Microorganisms and Detritus in the Water Column of a Subtidal Reef of Natal

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ABSTRACT: The parameters measured in this study were light and dark-bottle uptake of NaH\textsubscript{14}CO\textsubscript{3}, the chlorophyll content of seawater, direct counts of bacteria, microbial uptake of \textsuperscript{14}C glucose and a labelled algal extract, organic carbon in the seawater and the composition of debris washed onto the reef. The phytoplankton consisted largely of nannoplankton and primary production was low with an annual mean of $P = 12.82 \text{ mg C m}^{-3} \text{h}^{-1}$, while the mean chlorophyll a content of 2.13 mg m\textsuperscript{-3} was equivalent to a dry biomass of 170.1 mg m\textsuperscript{-3}. Microbial heterotrophic activity was relatively high with an annual mean glucose assimilation of $V_{\text{max}} = 0.612 \mu \text{g g}^{-1} \text{h}^{-1}$ and a total bacterial count of $2.02 \times 10^6 \text{ ml}^{-1}$, equivalent to a dry biomass of 20.3 mg m\textsuperscript{-3}. The largest proportion of this activity, measured by differential filtration and using antibiotic inhibitors, is performed by free bacteria as opposed to the smaller proportion conducted by bacteria attached to particles, or by microflagellates. Uptake of the labelled algal extract ($V_{\text{max}} = 13.2-72.6 \text{ mg C m}^{-3} \text{h}^{-1}$) indicates that microbial heterotrophic activity exceeds phytoplankton production. Carbon analyses gave an annual mean of $18.3 \text{ g m}^{-3}$ total organic carbon, of which 26% was particulate, the rest being dissolved organic carbon. Analysis of debris revealed that it consisted almost exclusively of seaweed, except when rivers were in flood, providing a large influx of terrestrial plant debris. These findings are compared with other published results, and their ecological significance is discussed in relation to the remainder of the environment. It is concluded that primary production by phytoplankton is not as important in supporting the biomass of filter feeders on the reef as is heterotrophic activity associated with organic detritus.

INTRODUCTION

The littoral shoreline of Natal (South Africa) is typified by long stretches of surf-washed beaches and sandy substrate interspersed by rocky reefs. Although these reefs are less extensive than the sandy substrate, they are highly productive. Biological studies have been completed on the predominant organisms resident on the reefs at several trophic levels, viz. mussels (Berry, 1978), rock lobsters (Berry, 1971a, b; Smale, 1978; Berry and Smale, 1980), octopus (Smale and Buchan, in prep.) and teleost fish (Berry et al., 1979b; Joubert, 1980; Joubert and Hanekom, 1980). Suspension feeders comprise the dominant faunistic component (Jackson, 1976; Berry et al., 1979a); a study of their particulate detrital and planktonic food sources, crucial to a complete understanding of the reef systems, was lacking until the present work.

Jackson (1976) and Berry (1978) discuss prevalent physical and climatic conditions on the Natal coast that influence the reef habitat; the latter describes the large physical energy subsidies contributed to reef systems by continuous swell and surf action and longshore currents. Large quantities of seaweed and terrestrial macrophyte debris transported by these currents accumulate in the lee of the reefs and are rapidly broken down by turbulence and sand abrasion into fine detrital particles (Berry, 1978; Berry et al., 1979a; Schleyer, 1979, 1980). Berry (1978) suggested that phytoplankton production was too low in Natal waters to support the large populations of suspension-feeders and postulated that they feed on these detrital particles. The purpose of this study was thus to establish the origin and relative abundance of the different particulate fractions available to the suspension-feeders. It forms part of a programme in which the ecology of a shallow subtropical reef community was investigated in terms of energy flow.

MATERIALS AND METHODS

Study Area

A small, isolated reef situated in front of the Oceanographic Research Institute (ORI) in Durban, Natal (South Africa), was chosen as study area. This artificial reef, known as ORI Reef (Berry, 1978; Smale,
Production. Fixation of $^{14}$CO$_2$ was measured, using the traditional light and dark-bottle method (Vollenweider, 1971) as detailed by Allanson and Hart (1975), except that samples were collected on 0.2 $\mu$m pore-size polycarbonate membranes (Nuclepore Inc.). These membranes were used in all the tracer work described in this paper and were dissolved with 0.1 ml phenylethylamine before counting in Instagel. This eliminated problems such as chemiluminescence and phosphorescence, which are associated with the use of dioxane, the scintillation cocktail used to dissolve cellulose acetate/nitrate membranes previously employed by certain workers (Ward and Nakanishi, 1971, 1973).

Silver and Davoll (1978) found that loss of fixed $^{14}$C, a problem caused by addition of a chemical fixative to stop incubation, was reduced if a solution of $I_2 + KI$ was added instead of formalin. The converse proved true when tested in the ORI Reef experiments, possibly because filtration proceeded soon after incubation was stopped and formalin thus remained the chemical fixative of choice.

At the start of the 6-h incubation period, the samples were placed in an outdoor bath supplied with a constant flow of seawater and screened with plankton and fish net to reduce light intensity to the level recorded in the field at 1 m depth. Light intensity fluctuated during incubation according to cloud cover and sun elevation, as did the temperature in the water bath according to sea temperature. Computations of production are based on the formulae and tables of Strickland and Parsons (1968).

