All authors contributed to the article and approved the submitted version.

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sampling in the Scattered Islands and providing logistical support during the scientific cruise and our stay on Europa Island.

### SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fevo.2022. 839160/full#supplementary-material

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