AS WE SEE IT

Isolates as models to study bacterial ecophysiology and biogeochemistry

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ABSTRACT: Here, we examine the use of bacterial isolates growing in artificial media or seawater as a means to investigate bacterial activity in the upper ocean. The discovery of a major role of bacteria in the ocean’s carbon cycle owes greatly to the development of culture-independent assemblage-level approaches; however, this should not detract from the recognition of model isolates as representing the environmental microbiome. A long-established tool for culturing bacteria, in medicine and general microbiology, has been agar plates. In addition, a great variety of liquid substrates including seawater have been used to successfully identify and cultivate important bacteria such as Pelagibacter ubique. Yet, the discrepancy between microscopic counts and plate counts, the great plate count anomaly, has led to a biased perception of the limited relevance of isolated bacteria. Linking isolates to whole-genome sequencing, phylogenetic analysis and computational modeling will result in culturable model bacteria from different habitats. Our main message is that bacterial ecophysiology, particularly growth rates in seawater, and functionalities inferred through the identity, abundance and expression of specific genes could be mechanistically linked if more work is done to isolate, culture and study bacteria in pure cultures. When we rally behind a strategy aimed at culturing targeted phenotypes, we are not saying that culture-independent studies of bacteria in the sea are not informative. We are suggesting that culture-based studies can help integrate the ecological and genomic views.

KEY WORDS: Bacteria · Culture · Physiology

INTRODUCTION AND HISTORICAL BACKGROUND

Marine microbiology has undergone major advances during the past 4 decades due to progress in techniques to observe and culture bacteria and further through the application of molecular methods (Temperton & Giovannoni 2012). However, fundamental questions on in situ growth physiology and viability remain unresolved in part because of a lack of consensus towards a conceptual framework on bacterial life strategies and in part because of a lack of suitable model organisms to study environmental bacterial physiology.

Ocean surface water as a substrate for bacteria

In retrospect, it is remarkable that until the end of the 1970s, it was thought that bacteria in the ocean could not grow in unenriched seawater (Jannasch 1979). This view was challenged through direct observation and laboratory studies.
Bacterial growth in seawater

SWC became an important model system for the study of bacterial growth in the pelagic ocean, for instance in efforts to determine bacterial growth rate in situ. Both methods using radioactive precursors to DNA and protein as well as growth cycle indicators such as frequency of dividing cells were calibrated using SWCs (Hagström et al. 1979, Fuhrman & Azam 1980, Kirchman et al. 1986). These new techniques to measure growth rate subsequently spurred extensive fieldwork to determine bacterial biomass turnover in different aquatic environments but with a somewhat disappointing outcome. It soon became clear that while separate estimates of growth rate, cell numbers and cell mass combined would provide the most accurate measure of bacterial production, integrated measures of production became the practical solution (Ducklow et al. 1986, Kirchman et al. 1986). Thus, the actual number of measurements of growth rate in situ for bacteria is few in the marine or other aquatic environments. In the following sections, we provide examples of growth rates for different bacterial isolates in artificial media or seawater. In particular, we examine different strategies of bacteria to proliferate in seawater and the effect of temperature as a major regulator of growth rate, as suggested from the ‘Tara’ Oceans expedition (Sunagawa et al. 2015). We finally discuss the current knowledge on how the diversity of isolated marine bacteria compares to the genomic diversity in the wild, i.e. natural microbial assemblages.

Bacterial culturability

Let us make some generalizations and examine a small volume of surface seawater. The sample we imagine we are looking at is fixed in formaldehyde, collected on a polycarbonate filter, stained with for example DAPI or SYBR Green® and viewed under an epifluorescence microscope. We will assume the