Bacteria

Total counts and calculations of bacterial biomass. Total bacterial counts were performed on 0.2 $\mu$m pore size polycarbonate membranes using the AODC technique already mentioned. The numbers of bacteria attached to detrital particles were recorded, as were the numbers of cocci (including ovoid forms), rods, commas and spirals. Notes were made on the nature, size and abundance of the detrital particles. Scanning electron micrographs provided linear dimensions of the different morphological forms of bacteria, and these were used to calculate mean volumes from which total counts were converted to values of biomass according to Sorokin and Kadota (1972). For this purpose, 10 samples collected over a 5 month period were fixed with glutaraldehyde (1–2 %), and the bacteria were filtered on 0.2 $\mu$m pore size polycarbonate membranes. The filters were then dehydrated at half-hour stages in a series of mixtures of ethanol, distilled water and sterile filtered seawater in the following proportions: 1:1:8 → 2:1:7 → 3:1:6 → etc.
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Subsequently they were critical-point-dried with CO₂ and routinely mounted and gold-coated for SEM viewing. The addition of distilled water at the dehydration stages proved necessary to prevent salt crystallisation which obscured the bacteria. The bacteria were well preserved (Fig. 1), but since dehydration followed by critical-point-drying is known to cause specimen shrinkage, the linear dimensions were increased by a factor of 10%. This rather arbitrary figure is based on a study of the problem by Boyde et al. (1977) and is commonly used (Evers, pers. comm.). When observed, microflagellates in these samples were also photographed for measurement.

Assimilation of ¹⁴C glucose. Dark assimilation of ¹⁴C glucose was measured, using the heterotrophic potential technique of Hobbie and Crawford (1969). A range of 8 concentrations (0.025 to 0.2 μCi) of D-[U-¹⁴C] glucose (specific activity 292 mCi mmol⁻¹ or 1.55 mCi mg⁻¹) was added to 25 ml subsamples of seawater, which were incubated for 1 or 2 h at sea temperature. A modification to the technique involving improved collection of respired ¹⁴CO₂ and advantages in the use of polycarbonate membranes in reducing the filter error are discussed by Schleyer (1980).

100 ml seawater samples were also incubated with 0.2 μCi labelled glucose and differentially filtered through 3.0 μm and 0.2 μm pore size polycarbonate membranes after incubation. This procedure was performed during the latter half of the year's sampling to establish whether the fraction of bacteria attached to detrital particles was more active than free bacteria.

Carbon Analysis

In order to measure dissolved organic carbon (DOC) and particulate organic carbon (POC), unfiltered seawater samples and membrane-filtered samples clear of POC were analysed on a Beckman Carbon Analyser. This gave results of total organic carbon (TOC) and DOC, the difference of which was assumed to comprise POC.

Debris Washed onto the Reef

'Mats' of debris were observed moving above the sand and accumulating adjacent to the reef in areas of reduced water circulation when sea conditions were calm. Samples of this debris were taken when possible and separated into constituents to establish their origin. These were oven-dried (60°C) to constant weight and the relative proportions were determined. The calorific content of the constituents of one sample was measured, using an adiabatic bomb calorimeter. For most of the year, conditions were too rough for the mats to form and the debris was churned in the surf and rapidly broken up.

Sampling on a Transect Across the Reef

Fortnightly samples were taken over a year for total counts of bacteria on a transect across the OR1 Reef at stations up- and down-current of the reef and on the reef itself. Samples were taken shortly after the turn to an incoming tide to facilitate collection of the samples and to detect current direction, so that the degree to which the reduced volume of water moving across the reef was filtered by suspension-feeders could be ascertained. Flow of surf onto the reef usually exceeded the flow of longshore currents across it, so that these counts can only be considered an indication of the filtering activity.

Identification of Heterotrophic Microbial Components and an Assessment of their Importance

The uptake of a labelled algal extract and of ¹⁴C glucose was measured over a 3 month period, using the heterotrophic potential technique in 5 experiments reported by Schleyer (1980). The labelled algal extract consisted of dissolved organics extracted from a ¹⁴C labelled Chlorella culture after sonication. It was added to samples at final concentrations of 0.656 to 3.282 μg C m⁻³ in an attempt to provide microbial heterotrophs with a tracer similar to the natural substrate and to facilitate measurement of their production. Labelled glucose concentrations were as previously described. Antibiotics were added to duplicate samples to inhibit bacterial activity and assess non-bacterial heterotrophy or uptake of the labelled substrate by passive diffusion. The samples were pre-incubated with antibiotics for an hour in the dark for these to take effect, whereafter normal incubation of an hour proceeded at sea temperature, again in the dark.

The antibiotics used were gentamycin (20 μg ml⁻¹) in 3 of the experiments and a mixture of penicillin G (400 μg ml⁻¹), streptomycin (200 μg ml) and chloramphenicol (80 μg ml⁻¹) in 2 of the experiments. Gentamycin was found by Chröst (1978) to be effective in totally inhibiting bacterial activity after a pre-incubation of an hour at the above concentration, without affecting photosynthetic carbon fixation by freshwater phytoplankton for the next 4 h. It proved ineffective in procuring total bacterial inhibition in this study, but its use is reported since its mode of inhibition was interesting. The other antibiotic mixture described above is recommended for culturing microflagellates axenically (Guillard,
1973); it proved effective for this purpose with a local species at the above concentrations (Aken, pers. comm.). Nevertheless, it was tested at two and three times these concentrations in a study of its effectiveness and for eukaryotic toxicity in the present work, using differential filtration through 1.0 μm and 0.2 μm pore size polycarbonate membranes. Other modern antibiotics were tested in synergistic combinations, viz. carbenacillin with gentamycin and with cefoxitin, but were found to be almost as ineffective as gentamycin on its own.

