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Abstract: Planktonic foraminiferal assemblages were studied at highresolution in core KCO1B from the Ionian Sea. Quantitative analysis allowed us to distinguish the main climatic features and associated paleoceanographic changes, that occurred between Marine Isotopic Stages (MIS) 13 and 9 (~ 500-300 ka).

MIS 12 and MIS 10 are characterized by relatively temperate conditions and an oligotrophic oceanographic regime in the early part and by colder conditions and nutrient supply in the sub-surface water masses in the upper part. During these intervals, small but distinct peaks of Neogloboquadrina pachyderma sinistral (sin) are detected at times of extremely negative values of the planktonic foraminifera paleoclimatic curve. Their co-occurrence with similar episodes in the Atlantic suggests that the climate in the Central Mediterranean was associated with north-Atlantic millennial-scale climate instability. MIS 11 and MIS 9 are dominated by surficial warm-water taxa. The climate optimum is reached in the middle part of each of these stages, as denoted by the presence of Globigerinoides sacculifer, and persists for approximately 20 and 6 ka during MIS 11 and MIS 9 respectively. This warming is not constant but is characterized by three distinct intervals with elevated winter temperatures and/or weak winter mixing.

Distribution of Globigerina bulloides, Turborotalita quinqueloba and Neogloboquadrina pachyderma dextral (dex) indicates that significant environmental changes occur across the transitions from glacial to interglacial MIS 12/MIS 11 (Termination V) and MIS 10/MIS 9 (Termination IV).

The studied record documents a close linkage between Mediterranean climate evolution and higher- and lower-latitude climate change throughout MIS 13-9.

Cover Letter, for Editor only

November, 2, 2015

Dear Editor,

I attach you the revised version of the manuscript entitled "*Central Mediterranean Mid-Pleistocene paleoclimatic variability and its association with global climate*" submitted by Lucilla Capotondi, Angela Girone, Fabrizio Lirer, Caterina Bergami, Marina Verducci, Mattia Vallefuoco, Angelica Afferri, Luciana Ferraro, Nicola Pelosi, Gert J De Lange. (REF PALAEO8460R1).

The "apparently minor mistakes" evidenced by the referee have been corrected. However I realized that the mistakes (many words linked without spaces between them) present in the pdf version are not present in the submitted Word version. The changes are in the file.pdf generated by the system. I think this is due to the fact that I have submitted the manuscript as a file "word.docx" and not "word.doc". I'm warning that probably the system does not convert well the version word.docx.

I add the point to point response to the reviewers, a version of the revised manuscript showing the new/changed text and a clean version of the revised manuscript.

Kind Regards

Lucilla Capotondi

Highlights

High-resolution Mid-Pleistocene climate reconstruction in the Central Mediterranean.Significant changes in temperature and productivity occurred during MISs 12 and 10.The climatic optimum during MIS 11c shows three warmest phasesT-V and T-IV are characterized by complex paleoceanographic changes

1 Central Mediterranean Mid-Pleistocene paleoclimatic variability and its association with

- 2 global climate
- 3
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- *Keywords:* Planktonic foraminifera, Middle Pleistocene, Central Mediterranean, Paleoceanographic
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- 19
- 20 Abstract

21 Planktonic foraminiferal assemblages were studied at high-resolution in core KC01B from the 22 Ionian Sea. Quantitative analysis allowed us to distinguish the main climatic features and 23 associated paleoceanographic changes, that occurred between Marine Isotopic Stages (MIS) 13 24 and 9 (~ 500-300 ka).

MIS 12 and MIS 10 are characterized by relatively temperate conditions and an oligotrophic oceanographic regime in the early part and by colder conditions and nutrient supply in the subsurface water masses in the upper part. During these intervals, small but distinct peaks of *Neogloboquadrina pachyderma* sinistral (sin) are detected at times of extremely negative values of the planktonic foraminifera paleoclimatic curve. Their co-occurrence with similar episodes in the

Atlantic suggests that the climate in the Central Mediterranean was associated with north-Atlantic millennial-scale climate instability. MIS 11 and MIS 9 are dominated by surficial warm-water taxa. The climate optimum is reached in the middle part of each of these stages, as denoted by the presence of *Globigerinoides sacculifer*, and persists for approximately 20 and 6 ka during MIS 11 and MIS 9 respectively. This warming is not constant but is characterized by three distinct intervals with elevated winter temperatures and/or weak winter mixing.

Distribution of *Globigerina bulloides*, *Turborotalita quinqueloba* and *Neogloboquadrina pachyderma* dextral (dex) indicates that significant environmental changes occur across the transitions from
 glacial to interglacial MIS 12/MIS 11 (Termination V) and MIS 10/MIS 9 (Termination IV).

The studied record documents a close linkage between Mediterranean climate evolution and
 higher- and lower-latitude climate change throughout MIS 13–9.

41

42 1. Introduction

43 To improve our understanding of natural climate variability and our abilities to forecast future 44 climate change, it is essential to investigate geological climate archives with relevant climate 45 change events. Accordingly, this paper focuses on climate variability that occurred before and after 46 the Mid-Brunhes event (MBE) (Jansen et al., 1986) in the Mediterranean Sea. The investigated 47 time interval (MIS 13-MIS 11; ~ 500-300 ka) is characterized by substantially warmer interglacials, (Epica Members, 2004; Jouzel et al., 2007) and enhanced atmospheric CO₂ content, at levels 48 49 similar to those for the pre-industrial Holocene (Siegenthaler et al., 2005). In particular, it includes 50 the MIS 11c, traditionally considered as potential analogue for future climate evolution because of 51 relatively similar orbital climate forcing (Loutre and Berger 2000; Masson-Delmotte et al., 2006; 52 Tzedakis et al., 2012). Therefore, a thorough study of this interval will provide information on type 53 and magnitude of climate variability under non-anthropogenic but otherwise comparable conditions 54 to the present.

In addition, the studied interval also includes the MIS 12, the most extreme glacial of the last 500
ka (Shackleton, 1987; Rohling et al., 1998; Lisiecki and Raymo, 2005) characterised by a sea level
of about 125 m lower than today (Rohling et al. 2009; 2014). The MIS 12-11 transition (Termination)

58 V) is also part of this peculiar interval. It represents a glacial-interglacial transition that is long 59 compared to later Pleistocene terminations (Oppo et al., 1998; Bauch et al., 2000; Thunell et al., 60 2002; Kandiano and Bauch, 2007; Helmke et al., 2008).

Notwithstanding the huge literature about the aforementioned climatic intervals, some features deserve additional clarifications. Two of the most intriguing aspects are the protracted deglaciation during Termination V and the cause of a long period of interglacial warmth during the MIS 11 (longer than any other mid-to late Pleistocene interglacial) with contrasting SST dynamics between polar- and mid-latitudes (Helmke and Bauch, 2003; Kandiano and Bauck, 2007; Kandiano et al., 2012 Milker et al. 2013 and references therein).

These issues could be addressed by analysing paleo-data from the Mediterranean Sea, a region highly sensitive to atmospheric and climatic system modifications due to its intermediate latitudinal position, where Euro-Asian and North-African climate regimes strongly interact (Roether et al., 1996; Béthoux et al., 1999; Pinardi and Masetti 2000; Trigo et al., 2004; Lionello et al., 2006).

71 Moreover the Mediterranean climate is exposed to the South-Asian Monsoon in summer and the 72 Siberian High-Pressure-System in winter (Luterbacher, 2005; Lionello et al., 2006). Today, the 73 southern part of the Mediterranean region is mostly influenced by the descending branch of the 74 Hadley cell, while the northern part is more linked to mid-latitude variability, characterized by the 75 North-Atlantic Oscillation (NAO) and other mid-latitude teleconnection patterns (Hurrel et al., 2004). 76 Thus, climate investigations of geological archives of the Mediterranean region reflect paleo-77 changes in the intensity and extension of global-scale climate patterns. In addition, the 78 Mediterranean sedimentary sequences are characterized by the (quasi-) periodical occurrence of 79 episodes of deep-sea oxygen depletion (sapropel layers) (Olausson, 1991; Rohling et al., 2015 80 and references therein). Based on their link with the astronomical parameters (Hilgen 1991; 81 Rossignol-Strick, 1983; Lourens 2004; Konijnendijk et al., 2014) these discrete levels represent a 82 useful constraint to establish accurate age models for marine and land sections.

We present a new high-resolution quantitative study of planktonic foraminifera distribution
throughout MIS 13-9 for sediment core KC01B collected in the Ionian Sea, central Mediterranean
Sea. Planktonic foraminifera are amongst the most commonly used proxies for paleoceanographic

and paleoclimate sea-surface reconstructions. Their distribution and abundance is strongly linked to surface-water properties. In addition, the physical and chemical properties of their shells reflect past environmental conditions of the water masses in which they lived (Kucera 2007 and references therein).

The investigated deep-marine sequence of core KC01B represents a key site for stratigraphic and paleoclimatic investigations. This is not only because of its strategic location but also because it was used for the construction of a sapropel-based astronomical timescale for the last 1.1 My (Lourens, 2004). Moreover, in this work, we update the studied time interval by using the recent chronological constraints on Pleistocene sapropel deposition (Konijnendijk et al. 2014) and a new oxygen isotopic record of *N. pachyderma*.

96 Our main aim is to explore how the ecosystem responded to climate variability during glacial and 97 interglacial intervals throughout MIS 13–9 in order to discuss the possible mechanisms through 98 which climate acts at the regional and global scale.

In detail, we focus on the main environmental and paleoceanographic processes occurring a) at
 times of glacial and interglacial MIS; b) during Termination V (T-V) and Termination IV (T-IV).

