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Longitudinal and Vertical Trends of Bacterial Limitation by Phosphorus and Carbon in the Mediterranean Sea

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ABSTRACT

The effect of phosphate (P), nitrate (N), and organic carbon (C, glucose) enrichment on heterotrophic bacterial production was examined along two longitudinal transects covering the whole Mediterranean Sea during June and September 1999. During these cruises, integrated bacterial production ranged from 11 to 349 mgC m⁻² d⁻¹ for the 0–150 m layer. P was found to stimulate bacterial production (BP) in 13 out of 18 experiments, in the eastern and in the western Mediterranean Sea. Organic carbon stimulation of bacterial production was observed at two stations in the Alboran Sea, where the highest bacterial production was recorded (216 and 349 mg C m⁻² d⁻¹) and in the Sicily Strait. Maximum rates of alkaline phosphatase (AP) increased from the Alboran to the Levantine Sea whereas AP turnover time decreased. Moreover, alkaline phosphatase activity was not systematically reduced following additions of P. In cases of P limitation, however, the alkaline phosphatase activity to bacterial production ratio was severely reduced in the P and NPC enrichments. Generally, the addition of the limiting factor—whether P or C—had a synchronous stimulating effect on bacterial production and ectoaminopeptidase activity and induced a decline in the amino acid respiration percentage. At two selected stations in the eastern and northwestern Mediterranean, response to enrichment was tested on vertical profiles. Bacteria shifted from P to C limitation at a depth where soluble reactive phosphorus was still undetectable, but corresponding to a strong increase in alkaline phosphatase turnover time. Our results showed that values of AP turnover time lower than 100 h corresponded to situations of P limitation of bacterial production.

Introduction

Bacteria play an important role in the flow of energy and materials through planktonic food webs. As a consequence of their high abundance, efficient nutrient uptake, and potentially rapid growth rates, bacteria may act as either a sink or a source of nutrients for higher trophic levels. Mechanisms controlling bacterial abundance and biomass have been the subject of intensive research in contemporary microbial ecology [1].

Studies on heterotrophic bacterial dynamics have shown phosphorus limitation of bacterial growth in a variety of environments [28, 37, 41, 46] and in the open ocean [4, 10, 30]. In the Mediterranean Sea, which is one of the world's most oligotrophic sea, mineral nutrients and particularly phosphorus have been shown to limit both phytoplankton and bacteria in certain areas [6, 24, 35, 36, 46, 44, 45].

An interesting feature of the Mediterranean is a broad west–east gradient of oligotrophic conditions, as shown by lower primary production, [31, 40], and lower sedimentation fluxes from west to east [26]. The increased importance of the microbial food web in the eastern Mediterranean Sea [7, 11] is consistent with an increase in bacterial production to primary production ratios [3, 40]. In the eastern, relative to the western Mediterranean, there is an increasing difference between the nitracline and the phosphacline depths in stratified conditions during summer [24, 26], and higher N/P ratios of mineral nutrients in deeper layers reaching 20 to 25 in the Aegean Sea [39].

Although the west–east gradient of oligotrophy in the Mediterranean is often evoked [7, 11, 40], the key processes regulating the planktonic foodweb, such as nutrient control on bacterial production, transformations of dissolved organic matter, and their biochemical consequences, have not been investigated. Thus, while the problem of bacterial growth rate limitation is a very general one, the Mediterranean with its trophic gradient is an ideal place for its study. The starting point of this study was the hypothesis that there might be a gradient of nutrient limitation on heterotrophic bacterial activities, corresponding to the longitudinal decline in nutrient availability. We studied relationships between the two main indicators of P-limitation: alkaline phosphatase (AP) activity—which enables plankton to remove inorganic P from dissolved organic matter—and the effect of nutrient amendments (glucose, nitrate, and phosphate) on bacterial production.

In order to provide a full spatial coverage of the area, we examined these variables along two longitudinal transects in June–July and September 1999, which included a broad range of trophic conditions. We also investigated these variables down vertical profiles to assess whether there was a gradient for P-limitation in the euphotic zone. Finally, in order to test for the biogeochemical implications of nutrient limitation on the transformations of dissolved organic matter, we examined the effect of carbon and nutrient enrichments on amino acid fluxes (assimilation, respiration), and ectoenzymatic activity (phosphatase, aminopeptidase).

Materials and Methods

Study Sites and Sampling

This study was carried out during the Trans Mediterranean Cruise (TMC) from June 5 to July 4 1999 on the RV *Aegaeo*, and the PROSOPE cruise from September 2 to October 4 1999 on the RV *Atalante*, in the framework of the EC program MATER and France-Proofs program PROSOPE, respectively. Water samples were collected from 9 (TMC) + 15 (PROSOPE) stations, which were selected to represent the main basins or physical features of the Mediterranean Sea, crossing a broad gradient of oligotrophy (Fig. 1). The phytoplankton biomass during the TMC cruise ranged from 15 to 61 mg chlorophyll *a* m⁻² (Pagou and Gotsis, pers comm.) and from 10 to 49 mg chlorophyll *a* m⁻² during PROSOPE cruise (Ras, Marty and Claustre, pers. comm). One of the stations, located outside of the Mediterranean in the Morocco upwelling (St. UPW), was sampled as representative of eutrophic conditions (100 mg chlorophyll *a* m⁻², Table 1, Fig. 1). Water samples were collected using a rosette equipped with 12 L Niskin bottles around noon.

Nutrients

Water samples for the determination of nitrite and nitrate were collected in 100 ml polyethylene bottles and kept frozen (−20°C) until analysis in the laboratory. Nitrate and nitrite were analyzed by a Bran+Luebbe Autoanalyzer II, according to the procedure of Treguer and Le Corre [38]. The precision is estimated at ± 0.05 μM for nitrate and ± 0.02 μM for nitrite.

Soluble reactive phosphorus was measured according to the method of Murphy and Riley [27]. During the TMC cruise, samples were analyzed on board with a Perkin Elmer Lambda 2S UV/VIS Spectrophotometer. The precision and detection limits of this method were 20 nM. During the PROSOPE cruise, soluble reactive P was analyzed after a 6-fold concentration step following the magnesium-induced coprecipitation (MAGIC) procedure [20], which brings the detection limits down to 5 nM. To avoid contamination, only one bottle (polycarbonate, 60 ml) was

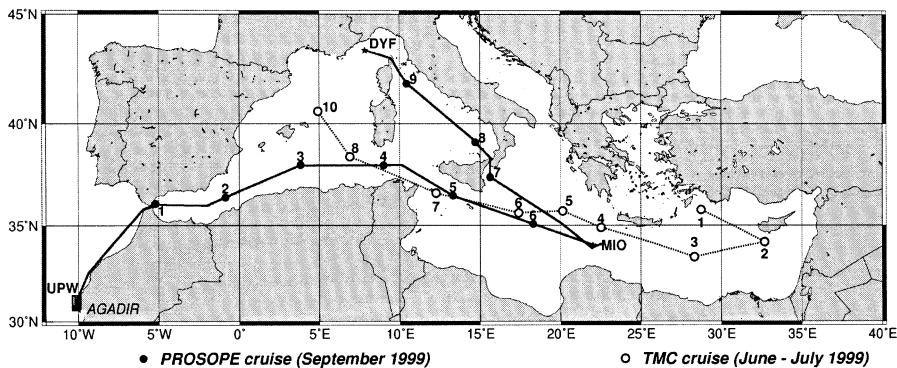


Fig. 1. TMC (June 1999) and PROSOPE (September 1999) cruise tracks with stations. Note that one station is located outside the Mediterranean, in the Marocco upwelling (St UPW) as representative of eutrophic conditions, while DYF (Dyfamed station, Ligurian Sea) and MIO (Ionian Sea) stations in Mediterranean were chosen to correspond to mesotrophic and highly oligotrophic conditions, respectively.

used for sampling, centrifugation, precipitate treatment, and analysis. Spectrophotometric analysis was performed with a CECIL CE 1011 spectrophotometer using a cell with a long optical length (10 cm) and small volume (10 ml). With this method precision was ± 3 nM. Dissolved and particulate organic phosphorus was determined using the persulfate wet-oxidation method [42].

Bioassays

Response of the bacteria to nutrient and/or carbon addition was examined in 15 experiments in the subsurface layer (TMC cruise, stations TM-01, -03, -06, -07, -08, and -10; PROSOPE Cruise, stations PRO-1 to PRO-9). During the PROSOPE cruise, enrichment experiments on vertical profiles were performed at three stations representative of eutro-, meso-, and oligotrophic conditions, abbreviated UPW (Marocco upwelling), MIO (Ionian Sea) and DYF (Dyfamed station, Ligurian Sea, Fig. 1).