Seawater samples of 100 ml were similarly incubated separately with labelled algal extract and 14C glucose (1.313 mg C l⁻¹ and 1.290 μg l⁻¹ respectively) and differentially filtered through 3.0 μm and 0.2 μm, 2.0 μm and 0.2 μm or 1.0 μm and 0.2 μm pore size polycarbonate membranes. These were counted and the percentage uptake of the different size fractions of the nannoplankton was calculated to establish their heterotrophic role.

### RESULTS

#### Phytoplankton

The mean temperature of samples collected over the year for these and the following studies was 22.6°C, with a range from 19.1°C to 26.5°C.

Primary production was variable and there was a wide range in light and dark-bottle carbon fixation, which had a coefficient of variation of ±66% around the annual mean of 12.82 mg C m⁻³h⁻¹ (Table 1). This variability was not clearly related to environmental factors; even though low primary productivity was recorded under turbid and turbulent conditions on most occasions, the two highest measurements were

<table>
<thead>
<tr>
<th></th>
<th>mg C m⁻³h⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual mean light-bottle fixation</td>
<td>12.92</td>
</tr>
<tr>
<td>Annual mean dark-bottle fixation</td>
<td>0.10</td>
</tr>
<tr>
<td>Annual mean production</td>
<td>12.82 ± 8.51</td>
</tr>
<tr>
<td>Maximum production</td>
<td>31.29</td>
</tr>
<tr>
<td>Minimum production</td>
<td>1.69</td>
</tr>
</tbody>
</table>

Table 1. Primary production measured using the ¹⁴C light and dark bottle method. (n = 29)
taken under these conditions. Chlorophyll a analyses (Table 2) were not as variable, being less dependent on environmental conditions, and the annual mean biomass of phytoplankton was estimated from these analyses to be 170.1 mg m\(^{-3}\) (dry weight).

Using Cassel's (1965) graphical technique, phytoplankton counts were estimated to be precise to only ±35 % at a confidence level of 95 % and thus are not presented in detail except as seasonal means (Table 11). These counts fluctuated in general with increase or decrease in the previous parameters and were dominated by microflagellates, referred to as monads by Hobbie et al (1972). Only a few pennate diatoms, filamentous forms and planktonic colonies such as *Asterionella japonica* were counted.

### Bacteria

The microbes in samples prepared for SEM measurements of bacteria appeared to be well preserved, as there was no evidence of distortion (Figs 1 and 2). The weighted mean diameter of cocci in these samples was 0.476 \(\mu\)m (Table 3), and some cocci measured were as small as the maximum pore size of the filters used in the study (0.2 \(\mu\)m), indicating that a proportion of bacteria was probably lost during filtration. It would have been impossible to measure such small bacteria accurately using light microscopy; hence measurement of the actual bacteria counted by AODC was precluded. This necessitated the separate SEM measurement of bacteria for conversion of total counts to bacterial biomass. A final mean bacterial volume of 0.0516 \(\mu\)m\(^3\) was obtained for this purpose from the measurements, which provided weighted mean volumes of the morphological types of bacteria. These were adjusted to the final volume according to the percentage occurrence of the morphological types in the total counts (Table 3), which did not vary much from sample to sample. The coefficient of variation

<table>
<thead>
<tr>
<th></th>
<th>Chl a (mg m(^{-3}))</th>
<th>Dry biomass (mg m(^{-3}))</th>
<th>Phytoplankton C (mg m(^{-3}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual mean</td>
<td>2.13</td>
<td>170.1</td>
<td>85.0</td>
</tr>
<tr>
<td>Maximum</td>
<td>3.88</td>
<td>310.4</td>
<td>155.2</td>
</tr>
<tr>
<td>Minimum</td>
<td>0.76</td>
<td>60.8</td>
<td>30.4</td>
</tr>
</tbody>
</table>
around the weighted mean volumes was fairly low (± 30%), suggesting that the final mean volume was acceptable for the intended purpose. Measurement of a small sample (n = 16) of microflagellates provided a mean width for these organisms of 2.32 μm (range 1.41 μm to 3.16 μm), this being their smallest dimension.

Using Cassell's (1965) graphical technique, bacterial counts were estimated to be precise to ± 11.5% at a confidence level of 95%, and counts on the reef transect indicated that filtration of these organisms occurred as water moved across the reef. The annual mean differential count across the reef was 0.188 × 10^6 ml^-1 (1.218 mg m^-3 dry biomass) and on one occasion counts as high as 4.276 × 10^6 ml^-1 occurred, with a differential across the reef of 2.639 × 10^6 ml^-1 (2.723 mg m^-3 dry biomass, Table 4).

Counts performed on samples collected for determining the heterotrophic potential were consistent, with an annual mean of 2.02 ± 0.37 × 10^6 ml^-1 and a mean dry biomass of 20.85 mg m^-3 (Table 5). The percentage of bacteria attached to detrital particles was variable, being on average 21 ± 14% of the total count. Occasionally this percentage was underestimated when dense detrital particles were present in a sample and bacteria could be counted only on one side of the particles.

