Size-fractionated phytoplankton pigment groups in the NW Iberian upwelling system: impact of the Iberian Poleward Current

F. Rodríguez1,*, J. L. Garrido2, B. G. Crespo2, B. Arbones2, F. G. Figueiras2

1Centro Oceanográfico de Canarias (IEO), Carretera de San Andrés s/n, 38120 Santa Cruz de Tenerife, Spain
2Instituto de Investigaciones Marinas (CSIC), Eduardo Cabello 6, 36208 Vigo, Spain

INTRODUCTION

Coastal phytoplankton populations experience large variations in their abundance, composition and size structure due to the dynamic nature of their environment, with potential mixing of distinct water masses of estuarine, neritic and/or open-ocean origin (Bidigare et al. 1993, Pelegri et al. 2005). In coastal areas affected by seasonal upwelling, biological changes are even more pronounced, as nutrient-rich upwelled waters favor the dominance of large organisms (microplankton, usually diatoms) and high levels of primary productivity (Varela 1992, Prézelin et al. 2000, Tilstone et al. 2003).

Over the seasonal cycle, pico-cyanobacteria and nanoflagellates, typically haptophytes and chlorophytes (Bustillos-Guzmán et al. 1995, Rodriíguez et al. 2003a), can account for a significant fraction of the productivity and biomass in coastal regions (Tilstone et al. 2003). In recent years, a wealth of studies has brought to light the true taxonomic diversity of the smallest algae, especially the eukaryotic picoplankton (e.g. Not et al. 2004, Romari & Vaulot 2004). However, more work is needed to answer some fundamental ques-
tions, such as: What are the dominant groups and what are the environmental parameters controlling their distribution in coastal versus oceanic regions?

The western Iberia Basin, which is the northern limit of the eastern North Atlantic Upwelling Region, displays high levels of primary production in coastal areas and shelf waters boosted by the seasonal upwelling (Bode et al. 1996, Joint et al. 2002). Two principal yearly periods can be distinguished in the region (Tilstone et al. 2003): the seasonal upwelling between March and October triggered by northerly winds, and the winter poleward flow or Iberian Poleward Current (IPC; Álvarez-Salgado et al. 2003, Peliz et al. 2005) associated with southerly winds from winter until early spring. The IPC is a jet of saline and relatively warm waters of subtropical origin (Peliz et al. 2005) that flows from waters south of Portugal to the NW Iberian peninsula and the southern Bay of Biscay. During winter, particularly at the core of the IPC waters, pico-nanoplankton communities and low chl a levels are typically observed (Álvarez-Salgado et al. 2003, Tilstone et al. 2003, Calvo-Díaz et al. 2004).

The phytoplankton composition, abundance and physiology and size distribution off the NW Iberian coast have been extensively studied (Figueiras & Pazos 1991, Varela 1992, Tilstone et al. 2003). Recently, cyanobacteria and picoeukaryotes have been reported to be responsible for the fact that pico-nanoplanktonic biomass is higher on the shelf than in the Ría de Vigo (Lorenzo et al. 2005). Unlike the microplanktonic component, the annual variability and composition of pico- and nanoplanktonic eukaryotes, which also dominate IPC waters (Álvarez-Salgado et al. 2003), remain undescribed.

A previous pigment analysis of a single sample from the slope current off NW Spain (Rodríguez et al. 2003b) evidenced the presence of the cyanobacteria genera Synechococcus and Prochlorococcus with a mixture of diverse picoeukaryotic groups. Flow cytometry signatures along the coasts of the northern Iberian peninsula in winter reported that the general dominance of Synechococcus over Prochlorococcus and a major contribution of picoplankton to the total algal biomass (~60%) were accentuated in IPC waters (Calvo-Díaz et al. 2004).

In the present work, phytoplankton abundance and composition in 2 size classes (pico- and nanomicroplankton) were investigated at a fixed station off the NW Iberian coast, combining high-performance liquid chromatography (HPLC) pigment analyses and carbon (C) biomass derived from microscopic examination. The results obtained allowed, for the first time, characterization of the annual variability and composition of the pico- and nano-microplanktonic community, as well as of the size distribution and major algal groups in each size class associated with the IPC.

**MATERIALS AND METHODS**

**Study site and sample collection.** A fixed station located on the shelf near the Ría de Vigo (42° 07.8’ N, 9° 10.2’ W, 150 m depth) was visited once a week from 15 May 2001 to 24 April 2002 (Fig. 1). Sampling was conducted using a CTD attached to a fluorometer and a rosette equipped with 10 l Niskin bottles. Samples for determination of nitrate, chlorophyll, and phytoplankton levels were taken within the photic layer during the upcasts. Nitrate concentrations (µmol kg⁻¹) were determined by segmented flow analysis with Alpkem autoanalyzers according to Hansen & Grasshoff (1983), with some modifications following Mouriño & Fraga (1985).

**Phytoplankton identification and carbon biomass.** Microplankton samples were preserved in Lugol’s iodine and sedimented in 100 ml composite sedimentation chambers. Diatoms, dinoflagellates, flagellates >20 µm, and ciliates were identified and counted at the species level, when possible, using an inverted microscope. Differentiation of phototrophic species of dinoflagellates, flagellates, and ciliates basically followed Lessard & Swift (1986); however, epifluorescence microscopy was also used to assign phototrophs within these plankton groups. The biovolumes of each taxon
were calculated from dimensions and shapes according to Hillebrand et al. (1999). The plasmatic volume of diatoms and total cell volume of dinoflagellates, flagellates >20 µm, and ciliates were converted to cell carbon following Strathmann (1967) for diatoms and dinoflagellates, Verity et al. (1992) for flagellates, and Putt & Stoecker (1989) and Verity & Langdon (1984) for ciliates.

Phototrophic pico- (≤2 µm) and nanoflagellates (2 to 20 µm) were enumerated from subsamples of 10 ml, which were fixed with buffered formalin (2% final concentration) and filtered on 0.2 µm black Millipore polycarbonate filters placed on top of 0.45 µm Millipore backing filters. An epifluorescence microscope was used to identify and count *Synechococcus*-type cyanobacteria and phototrophic picoflagellates and nanoflagellates. Cell biovolumes of *Synechococcus* were converted to carbon biomass following Fry (1990). The carbon of other pico- and nanoflagellates was estimated according to Verity et al. (1992).