101

102 2. Modern Oceanographic Setting

103 Currently, the Mediterranean Sea is an evaporative basin where freshwater loss exceeds 104 freshwater input, forcing an anti-estuarine circulation (Borghini et al., 2014).

At the surface (the first 100-200 m), moderate-salinity Atlantic Water (AW) intrudes through the Strait of Gibraltar and flows to the easternmost part of the Levantine basin modifying its temperature and salinity properties (Modified Atlantic Water – MAW). In the intermediate layer (depth between 150–200 and 600 m), Levantine Intermediate Water (LIW) forms in the eastern basin, spreads westwards and continues its flow towards the Strait of Gibraltar, and then into the Atlantic Ocean (Béthoux et al., 1992; Robinson et al., 1991; Manca et al., 2004; Malanotte-Rizzoli et al., 2014).

Atmospheric forcing and basin topography determine a large number of local cyclonic and anticyclonic cells (Pinardi and Masetti, 2000). In wintertime, outbreaks of cold and dry continental

air masses lead to significant negative heat budgets and buoyancy losses, initiating deep and/or
intermediate dense water formation both in the western and in the eastern basins (MalanotteRizzoli and Bergamasco, 1991; Castellari et al., 2000).

117 The Ionian Sea is influenced by the transit and on-site transformation of the major water masses 118 previously described (e.g., Modified Atlantic Water, MAW; Levantine Intermediate water, LIW; and 119 Eastern Mediterranean Deep Water, EMDW, Malanotte-Rizzoli et al., 1997; Napolitano et al., 120 2000) (Fig. 1). At the near-surface level, most important for biological production, the MAW enters 121 in the western Ionian basin advected by the Atlantic Ionian Stream (AIS) (Fig. 1). Recently, the 122 upper-layer circulation in the Ionian Sea has been associated with the deep thermohaline 123 circulation through the Bimodal Oscillating System (BiOS): the Ionian upper-layer circulation 124 reverses from cyclonic to anticyclonic and vice versa on decadal time scale affecting the biological 125 productivity in the northern Ionian and southern Adriatic Sea (Civitarese et al., 2010; Gačić et al., 126 2010).

The present-day Mediterranean Sea is characterized by oligotrophic conditions (Béthoux, 1979; Sarmiento et al., 1988). The main factor that controls the seasonal change in primary production is linked to the dynamics of the water column with increasing biomass in late winter/early spring and decreasing in summer (Antoine et al., 1995; Bosc et al., 2004; D'Ortenzio and Ribera d'Alcalà, 2009).

A significant West-East trophic gradient exists with nutrient depletion (mainly phosphorus) and a
 reduction in primary productivity in the eastern basin (Krom et al., 1991, 2010).

Moreover, primary productivity reflects the hydrological fragmentation due to mesoscale variability
(D'Ortenzio and Ribera d'Alcalà, 2009).

An oligotrophic regime, characterized by a low production, occurs in summer, when a stable stratification takes place (Klein and Coste, 1984; Krom et al., 1992; Crispi et al., 1999; Allen et al., 2002). During this period, low standing stocks characterize surface waters with the dominance of predatory species. The Ionian planktonic foraminiferal association is generally dominated by *Globigerinoides ruber* pink (40-60%) and *G. ruber* alba (20-40%) with peaks of maximum abundance in the first 50 m of the water column (Pujol and Vergnaud Grazzini, 1995). Winter

142 convection, and less frequently frontal zone migration or upwelling, brings nutrients into the photic 143 zone (mesotrophic regime) (Klein and Coste, 1984). During winter, the assemblage is 144 characterized by the dominance of grazing species such as *Globorotalia truncatulinoides* (50%) 145 and by the presence of other non-spinose species such as Globorotalia inflata (20%), and 146 Globigerina bulloides (8%). Globigerinoides ruber alba (8%) and Hastigerina siphonifera (7%) are 147 also part of the association. In detail, G. inflata and G. ruber alba are more abundant in the first 148 100 m of the water column, while G. truncatulinoides peaks at 200 meters water depth (Pujol and 149 Vergnaud Grazzini, 1995).

The moderate mixing and ventilation processes, occurring during wintertime, bring the nutrients to the photic layer (Napolitano et al., 2000) as documented by the coccolithophorid occurrence in sediment traps collected in this area (Ziveri et al., 2000) and the satellite-derived surface chlorophyll concentration (D'Ortenzio and Ribera d'Alcalà, 2009). This hydrographic/oceanographic feature can also explain the presence of juvenile specimens of *G. inflata* and *G. truncatulinoides* in the surface layer (Pujol and Vergnaud Grazzini 1995).

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157 3. Sediment core

Sediment core KC01B was collected from a small ridge on the lower slope of the southern
Calabrian Ridge (Pisano Plateau, 36°15.250' N, 17°44.340' E, 3643 m water depth; Fig. 1) during
cruise MD69 of the R/V Marion Dufresne in 1991.

161 The lithology consists of hemipelagic marls, with intercalation of sapropels and the presence of a 162 number of thin tephra layers and few thin turbidite levels (Castradori 1993; Sanvoisin et al. 1993; 163 Langereis et al. 1997; Lourens 2004).

This 37 m thick sediment sequence represents an invaluable opportunity of investigating the early to late Pleistocene. Core KC01B has been intensively studied from different points of view, i.e., planktonic foraminifera, nannoplankton, stable isotopes, chemical and paleomagnetic analyses, tephra and sapropel presence (Castradori 1993; Sanvoisin et al. 1993; Dekkers et al. 1994, van Santvoort et al. 1997; Langereis et al. 1997; Rossignol-Strick et al. 1998; Rossignol-Strick and Paterne, 1999; Lourens, 2004; Maiorano et al., 2013; Insinga et al., 2014).

170 Moreover, Core KC01B is well known in the chronostratigraphic literature because it was used for 171 the construction of the Astronomical Time Scale (ATS) (Langereis et al., 1997) in the 172 Mediterranean region and to propose the Tyrrhenian as a regional stage for the Upper Pleistocene 173 (Cita et al., 2005). The ATS is based on the correlation of dominantly precession-controlled 174 sedimentary cycles (*i.e.* sapropels and carbonate cycles) to astronomical parameters. In particular, 175 this core was claimed to fill most of the gap between the oldest sapropel (S12) documented in 176 marine sediments (piston core RC9-181 - eastern Mediterranean Sea) dated at 483 ka (Lourens et 177 al., 1996a) and the youngest sapropel (v) exposed in the land-based marine successions of the 178 Vrica section (Southern Italy) dated at 1.280 Ma (Lourens et al., 1996b).

179 Concomitantly, Rossignol-Strick et al. (1998) proposed an alternative independent age model 180 based on tuning of the oxygen isotope record of KC01B with the ice sheet model of Imbrie and 181 Imbrie (1980).

Differences between both age models are in the order of 0-5 kyr and result from the choice of two different target curves and the adopted time lags between insolation forcing and climate response (Langereis et al., 1997) (for discussion see Hilgen et al., 1993; Lourens et al., 1996a). Largest differences (in the order of 10 kyr) between both age models occur around 618 and 785 ka.

Subsequently, Lourens (2004) established an improved sapropel-tuned age model for this core based on high-resolution color reflectance correlation with the Ocean Drilling Project (ODP) Site 964. This time-scale resulted from a revised chronology of the marine isotope record of Rossignol-Strick et al. (1998), implying a much more uniform change in sedimentation rate for the Ionian Sea cores and a good fit with other Mediterranean and open ocean marine isotope records.

We studied the sediment interval through sections 21-16 of the core (between 21.82 and 15.85 m composite depth, spanning the time interval from 507.3 to 292.1 ka - Lourens, 2004). This interval includes three sapropels (S10, S11 and S12) (Lourens, 2004) (Fig. 2).

194

195 4. Methods

Quantitative micropaleontological analyses were performed on 596 samples with a spacing of 1 cm
 (average time resolution of ~ 380yr). Samples were washed through 63 micron sieves and oven

dried at 50°C. Planktonic foraminiferal assemblage composition was determined analysing the >125 µm size fraction. For the micropaleontological census study, each sample was divided with a microsplitter to obtain unbiased aliquots with more than 300 planktonic foraminifers per sample. All taxa are quantified as percentages of the total number of planktonic foraminifers. The faunal data sets described in this paper have been archived, and are available in digital form, at PANGEA.

In this study, *Globigerinoides sacculifer* includes *Globigerinoides trilobus*, *Globigerinoides sacculifer* and *Globigerinoides quadrilobatus* (sensu Hemleben et al., 1989); *Neogloboquadrina pachyderma* sinistral (sin) has been counted separately from the dextral (dex) form.

We distinguished *N. incompta* from *N. pachyderma* by its development of a distinct apertural rim and a more lobulate outline. These taxa showed different vertical distribution and ecology. Studies performed on living planktonic foraminifers in Japan seas and in the eastern North Atlantic document the presence of *N. incompta* in shallower and warmer waters compared to *N. pachyderma* (Kuroyanagi and Kawahata, 2004; Schiebel et al., 2001).

Moreover, different morphotypes of *Globigerinoides ruber* (white) were identified using the morphotype concept of Wang (2000). The typical *Globigerinoides ruber* (d'Orbigny, 1839) is reported as *G. ruber* sensu stricto (s.s.), whereas *Globigerinoides elongatus* (d'Orbigny, 1826) and *Globigerinoides gomitulus* (Seguenza, 1880) are grouped and reported as *G. ruber* sensu lato (s.l.).