Table 1. Station location

	Station	Date	lat.	Long.	Depth (m)
TMC cruise	TM-01	7/6/99	35°78 N	28°69 E	3800
	TM-02	10/6/99	34°15 N	32°67 E	2695
	TM-03	14/6/99	33°39 N	28°29 E	3082
	TM-04	16/6/99	34°89 N	22°53 E	2982
	TM-05	18/6/99	35°70 N	20°15 E	3150
	TM-06	20/6/99	35°62 N	17°39 E	4032
	TM-07	23/6/99	36°61 N	12°24 E	1313
	TM-08	27/6/99	38°40 N	6°89 E	2854
	TM-10	1/7/99	40°58 N	4°95 E	2756
PROSOPE cruise	UPW	10/09/99	31°00 N	10°01 W	100
	PRO-1	14/09/99	36°05 N	5°11 W	775
	PRO-2	15/09/99	36°24 N	0°51 W	2556
	PRO-3	16/09/99	37°59 N	3°50 E	2704
	PRO-4	17/09/99	37°59 N	8°31 E	2292
	PRO-5	18/09/99	36°28 N	13°19 E	1688
	PRO-6	19/09/99	35°04 N	18°17 E	3699
	MIO	21/09/99	33°58 N	22°02 E	2112
	PRO-7	26/09/99	37°23 N	15°36 E	2096
	PRO-8	27/09/99	39°06 N	14°42 E	3288
	PRO-9	28/09/99	41°53 N	10°26 E	1051
	DYF	30/09/99	43°24 N	7°51 E	2280

Seawater samples (250 ml during TMC cruise, 40 ml during PROSOPE cruise) were poured into five Nalgene acid-cleaned polycarbonate bottles. NaNO_3 , Na_2HPO_4 , and D-glucose were added to give final concentrations as follows: Co (control) was unamended, N received 2 μM nitrate solution, P received 0.25 μM phosphate solution, C received 1.66 μM glucose (i.e. 10 μM glucose-C) solution, and NPC received all three components. The corresponding stoichiometric atomic ratios were $\text{C/N} = 5$ and $\text{N/P} = 8$. After adding the nutrients, the bottles were placed in an on-deck incubator with circulating surface seawater and covered with a neutral screen providing 50% incident light. Following 24 h incubation, subsamples were removed from each bottle for bacterial production measurements. Samples for bacterial abundance, amino acid fluxes, and ectoaminopeptidase and phosphatase activity were also taken at stations TM-03, -06, -07, -08, and -10 during the TMC cruise.

Reproducibility tests run on three occasions in the eastern and western basins showed that the coefficient of variation of bacterial production response within triplicate bottles was on average 17% (max. 25%). Variability within triplicates for ^{14}C methodology (used for amino acid uptake rates and respiration) was higher, averaging 20%, whereas for potential ectoaminopeptidase activity it was lower, averaging 6%.

Bacterial Abundance

Water samples for bacteria, *Synechococcus*-like cyanobacteria, and heterotrophic and autotrophic nanoflagellates were preserved with tetraborate-buffered formalin (2% final concentration). Within 24 h, samples were filtered and stained with DAPI, and the slides were stored frozen until they could be counted using epifluorescence microscopy as described in Christaki et al. [6].

Bacterial Production

Bacterial production was estimated using ^3H -leucine. The classical filtration method was used during the TMC cruise [22, 43] whereas the centrifugation method [34] was used during the PROSOPE cruise. During the TMC cruise, 10 ml duplicates were amended with 1 nM [$4,5\text{-}^3\text{H}$]leucine (Amersham, 141 Ci mmol^{-1})

and 19 nM unlabeled leucine. A third sample was killed with 1% formalin prior to isotope addition and was used as the control. Samples were incubated in the dark at the ambient temperature of the depth of collection ($\pm 1^\circ\text{C}$) for 2 h, which was tested to satisfy linear incorporation with time. The incubations were stopped using formalin (1% final concentration). The samples were then filtered through 0.2 μm cellulose ester filters (Millipore type GS), extracted and rinsed three times with 5% trichloroacetic acid (TCA) and stored frozen. Filters were later dissolved with 0.5 ml ethyl acetate prior to the addition of 10 ml of PCS Amersham Scintillation cocktail.

During the PROSOPE cruise, 1.5 ml samples were dispensed into 2 ml screw cap microcentrifuge tube. A mixture of [4,5- ^3H]leucine (Amersham, 155 Ci mmol^{-1}) and nonradioactive leucine was added to final concentrations of 16 and 4 nM, respectively. Samples were incubated like in the filtration method. Incubations were stopped by the addition of 50% TCA to give a final concentration of 5% TCA. Fifty μl of a bovine serum albumin (BSA, Sigma) solution was added to facilitate precipitation of the proteins (final concentration 100 mg BSA l^{-1}). The samples were centrifuged at 16,000 g for 10 min. The supernatant was discarded and 1.5 ml of 5% TCA was added. The samples were shaken vigorously using a vortex mixer and centrifuged again. The supernatant was discarded and 1.5 ml of PCS scintillation cocktail (Amersham) was added. Radioactivity was analyzed by a Packard LS 1600 Liquid Scintillation Counter.

Bacterial production values in samples treated with BSA addition were on average $96 \pm 13\%$ ($n=21$) of those obtained with the filtration technique, and $73 \pm 9\%$ of the values obtained without the BSA addition [34]. Concentration kinetic experiments showed that isotopic dilution was negligible for the 20 nM leucine additions used during both cruises. We thus used a conversion factor equal to 1, yielding 1.5 kg C per mole leucine incorporated [22].

Alkaline Phosphatase Activity

Alkaline-phosphatase (AP) activity was measured fluorometrically using methylumbelliferone phosphate (MUF-P [17]). Stock solutions of 5 mM in methylcellosolve were stored at -20°C and diluted with boiled-filtered seawater prior to use. The linear increase in fluorescence in seawater with added MUF-P was measured over 9 h (excitation at 365 nm and emission at 460 nm) with a Hitachi F-2000 spectrofluorometer during the TMC cruise and on a Kontron SFM 23B during the PROSOPE cruise. Calibration curves were made with MUF standards. The kinetic coefficients were determined using a range of concentrations of MUF-P from 25 to 1000 nM, which is reasonable considering the dissolved and particulate organic phosphorus concentrations that were measured during the PROSOPE cruise (excluding station UPW, 160–265 nM and 4.2–45 nM, respectively). The following transformation of the Michaelis-Menten equation was used to determine kinetic coefficients:

$$S/V = (K + S_n)/V_m + S/V_m \quad (1)$$

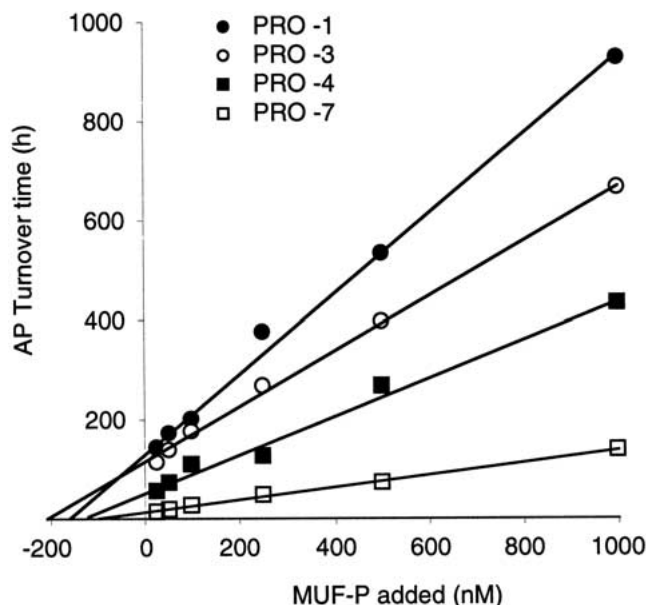


Fig. 2. Alkaline phosphatase (AP). Concentration dependence of turnover time, for stations PRO-1 (5m), PRO-3 (18m) PRO-4 (21m) and PRO-7 (35m). Fitted regression lines are shown.

where S is the MUF-P concentration and V the hydrolysis rate. Plotting S/V ratio versus S resulted in a straight line from which the maximum hydrolysis rate V_m (nM h^{-1}) is the inverse of the slope and the sum of the half saturation constant plus the natural concentration $K+S_n$ (nM) is the x -axis intercept (Fig. 2) and the turnover time is the y -axis intercept (expressed in h):

$$\text{Turnover time} = (K + S_n)/V_m \quad (2)$$

For depth profiles, we did not perform concentration kinetic experiments and thus neither V_m of AP nor $K + S_n$ (Eq. 1) values were measured. Alkaline phosphatase activity was measured along MIO and DYF profiles with only one concentration of added MUF-P (25 nM). From a comparison made on surface layers of the stations of the transect with turnover times ranging 14 to 182 h, we estimated that the turnover time measured with 25 nM MUF-P additions was only 5% higher than the value obtained from concentration kinetics ($n=11$, slope = 1.05 ± 0.01 , $r=0.99$). This 5% difference is close to the error of a turnover time estimated with a concentration kinetic (Table 1) and thus, on depth profiles, we assumed that the turnover times are reasonably correct. From Eq.(2), V_m can be calculated from the turnover time and an estimate of $K+S_n$, as discussed in the results section.

Alkaline phosphatase activity (with 0.25 μM MUF-P addition) was also measured on the bioassays processed at stations TM-03, -06, -07, -08, and -10.

