RESEARCH ARTICLE

Ecological traits of planktonic viruses and prokaryotes along a full-salinity gradient

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Abstract

Virus–prokaryote interactions were investigated in four natural sites in Senegal (West Africa) covering a salinity gradient ranging from brackish (10‰) to near salt saturation (360‰). Both the viral and the prokaryote communities exhibited remarkable differences in their physiological, ecological and morphological traits along the gradient. Above 240‰ salinity, viral and prokaryotic abundance increased considerably with the emergence of (1) highly active square haloarchaea and of (2) viral particles with pleiomorphic morphologies (predominantly spindle, spherical and linear shaped). Viral life strategies also showed some salinity-driven dependence, switching from a prevalence of lytic to lysogenic modes of infection at the highest salinities. Interestingly, the fraction of lysogenized cells was positively correlated with the proportion of square cells. Overall, the extraordinary abundance of viruses in hypersaline systems (up to $6.8 \times 10^8$ virus-like particles per milliliter) appears to be partly explained by their high stability and specific ability to persist and proliferate in these apparently restrictive habitats.

Introduction

Planktonic viruses are of tremendous ecological relevance because of their lytic capacities within each of the three domains of life (Eubacteria, Archaea and Eukarya), their extraordinary abundance and diversity, and their effects on biogeochemical cycles in the hydrosphere (Suttle, 2005). Among the physical factors that influence their distribution directly or indirectly, salinity is assumed to be among the most crucial (Oren, 2009), alongside UV radiation (Wilhelm et al., 2003) and temperature (Bettarel et al., 2009). Salinity was initially recognized for its remarkable relationship with the diversity and functions of prokaryotes, which are the usual hosts of aquatic viruses (Bouvier & del Giorgio, 2002; Cissoko et al., 2008; Boetius & Joye, 2009). Bacteria and Archaea were indeed thought to be physiologically, genetically and ecologically dependent on external osmolarity (Lozupone & Knight, 2007). For example, marine prokaryotes have long been known to be remarkably halotolerant, which allows them to live in habitats with a wide range of osmolarities and ion compositions (Forsyth et al., 1971). Typically, prokaryotes respond to osmotic stress by adjusting their cell turgor (Oren, 2002; Shabala et al., 2009), either by increasing the synthesis of a variety of organic ‘compatible’ osmolytes or by controlling the fluxes of ions across cell membranes (Sleator & Hill, 2001). However, at salinity levels in excess of 250‰, the osmotic stress becomes so high that only a very limited number of prokaryotic species can thrive, including the hyperhalophilic bacterium *Salinibacter ruber* (Antón et al., 2002; Rosselló-Mora et al., 2008) and the square haloarchaea of Walsby (SHOW), well known for its singular postage stamp morphology (Walsby, 2005; Cuadros-Orellana et al., 2007). Such extremophile prokaryotes have attracted considerable interest in recent decades, and they have been identified in various types of hypersaline waters such as solar salterns (Antón et al., 2002) and soda lakes (Sorokin & Kuenen, 2005), located in different regions of the world including Australia (Burns et al., 2007), Israel (Oren, 2002), Spain (Guixa-Boixareu et al., 1996) and Peru (Maturrano et al., 2006). Numerous reports regarding their cultivation, genetic features and metabolic properties have provided new insights into life at the biological limits of salt tolerance (Bolhuis et al., 2006; Burns et al., 2007).
Although the impact of salinity on prokaryote ecology has been fairly well documented, only a handful of studies have explored whether this parameter is also a strong determinant of viral abundance and diversity distribution in water. From recent reports, we know that viral abundance can increase drastically as the salt concentration increases, reaching up to $10^9$ viruses per milliliter in systems where NaCl is close to saturation, such as crystallizer ponds (Guixa-Boixareu et al., 1996) and hypersaline lakes (Oren et al., 1997; Brum et al., 2005; Bettarel et al., 2010).