216 Several investigations, based on molecular genetics and geochemistry, highlighted the need to 217 revise the taxonomy of G. ruber (d'Orbigny, 1839) which shows remarkable "morphological" 218 variations (inter alias Darling et al., 1999; Kuroyanagi et al. 2008). In addition, the discrimination of 219 the different morphotypes appears to be necessary because they have significantly different 220 habitat preferences and thus different stable isotopic signals (Wang, 2000; Kuroyanagi and 221 Kawahata, 2004; Loewemark et al., 2005; Kawahata, 2005; Lin and Hsieh, 2007; Numberger et al., 222 2009). On the other hand, the recent investigation in the Gulf of Mexico of Thirumalai et al. (2014) 223 reports no evidence for discrepancies in s.s.-s.l. calcifying depth habitat or seasonality. The 224 controversial outcomes reported suggest that additional studies on the relationship between living 225 foraminiferal distribution and oceanographic conditions (productivity, stratification) in different

basins are necessary to build a more extensive picture of the ecological requirements of different
foraminiferal genetic types. In this work, we distinguished between different morphotypes in order
to test if they presented different distribution patterns in the past.

229 The paleoclimate curve was calculated following Cita et al. (1977) and Sanvoisin et al. (1993). It 230 represents the algebraic sum of warm-water species percentages (expressed as positive values) 231 and cold-water species percentages (expressed as negative values) based on ecological 232 preferences and modern habitat characteristics reported in Hemleben et al. (1989), Rohling et al. 233 (1993), and Pujol and Vergnaud-Grazzini (1995). Warm water species are all G. ruber (white and 234 pink varieties), G. sacculifer, Globigerinoides tenellus, Globigerina rubescens, Hastigerina 235 siphonifera (including G. calida) and Orbulina universa. The cold water species are Globigerina 236 bulloides, Globigerinita glutinata, Globorotalia scitula, Turbototalita guingueloba and N. 237 pachyderma (dex). Negative and positive values of the curve allow qualitative estimates for cold 238 and warm surface water respectively.

The paleoproductivity curve is based on a combination of planktonic foraminiferal species: it is calculated as the sum of *G. bulloides*, *G. glutinata* and *T. quinqueloba* percentages. All these taxa are related to high productivity environments (inter alias Bé and Tolderlund, 1971; Fairbanks and Wiebe, 1980; Pujol and Vergaud Grazzini, 1989).

243 In order to reconstruct paleoenvironmental and paleoceanographical conditions, the relative 244 abundance of some species or groups considered in this study are plotted in percentages with 245 respect to the total foraminiferal assemblage versus time. We assume that the habitat 246 characteristics of the different species during the Pleistocene were similar to those observed today. 247 Stable isotope analyses were carried out on about 10-15 specimens of the planktonic foraminifers 248 N. pachyderma (dex) and G. ruber (s.s. and s.l.) (from the >150 µm fraction) using an automated 249 carbonate reaction device Kiel III coupled to a Thermo-Finnigan MAR253 Analytical precision was better than 0.03 and 0.05‰ for δ^{13} C and δ^{18} O respectively as deduced from international NBS-19, 250 251 and in-house Naxos standards.

252

5. Age control

For the time interval considered in this work (290-510 ka), the age model by Lourens (2004) was partially modified and fine-tuned, considering the new chronological constraints on Pleistocene sapropel depositions (Konijnendijk et al. 2014).

257 We adopted the revised sapropel chronological framework in the eastern Mediterranean (ODP 258 sites 967 and 968) provided by Konijnendijk et al. (2014) using the highly linear relation between 259 the elemental ratio of titanium and aluminium in the sediment and insolation. The late Pleistocene 260 sapropel chronology of Konijnendijk et al. (2014) presents deviation from Lourens (2004) down to ~400 ka, where sapropel S^b in ODP 967/968 (Emeis et al., 2000) is correlated to one insolation 261 cycle older. Consequently, Sapropel S11 in KC01B [corresponding to sapropel S^b in ODP 967/968] 262 263 (Konijnendijk et al., 2014)] does not correspond to insolation cycle 38 (as reported in Lourens, 264 2004 at 407 ka) but to cycle 40 with a corresponding age of 418.9 ka (Konijnendijk et al., 2014).

As additional constraint, we used the tuning of the new high-resolution δ^{18} O data from *N. pachyderma* (sin) with the open-ocean benthic oxygen isotopic stack from Lisiecki and Raymo (2005) (Fig. 2).

Interpolation between consecutive tie-points was carried out by a linear function, assuming a constant sedimentation rate between the consecutive tie-points (mean sedimentation rate of 0.035 m/ka) and resulted in a higher resolution age control than in previous investigations (Castradori 1993; Sanvoisin et al., 1993; Dekkers et al., 1994, van Santvoort et al., 1997; Langereis et al., 1997; Rossignol-Strick et al., 1998; Rossignol-Strick and Paterne, 1999). Data used for the age model construction are listed in table 1.

This new Mediterranean δ^{18} O stack of *N. pachyderma* (dex) in the core KC01B has been compared with the δ^{18} O data of *G. inflata* of Voelker et al. (2010) from the Atlantic Ocean and with the δ^{18} O records of *G. bulloides* from ODP-site 975 studied by Pierre et al. (1999) and modified by Lourens (2004) and Kandiano et al. (2012). The good visual comparison between these climatic records supports the adopted age model (Fig. 2).

279

280 6. Results

281 6.1 Foraminiferal assemblages

282 Foraminiferal assemblages are rich and well preserved. The Shannon-Weaver Index commonly 283 varies between 0.99 and 2.6 (Fig. 3) and exhibits a sharply decreasing trend in diversity during the 284 glacials MIS 12 and MIS 10. The more pronounced minima occur in the upper part of the glacial 285 periods, when more than the 80% of the assemblages is composed by T. guingueloba, 286 neogloboquadrinids (N. pachyderma (dex) and N. dutertrei) and G. bulloides (Fig. 3). The 287 remaining 20% of the assemblages is represented by G. scitula and G. glutinata. During the 288 interglacials the diversity is higher with dominance of G. ruber group (about 50%) and the presence 289 of other warm water taxa such as O. universa, H. siphonifera, G.rubescens and G. tenellus (10%).

G. bulloides, neogloboquadrinids and T*. quinqueloba* show frequencies ranging from 10 to 20%.

The presence of *G. inflata* appears not to be related to glacial-interglacial phases; however higher abundances occur during interglacials (Fig. 3).

A discontinuous pattern is observed in the distribution of *G. sacculifer* and *Globorotalia truncatulinoides*. *G. sacculifer* peaks during interglacial phases with a maximum value of about 20% (Fig. 3). Its distribution is discontinuous during MIS 13, more continuous during MIS 11 and MIS 9. *G. truncatulinoides* characterises the foraminiferal assemblages during the middle and upper part of interglacials. The maximum abundance of this taxon (20-25%) is recorded during MIS 11 (Fig. 3).

299

300 7. Discussion

301 7.1 Glacials

Based on the planktonic foraminifera climatic curve, full glacial conditions occur during the upper part of MIS 12 and MIS 10 (Fig. 4). The climate conditions detected during the initial part of MIS 12 and MIS 10 are warmer than expected for a glacial stage as they fall in the range of values prevailing during the previously interglacials MIS 11 and MIS 9 (Fig. 4). These temperatures are principally related to higher percentages of warm surficial waters *G. ruber* group (Fig. 4). At this time, the assemblages are characterized by the dominance of *G. ruber* (s.l.) relative to *G. ruber*

308 (s.s.). All these morphotypes occur in tropical-subtropical regions and prefer well-stratified waters 309 but they show different habitats and seasonal preferences. In general, G. ruber (s.l.) calcifies 310 deeper than G. ruber (s.s.) (Wang, 2000; Lowermark et al., 2005) and reflects different nutrient 311 availability due to stratification of the water column (Lin et al., 2004). Based on the ecological 312 divergence of the morphotypes, we interpret the higher relative abundance of G. ruber (s.l.) during 313 the early part of MIS 12 and 10 as indicative primarily of the deepening of the summer thermocline. 314 Probably G. ruber (s.l.) has shifted their habitat in order to avoid oligotrophic surface waters documented at this time by the low values of the productivity curve (Fig. 4). In the present 315 316 Mediterranean Sea, low levels of production occur in summer, when the summer thermocline 317 deepens to ~90 m leading to a stable stratification (Klein and Coste, 1984; Krom et al., 1992). The small differences between δ^{18} O of *N. pachyderma* and *G. ruber* show that the two taxa share a 318 319 similar habitat indicating more homogenous conditions during this interval between surface and 320 intermediate waters (Fig. 4).

321 The climatic conditions detected during early MIS 12 in core KC01B are coeval with the relative 322 wintertime warmer sea surface temperatures (SST) documented in the nearshore waters off 323 Portugal and in the western Mediterranean basin by Voelker et al. (2010) and associated with the 324 increased heat transport by the Azores Current across the Atlantic. Moreover, the warming 325 detected during the lower half of MIS 10 displays the same warm temperature anomaly in the SST 326 record of Site 980 (North Atlantic, Feni Drift) (McManus et al., 1999) and site 1089 (South Atlantic 327 Subtropical Front) (Cortese et al., 2007) during full glacial MIS 10 centered at ca. 350 ka and linked 328 by the authors to a stronger than usual Agulhas Current influence. These similarities can be 329 explained by the quick response of sub-surficial Mediterranean waters to atmospheric processes 330 during both glacial stages. The warming occurring during the glacial half of both glacial MIS 10 and 331 MIS 12 implies an extra-regional connection between the Mediterranean sea and the northern and 332 southern hemisphere.

