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Aurélie Pénaud, Frédérique Eynaud, María-Fernanda Sánchez-Goñi, B. Malaize, Jean-Louis Turon, L. Rossignol

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Contrasting sea-surface responses between the western Mediterranean Sea and eastern subtropical latitudes of the North Atlantic during abrupt climatic events of MIS 3

Penaud A. (a,b) *, Eynaud F. (b), Sánchez-Goñi M. (c), Malaizé B. (b), Turon J.L. (b), Rossignol L. (b)

(a) UMR 6538 Domaines Océaniques, IUEN-UBO, F-29280 Plouzané, France
(b) UMR 5805 EPOC, Université de Bordeaux 1, F-33405 Talence, France
(c) EPHE, UMR 5805 EPOC, Talence, F-33405 Talence, France

*Corresponding author. Tel.: +33-298-498-741; fax: +33-298-498-760.
E-mail address: aurelie.penaud@univ-brest.fr.
ABSTRACT

Dinoflagellate cyst (dinocyst) analysis was conducted on two cores from the SW Iberian margin and central Alboran Sea from which high quality records of Marine Isotope Stage 3 have been previously derived. Our aim in this study is to compare the dinocyst signature between 50 and 25 ka BP with existing datasets of foraminiferal and geochemical proxies related to hydrological parameters. Quantitative reconstructions of sea-surface temperatures (SSTs) and salinities (SSS) based on dinocysts are performed for the first time in this area. The results are compared to SSTs derived from planktonic foraminifera and alkenone measurements, and to SSS calculated from planktonic $\delta^{18}O$ and foraminiferal SST. Significant oscillations related to Dansgaard-Oeschger cycles are recorded in both cores. Dinocyst-derived hydrological parameters exhibit synchronous fluctuations and similar values to those derived from the other methods, in particular when considering quantitative reconstructions for February based on foraminifera and dinocysts. Our study shows that the influence of subpolar waters was felt during each Greenland Stadial (GS) off Portugal, and that the amplification of the Heinrich Stadial cooling in the Alboran Sea was related to the penetration of subpolar waters through the Strait of Gibraltar. During Greenland Interstadials (GI), we provide evidence for the occurrence of warm and nutrient-rich sea-surface waters in the Alboran Sea, probably due to gyre-induced upwelling. Finally, the difference between August and February dinocyst SST estimates suggests higher seasonal contrasts during GS compared to GI at the two core sites. Additionally, precession appears to have an imprint on dinocyst-derived long-term seasonality record. However, this observation needs to be confirmed by longer records.

KEYWORDS: Dinocysts; SST and SSS quantification; Dansgaard-Oeschger; Greenland and Heinrich stadials; Alboran Sea; Iberian margin.
1. Introduction

The climate of the last glacial was characterised by a distinctive oscillatory mode, the so-called Dansgaard-Oeschger (D-O) cycles, originally described and defined in Greenland ice cores (Johnsen et al., 1992; Dansgaard et al., 1993; Groottes et al., 1993) and paced by an empirical 1470-year factor, the nature of which being presently not known (Schulz et al., 1999; Moreno et al., 2005; Voelker et al., 2006). Greenland Interstadials (GI) and Greenland Stadials (GS) including Heinrich Stadials (HS, following Sánchez-Goñi and Harrison, 2010), have previously been observed in marine climate records worldwide (e.g. Voelker et al., 2002; Hemming, 2004; Clement and Peterson, 2008). The impacts of HS, which primarily correspond to large ice-sheet collapses, are now fairly well characterised in the NE Atlantic and Mediterranean areas. Their occurrence is associated with enhanced aridity in the adjacent borderlands (Combourieu-Nebout et al., 2002; Sánchez-Goñi et al., 2002), increases in northward Saharan dust transport (Moreno et al., 2002; Bout-Roumazeilles et al., 2007) and sea-surface temperature coolings in the western Mediterranean Sea (e.g. Cacho et al., 1999) and on the Portuguese margin (e.g. Cayre et al., 1999; Pailler and Bard, 2002; de Abreu et al., 2003). However, the forcing mechanisms behind the initiation of HS and the inferred reorganisation of Meridional Overturning Circulation within the last glacial period are still a matter of debate (Kageyama et al., 2009). Furthermore, a full understanding of this rapid transmission of millennial-scale climatic variability from boreal to subtropical latitudes, involving both atmospheric and oceanic processes, remains elusive. It has been demonstrated that convection in the western Mediterranean, and thus export of Mediterranean Outflow Water (MOW), was strengthened during GS of the last 50 ka (Cacho et al., 2000, 2006; Sierro et al., 2005; Voelker et al., 2006; Toucanne et al., 2007; Frigola et al., 2008). Consequently, it has been suggested that the Mediterranean could act as a major trigger for
the Meridional Overturning Circulation to switch from stadial to interstadial mode through the influence of MOW, providing saline water to the North Atlantic at times when the Meridional Overturning Circulation was partially interrupted (Johnson, 1997; Bigg and Wadley, 2001; Rogerson et al., 2006; Voelker et al., 2006). The production of MOW is mainly controlled by the salinity budget of the Mediterranean Sea which depends on the climate characteristics over the region and therefore on dominant modes of climatic variability, notably the North Atlantic Oscillation (NAO). At present, the NAO pattern, oscillating at decadal and centennial scales, has a strong influence on climate over a large part of Europe during winter, by affecting the storm tracks and the associated relative moisture over the Mediterranean and northern Europe (e.g. Hurrell, 1995; Serreze et al., 1997; Osborn et al., 1999). Over the eastern Atlantic subtropical sector, it has been argued that this oscillation also has an imprint at centennial or millennial timescales, and has been implicated in abrupt climatic events of the last glacial (Sánchez-Goñi et al., 2002; Moreno et al., 2005; Bout-Roumazeilles et al. 2007; Daniau et al., 2007; Naughton et al., 2009). Consequently, a change between two periods, each of them characterised by sustained frequencies of a particular atmospheric configuration over several centuries, may have caused significant variations of hydrological exchanges through time between the Atlantic and the Mediterranean Sea (Moreno et al., 2005).

Previous high resolution palynological analysis of dinoflagellate cysts (dinocysts) and pollen on core MD95-2042, collected off Portugal, indicate abrupt changes in sea surface and atmospheric conditions, respectively, in response to D-O cycles (Eynaud, 1999; Sánchez-Goñi et al., 2000). Additionally, a climatic contrast has been observed with respect to precipitation between the SW Iberian margin and the Alboran region, with dryer conditions in southeastern Iberia than in its southwestern part (Sánchez-Goñi et al., 2002). However, these studies do not quantify either changes in SST and SSS, or discuss the possible impact of the observed precipitation gradient on
the hydrological conditions of both sides of the Strait of Gibraltar between 25 and 50 ka BP. To fill this gap, we have analyzed dinocyst assemblages from core MD95-2043, located in the Alboran Sea, and applied transfer functions. Furthermore, we have tested, for subtropical latitudes, the robustness of quantitative paleotemperature and paleosalinity reconstructions inferred from dinocysts against other proxy reconstructions (i.e. dinocyst- versus foraminifera- versus alkenone- derived SST, and dinocyst- derived SSS versus SSS estimates calculated on the basis of the $\delta^{18}O$ of Globigerina bulloides associated with foraminiferal SST).
2. Core locations and present-day environmental settings

Calypso cores MD95-2043 (36°8.6’N; 2°37.3’W; 1841 m water depth; 36 m long) and MD95-2042 (37°48’N; 10°10’W; 3146 m water depth; 39.56 m long) were retrieved from the central Alboran Sea and the SW Iberian margin, respectively (Fig. 1), and were both collected by the oceanographic R/V Marion Dufresne during the 1995 International Marine Global Change Studies I (IMAGES I) cruise (Bassinot and Labeyrie, 1996).

The Mediterranean Sea is located between the path of the mid-latitude westerlies which dominate northern and central Europe, and the Azores High. At present, this basin experiences a typical seasonal cycle marked by cool-wet winters, and warm-dry summers. The resulting Mediterranean water budget is marked by a strong freshwater deficit due to a net excess evaporation (Béthoux, 1979, 1984). During winter and spring, intense cold and dry continental air outbursts induce strong evaporation and cooling in the northern Mediterranean Sea, and thus an increased density of surface water masses. As a result, surface waters sink in several specific Mediterranean regions, flow westward, and finally form the Mediterranean Outflow Water (MOW). The outflow current is exported at depth towards the Atlantic Ocean, through the narrow Strait of Gibraltar (main sill depth of 280 m) and is split into two stability levels: an upper core centred between 500 m and 800 m, and a more saline and dense lower core found between 1000 and 1400 m (Ambar et al., 2002). The MOW can be traced in the North Atlantic as a salinity and temperature maximum at about 1 km depth (e.g. Hill and Mitchelson-Jacob, 1993; Iorga and Lorzier, 1999; O’Neill-Baringer and Price, 1999), and mixes progressively with North Atlantic Intermediate Waters (NAIW; Mauritzen, 1996). In the opposite direction, a North-Atlantic low-salinity surface current penetrates the Alboran Sea and mixes with Levantine Intermediate Waters (which resurface in the westernmost part of the Alboran Sea), forming the so-called Modified Atlantic
Waters (MAW). The strong flow of MAW along the coast of Spain initiates the formation of two anticyclonic gyres, the Western and Eastern Alboran Gyres (WAG and EAG; Fig. 1) whose position and intensity fluctuate at a seasonal scale. The degree of development of the EAG, under which core MD95-2043 is located (Fig. 1), controls the position and intensity of the permanent Almeria-Oran Front (AOF; Fig. 1) (Tintoré et al., 1988; Rohling et al., 1995, 2009; Viúdez and Tintoré, 1995). Upwelling cells occur along the AOF, this front marking the deflection of MAW along the Algerian margin, forming the Algerian Current (AC; Fig. 1).