Conversely, the diversity of viruses together with that of their main prokaryotic hosts have been shown to decrease along increasing salinity gradients (Sandaa et al., 2003; Auguet et al., 2006). Currently, the reasons for the high densities of viruses in hypersaline waters are still poorly understood. The salt-forced extinction of potential virivorous nanoflagellates (Pedrós-Alió et al., 2000; Bettarel et al., 2005) is a possible explanation, but the ability of these viruses to better resist ambient UV radiations is not unrealistic either. Extreme halophilic Archaea and Bacteria could also be more susceptible to viral infection than their low-salinity counterparts. Furthermore, we still totally lack information on how viral life strategies may be influenced by salinity. The lytic, lysogenic and chronic cycles of infection are the main paths through which phages manipulate microbial processes, biogeochemical cycles and gene transfer (Weinbauer, 2004; Rohwer & Thurber, 2009). Because virus–prokaryote interactions are strongly dependent on the physiological state of the host (Maurice et al., 2010), we can anticipate that salinity is likely to exert a strong influence on the replication strategies used by prokaryotic viruses. However, it is still unclear whether hypersaline environments and their highly specific prokaryotic flora are more favorable to one or the other phage life cycles. These questions have never been addressed and they are fundamental to unveil the role played by salinity in structuring aquatic microbial food webs. Finally, previous studies on phage distribution along salinity gradients have been reported exclusively at temperate latitudes and mostly along restricted portions of the gradient. More systematic studies are thus required to elucidate host–virus interactions along full-salinity gradients, primarily in tropical ecosystems. Indeed, during the past two decades, biophysicochemical data provided by multidisciplinary studies have consistently revealed that aquatic ecology in temperate zones is somewhat different from that in tropical systems (Talling & Lemoalle, 1998). For instance, because of the typically high bacterial growth rates mostly resulting from the high temperatures encountered in tropical regions, trophic pathways within microbial food webs are not readily homologous in the two zones (Bouvy et al., 2004). The same conclusion is expected to be reached regarding virus-mediated processes.

In this study, we take advantage of four discontinuous aquatic sites located in Senegal, covering a full-salinity gradient from the same thalassohaline origin (i.e. the Senegalese coastal waters of the Atlantic Ocean), to examine virus–prokaryote interactions during the late dry season. By screening multiple physiological, ecological, morphological and phylogenetic traits of viral and prokaryotic communities, our main objective was to evaluate how these two communities can survive, interact and proliferate in water where the salt concentration increases up to saturation.

Materials and methods

Study sites

Samples were collected in Senegal (West Africa) from four aquatic sites contrasted in their salinity levels. These sites were sampled within an area encompassed between the Gambian border to the South and by the Senegal River to the north (Fig. 1). All the four sites share water from the same origin: the coastal Atlantic Ocean, and are therefore thalassohaline. A total of 14 sampling stations were identified within the four sites covering the full range of salinity from brackish (10‰) to near salt saturation (360‰).

The stations with salinity 10‰, 20‰ and 30‰ (named S.10, S.20 and S.30) were selected along the estuary of Senegal River (Fig. 1), which is among the longest rivers on the North West coast of Africa, running over 1800 km through four countries: Guinea, Mali, Mauritania and Senegal, where it flows into the Atlantic Ocean through a delta, which forms a complex canal system. Stations S.40, S.60, S.80, S.100 and S.120 were sampled in the Saloum River estuary, which has become an inverse estuary since the late 1960s with salinity that increases upstream and reaches 120‰ in some places. This unusual phenomenon is the result of the intrusion of coastal waters due to shallow estuarine slopes and a combination of insignificant freshwater flows (as a result of the Sahelian drought) and very high evaporation levels (more details in Diop et al., 1997). Stations S.140, S.190 and S.240 were investigated in artisanal saltern ponds located on the Saloum River bank, 3 km from the city of Kaolack. Finally, stations S.290, S.330 and S.360 were sampled in the hypersaline Lake Retba. This lake (surface area: ~400 ha) is located near Dakar (30 km), has shallow lagoon-like features with a maximum depth of 3 m and is located in close proximity (400 m) to the Atlantic Ocean. The high salinity of the Lake Retba (also known as the Pink Lake) results from severe evaporation that exceeded the input of water (from the sea) in a basin formerly connected to the sea (Garnier, 1978). The lake also receives some infiltration of freshwater, resulting locally in a steep salinity gradient ranging from c. 300‰ to salt saturation.

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at the end of the dry season. The geographical coordinates of the different stations are presented in Table 1.

**Sampling**

Samples were collected between 3 and 12 May 2007 (i.e. at the end of the dry season) from the subsurface waters (0.5 m) at the 14 stations. Water samples for nutrient and chlorophyll analysis, as well as for prokaryotic and viral enumeration, were collected using acid-cleaned sterile bottles. Samples intended for the determination of dissolved inorganic nutrients (NO$_3^-$-N, NH$_4^+$-N, PO$_4$-P) underwent preliminary filtration through Whatman GF/F fiberglass filters, stored at $-20^\circ$C and analyzed according to Strickland & Parson (1968). Chlorophyll a concentrations were determined fluorometrically following the filtration of samples through Whatman GF/F fiberglass filters, storage in liquid nitrogen and methanol extraction (Yentsch & Menzel, 1963). Analyses of nutrients and chlorophyll a concentrations were performed on a single sample.
Counts of microorganisms

