# Phytoplankton on the western coasts of Baja California in two different seasons in 1998

# DAVID U. HERNÁNDEZ-BECERRIL, ERNESTO BRAVO-SIERRA and JOSÉ A. AKÉ-CASTILLO

Laboratorio de Diversidad y Ecología del Fitoplancton Marino, Instituto de Ciencias del Mar y Limnología, Universidad Nacional Autónoma de México (UNAM), Apdo. postal 70-305, México, D.F. 04510 México. E-mail: dhernand@mar.icmyl.unam.mx

SUMMARY: Phytoplankton was studied in two different seasons of 1998 (March-April and December), during two cruises along the western coasts of Baja California, in three zones. Two different protocols for obtaining and studying phytoplankton were followed. In the March-April season, phytoplankton had relatively low species richness and was dominated in cell density (up to 93%) by coccolithophorids (mainly *Emiliania huxleyi*), together with nanoplanktonic centric and pennate diatoms, with abundances ranging from 5.4 10<sup>3</sup> to 1.2 10<sup>5</sup> cells L<sup>-1</sup>. In December, phytoplankton had higher species richness and was represented by larger, chain-forming diatom species, such as *Pseudonitzschia delicatissima* and *P. pungens*, which were widespread and numerically significant. There was a relative scarcity of coccolithophorids and thecate dinoflagellates, and densities were between 7 10<sup>2</sup> and 1.4 10<sup>6</sup> cells L<sup>-1</sup>. Hydrographic and oceanographic conditions in March-April were influenced by the occurrence of El Niño and the phytoplankton structure was found to be modified accordingly, with nanoplanktonic coccolithophorids and diatoms being significant contributors to the total abundance. In contrast, post-upwelling conditions might have favoured relatively high densities of *Pseudonitzschia* and other diatoms in December, 1998. Coccolithophorids have not been previously regarded as important contributors to the phytoplankton abundances in Baja California.

Keywords: Baja California, coccolithophorids, diatoms, El Niño, phytoplankton.

RESUMEN: FITOPLANCTON EN COSTAS OCCIDENTALES DE BAJA CALIFORNIA EN DOS TEMPORADAS DISTINTAS DE 1998. – Se estudió el fitoplancton en dos temporadas diferentes de 1998 (marzo-abril y diciembre), durante dos cruceros en tres zonas de las costas de Baja California. Se siguieron dos protocolos distintos para obtener y estudiar el fitoplancton. En marzo-abril, el fitoplancton tuvo una riqueza de especies baja y estuvo dominado en densidades celulares (de hasta 93%) por cocolitofóridos (principalmente *Emiliania huxleyi*), junto con diatomeas centrales y pennales nanoplanctónicas, con abundancias desde 5.4  $10^3$  a  $1.2 10^5$  céls L<sup>-1</sup>. Para diciembre, el fitoplancton incrementó la riqueza de especies y estuvo representado por diatomeas más grandes, formadoras de cadenas, que se distribuyeron ampliamente y fueron numéricamente importantes, como: *Pseudonitzschia delicatissima y P. pungens*, con relativa escasez de cocolitofóridos y dinoflagelados tecados, con densidades entre 7  $10^2$  y  $1.4 10^6$  céls L<sup>-1</sup>. Las condiciones oceanográficas en marzo-abril indicaron la presencia de El Niño y los resultados mostraron una estructura de fitoplancton modificada por ello, con una contribución importante de cocolitofóridos y diatomeas nanoplanctónicas a la abundancia total. En cambio, en diciembre, las condiciones de post-surgencia pudieron haber favorecido las relativas altas densidades de especies de *Pseudonitzschia* y otras diatomeas. Los cocolitofóridos no habeín sido documentados previamente como contribuyentes importantes a la abundancia fitoplanctónica en Baja California.

Palabras clave: Baja California, cocolitofóridos, diatomeas, El Niño, fitoplancton.