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SETTING THE SCENE

A long-standing issue in marine microbiology relates to the ability of bacteria in nature to grow in laboratory pure cultures. Marine microbiologists until recently had assumed that 99.9% of the marine bacteria seen under the microscope are unculturable. The basis for this assumption is the low number of bacteria that are able to form colonies on a plate. This ignores the fact that some strains can form biofilm/colonies or only grow as single cells (Simu & Hagström 2004). Also, in situ physiological conditions of bacteria such as phage infection and possible resting stages might render some bacteria unculturable. Addressing the problem of low culturability (typically 0.1 %) requires a basic understanding of the in situ physiological state of bacteria and their survival strategies.
field of view contains 100 bacteria-shaped objects. Further, we only consider chemoorganotrophic bacteria including methylotrophs and ask: What are these? What is their physiological state? What kind of interactions are they involved in? To keep track of the different life strategies, each strategy will be given a type number and a name. The names given to the numbered bacterial types/life strategies are added solely to help the reader by suggesting a stereotype. We are not trying to introduce a new classification. Instead, in the absence of valid phenotypic characterizations, we present nicknames that reflect the behavior of the described bacteria. After all, as more phenotypic characterization is done, we will eventually end up with a rainbow or a continuum of the expression of ecologically relevant phenotypes in varied environmental contexts.

**Solitary bacteria**

A large proportion of bacteria in the ocean are true oligotrophs, perhaps >30%, judging from counts based on genetic markers (Morris et al. 2002). They operate as single cells. These Type 1 ‘crumb picker’ bacteria occupy habitats that support slow growth, and they tend to be small in size. Their modest appearance does not mean they are unimportant, since for example the most common bacterium on earth, Pelagibacter ubicuitus (Morris et al. 2002, Rappé et al. 2002). However, they do not form biofilm or colonies, and therefore they have remained uncultured for a long time (Simu & Hagström 2004). These true oligotrophic bacteria have evolved an optimal access to the substrate in a low-nutrient environment, presumably due to the presence of high-affinity, low-capacity substrate transport systems. These seemingly free-living bacteria represent a relatively new concept that is currently being integrated in the ecological narrative of the ocean (Table 1). Under oligotrophic conditions, a premium is presumably placed on the ability to successfully compete for resources with other bacteria as well as other members of the same species, and thus colony formation should be non-adaptive (Simu & Hagström 2004).

In rich microenvironments, some bacteria can accomplish rapid growth as a result of behavioral and physiological adaptations to enhance their foraging success. These Type 2 ‘stalker’ bacteria show growth characteristics close to the opportunists, optimally positioning themselves in relation to a substrate gradient and thus achieving rapid growth (Blackburn et al. 1998). Examples of the Type 2 strategy are seen in bacterial cells hovering around algal cells exuding organic matter (Stock et al. 2008). In the case of a common diatom and a Sulfitobacter strain, metabolite and transcriptome analyses demonstrate that the bacterium promotes cell division of the diatom (Amin et al. 2015).

The frequent presence in marine metagenomes of bacterial genes coding for secretion mechanisms similar to those found in pathogenicity islands of animal and plant pathogens (Persson et al. 2009) suggests a life strategy expressed by Type 3 ‘killer’ bacteria. The action of such Type 3 bacteria attacking and killing ageing algal cells towards the end of

<table>
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<tr>
<th>Year</th>
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<tr>
<td>1979</td>
<td>Kuznetsov et al. (1979)</td>
<td>Defined oligotrophic bacteria as those that grow in media with a content of organic matter of 1 to 15 mg C l⁻¹.</td>
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<tr>
<td>1980</td>
<td>Ishida et al. (1986)</td>
<td>Distinguished obligate and facultative oligotrophs.</td>
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<tr>
<td>1984</td>
<td>Martin &amp; Mcleod (1984)</td>
<td>Questioned the existence of 2 classes of bacteria based on their ability to grow on high versus low concentrations of organic matter.</td>
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<tr>
<td>2004</td>
<td>Cho &amp; Giovannoni (2004)</td>
<td>Reported that uncultured and oligotrophic bacteria are the same.</td>
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<tr>
<td>2004</td>
<td>Simu &amp; Hagström (2004)</td>
<td>Suggested that oligotrophic bacteria are characterized in terms of behavior and choice of life strategy, including nutrient preference.</td>
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<tr>
<td>2013</td>
<td>Carini et al. (2013)</td>
<td>Demonstrated a defined medium for growth of <em>Pelagibacter ubique</em>.</td>
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blooms has been recorded repeatedly, as summarized in a review on algicidal bacteria in the sea by Mayali & Azam (2004).