For the prokaryotic and viral enumeration, aliquot samples of 1.5 mL were fixed with prefiltered (0.02 μm) buffered formaldehyde (2% final concentration), flash frozen in liquid nitrogen [as recommended by Patel et al. (2007)], stored at −20 °C and analyzed within a maximum of 4 weeks after collection. The abundances of planktonic prokaryotes and viruses were determined by standard techniques using SYBR Gold and epifluorescence microscopy (Patel et al., 2007). The number of virus-like particles and prokaryotes contained in triplicate samples of 50–300 μL were determined after particle retention of the particles on 0.02-μm pore-size membranes (Anodisc) and staining with SYBR Gold. On each slide, 300–600 prokaryotes and viruses were counted under an Olympus Provis-AX70 epifluorescence microscope with blue excitation, in 20 fields.

Water samples (100 mL) dedicated to heterotrophic nanoflagellates (HNF) enumeration were fixed with glutaraldehyde (1% final concentration) and stored at 4 °C. Subsamples of 15–20 mL of preserved water were stained with 4′,6-diamidino-2-phenylindole (DAPI) (final concentration, 15 μg mL⁻¹) for 15 min, filtered onto black Nucleopore filters (0.8 μm pore size), stored at −20 °C and counted under the epifluorescence microscope with UV excitation (Sherr et al., 1993). On each slide, 50–200 fields were observed according to cell densities.

Subsamples (50 mL) were fixed with a solution of formaldehyde and sodium borate (final concentration, 4%), and phytoplankton species were analyzed and counted using an inverted microscope Olympus IMT-2 (Utermöhl, 1958). Species were identified according to Bourrely (1990).

Viral morphology

The morphology of viruses was determined using transmission electron microscopy (TEM) after they were concentrated using the pegylation method (Colombet et al., 2007). Briefly, following successive prefiltration steps (0.2-μm pore-size filter at the end), viruses contained in 30-mL subsamples were reconstituted by polyethylene glycol (PEG) precipitation immediately after sampling. PEG 8000 (Sigma) was added to the water sample (final concentration, 10%) and incubated at 4 °C in the dark for 2 weeks. The white phase containing crystallized viruses was pipetted, centrifuged at 8000 g for 25 min at 4 °C and resuspended in 0.02 μm filtered water. KCl (1 M) was then added to this solution, incubated on ice for 20 min and centrifuged (12 000 g) for 10 min at 4 °C. The supernatant comprised of clean viruses was finally used for morphological diversity analysis by TEM. Viruses were collected onto a 400 mesh Cu electron microscopy grid supported with a carbon-coated Formvar film (Pelanne Instruments, Toulouse, France). The viral concentration was then centrifuged at 120 000 g for 2 h, 4 °C using an SW 40 Ti rotor (LE 80K, Beckman). Each grid was stained at room temperature for 30 s with uranyl acetate (2% w/v), rinsed twice with 0.02 μm filtered distilled water and dried on a filter paper. Grids were examined using a JEM 1200EX transmission electron microscope (JEOL) operated at 80 kV at a magnification of ×40 000–80 000. The different viral morphotypes were distinguished on the basis of the size and shape of the tail that typically makes it possible to identify phages belonging to the families of Myoviridae, Siphoviridae and Podoviridae. Tailess viruses were classified as icosahedral-, spherical-, filamentous- and lemon-shaped morphotypes.

Prokaryotic heterotrophic production (PHP)

PHP was estimated using the [³H]-thymidine incorporation method as developed by Fuhrman & Azam (1982) and modified by Bouvy et al. (2004) for tropical systems. For each sample, two 3 mL replicates and one formalin-fixed control were incubated with 100 μL of [³H]-thymidine (47 Ci mmol⁻¹, Amershams, UK) and kept in the dark at in situ temperature. The final total thymidine concentration was 20 nM. Incubations were stopped after 30 min by adding trichloroacetic acid (final concentration of 5%). Samples were precipitated on ice for 15 min and then filtered through cellulose nitrate filters (pore size, 0.2 μm, Whatman). The filters were then rinsed five times with 3 mL volumes of 5% trichloroacetic acid. The filters were placed in scintillation vials and solubilized with 0.5 mL of ethyl acetate. Six millilitters of a scintillation cocktail (Ready Save, Beckman) was added to each vial, and the radioactivity was measured using a liquid scintillation counter.

