

Osmoregulatory performance and immunolocalization of Na+/K+-ATPase in the branchiopod Artemia salina from the Sebkha of Sidi El Hani (Tunisia)

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1 2	Osmoregulatory performance and immunolocalization of Na ⁺ /K ⁺ -ATPase in the branchiopod <i>Artemia salina</i> from the Sebkha of Sidi El Hani (Tunisia)
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20 21	Abstract Artemia salina is an extremophile species that tolerates a wide range of salinity, especially
22	hypertonic media considered lethal for the majority of other aquatic species. In this study, A.
23	salina cysts were hatched in the laboratory and nauplii were acclimated at three different
24	salinities (60, 139 and 212 ppt). Once in the adult phase, their hemolymph osmolality was
25	measured. The animals were strong hypo-osmoregulators in the entire range of tested
26	salinities, with up to 10 fold lower hemolymph osmolalities than their surrounding
27	environment. Immunostaining of Na ⁺ /K ⁺ -ATPase was done on sections and on whole body
28	mounts of adults in order to localize the ionocytes in different organs. An intense Na ⁺ /K ⁺ -
29	ATPase immunostaining throughout the cells was observed in the epithelium of the ten pairs
30	of metepipodites. A positive immunoreactivity for Na ⁺ /K ⁺ -ATPase was also detected in the
31	maxillary glands, in the epithelium of the efferent tubule and of the excretory canal, as well as
32	in the anterior digestive tract. This study confirms the strong hypo-osmotic capacity of this
33	species and affords an overview of the different organs involved in osmoregulation in A.
34	salina adults.
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37 38 39	<i>Keywords:</i> Artemia salina, hypo-osmoregulation, Na ⁺ /K ⁺ -ATPase, immunolocalization, metepipodites, maxillary glands
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41 **1. Introduction**

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Artemia salina (Linnaeus, 1758) is part of a complex of euryhaline species of 43 branchiopod crustaceans qualified as an animal extremophile with an exceptionally wide 44 salinity tolerance among animals, and particularly an ability to survive and thrive in 45 hypertonic media, a "forbidden environment" from which most metazoans are excluded as 46 stated by Eads (2004) (Abatzopoulos et al., 2002; Browne and Bowen, 1991; Clegg and 47 Trotman, 2002). These animals have attracted the attention of scientists over decades and 48 49 have been the subject of extensive studies in order to understand the mechanisms allowing 50 them to tolerate ionic and osmotic stresses imposed by their environment.

51 These remarkable crustaceans are able to withstand ionic contents and salinities that, in their upper values, are lethal for the majority of other aquatic species, ranging from 9 ppt 52 53 (Brisset et al., 1982) to 340 ppt (Gajardo and Beardmore, 2012; Post and Youssef, 1977) and up to crystallizing brine close to 600 ppt (Croghan, 1958a, b). Artemia salina is a highly 54 potent hypo-osmotic regulator in media more concentrated than one third of seawater (salinity 55 of about 11 ppt corresponding to an osmolality of 300 mOsm.kg⁻¹). For instance, early studies 56 had shown that its hemolymph osmolality is maintained between 400 and 800 mOsm.kg⁻¹ for 57 salinities ranging from 34 to 340 ppt (1000 to 10 000 mOsm.kg⁻¹) (Croghan, 1958b). 58

Such levels of hypo-osmoregulation suppose the existence of powerful ion excretion 59 abilities in order to compensate the passive invasion of ions from the concentrated external 60 medium. Moreover, animals have to limit water loss by osmosis, but water uptake 61 mechanisms are not well known in crustacean, notably in those high-salinity environments. In 62 larvae, Conte (1984) has extensively described the mechanisms of osmoregulation in the 63 nauplii of A. salina that use a special salt-secreting gland, also called salt gland or neck or 64 nuchal organ, to excrete salts that penetrate by diffusion. Similarly, Russler and Mangos 65 (1978) have shown that this organ is the main route for sodium excretion. 66

In adult A. salina, different studies have shown a similarity between their mechanisms 67 68 of osmoregulation and those proposed for marine teleosts and hypo-regulating crustaceans (Croghan, 1958b, c, d; Copeland, 1967; Smith, 1969a, b). Brine shrimps have well-developed 69 active mechanisms for absorbing NaCl from the gut lumen to the hemolymph. Hence, water 70 follows passively to compensate water lost by osmosis to the external concentrated medium 71 (Croghan, 1958d). Also in adults, sodium and chloride, which enter the body through 72 73 diffusion given their high concentration in the external medium, are excreted to the medium 74 by specialized organs acting as gills, the metepipodites. The ten pairs of flattened, leaf-like metepipodites borne by the phyllopods have been described as the sites of active outward
transport of ions (Copeland, 1967; Croghan, 1958c).