## INTRODUCTION

The western coastal area of Baja California, Mexico has traditionally been considered a very rich and productive system, in which the influence of the California Current is remarkable (Parrés Sierra *et al.*, 1997) and upwellings play an important role in keeping this high productivity (Longhurst *et al.*, 1967; Wang and Walsh, 1976; Parrés Sierra *et al.*, 1997). The phytoplankton of this area have been relatively well-studied: there are studies on composition, distribution, primary productivity and variability of some of its features, as well as the main factors controlling the natural variation of phytoplankton (Smayda, 1975; Estrada and Blasco, 1979; Hernández-Becerril, 1985; Cardona *et al.*, 1990; Millán-Núñez and Loya Salinas, 1993; Cohen Fernández, 2000).

Diatoms and "armoured" dinoflagellates have been studied in greater detail than other taxonomic groups and they are usually regarded as the dominant phytoplankton in the area (Allen, 1923, 1924, 1934; Balech, 1960), and more recently (Hernández-Becerril, 1988, 1989, 1995, 1998; Hernández-Becerril and Tapia, 1995). Other groups (e.g. coccolithophorids and other flagellates) or size fractions (e.g. nanoplankton and picoplankton) have not been recognized as important contributors to the diversity, biomass and primary productivity. In nearby locations, in the California current system, coccolithophorids and other flagellates have been identified and counted (in terms of cell numbers and cellular volume) (Reid et al., 1970), the relative importance of microplankton (netplankton) and nanoplankton has been stressed (Malone, 1971; Mullin, 1998), and the contribution to biomass and productivity of fractions of phytoplankton (smaller and larger than 8  $\mu$ m) in a coastal area off Baja California has been studied (González Morales and Gaxiola-Castro, 1991).

The El Niño 1997-1998 event caused a pronounced effect on the climate, the hydrographic and oceanographic conditions, and the biological communities of the Pacific Ocean (Lynn *et al.*, 1998; Hayward *et al.*, 1999). This event reached a second peak of intensity in early 1998 and relaxed considerably by the middle of 1998. By the end of 1998 it disappeared completely from Baja California waters (Lavaniegos *et al.*, 2002), giving way to the strong La Niña event (Hayward *et al.*, 1999).

In this paper, results of the structure of the phytoplankton (in terms of species composition, abundance and distribution) collected during two cruises along the western coasts of Baja California, in two different seasons of 1998, one under the influence of El Niño and the other in contrasting conditions, are presented and discussed.

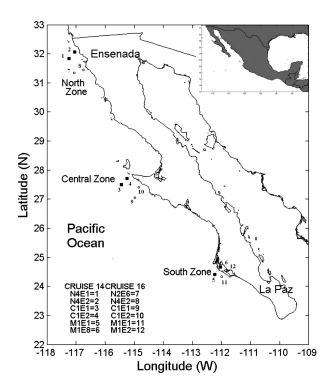


FIG. 1. – Map of the study area, showing sampling station during two cruises: Stations 1-6 from the cruise SIMSUP 14 (March-April, 1998), Stations 7-12 from the cruise SIMSUP 16 (December, 1998).

### MATERIAL AND METHODS

Sampling for this study was carried out during two cruises, on board "El Puma": SIMSUP XIV (March-April, 1998) and SIMSUP XVI (December, 1998), along the western coasts of Baja California, Mexico, in three different zones (north, centre and south) (Fig. 1). Samples were taken basically following two distinct protocols: (1) In March-April, 1998, which is considered the decline of the El Niño 1997-1998 event, phytoplankton was collected with a Niskin bottle (usually 4 L) from three depths at 6 fixed stations (two in the three different zones) and filtered through Millipore 0.45 µm meshed-filters with a vacuum pump (less than 100 mm Hg). The filters were immediately rinsed with a buffered (pH 7.5-8) solution, air dried at room temperature, and then stored in plastic Petri dishes (Winter et al., 1999; Bollmann et al., 2002). (2) In December, 1998, phytoplankton was taken with a Niskin bottle (4 L) from three depths at 14 fixed stations. Approximately 250 ml was then fixed with Lugol's solutions and placed in dark bottles until analysis. Variables measured in both cruises were obtained by a CTD instrument, including temperature, salinity and dissolved oxygen (vertical profiles).