Catalyzed reporter deposition (CARD)-FISH analyses of phylogenetic diversity

Five milliliter samples were fixed with formaldehyde (2% final concentration), filtered on 0.2-μm polycarbonate filters (Whatman) and kept at −20 °C until hybridization. Several horsederish peroxidase probes (Biomers) were used, following Pernthaler et al. (2004): EUB 338III, ALF968, BET42a, GAM42a and ARCH915, targeting, respectively, the Eubacteria, the Alpha-, Beta- and Gammaproteobacteria (ALPHA, BETA, GAMMA), the Bacteroidetes and the Archaea domain. The NON338 probe was used as a negative control. After permeabilization with lysozyme (10 mg mL⁻¹, 1 h at 37 °C; Euromedex) and achromopeptidase (60 U mL⁻¹, 30 min at 37 °C; Sigma), seven filter sections, corresponding to the seven selected probes, were cut and hybridized for 2 h at 35 °C (Pernthaler et al., 2004). Probes BET42a and GAM42a were used with competitor oligonucleotides as described in Manz et al. (1992). Filter portions were then counterstained with DAPI (1 μg mL⁻¹; Euromedex) before enumeration with the epifluorescence microscope. A
minimum of 10 fields per filter-portion was counted. The selected groups were counted and expressed as a percentage relative to the DAPI-stained bacterial cells. The error associated with replicate CARD-FISH counts ranged from 5% to 25% (mean = 10.3%) based on a subset of three samples for which we conducted independent replicate CARD-FISH counts. The error on the percentage of bacteria was 15.3%, and corresponded to the sum of the errors for the bacterial group counts and for the total bacterial counts (average 5%), according to standard propagation of error. This error was consequently applied to all CARD-FISH analyses.

**CTC-positive (CTC+) cells**

The proportion of respiring bacteria that have high rates of metabolism was determined using 5-cyano-2,3-ditolyl tetrazolium chloride (CTC), an indicator of the respiratory electron transport system activity (Sherr et al., 1999). Active cells reduce the tetrazolium salt CTC to its fluorescent formazan form, and the cells with a high respiration rate or metabolism produce enough intracellular red fluorescence to allow detection and enumeration by flow cytometry (del Giorgio et al., 1997). A stock solution of 50 mmol L\(^{-1}\) CTC (tebu-bio SAS) was prepared daily, filtered through 0.1-μm filters and kept in the dark at 4 °C until use. CTC stock solution was then added to 0.45 mL of sample (5 mmol L\(^{-1}\) final CTC concentration) and incubated for 3 h at room temperature in the dark. At the end of the incubation, fluorescent 0.94-μm-diameter beads (Polyscience Inc.) were added as an internal standard before analysis on the cytometer. The red fluorescence of CTC (FL3) and the light scatter SSC were used to discriminate the CTC+ cells from other cells or weak fluorescent particles. The percentage of CTC+ cells, based on triplicate analyses, was calculated relative to the total bacterial counts obtained by epifluorescence microscopy.

**Viral infection of prokaryotes**

Prokaryotes contained in duplicate 8 mL aliquots of formalin-fixed samples were harvested by ultracentrifuging at 70 000 g for 20 min onto 400 mesh Cu grids, stained for 30 s with uranyl acetate (2% w/w) and examined at \(\times 40\ 000\) by TEM operated at 80 kV to distinguish between visibly infected and uninfected prokaryotes (Bettarel et al., 2004). At least 600 prokaryote cells were inspected per grid. Burst size (BS, viruses per bacteria) was estimated for every single sample as the average number of viral particles in all visibly infected prokaryotes totally filled with viruses. The fraction of lytically infected cells (FIC) was calculated from the frequency of visibly infected cells (FVIC) (as a percentage) using the formula: FIC = 7.11 × FVIC (Weinbauer et al., 2002).

**Fraction of lysogenic prokaryotes**

We used the method of Jiang & Paul (1996) to initiate prophage induction in prokaryotes. Mitomycin-C (1 μg mL\(^{-1}\) final concentration, Sigma Chemical Co., No. M-0503) was added to duplicate 10 mL volumes of water. Duplicate untreated samples served as the control. All samples were formalin fixed after being incubated for 12 h in the dark, at in situ temperature. Prophage induction was calculated as the difference in viral abundance between the mitomycin-C treated \((V_m)\) and control incubations \((V_c)\). The fraction of lysogenic prokaryote cells (FLC) was calculated as

\[
FLC(\%) = 100\left[\frac{(V_m - V_c)}{(BS \times PA_{10})}\right]
\]

where BS is the burst size (virus per bacteria) and PA\(_{10}\) the prokaryote abundance at the start of the experiment, i.e. before adding mitomycin-C (Weinbauer et al., 2003).