**Pigment analysis.** Seawater samples (2 l) obtained at 4 depths within the photic layer were filtered sequentially through 47 mm diameter Whatman GF/D and GF/F filters (under vacuum pressure ≤100 mmHg) and stored at −20°C until analysis. Two phytoplankton size classes were operationally defined: (1) nanophytoplankton, constituted by organisms retained on a GF/D filter (2.7 µm nominal pore size), and (2) pico-phytoplankton, consisting of organisms which passed through a GF/D but were retained on GF/F filters (0.7 µm nominal pore size). It should be mentioned that a slight underestimation of the contribution of pico-phytoplankton using GF/D filters may have resulted in comparison with Nuclepore 3 µm filters. A comparative test using both filter types would be necessary in further studies to quantify these differences in the retention capabilities and size-fractionated phytoplankton groups.

The frozen filtered samples were extracted in 5 ml of 90% acetone using a spatula for filter grinding and further sonication over 5 min at low temperature (~5°C). Extracts were then centrifuged (4000 rpm [2862 × g], 5 min, 4°C) to remove cell and filter debris, and supernatants were filtered through Gelman GHP 0.2 µm filters. The HPLC pigment analyses were carried out with a Waters Alliance HPLC System (2695 separation module with a 996 photodiode array detector and a 474 scanning fluorescence detector). The autosampler chamber was kept at 4°C. Pigment separation was performed by HPLC according to Zapata et al. (2000), but slightly modified to reduce the pyridine concentration in Mobile Phase A. The stationary phase was a C8 column (Waters Symmetry: 150 × 4.6 mm, 3.5 µm particle size, 100 A pore size) thermostated at 25°C. Mobile phases were: A — methanol:acetonitrile:aqueous pyridine solution (0.025 M pyridine, pH adjusted to 5.0 with acetic acid) (50:25:25, v/v/v) and B — acetonitrile: methanol:acetonitrile:aqueous pyridine solution (60:20:20, v/v/v). A linear gradient from 0 to 40% B was pumped for 22 min, followed by an increase to 95% at Minute 28 and an isocratic hold at 95% B for a further 12 min. Initial conditions were reestablished by a reversed linear gradient. The flow rate was 1 ml min⁻¹. Chlorophylls and carotenoids were detected by diode-array spectroscopy (350 to 750 nm). Chlorophylls were also detected by fluorescence (excitation and emission wavelengths were 440 and 650 nm, respectively). Pigments were identified by co-chromatography with authentic standards (see Zapata et al. 2000) and by diode-array spectroscopy (wavelength range: 350 to 750 nm, 1.2 nm spectral resolution). Each peak was checked for spectral homogeneity using Waters Millennium³² software algorithms, and the absorption spectrum was compared with a spectral library previously created. Pigments were quantified by using external standards and extinction coefficients compiled by Jeffrey (1997).

Chlorophyll a was also determined using conventional fluorimetric procedures. Seawater samples of 100 ml were filtered under low vacuum pressure through 25 mm Whatman GF/F. Filters were then immediately frozen at −20°C until pigment extraction in 90% acetone over 24 h, in darkness at 4°C.

**Pigment data processing (CHEMTAX).** HPLC pigment data were processed by means of the Chemical Taxonomy program (CHEMTAX; Mackey et al. 1996). Pigment groups or chemotaxonomic classes were characterized using pigment ratios available from the literature (Table 1) and were processed using 12 marker pigments in the corresponding CHEMTAX input matrix of each size class (see Table 2). To consider changes in pigment ratios due to variable light conditions, incident photosynthetically available radiation (E_{PAR}) was used to distinguish 2 data sets within the CHEMTAX analyses, low light (LL, E_{PAR} < 200 µmol photons m⁻² s⁻¹, n = 136) and high light (HL, E_{PAR} > 200 µmol photons m⁻² s⁻¹, n = 30) sample groups. Two initial ratio matrices adjusted for LL and HL conditions were used in the subsequent CHEMTAX analyses of each light group and size fraction. The initial ratio matrices included LL and HL pigment:chl a ratios gathered from the literature (Table 1) and from phytoplankton cultures (*Chaetoceros socialis*, *Leptocylindrus danicus*, and *Micromonas pusilla*) isolated in a neighboring coastal area (Ria de Arousa, NW Spain). CHEMTAX was run 2 consecutive times for each light group and size fraction—the first output ratio matrix being used as the initial matrix in the second analysis. The HPLC method allowed the separation of divinyl (DV) chl a (the exclusive marker of the cyanobacterium *Prochlorococcus*) from chl a. Thus, zeaxanthin (Zea)
contributed by *Prochlorococcus* was estimated prior to performing CHEMTAX analyses and subtracted from total *Zea* values.

**Statistics.** The entire data set (including pigments, biomass, physico-chemical variables, and pigment groups) was examined using 1-Pearson *r* correlation to identify co-variations among variables. All statistical analyses were performed using STATISTICA for Windows, Release 5.0, StatSoft (1995).

**RESULTS**

**Hydrography and chl *a***

Based on temperature and nitrate distribution (Fig. 2A, B) 8 hydrographic periods could be distinguished at the station studied: (I) spring–summer stratification (May to July 2001), (II) summer relaxation (August 2001), (III) late summer upwelling (September 2001), (IV) downwelling and stratified poleward current (October to November 2001), (V) transitional upwelling (November 2001), (VI) homogeneous poleward current (December to February 2001/2002), (VII) winter mixing (March 2002), and (VIII) onset of spring conditions (April 2002). Most of these periods corresponded with significant changes in the chemotaxonomic composition of phytoplankton, particularly in the winter IPC (VI). The biological transition between periods usually took place a week after the physicochemical changes.