**Aggregating opportunists**

A small fraction of the bacteria are opportunistic, expressing strategies that render them culturable on rich media. Such Type 4 ‘fast handler’ bacteria are found in a highly variable fraction (0.2 to 20%) that nevertheless can respond quickly to organic input, e.g. attach to a colloid or a particle (Pinhassi et al. 1997). They grow rapidly to form microcolonies (cell aggregates) or biofilms, which means they have the ability to form colonies on agar plates. These bacteria are able to exploit organic matter in both dissolved and particulate form. Being able to react quickly and to outgrow the competition, they can form blooms in the wake of trophic events (Rehnstam et al. 1993). Thus, at times they are found in high abundance but relatively quickly return to low numbers (Fuhrman & Hagström 2008). Usually, such bacteria can be grown on agar plates and in rich liquid cultures; because of the ease with which they can be isolated, these are the marine isolates most commonly deposited into culture collections. Surprisingly, there seems to exist a remarkable redundancy in substrate use among these bacteria, although taxonomic analysis shows considerable diversity among the opportunistic bacteria of the upper ocean (Pommier et al. 2007). No doubt, conditions do change over time and space, but on a given occasion removal of 1 or several bacterial types does not seem to reduce the ability to oxidize the organic matter present in the water (Sjöstedt et al. 2012). This seems to suggest that to consume the brunt of the DOC, no cooperation among these different bacterial strains seems essential, and a single strain might do the job equally well (Pedler et al. 2014).

While colony formation may seem like a simple and primary trait of bacteria, this is not the case (Davies et al. 1998, Li et al. 2002). The formation of colonies or biofilm requires an active process through quorum sensing. The flagella motor has to be shut down for the cells to aggregate, optimal cell-to-cell contact has to be established to allow good cellular communication and the production of extracellular enzymes must be directed towards the joint goal of the colony to exploit the organic substrate beneath the culture. Type 5, the ‘fast builder’ bacteria, have adopted a swarming behavior that can be seen as a special case of colony formation, but since the lateral invasion of the substrate surface is an actively coordinated process that requires quorum sensing, this behavior represents a life strategy of its own (Lindum et al. 1998).

**Dead or resting bacteria**

A confusing component in the microbial scenario is the significant fraction, often >30%, of most likely dead or inactive bacterial cells. These cells are not part of any specific life strategy but can cause conflicting results in various kinds of measurements. Under the light microscope, dead cells are indistinguishable from living cells but may consist of virus-infected, moribund cells or cell capsule remains (‘ghosts’). The ghosts are distinguishable from live cells since the ghosts disappear when counter-stained (Zweifel & Hagström 1995). Bacterial cells infected with lytic virus will fail to divide, yet they are biochemically active, and this may create results that are difficult to interpret. One example is the question of how to judge low-activity uptake of nucleotide precursors that can also be attributed to a resting state (Samo et al. 2014). The initial observation of bacterial ghosts has been investigated and questioned by several authors (Heissenberger et al. 1996, Luna et al. 2002, Jones & Lennon 2010), and a reliable protocol to determine the fraction of live and functional cells is still lacking. Because of the renewed interest in phage therapy instead of treatments using antibiotics, the molecular tools used by phages to open the bacterial cell now start to shed light on the mechanisms involved in ghost formation (Haidinger et al. 2003, Letchumanan et al. 2016). These studies may help distinguish between phage-lysed ghost cells and intact cells including the actively growing cells in situ.

**ISOLATE IDENTITY AND THE SPECIES CONCEPT**

In the previous 2 sections, we have aimed to establish an overall setting for bacterial heterotrophic growth in the upper ocean, including the long-standing issue of how and why bacteria may be seen as culturable or otherwise. Before we turn to conclusions that may be drawn from studying bacterial isolates, we need to address the fundamental question of how to reconcile 2 images of the bacterial species concept. On the one hand, it is an empirically proven fact that taxonomically and functionally consistent strains of widely known bacteria such as Escherichia coli and Vibrio cholera can be isolated repeatedly.
(manuals found in any bacteriology textbook). On the other hand, a tremendous bacterial diversity can be anticipated from surveys based on molecular methods (Temperton & Giovannoni 2012). In relation to the supposition that bacterial isolates might be useful as model bacteria, the question must therefore be raised as to how information from a single isolate may be of any relevance if the species richness is close to endless.