**Persistence of free viruses in water**

In 100-mL samples, free viruses were isolated from their hosts by successively filtrating the water through 5.0- and 0.2-μm polycarbonate membranes (47 mm in diameter) to remove prokaryotes and larger organisms. Each 0.2 μm filtrate was placed in a 100-mL polyethylene UV-permeable sterile Whirl-Pack\® bag and was incubated for 12 h in the dark. The incubation time of 12 h was chosen to limit the potential repopulation from the small fraction of prokaryotes that may have passed through the 0.2-μm membrane filters (this was checked by an epifluorescence microscopic prokaryote count). The survival rates of viruses (SR%) were calculated as follows

\[
SR(\%) = 1 - \left[\frac{\left(V_{I12h} - V_{T12h}\right)}{V_{I12h}}\right] \times 100
\]

where VIR is the viral abundance (expressed in \(10^7\) particles mL\(^{-1}\)).

**Statistical analyses**

Data were log transformed to satisfy the requirements of normality and homogeneity of variance necessary for parametric analyses. Simple relationships between original data sets were tested using Pearson correlation analysis. All statistical analyses were performed using SIGMASTAT software.

**Results**

**Abiotic parameters**

Nutrients exhibited different patterns throughout the gradient, reaching their maximum concentration at salinities 30‰, 190‰ and 360‰, for nitrite + nitrate, ammonium and phosphate, respectively (Table 1). Chlorophyll a concentrations were also highly variable, peaking at salinities 30‰, 190‰ and 290‰. Temperature ranged from 22.9 to 29.1 °C,
with the highest values recorded in the shallow crystallizer ponds of Kaolack (Table 1).

**Abundance of viruses, prokaryotes and HNF**

Viral abundance increased up to 50-fold between both the extremities of the gradient (Fig. 2a). Although the concentrations were relatively stable between 10‰ and 240‰ of salinity, a sharp increase was observed at the top end of the gradient, with concentrations reaching up to $6.8 \times 10^7$ viruses mL$^{-1}$ at 360‰ (Fig. 2a). A similar pattern was observed for prokaryotes, which increased regularly from salinity 240‰ to attain $3.4 \times 10^8$ cells mL$^{-1}$ at 360‰ (Fig. 2b). The virus and prokaryote counts were highly significantly correlated ($r = 0.99$, $P < 0.001$, Table 2). Conversely, the abundance of HNF declined steadily from 564 to 7 cells mL$^{-1}$, as the salinity increased from 10‰ to 360‰ (Fig. 2c). Phytoplankton abundance increased steadily from 1.5 to $9.8 \times 10^7$ cells L$^{-1}$ between salinity 10‰ and 140‰ where picophytoplankton species were dominant (data not shown). Algal abundance showed marked variations throughout the end of the gradient, with values reaching $1.9 \times 10^6$ cells L$^{-1}$ in water approaching salt saturation (Fig. 2d). From salinity 240–360‰, the phytoplankton assemblage was mainly comprised of the green algae *Dunaliella salina* (18–99% of the total abundance) and of *Cyanophyceae* (data not shown).

**Prokaryote activity and physiological state**

The PHP and the percentage of CTC+ cells followed the same pattern characterized by a general decrease in the values between salinities 10–20‰ and 240‰, followed by a moderate increase from 240‰ to 340‰ (Fig. 3). The proportion of CTC+ cells was between 1.0% and 32.5%; the highest percentage was recorded at salinity 20‰ in the Senegal River, and then CTC+ cells declined steadily down to 1.0% at salinity 120‰. Over the second part of the salinity gradient (120–360‰), CTC+ then increased again to attain 15.9% in Lake Retba. PHP followed the same trend; however, its decrease continued up to salinity 240‰ and then increased again up to 360‰ (Fig. 3).

**Prokaryote community composition and distribution**

The proportion of Eubacteria ranged between 15.9% and 92.4% of the total prokaryotic assemblage. Eubacteria sharply declined at salinity 140‰, and then increased throughout the end of the gradient (Fig. 4a). Among the Eubacteria, there was a change in the phylogenetic dominance of the Alphaproteobacteria, Gammaproteobacteria and Bacteroidetes at salinities 30‰, 80‰ and 330‰, respectively (Fig. 4b). Archaea represented from 0.4% to 57% of the total cell counts. Their proportion was marginal up to salinity 110‰, and then increased continuously with increasing salinity up to near-saturation levels (360‰). Square-shaped cells emerged at salinity 190‰, where they accounted for up to 58% of the prokaryotes. From there on, along the gradient, these atypical cells constituted from 28% to 40% of the prokaryotes (Fig. 4c). The distribution of square-shaped cells was closely correlated with that of Archaea ($r = 0.87$, $P < 0.05$) and with salinity levels ($r = 0.79$, $P < 0.05$) (Table 2). Total percentages of Eubacteria and Archaea higher than 100% were due to fluorescent hybridization count errors. On one single occasion (S.190), the proportion of SHOW cells was higher than that of the archaean cells detected using the ARCH915 probe. However, this probe has been shown to efficiently hybridize square cells inhabiting crystallizer ponds (Antón et al., 1999), dismissing possible specificity problems. This discrepancy is thus more likely caused by the presence of organic aggregates in that station, which may have covered and hidden a significant fraction of the hybridized cells.