The detrital particles were generally amorphous aggregates, rarely including recognisable material derived from living organisms. Most were roughly 5 to 10 μm or larger in size, but they could be as small as 1 μm.

### Heterotrophic Activity

The annual mean of ^14^C glucose assimilation measured using the heterotrophic potential technique was \( V_{\text{max}} = 0.612 \mu g \, l^{-1} h^{-1} \) (0.263 μg C l^-1 h^-1) with a maximum uptake of 2.326 μg l^-1 h^-1 which occurred in spring. These values are listed with other parameters relating to heterotrophic glucose uptake in Table 6.

In the supplementary experiments in which the uptake of labelled algal extract and labelled glucose were compared, there was no correlation between the uptake of these two substrates (Table 7). An increase or decrease in the uptake of one was generally accompanied by a similar change in the uptake of the other, but the second highest uptake of the labelled algal extract was accompanied by the second lowest uptake of glucose in the 5 experiments. The previous spring maximum uptake of glucose was exceeded in one of the experiments by a \( V_{\text{max}} \) of 2.9 μg l^-1 h^-1 (1.247 μg C l^-1 h^-1) at the end of winter. The other 4 experiments were performed in spring and the mean experimental temperature was 22.0 °C.

Berman (1975) found gentamycin inconsistent in its effectiveness on marine bacteria; it barely reduced microbial activity (by < 30%) in 2 of the first 3 of these experiments. The antibiotic mixture (A. M. in Table 7) was far more effective and in most instances reduced...
microbial activity to below 50% of normal activity. In action, gentamycin displayed competitive inhibition and the antibiotic mixture non-competitive inhibition (Figs 3 and 4); the significance of this is not understood.

The efficacy of the antibiotic mixture was better revealed by the differential filtration experiments designed for this purpose (Table 8), than by the results of the comparative uptake studies which were low and lacked statistical validity. Since the mean diameter of cocci was only 0.476 μm, most free bacteria would have

Table 7. Results of separate uptake of labelled algal extract and labelled glucose measured in parallel experiments using the heterotrophic potential technique, with and without antibiotics added. All results proved significant at the 95% level using the F ratio test, except those marked +. Those marked * were predictive as well as significant, the former being defined as having an F ratio greater than 4 times the critical F value. The results are partially taken from Schleyer (1980); sample numbers refer to the same samples as those in Tables 8 and 9. The mean temperature of the experiments was 22°C.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Labelled algal extract</th>
<th>Labelled glucose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$V_{max}$ T K+S</td>
<td>$V_{max}$ T K+S</td>
</tr>
<tr>
<td></td>
<td>(μg C l$^{-1}$ h$^{-1}$)</td>
<td>(μg C l$^{-1}$ h$^{-1}$)</td>
</tr>
<tr>
<td>1</td>
<td>72.9* 14.5* 1049.7* 2.9* 3.2* 9.2*</td>
<td></td>
</tr>
<tr>
<td>+ gentamycin</td>
<td>52.1* 15.8* 820.8* 2.4+ 3.6+ 8.5+</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>13.2* 61.2* 607.6* 0.1* 10.6+ 1.3+</td>
<td></td>
</tr>
<tr>
<td>+ gentamycin</td>
<td>9.8 65.8 646.2 0.1+ 12.0+ 1.3+</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>48.9* 82.3* 402.7* 1.2+ 41.8+ 48.2+</td>
<td></td>
</tr>
<tr>
<td>+ gentamycin</td>
<td>18.1* 88.9* 1612.1* 0.3+ 57.8+ 15.2+</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>63.1 32.8 2066.6 0.6* 4.6* 2.6*</td>
<td></td>
</tr>
<tr>
<td>+ A.M.</td>
<td>28.7+ 79.6+ 2283.1+ 0.7* 13.9* 3.0*</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>40.2 15.0 604.0 1.4* 4.0 5.7</td>
<td></td>
</tr>
<tr>
<td>+ A.M.</td>
<td>18.6+ 42.4+ 787.0+ 2.3+ 17.9+ 40.5+</td>
<td></td>
</tr>
</tbody>
</table>

A.M. = antibiotic mixture described in text

Table 8. Effectiveness of the mixture of penicillin G, streptomycin and chloramphenicol in inhibiting bacterial uptake of the labelled algal extract. Concentrations of the antibiotic used were those given in the text or double and treble these concentrations. Samples were differentially filtered through 1.0 μm and 0.2 μm pore size polycarbonate membranes; results are presented as percentages of total uptake of the control. Samples numbers refer to the same samples as those in Tables 7 and 9.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Particle size</th>
<th>Control</th>
<th>Standard</th>
<th>Antibiotic Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Standard</td>
<td>×2</td>
</tr>
<tr>
<td>5</td>
<td>&gt; 1.0 μm</td>
<td>46.1</td>
<td>22.8</td>
<td>23.1</td>
</tr>
<tr>
<td></td>
<td>0.2–1.0 μm</td>
<td>53.9</td>
<td>6.9</td>
<td>8.4</td>
</tr>
<tr>
<td></td>
<td>Total uptake</td>
<td>100.0</td>
<td>29.7</td>
<td>31.5</td>
</tr>
<tr>
<td>4</td>
<td>&gt; 1.0 μm</td>
<td>47.8</td>
<td>33.0</td>
<td>33.0</td>
</tr>
<tr>
<td></td>
<td>0.2–1.0 μm</td>
<td>52.2</td>
<td>14.2</td>
<td>14.2</td>
</tr>
<tr>
<td></td>
<td>Total uptake</td>
<td>100.0</td>
<td>47.2</td>
<td>47.2</td>
</tr>
</tbody>
</table>
Table 9. Percentage uptake of labelled algal extract and labelled glucose by the different size fractions of the nannoplankton, measured by differential filtration. Results presented as percentages of total uptake; sample numbers refer to the same samples as those in Tables 7 and 8.