Relevance of single genomes/isolates and 16S rRNA

To overcome the difficulty of cultivation, Swan et al. (2013) used a protocol for single amplified genomes (SAGs) to identify genomes from single bacterial cells. In their study, 56 SAGs were used to analyze genome streamlining and latitudinal divergence of planktonic bacteria. However, the same question of relevance to the natural assemblages may be raised towards the SAG studies as with bacterial isolates serving as model organisms. Can single genomes/isolates represent wide taxa? In this context, the SAGs represent a biased selection, in the same way that isolation of bacteria on agar plates selects for opportunistic bacteria irrespective of their abundance. In the case of the SAGs, the relative abundance of the bacteria determines which cells preferentially become sequenced in the sample. Still, the larger question is not sampling bias per se but whether single genomes from either SAGs or bacterial isolates can be seen as representative model organisms. To answer this question, the following may be argued: (1) Based on DNA−DNA hybridization results, it is demonstrated that annually recurring bacterial blooms (Rehnstam et al. 1993) and seasonal succession of marine bacteria (Pinhassi & Hagström 2000) can be recorded. Thus, DNA from single bacterial isolates can be used to find and quantify closely related DNA in environmental samples through DNA−DNA hybridization. (2) Similarly, global distribution of SAG-related microorganisms (Swan et al. 2013) and insights into an abundant and uncultivated marine bacterial lineage SAR86 ( Dupont et al. 2012) can be obtained using metagenomic fragment recruitment. Thus, genetic markers identified from a single-cell genome can be used to determine identity and abundance of defined taxa in metagenome samples using a phylogenomic analysis.

There are useful techniques available to identify and quantify bacterial taxa in the environment either directly or in silico from metagenomes. So, if identifying means recognizing and knowing the names of bacterial isolates, we can give a positive answer. This means microbial ecologists should be able to handle the species concept pragmatically, as we do with insect species or human pathogens that are the subjects of constant evolution. We need to recall that pathogenic bacteria can be recognized through isolation from the patient’s body. Bacteriologists have thus learned to recover and name bacteria responsible for particular illnesses by experience and pragmatic means (Fraser et al. 2009). However, for this to happen in bacterial ecology, focus has to be turned away from bacterial phylogeny and be directed towards the identity of functional bacterial taxa. Then, we may see a totally different description of marine bacteria in the future.

16S rRNA species specificity

For the past 3 decades, the majority of all phylogenetic analyses of bacterial clones or isolates have been based on sequencing the 16S rRNA gene. Because of its very specific function, this gene displays conserved regions, and it has become generally accepted that these regions carry some degree of specificity towards different kinds of bacteria. Since the suggestion by Woese that relationships between bacteria may be inferred from sequence similarity of the 16S rRNA, new bacteria, new habitats and global patterns of bacterial distribution have been described (Woese 1987, Fuhrman & Hagström 2008). It is therefore an undisputed fact that 16S rRNA-based studies in the sea have been very valuable. Yet, we may question parts of the original argument, since over the last few decades the effort to produce plausible lists of bacterial species for specific habitats has largely failed. Not only are we lacking a valid species definition (Fraser et al. 2009), but the whole taxonomic framework also suffers from undocumented assumptions. Experimental lateral transfer of intact 16S rRNA genes (Kitahara et al. 2012) and genetic engineering of specific fragments of the 16S rRNA gene between phylogenetically distant bacteria (Kannan et al. 2016) show fully functional ribosomes. This implies that over long periods of time, 16S rRNA genes could have been exchanged between core genomes. These results thus undermine the original assumption and add support to Kitahara et al. (2012), who conclude, ‘The extreme ribosomal complexity is believed to be the basis for the species specificity in 16S rRNA, although solid experimental evidence is not available in this regard’. From this we realize and
agree with Swan et al. (2013) that insufficient phylogenetic resolution might explain the difficulties in detecting consistent differentiation of bacterioplankton when performing small subunit rRNA gene surveys, for instance, along longitudinal gradients, although the more pronounced differences between polar and tropical bacterioplankton have been reported from such studies (Pommier et al. 2005, 2007, Ghiglione et al. 2012).