**Viral life strategies**

Between salinity levels of 10‰ and 140‰, the FIC oscillated from 0.8% to 20.1%, peaking twice at salinities 10‰ and 120‰, respectively (Fig. 5a). Then, in the terminal part of the salinity gradient (i.e. from 190‰ to 360‰), no visibly infected cells were detected among the 9453 prokaryotes that...
Table 2. Correlation relationship of basic parameters estimated in the surface water of the 14 study stations

<table>
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<th>Salinity</th>
<th>VIR</th>
<th>PROK</th>
<th>HNF</th>
<th>CHLORO</th>
<th>SR</th>
<th>PO</th>
<th>FIC</th>
<th>FLC</th>
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Viral persistence in water

The survival of free viruses after 12 h slowly decreased between the salinities 10% and 140%, with survival rates ranging from 53.4% to 30.2% (Fig. 6b). Viral persistence then increased steadily up to 97.5% at the highest salinities in Lake Retba. Positive and significant correlations were found between viral persistence and (1) salinity, (2) the proportion of square cells and (3) the proportion of Archaea (Table 2).

Viral morphological diversity

Although tailless icosahedral viruses declined between 10% and 240% of salinity, they remained highly dominant, accounting for 58–86% of virioplankton standing stocks (Fig. 6). At salinities > 240%, the incidence of these viruses declined sharply and finally disappeared at near salt saturation, in Lake Retba. The same situation was observed for Myoviridae, Siphoviridae and Podoviridae, whose presence was quasi negligible between salinities 240% and 360%. Conversely, spindle-shaped viruses emerged in moderate amounts at salinity 240% (mean = 3%), but were dominant along the end of the salinity gradient, in Lake Retba (Fig. 6). In this hypersaline lake, other viral morphotypes typical of archaean viruses were also detected at high levels of abundance: filamentous and spherical viruses that accounted for up to 20% and 33%, respectively (Figs 6 and 7).

Discussion

Ecological traits of prokaryotes along the salinity gradient

As reported frequently in the literature, the abundance of the prokaryotes gained two orders of magnitude from freshwater to hypersaline stations, reaching up to $3.4 \times 10^8$ cell mL$^{-1}$, which is roughly 100-fold higher than what is generally found in marine environments (Suttle, 2005). In this study, while prokaryotic abundance increased steadily up to salinity 240%, conversely, their physiological state, as reflected by the proportion of CTC+ cells, and the PHP both decreased. We can surmise that the osmotic stress exerted on prokaryotes may directly limit their metabolic
activity. Halophilic prokaryotes can avoid water loss and adjust their turgor pressure in a highly saline environment by accumulating osmotically active compounds such as glycine betaine, glutamate, glutamine and proline (Sleator & Hill, 2001; Burkhardt et al., 2009). When salt stress increases, bacteria may be forced to use energy-consuming mechanisms to maintain the osmotic balance in their cells instead of being able to use this energy for growth. On the basis of our observations, this scenario might be accurate up to salinity 250‰, but beyond this, the tremendous abundance of highly active prokaryotes in the Lake Retba may be attributable to high selection pressure, leading to the development of a unique assemblage of highly adapted, fast-growing and active cells, as reported previously by Guixà-Boixareu et al. (1996). It is also possible that most of the bacterial populations that survived at salinity 200‰ were still somewhat reminiscent of those found in coastal marine waters (Pedrós-Alió et al., 2000), but that only the halophilic species remained active. Although salinity 250‰ seems to correspond to the threshold beyond which both prokaryote abundance and metabolism intensify, we observed that marked phylogenetic changes occurred earlier in the gradient, i.e. at salinity 150‰ (Fig. 4a and b). Between salinities of 0‰ and 150‰, the prokaryote assemblage was dominated by Eubacteria (mostly Alphaproteobacteria at salinity 30‰, then Gammaproteobacteria at 80‰), and Archaea