In particular, a detailed study of the metepipodites of A. salina was conducted by 77 Copeland (1967). Using silver nitrate, he confirmed that the metepipodites were the site of ion 78 (chloride) exchanges, a finding reported earlier by Croghan (1958c) and later confirmed by 79 other authors (Holliday et al., 1990). Copeland's ultrastructural exploration revealed the 80 presence and association of two types of cells in the metepipodites, the "light" and "dark 81 cells". The latter are columnar cells with projected stellate flanges; they extend from the 82 83 apical cuticle to the basal membrane lining the hemolymph, i.e. they face the external medium as well as the hemolymph, and they present deep interdigitations with the light cells. Stacks of 84 mitochondria, called "mitochondrial pumps" by Copeland, are located in the dark cells in 85 close association with their membrane, particularly in their projections. From these 86 87 observations, Copeland concluded that the metepipodites represent "a highly specialized tissue for the secretion of salt". Given the location of the dark cells, their ultrastructural 88 89 features and particularly the abundance of mitochondria in their cytoplasm, "it would appear likely that the dark cell is responsible" for the "release of salt to the external environment" 90 91 (Copeland, 1967). This hypothesis was confirmed by later experimental findings. In fact, the crude homogenates of metepipodites showed a very high specific enzyme activity of Na⁺/K⁺-92 ATPase, which increased proportionally with the salinity of the external medium (Holliday et 93 al., 1990). The same author found that the digestive tract and maxillary glands also had a high 94 Na⁺/K⁺-ATPase content, which suggests that these organs are also involved in ion transports 95 and probably for a part in osmoregulatory processes. Membrane-bound Na⁺/K⁺-ATPase was 96 partially purified from A. salina nauplii (Morohashi and Kawamura, 1984) and the α isoform 97 of the enzyme was isolated in salt glands and intestine of nauplii (Cortas et al., 1989). Also in 98 naupliar larvae, Na⁺/K⁺-ATPase α and β isoforms were immunolocalized in the basal 99 membranes of the salt gland cells (Sun et al., 1991) and the mRNA expression of their $\alpha 1$ and 100 101 α 2 isoforms was quantified in the same organs (Conte, 2008; Escalante et al., 1995; Sun et al., 1992). While the localization of the enzyme is known in the larval salt gland, to our 102 knowledge surprisingly few reports are available regarding the localization of Na⁺/K⁺-ATPase 103 in adult Artemia. Recently, an illustration of the presence of Na⁺/K⁺-ATPase in a 104 105 metepipodite of A. franciscana has been made available in a preliminary report (Drenth, 2017). 106

107 The present study aims to confirm the ability of *A. salina* to osmoregulate at very high 108 salinities, and to determine the localization of Na⁺/K⁺-ATPase in the metepipodites, maxillary 109 glands and digestive tract of adults, in order to improve the understanding of osmoregulatory 110 mechanisms allowing a Tunisian *Artemia salina* population (from the hyper-saline lagoon 111 Sebkha of Sidi El Hani) to withstand high salinities.

- 112
- 113 **2.** Materials and methods

2.1. Cysts sampling

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Artemia cysts were collected in spring 2017 from the banks of the hypersaline Sebkha
of Sidi El Hani where the salinity exceeded 280 ppt.

119 The Sebkha of Sidi El Hani is a NW-SE lengthened depression in the Sahel area (eastern Tunisia) (Fig. 1). The Sidi El Hani discharge area is approximately 370 km² and its average 120 121 water depth is 0.4 m, and 0.8 m at some locations (Fig. 1). The seasonal temperature fluctuates between 2 and 13 °C in winter and between 33 and 39 °C in summer (Ali et al., 122 123 2013). The seasonal average of salinity oscillates between a minimum of 180 ppt in winter registered after rainfall and up to 320 ppt in summer. The ionic composition of Sidi El Hani is 124 mostly based on Na⁺ (115 gL⁻¹), Cl⁻ (191 gL⁻¹), Mg²⁺ (8 gL⁻¹) and SO₄²⁻ (13 gL⁻¹). The 125 presence of Artemia in the site was first reported by (Gauthier, 1928). 126

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2.2. Culture experiments

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Upon collection, cysts were mixed with salt collected in the Sidi El Hani area, for 130 conservation. Once in the laboratory, cysts of Artemia salina were cleaned, separated and 131 132 stored according to the protocol of Sorgeloos et al. (1986). Cysts were incubated over 48 hrs in 1 liter of filtered seawater at a salinity of 35 ppt and a temperature of 25 °C, under constant 133 illumination (2000 lux) and continuous aeration to keep them in suspension (Lavens and 134 Sorgeloos, 1996). Following hatching, the nauplii were transferred to 3 containers of 5L filled 135 with water at different salinities, 60, 139 and 212 ppt. These media had been prepared from a 136 mixture of seawater sterilized in an autoclave and raw salt (harvested at the Sebkha of Sidi El 137 Hani). In each container, the salinity of the water was checked using a salinometer (Lovibond 138 SensoDirect con110), and the osmolality was measured with an Advanced Instruments 3300 139 micro-osmometer in later experiments. The containers were placed at a temperature of 25-27 140

°C and a photoperiod of 16 hrs L / 8 hrs D. Artemia nauplii density was adjusted to about 50 141 individuals per L. These larvae were fed twice a week by adding 100 ml of Chlorophycea 142 Dunaliella salina culture at an approximate density of 100,000 cells.ml⁻¹ (the salinity of the 143 microalga culture was the same as that used for the culture of nauplii). Water was renewed 144 once a week to ensure the elimination of Artemia salina waste and agglomerated microalgae. 145 Once the adult phase was reached, some animals were fixated for histology, others were 146 transported over 24 hrs in sealed containers to the Marbec laboratory (Montpellier, France) 147 148 for further experiments.