**BACTERIAL GROWTH RATE ADAPTATION TO TEMPERATURE — A CASE STUDY**

In the following section, the aim is to illustrate that ecophysiological insights can be gained by studying suitably selected bacterial isolates. For this, we use a 13 yr old and previously unpublished data set containing 55 bacterial isolates from 3 marine areas; this data set has been revisited, since it allows a comment on how the marine microbiology community might assimilate this information in thinking about the use of isolates in ecophysiology and biogeochemistry. Each isolate was grown at steady state in Zobell liquid media at 10 different temperatures from 5 to 50°C, and the respective growth rates were recorded (see the Supplement at www.int-res.com/articles/suppl/a080p015_supp.pdf for details of experimental conditions and statistical analyses). Based on these results, and in agreement with those discussed in the previous section from global surveys showing a latitudinal differentiation of marine bacterioplankton, the following framework can be formulated: The growth rate for different isolates, at their optimum temperature, depends on phylogenetic affiliation rather than geographic origin. Thus, temperature would drive annual species succession, as also suggested by results of culture-independent analysis from the ‘Tara’ expedition (Sunagawa et al. 2015). A major conclusion from the ‘Tara’ work is summarized in the following quote: ‘Taken together our data suggest that geographic distance plays a subordinate role and reveals temperature to be the major environmental factor shaping taxonomic and functional microbial community composition in the photic open ocean’.

**Bacterial isolate identity**

The bacteria in our analysis included 2 Alphaproteobacteria, 4 Gammaproteobacteria and 6 Bacteroidetes from the Mediterranean Sea; 4 Alphaproteobacteria, 3 Gammaproteobacteria and 7 Bacteroidetes from the Southern California Bight; and 5 Alphaproteobacteria, 1 Gammaproteobacteria and 6 Bacteroidetes from the Baltic Sea. Details of these isolates are compiled in Table S1 in the Supplement. When comparing the identities of the isolates based on 16S rRNA sequence similarity generated in 2003 and 2016, a substantial discrepancy is apparent. It is clear that over time, identity defined as resemblance to other bacteria based on partial sequences of a single gene does not provide sufficient resolving power to go beyond phylum or subclass. However, the distribution of the bacterial isolates over a general phylogenetic tree is sufficient to allow us to determine that the isolates do not represent a closely related cluster (see Fig. S1 in the Supplement).

**Growth characteristics of selected opportunistic bacteria**

Each isolate demonstrated a characteristic temperature-to-growth rate relationship as exemplified in Fig. 1, showing a number of isolates from the Southern California Bight. Growth rate at optimum growth temperature was determined from the resulting growth curves, as well as minimum and maximum temperature.
tures for growth. Typically, growth rates at temperatures above the optimum temperature shape the curve into a steep downward slope. The results are summarized in Fig. 2, where the average growth rates of the isolates at each temperature are plotted for each of the 3 different bacterial divisions. The large variability in growth rate at higher temperature is caused by the different optima displayed by the isolates, causing the growth rate to drop within a wide upper range of temperatures (see Fig. 1). The growth rates were significantly related to growth temperature and taxonomic affiliation of the investigated strains.

Fig. 2. (A) Mean growth rates divided after taxonomic affiliation of isolates at different temperatures; error bars represent SD. (B−D) Box plots showing growth rates of Alphaproteobacteria (red), Gammaproteobacteria (blue) and Bacteroidetes (green). Horizontal line inside the box represents the median, which is also connected by curves throughout the temperature range (line plots). Upper and lower margins of the boxes are the 75th (Q3) and 25th (Q1) percentiles; dots represent outliers defined as being less than Q1 − 1.5 × IQR or greater than Q3 + 1.5 × IQR; IQR = Q3 − Q1. IQR: interquartile range; Q: quartile

vs. Alpha: p = 0.0023*
vs. Bacteroidetes: p = 0.0128*

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vs. Alpha: p = 0.0128*
vs. Gamma: p = 0.5264
(p = 5.9 × 10⁻⁹ and p = 7 × 10⁻⁴, respectively). In contrast, growth rate was not related to geographical origin (p = 0.47). Grouping the strains by their taxonomic origin, the Alphaproteobacteria had a significantly lower growth rate than the Bacteroidetes (p = 0.0128, Tukey’s HSD post hoc test) and the Gammaproteobacteria (p = 0.0023). In contrast, growth of Gammaproteobacteria was not significantly different from that of Bacteroidetes (p = 0.53). From these results, we find weak but statistically significant support for the hypothesis that major phylogenetic groups on average show different growth rates.