Ectoaminopeptidase Activity

The ectoenzymatic aminopeptidase activity was estimated by measuring the hydrolysis rate of L-leucine-4-methyl-7-coumarinylamide (leu-MCA, Sigma), which is used as an analog substrate

for natural peptides [17]. Stock solutions of 10 mM leu-MCA were prepared in methylcellosolve and kept at -20°C . Potential ectoaminopeptidase activity was measured with a substrate addition of 200 μM . This concentration represented the saturation level as determined with preliminary concentration kinetics. The increase of fluorescence was measured with the excitation set at 380 nm and the emission at 440 nm. Incubation was performed at *in situ* temperature in the dark. Blanks were run by adding the leu-MCA to filtered boiled seawater and were shown to be insignificant. Relative fluorescence units were calibrated with 4 methyl-coumarin standards. Potential ectoaminopeptidase activity (PEA, expressed in nmol leu hydrolyzed $\text{L}^{-1} \text{h}^{-1}$) was measured along the surface layers studied for the longitudinal transects and on the bioassays processed at stations TM-03, -06, -07, -08, and -10.

In order to estimate the size distribution of bacterial activity, a size fractionation experiment was performed at station DYF for a 15 m water sample. Reverse filtration with in a Teflon apparatus without vacuum was used [2]. Bacterial production, alkaline phosphatase activity, and ectoaminopeptidase activity were measured in each fraction: total, $<2 \mu\text{m}$, $<0.6 \mu\text{m}$, and $< \text{GF/F}$ (glass fiber filter) in triplicate.

Amino Acid Uptake and Respiration

Potential assimilation rates and percentage of respiration of amino acids (PAA and % resp, respectively) were measured in subsamples taken from the bioassays, 24 h after enrichments, at stations TM-03, -06, -07, -08, and -10. PAA and % resp were

measured by adding a mixture of ^{14}C -amino acids at saturating substrate concentrations (final concentration of 200 nM) to 20 ml duplicates and a formalin killed control. The ^{14}C amino acid mixture (Amersham, 52 mCi mmole C^{-1}) included 16 amino acids, and their individual proportion (by activity) ranged from 0.6 (meth) to 13% (leu). Samples were incubated for 2 hours at *in situ* temperature in the dark and incubation was stopped by the addition of formalin (1% final concentration) buffered with sodium tetraborate. Following fixation, the samples were filtered through 0.2 μm polycarbonate Nuclepore filters, which were then rinsed 3 times with 0.2 μm filtered seawater.

The filtrates were acidified (200 μl HCl, 6 N) and the released ^{14}C - CO_2 was flushed using N_2 (100 ml min^{-1}) for 10 min. CO_2 was trapped in two successive scintillation vials each containing 9 ml of a cocktail of ethanolamine-methanol-scintillation liquid (PCS, Amersham) in a ratio 1:1:7 v/v [13] and counted on a Packard 1600 TR scintillation counter.

Results

Water Column Structure and Trophic Gradient

In June–July, mixed layer depths at the stations investigated ranged from 6 to 25 m (Table 2). The lowest values of bacterial production and nutrients were measured in the Irapetra anticyclonic gyre in the Levantine basin (Table 2; stations TM-02 and TM-03). At these two sta-

Table 2. Main characteristics of the stations (depth of the mixed layer, depth of the chlorophyll maximum (CHL MAX), depth of the 50 nM soluble reactive phosphorus (SRP) level

		Mixed layer depth (m)	Chl max depth (m)	0–150m IBP (mg C m ⁻² d ⁻¹)	0–150m I-SRP (mmol m ⁻²)	Depth of 50 nM SRP (m)
TMC cruise	TM-01	12	115	24.5	6.5	128
	TM-02	8	140	11.4	3.3	264
	TM-03	15	135	16.4	3.2	228
	TM-04	10	90	29.8	5.4	127
	TM-05	12	65	35.2	6.7	114
	TM-06	6	95	18.7	3.4	220
	TM-07	25	95	50.4	4.4	122
	TM-08	15	70	70.7	15.3	58
	TM-10	20	62	72.1	7.8	106
PROSOPE cruise	UPW	5	10	394 ^b	63.0 ^b	surf ^c
	PRO-1	5	25 & 60	349	26.8	57
	PRO-2	15	45	216	25.2	42
	PRO-3	18	55 & 95	121	12.1	41
	PRO-4	21	75	49.2	12.3	76
	PRO-5	15	55 & 65	71.3	8.9	77
	PRO-6	31	95	52.6	3.9	155
	MIO	30	95	62.6	2.1	171
	PRO-7	35	70	68.8	3.1	157
	PRO-8	20	70	53.2	5.9	112
	PRO-9	18	65	56.2	9.8	91
	DYF	20	45	76.5	21.4	63

^aIntegrated bacterial production (IBP), integrated SRP (I-SRP).

^bUPW station was integrated 0–90m.

^cfor UPW all SRP concentrations were >100 nM.

tions, the chlorophyll maximum was recorded at 140 and 135 m while a concentration of 50 nM of soluble reactive P was recorded at 264 and 228 m, respectively. Bacterial production ranged from 11.4 (TM-02) to 72.1 (TM-10, 4.95°E, North of Balearic Islands) mg C m⁻² d⁻¹, and integrated soluble reactive P ranged 3.3 to 15.3 mmol m⁻².

In September, the depth of the mixed layer varied from 5 to 35 m. The chlorophyll maximum depth increased from the west (10°W) to the east (22°E) from 25 to 95 m. Integrated soluble reactive P decreased from 26.8 to 2.1 mmol m⁻² from west to east and the depth where 50 nM soluble reactive P was reached deepened from 40 to 170 m. In the Tyrrhenian Sea (stations MIO to DYF, Fig. 1), the integrated soluble reactive P remained low (2–6 mmol m⁻²), with slightly higher values at the North Tyrrhenian Sea (PRO-9) and DYF Stations (10 and 21 mmol soluble reactive P m⁻², respectively). Bacterial production decreased from 349 to 59 mg C m⁻² d⁻¹ from the east to the central Mediterranean (from PRO-1 to PRO-6, Fig. 1) and increased along the Tyrrhenian sea transect up to 76 mg C m⁻² d⁻¹ (DYF station). The eutrophic status of the station UPW in the Maroco upwelling was evident from the high values of integrated soluble reactive P (63 mmol m⁻²) and bacterial production (394 mg C m⁻² d⁻¹), which

were clearly out of the range of values measured in the Mediterranean.

Combining all the data from the Mediterranean stations (i.e., excluding the station UPW), integrated bacterial production correlated with chlorophyll ($r = 0.51, p < 0.05$) and with soluble reactive P ($r = 0.73, p < 0.001$) whereas correlation was insignificant between chlorophyll and soluble reactive P ($r = 0.3, p > 0.05$).

Environmental and Biological Variables in Surface Layer

During both cruises, soluble reactive P concentrations in the surface layer were undetectable (excluding UPW station). The sum of nitrate + nitrite ranged from 0.18 to 0.45 μM in June and 0.03 to 0.14 μM in September.

The number of *Synechococcus*-like cyanobacteria increased from the east to the west Mediterranean Sea during both cruises, with slightly higher abundances in September (from 2.9 to 13 × 10³ ml⁻¹ and from 9.6 to 85 × 10³ ml⁻¹, respectively; Table 3). Excluding the UPW station outside the Mediterranean, phototrophic nanoflagellates, heterotrophic nanoflagellates and bacterial abundances varied 5-fold (0.4 to 2.2 × 10³ ml⁻¹, 0.5 to 2.3 × 10³ ml⁻¹, and 2.4 to 10.7 × 10⁵ ml⁻¹, respectively; Table 3).

Table 3. Biological variables at the depths sampled for bioassays and/or APA kinetics

	Station	Depth (m)	TEMP (°C)	SAL (PSU)	CYANO × 10 ³ mL ⁻¹	PNAN × 10 ³ mL ⁻¹	HNAN × 10 ³ mL ⁻¹	NB × 10 ⁵ mL ⁻¹	BP ngCL ⁻¹ h ⁻¹	PEA nmol L ⁻¹ h ⁻¹
TMC cruise	TM-01	3	22.7	39.01	4.3	0.94	0.77	2.68	10.0	1.1
	TM-02	3	24.2	38.94	3.6	0.64	0.51	2.81	21.7	6.1
	TM-03	3	22.9	38.76	2.9	0.59	0.65	2.38	15.6	4.6
	TM-04	3	23.9	38.60	4.6	0.89	1.00	2.6	14.5	5.2
	TM-05	3	24.9	38.59	4.4	1.03	0.84	2.72	18.8	4.4
	TM-06	3	24.5	38.54	4.1	0.72	1.09	2.71	16.8	5.2
	TM-07	3	21.3	37.28	11.0	1.05	1.27	5.72	26.9	6.8
	TM-08	3	22.2	37.44	13.0	1.05	1.64	5.73	40.1	6.0
	TM-10	4	22.7	37.49	7.7	1.99	0.82	5.50	33.7	9.3
	UPW	16	16.5	36.11	6.7	2.20	2.47	14.9	1153	70
PROSOPE cruise	PRO-1	11	20.3	36.69	84.8	1.16	1.03	10.7	110.8	15.1
	PRO-2	15	24.2	36.78	40.3	0.76	1.65	7.68	74.6	12.8
	PRO-3	10	25.4	36.77	21.9	0.75	0.51	7.64	65	10.6
	PRO-4	13	25.5	37.86	12.3	0.40	1.27	5.85	34.8	8.6
	PRO-5	11	24.5	37.93	16.1	0.51	1.03	6.68	43.8	5.4
	PRO-6	10	27.4	38.64	9.6	0.72	1.13	4.45	30.1	7.8
	MIO	15	25.8	38.89	10.7	0.67	1.36	4.61	29.1	5.0
	PRO-7	14	25.6	38.53	19.8	0.61	1.17	5.29	40.6	8.2
	PRO-8	15	25.3	38.02	21.9	0.68	1.33	7.33	24.9	14.1
	PRO-9	15	23.4	38.14	16.4	1.19	2.01	6.20	41.2	9.9
	DYF	10	22.3	38.32	47.2	1.13	2.27	8.21	42	10.1