<table>
<thead>
<tr>
<th>Sample</th>
<th>% uptake of labelled algal extract</th>
<th>% uptake of labelled glucose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.2-1.0</td>
<td>1.0-2.0</td>
</tr>
<tr>
<td>1</td>
<td>74.1</td>
<td>12.2</td>
</tr>
<tr>
<td>2</td>
<td>40.8</td>
<td>22.3</td>
</tr>
<tr>
<td>3</td>
<td>53.4</td>
<td>21.2</td>
</tr>
<tr>
<td>4</td>
<td>52.2</td>
<td>21.5</td>
</tr>
<tr>
<td>5</td>
<td>53.8</td>
<td>14.3</td>
</tr>
</tbody>
</table>

been in the particulate fraction under 1.0 μm, which was separated from the fraction containing microflagellates (mean width 2.32 μm) and detrital particles with attached bacteria during differential filtration. Uptake of labelled algal extract by the free bacterial fraction was almost fully depressed by the antibiotic mixture. Uptake by the larger particulate fraction was only partially depressed; since the activity of bacteria attached to detritus was probably also inhibited, this uptake must have been performed mainly by heterotrophic microflagellates. Doubling and trebling the concentration of the mixture did not change the results, and it is assumed that the concentration used in the heterotrophic uptake studies was therefore not toxic to eukaryotes and did not impair their metabolism.

The mean percentage of uninhibited assimilation of labelled glucose by particles >3.0 μm was 11.75% (range 3.4% to 38.4%), this being established by differential filtration during the routine sampling adjacent to the reef. Once again, this indicates that most of the uptake was performed by particles <3.0 μm, predominantly free bacteria, as opposed to the uptake performed by larger particles, including bacteria attached to detritus. Even though they had a mean width of 2.32 μm, few microflagellates would have been included in the free bacterial fraction, since their other dimensions would have prevented most of them from passing through a 3.0 μm pore size filter.

No doubt was left as to the heterotrophic role of the different size fractions of the nannoplankton in the differential filtration study in which samples were filtered through a number of filter sizes after incubation with labelled algal extract and with labelled glucose (Table 9). Free bacteria in the size range 0.2 μm to 1.0 μm were the most important, particularly in the case of glucose where they accounted for more than 75% of the uptake. Particles in the size range 1.0 μm to 2.0 μm were less important and were identified from SEM measurements as comprising large cocci. The other less active fraction consisted of particles >3.0 μm; microflagellates and detritus colonised by bacteria.

**Carbon Analysis**

The annual mean values obtained from carbon analyses were: TOC, 18.28 g m⁻³; DOC, 14.75 g m⁻³; by subtraction POC, 3.53 g m⁻³.

**Analysis of Debris**

The major component of debris samples collected adjacent to the reef during most of the year was seaweed, except after summer rains when rivers came down in flood and washed quantities of terrestrial macrophytes such as reeds into the sea (Table 10).

**Seasonal Means**

Seasonal means (Table 11) were calculated from the annual results by grouping data in seasons determined from monthly means of surf temperatures measured twice daily at Durban from 1957 to 1974. Most parameters increased slightly or markedly in spring and summer.

**DISCUSSION**

Uncertainty exists as to the exact nature of the results of ¹⁴CO₂ fixation experiments, whether they provide
Table 11: Seasonal means of various parameters measured during OR1 Reef study

<table>
<thead>
<tr>
<th>Season</th>
<th>Spring</th>
<th>Summer</th>
<th>Autumn</th>
<th>Winter</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Oct- Dec</td>
<td>Jan- Mar</td>
<td>Apr- Jun</td>
<td>Jul- Sep</td>
</tr>
<tr>
<td><strong>Temperature (°C)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Previous records (15 years)</td>
<td>22.0</td>
<td>21.4</td>
<td>19.8</td>
<td>20.2</td>
</tr>
<tr>
<td>OR1 Reef samples</td>
<td>22.4</td>
<td>24.4</td>
<td>22.4</td>
<td>20.2</td>
</tr>
<tr>
<td><strong>Bacteria</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Counts (× 10⁶ ml⁻¹)</td>
<td>2.05</td>
<td>2.13</td>
<td>1.91</td>
<td>1.93</td>
</tr>
<tr>
<td>% attached to particles</td>
<td>18</td>
<td>27</td>
<td>20</td>
<td>19</td>
</tr>
<tr>
<td>Dry biomass (mg m⁻³)</td>
<td>21.09</td>
<td>22.01</td>
<td>19.65</td>
<td>19.96</td>
</tr>
<tr>
<td><strong>14C glucose assimilation:</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$V_{max}$ (µg l⁻¹ h⁻¹)</td>
<td>1.30</td>
<td>0.54</td>
<td>0.48</td>
<td>0.13</td>
</tr>
<tr>
<td>T (h)</td>
<td>5.4</td>
<td>13.7</td>
<td>7.6</td>
<td>50.1</td>
</tr>
<tr>
<td>K + S (µg l⁻¹)</td>
<td>7.66</td>
<td>4.95</td>
<td>2.71</td>
<td>3.94</td>
</tr>
<tr>
<td><strong>Phytoplankton</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Production (mg C m⁻³ h⁻¹)</td>
<td>15.92</td>
<td>11.45</td>
<td>10.73</td>
<td>6.23</td>
</tr>
<tr>
<td>Counts (×10⁶ ml⁻¹)</td>
<td>5.1</td>
<td>5.4</td>
<td>2.4</td>
<td>1.5</td>
</tr>
<tr>
<td>Chlorophyll a (mg m⁻³)</td>
<td>2.23</td>
<td>2.20</td>
<td>1.77</td>
<td>2.29</td>
</tr>
<tr>
<td>Dry biomass (mg m⁻³)</td>
<td>178.4</td>
<td>175.8</td>
<td>141.7</td>
<td>183.5</td>
</tr>
</tbody>
</table>