HPLC total chl *a* (TotChl *a* = chl *a* + DV chl *a*) values ranged between 1 and 2 µg l⁻¹ throughout the year. Only during the September upwelling and in spring (April 2002) did chl *a* values rise to ~3 µg l⁻¹, when diatoms were more abundant. DV chl *a* (*Prochlorococcus*) never exceeded 100 ng l⁻¹.

Overall, picoplankton and nano-microplankton represented 20 and 80% of the TotChl *a* measured between May 2001 and April 2002. The annual trends in chl *a* concentration differed between size fractions (Fig. 2C, D). Picoplankton chl *a* showed smoother variations (average ± SD, 89.4 ± 60.1 ng chl *a* l⁻¹) than the nano-microplankton fraction (359.8 ± 433.8 ng chl *a* l⁻¹). For nano-microplankton (Fig. 2C), maximum chl *a* values occurred during the September upwelling (2.95 µg chl *a* l⁻¹, 25 September 2001, 15 m depth), with other chl *a* maxima registering in July and April 2002. In contrast, the maximum picoplankton chl *a* was measured in March (306 ng chl *a* l⁻¹, 26 March 2002, 3 m depth; Fig. 2D), while from July to September and in April 2002 other secondary chl *a* maxima were observed. Maximum contribution of picoplankton to TotChl *a* was measured during the October downwelling and in the winter IPC (~40% of mean TotChl *a* for each period). However, both size classes showed minimum chl *a* values during the same IPC period (VI). DV chl *a*, the marker pigment for *Prochlorococcus*, showed its maxima in October and November (Periods IV and V, 90.1 ng DV chl *a* l⁻¹, 30 October 2001, 15 m depth; Fig. 2E).

<table>
<thead>
<tr>
<th>Pigment groups</th>
<th>Main marker pigments</th>
<th>Source species</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diatoms I</td>
<td>Chl <em>c</em>₁, Fuco</td>
<td><em>Chaetoceros socialis</em></td>
<td>M. Zapata (pers. comm)</td>
</tr>
<tr>
<td>Diatoms II</td>
<td>Chl <em>c</em>₂, Fuco</td>
<td><em>Leptocylindrus danicus</em></td>
<td>M. Zapata (pers. comm)</td>
</tr>
<tr>
<td>Dinoflagellates</td>
<td>Per</td>
<td><em>Prorocentrum micans</em></td>
<td>Schlüter et al. (2000)</td>
</tr>
<tr>
<td>Haptophytes I</td>
<td>Chl <em>c</em>₂-MGDG [14:0/18:4], 19 Hex</td>
<td><em>Emiliania huxleyi</em></td>
<td>Schlüter et al. (2000), Zapata et al. (2004)</td>
</tr>
<tr>
<td>Haptophytes II</td>
<td>Chl <em>c</em>₂-MGDG [14:0/14:0], 19 Hex</td>
<td><em>Chrysochromulina polylepis</em></td>
<td>Henriksen et al. (2002), Zapata et al. (2001)</td>
</tr>
<tr>
<td>Chlorophytes</td>
<td>Chl <em>b</em></td>
<td><em>Brachiononas submarina</em></td>
<td>Schlüter et al. (2000)</td>
</tr>
<tr>
<td>Prasinophytes II</td>
<td>Chl <em>b</em>, Pras, Uriol</td>
<td>Picoplankton: <em>Pycnococcus provasoli</em></td>
<td>Schlüter et al. (2000)</td>
</tr>
<tr>
<td>Pelagophytes</td>
<td>19 buta</td>
<td><em>Micrononas pusilla</em></td>
<td>M. Zapata (pers. comm.)</td>
</tr>
<tr>
<td>Cryptophytes</td>
<td>Allox</td>
<td><em>Pelagococcus subviridis</em></td>
<td>Schlüter et al. (2000)</td>
</tr>
<tr>
<td>Synechococcus</td>
<td>Zea</td>
<td><em>Rhodomonas salina</em></td>
<td>Schlüter et al. (2000)</td>
</tr>
<tr>
<td><em>Prochlorococcus</em></td>
<td>DV Chl <em>a</em></td>
<td><em>Synechococcus sp.</em></td>
<td>Henriksen et al. (2002)</td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Prochlorococcus marinus</em></td>
<td>Goericke &amp; Repeta (1992)</td>
</tr>
</tbody>
</table>

Table 1. Marker pigments and reference source for the pigment groups included in the CHEMTAX analyses. For pigment abbreviations see Fig. 4 legend.
Picoplankton, nanoplanckton, and microplankton accounted for 9.7, 63.0, and 27.3% of the total autotrophic carbon, respectively. When examining the percentage of these size classes for each seasonal period, microplankton and nanoplanckton were co-dominant during the August relaxation and September upwelling periods (Fig. 3). Microplankton carbon showed its lowest contribution during the poleward flow (Period VI), while picoplankton and nanoplanckton showed maximum values (18 and 76% autotrophic C, respectively; Fig. 3).

After examining the phytoplankton groups, the major contributors to autotrophic C were unidentified nanoflagellates (63%) and diatoms (22%). Major diatom species were *Leptocylindrus minimus* and *L. danicus*, small centric unidentified species and *Skeletonema* spp., which accounted for 35, 14, and 10%, respectively, of the total diatom C.

The autotrophic species of dinoflagellates represented 3% of the total autotrophic C. The most relevant species of the total autotrophic dinoflagellate C were *Prorocentrum minimum*, *Gymnodinium nanum*, and *Karenia mikimotoi*, at 24, 12, and 11%, respectively. Cryptophytes and other minor groups (chrysophytes, haptophytes, prasinophytes, and chlorophytes) also contributed less than 1% to total autotrophic C.

**Phytoplankton assemblages: light microscopy and carbon biomass**

Picoplankton, nanoplanckton, and microplankton were counted for 9.7, 63.0, and 27.3% of the total autotrophic carbon, respectively. When examining the percentage of these size classes for each seasonal period, microplankton and nanoplanckton were co-dominant during the August relaxation and September upwelling periods (Fig. 3). Microplankton carbon showed its lowest contribution during the poleward flow (Period VI), while picoplankton and nanoplanckton showed maximum values (18 and 76% autotrophic C, respectively; Fig. 3).