TEMP: temperature, SAL: salinity, abundances of CYANO: *Synechococcus*-like cyanobacteria, PNAN: phototrophic nanoflagellates, HNAN: heterotrophic nanoflagellates, NB: heterotrophic bacteria, BP: bacterial production, PEA: potential ectoaminopeptidase activity (measured with 200 μM MCA-leu additions)

Bacterial production varied 10 fold, from 10 to 110 ng C L⁻¹ h⁻¹, whereas surface water temperature ranged from 20 to 27°C (Table 3). Potential ectoaminopeptidase activity was 1.1 nmol L⁻¹ h⁻¹ at station TM-01 and 9.3 nmol L⁻¹ h⁻¹ at station TM-10, but there was not a clear east-west gradient (mean 5.4 ± 0.8 nmol L⁻¹ h⁻¹ for stations TM-02 to TM-08). In September, values ranged from 5 to 14 nmol L⁻¹ h⁻¹; decreasing from the Gibraltar Strait to station MIO and increasing again toward station DYF. The outlying status of the UPW station was characterized by most variables: lower temperature, higher abundances of flagellates (both auto and heterotrophs), bacteria, bacterial production, and aminopeptidase activities.

In Situ Kinetics of Alkaline Phosphatase Activity

Turnover time of alkaline phosphatase (AP) decreased from the west to the east Mediterranean, i.e., from 182 h at station PRO-2 to 11 h at station PRO-7, whereas it reached 786 h at station UPW (Table 4). Maximum rates of AP ranged from 0.62 to 12 nmol L⁻¹ h⁻¹, including a value of 1.1 nmol L⁻¹ h⁻¹ at station UPW. K+ Sn (Eq. 1) ranged from 76 to 206 nM in the Mediterranean surface samples and reached 848 nM at station UPW. K+Sn values of AP activity varied to a much lesser degree than AP maximum rates (Table 4). We computed a mean K+Sn of 96 ± 13 nM for stations PRO-5 to PRO-9 and a mean of 183 ± 20 nM for stations PRO-1 to PRO-3.

Table 4. Kinetic parameters of alkaline phosphatase.^a TT: turnover time, Vm: maximum velocity, K+Sn: X intercept of the regression (see Methods).

		Depth		Vm (nmol	K + Sn
Station		(m)	TT (h)	L ⁻¹ h ⁻¹)	(nM)
TMC cruise	TM-02	3	84 ± 9	1.65 ± 0.05	139 ± 18
	TM-04	3	31 ± 5	3.0 ± 0.2	91 ± 25
	TM-07	3	121 ± 26	0.62 ± 0.02	76 ± 18
PROSOPE cruise	UPW	15	786 ± 61	1.1 ± 0.2	848 ± 165
	PRO-1	11	135 ± 13	1.25 ± 0.04	168 ± 21
	PRO-2	15	182 ± 17	0.96 ± 0.03	175 ± 20
	PRO-3	10	114 ± 6	1.80 ± 0.04	206 ± 15
	PRO-4	12	51 ± 10	2.6 ± 0.2	133 ± 31
	PRO-5	11	52 ± 8	1.63 ± 0.04	86 ± 14
	PRO-6	10	31 ± 3	2.60 ± 0.04	83 ± 8
	MIO	15	27 ± 4	4.0 ± 0.1	111 ± 16
	PRO-7	14	11 ± 1	7.91 ± 0.09	88 ± 6
	PRO-8	15	21 ± 2	4.57 ± 0.08	95 ± 9
	PRO-9	15	19 ± 2	5.9 ± 0.2	114 ± 13
	DYF	10	14 ± 1	12.6 ± 0.3	178 ± 13

Response to Enrichment

Longitudinal Trends. Bacterial numbers for the enrichment experiments during the TMC cruise revealed no significant increase in the unamended control after 24 h incubation. Some increase in bacterial abundance was observed in P and C amended bottles at TM-06 (1.5-fold and 2.4-fold, respectively). NPC enrichment resulted in significant increase in bacterial numbers at stations TM-03 (× 2.3), and TM-06 (× 3.3).

Bacterial production showed an increase of × 4.4 on average (range from × 1.6 to × 7.5) in the unamended controls after 24 h of incubation (Table 5). The highest increase in BP relative to the control was recorded in NPC amended bottles, varying from × 2.0 (station UPW) to × 40 (station TM-10). Bacterial production increase relative to the control varied from × 0.9 to × 4.6 in P, from × 0.7 to × 2.1 in N, and from × 0.7 to × 4.7 for C amendments.

Overall, positive effects for P were detected at 13 of the 18 stations, and for C at 5 of the 18 stations, while N stimulated bacterial production only at stations PRO-1 and PRO-8 (× 1.9 and × 2.1, respectively). Finally, bacterial production was stimulated by addition of two different nutrients at three stations (PRO-1, -6, and -8, Table 5) and by addition of the three only at station PRO-2.

Table 5. Bacterial production in the different bioassays along longitudinal transects of TMC and PROSOPE cruise

Station Depth (m)			xP	xN	xC	xNPC
TMC cruise	TM-01	3	3.1	0.8	1.1	16.9
	TM-03	3	3.0	0.8	0.9	16.9
	TM-06	3	1.1	1.3	2.6	2.1
	TM-07	3	1.3	1.1	4.7	3.4
	TM-08	3	2.7	1.1	1.2	10.0
	TM-10	3	2.0	1.3	1.1	40.0
PROSOPE cruise	UPW	15	0.9	0.7	1.6	2.0
	PRO-1	11	1.0	1.9	1.6	5.2
	PRO-2	15	1.0	0.8	0.9	5.7
	PRO-3	10	1.4	1.3	0.9	4.7
	PRO-4	12	2.1	0.8	0.7	8.7
	PRO-5	11	4.0	0.8	0.9	17.7
	PRO-6	10	4.6	0.9	2.5	19.1
	MIO	15	2.9	1.1	0.9	11.8
	PRO-7	14	1.8	1.2	0.8	7.9
	PRO-8	15	2.8	2.1	1.0	11.9
	PRO-9	15	3.3	1.1	0.8	11.1
	DYF	10	4.3	0.9	0.9	13.0

Bacterial production after 24 hours in P alone, N alone, C alone or the NPC enrichment is relative to the bacterial production after 24 h in the unenriched control (xP, xN, xC, xNPC, respectively). P: addition of 0.25 μM SRP, N: addition of 2 μM nitrate, C: addition of 10 μM C-glucose, NPC: addition of the three.

UPWELLING

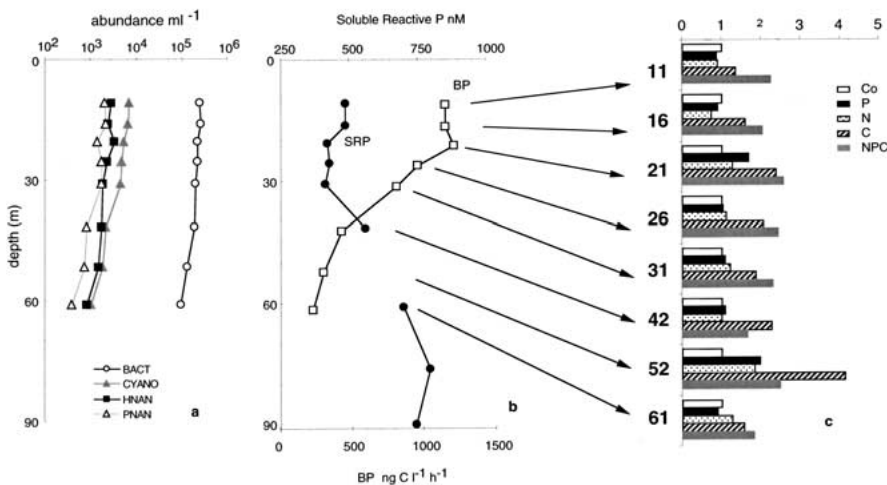


Fig. 3. Depth profiles of station UPW. (a) Abundance of heterotrophic bacteria (BACT), *Synechococcus*-like cyanobacteria (CYANO), heterotrophic nanoflagellates (HNAN), and autotrophic nanoflagellates (PNAN). (b) Soluble reactive phosphorus (SRP) and bacterial production (BP). (c) Bacterial production response in bioassays relative to the control, after 24 h incubation. Co: control, P: phosphorus enriched, N: nitrate enriched, C: glucose enriched, NPC: threefold enrichment. The results are normalized to the control, with values for Co set to 1.