Phytoplankton was analyzed with protocol 1 as follows: small pieces (approximately 1 cm<sup>2</sup>) of the filters containing phytoplankton were cut then mounted onto slides. An oil immersion was added to clear the filters, then they were covered by a coverslip and observed with a light microscope (LM), using bright field and total magnifications of 600× or 1250×, to count and confirm species identification respectively (Bollmann et al., 2002, in most details, except the use of immersion oil and the bright field microscopic technique). Preliminary species identification and cell counting were made in LM. The number of cells per litre was estimated from data such as the number of organisms counted, areas of counting and filter, and volume filtered (Bollmann et al., 2002). Observations were also made by scanning electron microscopy (SEM) to confirm identification of tiny species, using small pieces of filters treated conventionally for SEM. Very few cells were found to be collapsed (due to the possible effect of changing pH or vacuum pressure). Samples fixed with Lugol's solution (protocol 2) were studied using an inverted microscope, following the usual settling in chambers and routine counting procedures (Hasle, 1978).

We used two different protocols to study the phytoplankton in this study, which may prevent direct comparisons being made between different seasons (March-April, 1998 and December, 1998). Our basic purpose when filtering samples (protocol 1) was to obtain the preservable fraction of the phytoplankton, especially the coccolithophorids, although the filters also contained diatoms, thecate (and resistant naked) dinoflagellates, silicoflagellates and other groups which have siliceous or carbonate coverings. This method underestimates the contribution of fragile "phytoflagellates", such as forms of Prasinophyceae, Cryptophytes, other Haptophytes, and also some picoplanktonic forms.

The number of coccolithophorids is estimated more accurately using this method (Bollmann *et al.*, 2002), as we can confirm identification and counting by SEM. Therefore, we were able to establish a phytoplankton community dominated by this group, and the diatoms, the group considered the most significant in terms of species diversity, abundance, and biomass in the study area, are not excluded by this method.

Samples taken and studied following the method for inverted microscope were analyzed very soon after collection, which prevents a possible dissolution of coccoliths from coccolithophorids. Hence, we were able to count this group when analyzing samples. Observation of other flagellates and nanoplanktonic diatoms (i.e. *Chrysochromulina* sp., *Fragilariopsis pseudonana*, *Meringosphaera mediterranea*, *Myrionecta rubra*, *Nitzschia bicapitata*) following this protocol allowed us to be confident with the phytoplankton composition and estimated abundances.

# RESULTS

# Hydrographic conditions and phytoplankton, March-April, 1998

Surface temperature data for March-April, 1998, were higher (by more than 1°C) than those usually reported in other studies for this season (Lynn *et al.*, 1998). The thermocline in most stations was located between 50 and 60 m, except in a shallow station (Fig. 2). There was only a significant change in salinity in a station situated in Bahia Magdalena (M1 E8), and in the remaining stations it did not change considerably (Fig. 2). Dissolved oxygen showed variation similar to temperature (Fig. 2).

The phytoplankton showed low species diversity, with up to 82 species identified. Its composition exhibited a community predominated by nanoplanktonic forms, in which coccolithophorids were important contributors to the abundance, mainly the species Emiliania huxleyi and Gephyrocapsa oceanica (Table 1). Diatoms were represented by smallsized, solitary and lanceolate/centric species, some of them of tropical affinity such as Asteromphalus sarcophagus, Fragilariopsis pseudonana, Nitzschia bicapitata and some Thalassiosira species, which were numerically significant. A few diatoms of fresh-water/brackish habitats also occurred in some samples. Thecate dinoflagellates appeared to be rather scarce, whereas the numbers of "naked"/unarmoured dinoflagellates were slightly higher. The heterotrophic flagellate Meringosphaera mediterranea was a very common, widespread species, and silicoflagellates were very common too, although not abundant.