The main modern hydrological structures and currents of the western Iberian margin form part of the North Atlantic Eastern Boundary Current (Fig. 1; Peliz et al., 2005; Relvas et al., 2007). They are driven by the North Atlantic subtropical gyre intensity in relation to the seasonal displacement of the Azores High (e.g. Fiúza, 1984; Fiúza et al., 1998). Core MD95-2042 is located in the seasonal coastal upwelling band of the Portugal-Canary eastern boundary upwelling system that is active mainly from April to October (Aristegui et al., 2005; Peliz et al., 2005). The upwelling predominantly receives North Atlantic Central Waters (NACW) and, in part, also MOW (Sánchez and Relvas, 2003). Occasionally, during winter months, a warm northward-flowing surface current known as the Iberian Poleward Current or Portugal Coastal Counter Current is formed by coastal convergence along the western Iberian margin. This phenomenon occurs when weak northerlies are interspersed with strong south-southwesterly winds (e.g. Fiúza et al., 1998; Peliz et al., 2005). This current can be traced as far as the Bay of Biscay where it is known as the “Navidad Current”. Winter warming in the southern Bay of Biscay during Navidad years has been correlated with low values of the NAO index and this current was found to extend from Portugal to Norway in exceptional Navidad years (Garcia-Soto et al., 2002).
Modern sea-surface conditions of the SW Iberian margin and Alboran Sea are characterised respectively by mean annual salinities of 36.5 and 37.1 psu (world dataset atlas compiled by Schmidt, 1999 and Bigg and Rohling, 2000, http://data.giss.nasa.gov/o18data: Craig and Gordon dataset for the Iberian margin and C. Pierre dataset for the Alboran Sea), February SSTs are around 15.2 and 14.7°C, respectively, and August SSTs are around 20 and 23.7°C (WOA 2001).
3. Methodology

3.1. Chronostratigraphy

Cores MD95-2043 (central Alboran Sea) and MD95-2042 (SW Iberian margin), characterised by high sedimentation rates, are composed mainly of calcareous hemipelagic clays and have yielded high resolution paleoceanographic records (e.g. Cacho et al., 1999 and Shackleton et al., 2000, respectively).

The age model of core MD95-2042 is derived from 16 AMS $^{14}$C dates and, for the period beyond AMS $^{14}$C range, by graphical tuning of the MD95-2042 planktonic $\delta^{18}$O record with Greenland ice core $\delta^{18}$O records. The GISP 2 chronology (Bard et al., 2004) was used between 26 and 47 ka, and the GRIPSS09sea chronology (Shackleton et al., 2004) was applied between 47 and 77 ka. Further details of the age models are given in Daniau et al. (2007) and Sánchez-Goñi et al. (2008).

The age model of core MD95-2043 is based on 21 AMS $^{14}$C dates and graphical tuning of the MD95-2043 alkenone SST record to GISP2 $\delta^{18}$O (Cacho et al., 1999). Due to the recent improvement of the chronology of the SW Iberian margin core, some discrepancies in the two age models appear in the dates of the climatic events at around 40 ka BP. In these cases, we have slightly modified the chronology of the Alboran Sea record for the time interval between 40 and 50 ka, according to Sánchez-Goñi et al. (2009), in order to align the latter sedimentary sequence with that of the SW Iberian margin.

3.2. Dinoflagellate analysis
Dinoflagellates are flagellate protists that occur in both marine and freshwater environments, and which thrive in the depth range of 18-100 m in oceanic domains (Dodge and Harland, 1991; Raine et al., 2002) and 0-10 m at the coastline. Water turbulence is greater in the neritic zone and represents a limiting factor for light penetration and thus the maximum depth of dinoflagellate habitats. Dinoflagellates reproduce primarily through fission, but sexual reproduction also occurs resulting in a resting cyst which is preserved in sediments (Dodge et Harland, 1991; Head, 1996).

The distribution of dinoflagellate cysts (dinocysts) reflects physico-chemical parameters of the overlying water masses (temperature, salinity, sea-ice cover, seasonality and nutrient availability) (e.g. Turon, 1984; Mudie, 1992; Matthiessen, 1995; Rochon et al., 1999; Devillers and de Vernal, 2000; Zonneveld et al., 2001; Marret and Zonneveld, 2003; de Vernal and Marret, 2007).

Dinocyst analysis is therefore an essential tool for reconstructing Quaternary paleoenvironments (e.g. Turon, 1984; Turon and Londeix, 1988; Eynaud et al., 2000, 2004, 2009; de Vernal et al., 1997, 2001, 2005; Mudie et al., 2002, 2004; Grøsfjeld et al., 2006; Penaud et al., 2008, 2009, 2010).

Dinocyst assemblages were characterised at the species level on the sediment fraction smaller than 150 µm on 61 palynological slides for core MD95-2043 (this study) and on 71 slides for core MD95-2042 (Eynaud, 1999; Sánchez-Goñi et al., 2000). The preparation technique followed the protocol described by de Vernal et al. (1999) and Rochon et al. (1999), slightly modified at the EPOC laboratory (Castera and Turon, http://www.epoc.u-bordeaux.fr/index.php?lang=fr&page=eq_paleo26). Each subsample of 8 cm³ was weighed, dried overnight and then weighed again to obtain dry weight. Subsamples were then washed through a 150 µm sieve and the fraction smaller than 150 µm was used for palynological analysis. After chemical and physical treatments (cold HCl, cold HF and sieving through single-use 10 µm nylon
mesh screens), the final residue was mounted between slide and coverslip with glycerine jelly
coloured with fuschin. Identifications and counts were performed using a Leica DM 6000
microscope at 400 × magnification, and counts aimed to reach 300 specimens wherever possible
for each sample (cf. Appendix A). Taxonomic identifications are consistent with those of
Fensome et al. (1998) and Fensome and Williams (2004). *Brigantedinium* cysts are grouped
together and include all spherical brown cysts, since it is rarely possible to identify them at the
species level due to their crumbled aspect which masks the archeopyle.

Dinocyst assemblages were described by the percentages of each species calculated on the basis
of the total dinocyst sum including the few unidentified taxa and excluding pre-Quaternary cysts.
Palynomorph concentrations were calculated using the marker grain method (de Vernal et al.,
1999). Aliquot volumes of *Lycopodium* spores were added to each sample before chemical
treatments in order to obtain palynomorph concentrations.

**3.3. Quantitative reconstructions of sea-surface parameters**

**3.3.1. Dinocyst SST and SSS reconstructions**

We used a transfer function based on the Modern Analogue Technique (MAT) to reconstruct sea-
surface hydrological parameters from dinocysts. The MAT principally uses the statistical distance
between fossil (paleoceanographic record) and current (modern database) assemblages. The
calculation of past hydrological parameters relies on a weighted average of the SST values of the
best modern analogues found (minimum and maximum number of analogues imposed in the
transfer functions are 5; cf. Appendix B for the list of analogues found for each sample). The
maximum weight is given for the closest analogue in terms of statistical distance. The reader is
referred to Guiot and de Vernal (2007) for a review of theory of transfer functions and to de
Vernal et al. (2001, 2005) for a step by step description of the application of transfer functions to
dinocysts, including discussion about the degree of accuracy of the method.

The dinocyst transfer function used in this work (cf. de Vernal et al., 2005; GEOTOP website:
http://www.unites.uqam.ca/geotop/monographie_n940/eng/index.shtml) is derived from a
modern database comprising 60 dinocyst species and 940 stations from the North Atlantic, Arctic
and North Pacific oceans and their adjacent seas, including the Mediterranean Sea (84 stations
including station “M1039”, cf. Fig. 1; Mangin, 2002), as well as epicontinental environments
such as the Estuary and Gulf of St. Lawrence, the Bering Sea and the Hudson Bay. The transfer
function (n=940) is run under the “3Pbase” software (Guiot and Goeury, 1996). This software
was originally developed for pollen-based quantitative climate reconstruction (http://www.imep-
cnrs.com/pages/3pbase.htm) and was subsequently applied to dinocyst assemblages (e.g. de
Vernal et al., 2001, 2005). An index “Dmin”, provided by the software “3PBase”, allows testing
the reliability of the reconstructions (cf. de Vernal et al., 2005). This index describes, for each
sample analyzed, the distance between the closest analogue found by the transfer function and the
fossil assemblage. A threshold value is calculated from the calibration of the database for the
identification of non-similar or very bad analogues. This threshold value provided by the
software “3PBase” is 71.72 and, below this value, the similarity between the modern data and the
fossil record is considered significant (cf. Appendix B for the Dmin values calculated for each
sample). The authors caution that the full reference set has been used for the calculations of
hydrological parameters without any regional selection of samples within the modern database. In
this study, we present February and August mean sea-surface temperatures (SST, with prediction
errors of ±1.2°C and ±1.8°C respectively), and February and August mean sea-surface salinities (SSS, with prediction errors of ±1.7 for both).