The subsequent increase in abundances of cold-water indicators *T. quinqueloba*, *N. pachyderma* (dex) and *G. bulloides* document the establishment of full glacial conditions during MIS 12 and MIS 10 at about 455 ka and 360 ka, respectively (Fig. 4). Well-known environmental preferences of

336 these species for nutrient-rich environments (Be and Tolderlund, 1971; Fairbanks and Wiebe, 337 1980; Reynolds and Thunell, 1986; Pujol and Vergnaud-Grazzini, 1995; Sierro et al., 2003) 338 suggest productive sea surface waters as also supported by the highest values of the 339 paleoproductivity curve (Fig. 4). At this time the fertilization can be triggered by the concurrence of 340 different factors. One is represented by eolian input due to enhanced North-African dust deposition 341 in the eastern Mediterranean (Roberts et al., 2011) during the upper part of the last five glacial 342 stages. However, eolian dust in general does not seem to provide an adequate and/or continuous 343 source of nutrients to enhance primary production (Krom et al., 2005; Incarbona et al., 2008). 344 Another important factor may be the higher buoyancy gradient due to the reduced Atlantic surface-345 waters inflow that has altered the equilibrium of vertical mixing in the water column.

346 The glacial sea-level lowstand at the time of MIS 12 and MIS 10, leads to a reduced Atlantic 347 surface-water influx and thus to shoaling of the density gradient (pycnocline) between intermediate 348 and surface waters within the Mediterranean. This shoaling is similar to what has been suggested 349 for other glacial sea-level lowstands (Rohling and Gieskes, 1989; Rohling, 1991; Rohling and 350 Bryden, 1994; Myers et al., 1998). This factor is more evident during MIS12 when sea level was 351 about 125 m below present (Rohling et al., 2009) (Fig. 4). MIS 12 is generally dominated by higher abundances (compared to MIS 10) of N. pachyderma (dex) and N. dutertrei reaching up to 60% of 352 353 the total assemblage. These taxa are indicative for the intensity of the deep chlorophyll maximum 354 (DCM), occurring when the upper part of the water column is isothermal and cold (Rohling and 355 Gieskes, 1989; Pujol and Vergnaud Grazzini, 1995). Such DCM may develop during periods of 356 reduced deep mixing. Similar evidence comes from the offset observed between N. pachyderma 357 and G. ruber oxygen isotope values (Fig. 4) that document stratification between the surficial and 358 the lower part of the photic zone. This is consistent with a different density gradient in the water 359 column due to the large decrease in Atlantic surface inflow.

The colder intervals of the investigated glacials are characterized by peaks in abundance of *N. pachyderma* (sin) (with values of 18%) at times of extremely negative, cold values of the climate curve (Fig. 5). *N. pachyderma* (sin) is known not only to prefer polar-subpolar waters (Hemleben et al., 1989; Bé and Tolderlund, 1971; Reynolds and Thunell, 1986; Dieckmann et al., 1991;

Johannessen et al., 1994), but also to be the dominant planktonic foraminiferal species during Heinrich events recorded in the North Atlantic Ocean (Heinrich, 1988; Bond et al., 1992).

366 Very rare specimens of N. pachyderma (sin) have been towed at the end of summer in the Ionian Sea (Pujol and Vergnaud-Grazzini, 1995); however this taxon is generally uncommon (<5%) in the 367 368 Mediterranean during the Quaternary (Thunell, 1978; Rohling and Gieskes, 1989; Rohling et al., 369 1993; Hayes et al., 1999; Sprovieri et al., 2003; Hayes et al., 2005; Triantaphyllou et al., 2009; 370 Siani et al., 2010; Sprovieri et al., 2012). Significant percentages of N. pachyderma (sin) in the 371 western Mediterranean Sea during the last glacial period have been interpreted as the result of 372 polar water intrusions into the Mediterranean via the Strait of Gibraltar at the time of the Atlantic 373 Heinrich events (Cacho et al., 1999; Pérez-Folgado et al., 2003; Sierro et al., 2005). The increases 374 in the relative abundance of N. pachyderma (sin), coeval with a SST drop comparable for 375 distribution and amplitude to the Heinrich-type events, have also been documented during MIS5 376 (Combourieu Neobout et al., 2002; Martrat et al., 2004), throughout MIS 15-9 (Girone et al., 2013) 377 and during MIS 100 (Becker et al., 2006) in the western and eastern basin.

378 At the investigated site (Fig. 5), we observe that the distribution of *N. pachyderma* (sin) is positively 379 correlated with that of neogloboguadrinids and that, despite the different time-resolution of the 380 data, the significant increase in abundance of N. pachyderma (sin) correlates well with that 381 observed in the western Mediterranean (ODP Site 975) (Girone et al., 2013) and North Atlantic Heinrich-type events (Stein et al., 2009; Rodrigues et al., 2011). In the Mediterranean Sea high 382 383 abundances of neogloboquadrinids are often associated with low-density water masses as 384 observed in sapropel deposition during colder climatic phases (Capotondi and Vigliotti, 1999; Negri 385 et al., 1999; Stefanelli et al., 2005). We can speculate that the cooling of Atlantic inflowing waters 386 due to iceberg melting, already documented in the western Mediterranean (Sierro et al., 2005; 387 Frigola et al., 2008), could also have affected the hydrological setting of the Ionian Sea. On the 388 other hand, this may have occurred because of the intense freshwater discharges into the 389 Mediterranean through the Black Sea from meltwater of lakes dammed by the Scandinavian ice 390 sheet (Sprovieri et al., 2012) and/or from NW African (Atlas) mountain glaciers (Rogerson et al., 391 2008). At present, we do not have clear evidence for local sources of water; further high-resolution

studies are necessary to understand the occurrence and distribution of *N. pachyderma* (sin) in the
 Mediterranean Sea.

394

395 7.2 Interglacials

396 From the paleoclimatic curve typical full interglacial conditions can for the first time be recognized 397 during MIS 11 at ~420 ka. The warm conditions culminate between 415 to 395 ka when the 398 microfauna is characterized by the presence of tropical-subtropical species G. sacculifer that 399 prefers warmer and shallower waters than G. ruber (Kuroyanagi and Kawahata, 2004). At present, 400 this species lives in the western Mediterranean at the end of summer (Pujol and Vergnaud-401 Grazzini, 1995) and its distribution in surface sediments from west to east reflects its strong 402 temperature dependence (Thunel, 1978). The increase in abundance of G. sacculifer in the 403 Mediterranean has been reported to be associated to "Climatic Optima" during the late Holocene 404 intervals e.g. Medieval Warm Period, the Roman Age, the late Bronze Age and the Copper Age 405 (Piva et al., 2008; Lirer et al., 2013; 2014). According, we interpret the occurrence of this species in 406 core KC01B during interglacial MIS 11 and MIS 9 as indicative of the "Climatic Optimum 407 interval/Thermal maximum". This interpretation concurs with the paleoclimate reconstruction of the 408 same period and for the same core, based on calcareous nannofossil assemblages (Maiorano et 409 al., 2013). The presence of an extended "Climatic Optimum", lasting ~30 ka, has been documented 410 in SST records across the world ocean (McManus et al., 1999; Hodell et al., 2000; De Abreu et al., 411 2005; Kandiano and Bauch, 2007; Dickson et al., 2009; Stein et al., 2009; Voelker et al., 2010), 412 from Antarctic ice cores (Petit et al., 1999; Jouzel et al., 2007) and in terrestrial pollen records 413 (Tzedakis, 2010). It occurs after a prolonged deglacial warming (Termination V) that extends into 414 MIS 11 with global and regional climate variability (Milker et al., 2013 and references therein).

The present high-resolution database provides new climatostratigraphic insights for the "Climatic Optimum" of the central Mediterranean region. It is interesting to note that, during MIS 11, *G. sacculifer* shows three different peaks of abundance concomitant with the decrease/absence of *G. truncatulinoides* (Fig. 4). *G. sacculifer* reaches its maximum quantity in the Eastern Basin, where

surface waters remain relatively warm, and low nutrient contents prevail throughout the year due to
the a relatively stable pycnocline at depth (Pujol and Vergnaud Grazzini, 1995).

Higher values of living *G. truncatulinoides* have been observed in the Mediterranean in areas of
intense water mixing during winter, i.e. the North Western Basin (Pujol and Vergnaud Grazzini,
1995), while in the more stratified waters of the Eastern Basin it is rarely found (Core top database
of Kallel et al. (1997).

Distributional trends of these taxa document that the "Climatic Optimum" of MIS 11 was not uniform but characterized by three distinct intervals with a year-around water column stratification and/or a weaker winter mixing with a consequent limited food supply. At the same time, higher values of the paleoclimate curve imply warmer superficial conditions or elevated winter sea surface temperatures (i.e. increase of the warm water taxa). The three warmer intervals present an estimated age of 415-410 ka, 408-403 ka, and 400-395 ka, respectively.

An interesting remark is that the peak from 400 to 395 ka is coeval with and shows the same length of the warm interval (400-395 ka Fig. 4) recorded in marine paleoclimate data off Iberia at the end of the MIS 11c (Koutsodendris and Zahn, 2014). Since that interval corresponds to an insolation minimum, this implies that orbital insolation forcing is not the warming cause.

Koutsodendris and Zahn (2014) motivate the North-Atlantic warmth as an interhemispheric teleconnection between strong leakage in the South Atlantic and Atlantic Meridional Overturning Circulation (AMOC)-driven warmth in the North Atlantic maintaining temperate conditions off Iberia and the continental Europe during the MIS11c. Accordingly, we speculate that the warmer intervals identified in our record are the result of an ocean-climate teleconnection between the high- and low- latitudes. Nevertheless additional sites are necessary in order to understand the exact mechanisms and extent of such climatic variability at time of the "Climatic Optimum".