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2.3. Measure of hemolymph osmolality

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Hemolymph osmolality was measured in 8-12 adult A. salina per condition. According 152 153 to previous results on different crustacean species (Charmantier, 1998), the animals were exposed directly to the experimental media for 48 hrs in covered beakers filled with water at 154 155 three different salinities maintained at 24 °C, aerated and covered with parafilm to prevent evaporation. The salinities of the media, 60, 139 and 212 ppt, corresponded to osmolalities of 156 1737, 4064 and 6192 mOsm.kg⁻¹, according to measures made on an Advanced Instrument 157 Model 3300 micro-osmometer. Prior to sampling, the specimens were carefully rinsed with 158 deionized water, dried on filter paper and quickly immersed into mineral oil to prevent 159 evaporation and desiccation. Remaining adherent water was removed using a hand-made 160 glass micropipette. A second micropipette was then inserted dorsally into the heart to obtain 161 hemolymph samples, which were then immediately measured with reference to a 300 162 mOsm.kg⁻¹ standard solution on a Kalber-Clifton nano-osmometer (Clifton Technical Physics, 163 Hartford, NY, USA) requiring about 30 nl. 164

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2.4. Histology and immunolocalization of Na⁺/K⁺-ATPase

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In order to localize the ionocytes, and to visualize and semi-quantify Na⁺/K⁺-ATPase, we performed histology, immunofluorescence staining, whole-mounts and microscopic observations. About twenty adult brine shrimp from each salinity condition were fixed for 24 hrs by immersion in Bouin's fixative. Once rinsed in several baths of 70 % ethanol, samples were dehydrated in a graded ethanol series and finally embedded in Paraplast (Sigma). Longitudinal and transverse sections of 4 μm were cut on a Leitz Wetzlar microtome, collected on poly-L-lysine coated glass slides and stored at 37 °C for 48 hrs. One series of

slides was stained using the Masson's Trichrome staining protocol and observed under a Leica 175 Diaplan microscope. The other series was used for in situ immunolocalization of Na⁺/K⁺-176 ATPase. Slides were dewaxed (butanol and LMR), hydrated through a descending series of 177 ethanol baths (from 100 % to 50 %) and rinsed in phosphate-buffered saline (PBS; 137 mM 178 NaCl, 2.7 mM KCl, 10 mM phosphate buffer, pH 7.4, Sigma). The slides were then immersed 179 for 10 min into 0.02 % Tween 20, 150 mM NaCl dissolved in PBS, pH 7.3. After blocking in 180 5 % skimmed milk (SM) in PBS at 37 °C for 20 min, the slides were rinsed twice with PBS. 181 Primary labelling was carried out overnight at 4 °C in a wet chamber with the primary 182 monoclonal antibody (α5) mouse anti-Na⁺/K⁺-ATPase (Hybridoma Bank, University of Iowa) 183 diluted in 0.5% SM-PBS at 10 μ g.mL⁻¹. Using a procedure similar to that of this study, this 184 antibody was shown to specifically react with Na⁺/K⁺-ATPase in several crustacean species, 185 such as Porcellio scaber (Ziegler, 1997), Homarus gammarus (Lignot et al., 1999), Astacus 186 leptodactylus (Lignot et al., 2004), Carcinus maenas (Cieluch et al., 2004), Crangon crangon 187 (Cieluch et al., 2005), Eriocheir sinensis (Cieluch et al., 2007), Macrobrachium amazonicum 188 (Boudour-Boucheker et al., 2014) and Eurytemora affinis (Gerber et al., 2016). Once rinsed 189 three times in PBS to remove unbound primary antibody, the sections were incubated for 1 hr 190 with a secondary antibody (donkey anti-mouse Alexa Fluor® 488 (Invitrogen, Life 191 Technologies) at 10 μ g·mL⁻¹ in SM-PBS) and rinsed again three times in PBS. Nuclei of 192 193 some slides were counterstained using DAPI at 1µg/ml for 6 min, followed by three washes in PBS. 194

Several protocols of preparation and immunostaining of whole mounts of A. salina 195 have been tried. A limitation of antibody penetration was encountered probably due to the 196 cuticle barrier. EDTA, ultrasonic treatment from 5 s to 2 min, and Triton X-100 were tested in 197 different combinations. Positive staining was obtained using ultrasonic pulses at 35 KHz from 198 10 s to 20 s as indicated thereafter. Even in the best preparations, only some metepipodites of 199 the animals showed positive staining. Following pretreatment with ultrasonic pulses (from 10 200 s to 20 s) to enhance probe accessibility, whole mounts of A. salina adults were pre-incubated 201 in 5 % skimmed milk (SM) in PBS and 0.3 % Triton X-100 (PBS-TX) for 2 hrs at room 202 203 temperature. After blocking, the animals were rinsed several times with PBS. Primary labelling was carried out overnight at 4 °C in a wet chamber with the primary monoclonal 204 205 antibody (a5) mouse anti-Na⁺/K⁺-ATPase (Hybridoma Bank, University of Iowa) diluted at 10 µg.mL⁻¹ in 0.5 % SM-PBS. Following three rinses in PBS, the specimens were incubated 206 207 for 3 hrs with a secondary antibody (donkey anti-mouse Alexa Fluor® 488 (Invitrogen, Life

Technologies) at 10 μ g.mL⁻¹) and rinsed again in PBS. Nuclei of some mounts were counterstained using DAPI at 1 μ g.mL⁻¹ for 6 min, followed by three washes in PBS.