3.3.2. Foraminiferal SST and SSS reconstructions

In the same way, we used a transfer function based on the MAT to reconstruct foraminiferal SST from the foraminiferal assemblages of cores MD95-2042 (Cayre et al., 1999) and MD95-2043 (Pérez-Folgado et al., 2003). Calculation of past hydrological parameters relies on a weighted average of the SST values of the best 5 modern analogues found systematically. This transfer function has been developed at EPOC laboratory (“Environnements et Paléoenvironnements OCéaniques”, Bordeaux1 University, France). The MAT (e.g. Kucera, 2007) is run under the “R” software using a script first developed for dinocyst transfer functions (http://www.cerege.fr/IMG/pdf/ECCOR_StatRAvr08.pdf). The modern database relies on a modern database of 1007 modern assemblages and is derived from the ones developed separately for the North Atlantic and the Mediterranean seas during the MARGO project (Kucera et al., 2005; Hayes et al., 2005). These databases were merged together to offer a larger set of analogues for subtropical reconstructions over the last glacial period notably (Eynaud et al., 2009; Matsuzaki et al., in press). Modern hydrological parameters were requested from the WOA 2008 database using the tool developed during the MARGO project (http://www.geo.uni-bremen.de/geomod/staff/csn/woasample.html). This method allows the reconstruction of annual and seasonal (winter, spring, summer and fall) SST. In this paper, we present winter and summer mean SST with a prediction error of ±1.2°C and ±1.3°C respectively, and annual SST with a prediction error of ±1.1°C (Eynaud et al, 2009; Matsuzaki et al., in press).
SSS estimates have been derived using the approach summarised in Malaizé and Caley (2009) for calibration of the salinity-water isotope relationship. Craig and Gordon (1965) established the first salinity-water isotope relationship as follows:
\[ \delta^{18}\text{O}_{\text{sw}} = 0.66 \text{SSS} - 23.5 \]

Since this pioneer work, many measurements have been performed and many different relationships, with different slopes, have been derived for different oceans (Ostlund et al., 1987, Schmidt, 1999; LeGrande and Schmidt, 2006). Discrepancies in slope and in intercept values are due to local characteristics, on spatial and temporal scales. For the Atlantic Ocean, an important dataset, established from several decades of oceanographic measurements, allows revision of the Craig and Gordon (1965) calibration and the determination of a new relationship, with a mean slope component of 0.558.

In order to estimate past changes in oceanic \( \delta^{18}\text{O}_{\text{sw}} \), Epstein et al. (1953) established a paleotemperature equation which links temperature with the isotopic composition of calcite \( (\delta^{18}\text{O}_{\text{c}}) \) in calcareous shells (e.g. foraminifera) and the ambient waters \( (\delta^{18}\text{O}_{\text{sw}}) \). Shackleton and Opdyke (1973) have adapted the equation of Epstein et al. (1953) as follows:
\[
T = 16.9 - 4.38 (\delta^{18}\text{O}_{\text{c}} - \delta^{18}\text{O}_{\text{sw}}) + 0.13 (\delta^{18}\text{O}_{\text{c}} - \delta^{18}\text{O}_{\text{sw}})^2
\]

Stable isotope \( \delta^{18}\text{O}_{\text{c}} \) measurements in cores MD95-2042 (Cayre et al., 1999; Shackleton et al., 2000) and MD95-2043 (Cacho et al., 1999) were carried out on \( G. \text{bulloides} \) monospecific samples. Following the pioneer study of Duplessy et al. (1991), we corrected the summer temperature by 1°C for \( G. \text{bulloides} \) species. By solving the Shackleton and Opdyke (1973) equation, we obtain \( \delta^{18}\text{O}_{\text{sw}} \) variations, which integrate the signal of both local and global variations. To remove the global influence of continental ice volume, we used estimations of past global \( \delta^{18}\text{O}_{\text{sw}} \) changes based on benthic isotopic records and coral terrace growth (Waelbroeck et
al., 2002), and used the modern δ\textsuperscript{18}Osw values (1.2 ‰ for the Alboran Sea and 0.97 ‰ for the Iberian margin) extracted from the world dataset atlas compiled by Schmidt (1999) and Bigg and Rohling (2000) (C. Pierre dataset for the Alboran Sea, and Craig and Gordon dataset for the Iberian margin, from http://data.giss.nasa.gov/o18data). The residual δ\textsuperscript{18}Osw “ice-corrected” signal is then converted into quantitative SSS values for both cores, using a mean slope of 0.558. SSS uncertainties of around 1 psu derive from the prediction error of the foraminiferal SST reconstructions (1.8°C uncertainty linked with August SST reconstructions corresponds to a 0.45 ‰ uncertainty on a δ scale), as well as on uncertainties in the global δ\textsuperscript{18}Osw changes linked to sea level changes (0.15 ‰).
4. Dinocyst assemblages through time

Concerning the species of the genus *Brigantedinium* spp., round brown cysts formed by heterotrophic dinoflagellates, it is important to take into account the fact that high occurrences of *Brigantedinium* spp. can be linked to better preservation under hypoxic or anoxic bottom conditions (Combourieu-Nebout et al., 1998; Zonneveld et al., 2001; Kodrans-Nsiah et al., 2008). We can not exclude that *Brigantedinium* spp. may have been subjected to dissolution since deep ventilation in the western Mediterranean Sea (and thus oxidation processes) during GS was stronger than during GI of the last glacial (cf. benthic $\delta^{13}$C in Fig. 4; Cacho et al., 2006).

However, *Brigantedinium* spp. are present throughout the Alboran core without any clear trend between few percentages during GS and high percentages during GI. We can thus expect to reconstruct dinocyst assemblages without significant preservational problems.

When comparing both cores (Fig. 2), we note extremely high relative abundances of *Brigantedinium* spp. (30-89%) on the SW Iberian margin (Fig. 2b). This may reflect the high productivity induced by permanent or intensified upwelling cells along the Portuguese margin during the last glacial (e.g. Abrantes, 2000; Lebreiro et al., 1997); this zone today is characterised by a seasonal upwelling system (Peliz et al., 2005) and *Brigantedinium* spp. percentages only reach a maximum of 2-5% in the modern database. Percentages of these cysts fluctuate more in the Alboran Sea, with pronounced oscillations (average of 19% and peaks reaching 30 to 45%) (Fig. 2a).

4.1. Dinocysts occurring during Greenland Stadials
Dinocysts which feature prominently during GS include *Bitectatodinium tepikiense*, *Spiniferites lazus*, *Spiniferites elongatus*, *Impagidinium aculeatum*, and *Operculodinium centrocarpum* in the Alboran Sea (Fig. 2a), and *B. tepikiense*, *S. elongatus*, cysts of *Pentapharsodiniumdalei*, *Lingulodinium machaerophorum* and *Nematosphaeropsis labyrinthus* on the SW Iberian margin (Fig. 2b). It is also interesting to note that maximal dinocyst concentrations occur during GS off Portugal (most notably during HS 4; Fig. 2b) contrarily to what we observe in the Alboran record (Fig. 2a).

In the Alboran Sea, *B. tepikiense* increases during HS events, especially HS 3 and HS 4 (Fig. 2a), whereas it occurs during each GS on the SW Iberian margin, where it consistently makes up almost 10% of the total dinocyst assemblage (Fig. 2b). Today, *B. tepikiense* is mainly distributed between 40°N and 60°N in temperate to sub-arctic environments of the North Atlantic, with the highest abundances found south of the Gulf of St. Lawrence in coastal environments of Nova Scotia and the Gulf of Maine (Wall et al., 1977; Mudie, 1992). This species is characteristic of areas marked by strong seasonal contrasts, with freezing winter SSTs and up to 16°C summer SST (Rochon et al., 1999; de Vernal et al., 2005), and enhanced surface water stratification (Rochon et al., 1999; Marret and Zonneveld, 2003). Previous results from the North Atlantic (Zaragosi et al., 2001; Penaud et al., 2009), the SW Iberian margin (Sánchez-Goñi et al., 2000; Turon et al., 2003), and the western Mediterranean Sea (Turon and Londeix, 1988; Combourieu-Nebout et al., 2002) have shown increased abundances of *B. tepikiense* during HS.

*S. elongatus* develops during each GS with low percentages reaching 2-5% in the Alboran Sea (Fig. 2a) and 1-2% on the western Iberian margin (Fig. 2b). Maximum present-day occurrences of this species are observed in the Baffin Bay and Barents Sea, and this taxon is generally related
to cool to temperate conditions (Rochon et al., 1999). The significant occurrence of *B. tepikiense* and *S. elongatus* is attributed to strong seasonality characterised by winter sea-surface temperatures probably less than 5°C (Marret et al., 2004).