The observed post-glacial temperature increase after Termination V does not match the Holocene SST trend when the "Climatic Optimum" already began after Termination I coincident with Sapropel S₁ deposition (Rohling and De Rijk, 1999; Capotondi et al., 1999; Cacho et al., 2001; Triantaphyllou et al., 2009). However a direct comparison between MIS 11 and MIS 1 is difficult to make because of a different phase in the orbital parameters. In fact, the present interglacial

(Holocene) spans a single summer insolation maximum (summer at 65°N), while MIS 11
interglacial optimum spans two (weak) insolation maxima (Laskar et al., 2004).

449 From 392 ka onward, the MIS11 in core KC01B is characterized by oligotrophic surface waters 450 during summer (Fig. 4) and eutrophic during winter, when deep-water masses are well-ventilated, 451 as testified by the presence of G. inflata and G. truncatulinoides. These conditions are comparable 452 to those in the present-day Ionian Sea. At present, G. inflata and G. truncatulinoides dominate the 453 fauna of the winter assemblages in the Ionian Sea (Pujol and Verganud Grazzini, 1995). 454 Distribution of planktonic foraminifera during MIS 9 document similar general trends as described for MIS 11, with a thermal maximum between 326 and 320 ka (Fig.4). However, MIS9 is 455 456 characterized by lower percentages of warm-water taxa compared to MIS 11, thus suggesting that 457 SSTs were lower during MIS9 than during MIS11.

458

459 7.3 Sea surface changes during transition T-V and T-IV

460

Termination V in core KC01B is marked by a decrease of δ^{18} O of *N. pachyderma* from +2.5‰ at 462 430 ka to +0.8‰ at 420 ka (Fig. 6). During this transition the planktonic foraminifera content shows 463 significant changes related to different environmental conditions (Fig. 6) and provides new data on 464 how the transitions evolved from glacial to interglacial.

465 The high abundance of the herbivorous species G. bulloides and T. guingueloba from ~430 to ~425 ka (Fig. 6) suggests enhanced nutrient supply in the sub-surface water masses. As these 466 467 species reach the highest concentrations in upwelling regions or in areas of vigorous vertical 468 mixing in the water column (Reynolds and Thunell, 1986), where high phytoplankton productivity 469 prevails, we can hypothesize the presence of an upwelling regime or continental coastal input at 470 this time, probably induced by the presence or intensification of the gyre similar to the one 471 observed at present in the Ionian Sea (Civitarese et al., 2010, Gacic et al., 2010). The absence of 472 G. inflata (Fig. 4) supports this oceanographic scenario, as this species prefers temperate and high 473 nutrient waters not affected by upwelling processes (Giraudeau 1993. The increase in primary 474 productivity is also documented by the calcareous nannofossil data (Maiorano et al., 2013). This 475 phase is coeval with a peak of iron-rich terrigenous dust at ODP Site 958 (Helmke et al., 2008),

476 related to the strengthening of trade winds over northwestern Africa. Probably, this strengthening 477 of westerlies is also associated with the strengthening of the Atlantic Ionian Stream jet leading to a 478 dynamic activity of mesoscale features such as meanders and eddies. The latter results in mixing 479 thus in enhanced nutrient supply, and leads to increased primary production. Several 480 oceanographic studies performed in different areas of the Mediterranean document that variability 481 in mesoscale hydrographic features leads to an increase of biological productivity (Estrada, 1996; 482 Christaki et al., 2011).

The relatively low abundance of *G. bulloides* between 424 and 420 ka, together with the increasing percentages of *N. pachyderma* (dex) and *N. dutertrei* (Fig. 6) documents the transition to stratified water conditions and the development of a DCM. The co-occurrence of *N. incompta, a species* that prefers shallower and warmer waters than *N. pachyderma* (Kuroyanagi and Kawahata, 2004), also suggests amelioration of climate (Fig. 6).

We interpret this microfaunal assemblage as the result of fresh-water input/land-derived nutrients associated with the climatic transition from glacial conditions of MIS 12 to the interglacial conditions of MIS 11. Further evidence is provided by the steadily decreasing δ^{18} O of *N. pachyderma* during the same period (Fig. 6), which might be explained by a SST increase or a salinity decrease due to surface water freshening during this phase.

The increase in *G. ruber* abundance at around 420 ka together with lighter oxygen isotopic values in *N. pachyderma* reflects the influence of the African humid phase in the Ionian sea that culminates with sapropel layer S11 deposition at 418.9 ka (Konijnendijk et al., 2014) (Fig. 6).

This is suggested by the timing that is coherent with the onset of the wet phase over North-West Africa (Helmeke et al., 2008), when the enhanced influence of the West African Monsoon system on the Saharan-Sahel region led to higher fresh-water input into the Mediterranean. During the early MIS 11 the African monsoon system intensification is also documented in marine records from the North Atlantic and western Mediterranean Sea and it has been related to the northwardmoving of the Intertropical Convergence Zone (ITCZ) (Kandiano et al., 2012).

502 In general, changes in the planktonic foraminiferal distribution observed during transition T-IV lead 503 to a similar climatic reconstruction as outlined for T-V (Fig. 6). The dominance of *G. bulloides* and

T. quinqueloba from 342 to 337 ka points to enhanced nutrient content and mixing/upwelling during
the first part of the transition from MIS 10 to MIS9. The subsequent replacement (~ 336-333 ka) of *N. pachyderma* and *N. dutertrei* suggests stratified conditions with a shallow mixed layer. Then,
Sapropel S10 occurs at 332 ka (Konijnendijk et al. 2014) (Fig. 6).

508 Based on our data, the dynamics of the sea surface property during Termination V are due to the 509 deglaciation and wind system variability. The timing and modalities of climate dynamics during the 510 Terminations in different regions are as yet not fully understood. Recent investigations focus on the 511 relative position of the Heinrich cold events that characterize all last five Terminations at a global 512 scale (Cortese et al., 2007; Cheng et al., 2009; Barker et al., 2011; Marino et al., 2015). During the 513 most recent Terminations I (MIS 2/MIS 1) and II (MIS 6/MIS 5) their positions show strong 514 differences in these two deglaciations and point to a bipolar seesaw control mechanism for 515 Termination II (Marino et al., 2015).

516 Observing the distributional trend of N. pachyderma (sin) in core KC01B, a "Heinrich-like" cold 517 event (see paragraph above) is clearly detected within both the transitions T-V and T-IV (Fig. 6) in 518 correspondence with a small drop in the oxygen isotope record (Fig. 6). Based on 519 paleoceanographic inferences provided by planktonic foraminifera, these cold episodes occur 520 approximately in the mid-point of the deglaciations, coincident with the increase in trade wind 521 intensity off NW Africa (Fig. 6). This suggests that they are probably linked to the wind influence 522 and thus to atmospheric conditions. This interpretation is also consistent with reported results from 523 other ocean basins, indicating that Heinrich-like events are associated with stronger winds (e.g. 524 Wang et al., 2001; Moreno et al., 2002; Itambi et al., 2009; Roberts et al., 2011). These are 525 probably induced by southward shifts of the Inter Tropical Convergence Zone (Jullien at al., 2007). 526 Accordingly, we conclude that the T-V and T-IV observed in the Mediterranean are not only 527 regional events but are associated with a dynamic reorganization of global atmospheric conditions. 528 Our environmental scenarios are consistent with the sequence of major events documented in the 529 last four Terminations that link the displacements of the ITCZ, the AMOC and the North Atlantic 530 cooling (Cheng et al., 2009; Schneider et al., 2014; Marino et al., 2015).

531 During glacial-interglacial transitions T-IV and T-V the climate/ocean interaction was probably 532 related to strong feedback processes: the weakening or shutdown of the AMOC due to enhanced 533 freshwater input to the North Atlantic resulted in an increase in sea surface temperature within the 534 tropics as well as in cooling of the North Atlantic and in the geographical shift of the wind system 535 over North Africa.

536 Clearly, more data with good age control are needed from a wider area so to substantiate and 537 evaluate extent and intensity of these events. This is not only needed to better understand 538 mechanisms of paleo-climate change but is also relevant for our abilities to forecast potential future 539 climate change processes.

540

541 8. Conclusions

A detailed study on planktonic foraminiferal assemblages from sediment core (KC01B) collected in the Ionian Basin (central Mediterranean Sea) allowed us to reconstruct the climate variability during glacial- interglacial periods between 500 and 300 ka (MIS 13 - MIS9). The main results can be summarized as follows:

- The early part of MIS 12 and MIS 10 is characterized by relatively "warm conditions" with a deepening of the summer thermocline derived from the quantitative distribution of *G. ruber* s.l. with respect to *G. ruber* s.s. Glacial conditions and eutrophic regimes are established in the upper half of the interval evidenced by significant increase of *T. quinqueloba*, N. *pachyderma* (dex) and *G. bulloides*. The colder intervals are interrupted by peaks in abundance of *N. pachyderma* (sin) coeval with north-Atlantic Heinrich-type cold events suggesting the close association of Central Mediterranean climate and North-Atlantic millennial-scale climate instability.

- Interglacials MIS 11 and MIS 9 have a prolonged "Climatic Optimum" lasting ~20 and 6 Ka respectively, as documented by the increase of the warm species *G. sacculifer*. Here for the first time we document that the extended warmth during the MIS 11c is characterized by three intervals with elevated winter sea surface temperatures and a weaker winter mixing.

557 - Complex paleoceanographic changes occurred during the glacial - interglacial transitions (T-V

and T-IV) consistent with the sequence of major events documented in the last four Terminations

that link the displacements of the ITCZ, the AMOC and the North Atlantic cooling.