**Fig. 3.** Dynamics of PHP and CTC+ cells, along the salinity gradient.

**Fig. 4.** Distribution of the Eubacteria and Archaea along the salinity gradient (a). Phylogenetic distribution among the Eubacteria of the Bacteroidetes and the Alpha-, Beta- and Gammaproteobacteria along the salinity gradient (bars). Pies indicate the representative eubacterial community composition at salinities 30‰, 80‰ and 330‰ (b). Micrograph and contribution of the SHOW to the prokaryote assemblage (c). Scale bar = 500 nm.
were relatively rare (< 10% of DAPI counts). Above 150‰ of salinity, the proportion of Archaea increased continuously up to the top of the gradient, where they constituted up to 60% of the prokaryote assemblage, along with the emergence and proliferation of cells with an atypical square shape (Fig. 4c). In two different studies conducted in crystallizer ponds, Antón et al. (1999), and then Casamayor et al. (2000) also reported that Archaea can account for 467% of the total DAPI counts where salinity reaches saturation. Cells belonging to the Bacteroidetes exhibited a similar pattern characterized by a marked increase at the most salinized sites, constituting up to 28% of the total DAPI counts in the Lake Retba (Fig. 4b). Previous studies have reported that microbial communities in hypersaline environments have a low diversity (Benlloch et al., 2002; Maturrano et al., 2006), and are typically dominated by a very small number of prokaryote species while maintaining a large biomass: the extreme halophilic rod-shaped bacterium belonging to the Bacteroidetes group, S. ruber (Antón et al., 2002; Rosselló-Mora et al., 2008; Sime-Ngando et al., 2011), and the square red halophilic archaea, recently isolated and described as SHOW (Walsby, 2005; Burns et al., 2007; Cuadros-Orellana et al., 2007). In the tropical sites studied, we found strong evidences that the bacterial community were also comprised of these dominant prokaryotes. Indeed, the strong correlation between the abundance of square cells and the proportion of Archaea seemed to indicate the presence of SHOW. From a culture-independent analysis of the microbial diversity in a sample collected in Lake Retba in April 2008, few of the 16S sequences corresponded to known archaeal general (Halococcus, Halorubrum and Natronomonas), whereas the majority represented novel archael clades. The cells in our samples resembled thin, square or rectangular sheets with sharp corners, measuring 1.5–3.0 μm across and contained numerous gas vesicles (see picture in Fig. 4c), which are typical characteristics of the Halobacteriaceae (Walsby, 2005).

Viral dynamics along the salinity gradient
Viral abundance was tightly linked to that of prokaryotes along the salinity gradient with the concentration reaching
up to $6.8 \times 10^8$ particles mL$^{-1}$ at salinity 360‰. Such abundances come as no surprise because hypersaline environments have been well described to harbor huge numbers of viruses, as high as 2 billion particles per milliliter of water (Guixa-Boixareu et al., 1996; Oren et al., 1997; Bettarel et al., 2006; Schapira et al., 2009). Intuitively, such high numbers of halophages might be explained by the high abundance of their prokaryotic hosts, which in turn might be due to the very low abundance of their nanoflagellate predators. This scenario is supported by a previous study where bacteriophages could not be detected at salinities $> 200$‰ (Pedrós-Alió et al., 2000). Another potential explanation is that halophages might be more resistant to ambient virucidal agents than their freshwater or marine counterparts. There is some confirmation of this hypothesis in our experiment, where we found that the survival rates of free viruses (i.e. without their hosts) increased significantly from salinity 200‰ and peaked at 340‰. The reasons for such better survival rates in hypersaline habitats remain unclear. We know that viruses are much more resistant to changes in ionic strength conditions than their host cells (Kukkaro & Bamford, 2009), but we can also envisage that viruses inhabiting seemingly inhospitable niches may be genetically distinct, and specifically niche adapted as suggested by Prangishvili et al. (2006) and Le Romancer et al. (2007). In such high-salinity habitats where potentially virivorous HNF or ciliates cannot live, viruses might also be preserved from predation and this may also contribute to their high abundance. Furthermore, the unicellular Dunaliella salina was one of the last Eukarya capable of surviving and proliferating at salinities higher than 240‰. Dunaliella is known to produce massive amounts of glycerol to ensure osmotic stabilization of the cytoplasm, and this compound is often postulated to be the main source of organic carbon for the heterotrophic prokaryotes in hypersaline ecosystems (Elevi Bardavid et al., 2008). Questionably, glycerol, as demonstrated by Peak & Peak (1980), might also represent a protective substance for viral capsids against virucidal agents such as UV radiation, but also presumably from the ambient proteases or the degradative effects of temperature.