Cysts of *P. dalei* occur sporadically in the Alboran Sea core (Fig. 2a) but mark clearly the cold events in core MD95-2042, most notably HS 2 and HS 4 (Fig. 2b). Cysts of *P. dalei* are well represented in modern sediments from polar to subpolar environments that experience summer sea-surface temperatures higher than 4°C (Rochon et al., 1999; Matthiessen, 1995; Marret et al., 2004; de Vernal et al., 2005). They are particularly prevalent as part of the spring bloom within North Atlantic fjord systems (Dale, 1977; Harland et al., 2004a,b).

In the Alboran Sea, we show that *S. lazus* is associated with cold HS and especially HS 5 (Fig. 2a). Today, the distribution of *S. lazus* is restricted to coastal regions of western Europe, always with low abundances (less than 2% of the dinocyst assemblages) (Reid, 1974). This species can be regarded as a neritic temperate species of regions characterised by oligotrophic to eutrophic surface water conditions (Reid, 1974; Harland, 1983).

Finally, the most striking feature visible in the dinocyst distribution during the D-O cycles is the systematic occurrence of *Impagidinium aculeatum* during GS in the Alboran Sea, with low percentages reaching a maximum of 5% (Fig. 2a). On the western Iberian margin, this species peaks during GI, although a local maximum is also observed during HS 5 (Fig. 2b). At present, this taxon is associated with warm water dinocyst assemblages; high relative abundances are found in tropical/subtropical oligotrophic open oceanic sites (Turon, 1984). In SW Iberian and western Mediterranean paleoclimate records, this species is characteristic of the Holocene with
percentages close to 20% (Combourieu-Nebout et al., 1998; Turon et al., 2003; Rouis-Zargouni et al., 2010). However, percentages close to 5% were also previously observed during cold stadials (HS 1 and HS 2) in a core from the Sicilian-Tunisian Strait (Rouis-Zargouni et al., 2010).

### 4.2. Dinocysts occurring during Greenland Interstadials

Dinocysts which feature prominently during GI in the Alboran Sea include *Spiniferites mirabilis*, *Impagidinium patulum*, *L. machaerophorum*, *N. labyrinthus* as well as some heterotrophic species (Protoperidinioids, *Selenopemphix nephroides* and *Selenopemphix quanta*, but excluding *Brigantedinium* spp. whose ecology is rather complex) (Fig. 2a). On the SW Iberian margin they include *Impagidinium* species (*I. aculeatum* and *I. patulum*), *S. mirabilis*, *O. centrocarpum* and *S. nephroides* (Fig. 2b). Unlike the SW Iberian margin (Fig. 2b), maximal dinocyst concentrations occur during GI in the Alboran Sea (especially during GI 8 and GI 12; Fig. 2a).

In the Alboran Sea, GI are characterised by the species *S. mirabilis* (Fig. 2a) with percentages on average four times higher than on the SW Iberian margin (Fig. 2b). The highest relative abundances of *S. mirabilis* are recorded during GI 7 and GI 8. Today, *S. mirabilis* is mainly distributed between 35°N and 50°N in warm temperate to temperate environments of the North Atlantic with highest occurrences found off the coast of Portugal and in the Bay of Biscay (Rochon et al., 1999). This species extends as far south as 10°N and is generally absent from areas with summer SST below 12°C and salinity below 28.5, and thrives optimally when winter SSTs are between 10°C and 15°C and summer SSTs are above 15°C.

On the SW Iberian margin, GI are characterised by *I. patulum* (Fig. 2b) with abundances four
times higher than in the Alboran Sea (Fig. 2a). Most *Impagidinium* species occur today with
maximum frequencies in tropical to warm temperate waters between 20°N and 35°N and are
representative of full-oceanic conditions (Harland, 1983; Turon, 1984; Bouimetarhan et al.,
2009).

*Spiniferites delicatus* is common on the SW Iberian margin and is mainly linked with interstadial
conditions (Fig. 2b), while it is rare in the Alboran Sea (Fig. 2a). *S. delicatus* is a temperate
species adapted to warm and neritic conditions (Wall et al., 1977; Harland, 1983; Marret, 1994).

*S. quanta*, *S. nephroides* and some Protoperidinioids show a distribution pattern closely linked to
GI in the Alboran Sea (Fig. 2a). This association also holds for *S. nephroides* on the SW Iberian
margin, but is less obvious for *S. quanta* and Protoperidinioids as they also increase during GS
(Fig. 2b). These latter taxa, represented by low relative abundances in the modern database and a
scattered distribution in the North Atlantic (Rochon et al., 1999), are derived from heterotrophic
dinoflagellates and are mainly related to high food resources. Their heterotrophic strategy of
nutrition probably links them to the presence of higher concentrations of nutrients in surface
waters. *S. quanta* has previously been linked to the dynamics of upwelling cells off NW Africa
(Dodge and Harland, 1991; Penaud et al., 2010).

### 4.3. Opposite dinocyst patterns between Iberian and Alboran environments over D-O cycles

Other dinocyst species not described above, *L. machaerophorum*, *N. labyrinthus*, and *O.
 centrocarpum*, are an important component of the dinocyst assemblages and show opposite
patterns in both cores
*N. labyrinthus* shows very high percentages in the Alboran Sea (average 24%), where it occurs generally within GI (Fig. 2a). Peaks of this species systematically exceed 40% and can reach 60 to 70% of the total dinocyst assemblage. On the SW Iberian margin, this taxon is less prevalent (average of 4.5%) with peaks generally reaching 10% during GS (except during HS) (Fig. 2b). At present, *N. labyrinthus* is a typical open-ocean species found predominantly between 45°N and 65°N in the North Atlantic Ocean (Rochon et al., 1999; Matthiessen, 1995; Marret et al., 2004; de Vernal et al., 2005). Maximum abundances of this species are recorded off southern Greenland where cold waters from the East Greenland and Labrador currents mix with warm North Atlantic waters of the Irminger Current (Rochon et al., 1999; Marret et al., 2004). This species, in association with *B. tepikiense*, was previously related to polar water incursions during MIS 5 cold substages off Portugal (Sánchez-Goñi et al., 1999; Eynaud et al., 2000), and during the Younger Dryas off Portugal (Turon et al., 2003), in the Mediterranean (Turon and Londeix, 1988; Rouis-Zargouni et al., 2010) and off NW Morocco (Marret and Turon, 1994; Penaud et al., 2010). *N. labyrinthus* has also been positively correlated with nutrient-rich and cool waters (Turon and Londeix, 1988; Devillers and de Vernal, 2000).

*L. machaerophorum* occurs during GI in the Alboran Sea, particularly GI 8, 9 and 12 (Fig. 2a), while it occurs during GS on the SW Iberian margin (Fig. 2b). *L. machaerophorum* is today considered to be a temperate to tropical coastal euryhaline species (Mertens et al., 2009). It dominates associations from the northern African and southern European Atlantic shelves, and it is also found in abundance in North African coastal upwelling regions (Targarona et al., 1999; Sprangers et al., 2004) and near the Congo outlet (Marret, 1994). Extremely high concentrations of these cysts have been found in areas with typical seasonally stratified water columns such as
fjords, bays, and estuaries (e.g. Reid, 1972; Bradford and Wall, 1984; Dale, 1985; Lewis, 1988; Morzadec-Kerfourn, 1988; Dale et al., 1999). This taxon has often been related to warm and stratified surface waters (Marret and Zonneveld, 2003), and has also been used as a proxy for fluvial inputs towards the ocean (Zaragosi et al., 2001; Holzwarth et al., 2010).

*O. centrocarpum* occurs frequently in the Alboran Sea record with percentages reaching 20 to 40% during almost every GS (Fig. 2a). On the SW Iberian margin, this species shows relatively low percentages, with peaks generally less than 10% during GI and reaching a maximum of 20% during GI 3 (Fig. 2b). This species is considered to be a cosmopolitan, cool to temperate taxon (Turon, 1984; Rochon et al., 1999; Marret and Zonneveld, 2003), and an abundance pattern following the route of the North Atlantic Drift (NAD) has been identified from its present geographical distribution in North Atlantic surface sediments (Turon, 1984; Rochon et al., 1999). This observation was previously used to interpret the presence of an active NAD at times when this species was abundant in the Quaternary North Atlantic sediments (Zaragosi et al., 2001; Eynaud et al., 2004; Penaud et al., 2008, 2009).