560 The high-resolution investigations allow us to provide the timing of these changes occurring in the

561 Mediterranean region and to link these to global climate events.

562

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Figure captions

Fig. 1. (a) Location map of the investigated core site KC 01B and the discussed references cores; (b) Bathymetric map of the Ionian Sea (DTM 450 m resolution retrieved from http://portal.emodnethydrography.eu/EmodnetPortal/index.jsf#_); (c) mesoscale surface water circulation features of the Ionian Sea (from Napolitano et al. 2000 modified) are indicated (AIS= Atlantic-Ionian Stream, MAW= Modified Atlantic Water, MIJ= Mid Ionian Jet).

Fig. 2 - Comparison in time domain between the benthic oxygen isotopic stack of Liesicki and Raymo, (2005), the δ 18O *G. inflata* record of Voelker et al. (2010), the δ 18O *G. bulloides* record of Pierre et al. (1999) modified by Lourens (2004) and Kandiano et al. (2012), the δ 18O *Globigerinoides ruber* and *Neogloboquadrina pachyderma* dex records from the study KC01B core [3 points moving average (thick black curve) are superimposed on the δ 18O raw data], and the astronomical insolation curve of Laskar et al. (2004).

Fig 3 (a) Lithological log of sedimentary investigated time interval of core KC 01B, (b) planktonic oxygen isotope records, (c) diversity and (d) foraminiferal distribution (in percentages of total assemblages *versus age*) of the identified species. As "other" we grouped taxa with percentages <0.5 %. Chronology of sapropel (S10-S12) are according to Konijnendijk et al. (2014).

Fig. 4 Quantitative distribution patterns of selected planktonic species (percentage values plotted *versus age*) during MIS 13 and MIS 9 from core KC01B. Curve of Summer insolation at 65°N is from Laskar et al. (2004). Relative sea level record is from Rohling et al. (2009). Grey bands indicate interglacial intervals according to oxygen isotope chronology of Lisiecki and Raymo (2005); yellow bands correspond to the climatic optimum intervals. The MIS 13-9 and Termination V (T-V) and IV (T-IV) are indicated.

Fig. 5. Comparison between *N. pachyderma* (dex) δ 18O, foraminiferal paleoclimate curve; neogloboquadrinids and relative abundance of *N. pachyderma* (sin) records at core site KC01B and foraminiferal SST (f) and plot of *N. pachyderma* (sin.) at Mediterranean ODP Site 975 (Girone et al., 2013). For the Atlantic climate records are indicated: Alkenone-based SST and relative proportion of ice ice-rafted detritus (IRD) at Core MD03-2699 (Rodrigues et al., 2011). *N. pachyderma* (sin) distribution record and (m) proportion of IRD at core MD 01-2446 (Marino et al., 2014 and Voelker et al., 2010), the dolomite relative intensity at IODP Site U1313 (Stein et al., 2009). Light blu bands mark abrupt fluctuations of *N. pachyderma* (sin.) correlated to Heinrich-type events recorded at the western Mediterranean and Atlantic sites.

Fig. 6. Down core distribution of selected planktonic species during Termination V and IV from core KC01B. Curve of Summer insolation at 65°N is from Laskar et al. (2004). The colour bands indicate the different phases (see text for the descriptions). Position of sapropel S10 and S11 and Termination V (T-V) and IV (T-IV) are indicated. H1 and H4 refer to Heinrich-like event 1 and 4 following the nomenclature adopted by Girone et al. (2013).

1	Central Mediterranean Mid-Pleistocene paleoclimatic variability and its association with
2	global climate
3	
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17 18	Keywords: Planktonic foraminifera, Middle Pleistocene, Central Mediterranean, Paleoceanographic
19	changes.
20	
21	Abstract
22	Planktonic foraminiferal assemblages were studied at high-resolution in core KC-01B from the
23	Ionian Sea. Quantitative analysis allowed us to distinguish the main climatic features and
24	associated paleoceanographic changes, that occurred between Marine Isotopic Stages (MIS) 13
25	and 9 (~ 500300 ka).
26	MIS 12 and MIS 10 are characterized by relatively temperate conditions and an oligotrophic
27	oceanographic regime in the early part and by colder conditions and nutrient supply in the sub-
28	surface water masses in the upper part. During these intervals, small but distinct peaks of
29	Neogloboquadrina pachyderma sinistral (sin) are detected at times of extremely negative values of
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the planktonic foraminifera paleoclimatic curve. Their co-occurrence with similar episodes in the Atlantic suggests that the climate in the Central Mediterranean was associated with north-Atlantic millennial-scale climate instability. MIS 11 and MIS 9 are dominated by surficial warm-water taxa. The climate optimum is reached in the middle part of each of these stages, as denoted by the presence of *Globigerinoides sacculifer*, and persists for approximately 20 and 6 ka during MIS 11 and MIS 9 respectively. This warming <u>was-is</u> not constant but <u>is</u> characterized by three distinct intervals with elevated winter temperatures and/or weak winter mixing.

Distribution of *Globigerina bulloides*, *Turborotalita quinqueloba* and *Neogloboquadrina pachyderma* dextral (dex) indicates that significant environmental changes occur across the transitions from
 glacial to interglacial MIS 12/MIS 11 (Termination V) and MIS 10/MIS 9 (Termination IV).

40 The studied record documents a close linkage between Mediterranean climate evolution and
41 higher_ and lower_-latitude climate change throughout MIS 13—9.

42

43 1. Introduction

44 To improve our understanding of natural climate variability and our abilities to forecast future 45 climate change, it is essential to investigate geological climate archives with relevant climate 46 change events. Accordingly, this paper focuses on climate variability that occurred before and after 47 the Mid-Brunhes event (MBE) (Jansen et al., 1986) in the Mediterranean Sea. The investigated 48 time interval (-MIS 13-MIS 11; ~ 500-300 ka) was-is_characterized by substantially warmer 49 interglacials, (Epica Members, 2004; Jouzel et al., 2007) and an increase inenhanced atmospheric 50 CO₂ content, at levels similar to those for the pre-industrial Holocene (Siegenthaler -et al., 2005). In 51 particular, it includes the MIS 11c, traditionally considered as potential analogue for future climate 52 evolution because of relatively similar orbital climate forcing_(Loutre and Berger 2000; Masson-53 Delmotte et al., 2006; Tzedakis et al., 2012). Therefore, a thorough study of this interval will 54 provide information on type and magnitude of climate variability under non-anthropogenic but 55 otherwise comparable conditions to the present conditionspresent conditions.

In addition, the studied interval also includes the MIS 12, the most extreme glacial of the last 500
ka (Shackleton, 1987; Rohling et al., 1998; Lisiecki and Raymo, 2005) characterised by a sea level

of about 125 m lower than today_today (Rohling et al. 2009; 2014). The MIS 12---11 transition
(Termination V) is also part of this peculiar interval. It represents a glacial-interglacial transition that
is long compared to later Pleistocene terminations (Oppo et al., 1998; Bauch et al., 2000; Thunell
et al., 2002; Kandiano and Bauch, 2007; Helmke et al., 2008).

Notwithstanding the huge literature about the aforementioned climatic intervals, some features deserve additional clarifications. Two of the most intriguing aspects are the protracted deglaciation during Termination V and the cause of a long period of interglacial warmth during the MIS_11 (longer than any other mid-to late Pleistocene interglacial) with contrasting SST dynamics between polar- and mid- latitudes_(Helmke and Bauch, 2003; Kandiano and Bauck, 2007; Kandiano et al., 2012 Milker et al. 2013 and references therein).

These issues could be addressed by analysing paleo-data from the Mediterranean Sea, a_-region highly sensitive to atmospheric and climatic system modifications due to its intermediate latitudinal position, where Euro-Asian and North-<u>african_African_climate</u> regimes strongly interact <u>(Roether et</u> al., 1996; <u>Bethoux-Béthoux_et al., 1999; Pinardi and Masetti 2000; Trigo et al., 2004; Lionello et al.,</u> 2006).

Moreover the Mediterranean climate is exposed to the South_-Asian Monsoon in summer and the
Siberian High-Pressure_-System in winter (Luterbacher, 2005; Lionello et al., 2006). Today, the
southern part of the Mediterranean region is mostly influenced by the descending branch of the
Hadley cell, while the northern part is more linked to mid-latitude variability, characterized by the
North_-Atlantic Oscillation (NAO) and other mid-latitude teleconnection patterns (Hurrel et al.,
2004).

Thus, <u>climate_climate_investigations_-of geological archives of the Mediterranean region reflect</u> paleo-changes in the intensity and extension of global-scale climate patterns. In addition, the Mediterranean sedimentary sequences are characterized by the (quasi-)_periodical occurrence of episodes of deep-sea oxygen depletion (sapropel layers) (Olausson, 1991; Rohling et al., 2015 and references therein). Based on their link with the astronomical parameters (–Hilgen 1991; Rossignol—Strick, 1983; Lourens 2004; Konijnendijk et al., 2014) these discrete levels represent a useful constraint to establish accurate age <u>models_models</u> for marine and land sections. Formatted: Font: Not Italic

We present a new high-resolution quantitative study of planktonic foraminifera distribution throughout MIS 13-9 for sediment core KC01B collected in the Ionian Sea, central Mediterranean SeaMediterranean Sea. Planktonic foraminifera are amongst the most commonly used proxies for paleoceanographic and paleoclimate sea-surface—_reconstructions.—_Their distribution and abundance is strongly linked to surface-water properties. In addition, the physical and chemical properties of their shells reflect the-past environmental conditions of the water masses in which they lived (Kucera 2007 and references therein).