Vertical Trends. At UPW station, the mixed layer extended to 5 m depth, the 1% light penetration depth was at 61 m, and the chlorophyll maximum was at 10 m. Soluble reactive phosphorus was detectable at all depths, ranging from 0.48 to 0.79 μM (mean $0.57 \pm 0.16 \mu\text{M}$, Fig. 3). While bacterial abundance was relatively constant with depth, bacterial production decreased below 20 m depth. Organic carbon addition stimulated bacterial production at all depths. Bacterial production increased following C and NPC additions, and ranged from $\times 1.4$ to $\times 4.1$ and from $\times 1.6$ to $\times 2.6$ relative to the control, respectively (Fig. 3).

At MIO station the mixed layer extended to 30 m depth, the 1% light penetration depth was at 131 m, and the chlorophyll maximum was at 95 m. Bacterial production

values at this station were low and decreased with depth. Soluble reactive P was undetectable in the surface and increased slowly with depth (50 nM concentration at 171 m, Fig. 4). Minimum values of turnover time of alkaline phosphatase were detected in surface layers (25 h) and increased with depth.

At DYF station the mixed layer extended to 20 m depth, the 1% light penetration depth was at 70 m, and the chlorophyll maximum was at 45 m. Bacterial production was slightly higher at station DYF compared to MIO (Table 2) and showed a maximum within the chlorophyll maximum (Fig. 5). Soluble reactive P was undetectable in the surface and increased sharply around 60 m depth. The AP turnover time was minimum in the surface and in-

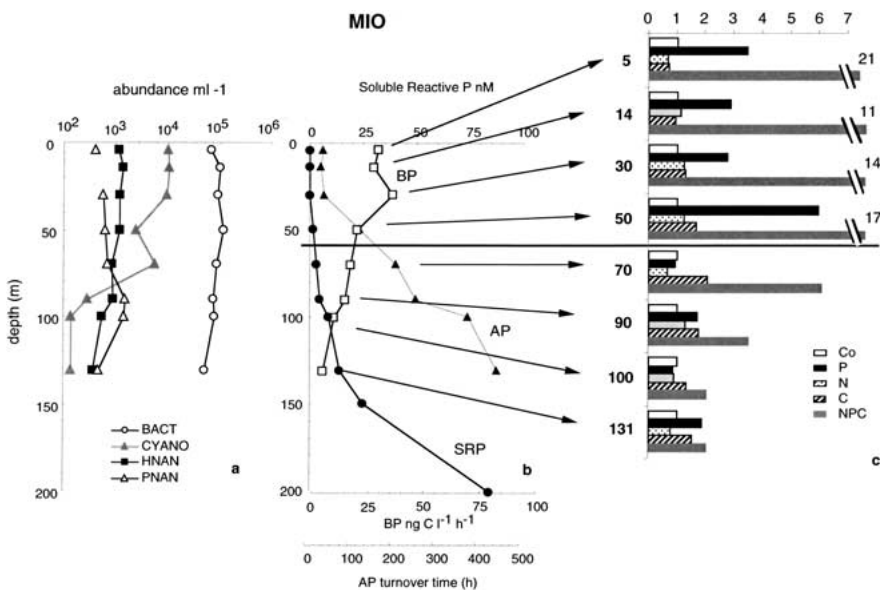


Fig. 4. Depth profiles of station MIO. Legend as in Fig. 3 except that the vertical profile of the AP turnover time is also plotted.

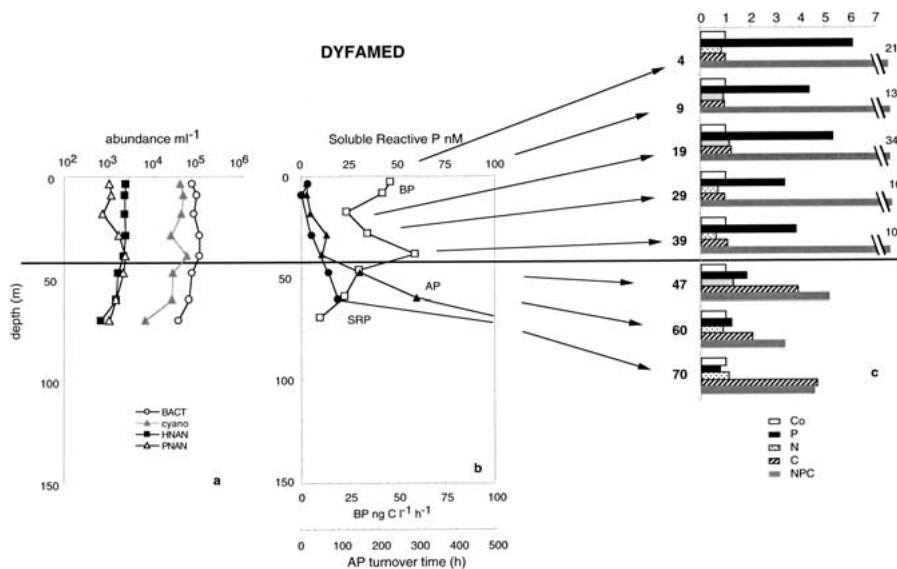


Fig. 5. Depth profiles of station DYF. Legend as in Fig. 3 except that the vertical profile of the AP turnover time is also plotted.

creased with depth, but this increase was sharper and occurred at a shallower depth than at the MIO station.

At stations MIO and DYF, addition of P alone increased bacterial production in the surface layer (Figs. 4 and 5) and glucose addition stimulated bacterial production in deeper layers (Figs. 4 and 5). At both stations DYF and MIO, the depth of transition from phosphorus to carbon limitation for bacterial production coincided with a significant increase in AP turnover time (more than 100 h) and was shallower than the depth for significant soluble reactive P increase (Figs. 4 and 5).

Bacterial production increase, relative to the control, following NPC additions were generally higher at stations MIO and DYF than at station UPW (Figs. 3, 4, 5). This increase was particularly important for the upper layers where addition of P alone stimulated bacterial production.

On vertical profiles, we did not perform concentration kinetics measurements. Estimates of V_m were determined using Eq. (2) and assuming constant values for $K+S_n$. The latter assumption was reasonable as results from surface samples along the horizontal transect, where the enzymatic properties of alkaline phosphatase activity were measured, showed that turnover time and V_m varied to a wider extent than $K+S_n$. The two sets of $K+S_n$ values obtained along the horizontal transect (96 ± 13 and 183 ± 20 nM) corresponded to C and P limitation of bacterial production, respectively, as revealed from enrichment experiments (see above, Table 2 and Table 4). We thus extrapolated V_m of AP in depth profiles from Eq. (2), with a constant $K+S_n$ value of 96 nM for depths where bacterial production was stimulated by P additions, and a value of 183 nM for depths where bacterial

production was stimulated by glucose. Based on these assumptions, V_m of AP decreased with depth, from 3 nM h^{-1} at 5 m to 0.42 nM h^{-1} at 130 m at station MIO and from 6 nM h^{-1} at 5 m to 0.07 nM h^{-1} at 70 m at station DYF.

Effects of Nutrient and Glucose Amendments on Ecto-enzymatic Activities and Utilization of Amino Acids

Potential ectoaminopeptidase activities measured after 24 h in enriched and control bottles ranged from 4 to $264 \text{ nmol L}^{-1} \text{ h}^{-1}$. Results at stations TM-03, TM-07, and TM-08 suggested that addition of the limiting factor had a synchronous stimulating effect on bacterial production and ectoaminopeptidase activity. However, this was not the case at stations TM-06 and TM-10 (Figs. 6a, 6b). Potential assimilation rates of amino acids after 24 h in enriched and control bottles ranged from 0.5 to $45 \text{ nmol L}^{-1} \text{ h}^{-1}$. Amino acid assimilation (relative to the control) followed the same trend as bacterial production: P stimulation at stations TM-03, -08, and -10 and C stimulation at station TM-06 and TM-07 (Figs. 6a, 6c). Respiration percentages of amino acids decreased in all the NPC amended bottles compared to those obtained in the unamended control (Fig. 7). Moreover, respiration percentage of amino acids decreased in response to the same factors limiting bacterial production: i.e., at TM-06 and TM-07, it decreased in C amended bottles and at TM-03, TM-08, TM-10 in P amended bottles.