5. Temperature and salinity records: convergences and discrepancies

5.1. SST: dinocysts versus planktonic foraminifera

Quantitative reconstructions derived from the two transfer functions (dinocyst and foraminifera) are not directly comparable since the reconstructed parameters are not exactly the same. The dinocyst transfer function 3PBase-940 only provides February and August SST reconstructions while the foraminiferal transfer function R-1007 only provides annual and seasonal-mean (i.e.
winter, spring, summer and fall) SST reconstructions. However, February and August
foraminiferal SST can be estimated using the MAT transfer function derived from Pflaumann et
al. (1996) which relies on a modern database of 692 modern assemblages (only Atlantic stations),
improved during the MARGO project (Kucera et al., 2005). It is, however, more appropriate for
us to discuss the foraminiferal data from the R-1007 transfer function (Atlantic and
Mediterranean modern databases) for the Alboran Sea core which is located at the boundary of
the two basins. The comparison of winter versus February SST values on the 664 common
stations from the foraminiferal north Atlantic databases (n=1007 versus n=692) generates a mean
difference of 0.26°C (with a maximum difference of 1.95°C). For the summer versus August SST
values, the mean difference is 0.37°C (with a maximum difference of 1.1°C). These values are
within the range of the error bars of the foraminiferal reconstructions and therefore encourage us
to consider that monthly and seasonal foraminiferal SST values can be discussed interchangeably.
Concerning dinocysts, the n=940 database also includes Mediterranean stations. It is worth noting
that, among the 84 Mediterranean analogues (including 17 stations in the Alboran Sea), the only
Mediterranean analogue found by the transfer function is located in the Alboran Sea and named
“M1039” (Fig. 1; Appendix B). This is mainly the case during GI conditions (Gls 3, 5, 7, 8, 9,
11, 12 and 13; cf. Appendix B), suggesting that GI conditions in the Alboran Sea between 25 and
50 ka were roughly equivalent to conditions prevailing at the “M1039” site today (Fig. 1).
When comparing SST reconstructions from both micropaleontological proxies on the two cores,
one can note that February versus winter SST values obtained with dinocysts (3PBase-940) and
foraminifera (R-1007), respectively, are closer than August versus summer ones (Fig. 3).
However, the general good consistency of February versus winter SST is not valid for the SW
Iberian margin during HS 5 with about 8°C difference between both micropaleontological
reconstructions (dinocyst SST estimates around 13°C and foraminiferal ones close to 5°C; Fig. 3). This offset is unrealistically large given the prediction error for each method (around 1.2°C).

This minor SST drop expressed in dinocyst populations may be underestimated, as also revealed by the annual alkenone-derived SST values warmer than the February dinocyst-based SST ones (Fig. 3). One reason may come from the occurrence of the dinocyst species *Impagidinium aculeatum* during HS 5 (Fig. 2). Indeed, on the western Iberian margin, this species peaks during GI, although a local maximum is observed during HS 5 (percentages slightly higher than 5%). However, this species is associated with warm water dinocyst assemblages and is found today in tropical/subtropical oligotrophic open oceanic sites (Turon, 1984). Although the occurrence of this taxon during a cold HS is not understood, it may switch the transfer function towards warmer SST values.

August versus summer SST reconstructions for both cores show similar overall trends but differ more in terms of amplitudes, with dinocyst SSTs being 5 to 10°C higher than foraminiferal SST estimates (Fig. 3). Two hypotheses can be put forward with respect to this observation. Either the August/summer (dinocyst or/and foraminiferal) quantifications are less reliable or the discrepancy reflects an ecological bias. Dinocysts are indeed produced by dinoflagellates that thrive in the photic zone while foraminifera can migrate deeply in the water column with living depths ranging from 0 to 1000 m. Dinoflagellates, being found in shallower water, would thus record warmer SST consistently with a higher stratification during the warmest month, i.e. August. Various biases between different micropaleontological reconstructions can thus occur in relation to the ecological strategy (depth of habitat and growth seasons) of the different planktonic populations (de Vernal et al., 2005). The following comparison with alkenones will help us to decipher whether the foraminiferal or dinocyst signal is too cold or warm, respectively.
Finally, concerning the Alboran Sea core, one can note that dinocysts and foraminiferal show closer August versus summer SST values between 31 and 38 ka and larger offsets between 38 and 50 ka. This point will be further discussed in section 6.2.3. of this manuscript.

5.2. SST: transfer functions versus alkenones

Alkenone-derived SST from cores MD95-2043 (Cacho et al., 1999) and MD95-2042 (Pailler and Bard, 2002) has previously been compared with ice-core records, demonstrating a close linking between SSTs in the western Mediterranean and temperature developments over the wider North Atlantic region, including Greenland. Here, it appears that our dinocyst-based SST reconstructions for both cores match peak to peak with the SSTs derived from alkenones, showing minimum values during GS (Fig. 3). Furthermore, the alkenone-based SST, which records an annual signal of temperature, fluctuates systematically in between the temperature range given by seasonal dinocyst SST reconstructions and is closer to February than August SST (Fig. 3). A co-variation between February dinocyst SST and alkenone-derived SST reconstructions was previously observed off NW Morocco during the last glacial over the last 30 ka (Penaud et al., 2010). This would confirm the hypothesis that switches in mean annual temperatures were dominated by, and thus weighted towards, the winter season during the last glacial cycle as suggested by Denton et al. (2005).

The foraminiferal transfer function n=1007 also provides annual SST reconstructions that can directly be compared with the alkenone signal. One can note that annual SST values reconstructed with both proxies are closer during GI than during GS (Fig. 3). Foraminifera routinely provide much colder temperatures during GS, and especially during HS that are
characterised by significant SST anomalies between 4 and 7°C (Fig. 3). This shift of annual SST values towards cold SSTs is mainly due to extremely cold summer SSTs reconstructed with foraminifera during GS (Fig. 3). This contrast has parallels to the previous observation on the considerable offsets between dinocyst and foraminiferal August versus summer SST. Since alkenones are synthesised by coccolithophorids which are single-celled algae, protists and phytoplankton such as dinoflagellates, we can assume that they both give a signature of sea-surface hydrological changes occurring in the photic zone, while foraminifera may yield a bias towards colder SSTs, especially during GS. However, it will be crucial to understand if this finding reflects a real ecological strategy of foraminifera (different depths of habitat following different seasons) or a problem associated with the transfer function.

5.3. SSS: dinocysts versus planktonic foraminifera

Quantifications of salinity are of critical interest as they are fairly rare in paleoclimate studies. A major challenge in paleoceanography is to increase the availability of SSS reconstructions for comparison with climate models (e.g. MARGO project members, 2009). In our records, similar changes, both in amplitude and timing, are revealed through the comparison of February dinocyst-based SSS versus winter foraminiferal derived ones (Fig. 3). Numerical results are always very close or at least within the prediction error of 1-1.8 psu, except during HS 5 and GI 12 at the SW Iberian margin which are marked by differences of 2 to 3 psu between both micropaleontological proxies (Fig. 3). Comparison of the February/winter SSS data is particularly striking because the methods of quantification are different. One method requires MAT calculation of SST based on planktonic foraminifera and δ¹⁸O analysis (e.g. Malaizé and
Caley, 2009), while the other one is obtained on the basis of MAT transfer function applied to dinocyst assemblages (e.g. de Vernal et al., 2005). However, correlations between fluctuations in August dinocyst-based SSS and summer foraminiferal derived ones are less evident with fluctuations similar in timing but divergent in terms of amplitudes, especially for the Alboran Sea (Fig. 3).

For the SW Iberian margin, our study shows that major low salinity events are recorded during HS (Fig. 3b). HS 4 displays the maximum drop in February/winter SSS with values reaching 31 psu, corresponding to the largest fluxes of freshwater to the ocean over our study period. Our data for HS 3 (around 31.5-32.5 psu, 4 psu lower than the modern value) also show a major low salinity event and are consistent with previous results acquired further north along the Celtic margin over the last 35 ka BP where this interval was also characterised by a 4 psu depletion in SSS (Eynaud et al., submitted).

For the Alboran Sea, the difference in the resolution of analysis between dinocyst and foraminiferal/isotope reconstructions makes it difficult to compare some sections (Fig. 3a). This is mainly due to the fact that foraminiferal SSS is calculated from both foraminiferal SST and planktonic δ¹⁸O data and the resolution of the δ¹⁸O record is lower than the foraminiferal SST record. For the whole Alboran Sea record, February/winter SSS shows changes of similar order of magnitude between each GS, with the strongest signal of freshening recorded during HS 4 reaching 33 psu (4 psu lower than the modern value).

6. Paleohydrological changes affecting subtropical Atlantic latitudes through D-O cycles
6.1. The new contribution of dinocyst assemblages in the Alboran Sea

6.1.1. Warm intervals (GI) in the Alboran Sea

During GI, we observe an expansion of warm temperate to tropical species with a trend of decreasing abundance, and decreasing February and alkenone SST, from immediately prior to a Heinrich Stadial to the next HS (Fig. 4). This pattern is similar to that seen in the Greenland ice core isotope records showing progressively shorter GI and smaller increases in Greenland air temperatures between HS 5 and HS 4 and between HS 4 and HS 3 (Fig. 4, NGRIP GICC05). This demonstrates a similar trend between Alboran SST and atmospheric temperatures over Greenland, involving a rapid transmission of Northern Atlantic climate changes into the western Mediterranean region.