Sections and whole mount preparations were then mounted in an anti-bleaching mounting medium (Immunohistomount, Santa Cruz Biotechnology) and observed with a Leica Diaplan microscope equipped with a special filter for fluorescence (450-490 nm) and coupled to a Leica DC300F digital camera and FW4000 software. Observations and photography of selected slides were conducted using a Leica TCS SP2 confocal microscope of the MRI platform, Montpellier University.

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2.5. Quantification of the Na^+/K^+ -ATPase (NKA) immunostaining intensity in phyllopods

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To determine the relative importance of different phyllopods in ion transport in each salinity condition, we compared the relative fluorescence intensity of immunostaining in metepipodites of *Artemia* adults. Several photographs were taken in 5 individuals for each studied salinity using a constant exposure time. Longitudinal sections were used to visualize the different metepipodites and their Na⁺/K⁺-ATPase content within an animal.

The photographs were analyzed using the public domain ImageJ software (version 1.49, v), 225 according to a protocol successfully used in other crustaceans (Boudour-Boucheker et al., 226 2013; Gerber et al., 2016; Issartel et al., 2010) and in fish (Ouattara et al., 2009; Riou et al., 227 2012; Sucré et al., 2012). The intensity of immunostaining, i.e. the staining intensity of the 228 immunolabelled area, was measured by quantifying the pixel intensity of the immunolabeled 229 230 area using ImageJ software. These measurements were performed using the ImageJ software's "Average Gray Value" parameter, which determines the sum of the gray values of all pixels in 231 the selected area relative to the number of pixels, expressed in calibrated units (optical 232 density). In a second step, the ImageJ software was used to quantify the difference in average 233 pixel intensity of fluorescence between different salinities among metepipodites within the 234 same adults. 235

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2.6. Statistics

Statistical analyses (hemolymphe osmolality and fluorescence intensity) were performed
using Graphpad Prism (version 6, GraphPad Software Incorporated, La Jolla, CA 268, USA).
Normality and homogeneity of variance were respectively checked using D'Agostino-Pearson
test and Barlett's test. For data fitting homogeneity of variance requirement, a one-way

ANOVA regarding or regardless salinity as the main factor was performed; critical differences between groups were appraised using the Tukey's multiple comparisons test. For data not fitting homogeneity of variance and data due to the small sample size, a nonparametric KrusKal-Wallis test followed by a multiple comparisons Dunn's test was used. Data are presented as means \pm SD, and the level of statistical significance was set at p < 0.05.

- 248
- 249 **3. Results**
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251 The main results of this study are summarized in Fig. 1S.

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- 253 *3.1. Hemolymph osmolality*
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Artemia adults were extremely strong hypo-osmoregulators. Their hemolymph osmolality, which ranged between 406±21 and 608 ±16 mOsm.Kg⁻¹ in our experiment, was significantly different according to salinity (Fig. 2; one-way ANOVA, p<0.001). It is worth noting that hemolymph osmolalities differed by only 202 mOsm.Kg⁻¹ between the lower and higher tested salinities that themselves differed by more than 4450 mOsm.Kg⁻¹.

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261 *3.2. Morphological description*

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The morphology of *Artemia* adults is illustrated in Fig. 3. Eleven pairs of phyllopods (Ph1-11) were observed. The ten pairs of flattened leaf-like metepipodites are attached at the middle part of the ten phyllopods, (Me1-10). The eleventh pair of phyllopods is sexually dimorphic between the sexes and does not bear a metepipodite (Fig. 3A, 4A, B, C, 5A, B). Each metepipodite is a flattened leaf-like oval structure (Fig. 3A, C, 5G), with average minimum and maximum dimensions of 225±50 µm and 486±84 µm. Metepipodites 1 and 2 are smaller (Fig. 3A).

- Internally, we observed a longitudinal digestive tract, and a pair of maxillary glandslocated on each side of the digestive tract in the anterior part of the body (Fig. 6).
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273 *3.3. Histology and immunolocalization of Na⁺/K⁺-ATPase (NKA)*

Metepipodites are flat structures formed of two identical facing monolayered epithelia 275 covered by a thin cuticle, limiting a hemolymph lacuna (Fig. 5). A hemolymphatic space at 276 the base of each metepipodite allows a communication with the phyllopod (Fig. 5C, D). NKA 277 localization was visualized using immunohistochemical analysis of longitudinal and 278 279 transverse sections. An intense immunostaining throughout most of the cells was observed in the epithelium of the ten pairs of metepipodites (Figs 3B, C, 4C, 5). The immunopositive cells 280 were large (57±3 µm cellules) and contained a voluminous nucleus with average diameter of 281 10±1 µm (Fig. 5F, F'). Immunostaining intensity appears lower in some cells (Figs 3C, 5G). 282

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The semi-quantification of fluorescence intensity of metepipodites showed no significant difference between the different metepipodites and among the analyzed salinities and an overall high variability was observed between animals (Fig. 2S).