The ratios of potential ectoaminopeptidase activity to bacterial production (PEA/BP) and potential amino acid assimilation rates to bacterial production (PAA/BP) were

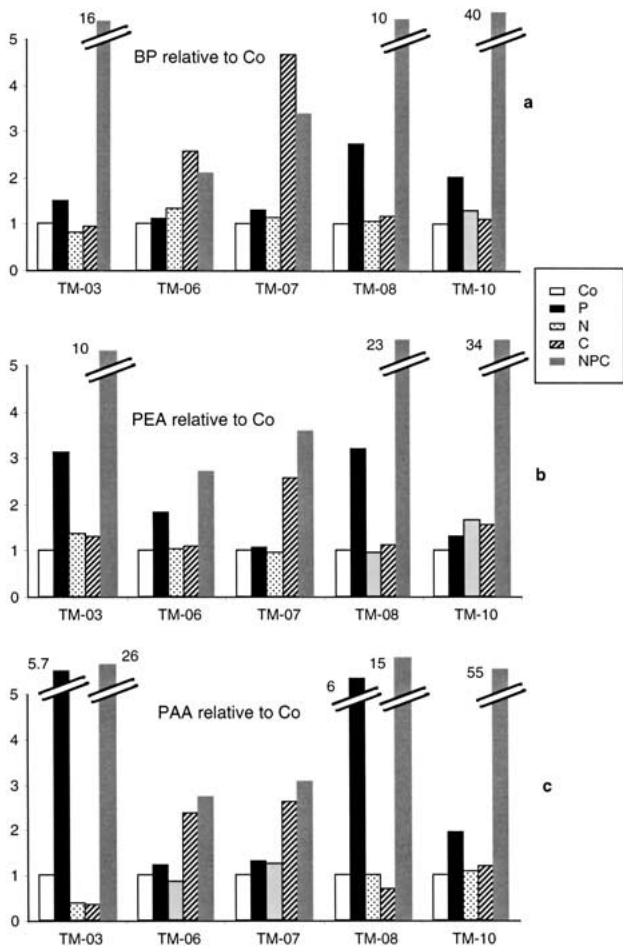


Fig. 6. (a) Bacterial production (BP), (b) potential ectoamino-peptidase activity (PEA), and (c) potential amino acid assimilation rates (PAA) in enrichment experiments during TMC cruise. Results are expressed relative to the unamended control. Co: control, P: phosphorus enriched, N: nitrate enriched, C: glucose enriched, NPC: threefold enrichments.

compared to those in the unamended control. PEA/BP ratios relative to the unamended control ranged from $\times 0.55$ to $\times 2.3$ (Fig. 8a) and PAA/BP ratios relative to the control

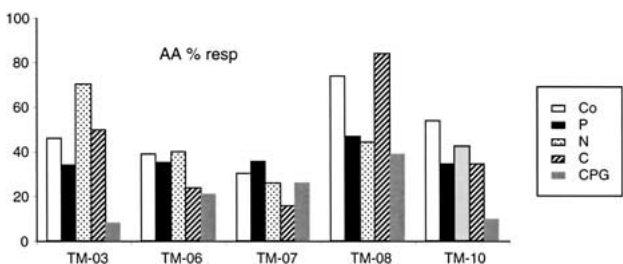


Fig. 7. Enrichment experiments during TMC cruise, station TM-03, TM-06, TM-07, TM-08, TM-10. Percentage of respiration of amino acid after 24 h incubation.

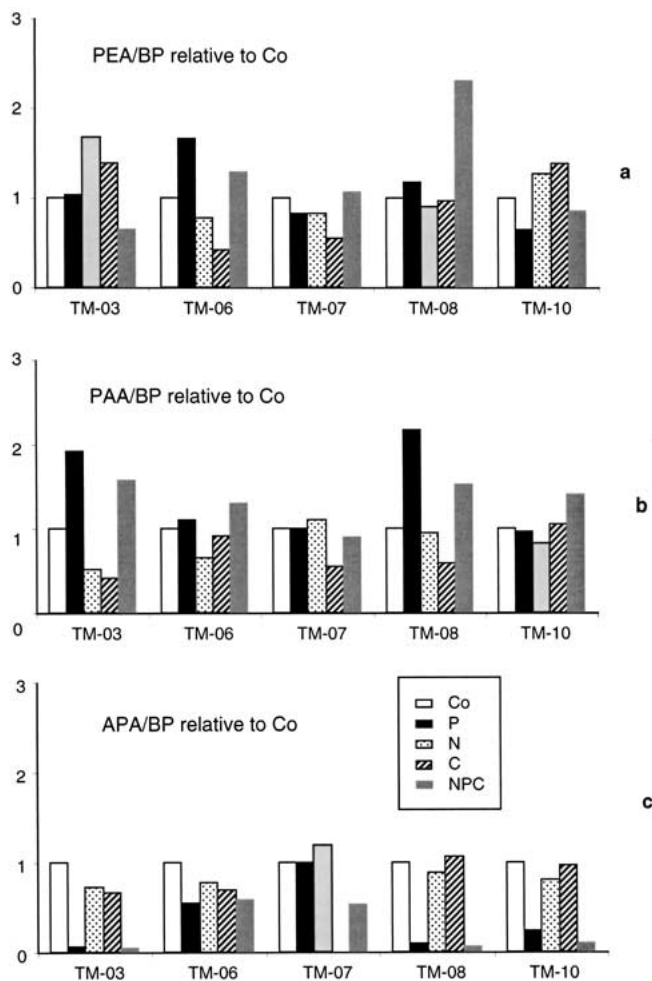


Fig. 8. Ratios of PEA/BP, PAA/BP and APA/BP relative to the same ratios in the control measured after 24 h incubation. PEA: potential ectoamino-peptidase activity PAA: potential assimilation rates of amino acids APA: alkaline phosphatase activity and BP: bacterial production.

ranged from $\times 0.41$ to $\times 2.2$ (Fig. 8b). In C amended bottles, the PEA/BP ratio was reduced about 50% when glucose stimulated bacterial production, whereas it was maintained at the same levels or even increased when phosphorus stimulated bacterial production. In two out of the three cases where P stimulated bacterial production, the PAA/BP ratio also increased relative to the control (TM-03, TM-08).

Relationships between Alkaline Phosphatase Activities and Response of Bacterial Production to Enrichments

AP turnover time was negatively related with the bacterial production response in P amendments. The data resulted in a significant $Y = aX^{-b}$ relationship ($r^2 = 0.47$, $n = 26$, Fig. 9). Interestingly, the four cases of simultaneous stimulation of

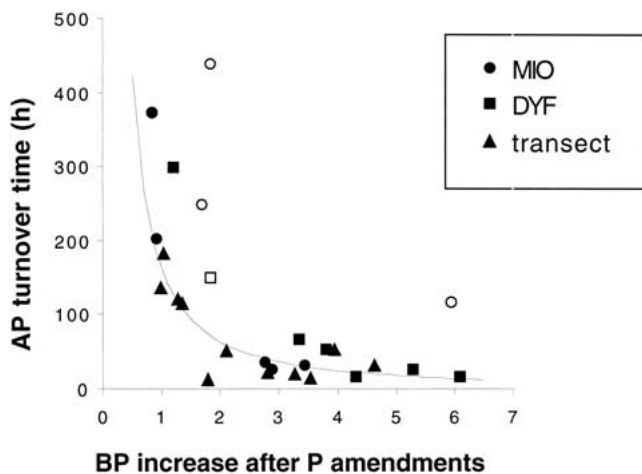


Fig. 9. Relationships between turnover time of alkaline phosphatase (AP turnover time) and bacterial production in P-amended bottles relative to the control (BP increase after P amendments). Open symbols coincide with bioassays situations where bacterial production increased both after addition of P alone and C alone. The relationship represented was computed excluding these double limitation cases: $y = 163 \times x^{-1.37}$, $r^2 = 0.64$, $n = 22$.

bacterial production in P and C amended flasks (MIO profile: 50 m, 90m, and 130 m; DYF profile: 50 m) are outliers of the relationship (Fig. 9). Excluding these outliers, the significance of the regression increased ($r^2 = 0.64$, $n = 22$).

When P stimulated bacterial production (Fig. 6a), AP activity to BP ratios (relative to the control) were severely reduced in P and in NPC enrichments (TM-03, TM-08, TM-10, Fig. 8c). In contrast, when bacterial production was stimulated by glucose (TM-06, TM-07, Fig. 6a), this ratio did not decrease or decreased to a lesser degree in P and NPC amended bottles (Fig. 8c).

Discussion

A Gradation in P Limitation?

A positive relationship between integrated soluble reactive P and integrated bacterial production was obtained across the Mediterranean Sea. This relationship could be simply explained by decreasing primary production (i.e., the main source of organic carbon for heterotrophic bacteria in the open sea) related to phosphorus deficiency. However, the enrichment experiments confirmed that bacterial production in the surface layer of the Mediterranean Sea appeared to be primarily controlled by phosphorus and secondarily by carbon during the stratification period

(summer and autumn). Thus, the present results confirm that phosphorus limitation of bacterioplankton is a generalized phenomenon in the Mediterranean sea, as has been hypothesized from scattered data collected at selected Mediterranean sites [6, 36, 45]. Most of the studies dealing with alkaline phosphatase activity were conducted in freshwater eutrophic conditions, e.g., [8, 41], whereas data in marine oligotrophic environments are scarce [4, 10, 36]. With a set of concentrations of 10–1000 nM MUF-P, Cotner et al. [10] determined maximum rates of alkaline phosphatase (AP) activity from 1.39 to 2.70 nmol L⁻¹ h⁻¹ in the upper 80 m in the Sargasso Sea in spring and summer. These values are of the same range as those found in this study under P limitation conditions (Table 4). In the Bay of Villefranche in summer Thingstad et al. [36] found that the maximum rate of AP was 28 nmol L⁻¹ h⁻¹, i.e., two fold higher than our greatest value (12 nmol L⁻¹ h⁻¹).