When comparing the amplitude of warm taxa development during individual D-O warming (Fig. 4), we note less frequent occurrences of thermophilous taxa during the earlier interstadials (GI 12-9) than during the subsequent interstadials (GI 8-5). This is consistent with pollen analysis conducted on the same core (Fletcher and Sánchez-Goñi, 2008) demonstrating high values for Mediterranean forest during GI 8, 7, 6 and 5 reflecting the maximum in subtropical summer insolation associated with the precession minimum centred around 30-35 ka (Fig. 4). Such a development is also seen in paleo-vegetation records at nearby Alboran site ODP 976 (Combourieu-Nebout et al., 2002) and on the SW Iberian margin (MD95-2042: Sánchez-Goñi et al., 2000).

The interstadials GI 8 and GI 12, immediately succeeding HS 4 and HS 5, respectively, exhibit particularly long and warm periods. They are both marked by high relative abundances of warm water dinocysts, high SST estimates (especially clear with alkenones; Fig. 4), and by the highest
paleoproduction conditions indicated by heterotrophic taxa in the early interstadials and high
dinocyst concentrations in the second half of GI 8 and GI 12 (Fig. 4). In terms of sea-surface
paleohydrology, GI 8 and GI 12 are thus very similar with a comparable bipartite structure (Fig.
4). Palynological investigations of GI 8 and GI 12 in the same Alboran Sea core showed that
these periods were characterised by the strongest expansions of mixed oak forest between 48 and
15 ka (Fletcher and Sánchez-Goñi, 2008). Furthermore, Fletcher and Sánchez-Goñi (2008)
identified an Atlantic oceanic character during GI 12, in contrast to a markedly Mediterranean
character during GI 8, linked to the precession minimum that likely enhanced the Mediterranean
climate and caused an enhanced seasonal contrast between dry summers and wet winters
(Sánchez-Goñi et al., 2008, 2009; Fletcher and Sánchez-Goñi, 2008). Our dinocyst data also
reveal a strong expansion of the species *L. machaerophorum* in the second half of GI 8,
representing between 20 and 40% of the total dinocyst assemblage (Fig. 4). This species has
sometimes been used to trace fluvial inputs (Zaragozi et al., 2001; Holzwarth et al., 2010) and
could reflect higher river run-off to the Alboran Sea, providing further evidence for increased
winter precipitations during this interval.
Throughout our record, heterotrophic dinocyst species (*S. nephroides*, *S. quanta*, and
*Protoperidinioids cysts*) occur during each GI in the Alboran Sea (Fig. 4). Their occurrences
suggest sea-surface conditions characterised by increased productivity (Rochon et al., 1999), as
also indicated by increased total dinocyst concentrations (Fig. 4). This general pattern of higher
productivity during GI relative to GS has previously been discussed based on geochemical
evidence (calcium carbonate, barium excess, and total organic carbon) from the same core
(Moreno et al., 2004). Today, the two semi-permanent anticyclonic gyres found in the Alboran
Sea represent energetic mesoscale features and the main forcing maintaining these gyres is the
Atlantic jet which enters through the Strait of Gibraltar (Bormans and Garret, 1989; Benzohra
and Millot, 1995; Garcia-Lafuente et al., 1998; Macias et al., 2008). The intensity of the jet is typically modulated by atmospheric pressure variations over the western Mediterranean. Indeed, when atmospheric pressures are lower than average, configuration close to a NAO negative mode, westerlies prevail above the Mediterranean, the Atlantic jet flows northward near the Spanish coast and the western Alboran gyre is well developed in the entire western Alboran Sea. The opposite (NAO positive mode) occurs when easterly winds prevail, the inflow of Atlantic waters is lower and the Atlantic jet is directed southward, reducing the western Alboran gyre extent (Candela et al., 1989; Garcia-Lafuente et al., 2002; Macias et al., 2008). At an annual scale, on the basis of satellite imagery analysis (e.g. Garcia-Gorriz and Carr, 1999; Baldacci et al., 2001; Macias et al., 2007, 2008) maximum surface chlorophyll concentrations were usually found in winter and minimum values were observed in summer (July and August). Biological patterns are thus also tightly coupled to atmospheric pressure above the Mediterranean Basin with westerlies being shifted southward during winter. At the millennial-scale resolution of our study and in agreement with Moreno et al. (2004), our data suggest that GI conditions would exhibit a prolonged southward shift of the westerly wind belt, inducing a more intense Atlantic surface jet that favoured gyre-induced upwelling in the Alboran Sea.

6.1.2. Cold intervals (GS) recorded in the Alboran Sea

The association *B. tepikense* - *S. elongatus* represents an indicator for the incursion of subpolar water masses at the Alboran site that is highly evident during HS (Fig. 4). This latter pattern is similar to that of the subpolar foraminifera *Neogloboquadrina pachyderma* s. recorded in the same core (Cacho et al., 1999) that shows higher percentages during HS (Fig. 4). This foraminifer displays the highest percentages during HS 4, then HS 3 and finally HS 5, while *B.*
tepikiense shows higher percentages during HS 3 compared to HS 4 (cf. Figs. 2 and 4). B. tepikiense is absent from the Mediterranean Sea today, and highest abundances of this species occur in areas characterised by high-amplitude (10°C) seasonal temperature shifts (Rochon et al., 1999). Therefore, its occurrence during HS in Alboran Sea surface waters implies enhanced seasonal temperature contrast (15°C; cf. Fig. 5) compared to the present-day one (i.e. 10°C; 23.5°C in summer and 14.5°C in winter; cf. Fig. 5), caused by a strong decrease of winter SST (Fig. 4). Our February dinocyst SST reconstructions reinforce this hypothesis by showing low SST with values around 5.5°C and 6.5°C during HS 4 and HS 3, respectively, i.e. 9°C less than at present (Fig. 4). It is thus important to note that, in contrast to N. pachyderma s., B. tepikiense does not mark systematically colder intervals but intervals with larger seasonal contrasts. Our results are also in agreement with other observations made at ODP Site 976 (Alboran Sea) that reflect coeval increases of B. tepikiense with N. pachyderma s. percentages during HS, over the last 50 ka BP (Turon and Londeix, 1988; Combourieu-Nebout et al., 2002). Our data thus confirm that sea-surface cooling in the Alboran Sea was mainly linked to the advection of cold Atlantic water to the western Mediterranean (Cacho et al., 1999). Such cold-water advection occurred synchronously with regional cooling related to atmospheric conditions over the western Mediterranean (Combourieu-Nebout et al., 2002; Sánchez-Goñi et al., 2002). Indeed, on the adjacent continent, the Mediterranean forest (i.e. temperate taxa) collapsed (Fig. 4) and estimated winter atmospheric conditions indicate a 10°C lowering and a decrease of 400 mm in precipitation (Sánchez-Goñi et al., 2002).

The HS configuration contrasts with the other GS. In the Alboran Sea, before or during each GS (especially those not associated with HS conditions) we note pronounced peaks of O. centrocarpum (Fig. 4). The occurrence of O. centrocarpum in the Alboran Sea, whose present-day distribution directly mirrors the flow path of the NAD, may result from the influx of cool
North Atlantic waters entering into the Mediterranean; temperatures of these waters, while presumably low, remained above those of the subpolar waters that entered the Mediterranean during HS. GS events of the last glacial have been demonstrated to coincide with intensification of the deep circulation in the Mediterranean (Cacho et al., 2000, 2006; Sierro et al., 2005; Frigola et al., 2008). Our data suggest that the intensification of deep ventilation in the Alboran Sea, reflected in benthic $\delta^{13}C$ data (Fig. 4; Cacho et al., 2006), was synchronous with the advection of North Atlantic waters to the Mediterranean, as reflected by highest relative abundances of cool-water dinocyst taxa (Fig. 4).

We demonstrate two different patterns during GS: those associated with HS are marked by increased abundances of cold water species associated with subpolar waters, and those not associated with HS are marked by an expansion of cool North Atlantic species. Interestingly, it has been demonstrated that the densest Western Mediterranean Deep Water was formed during GS not associated with HS (Cacho et al., 2006; Frigola et al., 2008). More precisely, it has been suggested that a strong mode of overturning prevailed during GS not associated with HS, an intermediate mode of overturning during HS and a weak mode of overturning during GI (Sierro et al., 2005; Frigola et al., 2008). Strong overturning was expected during HS since strong and cold northern continental winds prevailed over the Mediterranean resulting in dry-cold conditions on land (Fig. 4; Combourieu-Nebout et al., 2002; Sánchez-Goñi et al., 2002). However, the HS intermediate mode has been linked with a strong influence of subpolar waters that lowered sea-surface salinity thus reducing deep water formation and favouring water column stratification (Sierro et al., 2005). The observation of subpolar species (*N. pachyderma* s. and *B. tepikiense*) during HS (especially HS 3 and HS 4), as well as the low winter and summer salinities reconstructed from dinocysts, lowered by around 0.5 to 1.5 psu compared to the other GS in both
seasonal configurations (Fig. 3), within low precipitation phases, reinforce the idea of sustained cold conditions with subpolar water masses advection towards the Mediterranean Sea and decreased deep water formation.