**Sampling sites and isolates**

The different sea areas chosen to isolate bacteria from are different in many aspects, but here 2 particular aspects are noted, temperature range and overall nutrient regimes. The brackish Baltic Sea is temperate, with water temperature ranging from 0 to 20°C over the year and with a salinity gradient from north to south of 3 to 10‰. Phosphorus limits growth in the north, while nitrogen limitation prevails in the south, i.e. in the Baltic proper (Zweifel et al. 1993). The northeastern Mediterranean Sea and the Southern California Bight, on the other hand, both exhibit similar salinity and temperature regimes, yet the nutrient status differs, the Mediterranean Sea being phosphorus limited and the Southern California Bight being mainly nitrogen limited (Fuhrman et al. 1988, Thingstad & Rassoulzadegan 1995).

The Baltic strains showed an optimum growth temperature between 15 and 35°C, with an average of 26 ± 6°C (SD) (Table 2). All strains could grow at 5°C, and 2 strains had an optimum growth temperature <20°C (Fig. 2). The growth rate at optimum growth temperature averaged 0.40 h⁻¹. The isolates from the Mediterranean Sea had an optimum growth temperature between 30 and 35°C, with an average of 28 ± 4°C. Three of 7 strains did not grow below 10°C. The growth rate at optimum growth temperature averaged 0.50 h⁻¹. Among the isolates from the Southern California Bight, optimum temperature for growth varied between 25 and 40°C, with an average of 29 ± 4°C (Table 2). Fifteen of 19 strains were capable of growth at 5°C, while 3 strains did not grow below 10°C, and 1 strain did not grow below 15°C. The growth rate at optimum temperature for growth averaged 0.54 h⁻¹. Bacteria able to grow at 5°C were only found among the Baltic Sea isolates, and 2 psychrophiles (bacteria with an optimum growth temperature <20°C) were found among these isolates.

This suggests adaptation to the temperature environment from which the bacteria were isolated. However, growth rates did not differ significantly (p = 0.46 to p = 0.95, 2-way ANOVA and Tukey HSD post hoc test) between isolates from the 3 different sea areas. In nature, bacteria will not experience the optimum growth temperature, since this temperature averaged 30°C for all isolates tested. Thus, the isolated marine bacterioplankton in this study usually grow at temperatures >10°C below their optimum temperature for growth.

**Motility**

In an attempt to find further characteristics between different groups of isolates, motility was recorded. While it seems the true oligotrophs are non-motile, a total of 58% of the tested opportunistic bacteria were motile (Table 2). Different swimming modes were visually recorded and ranged from a comparably slow pace to intense swarming. In an earlier study of many of the same bacterial isolates as in this study, Alphaproteobacteria were found to have the slowest swimming speed (Johansen et al. 2002). Of the 3 major divisions of bacteria studied, the Alpha- and Gammaproteobacteria included a higher number of motile bacteria (82 and 75%, respectively) than the Bacteroidetes (37%). The isolates from the Baltic included a higher number of motile bacteria (83%) than those from the Mediterranean Sea and Southern California Bight (50 and 43%, respectively).

**CONSEQUENCES AND SUGGESTIONS**

Seawater may seem to be a poor substrate, and compared to a rich broth it is certainly dilute; how-

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**Table 2.** Average phenotypic characteristics of isolates compared between affiliation and origin. *Significantly higher than Alphaproteobacteria (p < 0.05). Alpha: Alphaproteobacteria; Bact: Bacteroidetes; Gamma: Gammaproteobacteria; BAL: Baltic Sea; MED: Mediterranean Sea, SCB: Southern California Bight; Topt: temperature optimum

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<th>Phenotypic test</th>
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<tr>
<td>Topt (°C)</td>
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<tr>
<td>Growth rate at Topt (h⁻¹)</td>
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<td>Motility (%)</td>
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<td>Topt (°C)</td>
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<tr>
<td>Growth rate</td>
<td>0.30 0.50* 0.64* 0.40 0.50 0.54</td>
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<tr>
<td>Motility (%)</td>
<td>82 37 75 83 50 43</td>
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ever, the quality of the substrate, that is, the proportions of macromolecules to monomers and organic to inorganic ratios, is much the same in seawater as in a broth. Thus, having an efficient uptake system is what determines whether a bacterial strain can grow in seawater, and the cell abundance, of course, can never be high.