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A positive immunoreactivity for Na⁺/K⁺-ATPase was detected in the maxillary glands, where a strong immunostaining was observed in the epithelium of the efferent tubule and in the excretory canal (Fig. 6C-H). The proximal coelomic sac was not immunostained (Fig. 6C). The anterior digestive tract revealed a positive immunolabelling in the basal cell part as shown in Fig. 6D-F.

Negative control sections without the primary antibody did not show immunolabellingin any organ or tissue (not shown).

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297 **4. Discussion**

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299 Brine shrimps, Artemia species, are usually considered as extremophiles especially for 300 their ability to face challenging salinities (up to 10-fold that of standard seawater) considered 301 lethal for the majority of other aquatic species (Gajardo and Beardmore, 2012). Such 302 environmental pressures demand an ability to regulate hemolymph osmolytes according to the external medium via powerful osmoregulatory mechanisms (Campbell et al., 2012; 303 304 Charmantier et al., 2009; Freire et al., 2003). Blood osmolality change is often used as an indicator for osmoregulatory acclimation and tolerance to salinity stress (Varsamos et al., 305 306 2005). This study has confirmed the hypo-osmotic capacity of Artemia adults that can maintain their hemolymph osmolality up to 10 fold lower than their surrounding environment 307 (for example hemolymph osmolality of $608 \pm 16 \text{ mOsm.Kg}^{-1}$ in a medium at $6192 \text{ mOsm.Kg}^{-1}$ 308

¹, 212 ppt). Due to these exceptional physiological abilities, brine shrimps are considered the 309 most powerful hypo-osmoregulators among aquatic metazoans, with the highest tolerance to 310 salinity (Cole and Brown, 1967). Extensive similar studies have concluded that the 311 312 hemolymph osmotic pressure was markedly hypotonic even in the most concentrated media (Medwedewa, 1927; Plattner, 1955). Holliday et al. (1990) stated that A. salina is a weak 313 hyporegulator in 50 % SW and an increasingly strong hyporegulator in 100 %, 200 % and 400 314 % SW. Different studies have also shown a similarity between the mechanisms of 315 osmoregulation in Artemia adults and those proposed for marine teleosts and hypo-regulating 316 317 crustaceans (Croghan, 1958b, c, d; Copeland, 1967; Thuet et al., 1968; Smith, 1969a, b). Croghan (1958b, c, d), in an extensive study of osmoregulation in brine shrimps, has shown 318 that they have well-developed active mechanisms for absorbing NaCl from the gut lumen to 319 the hemolymph, resulting in a passive influx of water to hemolymph that compensates water 320 321 lost by osmosis to the external concentrated medium (Croghan, 1958d). Sodium and chloride, which also enter the body through diffusion given their high concentration in the external 322 323 medium, are excreted to the medium by specialized organs.

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325 Na⁺/K⁺-ATPase is known as the major driving force that ensures ion exchanges (Cieluch et al., 2004; Geering, 2008; Lignot et al., 1999; Lignot and Charmantier, 2001; Lucu and Towle, 326 327 2003; Thabet et al., 2016; Thuet et al., 1988) as well as generating an electrical gradient fuelling other transporters (Campbell et al., 2012; Esbaugh et al., 2019; Pivovarov et al., 328 329 2019; Sáez et al., 2009). Its abundance in an organ suggests an involvement in ion transport. The present study affords an overview of the different organs involved in osmoregulation in 330 331 A. salina adults. Immunofluorescence staining of Na⁺/K⁺-ATPase was used in order to 332 localize ionocytes in these organs.

333 Previous studies first addressed osmoregulation in embryos and during early post-334 embryonic development. Several extensive studies have reported that active ion excretion mediated by Na⁺/K⁺-ATPase is carried out by the naupliar salt gland (or dorsal / neck / nuchal 335 organ) that develops in late pre-naupliar embryonic stages; this organ becomes apparently 336 functional shortly before hatching. Later in development, thoracic epipodites take over ion 337 transport according to the stage of development (Conte, 1984; Conte et al., 1977, 1972; 338 Mitchell and Crews, 2002; Peterson et al., 1978; Russler and Mangos, 1978; Thuet, 1982). 339 Working in naupliar larvae, Sun et al. (1991) reported that Na⁺/K⁺-ATPase α and β subunits 340 were immunolocalized in the basal membranes of the salt gland cells. The mRNA expression 341

of α1 and α2 paralogs was quantified in the same organs (Conte, 2008; Escalante et al., 1995;
Sun et al., 1991).