It is known that both phytoplankton and heterotrophic bacteria contribute to AP activity [8, 16]. Indeed, in surface waters of station DYF, 47% of the AP activity and 62% of the bacterial production were associated with the >0.6 μm fraction. In order to consider both populations and compare different situations, maximum rates of AP are generally related to particulate organic phosphorus (POP) concentration [10, 36]. Particulate organic phosphorus was determined only during the September cruise [25]. In surface samples taken along the transect, ratios of Vm of alkaline phosphatase to particulate organic phosphorus (VmAP / POP) ranged from 0.11 to 0.68 h⁻¹ in conditions of P-limitation and from 0.003 to 0.03 h⁻¹ in conditions of C-limitation. Along the depth profiles, they fell in the same range as those measured along the transect, ranging from 0.05 to 0.67 h⁻¹ in the upper layers where bacterial P limitation was found. VmAP/POP ratio reached a maximum of 0.45–0.65 h⁻¹. These values are similar to those obtained in conditions of P limitation in Villefranche, 30 miles further north of station DYF in June (0.6 h⁻¹ [36]), and in the Sargasso Sea in spring and summer (0.3 and 0.35 h⁻¹, respectively [10]).

In cases of bacterial P limitation, the ratio of alkaline phosphatase activity to bacterial production (AP / BP ratio) was strongly reduced after P amendments as a logical consequence of physiological (presence of excess phosphate, the by-product of phosphatase) and genetic regulation (inducible phosphatase). Surprisingly, this ratio was not always reduced after P additions when carbon was limiting bacterial production. In this case, bacterial

production and alkaline phosphatase activity could be maintained because hydrolysis of organophosphoric compounds also provided organic carbon sources. The latter process has been hypothesized for deep oceanic waters [18] and freshwater [8].

The stimulation of bacterial production by a factor $\times 1.4$ relative to the control after 24 h of incubation appeared to be the threshold of a significant positive effect of amendments on bacterial activity and was consistent with low turnover times of AP (Fig. 9). Our tests of reproducibility were 17% on average, which is similar to other studies: 15% [45] and 12% [37]. However, the comparison of bacterial production between an amended flask and the control is sometimes difficult due to variable responses of bacterial production in the unamended control itself, which are not clearly understood [28, 32, 45]. The trend that we observed in vertical profiles was that the largest increases in the unamended controls were obtained in the deepest part of the vertical profiles where bacterial production to bacterial biomass ratios were lowest. This result suggests that the inactive fractions of the bacterial community react rapidly to confinement. On the contrary, in the upwelling station (UPW) where this ratio was high, bacteria reacted to a much lesser degree to confinement.

Thus, we think that the absence of a clear gradient in P limitation is probably due to the above mentioned sources of variability during enrichment experiments. However, from the relationship between increase of bacterial production after 24h of P addition and turnover time of AP (Fig. 9), two interesting points emerged: i) turnover times of AP less than 100 h, associated with a minimum of $\times 1.5$ to $\times 2$ increase of bacterial production relative to the control in P amended bioassays, indicated P limited bacteria and ii) turnover times greater than 100 h corresponded to an insignificant bacterial growth following P amendments indicating no P limitation.

Switch from P to C limitation of bacterial production with depth

Studies focusing on factors limiting bacterial production along depth profiles are scarce. In the Gulf of Mexico, Skoog et al. [33] reported that N limitation of bacterial activity could occur only in the vicinity of the chlorophyll maximum. Gurung and Urabe [15] observed P limitation in surface waters of Lake Biwa, and argued that temperature limited bacterial growth at greater depths. At both DYF (western Mediterranean) and MIO (eastern Mediterranean) profiles, the turnover time of alkaline phos-

phatase increased with depth. Moreover, at station DYF, bacteria switched from phosphorus to carbon limitation at around 50 m. At station MIO, however, the switch was not very clear. Although P limitation was detected at the surface, and carbon limitation below 40 m, we observed that at 50, 90, and 130 m bacterial production increased after both P alone and C alone enrichments. Furthermore at station MIO, at the depths where C limitation was found, the ratio of maximum rates of alkaline phosphatase to particulate organic phosphorus (VmAP/POP) were on average higher (0.075 to 0.1 h^{-1}) than those measured at the C limited depths of station DYF (0.006 – 0.056 h^{-1}). These two observations suggest that the C limited bacteria below 40 m at station MIO were also still P limited. Modest stimulations of bacterial growth after addition of nutrients alone or glucose alone were also reported by Felip et al. [12] in eutrophic lakes, and these authors suggested that, at *in situ* conditions, both nutrients and organic carbon might limit bacterial growth. Theoretically, a real colimitation of heterotrophic bacteria by P and C (i.e., the same bacteria are limited both in C and P) bacterial production should be stimulated in PC enrichments, but not in P alone and C alone. Some possible explanations are as follows:

The increase in leucine incorporation rates in P amended bottles could be due to both heterotrophic bacteria and cyanobacteria, because cyanobacteria can assimilate leucine at low concentrations [19]. However, this explanation is unlikely in the Mediterranean, where *Synechococcus*-like cells, which are known for their low capability of leucine incorporation [19], dominated the cyanobacterial populations. In fact, applying the maximal rate of leucine incorporation reported in Kamjunke and Jahnichen [19], i.e., $37 \mu\text{mol leu (g chl } a)^{-1} \text{ h}^{-1}$ to the maximum *Synechococcus* concentration observed during our study (station DYF, $27 \times 10^3 \text{ cyano ml}^{-1}$), leucine incorporation by *Synechococcus* would represent only 1.8% of the total incorporation rate in P-amended bottles.

Another explanation could be related to phytoplankton-bacteria interactions during the bioassays. P addition alone may stimulate both P limited autotrophs and heterotrophic bacteria. The latter may be stimulated because, as dissolved organic carbon due to sloppy feeding, lysis, and excretion by phytoplankton increases, labile carbon becomes more available to bacteria. In contrast, glucose addition alone can stimulate only heterotrophic bacteria. In this case, glucose addition may shift the competition between phytoplankton and heterotrophic bacteria for P in

favor of bacteria. Consequently, stimulation of bacterial production by addition of P alone or C alone is possible even if heterotrophic bacteria are under P and C colimitation conditions. At least, the two types of bacterial populations (one P limited, one C limited) could coexist, as models allowing such coexistence can now be constructed (Thingstad, pers. comm).

Effect of Enrichment on Ectoaminopeptidase Activity and Utilization of Amino Acids

Potential ectoaminopeptidase activity and potential assimilation rates of amino acids behaved like bacterial production after enrichments. Ratios of these activities to bacterial production, (PEA/BP and PAA/BP) did not differ more than 2-fold from those obtained for the unamended control (Fig. 8), even though bacterial production response was usually much greater (in NPC amended bottles, bacterial production relative to the control increased up to $\times 40$; Fig. 6).

Changes within these ratios are also due to probable changes in energy requirements and/or changes of biochemical pathways induced by added substrate or nutrients. For example, when the factor limiting bacterial production was carbon, amino acid respiration percentages decreased in C and NPC amended flasks (stations TM-6 and TM-7, Fig. 7). Glucose is known to support a large fraction of the bacterial carbon requirement and is also an important energy source [29]. Consequently, bacteria used glucose to restore their energetic capacity and thus the percentage of respired amino acids was reduced, since amino acids could be assimilated directly into cellular constituents [9, 14].

When the factor limiting bacterial production was phosphorus, P addition apparently favored bacterial populations adapted to use amino acids at high concentrations (i.e., PAA/BP ratios increased, stations TM-3 and TM-8, Fig. 8b), whereas the potential to degrade proteins was more or less maintained (Fig. 8a). Under phosphorus limitation, populations favored after P amendments are thus, those adapted to rapidly use the pool of amino acids present. Additionally, the percentage of respiration of amino acids decreased after addition of P when phosphate limited bacterial production (station TM-3, TM-8, and TM-10). Indeed, it has been previously shown that bacterial growth efficiency decreases with nutrient limitation, both experimentally and *in situ* [23, 37, 46]. Our results on respiration percentage of amino acids suggest that both glucose and

phosphorus limitation induced an energy limitation [5, 21] within bacterial communities in the Mediterranean.

Phosphorus limitation of bacterioplankton production was widespread in the surface layers of the Mediterranean Sea in summer and autumn. Along depth profiles, carbon limitation and/or phosphorus and carbon colimitation were observed below analytical detection limits of soluble reactive P. A future challenge is to produce P budgets in the water column, through estimates of size-class components of particulate P, dissolved organic P, soluble reactive P, and uptake and regeneration of P. This budget would help to clarify interactions between C and P cycles, and this task is currently in progress.