After examining the phytoplankton groups, the major contributors to autotrophic C were unidentified nanoflagellates (63%) and diatoms (22%). Major diatom species were *Leptocylindrus minimus* and *L. danicus*, small centric unidentified species and *Skeletonema* spp., which accounted for 35, 14, and 10%, respectively, of the total diatom C.

The autotrophic species of dinoflagellates represented 3% of the total autotrophic C. The most relevant species of the total autotrophic dinoflagellate C were *Prorocentrum minimum*, *Gymnodinium nanum*, and *Karenia mikimotoi*, at 24, 12, and 11%, respectively. Cryptophytes and other minor groups (chrysophytes, haptophytes, prasinophytes, and chlorophytes) also contributed less than 1% to total autotrophic C.

**Fig. 2.** Temporal distribution (2001 to 2002) of: (A) temperature (°C), (B) nitrates (µmol kg⁻¹), (C) nano-microplankton Chl a (ng chl a l⁻¹), (D) picoplankton chl a (ng chl a l⁻¹) and (E) divinyl chl a (ng DV chl a l⁻¹). The lengths of each of 8 hydrographic periods (see 'Results' for description) are shown in (A). The lengths of each of 8 hydrographic periods (see 'Results' for description) are shown in (A).

**Fig. 3.** Percentage contribution of picoplankton, nanoplanckton, and microplankton to carbon biomass determined by light microscopy in each hydrographic period. IPC: Iberian Poleward Current. Overall means (±SD) for the size fractions are also given.
phytes, raphidophytes, and euglenophytes) contributed only ~1% of the autotrophic C biomass.

**HPLC pigment composition**

The major pigments in each fraction are shown in Fig. 4. Nano-microplankton and picoplankton were dominated quantitatively by different chlorophylls and carotenoids. The latter were higher in proportion in the nano-microplankton than in the picoplankton (70 and 54% total accessory pigments, respectively; Fig. 4). Log accessory pigments and log TotChl \(a\) were statistically related (slope = 1.180, \(r^2 = 0.954, p < 0.001\)), and, in spite of the distinct contribution of chlorophylls and carotenoids, no significant differences were found between picoplankton and nano-microplankton.

Some pigments were exclusive to a single size group: chl \(c_1\) and peridinin from the nano-microplankton and uriolide, micromonol, micromonal (the 2 latter only detected in trace amounts), and DV chl \(a\) in the picoplankton. For nano-microplankton, chl \(c_2\) (50% total accessory chlorophylls) and fucoxanthin (58% total carotenoids), both shared by multiple taxonomic groups (diatoms, haptophytes, pelagophytes, chrysophytes, etc.), were the main accessory chlorophyll and carotenoid, respectively (Fig. 4A).

In the picoplankton, chl \(b\) (~40% total accessory chlorophylls, contributed by green algae) and zeaxanthin (34% total carotenoids, from cyanobacteria Synechococcus and Prochlorococcus) were the most abundant pigments (Fig. 4B). Non-polar chls \(c\) (i.e. chl \(c_2\)-MGDG [14:0/18:4] and chl \(c_2\)-MGDG [14:0/14:0]) were relatively abundant in the nano-microplankton (11% accessory chlorophylls), but were barely detected in the picoplankton (only 1%). The HPLC method used in this study did not allow separation of DV chl \(a\) (exclusive to Prochlorococcus) from chl \(b\). However, the similarity between the visual ultraviolet spectrum of chl \(b\) in samples with maxima of DV chl \(a\) and that of chl \(b\) allowed us to conclude that little or no DV chl \(b\) was ever present.

**CHEMTAX analysis**

The final ratio (LL and HL) matrices for nano-microplankton and picoplankton are shown in Tables 2 & 3, with the corresponding percentage of variation relative to the initial estimations. The annual average of chl \(a\) contributed by each pigment group is shown in Fig. 5A,B. Nano-microplankton was ‘brown’, as groups containing chl \(c\) dominated the larger fraction; picoplankton was ‘green’, as chl \(b\)-containing groups
together with cyanobacteria (Synechococcus-like pigments) were the major components. For comparison, diatoms I + II and haptophytes I + II in the nano-microplankton fraction accounted for 67% of the total chl $a$ in the annual term, whereas in the picoplankton these represented only 13%.

Significant changes were found in the contribution of each pigment group to the chl $a$ during the sampled period (Fig. 5C, D). In the nano-microplankton, diatoms I + II increased in proportion from the spring–summer stratification to the September upwelling (Periods I to III), but dropped from the October downwelling to the winter poleward current (IV to VI). Haptophytes I + II followed inverse trends during these periods. The other pigment groups were minor throughout the year (<40% of the total chl $a$), with the exception of 2 seasons: the transitional upwelling in November (V), showing a maximum contribution of dinoflagellates, and the homogeneous poleward current (VI), where cryptophytes and chlorophytes co-dominated together with haptophytes II.

The picoplankton was dominated most of the year by chlorophytes and *synechococcus* (~60% of the total chl $a$), whereas prasinophytes II (urilide-containing prasinophytes, which, with a few exceptions such as the genus *Prasinococcus*, belonged exclusively to the order Mamiellales, e.g. *Micromonas pusilla*, *Bathycoccus prasinos*, *Ostreococcus tauri*; Latasa et al. 2004), haptophytes, and pelagophytes accounted for approximately 25% of the total chl $a$. The most significant changes in pigment group composition were estimated during Periods IV to VI. During the October downwelling (IV) DV chl $a$ from *Prochlorococcus* appeared first, with maximum contributions (together with pelagophytes)