The abundances of phytoplankton were low to relatively high, ranging between 5.4  $10^3$  and 1.2  $10^5$ cells L<sup>-1</sup>, with the coccolithophorids making up to a maximum of 93% of the total abundance (Table 2), and in some points reaching a ratio of 21:1 with 738 • D.U. HERNÁNDEZ-BECERRIL et al.

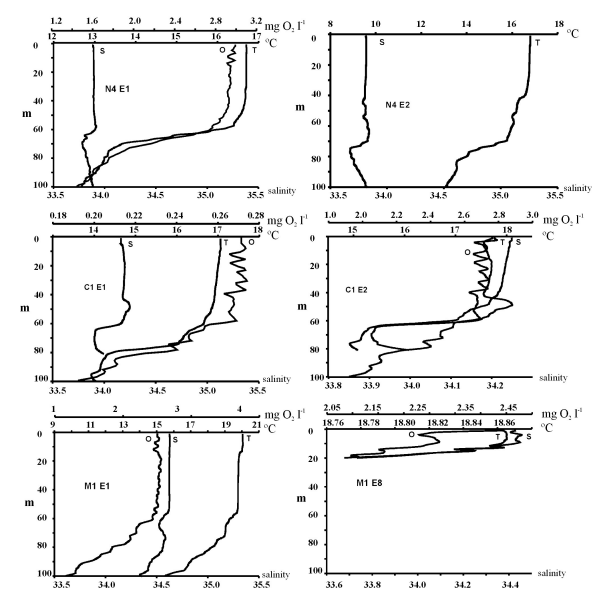


FIG. 2. - Vertical profiles of temperature, salinity and dissolved oxygen at stations from the cruise SIMSUP 14 (March-April, 1998).

regard to diatoms. These abundances were rather homogeneous (about the same magnitude order) with respect to both horizontal and vertical distribution of phytoplankton in most stations, except C1 E1 (zone centre), where we found the maximum values of phytoplankton abundances at the surface and subsurface, with a high contribution from the coccolithophorids (1.1  $10^5$  and 0.95  $10^5$  cells L<sup>-1</sup> respectively) (Table 2).

# Hydrographic conditions and phytoplankton, December, 1998

Surface temperatures were usual for this time of the year. Well-defined thermoclines were only detected at stations of the northern and central zones. They were not found below 45 m (Fig. 3). Salinity showed no considerable variation (not even in a station at Bahia Magdalena, M1 E2), and dissolved oxygen again varied in parallel to temperature, except in station N2 E6 (Fig. 3).

The number of species in the phytoplankton was relatively higher than in the precedent season, with up to 121 species identified, with somewhat homogeneous communities containing large- and medium-sized chain forming diatoms. Two species, *Pseudonitzschia delicatissima* and *P. pungens* were widespread and significant contributors to the total phytoplankton abundance (Table 1). Other traditionally common diatoms, such as species of *Chaetoceros* were not as abundant as in other sea-

Species M.	laximum density (cells L <sup>-1</sup> )	Mean
MARCH-APRIL, 1998		
Emiliania huxleyi	73000	22036
Gephyrocapsa oceanica	35200	9344
Nitzschia bicapitata	2500	980
Fragilariopsis pseudonana	1420	640
Syracosphaera pulchra	1230	560
Thalassiosira oestrupi var. veni	rickae 820	230
Total number of species: 82		
December, 1998		
Pseudonitzschia delicatissima	114681	14071
Pseudonitzschia pungens	95567	8107
Chaetoceros curvisetus	99815	7056
Guinardia delicatula	165650	6917
Guinardia striata	116804	6454
Detonula pumila	146536	5972
Chaetoceros lorenzianus	82825	5037
Chaetoceros debilis	110433	4591
Chaetoceros compressus	114681	3911
Chaetoceros laciniosus	46722	2481
Chaetoceros brevis	31416	2327
Eucampia cornuta	29732	1673
Skeletonema cf. costatum	31856	1616
Chaetoceros radicans	35343	1464
Chaetoceros tortissimus	26546	1242
Dactilyosolen fragilissimus	27915	1119
Prorocentrum triestinum	11781	1158
Nitzschia bicapitata	23562	983
Guinardia flaccida	24313	983
Ceratium furca	1815	84
Prorocentrum micans	1062	25
Ceratium fusus	708	16
Total number of species: 121		

TABLE 1. – Dominant species in cell densities during March-April,
1998, and December, 1998 respectively.