Acknowledgments

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References

1. Azam F (1998) Microbial control of oceanic carbon flux: The plot thickens. *Science* 280:694–696
2. Belviso S, Claustre H, Marty J-C (2001) Evaluation of the utility of chemotaxonomic-pigments as a surrogate for particulate DMSP. *Limnol Oceanogr* 6:989–995
3. Bianchi M, Turley C, Van Wambeke F, Stutt E, Christaki U, Psara S, Conan P (1996) A first and preliminary description of a relationships between the cycling of the primary production by bacteria in the photic zone and the extent of particles flux measured during the MTP pilot phase. *Mediterranean Target Project News* :8–9
4. Caron F, Lin Lim E, Sanders RW, Dennet MR, Berninger UG (2000) Responses of bacterioplankton and phytoplankton to organic carbon and inorganic nutrient additions in contrasting oceanic ecosystems. *Aquat Microb Ecol* 22:175–184
5. Cherrier J, Bauer JE, Druffel ER (1996) Utilization and turnover of labile dissolved organic matter by bacterial heterotrophs in eastern North Pacific surface waters. *Mar Ecol Prog Ser* 139:267–279
6. Christaki U, Van Wambeke F, Dolan JR (1999) Nanoflagellates (mixotrophs, heterotrophs and autotrophs) in the oligotrophic Eastern Mediterranean: Standing stocks, bacte-

- rivory and relationships with bacterial production. *Mar Ecol Prog Ser* 187:297–307
7. Christaki U, Van Wambeke F, Giannakourou A, Gregori G (2001) Nanoflagellate predation on auto- and heterotrophic picoplankton in the oligotrophic Mediterranean Sea. *J Plankt Res* 23:1297–1310
 8. Chrost RJ, Overbeck J (1987) Kinetics of alkaline phosphatase activity and phosphorus availability for phytoplankton and bacterioplankton in lake Plubsee (North German eutrophic lake). *Microb Ecol* 13:229–248
 9. Church M, Hutchins D, Ducklow H (2000) Limitation of bacterial growth by dissolved organic matter and iron in the Southern Ocean. *Appl Environ Microbiol* 66:455–466
 10. Cotner J, Ammermann J, Peele E, Bentzen E (1997) Phosphorus-limited bacterioplankton growth in the Sargasso Sea. *Aquat Microb Ecol* 13:141–149
 11. Dolan JR, Vidussi F, Claustre H (1999) Planktonic ciliates in the Mediterranean Sea: longitudinal trends. *Deep-Sea Res I* 46:2025–2039
 12. Felip M, Pace ML, Cole JJ (1996) Regulation of planktonic bacterial growth rates: The effects of temperature and resources. *Microb Ecol* 31:15–28
 13. Garabétian F (1991) ^{14}C -glucose uptake and ^{14}C - CO_2 production in surface microlayer and surface-water samples: influence of UV and visible radiation. *Mar Ecol Prog Ser* 77:21–26
 14. Gardner WS, Benner R, Amon RM, Cotner JB, Cavaletto JF, Johnson JR (1996) Effect of high-molecular weight dissolved organic matter on nitrogen dynamics in the Mississippi River plume. *Mar Ecol Prog Ser* 133:287–297
 15. Gurung TB, Urabe J (1999) Temporal and vertical difference in factors limiting growth rate of heterotrophic bacteria in Lake Biwa. *Microb Ecol* 38:136–145
 16. Hoppe HG (1986) Relations between bacterial extracellular enzyme activity and heterotrophic substrate uptake in a brackish water environment. In: IFREMER (Ed) *Deuxième Colloque Int. de bactériologie Marine*, Brest, 1–5 Octobre 1984, Actes et Colloques 3, Ann Arbor, pp. 119–127
 17. Hoppe HG (1993) Use of fluorogenic model substrates for extracellular enzyme activity measurement of bacteria. In: PF Kemp, BF Sherr, EB Sherr, JJ Cole (eds) *Handbook of Methods in Aquatic Microbial Ecology*. Lewis, Boca Raton, pp 423–431
 18. Hoppe HG, Ullrich S (1999) Profiles of ectoenzymes in the Indian Ocean: phenomena of phosphatase activity in the mesopelagic zone. *Aquat Microb Ecol* 19:139–148
 19. Kamjunke N, Jahnichen S (2000) Leucine incorporation by *Microcystis aeruginosa*. *Limnol Oceanogr* 45:741–743
 20. Karl D, Tien G (1992) MAGIC: a sensitive and precise method for measuring dissolved phosphorus in aquatic environments. *Limnol. Oceanogr.* 37:583–588
 21. Kirchman DL (1990) Limitation of bacterial growth by dissolved organic matter in the subarctic Pacific. *Mar Ecol Prog Ser* 62:47–54
 22. Kirchman DL (1993) Leucine incorporation as a measure of biomass production by heterotrophic bacteria. In: PF Kemp, BF Sherr, EB Sherr, JJ Cole (eds) *Handbook of Methods in Aquatic Microbial Ecology*. Lewis, Boca Raton, pp 509–512
 23. Kroer N (1993) Bacterial growth efficiencies on natural dissolved organic matter. *Limnol Oceanogr* 38:1282–1290
 24. Krom MD, Kress N, Brenner S, Gordon S (1991) Phosphorus limitation of primary productivity in the eastern Mediterranean Sea. *Limnol Oceanogr* 36:424–432
 25. Moutin T, Thingstad FT, Van Wambeke F, Marie D, Claustre H Is the predominance of synechococcus as primary producer in the Mediterranean Sea related to phosphate uptake competition with heterotrophic bacteria? Submitted
 26. Moutin T, Raimbault P (2002) Primary production, carbon export and nutrients availability in western and eastern Mediterranean Sea in early summer 1996 (MINOS cruise). *J Mar Syst*: in press
 27. Murphy J, Riley JP (1962) A modified solution method for determination of phosphate in natural waters. *Analyt Chim Acta* 27:31–36
 28. Pomeroy LR, Sheldon JE, Sheldon WM, Peters F (1995) Limits to growth and respiration of bacterioplankton in the Gulf of Mexico. *Mar Ecol Prog Ser* 117:259–268
 29. Rich JH, Ducklow HW, Kirchman DL (1996) Concentration and uptake of neutral monosaccharides along 140°W in the equatorial Pacific: contribution of glucose to heterotrophic bacterial activity and DOM flux. *Limnol Oceanogr* 41:596–604
 30. Rivkin RB, Anderson M (1997) Inorganic nutrient limitation of oceanic bacterioplankton. *Limnol Oceanogr* 42:730–740
 31. Sournia A (1973) La production primaire planctonique en Méditerranée—Essai de mise à jour, Newsletter of Cooperative Investigations in the Mediterranean. Monaco 5:1–127
 32. Sherr EB, Sherr BF, Sigmon C (1999) Activity of marine bacteria under incubated and *in situ* conditions. *Aquat Microb Ecol* 20:213–223
 33. Skoog A, Biddanda B, Benner R (1999) Bacterial utilization of dissolved glucose in the upper water column of the Gulf of Mexico. *Limnol Oceanogr* 44:1625–1633
 34. Smith DC, Azam F (1992) A simple, economical method for measuring bacterial protein synthesis rates in sea water using ^3H -Leucine. *Mar Microb Food Webs* 6:107–114
 35. Thingstad TF, Rassoulzadegan R (1995) Nutrient limitations, microbial food webs, and “biological C-pumps”: suggested interactions in a P-limited Mediterranean. *Mar Ecol Prog Ser* 117:299–306
 36. Thingstad TF, Zweifel UL, Rassoulzadegan F (1998) P limitation of heterotrophic bacteria and phytoplankton in the northwestern Mediterranean. *Limnol Oceanogr* 43:88–94
 37. J-P Torréton, Talbot V, Garcia N (2000) Nutrient stimulation of bacterioplankton growth in Tuamotu atoll lagoons. *Aquat Microb Ecol* 21:125–137
 38. Treguer P, Le Corre P (1975) Manuel d’analyses des sels nutritifs dans l’eau de mer (utilisation de l’Auto-Analyser II

- Technicon), 2nd ed., Laboratoire d'Océanographie chimique. Université de Bretagne Occidentale, Brest
39. Tselepides A, Zervakis V, Polychronaki T, Danovaro R, Chronis G (2000) Distribution of nutrients and particulate organic matter in relation to the prevailing hydrographic features of the Cretan Sea (NE Mediterranean). *Prog Oceano* 46:113–142
 40. Turley C, Bianchi M, Christaki U, Conan P, Harris J, Psarra S, Ruddy G, Stutt E, Tselepides A, Van Wambeke F (2000) Relationships between primary producers and bacteria in an oligotrophic sea—the Mediterranean. *Aquat Microb Ecol* 183:11–18
 41. Vadstein O Heterotrophic, planktonic bacteria and cycling of phosphorus. Phosphorus requirements, competitive ability and food web interactions. In: B Schink (ed) *Advances in Microbial Ecology*, vol 16. Kluwer Academic/Plenum, New York, pp 115–167
 42. Valderrama JC (1981) The simultaneous analysis of total nitrogen and total phosphorus in natural waters. *Mar Chem* 10:109–122
 43. Van Wambeke F, Christaki U, Bianchi M, Psarra S, Tselepides A (2000) Heterotrophic bacterial production in the Cretan Sea (NE Mediterranean). *Prog Oceanogr* 46:205–216
 44. Vaultot D, Le Bot N, Marie D, Fukai E (1996) Effect of phosphorus on the *Synechococcus* cell cycle in surface Mediterranean waters during summer. *Appl Environ Microbiol* 62:2527–2533
 45. Zohary T, Robarts RD (1998) Experimental study of microbial P-limitation in the eastern Mediterranean. *Limnol Oceanogr* 43:387–395
 46. Zweifel UL, Norrman B, Hagström ADA (1993) Consumption of dissolved organic carbon by marine bacteria and demand for inorganic nutrients. *Mar Ecol Prog Ser* 101: 23–32