### Table 2. Final ratio matrix (and percentage of variation relative to initial ratios in parentheses; arrows indicate higher or lower ratio) in the nano-microplankton fraction in low light samples and high light samples. (–): pigment absent. For other pigment abbreviations see Fig. 4 legend

|              | Chl $c_3$ | Chl $c_2$ | Chl $c_1$ | Chl $c_2$-EHUX | Chl $c_2$-CHRY | Per | 19 buta | Fuco | Pras | 19 Hex | Allox | Chl $b$
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Low light samples</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diatoms I</td>
<td>–</td>
<td>0.165</td>
<td>0.107</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>
|              |           |           |           |                |                |     |         |      |      |       |       | († 4%)
| Diatoms II   | 0.046     | 0.299     | –         | –              | –              | –   | –       | –    | –    | –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 60%)
| Haptophytes I| 0.249     | 0.171     | –         | 0.097          | –              | –   | –       | –    | –    | –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 12%)
| Haptophytes II| 0.059    | 0.149     | –         | –              | 0.134          | –   | –       | 0.009| 0.129| –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 61%)
| Chlorophytes | –         | –         | –         | –              | –              | –   | –       | –    | –    | –     | –     | –     |
| Prasinophytes| –         | –         | –         | –              | –              | –   | –       | –    | –    | –     | –     | –     |
| Pelagophytes | 0.151     | 0.316     | –         | –              | –              | 0.656| 1.241   | –    | –    | –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 11%)
| Dinoflagellates| –       | 0.111     | –         | –              | 0.442          | –   | –       | –    | –    | –     | –     | –     |
| Cryptophytes | –         | 0.060     | –         | –              | –              | –   | –       | –    | –    | –     | –     | 0.172 |
| **High light samples** |           |           |           |                |                |     |         |      |      |       |       |       |
| Diatoms I    | –         | 0.165     | 0.066     | –              | –              | –   | –       | –    | –    | –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 41%)
| Diatoms II   | 0.046     | 0.299     | –         | –              | –              | –   | –       | –    | –    | –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 61%)
| Haptophytes I| 0.268     | 0.171     | 0.097     | –              | –              | –   | –       | –    | –    | –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 21%)
| Haptophytes II| 0.058    | 0.155     | –         | 0.140          | –              | 0.009| 0.082   | –    | 1.074| –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 62%)
| Chlorophytes | –         | –         | –         | –              | –              | –   | –       | –    | –    | –     | –     | –     |
| Prasinophytes| –         | –         | –         | –              | –              | –   | –       | –    | –    | –     | –     | –     |
| Pelagophytes | 0.103     | 0.316     | –         | –              | –              | 0.510| 0.317   | –    | –    | –     | –     | –     |
|              |           |           |           |                |                |     |         |      |      |       |       | († 33%)
| Dinoflagellates| –       | 0.093     | –         | –              | 0.375          | –   | –       | –    | –    | –     | –     | –     |
| Cryptophytes | –         | 0.060     | –         | –              | –              | –   | –       | –    | –    | –     | –     | 0.172 |
in October and November. During establishment of the homogeneous poleward current (VI) chlorophytes attained their maximum (54\% total chl $a$).

### Chl $a$:C ratios

Chl $a$ estimated by the conventional fluorescence technique (FLUO) was, on average, twice as high as the corresponding HPLC chl $a$ values (FLUO = 2.58 × HPLC + 0.18, $r^2 = 0.83$, n = 165). Consequently, FLUO and HPLC chl $a$:C ratios (Fig. 6A, B) differed significantly, varying between 0.004 and 0.067 and 0.001 and 0.090, respectively.

When calculating chl $a$:C ratios for total ‘diatoms’ (including diatoms I + diatoms II + picoplanktonic diatoms II) estimated by CHEMTAX, we found that ~50 of the samples had chl $a$ data with no, or with unrealistically low, carbon correspondence (chl $a$;C < 0.001). The opposite (C data without chl $a$) occurred for dinoflagellates (Fig. 6C). This means that in one-third of the samples, chl $a$ contributed by diatoms in CHEMTAX did not correspond to any microscopic counts, whereas for dinoflagellates this was quite the opposite. Two distinct trends were revealed when plotting C versus chl $a$ for diatoms (Fig. 6C): chl $a$;C ratios were unusually low in a group of medium to high carbon values registered at the surface (3 m depth, belonging to the HL sample group), whereas other data (mostly from 15 m depth) exhibited higher chl $a$:C ratios.

In the nano-microplankton, some pigment groups showed significant relationships with available carbon

| Pigments (final ratio/initial ratio) | Chl $c_3$ | Chl $c_2$ | Uriol | 19 buta | Fuco | Pras | 19 Hex | Allox | Zea | Chl $b$
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Low light samples</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diatoms II</td>
<td>0.086</td>
<td>0.299</td>
<td>–</td>
<td>–</td>
<td>1.267</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>(† 26%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(† 63%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haptophytes</td>
<td>0.099</td>
<td>0.139</td>
<td>–</td>
<td>–</td>
<td>0.008</td>
<td>0.121</td>
<td>–</td>
<td>0.872</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>(† 35%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(† 7%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlorophytes</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.433</td>
<td></td>
</tr>
<tr>
<td>Prasinophytes</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.438</td>
<td>–</td>
<td>–</td>
<td>0.790</td>
<td></td>
</tr>
<tr>
<td>Prasinophytes II</td>
<td>–</td>
<td>–</td>
<td>0.210</td>
<td>–</td>
<td>–</td>
<td>0.276</td>
<td>–</td>
<td>–</td>
<td>0.914</td>
<td></td>
</tr>
<tr>
<td>Pelagophytes</td>
<td>0.130</td>
<td>0.316</td>
<td>–</td>
<td>0.530</td>
<td>0.437</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>(† 16%)</td>
<td></td>
<td></td>
<td></td>
<td>(† 121%)</td>
<td>(† 65%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cryptophytes</td>
<td>–</td>
<td>0.060</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.172</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td><em>Prochlorococcus</em></td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.457</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td><strong>High light samples</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diatoms II</td>
<td>0.076</td>
<td>0.299</td>
<td>–</td>
<td>–</td>
<td>1.186</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>(† 34%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(† 52%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haptophytes</td>
<td>0.091</td>
<td>0.139</td>
<td>–</td>
<td>0.008</td>
<td>0.073</td>
<td>1.035</td>
<td>–</td>
<td>–</td>
<td>0.336</td>
<td></td>
</tr>
<tr>
<td>(† 40%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(† 51%)</td>
<td></td>
<td></td>
<td>(† 8%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlorophytes</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.439</td>
<td>–</td>
<td>–</td>
<td>0.568</td>
<td></td>
</tr>
<tr>
<td>Prasinophytes</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.206</td>
<td>0.095</td>
<td>–</td>
<td>–</td>
<td>0.523</td>
<td></td>
</tr>
<tr>
<td>Prasinophytes II</td>
<td>–</td>
<td>–</td>
<td>0.206</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.523</td>
<td></td>
</tr>
<tr>
<td>Pelagophytes</td>
<td>0.059</td>
<td>0.308</td>
<td>–</td>
<td>0.471</td>
<td>0.249</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>(† 62%)</td>
<td></td>
<td></td>
<td></td>
<td>(† 208%)</td>
<td>(† 22%)</td>
<td></td>
<td></td>
<td>(† 70%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cryptophytes</td>
<td>–</td>
<td>0.060</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.339</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td><em>Synechococcus</em></td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.939</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td><em>Prochlorococcus</em></td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.760</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