TABLE 2. – Phytoplankton abundances (cells L<sup>-1</sup>) in March-April, 1998, and December, 1998, with dominance (%) of coccolithophorids (*Emiliania huxleyi* and *Gephyrocapsa oceanica*) and *Pseudonitzschia* species respectively. Stations as in Figure 1.

Station N	V4E1		Station N4	E2	
5 m	54 749		5 m	50 643	
30 m	56 118		30 m	71 995	88%
50 m	57 760	87%	50 m	62 120	93%
Station C1E1			Station C1E2		
5 m	126 744	87%	5 m	6 296	
30 m	105 666	91%	30 m	5 475	
50 m	20 257		50 m	7 938	
Station M1E1		Station M1E8			
5 m	20 530		5 m	13 961	
30 m	20 257	88%	10 m	9 857	
50 m	20 530		15 m	16 972	
Decembi	er, 1998				
Station N2E6		Station N2E8			
5 m	53 233		5 m	4 720	
25 m	36 216		25 m	43 733	
52 m	56 632		50 m	13 686	
	"1E1		Station C1	E2	
Station C			5 m	7 854	58%
Station C 5 m	84 430		J III		
			50 m	79 639	
5 m	84 430			79 639 11 892	53%
5 m 25 m	84 430 92 028 1 745		50 m	11 892	53%
5 m 25 m 50 m	84 430 92 028 1 745		50 m 100 m	11 892	53%
5 m 25 m 50 m Station M	84 430 92 028 1 745		50 m 100 m Station M1	11 892 E2	53%

sons, and species of *Rhizosolenia* and *Thalassiosira* only appeared sporadically. Flagellates were rather scarce (including coccolithophorids and thecate dinoflagellates).

Phytoplankton abundances varied notably, from 7  $10^2$  to 1.4  $10^6$  cells L<sup>-1</sup>, with the lowest abundances in the northern and central zones and the highest ones at the surface layer in the southern zone (Bahia Magdalena and surroundings), where the maximum density reached more than a million cells L<sup>-1</sup> (Table 2). Generally, phytoplankton cell numbers were higher in surface and subsurface waters (up to 50 m) (Table 2).

## DISCUSSION

# Hydrographic and oceanographic conditions: the El Niño effect

Available data (either obtained from the field, Fig. 2, or from other sources, e.g. Lynn *et al.*, 1998; Hayward *et al.*, 1999; Lavaniegos *et al.*, 2002) support the general condition: El Niño was occurring in Baja California waters at the time that collections were made for March-April, 1998. The temperature profiles showed that the thermocline was relatively deep (up to 50-60 m), with the sub-surface and surface water being relatively homogeneous (Fig. 2). Temperatures of the sub-surface and surface water were slightly higher than the general average for this time in the study area. Chlorophyll a data for January-February, 1998, showed an increase in surface concentration and a strong gradient onshoreoffshore, with higher values in the coastal zone. Whereas by summer and autumn, 1998 (July, September-October respectively), chlorophyll a concentrations were significantly lower (Lavaniegos et al., 2002, 2003).

We consider that these conditions had a strong influence on the phytoplankton community and were the reason for the composition of the phytoplankton with a strong numerical dominance of nanoplanktonic forms, especially coccolithophorids and small-sized diatoms with similar abundances from surface to 50 m.