6.2. Alboran Sea versus SW Iberian margin between 50 and 25 ka

6.2.1. SST reconstructions and paleoenvironmental signatures

In both cores, lowest SST estimates are observed during HS 4 which represents the most pronounced cold event between 25 and 50 ka, with February/winter SST of about 5.5°C and 4°C recorded in the Alboran Sea and the SW Iberian margin, respectively (Fig. 3). Previously, the coldest deep water temperatures in the Alboran Sea (Cacho et al., 2006), maximum concentrations of IRD off the Portuguese margin (Thouveny et al., 2000), and a significant increase in the transport of Saharan dust (Moreno et al., 2002) were observed to occur during HS 4. This confirms the magnitude of HS 4 (cf. Eynaud et al., 2009) in terms of expansion of polar waters towards the Iberian margin (i.e. February SST 11°C lower than modern ones) and even towards the western Mediterranean (i.e. February SST 9°C lower than modern ones), in phase with a nearly complete shutdown of the thermohaline circulation (Maslin et al., 1995; Elliot et al., 2002; Roche et al., 2004).

HS 3 is marked by a change in surface hydrological conditions with February/winter SST of about 6.5°C in the Alboran Sea and 5.5°C off Portugal (Fig. 3). Winter conditions seem thus to have been less severe during HS 3 than HS 4, although they are nevertheless characterised in both paleo-records by significant cooling.
Finally, HS 5 is the event which displays the weakest changes both in the Alboran Sea and off Portugal, where dinocyst February SST are about 9°C and 13°C, respectively (Fig. 4). However, foraminiferal winter SST estimates are close to 6°C and 5°C in the Alboran Sea and off Portugal, respectively. Quantifications derived from transfer functions appear less reliable during this interval. The alkenone annual signal (around 10.5-11°C in the Alboran Sea and 12-12.5°C off Portugal) probably provide an intermediate signature between dinocyst-based (too warm) and foraminiferal-based (too cold) February/winter SST estimates.

Micropaleontological evidence in the subpolar North Atlantic has shown that abrupt SST changes associated with the D-O events in Greenland were matched by SST variations of at least 3-5°C (Bond et al., 1992, 1993; Elliot et al., 2002). In the subtropical North Atlantic, SST excursions of 4-5°C across stadial-interstadial transitions of the last glacial were recorded from the Bermuda Rise (alkenones: Sachs and Lehman, 1999; isotopes: Keigwin and Boyle, 1999) and off Portugal (alkenones: Bard et al., 2000; Martrat et al., 2007). SST changes of up to 6°C are also documented in the western Mediterranean (alkenones: Martrat et al., 2004), due to southward shifts in the position of the Polar Front. Consistently with these previous results, we show here that the Alboran Sea and the SW Iberian margin experienced fluctuations of dinocyst SST of around 5-6°C between GI and GS, except for HS events and more specifically HS 4 with a SST drop of around 11°C off Portugal and of 8°C in the Alboran Sea. This demonstrates the extreme sensitivity of dinocysts to climate fluctuations in subtropical latitudes.

### 6.2.2. Multiproxy evidence for paleohydrological changes through time

During GI, the expansion of the Mediterranean forest (Fig. 4) was attributed to atmospheric
conditions close to present-day ones with warm summer and wet winter conditions over south-western Europe (e.g. Sánchez-Goñi et al., 2002; Combourieu-Nebout et al., 2002; Bout-Roumazeilles et al., 2007; Daniau et al., 2007). During those times, we show that warm temperate species *S. mirabilis* and *I. patulum* expanded in the Alboran Sea and on the SW Iberian margin, respectively (Fig. 4), and high SST is recorded at both sites during GI revealing sea-surface conditions closer to present-day ones (Figs. 3 and 4).

During GS, the decline of the Mediterranean forest (Fig. 4) and the development of steppe and semi-desert vegetation over the south-western European borderlands were attributed to intensified winter dryness with cold continental conditions affecting the western Mediterranean area (e.g. Sánchez-Goñi et al., 2002; Combourieu-Nebout et al., 2002; Moreno et al., 2002, 2005; Roucoux et al., 2005; Bout-Roumazeilles et al., 2007; Daniau et al., 2007). At that time, assemblages of cold-water dinocyst species, including *B. tepikiense* and *S. elongatus* developed, reflecting cold-water advection along the SW Iberian margin and towards the Alboran Sea (Fig. 4). However, unlike the western Iberian margin where *B. tepikiense* occurs during each GS, *B. tepikiense* expands principally in the Alboran Sea during HS, and especially during HS 3 and HS 4. In the western Mediterranean Sea, HS impacts were therefore greater than those associated with the other GS. Broecker (2006) has compiled data from several areas and has identified sites where the impact associated with North Atlantic Heinrich events is larger (eastern Brazil, central Florida, Arabian Sea, Chinese stalagmites, and western Mediterranean area), in contrast to sites where impacts are similar to those observed during the other stadials (Greenland ice and Cariaco Basin). To explain this discrepancy, Broecker (2006) involves the greater magnitude of north-hemispheric sea ice expansion associated with North Atlantic Heinrich events which exceeded that associated with the other stadials.
B. tepikiense was previously used to trace subpolar water masses on the western Iberian margin (Eynaud et al., 2000; Turon et al., 2003) and in the Alboran Sea (Turon and Londeix, 1988; Combourieu-Nebout et al., 2002) during HS. In our data, the observation of subpolar dinocysts in both cores, in phase and synchronous with peaks of N. pachyderma s. (Fig. 4), confirms the idea of the shift of the Polar Front (PF) towards southern latitudes during North Atlantic Heinrich events. Indeed, Eynaud et al. (2009) proposed a conceptual scheme for the position of the PF on the western Iberian margin during Heinrich events and its influence on the local hydrology, and noticed that the protrusion of subpolar waters extended until approximately 40°N. Core MD95-2042 (37°48’N), at the southern limb of the Ruddiman belt (i.e. between 40 and 55°N; Ruddiman, 1977), was affected by subpolar waters and iceberg discharges. However, it is surprising to observe that the peaks of B. tepikiense are even larger during GS not associated with HS on the SW Iberian margin. When considering dinocyst, foraminiferal, and especially alkenone-based SST reconstructions, HS are characterised by the coldest temperatures (Figs. 3 and 4). We can therefore assume that temperatures were probably too cold during summer months to permit the expansion of B. tepikiense. This hypothesis is reinforced by the large excursions towards cold summer temperatures observed with dinocyst and foraminiferal SST reconstructions (Fig. 3). The huge advection of subpolar waters down to the SW Iberian margin during each GS, and especially HS, is furthermore reinforced by the dinocyst SSS reconstructions that show a generally more pronounced influence of meltwater in this sector compared with the westernmost part of the Mediterranean Sea (average salinity offset of around 1 psu between the sectors; Fig. 3). This is especially true for HS 4, characterised by salinities of around 31 on the SW Iberian margin and 33 in the Alboran Sea (Fig. 3). We thus show the pronounced impact of subpolar waters affecting the SW Iberian margin during each GS and especially during HS, while the Alboran Sea is only impacted by huge freshwater discharges during HS.
6.2.3. A first attempt to interpret the dinocyst seasonality signal

Sea surface temperatures and precipitation are environmental parameters directly linked to seasonality in the study region, and the difference between August and February SST estimates derived from dinocysts may contain important information regarding the seasonality (Fig. 5). Overall, the two cores show excursions in seasonality reconstructions towards a higher seasonal contrast during GS due to extremely cold February SST recorded during these cold intervals. At the SW Iberian margin, seasonal contrasts during GI appear similar to the present-day range, while extremely pronounced seasonal contrasts of roughly similar magnitude are recorded during each GS. The pattern of seasonality is very close to the relative abundance curve of *B. tepikiense* (Fig. 5). In the Alboran Sea, higher seasonal contrasts are mainly noted during HS and also correspond to higher percentages of *B. tepikiense*. It is not surprising to find parallels between this species and the seasonal signal since highest abundances of this species are found in areas today characterised by high-amplitude (10°C) seasonal temperature shifts (Rochon et al., 1999). In the Alboran Sea, it confirms the establishment of an enhanced seasonal temperature contrast compared to the present-day caused by a strong decrease in winter SST during HS (Combourieu-Nebout et al., 2002). At the SW Iberian margin, it reveals very clearly the marked shift between seasonal temperatures during each GS.