In the present study, an intense Na⁺/K⁺-ATPase immunostaining throughout most of the 344 cells was observed in the epithelium of the ten pairs of metepipodites. Our results confirm 345 those found in earlier works that attributed active outward transport of ions to metepipodites 346 347 in adults (Copeland, 1967; Croghan, 1958c; Holliday et al., 1990). In a major study using electron microscopy, Copeland (1967) concluded that the metepipodites represent "a highly 348 specialized tissue for the secretion of salt and a special cell type found in these structures (the 349 350 'dark cell'), rich in mitochondria, is thought to be responsible for this ion transport". The numerous immunostained cells that we observed in metepipodites therefore correspond to the 351 "dark cells" as described by Copeland (1967). As we detected no difference in 352 immunostaining intensity among metepipodites, this suggests a similar cellular Na⁺/K⁺-353 ATPase content among different metepipodites and an involvement of all metepipodites in 354 active transport. Later experimental findings conducted on crude homogenates of 355 metepipodites revealed a very high specific enzyme activity of Na⁺/K⁺-ATPase, which 356 increased proportionally with the salinity of the external medium (Holliday et al., 1990). A 357 slight but not significant increase in immunostaining intensity has been observed between 60 358 359 and 139 ppt and could indicate a slightly increased Na⁺/K⁺-ATPase content within the cells lining phyllopods at 139 ppt. Recently, a preliminary report of Drenth (2017) confirmed the 360 presence of Na⁺/K⁺-ATPase in a metepipodite of A. franciscana and the salt extrusion occurs 361 in a mitochondrial rich, membraneous cell layer in the metepipodites of the brine shrimp. The 362 363 same investigator suggested that brine shrimp upregulate only the α^2 form of Na⁺/K⁺-ATPase in response to increasing salinity, while the $\alpha 1$ form remains relatively unchanged. 364

A positive immunoreactivity for Na⁺/K⁺-ATPase was also detected in the anterior digestive 365 tract and in the maxillary glands. In these paired glands, Na⁺/K⁺-ATPase immunostaining was 366 367 observed in the epithelium of the efferent tubule and of the excretory canal, while the coelomic sac remained unstained. Our results confirmed that these organs are also involved in 368 ion transport and probably for a part in osmoregulatory processes. In the same context, Tyson 369 (1969) had shown that the efferent duct of the maxillary gland of A. salina presents 370 ultrastructural features typical of transporting epithelia. Holliday et al. (1990) measured a 371 high Na⁺/K⁺-ATPase activity in the digestive tract. Studying the physiology of the gut in A. 372 373 salina and its implication in osmoregulation, Croghan (1958c) found that the concentration of both sodium and chloride ions in the gut fluids was always lower than that in the hemolymph, 374 pointing to an active uptake of NaCl across the gut epithelium, which in turn controls water 375

balance and prevents dehydration in hypertonic media. Indeed, Drenth (2017) confirmed the basolateral localization of Na⁺/K⁺-ATPase in the gut of *A. franciscana* adults, where ions and water are taken in from the environment into the hemolymph (Russler and Mangos, 1978).

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380 5. Conclusion

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In this work, we have focused on the hypo-osmoregulatory capacity of adult A.salina 382 and particularly on their main organs involved in active ion transport. We have confirmed that 383 384 brine shrimps are powerful hypo-osmoregulators as they keep their hemolymph osmolality strongly lower than even the most concentrated media. Immunofluorescence and whole 385 386 mounts staining of Na⁺/K⁺-ATPase have been used to illustrate the histological and cellular structure of metepipodites related to osmoregulation; despite their major function, their 387 388 functional histology had surprisingly not been re-addressed since the major works of the 1960's. We have also confirmed the involvement in osmoregulation of the maxillary glands 389 390 and of the anterior part of the digestive tract.

391 392

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- 397

398 List of References

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576	Legends of Figures
577	
578	Abbreviations. ADT: anterior digestive tract; Cs: coelomic sac; E: eye; EC: excretory canal; Me:
579	metepipodite; MG: maxillary gland; N: nucleus; Ph: phyllopod; PI: posterior intestine; T: tubule.
580	
581	Fig. 1. Geographical location of the Sebkha of Sidi El Hani (Tunisia). Google Earth Pro V. 7.3.2.5776.
582	
583	Fig. 2. Adult Artemia salina. Osmotic pressure of the hemolymph of animals maintained at three
584	different salinities (60 ppt: 1737 mOsm.Kg ⁻¹ , 139 ppt: 4064 mOsm.Kg ⁻¹ , 212 ppt: 6192 mOsm.Kg ⁻¹ ,
585	N=8-12). The dotted line is the isosmotic line. Different letters indicate significant differences
586	(p<0.001).
587	
588	Fig. 3. Adult Artemia salina. Morphology and localization of metepipodites (Me) and maxillary
589	glands (MG). (A) Dorsal view showing the eleven pairs of phyllopods (Ph) and the ten pairs of
590	metepipodites. (B), (C) Whole-mounts and immunolocalization of Na ⁺ /K ⁺ -ATPase; (B) Anterior view
591	with immunostaining in the two maxillary glands and in one metepipodite 1; (C) Median view with
592	several phyllopods and immunostained metepipodites. Scale bars: (A) 500 μ m; (B), (C) 200 μ m.
593	
594	Fig. 4. Adult Artemia salina. Localization of metepipodites (Me) on phyllopods (Ph). (A), (B)