**In situ growth rate versus culture data and life strategy**

The growth rates given in Table 2 for the opportunistic bacteria were recorded at temperature optimum for the respective strain, and to compare these rates to those of the oligotrophs grown at 15°C (Table 3), a correction is needed. From the curves in Fig. 2, the range of growth rate at 15°C can be seen to be between 0.09 and 0.20 h⁻¹ for the isolates of the opportunistic bacteria. If that is compared to growth rates of around 0.015 h⁻¹ for the oligotrophs (Table 3, growth rate d⁻¹), the former are more than 10 times faster than the oligotrophs. These group-specific traits may be important for the establishment and succession of bacterioplankton (as discussed in the following sections) in relation to properties of the core genome defining a bacterial species.

Comparing the growth rates of virus-free bacterial isolates in cultures, oligotrophic bacteria on average grew >10 times slower than most opportunistic bacteria (Fig. 2, Table 3). The real question, however, is what this difference in growth rate might be in the natural environment. For the sake of argument, we will claim that the oligotrophic strains in Table 3 grow in seawater, although the SAR11 cultures were amended with mineral nutrients. After all the talk earlier in this paper concerning SWC, it may come as an anticlimax that very few identified opportunistic bacterial isolates have been grown in seawater. In fact, the *Alteromonas* strain grown by Pedler et al. (2014) and *Dokdonia* sp. strain MED134 (Gómez-Consarnau et al. 2007), together with the SAR11 isolates, are the only published examples we could find. There exist, however, quite a few published data sets of seawater cultures from chemostats and batch cultures, although these are all mixed cultures. The *Alteromonas* and *Dokdonia* both grew at about 0.03 h⁻¹, which is twice as fast as the common oligotrophs (0.015 h⁻¹). In the seawater chemostats with mixed marine bacteria inoculates, the maximum recorded and published turnover rate is 0.11 h⁻¹ (Hagström et al. 1984), thus at the lower range of the growth rate of the opportunist strains in this paper. Still, if we consider that, in addition to the flow rate removing cells from the chemostat, an internal mortality factor caused by phages has to be compensated for, we can estimate the actual growth rate in the chemostat. From model simulation of a data set obtained in a multistage seawater chemostat, the true growth rate was shown to be 2 to 10 times higher for the non-infected cells depending on the turnover rate in the chemostat (Blackburn et al. 1996, Zweifel et al. 1996).

In terms of bacterial production, this means that while oligotrophic bacteria are very abundant in surface waters, opportunistic bacteria should easily outgrow them, and as their cell volume can be >10 times larger than the oligotrophic bacteria, this ability becomes even more pronounced in production terms. In this regard, Pedler et al. (2014) demonstrated complete removal of the labile DOC in coastal surface seawater by a single opportunistic taxon.

**Generating microdiversity in real time**

Bacteriophage resistance is a summary term for a large number of molecular strategies where the bacteria are able to fend off the attacks by viruses to some degree (Labrie et al. 2010). In every case, an evolutionary arms race of varying intensity can be expected to result in the development of microdiversity. In particular, the interaction between the outer membrane proteins and phage attachment