The mean primary production of 12.82 mg C m⁻³ h⁻¹ (Table 1) measured adjacent to the OR1 Reef was comparable to that found at a station near the mouth of Richards Bay at which the conditions were marine (Hemens et al., 1971), and in Lake Sibaya which was the least productive of tropical freshwater lakes studied at the time (Allanson and Hart, 1975), both these localities being north of Durban in Natal. It was higher than that measured on the continental shelf off Durban by Burchall (1968) who measured a mean surface production of 1.52 mg C m⁻³ h⁻¹ (range 0.01 to 9.52 mg C m⁻³ h⁻¹) over a period of 5 years, but considerably lower than that which occurs in areas of upwelling such as the south west Cape coast. For example, daily rates recorded by Henry and Mostert (1977) at a station exposed to the sea in the mouth of Langebaan lagoon indicated approximate primary production of 24 mg C m⁻³ h⁻¹ in winter (April to August), 54 mg C m⁻³ h⁻¹ in spring (September to November) and 66 mg C m⁻³ h⁻¹ in summer (December to February). In adjacent Saldanha Bay an annual mean primary production of 55.7 mg C m⁻³ h⁻¹ (range 28.0 to 125.7 mg C m⁻³ h⁻¹) was estimated from chlorophyll analyses. Thus the primary production measured during this work was relatively low, but it is not possible to specify whether it was close to the gross or net production.

Horne (1969) lists coastal variations in chlorophyll a for different localities with the comment that they are generally variable and high (mostly > 5 mg m⁻³, range 1 to 91 mg m⁻³). In comparison, the chlorophyll a values measured adjacent to the OR1 Reef (Table 2) were low, once again being similar to those found by Allanson and Hart (1975) in the oligotrophic Lake Sibaya (2.6 to 2.8 mg m⁻³). From the chlorophyll a determinations the annual mean dry biomass of the phytoplankton was estimated to be 170.1 mg m⁻³ and total counts revealed that this biomass consisted largely of microflagellates. Primary production in Lake Sibaya was likewise associated with a greater density of microflagellates (Allanson and Hart, 1975), as was the case in the Gulf Stream (Hobbie et al., 1972). This is not surprising in the case of the OR1 Reef, where turbulence in the rough surf environment could injure larger phytoplankton.

Benthic seaweeds on the reef form a turf in small patches not colonised by suspension-feeders. They provide an input to the system in that they are closely cropped by fishes (Joubert, 1980; Joubert and Hanekom, 1980), and they appear inconspicuous because of this. Faecal products from these fish enter the detrital food chain, and many species digest only the epiphytic diatoms, the seaweed being excreted in a macerated state. Thus the benthic algal production on the reef was indirectly measured on entry into the water column with detritus imported into the system.

In general the bacteria were small, and most were free-living cocci not attached to particles (Tables 3 and 5). These findings are in agreement with recently published work on aquatic bacteria (Daley and Hobbie, Schleyer: Microorganisms and detritus of a subtidal reef 315.
With the exception of Watson et al., total bacterial counts were higher in the present study than in the cited works but with a smaller volume. This is attributable to the technique used, since more small bacteria are filtered and counted on 0.2 μm pore size polycarbonate membranes than on the 0.45 μm porosity cellulose matrix membranes used by all these authors except Watson et al. The mean diameter of cocci in this study was almost the same as the latter pore size, and many cocci would have been lost in the filtrate or matrix of the filter using cellulose membranes.

Hoppe (1978) cites \( V_{\text{max}} \) values of \(^{14}\text{C} \text{glucose assimilation} \) expressed in μg C h\(^{-1}\) for coastal and polluted areas as follows: Woods Hole Dock, 0.11; Tokyo Bay, 7.2; Pamlico River Estuary, 1.3 to 3.8 and Shimoda Bay, 9.2. When expressed in the same units, the annual mean and maximum uptake of this substrate by the ORI Reef samples (Tables 6 and 7), \( V_{\text{max}} = 0.263 \) and 1.247 μg C h\(^{-1}\) respectively, reveals that microbial activity adjacent to the reef approaches that of polluted areas.