In the Alboran Sea, another climatic trend is superimposed on the general observations made above with, in broad terms, stronger seasonal contrasts during the early part of the record between 50 and 38 ka, compared to the period 38-31 ka (Fig. 5). Since seasonality and precession are closely linked, we show the precession curve (Berger and Loutre, 1991) alongside the reconstruction of seasonality (Fig. 5). It appears that lower seasonal contrasts are observed during
the precession minimum and *vice versa* (Fig. 5). We would have expected higher seasonal contrasts during the precession minimum since this orbital parameter enhanced the Mediterranean climate with warmer summer and wetter winters (Meijer and Tuenter, 2007). When looking at February SST reconstructions in the Alboran Sea, one can note that fluctuations during GS and GI are of similar magnitude (Fig. 5). The contrast is mainly due to August SST reconstructions that show colder values between HS 4 and HS 3 than between HS 5 and HS 4. Our data would suggest a link between the Alboran Sea paleohydrology and the precession signal through a climatic forcing acting on August temperatures. A first hypothesis could involve the general strengthening of the gyre-induced upwelling in the Alboran Sea during the time interval 38-31 ka favouring a cooling of sea-surface waters compared to the period 50-38 ka. However, the functioning of the gyres is mainly controlled today by winter conditions. Furthermore, no large differences are observed between 50-38 ka and 38-31 ka in terms of total dinocyst concentrations and heterotrophic dinocyst species, these latter proxies indicating paleoproductivity conditions probably connected to gyre-induced upwelling intensity. Greater stratification of the Alboran water column may also be suggested for the period 50-38 ka compared to the period 38-31 ka on the basis of the observation of dinocyst and foraminiferal August/summer SST (Fig. 3). Indeed, SSTs in August and summer are similar between 38 and 31 ka and are more distinct between 50 and 38 ka (Fig. 3). This might suggest a greater stratification of water masses during summers within the interval 50-38 ka and a stronger mixing of water masses during summers within the interval 38-31 ka. Warmer dinocyst SSTs might relate to sea-surface conditions while foraminiferal SSTs would incorporate SST signals at greater depths in the water column for the period 50-38 ka. Further investigation will be necessary to verify and explore on longer time-scales: a) the imprint of precession on the seasonality changes inferred from dinocysts, and b) the functioning of the mesoscale energetic features in the Alboran Sea represented by two
anticyclonic gyres today.
7. CONCLUSION

We have characterised glacial climate variability between 25 and 50 ka BP by comparing surface paleohydrology signals on both sides of the Strait of Gibraltar (Alboran Sea and SW Iberian margin). Comparison of dinocyst assemblages enables the reconstruction of hydrological features at both locations and we present, in this study, the first quantitative dinocyst reconstructions (SST and SSS) for MIS 3 obtained at mid-latitudes. Our hydrological quantifications acquired for both cores reproduce millennial-scale changes correlated to the D-O climatic variability, with a pattern of marked decrease in SST accompanied by a strong freshening of sea-surface waters evident during each GS on the SW Iberian margin and during HS in the Alboran Sea. Furthermore, we show similar patterns and amplitudes in SST reconstructions based on dinocysts, foraminifera and alkenones, and in SSS derived from dinocysts and foraminiferal SST coupled with planktonic δ¹⁸O. Larger discrepancies between dinocyst and foraminiferal estimates occur for summer reconstructions than for winter reconstructions, which are very similar in amplitude. We furthermore show the occurrence of cold taxa (including B. tepikiense and S. elongatus) during GS and the presence of thermophilous ones (S. mirabilis and Impagidinium spp.) during GI. However, the amplitude of variation in these taxa is not equivalent at both sites. B. tepikiense characterises each GS in the SW Iberian margin while it only develops during HS in the Alboran Sea. This pattern reflects, in the Alboran Sea, the maximum climatic deterioration during HS and the incursion of low-salinity subpolar waters to the western Mediterranean. During other GS (i.e. excluding HS), conditions were less severe in the Alboran Sea as is reflected by the occurrence of O. centrocarpum at very high relative abundances in the Alboran Sea and low values on the SW Iberian margin. This species conceivably reflects the inflow of cool North Atlantic waters to the
western Mediterranean, synchronously with stronger deep convection occurring at that time in the western Mediterranean. Finally, expansions of temperate to tropical taxa testify to the installation of warm sea-surface waters during GI. In the Alboran Sea, a high primary productivity pattern is deduced from heterotrophic species and high total dinocyst concentrations, suggesting gyre-induced upwelling due to prevailing southward-shifted westerlies above the Mediterranean at that time. The functioning of the gyre on multi-millennial timescales has also been discussed in light of seasonality reconstructions based on dinocysts. However, the relationships between gyre dynamics, seasonality and precession need to be explored in longer records covering several precession cycles. The dinocyst data thus exhibit regional trends and suggest distinct oscillations of sea-surface temperature and salinity, documenting the combined influence of atmospheric and hydrologic processes impacting on the western Mediterranean Sea and eastern subtropical latitudes of the North Atlantic during the abrupt climatic events of MIS 3.
8. Acknowledgements

Thanks to the French polar institute IPEV (*Institut Paul Emile Victor*), the captain and the crew of the Marion Dufresne and the scientific team of the 1995 IMAGES cruise. We wish to thank Mr. Y. Balut for his assistance at sea and M. Castera and O. Ther for invaluable technical assistance at the laboratory. We gratefully acknowledge the reviewers, whose comments have enabled us to greatly improve this manuscript. We thank W. Fletcher for improving the English language of the manuscript. This study was supported by the French CNRS and contributes to the EuroCLIMATE project RESOLuTION.

9. References


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10. Figure captions

**Figure 1:** Area of interest with major sea-surface features. The studied cores MD95-2043 (Alboran Sea; 36°8.6’N; 2°37.3’W; 1841 m water depth) and MD95-2042 (Iberian margin; 37°48’N; 10°10’W; 3146 m water depth) are located on the large map, depicting also the bathymetry of the study area and the major surface currents within the Alboran sea; WAG: Western Alboran Gyre; EAG: Eastern Alboran Gyre; AOF: Almeria-Oran Front; AC: Algerian Current. The small map on the left shows large scale North-Atlantic currents with: the North Atlantic Drift (NAD), the Portugal Current (PC) flowing southward from 45°N to 30°N, the Azores Current (AzC) derived from the southern branch of the Gulf Stream and flowing eastward to the Gulf of Cadiz at about 35°N, and the Canary Current (CC) fed by both the AzC and the PC. Together, these currents form the Eastern Boundary Current of the North Atlantic subtropical gyre. A Mediterranean analogue of the dinocyst modern database has been located on the map (M1039). Colors from red to dark blue on the map reflect growing bathymetry towards the deepest areas.

**Figure 2:** MD95-2043 (a) and MD95-2042 (b). Core depths are displayed in centimetres along the vertical axis. The relative abundances of selected dinocyst species are compared with planktonic $\delta^{18}$O data and $U^{\delta^{37}}$-SST (Core MD95-2043: Cacho et al., 1999; Core MD95-2042: Cayre et al., 1999; Shackleton et al., 2000; Pailler and Bard, 2002), providing the stratigraphical framework for the two cores. Quaternary dinocyst concentrations are also illustrated. HS: Heinrich Stadial; GI: Greenland Interstadial.
Figure 3: MD95-2043 (a) and MD95-2042 (b). Quantitative dinocyst reconstructions (February and August SST and SSS, 3PBase-940) compared with SST provided by foraminifera (winter, summer and annual, R-1007) and alkenones (U$_{37}^{k}$ - SST), and SSS provided by calculations based on planktonic $\delta^{18}$O (G. bulloides) and foraminiferal SST estimates (winter and summer, R-1007). Error bars are shown in the figure for the different reconstructions. HS: Heinrich Stadial; GI: Greenland Interstadial.

Figure 4: Comparison between cores MD95-2043 and MD95-2042 of a multi-proxy compilation including: selected dinocyst species, total dinocyst concentrations, February dinocyst (3PBase-940) and alkenone SST (Cacho et al., 1999; Pailler and Bard, 2002) reconstructions, percentages of N. pachyderma s. (Cacho et al., 1999; Pérez-Folgado et al., 2003), benthic $\delta^{13}$C (Cacho et al., 2006), and percentages of the Mediterranean forest (Sánchez-Goñi et al., 2000, 2002; Fletcher and Sánchez-Goñi, 2008). These data are compared with $\delta^{18}$O ice core records and the precession signal (Berger and Loutre, 1991). HS: Heinrich Stadial; GI: Greenland Interstadial.

Figure 5: MD95-2043 (a) and MD95-2042 (b). Estimation of a parameter of seasonality based on the difference between February and August dinocyst SST reconstructions (3PBase-940), compared with percentages of the dinocyst species Bitectatodinium tepikiense, the precession signal (Berger and Loutre, 1991) and planktonic $\delta^{18}$O records (G. bulloides). Dinocyst and alkenone SST records are also represented, as well as modern SST values. HS: Heinrich Stadial; GI: Greenland Interstadial.

11. Appendices
Appendix A: MD95-2042 (SW Iberian Margin) and MD95-2043 (Alboran Sea): individual counts of dinocyst species, total dinocysts counted and total dinocyst concentrations per sample.

Appendix B: MD95-2042 (SW Iberian Margin) and MD95-2043 (Alboran Sea): lists of the five best analogues found with the transfer function 3PBase-940. The geographical coordinates of each named analogue can be found on the GEOTOP website (http://www.geotop.ca/index.php?option=com_content&task=view&id=762&Itemid=226). The five analogues found by 3PBase-940 have systematically been used for the calculations, the threshold (Dmin value of 71.72) may then be considered to judge the good reliability of the reconstructed hydrological parameters. The M1039 analogue, highlighted in the tables in bold, is a Mediterranean analogue located in Figure 1.
Figure 1
a) MD95-2043 - Alboran Sea

Figure 2
### a) MD95-2043 - Alboran Sea

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**Figure 3**

### b) MD95-2042 - Iberian Margin

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**Figure 3**
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