The investigated deep-marine sequence of core KC01^B represents a key site for stratigraphic and paleoclimatic investigations. This is not only because of its strategic location but also because it was used for the construction of a sapropel-based astronomical timescale for the last 1.1 My (Lourens, 2004). Moreover, in this work, we update the <u>studied time interval studied by using the</u> recent chronological constraints on Pleistocene sapropel deposition (Konijnendijk et al. 2014) and and a new oxygen isotopic record of *N. pachyderma*.

99 Our main aim is to explore how the ecosystem responded to climate variability during glacial and 100 interglacial intervals throughout MIS 13—9 in order to discuss the possible mechanisms through 101 which climate acts at the regional and global scale.

In detail, we focus on the main environmental and paleoceanographic processes occurring a) at
 times of glacial and interglacial MIS; b) during Termination V (T-V) and and Termination IV (T-IV).

104

105 2. Modern Oceanographic Setting

Currently, the Mediterranean Sea is an evaporative basin where freshwater loss exceeds
freshwater input, forcing an anti-estuarine circulation (Borghini et al., 2014).

At-<u>At</u> the surface (the first 100-200 m), moderate-salinity Atlantic Water (AW) intrudes through the Strait of Gibraltar and flows to the easternmost part of the Levantine basin while changingmodifying its temperature and salinity properties (Modified Atlantic Water – MAW). In the intermediate layer (depth between 150–200 and 600 m), Levantine Intermediate Water (LIW) forms in the eastern basin, spreads westwards and continues its flow towards the Strait of Gibraltar, and then into the Atlantic Ocean (Bethoux_Béthoux_et al., 1992; Robinson et al., 1991; Manca et al.,
2004; Malanotte-Rizzoli et al., 2014).

Atmospheric forcing and basin topography determine a large number of local cyclonic and anticyclonic cells (Pinardi and Masetti, 2000). In wintertime, outbreaks of cold and dry continental air masses lead to significant negative heat budgets and buoyancy losses, initiating deep and/or intermediate dense water formation both in the western and in the eastern basins (Malanotte-Rizzoli and Bergamasco, 1991; Castellari et al., 2000).

120 The Ionian Sea is influenced by the transit and on-site transformation of the major water masses 121 previously described (e.g., Modified Atlantic Water, MAW; Levantine Intermediate water, LIW; and 122 Eastern Mediterranean Deep Water, EMDW, MalonotteMalanotte--Rizzoli et al., 1997; Napolitano et al., 2000) (Fig. 1). At the near-surface level, most important for biological production, the MAW 123 124 enters in the western Ionian basin advected by the Atlantic Ionian Stream (AIS) (Fig. 1). Recently, 125 the upper-layer circulation in the Ionian seaSea has been associated with the deep thermohaline 126 circulation through the Bimodal Oscillating System (BiOS): the Ionian upper-layer circulation 127 reverses from cyclonic to anticyclonic and vice versa on decadal time scale affecting the biological productivity in the northern Ionian and southern Adriatic Adriatic Sea (Civitarese et al., 2010; Gačić 128 129 et al., 2010).

The <u>The</u> present-day Mediterranean Sea is characterized by oligotrophic conditions
(BethouxBéthoux, 1979; Sarmiento et al., 1988).-). The main factor that controls the seasonal
change in primary production is linked to the dynamics of the water column with increasing
biomass in late winter/early spring and decreasing in summer (Antoine et al., 1995; Bosc et al.,
2004; D'Ortenzio and Ribera d'Alcalà, 2009).

A significant West–East trophic gradient exists with nutrient depletion (mainly phosphorus) and a reduction in primary productivity in the eastern basin (Krom et al., 1991, 2010).

Moreover, primary productivity reflects the hydrological fragmentation due to mesoscale variability
(D'Ortenzio and Ribera d'Alcalà, 2009).

An-<u>An</u> oligotrophic regime, characterized_-by a low_-production, occurs_-in summer, when_-a_-stable
 stratification takes place (Klein and Coste, 1984; Krom et al., 1992; Crispi et al., 1999; Allen et al.,

141 2002). During this -period, low standing stocks characterize surface waters with the dominance of 142 predatory species. The Ionian planktonic foraminiferal association is generally dominated by 143 Globigerinoides ruber pink (40-60%) and G. ruber alba (20-40%) with -peaks -of -maximum 144 abundance in the first 50 m of the water column (Pujol and Vergnaud Grazzini, 1995). Winter 145 convection, and less frequently frontal zone migration or upwelling, brings nutrients into the photic 146 zone (mesotrophic regime) (Klein and Coste, 1984). During winter, the assemblage is 147 characterized by the dominance of grazing species such as Globorotalia truncatulinoides (50%) 148 and by the presence of other non-spinose species such as Globorotalia inflata (20%), and 149 Globigerina bulloides (8%). Globigerinoides ruber alba (8%) and Hastigerina siphonifera (7%) are 150 also part of the association. In detail, G. inflata and G. ruber alba are more abundant in the first 151 100 m of the water column, while G. truncatulinoides peaks at 200 meters water depth (Pujol and 152 Vergnaud Grazzini, 1995).

The moderate mixing and ventilation processes, occurring during wintertime, bring the nutrients to the photic layer (Napolitano et al., 2000) as documented by the coccolithophorid occurrence in sediment traps collected in this area (Ziveri et al., 2000)_—and the satellite-derived surface chlorophyll <u>concentration</u> (D'Ortenzio and Ribera d'Alcalà, 2009). This hydrographic/oceanographic feature can also explain the presence of juvenile specimens of *G. inflata* and *G. truncatulinoides* in the surface layer (Pujol and Vergnaud Grazzini 1995).

159

160 3. Sediment core

Sediment core KC01B was collected from a small ridge on the lower slope of the southern
Calabrian Ridge (Pisano Plateau, 36°15.250' N, 17°44.340' E, 3643 m water depth; Fig. 1) during
cruise MD69 of the R/V Marion Dufresne in 1991.

The lithology consists of hemipelagic marls, with intercalation of sapropels and the presence of a number of thin tephra layers and few thin turbidite levels (Castradori 1993; Sanvoisin et al. 1993; Langereis et al. 1997; Lourens 2004).

167 This 37 m thick sediment sequence represents an invaluable opportunity of investigating the early
168 to late Pleistocene. Core KC-01B has been intensively studied from different points of view, i.e.,

planktonic foraminifera, nannoplankton, stable isotopes, chemical and paleomagnetic analyses,
tephra and sapropel presence (Castradori 1993; Sanvoisin et al. 1993; Dekkers et al. 1994, van
Santvoort et al. 1997; Langereis et al. 1997; Rossignol-Strick et al. 1998; Rossignol-Strick and
Paterne, 1999; Lourens, 2004; Maiorano et al., 2013; Insinga et al., 2014, 2014).

173 Moreover, Core KC01B is well known in the chronostratigraphic literature because it was used for 174 the construction of the Astronomical Time Scale (ATS) (Langereis et al., 1997) in the 175 Mediterranean region and to propose the Tyrrhenian as a regional stage for the Upper Pleistocene 176 (Cita et al., 2005). The ATS is based on the correlation of dominantly precession-controlled 177 sedimentary cycles (i.e. sapropels and carbonate cycles) to astronomical parameters ... In 178 particular, this core was claimed to fill most of the gap between the oldest sapropel (S12) 179 documented in marine sediments (piston core RC9-181 - eastern Mediterranean Sea) dated at 483 180 ka (Lourens et al., 1996a) and the youngest sapropel (v) exposed in the land-based marine 181 successions of the Vrica section (Southern Italy) dated at 1.280 Ma (Lourens et al., 1996b).

182 Concomitantly, Rossignol-Strick et al. (1998) proposed an alternative independent age model 183 based on tuning of the oxygen isotope record of KC01B with the ice sheet model of Imbrie and 184 Imbrie (1980).

Differences between both age models are in the order of 0-5 kyr and result from the choice of two different target curves and the adopted time lags between insolation forcing and climate response (Langereis et al., 1997) (for discussion see Hilgen et al., 1993; Lourens et al., 1996a). Largest differences (in the order of 10 kyr) between both age models occur around 618 and 785 ka.

Subsequently, Lourens (2004) established an improved sapropel-tuned age model for this core based on high-resolution colour reflectance correlation with the Ocean Drilling Project (ODP) Site <u>Site</u> 964. This time-scale <u>resulted fromresulted from</u> a revised chronology of the marine isotope record of Rossignol-Strick et al. (1998), implying a much more uniform change in sedimentation rate for the Ionian Sea cores and a good fit with other Mediterranean and open ocean marine isotope records.

7

We studied the sediment interval through sections 21-16 of the core (between 21.82 and 15.85 m composite depth, spanning the time interval from 507.3 to 292.1 ka - Lourens, 2004). This interval includes three sapropels (S10, S11 and S12) (Lourens, 2004) (Fig. 2).

- 198
- 199 4. Methods

200 Quantitative micropaleontological analyses were performed on 596 samples with a spacing of 1 cm 201 (average time resolution of ~ 380yr). Samples were washed through 63 micron sieves and 202 ovendriedoven dried at 50°C. Planktonic foraminiferal assemblage composition was determined 203 analysing the >125 µm size fraction. For the micropaleontological census study, each sample was 204 divided with a microsplitter to obtain unbiased aliquots with more than 300 planktonic foraminifers 205 per sample. All taxa are quantified as percentages of the total number of planktonic foraminifers. 206 The faunal data sets described in this paper have been archived, and are available in digital form, 207 at PANGEA.