The ‘infectivity’ paradox in hypersaline habitats

Certainly, the most intriguing finding in this study is the paradox of high viral concentrations, but low FVIC in hypersaline environments. Indeed, we observed that progressively fewer prokaryotes were visibly infected along the gradient, and infected cells even became undetectable beyond salinity 170‰. From a study conducted in a salt pond in Jamaica, Daniels & Wais (1998) reported that phages of low virulence predominated in culturable populations. The same authors also demonstrated that lytic phages infecting extreme halophilic prokaryotes in hypersaline lagoons do not produce active infections at saturating concentrations of NaCl (Daniels & Wais, 1980). By contrast, a study conducted in solar salterns located in Spain reported that between 1% and 10% of the square cells were filled with lemon-shaped phage particles (Guixa-Boixareu et al., 1996). Therefore, we cannot exclude the possibility that the latent period (during which mature viruses are not yet visible within the host cell) of tropical halophages could be much longer than that is generally reported in the literature for temperate marine or freshwater viruses (Weinbauer et al., 2002). This might partly explain why observations of infected cells in water were so rare in the tropical hypersaline stations, but this has to be further explored.

According to the ‘kill-the-winner’ hypothesis (Winter et al., 2010), the typically low-diversity prokaryote assemblage in hypersaline habitats (Benlloch et al., 2002) provides a virtually ideal environment for large viral populations (Pedrós-Alió et al., 2000). On the other hand, if the ‘kill-the-winner’ hypothesis is true, then this dense phage population could rapidly induce dramatic effects in this high-biomass prokaryote community. Guixa-Boixareu et al. (1996) have suggested that the occurrence of such abundant pathogens is inconceivable without ‘an evasion strategy that promotes variations at the phage attachment sites’. Daniels & Wais (1998) also suggested that the low virulence of halophages probably arises as a result of slow adsorption processes occurring in hypersaline environment. Interestingly, there have been several reports of high salt concentrations inhibiting the binding of viruses to their hosts (Kukkaro & Bamford, 2009) and this is certainly a promising avenue for further researches to unravel halovirus–haloarchaeon interactions in nature.

The addition of mitomycin-C as a prophage inductor revealed that the percentage of cells at the lysogenic stage of infection was strongly and positively correlated with the proportion of the square cells. Although we know that few of the haloviruses identified have been reported to be strictly lytic (Porter et al., 2007), this study is among the first to report lysogenic-oriented life strategies in haloarchaeophages along salinity gradients. This seems to indicate, as suggested previously by Porter et al. (2007), that a substantial fraction of the haloarchaeophages may be temperate rather than virulent in habitats where the salinity exceeds 250‰. This is in support of Daniels & Wais (1998) hypothesis that halophages have probably evolved to exert minimal selective pressure on sensitive hosts, by producing active infections mainly when the host is likely to be destroyed by changing environmental conditions. Our samples were collected in late April, at the end of the dry season after 6 months of stable climatic conditions characterized by a total absence of rain and a temperature of between 28 and 34°C (data not shown). Such stable conditions may thus partly explain why lysogens were detected in Lake Retba before the
occurrence of any natural induction event such as the arrival of the rainy season and the subsequent decline in salinity. Interestingly, phages infecting hyperthermophilic archaea have also been recognized for their lysogenic rather than lytic lifestyle (Prangishvili et al., 2006).

Moreover, during halovirus infection, archaeons that have a distinct thin, glycoprotein surface layer protecting the cell membrane (unlike Bacteria with their rigid peptidoglycan cell walls) have been shown to release viruses continuously without cell lysis, in a manner similar to the situation in chronic infection (Prangishvili et al., 2006; Porter et al., 2007). This strategy could also allow haloarchaea to continue to divide, thus maintaining high abundances of prokaryotes along with their viral pathogens, as hypothesized previously by Daniels & Wais (1980). Therefore, the nearly 1 billion viral particles per milliliter recorded at salinity higher than 300‰ may not result from the release of lytic viruses, but rather from the chronic-like interactions or from sporadic prophage induction, likely coupled with good conditions for viral preservation.

The morphology of planktonic viruses also showed strong salinity-driven dependence with the emergence, > 150‰, of spindle shaped, linear or spherical morphotypes. Although some morphologies found in the Lake Retba were similar to known viruses of hypersaline environment (Dyall-Smith et al., 2003; Santos et al., 2007) or hyperthermophilic archaea (Prangishvili et al., 2006), an unexpected morphological diversity comprising uncommon viruses that did not resemble any known group of virus (hairpin-shaped or bacilliform particles for example) was revealed from Lake Retba that appears to exceed that observed in other saline ecosystems (Sime-Ngando et al., 2011). Currently, about 17 haloarchaegal viruses have been isolated, with the great majority resembling tailed dsDNA bacteriophages, such as λ phases (Porter et al., 2007).

Finally, the constraining pressure that is typically exerted by salt on life forms was successfully faced by prokaryotes and their viral parasites. Both communities displayed remarkable changes in their ecological traits along the salinity gradient and became highly abundant at or near salt saturation. Further studies are now necessary to understand how the coevolution of both communities in hypersaline environments has led to this peculiar type of relationship, seemingly favoring both viral and prokaryotic proliferation.

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