*a Zea:DV chl $a$ ratio of *Prochlorococcus* was not included in CHEMTAX analyses.
biomass. This was the case for diatoms I and diatoms II with diatom C ($r^2 = 0.41$, $p < 0.001$, $n = 167$, in both cases) and dinoflagellates with autotrophic dinoflagellate C ($r^2 = 0.56$, $p < 0.001$, $n = 167$). In the picoplankton, chl a contributed by *Synechococcus* did not match cyanobacterial C, but a significant relationship between zeaxanthin and cyanobacterial C was found ($r^2 = 0.61$, $p < 0.001$, $n = 167$).

**DISCUSSION**

**Size-fractionated phytoplankton biomass**

In open ocean regions of the NASE ecological province (North Atlantic Subtropical Gyral East; Li & Harrison 2001) phytoplankton biomass is largely dominated by picoplankton, specifically by the cyanobacterium *Prochlorococcus* (Zubkov et al. 2000, Morán et al. 2004). Our study, although in the northeasternmost area of NASE, was conducted in temperate neritic waters influenced by the seasonal upwelling and continuous exchange of coastal phytoplankton populations (shortened only during the winter IPC; Álvarez-Salgado et al. 2003). In this region, as in other temperate areas of the world’s oceans (Barlow et al. 2004, Dandonneau et al. 2004), nanoplankton communities usually dominated the autotrophic biomass year-round (Tilstone et al. 2003), while picoplankton remained a low fraction of the total autotrophic C that attains greater importance during the stratified (IV) and homogeneous IPC (VI) (Fig. 3). Only during the summer upwelling, when annual maxima of biomass are registered (Varela 1992), do microplanktonic diatoms sometimes dominate the autotrophic biomass, as occurred in the present study. Teira et al. (2001) reported a similar dominance of nano-microplankton C biomass during the summer upwelling for the same study site. The latter authors also found a major contribution of picoplankton in autumn and
during stratified IPC conditions. Calvo-Díaz et al. (2004) reported 12% picoplankton to total autotrophic C between March and April 2002 at their westernmost stations, slightly north of our study. During the same months (Periods VII & VIII in the present study) we determined ~8% mean contribution of picoplankton to total autotrophic carbon (Fig. 3).

**Seasonal succession of phytoplankton groups: impact of IPC**

Size-fractionated pigments, light microscopy, and carbon analyses enabled us to describe the seasonal succession and concomitant changes in the composition and size structure of phytoplankton communities on the NW Iberian shelf. Spring–summer stratified waters were dominated by nanoplanckton (hapto-

Fig. 6. Carbon (µg C l⁻¹) versus chl a (µg chl a l⁻¹) for: (A) high light (HL) samples, high-performance liquid chromatography (HPLC)- and in situ fluorescence (FLUO)-determined estimates of chl a, (B) low light (LL) samples, HPLC- and in situ FLUO-determined estimates of chl a, and (C) diatoms (including total chl a estimated by CHEMTAX of diatoms I + diatoms II + picoplanktonic diatoms; (●) high chl a:C; (○) low chl a:C dataset) and dinoflagellates. P-levels (at 0.05 significance) and SE for the intercept and slope are shown in each panel, except for diatoms (SE intercept = 0.00880***, SE slope = 0.00041*** for high chl a:C series [●]; SE intercept = 0.17860ns, SE slope = 0.00091** for low chl a:C series [○]).
The homogenous winter IPC (Period VI) should be discussed separately, due to its overall influence on the composition and size distribution of phytoplankton (Figs. 3 & 5C, D), with the lowest chl a and C concentrations during the annual cycle. Low chl a values (Tilstone et al. 2003, Calvo-Díaz et al. 2004) and the dominance of tiny cells have already been reported for the poleward current (Álvarez-Salgado et al. 2003, Calvo-Díaz et al. 2004). The convergence front on the slope created by the IPC effectively isolates coastal phytoplankton from IPC populations (Álvarez-Salgado et al. 2003). It is known that the winter IPC is dominated by small flagellates, while large diatoms and autotrophic dinoflagellates are confined for the most part to inner coastal waters. In accordance with these observations during the homogenous IPC (Period VI), diatoms greatly declined and dinoflagellates were absent. Microplankton was almost absent from the homogenous IPC; marker pigments allowed nanoplankton (chlorophytes, cryptophytes, and Chrysochromulina-type haptophytes II) and picoplankton groups (chlorophytes overwhelming other eukaryotic groups and Synechococcus) associated with the IPC to be identified. Interestingly, prasinophytes II (Order Mamiellales), widely abundant throughout most of the year, disappeared in the winter IPC.

Picoplanktonic populations

The Eastern North Atlantic Central Water of subtropical origin (ENACW$_{st}$), as the source of the poleward flow, appears to provide the inoculums of small cells, low nutrients, low chl a, and the biomass characteristic of the IPC (Álvarez-Salgado et al. 2003). We might expect significant numbers of prochlorophytes to be associated with the poleward flow, given its subtropical origin, but, instead, these organisms fade out during Period VI.