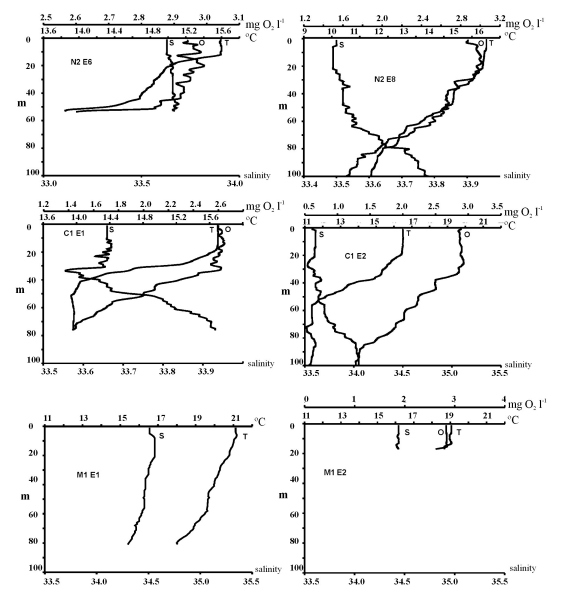


FIG. 3. - Vertical profiles of temperature, salinity and dissolved oxygen at stations from the cruise SIMSUP 16 (December, 1998).

After a period of weaker than normal upwelling through late spring and summer 1998 from San Francisco to southern Baja California, the positive index anomaly indicates slightly above normal upwelling in late 1998 and early 1999 (Hayward *et al.*, 1999). In fact, typical colder surface waters associated with upwelling in coasts off Baja California were observed during October 1998 (Aguirre-Hernández *et al.*, 2004). Our hydrographic data also support these observations, indicating upwellings still occurring during December 1998, with post-upwelling conditions at different zones throughout the study area (Fig. 3). These circumstances may have favoured the spread and relatively high abundances of certain diatom species, such as *Pseudonitzschia delicatissima* and *P. pungens*, in December, 1998. A closely related species, *Pseudonitzschia australis*, exhibited high abundances in Southern California from February to August, even forming blooms by March, 1991 (Lange *et al.*, 1994). This has been associated with the intrusion of cool, nutrient-rich waters, which are a product of upwellings (Lange *et al.*, 1994).

The extensive spread of the *Pseudonitzschia* species was evident from May, 1998 on the western coasts of the USA, such as Washington (Adams *et al.*, 2000; Horner *et al.*, 2000) and California, especially by July, 1998 (Bargu and Silver, 2003), including some cases of intoxication in Californian waters due to blooms of some toxic species (Trainer

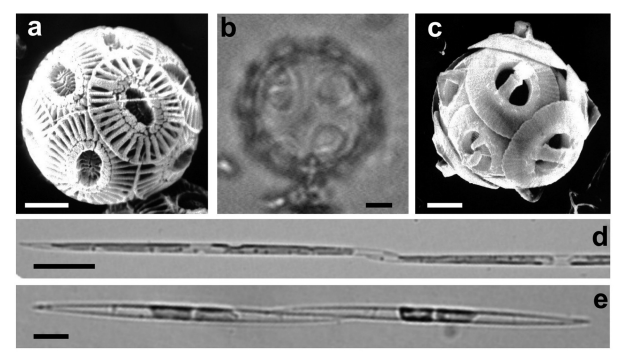


FIG. 4. – Important species during March-April and December, 1998; a, the coccolithophorid *Emiliania huxleyi*, SEM; b, c, *Gephyrocapsa oceanica*, another coccolithophorid, LM and SEM respectively; d, e, the chain-forming, pennate diatoms *Pseudonitzschia delicatissima* and *P. pungens* respectively, LM. Scale bars = 10 µm (d, e) and 2 µm (a-c).

*et al.*, 2000, 2001). This distribution was also documented in more tropical waters from the Mexican Pacific from November, 1998, until April, 1999, from Baja California to warmer waters around Mazatlán, México (Gómez-Aguirre, 1999; Meave *et al.*, 2000). This large-scale distribution may be explained in terms of local events (for instance post-upwellings and heavy precipitations, among others), but also on a larger meteorological and oceano-graphic scale as the effect of La Niña.