Fig. 4. Adult *Artemia salina*. Localization of metepipodites (Me) on phyllopods (Ph). (A), (B)
Longitudinal vertical and horizontal histological sections. (C) Longitudinal horizontal section:

599 Fig. 5. Adult Artemia salina. Histological structure and immunolocalization of Na⁺/K⁺-ATPase (NKA)

- and DNA staining of nuclei on phyllopods and metepipodites sections. (A), (B), (F), (F') Merged
- staining of Na⁺/K⁺-ATPase (green) and DNA (blue). (C), (D), (E) Phyllopds bearing metepipodites
- 602 with NKA immunostaining (D, E) or Trichrome Masson's staining (C). (G) Immunolocalization of
- 603 NKA on metepipodites in horizontal (Me4) and transverse (Me5, 6) sections. Scale bars: (A), (B) 40
- 604 μ m; (C), (D), (E), (F), (G) 20 μ m; (F') 5 μ m.
- 605

Fig. 6. Adult *Artemia salina*. Histological structure, whole-mounts and immunolocalization of Na⁺/K⁺-ATPase (NKA) in the anterior digestive tract (ADT), in the maxillary gland (MG), and in metepipodite (Me). (A), (B) Transverse histological sections of the anterior digestive tract and of the maxillary gland. (C), (D), (E) NKA localization in transverse sections of the anterior digestive tract and of the maxillary gland. (F) Horizontal section of the anterior digestive tract and of the two maxillary glands. (G), (H) Whole mounts showing NKA localization in the maxillary gland and metepipodite 1. Scale bars: (A), (B), (C), (D), (F) 20 μ m; (E) 40 μ m; (G) 200 μ m; (H) 50 μ m.

- 613
- 614 Fig. 1S. Present results (in red) and previous knowledge on osmoregulation in the branchiopod
- 615 *Artemia* at different development stages involving different osmoregulatory organs.
- 616

Fig. 2S. Adult *Artemia salina*. Relative immunostaining intensity for Na⁺ /K⁺-ATPase (NKA) in the different metepipodites (1-10) and its difference according to different salinities. (A) Comparison between the metepipodites (1-10) regardless of salinity (N=5-6). (B) Comparison between the studied salinities regardless of the location of metepipodite (N=3-12).

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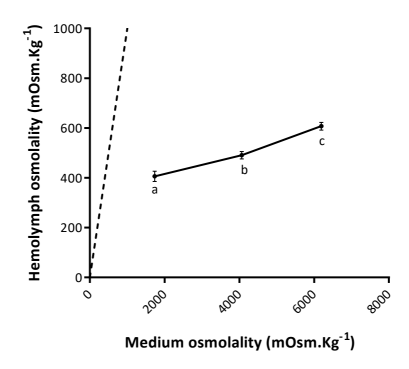
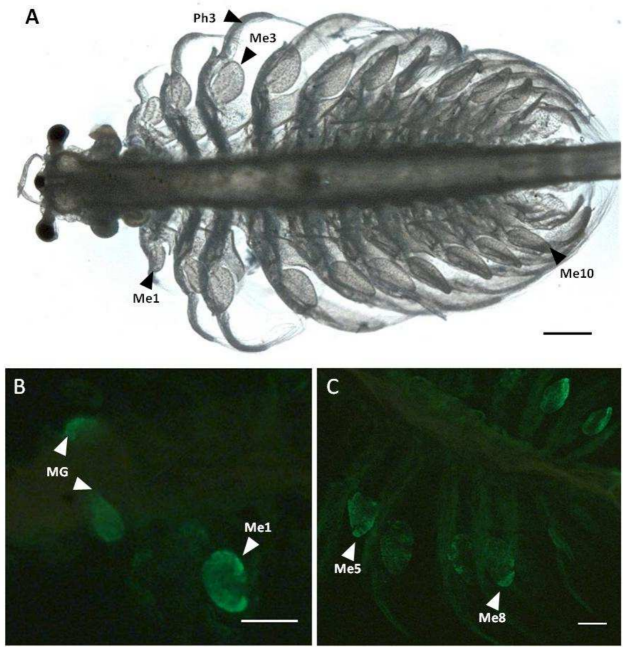


Fig. 2.