<table>
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<tr>
<th>Isolate</th>
<th>Accession no.</th>
<th>Division</th>
<th>Source</th>
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<th>Solid surface</th>
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<td>0.27</td>
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<td>SKA48</td>
<td>AY317115</td>
<td>Alphaproteobacteria</td>
<td></td>
<td>0.40</td>
<td>No growth</td>
<td>0.01</td>
<td>Simu &amp; Hagström (2004)</td>
</tr>
<tr>
<td>SAR 11</td>
<td>HTCC1002</td>
<td>Alphaproteobacteria</td>
<td></td>
<td>0.40–0.58</td>
<td>ND</td>
<td>ND</td>
<td>Rappé et al. (2002)</td>
</tr>
</tbody>
</table>
may serve as a rapidly evolving battlefield. The outer membrane of Gram-negative bacteria contains several general porins, various high-affinity substrate receptors and different iron complexes, which are used as phage receptors. Further examples are the TonB and TolA proteins, which are anchored to the inner membrane but span the periplasm and as such are important for infection by many phages (Guttman et al. 2005). Thingstad et al. (2015) have proposed a model to account for phage resistance and the number of bacterial taxa in a given habitat. Based on their model, we visualize the virus host interaction in a closed substrate controlled space as follows: Imagine an initial number of cells of a bacterial species that utilizes the available food source in full. A corresponding bacteriophage would then be able to infect and kill a large fraction of these cells. This will in all likelihood select for a modified bacterium (a strain) resistant to the phage, but with resistance follows a slight penalty in growth rate that does not prevent utilization of the food source not used by the original unmodified species. In this manner, new strains of both the host and the phage can be selected for, in an ongoing arms race, until there is no more room (substrate resource) for hosts to grow and produce large enough phage populations that can infect sensitive hosts. An alternative restriction to further host strains is if growth rate due to cost of resistance for new strains has been reduced below the bacterial clearance rate exerted by the protozoa. However, at any time during the process, a new species equipped with a different core genome can be introduced into the habitat and, if conditions like temperature have changed, may compete successfully for the available resources, since, on the scale of bacterial presence, conditions constantly change. This is due to diurnal variation, zooplankton activity and so on. Thus, the arms race within each bacterial species will rarely come to an end; instead, an arena with a diverse bacterial community is generated containing high viral numbers and many susceptible hosts.

**Marine isolates representing core genomes**

The results in Tables 2 & 3 originate from growth experiments involving a long list of marine isolates (Table S1 in the Supplement). To understand the integration of these isolates in the natural environment, we suggest that the pan-genome concept of Tettelin et al. (2005) be adopted. The pan-genome includes the core genome-containing genes present in all strains, where in addition certain strains harbor unique but dispensable genes (Medini et al. 2005). In seawater, a successful bacterial species carries a core genome well suited for the particular habitat, which is demonstrated by fast growth. To flourish, the species must have the correct temperature optimum, a shape and size suitable for osmotrophic life and the ability to display a correct salinity preference; to do even better, it should have acquired additional traits that match the existing local conditions. The core genome has an organized structure evolved through having experienced eons of habitat fluctuations (Rocha 2008, Caro-Quintero & Konstantinidis 2012). With this view, the core genome may be regarded as the evolutionary solution to the least unfavorable gene composition in relation to environmental change. The difference in growth characteristics seen between the different groups of bacteria in Table 2 may thus be an observable example of this overarching evolutionary outcome.

**CLOSING REMARKS**

While large numbers of bacterial strains have had their DNA sequenced partly or as whole-genome projects, no general information about the organisms can be found. Compared to animals and plants, where the species names carry information gathered over hundreds of years, most bacteria, except perhaps pathogenic strains, are totally anonymous. To advance the field of marine microbiology, we therefore rally behind the strategy of Browne et al. (2016). These authors, working on the human microbiota, suggest that when culturing targeted phenotypes and linking whole-genome sequencing with phylogenetic analysis and computational modeling, the results will demonstrate a substantial proportion of bacteria in different habitats to be culturable. The extensive work by Stephen Giovannoni and his group on *Pelagibacter ubique* serves as an excellent example of how to isolate, grow and investigate a new bacterium (Giovannoni & Stingl 2007, Carini et al. 2013). To move microbiology forward, we conclude that physiological data need to be included into molecular databases, and other recent papers have also highlighted this same need from different perspectives (Garcia 2016, Salcher & Šimek 2016). Examples of prokaryote behavior that can be recorded easily are growth strategies and motility. In the future, we believe that generating ecological information from cultured bacteria will revitalize marine microbiology.
Acknowledgements. This paper is based on a keynote address delivered by Å. Hagström at the EMBO Symposium on Aquatic Microbial Ecology (SAME) 14 meeting held in Uppsala, Sweden, 23−28 August 2015. This work was supported by the BLUEPRINT project, funded by the European Community’s Seventh Framework Programme (FP/2007-2013) under implementation agreement no. R&I/ I3/2012/014776. BONUS made with BONUS, the joint Baltic Sea research and development programme.

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Editorial responsibility: Paul del Giorgio, Montreal, Quebec, Canada

Submitted: November 17, 2016; Accepted: May 22, 2017

Proofs received from author(s): August 4, 2017