### Phytoplankton structure

In our results from March-April, 1998, the coccolithophorid *Emiliania huxleyi* (Fig. 4a) was the most significant contributor to the cell abundance of phytoplankton, although it did not reach spectacular densities (less than 1.2 10<sup>5</sup> cells L<sup>-1</sup>) or produce real blooms (as documented by Tyrell and Merico, 2004, in other parts of the world) (Tables 1 and 2). It was usually found together with *Gephyrocapsa oceanica* (Fig. 4b, c), which occurred in lower abundances (Table 1). *Emiliania huxleyi* appears as the species involved in most blooms of coccolithophorids world-wide and it is considered cosmopolitan (Tyrell and Merico, 2004). Some recent studies show that this species has high genetic variability, possibly with different physiological responses to different oceanographic conditions (Brand, 1982; Young and Westbroek, 1991; Medlin *et al.*, 1994).

Coccolithophorids are usually regarded as an important and abundant phytoplankton group in warm, stratified and oligotrophic waters (Brand, 1994). They often bloom in many areas, including coastal areas. They also reach high populations in mature upwelled waters (Balch, 2004). However, no previous report has been made on numerical dominance of coccolithophorids in the phytoplankton of Baja California, although a paper dealing with the floristic of this group in the same study area was recently published (Hernández-Becerril *et al.*, 2001), and previous plankton studies in La Jolla, California, USA, included coccolithophorids (with a maximum *Emiliania huxleyi* abundance of 5 10<sup>5</sup> cells L<sup>-1</sup>, in August, 1967, Reid *et al.*, 1970).

Nanoflagellates and naked dinoflagellates yielded considerable densities only at the surface, in the northern and central zones of Baja California in autumn and winter, 1998, in a previous plankton study during different seasons between September, 1997 and October, 1998, whereas centric diatoms were only abundant in autumn, 1997 (Lavaniegos *et al.*, 2003). This information is consistent with our results.

Diatoms have traditionally been regarded as the most significant group in the study area, as they are

usually dominant in these upwelling regions. The relative scarcity of this group in the March-April season and the presence and relative abundance of some rather unusual species in the study area, such as the diatom *Asteromphalus sarcophagus*, which is considered to be a tropical form, confirm the intrusion of warmer water into the system. Furthermore, the association of diatoms in the tropical-subtropical species complex *Nitzschia bicapitata/N. bifurcata* (both species were found in this season) has been previously recognized in Pacific waters (Lee and Fryxell, 1996; Fryxell, 2000).

Other evidence of changing oceanographic conditions and their effects on phytoplankton comes from micropaleontologists: terrigenous input into Santa Barbara Basin, California, USA, was significantly higher during El Niño 1997-1998, which reflects higher rainfall and runoff into the basin (Lange *et al.*, 2000). Diatom fluxes were an order of magnitude lower than in the 1996 non-El Niño period, the proportion of warm water flora increased significantly, and by winter of 1998 organic carbon fluxes were unusually high (perhaps indicating the significant populations of coccolithophorids) (Lange *et al.*, 2000). We also found indicators of a freshwater influence, such as the diatom *Cocconeis placentula*, in very low numbers.

In conclusion, the phytoplankton community found for March-April, 1998 along the coasts of Baja California was different to the usual community there, and reflects the particular conditions of El Niño 1997-1998. The usual composition of the phytoplankton for April in the system of Baja California/California is rich in diatoms. The community detected for December, 1998 is more typical and reflects more periodical environmental conditions. Our results of this season (concentrated in Table 1) are very similar to those provided by Venrick (1998) in her study of phytoplankton from a neighbouring area (southern California) in April 1993 and 1995, with similar significant species (within the maximum rank) such as the diatoms Pseudonitzschia spp. (Figs 4d, e), Chaetoceros compressus, C. debilis, C. radicans and Skeletonema costatum.

However, despite the structure of the phytoplankton community (composition, abundance and distribution) being different in the two periods studied on Baja Californian coasts, other ecological parameters seemed to remain the same during El Niño 1997-1998 in California, USA, such as the

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general relation between biomass of large phytoplankton and the depth of the nitricline (Mullin, 2000).

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