*Prochlorococcus* populations are known to decline north of 45° N, due to lower seawater temperatures (Partensky et al. 1999). Calvo-Díaz et al. (2004) uniquely detected prochlorophytes at their westernmost stations (<43° N), but not on the northern Iberian coast (>43° N), where picoplankton actually dominated TotChl a. These authors identified *Synechococcus* as the most abundant picoplankter in our study area, generally reflecting our results, although, as they did not study eukaryotic groups, chlorophytes were absent from their study.

*Prochlorococcus* was not found in the southern Bay of Biscay, North Spain (Rodríguez et al. 2003b) either inside or surrounding the mesoscale structure of a slope water oceanic eddy (SWODDY), derived from jet-like extensions of the poleward flow. Our annual dataset showed that *Prochlorococcus* was not primarily associated with the core of the winter IPC, as suggested by Calvo-Díaz et al. (2004) in a late winter survey. Rather, *Prochlorococcus* populations appeared on the inner shelf and on the western Iberian coast during the second part of the year, associated with autumn downwelling conditions (October to November in this survey). Later, once the winter IPC had set in, the main *Prochlorococcus* populations would be gradually washed offshore, with a weaker DV chl a signal still detected in the homogenous IPC. Finally, coastal *Prochlorococcus* populations disappeared in late winter, probably due to unfavorable environmental conditions (i.e., decreasing temperatures). Their absence would also be influenced by colder and less salty waters of continental origin, intruding on the inner shelf areas as a buoyancy plume (Peliz et al. 2005). In fact, DV chl a has never been reported in previous HPLC pigment analyses of the Rías de Pontevedra (Rodríguez et al. 2003a) or Arousa (M. Zapata pers. comm.).

Phytoplankton pigments at open ocean and coastal sites of the NE Atlantic Ocean have been reported elsewhere (e.g. Barlow et al. 2004, Gibb et al. 2000, Riegman & Kraay 2001). These studies generally indicate a dominance of 19'-hexanoyloxy-fucoxanthin (19 Hex) and fucxanthin (Fuco) (prymnesiophytes and diatoms) in the temperate open ocean (e.g. Gibb et al. 2000, Riegman & Kraay 2001). Coastal sites are more heterogeneous, and several groups (e.g. dinoflagellates, cryptophytes, chl b-containing algae, and *Synechococcus*) alternate with diatoms as the most abundant taxa (Schlüter et al. 2000).

Much less information is available for picoeukaryotes than for their prokaryotic counterparts (Veldhuis et al. 2005). A few authors who have performed size-fractionated pigment analyses on NE Atlantic samples (Brunet & Lizon 2003, Rodríguez et al. 2003a, Not et al. 2004) highlight the prevalence of chlorophytes (particularly prasinophytes; Rodríguez et al. 2003a, Not et al. 2004) in coastal picoeukaryotic populations, along with variable amounts of 19'-butanoloxy-fucoxanthin 19 Buta-, 19 Hex- (cryptophytes and haptophytes; Brunet & Lizon 2003), and Fuco-containing groups (diatom-like pigments; Rodríguez et al. 2003a).

Varela (1992) enumerates Chlorophyceae, Primsiophyceae, Chrysophyceae, and Cryptophyceae as some of the most common microflagellates (<10 µm) identified on the Galician shelf. However, to our knowledge, this is the first report detailing specific size-fractionated pigment groups and their abundance in the NASE, particularly for the NW Iberian peninsula. Until now the results available originated from estuarine samples (Ría de Pontevedra; Rodríguez et al. 2003a). The most important differences between estu-
arine and neritic waters are: (1) the contribution of picoplankton to TotChl a is double that found in our study (29% ± 17% in comparison with 13% ± 10% TotChl a in the Ria de Pontevedra); (2) the contribution of the picoplanktonic pigment groups diatoms I + II to chl a in the ría is 4 times higher (>25% TotChl a relative to 6% in the present study); and (3) the exclusive presence of Synechococcus-type cyanobacteria in the ría, whereas Prochlorococcus is also present in shelf waters. These differences are explained by the environmental conditions in the rías (semi-enclosed and shallower embayments influenced by river runoff and continental discharges), which increase the productivity and size of phytoplankton relative to offshore shelf waters (Tilstone et al. 2003). Most of the new pigment information we gained in this study originates from a variety of chl c compounds (chl c1, chl c2, chl c2-MGDG [14:0/18:4] & [14:0/18:4]). Our results demonstrate that chl c1 helps to discriminate a group called ‘diatoms I’ from other chl c-containing groups (see also Riegman & Kraay 2001, Veldhuis & Kraay 2004). Additional diagnostic pigments for haptophytes, the non-polar chls c, first reported in the NE Atlantic (Gieskes & Kraay 1986), are also readily detected in natural samples.

Chl c2-MGDG [14:0/14:0], the marker for haptophytes II and almost exclusive to the genus Chrysochromulina (Zapata et al. 2001, 2004), was widely present in the nano-microplankton fraction. This suggests that Chrysochromulina species were a dominant component of nanoplanctonic haptophytes, in agreement with a previous publication concerning the southern Bay of Biscay (Rodriguez et al. 2003b). Chl c2-MGDG [14:0/18:4] from haptophytes I, a typical winter group in our study, pointed towards Emiliania huxleyi, the genus Phaeocystis, and to certain other Chrysochromulina species in the same season (Zapata et al. 2004). In turn, non-polar chls c were barely detected in the picoplankton, and we could not hypothesize on a particular haptophyte genus.