Measurement of the uptake of simple compounds using the heterotrophic potential technique provides a comparative index and not an absolute measure of microbial heterotrophy (Hobbie et al., 1972). However, in evaluating the results of uptake of the labelled algal extract (Table 7), Schleyer, (1980) suggested the values for \( V_{\text{max}} \) were a realistic measure of microbial heterotrophic consumption, provided the labelled extract was representative of the environmental substrate. If this was the case, heterotrophic production on the reef exceeded the mean primary production by a factor of roughly 4 times. The significance of this is discussed later.

Most of the heterotrophic activity (\( > 50 \% \)) was performed by free bacteria smaller than 1 μm, this being demonstrated by the combination of differential filtration and antibiotic inhibition of bacteria in the uptake experiments (Tables 7 to 9). Similar findings are reported for marine bacteria by Williams (1970), Azam and Hodson (1977) and Cole and Likens (1979) who applied differential filtration techniques in uptake studies, using various labelled substrates in different areas. In the present study, glucose uptake was performed almost exclusively by particles smaller than 3.0 μm, as was found by Berman (1975) in the Gulf of California.

This activity of free bacteria is in contrast to the proposal of Wangersky (1977) to the effect that the small size of bacteria means that in the free state they are limited by Brownian movement to a small envelope of water which they impoverish before entering a dormant state, then needing up to six to twelve hours incubation at high nutrient levels for induction to activity. Turbulence on the ORI Reef is likely to preclude such dormancy. Stevenson (1978) further suggested that the prevalent small coccoid bacteria in the aquatic environment are in fact the dormant form. He made this controversial proposal with little supporting physiological evidence, welcoming agreement or disagreement from work it would provoke. Both he and Wangersky felt that dormancy would account for the enigmatic discrepancy between colony counts from plate cultures and total microscopic counts, the latter always greatly exceeding the former. This would appear to involve a contradiction, since if dormancy alone were involved in the discrepancy, on enriched culture the bacteria would re-enter an active state after a lag period, giving a 'full' plate count. As this is not the case, it seems more likely that the aquatic environment has not yet been adequately simulated in culture work.

Recently Kogure et al. (1979) have developed a technique in which direct viable counts of natural bacterial populations closer to total counts (\( > 35 \% \)) have been made possible by incubation of water samples with nalidixic acid and slight enrichment with yeast extract before performing AODC. This method provided 'a minimum estimate of viable cell numbers' approximately 1000 times greater than plate counts, further highlighting the inadequacy of the latter method.

Switching from dormancy to an active state would not only involve a lag phase but should also include increased respiration as substrate is oxidised for energy in returning to active metabolism. However, uptake of the labelled algal extract proceeded immediately on incubation and was linear over the incubation period, while little of the substrate was respired (Schleyer, 1980). In the same paper, only slight pleomorphic changes are reported as accompanying natural bacterial proliferation in an eight hour study, indicating little change from the predominant 'dormant form'.

Wangersky (1977) and Stevenson (1978) also contend that bacteria attached to particles are more active, because they live in a micro-environment richer in nutrients than surrounding water by virtue of nutrition intrinsic to particles or adsorbed to them. On a numerical pro rata basis, uptake of labelled substrates by attached bacteria was only slightly lower than uptake by their unattached counterpart in the antibiotic and differential filtration studies. This seems to indicate that attachment to particles did not afford bacteria near-independence from the surrounding water; their nutritional requirements were only slightly better met by the particulate substrate than their free counterparts more dependent on DOC, and they did not appear to
be more active. None of these results corroborates Wangersky’s and Stevenson’s theories of dormancy, and it appears that small coccoid bacteria are merely the most efficient morphological form for survival in an environment in which competition for substrates runs high.

The effective, non-competitive nature of the antibiotic mixture used in the inhibition studies (Fig. 4) proved non-toxic to eukaryotes despite the high application concentration. Since free bacteria were almost entirely inhibited, attached bacteria were probably similarly affected, and the heterotrophic uptake by larger particles was probably performed by eukaryotes, predominantly microflagellates (Tables 8 and 9).

Most phytoplankters appear capable of taking up organic substrates by passive diffusion only at concentrations much higher than natural concentrations (Wright and Hobbie, 1966) and the concentrations used in these experiments. However, true heterotrophic organic uptake has been detected in a small number of freshwater phytoplankters (Pollinger and Berman, 1976), and non-pigmented estuarine microflagellates incapable of organic heterotrophy were demonstrated to incorporate labelled organics indirectly by ingestion of bacteria which had assimilated the label (Haas and Webb, 1979). Whatever combination of the latter two mechanisms was involved in heterotrophic uptake by the OR1 Reef microflagellates, they accounted on average for 20% or less of the total uptake of organic substrates.

Seasonal variation in most of the parameters measured was not remarkable (Table 11). The increase in spring and summer of most of the data obtained was probably due to summer storms with their attendant detrital input by rough seas and flooded rivers. However, there was considerable variation in individual measurements that could not be related to the prevailing environmental conditions, since these were so changeable; a level of biological activity procured by a set of environmental conditions would often remain despite a change in the conditions. The greatest seasonal variation was found in the phytoplankton counts, but little accuracy was attributed to these counts (see Results).