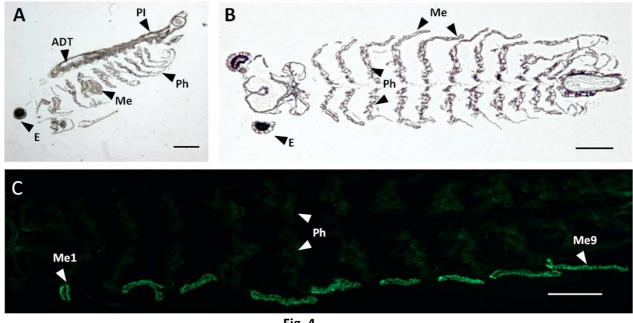
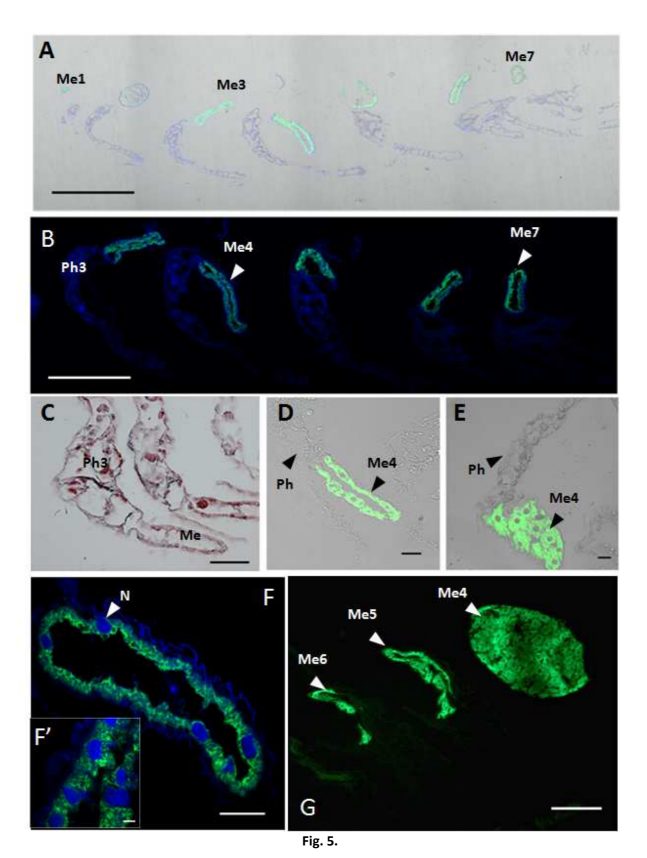


Fig. 4.



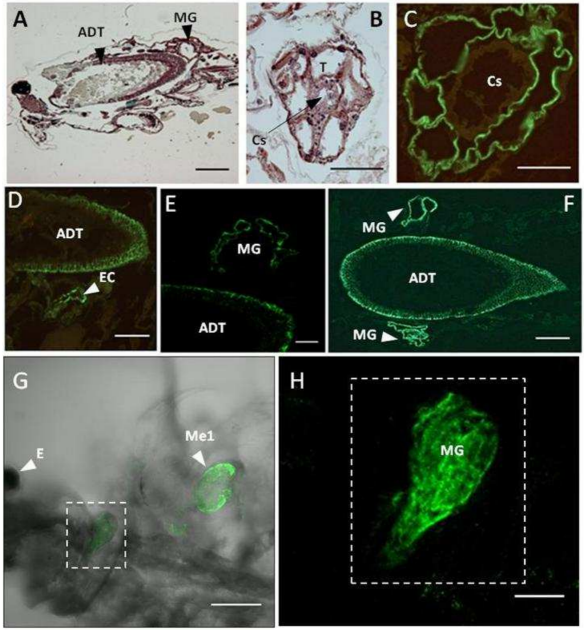
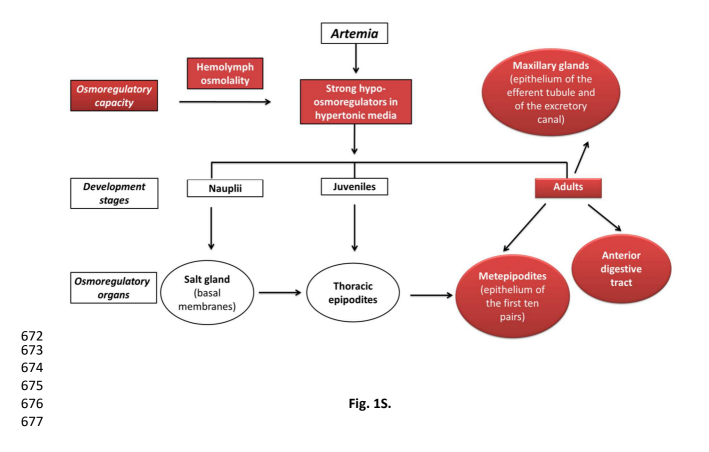
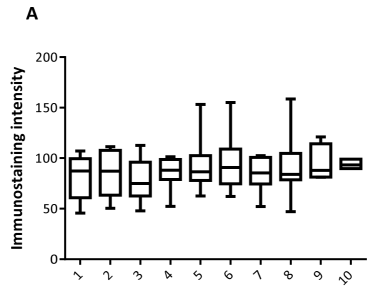




Fig. 6.





Metepipodite number

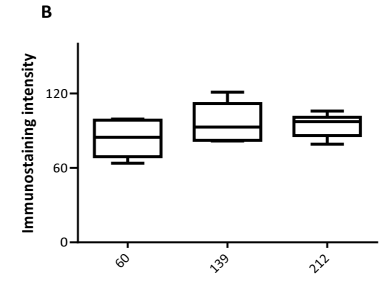




Fig. 2S.