**Pigment ratios**

**Accessory pigment:chl a**

As pigment:chl a ratios were determined based on initial estimates from cultures (Table 1), output CHEMTAX ratios (Tables 2 & 3) do not exclusively indicate photoacclimation trends. However, these values, on the whole, exhibited the expected light-dependent responses (i.e. lower accessory chlorophylls in HL vs. LL, although some exceptions have been reported; e.g. Schlüter et al. 2000). In the case of Fuco and 19 Hex, a general pattern from cultures (Schlüter et al. 2000, Henriksen et al. 2002) and CHEMTAX studies (Llewellyn et al. 2005) indicates that diatoms and haptophytes increase Fuco:chl a ratios in low light and deep samples, the converse being true for 19 Hex. This result was also obtained in our study, with the exception of haptophytes II, which retained higher 19 Hex:chl a ratios in LL samples. This pigment group (and picoplanktonic haptophytes) was estimated using Chrysochromulina ratios, a haptophyte genus that does not follow the general rule above (Henriksen et al. 2002). In a recent North Atlantic survey (Lutz et al. 2003), no significant differences were found in 19 Hex:chl a ratios between surface and deep samples. The photoacclimation patterns in haptophytes are thus species dependent and a likely explanation for the variable or constant 19 Hex:chl a ratios along vertical profiles in contrasting oceanic areas.

**Chl a:C**

The usefulness of chemotaxonomy to estimate phytoplankton composition and biomass has been assessed comparing pigment analyses with carbon estimated from light microscopy (Garibotti et al. 2003, Llewellyn et al. 2005) and flow cytometry (Veldhuis et al. 2005). Despite variations of pigment content relative to carbon and intrinsic discrepancies between techniques, more accurate characterization of phytoplankton assemblages and important ecophysiological information can be gained from chl a:C ratios, such as carbon-based ocean productivity from remote sensing (Garibotti et al. 2003). In the present paper, the range of variability in estimated chl a:C ratios was similar to that reported by Llewellyn et al. (2005) for NE Atlantic samples. Different subsets of chl a:C data have been distinguished in previous studies to accommodate for variations in pigment content per cell due to environmental factors and changes in species composition (Garibotti et al. 2003, Llewellyn et al. 2005). In the present study, a stronger linear relationship between chl a and C values was obtained for HL samples, as these included fewer fluctuating irradiances than LL samples. Lower variability in cellular pigment content relative to C would explain this stronger relationship in HL samples, as was also discussed by Llewellyn et al. (2005).

The slope of the HPLC–chl a:C relationship (Fig. 6) was in the lower range obtained for algal cultures (Cloern et al. 1995), the slope of FLUO–chl a:C ratios being closer to previous results for the NE Atlantic (Llewellyn et al. 2005). Apart from intrinsic errors in carbon determination, this may suggest that our HPLC values were somewhat lower than expected for this area. It must also be borne in mind that systematic HPLC overestimates of chl a, in comparison with HPLC estimates, are usually reported due to interferences with
DV chl a and/or high concentrations of chls c, as may also be the case in our study (Latasa et al. 1996, Stuart et al. 2004). Carbon and chl a biomass estimates of individual pigment groups agree in some cases, e.g. diatoms, but significant discrepancies have been reported for other classes (Garibotti et al. 2003, Llewellyn et al. 2005). In the case of diatoms, C and chl a biomass estimates match fairly well, except when significant contributions to Fuco by other groups are observed (Garibotti et al. 2003). The low chl a:C dataset for diatoms in our surface samples (Fig. 6C) corresponded with different dominant species, so photoc acculation reduction of chl a:C ratios in HL conditions would be the likely explanation for their distinct chl a:C ratios (Llewellyn et al. 2005). Thus, for diatoms only, and after splitting natural samples into 2 depth groups, carbon biomass estimates may be inferred from their contribution to chl a as calculated by CHEMTAX. In turn, the relationship between chl a and C estimates for dinoflagellates was blurred, probably due to difficulties in distinguishing some autotrophic and heterotrophic species and the presence of non-peridinin-containing species (e.g. Karenia mikimotoi, not included in the pigment group dinoflagellates), as discussed previously (Garibotti et al. 2003, Llewellyn et al. 2005).

CONCLUSIONS

Size-fractionated carbon analyses indicated that nanoplanctonic organisms accounted for a major portion of the autotrophic biomass in the NW Iberian shelf, and chemotaxonomy revealed that diatoms (chl c3-type pigment) and haptophytes (particularly Chrysochromulina-type pigment) dominated in terms of the relative chl a contribution. However, during some hydrographic periods, particularly once the IPC signal in autumn–winter has been detected, major changes in size structure and composition of phytoplankton populations were evidenced, with a simultaneous reduction of microplanktonic populations and higher abundance of picoplanktonic cells (namely chlorophytes) during the IPC period. We conclude that a seasonal succession in the smaller groups takes place in the study area, similar to that observed for larger organisms. However, the taxonomical and pigmented composition remains quite different in picoeukaryotes, which are dominated by chl b-containing groups (chlorophytes and prasinophytes), compared with nano-microeukaryotes, which characterized by chl c-containing organisms. The specific gene losses of green plastids relative to red plastids have been suggested as the main reason for their increased portability and the dominance of chl c-containing plastids in modern eukaryotic algae (Grzebyk et al. 2003). However, these reductive traits in green plastids would imply an evolutionary advantage in a picoplanktonic cell whose tiny size forces reduction and simplification at all levels. Elucidating the ecophysiological and molecular traits that underlie this apparent success of chl b- versus chl c-containing picoeukaryotes constitutes one of the most promising challenges in the coming era of molecular-ecological studies of phytoplankton.

Acknowledgements. We thank the members of the oceanography group at the Instituto de Investigaciones Marinas who participated in the sampling and analysis of nutrients, and M. Penabad for her skilful technical help in HPLC analysis. We thank E. A. Barbour for spelling and grammar checks. We also thank 2 anonymous reviewers whose comments significantly improved the final paper. Financial support for this work came from the Spanish DYBAGA project (MAR99-1039-C02-01) and from the EU HABILE project (EVK3-CT-2001-00063). B.G.C. was funded by a pre-doctoral I3P fellowship of CSIC-ESF.

LITERATURE CITED


Teira E, Serret P, Fernandez E (2001) Phytoplankton size-structure, particulate and dissolved organic carbon pro-


Editorial responsibility: Fereidoun Rassoulzadegan (Contributing Editor), Villefranche-sur-Mer, France

Submitted: July 27, 2005; Accepted: February 8, 2006
Proofs received from author(s): September 1, 2006