At this stage, the results can be synthesised with the findings and theories of other workers in a description of the lower levels of the food chain on the OR1 Reef. The nutrient source for most of the heterotrophic activity in the OR1 Reef water column is undoubtedly seaweed and terrestrial plant debris (Table 10) imported into the system by wave action and currents, these constituting energy subsidies. This is not surprising, since Mann (1972) concluded in a review on the subject that over 90% of the extensive marine macrophytic production in coastal waters enters the detrital food chain as dissolved or particulate organic matter. The debris on the OR1 Reef is rapidly broken down by turbulent surf action and sand abrasion and soluble constituents rapidly leach out of it as it is broken into fine detrital particles. While still organically rich, these are probably heavily colonised by successional pleomorphic stages of bacteria and are rapidly decomposed into relatively inert particles with a reduced microbial flora (Oláh, 1972). As this stage is shortlived, particles as centres of intense metabolic activity are relatively sparse (Azam and Hodson, 1977) and probably insignificant in relation to the activity of the prevalent free bacteria. The remaining inert particles are enriched by the presence of colonising bacteria, which slowly break them down, improving their nutritional value for detritivores rather than the converse (Odum and de la Cruz, 1967; Mann, 1972). It is possible that grazing pressure on these particles by coarse suspension feeders might be the cause of the greater abundance of free bacteria (Sorokin, cited by Azam and Hodson, 1977).

Free bacteria are dependent on DOC as a food source, and carbon analysis revealed that this is more abundant than POC. Although some DOC can be absorbed directly by suspension-feeders (Jørgensen, 1976), it is likely that most of it becomes available to them as particles by way of a complex interaction of biological (bacterial) and physico-chemical processes to which the reviews of Riley (1970) and Mann (1972) provide a good background. The DOC pool appears to consist of two components, the first being an assimilable component to which bacterial activity is related, so that high concentrations of it rarely occur (Andrews and Williams, 1971; Sayler and Gilmour, 1978). In the OR1 Reef system this component would be largely derived from leaching of organics from the allochthonous seaweed and plant debris, hence the choice of a labelled algal extract as the substrate in measurements of bacterial activity (Schleyer, 1980). The other component is larger, consisting of accumulated dissolved organics resistant to rapid microbial degradation (Jørgensen, 1976).

In the surf environment, air bubbles are vigorously and constantly introduced into the water. Such bubbles collect surface-active organic material at their gas-liquid interface and, on bursting of the bubbles, organic particles form by aggregation (Riley, 1970; Mann, 1972; Wangersky, 1977) in a process which can even include viruses and bacteria known as 'jet drop enrichment' (Blanchard, 1978). These particles are in equilibrium in a normal distribution with larger particles formed by electrostatic attraction, causing combination of the smaller particles into micelles, the process being irreversible if the micelle components are cemented together by bacteria (Wangersky, 1977). At
18 °C in 0.6 M NaCl, Wangersky cites these micelles as being ellipsoidal and larger than 0.1 μm. Particles in seawater, including bacteria, always have an electro-negative surface charge of varying strength (Neihof and Loeb, 1972), by means of which they may obtain a further organic coating. In this way inorganic particles also become nuclei of organic accretion and by continued bubbling, aggregates up to and over 1 mm eventually form (Wallace and Duce, 1978). These are probably the common amorphous particles with few attached bacteria, including little recognisable organic detritus, categorised as flocs and flakes by Wiebe and Pomeroy (1972). They appeared to be the most common type of particle observed in the OR1 Reef samples, although they never attained the large size found by Wallace and Duce, probably owing to turbulence. It seems likely that they are derived largely from the unassimilable dissolved organics, since they are so poorly colonised by bacteria.

Paerl (1974, 1976) demonstrated that bacteria harvest considerable quantities of DOC and rapidly convert significant amounts of it to particulate (non-respired) organic carbon, depositing most of it as extracellular material. Since free bacteria exhibited the greatest heterotrophic activity in the OR1 Reef experiments, they probably selectively take up assimilable dissolved organics and constitute the most important mechanism by which it is converted into particulate matter. Although microflagellates are apparently also capable of a small amount of such heterotrophic activity, their major contribution is probably in primary production which was low.

The relative abundance and productivity of the photosynthetic and detrital inputs available to suspension-feeders are summarised in Table 12. It is obvious that, although the bacterial population has a low standing biomass, it has a high turnover rate, evidence for this being the high V_{max} recorded in the OR1 Reef experiments, corroborated by the short generation time (10 min) observed for at least one marine bacterial species in culture (Eagon, 1962). Since heterotrophic activity appears to exceed primary production quite considerably, the bacteria should attain a greater biomass than the phytoplankton. This does not occur, and the bacterial population must thus be limited by a combination of availability of food, the assimilable dissolved organics being rapidly consumed by the predominant free bacteria, and by grazing pressure. Bacteria would provide protein-rich food for suspension-feeders on the OR1 Reef if these are adapted to filtering such small filtering particles. If this is not the case, suspension-feeders are limited to filtering out phytoplankton (largely microflagellates) and the less nutritious detrital particles with their attached bacteria.

$\text{DOC} = 14.75 \times 10^3 \text{ mg C m}^{-3}, \text{ Calorific value} = 23.074 \text{ (mg C m}^{-3}\text{h}^{-1}), \text{ Production} = 47.60 \text{ (mg C m}^{-3}\text{h}^{-1})$
LITERATURE CITED


Smale, M. J., Buchan, P. R. (in prep.). The biology of Octopus vulgaris Cuvier off the east coast of South Africa


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