In this study, *Globigerinoides sacculifer* includes *Globigerinoides trilobus*, *Globigerinoides sacculifer* and *Globigerinoides quadrilobatus* (sensu Hemleben et al., 1989); *Neogloboquadrina pachyderma* sinistral (sin) has been counted separately from the dextral (dex) form.

We distinguished *N. incompta* from *N. pachyderma* by its development of a distinct apertural rim and a more lobulate outline. These taxa showed different vertical distribution and ecology. Studies performed on living planktonic foraminifers in Japan seas and in the eastern North Atlantic document the presence of *N. incompta* in shallower and warmer waters compared to *N. pachyderma* (Kuroyanagi and Kawahata, 2004; Schiebel et al., 2001).

Moreover, different morphotypes of *Globigerinoides ruber* (white) were identified using the morphotype concept of Wang (2000). The typical *Globigerinoides ruber* (d'Orbigny, 1839) is reported as *G. ruber* sensu stricto (s.s.), whereas *Globigerinoides elongatus* (d'Orbigny, 1826) and *Globigerinoides gomitulus* (Seguenza, 1880) are grouped and reported as *G. ruber* sensu lato (s.l.).

221 Several investigations, based on molecular genetics and geochemistry, highlighted the need to 222 revise the taxonomy of *G. ruber* (d'Orbigny, 1839) which shows remarkable "morphological"

223 variations (inter alias Darling et al., 1999; Kuroyanagi et al. 2008). In addition, the discrimination of 224 the different morphotypes appears to be necessary because they have significantly different 225 habitat preferences and thus different stable isotopic signals (Wang, 2000; Kuroyanagi and 226 Kawahata, 2004; Loewemark et al., 2005; Kawahata, 2005; Lin and Hsieh, 2007; Numberger et al., 227 2009). On the other hand, the recent investigation in the Gulf of Mexico of Thirumalai et al. (2014) 228 reports no evidence for discrepancies in s.s.-s.l. calcifying depth habitat or seasonality. The 229 controversial outcomes reported suggest that additional studies on-on the relationship between 230 living foraminiferal distribution and oceanographic conditions (productivity, stratification) in different 231 basins are necessary to build a more extensive picture of the ecological requirements of different 232 foraminiferal genetic types-. In this work, we distinguished between different morphotypes in order 233 to test if they presented different distribution patterns patterns in the past.

234 The paleoclimate curve was calculated following Cita et al. (1977) and Sanvoisin et al. (1993). It 235 represents the algebraic sum of warm-water species percentages (expressed as positive values) 236 and cold-water species percentages (expressed as negative values) based on ecological 237 preferences and modern habitat characteristics reported in Hemleben et al. (1989), Rohling et al. 238 (1993), and Pujol and Vergnaud-Grazzini (1995). Warm water species are all G. ruber (white and 239 pink varieties), G. sacculifer, Globigerinoides tenellus, Globigerina rubescens, Hastigerina 240 siphonifera (including G. calida) and Orbulina universa. The cold water species are Globigerina 241 bulloides, Globigerinita glutinata, Globorotalia scitula, Turbototalita guingueloba and N. 242 pachyderma (dex). Negative and positive values of the curve-curve allow qualitative estimates for 243 cold and warm surface water respectively.

The paleoproductivity curve is based on a combination of planktonic foraminiferal species: it is calculated as the sum of *G. bulloides*, *G. glutinata* and *T. quinqueloba* percentages. All these taxa are related to high productivity environments (inter alias Bé and Tolderlund, 1971; Fairbanks and Wiebe, 1980; Pujol and Vergaud Grazzini, 1989).

In order to reconstruct paleoenvironmental and paleoceanographical conditions, the relative abundance of some species or groups considered in this study are plotted in percentages with respect to the total foraminiferal assemblage versus time. We assume that the habitat characteristics of the different species during the Pleistocene were similar to those observed today. Stable isotope analyses were carried out on about 10-15 specimens of the planktonic foraminifers *N. pachyderma* (dex) and *G. ruber* (s.s. and s.l.) (from the >150 µm fraction) using an automated carbonate reaction device Kiel III coupled to a Thermo-Finnigan MAR253 Analytical precision was better than 0.03 and 0.05-‰ for δ^{13} C and δ^{18} O respectively as deduced from international NBS-19, and in-house Naxos standards.

257

258 5. Age control

For the time interval considered in this work (290-510 ka), the age model by Lourens (2004) was partially modified and fine-tuned, considering the new chronological constraints on Pleistocene sapropel depositions (Konijnendijk et al. 2014).

262 We adopted the revised sapropel chronological framework in the eastern Mediterranean (ODP 263 sites 967 and 968) provided by Konijnendijk et al. (2014) using the highly linear relation between 264 the elemental ratio of titanium and aluminium in the sediment and insolation. The late Pleistocene 265 sapropel chronology of Konijnendijk et al. (2014) presents deviation from Lourens (2004) down to ~400 ka, where sapropel S^b in ODP 967/968 (Emeis et al., 2000) is correlated to one insolation 266 267 cycle older. Consequently, Sapropel S11 in KC01B [corresponding to sapropel S^b in ODP 967/968 268 (Konijnendijk et al., 2014)] does not correspond to insolation cycle 38 (as reported in Lourens, 269 2004 at 407 ka) but to cycle 40 with a corresponding age of 418.9 ka (Konijnendijk et al., 2014).

As additional constraint, we used the tuning of the new high-resolution δ^{18} O data from *N. pachyderma* (sin) with the open-ocean benthic oxygen isotopic stack from Lisiecki and Raymo (2005) (Fig. 2).

Interpolation between consecutive tie-points was carried out by a linear function, assuming a constant sedimentation rate between the consecutive tie-points (mean sedimentation rate of 0.035 m/ka) and resulted in a higher resolution age control than in previous investigations (Castradori 1993; Sanvoisin et al., 1993; Dekkers et al., 1994, van Santvoort et al., 1997; Langereis et al.,

1997; Rossignol-Strick et al., 1998; Rossignol-Strick and Paterne, 1999). Data used for the age
model construction are listed in table 1.

This new Mediterranean δ^{18} O stack of *N. pachyderma* (dex) in the core KC01B has been compared with the δ^{18} O data of *G. inflata* of Voelker et al. (2010) from the Atlantic Ocean and with the δ^{18} O records of *G. bulloides* from ODP-site 975 studied by Pierre et al. (1999) and modified by Lourens (2004) and Kandiano et al. (2012). The good visual comparison between these climatic records supports the adopted age model (Fig. 2).

- 284
- 285 6. Results

286 6.1 Foraminiferal assemblages

287 Foraminiferal assemblages are rich and well preserved. The Shannon-Weaver Index commonly 288 varies between 0.99 and 2.6 (Fig. 3) and exhibits a sharply decreasing trend in diversity during the 289 glacials MIS 12 and MIS 10. The more pronounced minima occur in the upper part of the glacial 290 periods, when more than the 80% of the assemblages is composed by T. quinqueloba, 291 neogloboquadrinids (N. pachyderma (dex) and N. dutertrei) and G. bulloides (Fig. 3). The 292 remaining 20% of the assemblages is represented by G. scitula and G. glutinata. During the 293 interglacials the diversity is higher with dominance of G. ruber group (about 50%) and the presence 294 of other warm water taxa such as O. universa, H. siphonifera, G.rubescens and G. tenellus (10%).

295 G. bulloides, neogloboquadrinids and T. quinqueloba show frequencies ranging from 10 to 20%.

The presence of *G*. *inflata* <u>inflata</u> appears not to be related to glacial-interglacial phases; however higher abundances occur during interglacials (Fig. 3).

A discontinuous pattern is observed in the distribution of *G. sacculifer* and *Globorotalia truncatulinoides. G. sacculifer* peaks during interglacial phases with a maximum value of about 20% (Fig._3). Its distribution is discontinuous during MIS 13, more continuous during MIS 11 and MIS 9._-*G. truncatulinoides* characterises the foraminiferal assemblages during the middle and upper_-part of interglacials. The maximum abundance of this taxon (20-25%) is recorded during MIS_11 (Fig._3).

304

305 7. Discussion

306 7.1 Glacials

307 Based on the planktonic foraminifera climatic curve, full glacial conditions occur during the upper 308 part of MIS 12 and MIS 10 (Fig. 4). The climate -conditions detected during the initial part of MIS 309 12 and MIS 10 are -warmer than expected for a glacial stage as they fall in the range of values 310 prevailing during the previously interglacials MIS 11 and MIS 9 (Fig. 4). These temperatures are 311 principally related to higher percentages of warm surficial waters G. ruber group (Fig. 4). At this 312 time, the assemblages are characterized by the dominance of G. ruber (s.l.) relative to G. ruber 313 (s.s.). All these morphotypes occur in tropical-subtropical regions and prefer well-stratified waters 314 but they show different habitats and seasonal preferences. In general, G. ruber (s.l.) calcifies 315 deeper than G. ruber (s.s.) (Wang, 2000; Lowermark et al., 2005) and reflects different nutrient 316 availability due to stratification of the water column (Lin et al., 2004). Based on the ecological 317 divergence of the morphotypes, we interpret the higher relative abundance of G. ruber (s.l.) during 318 the early part of MIS 12 and 10 as indicative primarily of the deepening of the summer thermocline. 319 Probably -G. ruber (s.l.) has shifted their habitat in order to avoid oligotrophic surface waters 320 documented at this time by the low values of the productivity curve (Fig. 4). In the present 321 Mediterranean Sea, low levels of production occur in summer, when the summer thermocline 322 deepens to ~90 m leading to a stable stratification (Klein and Coste, 1984; Krom et al., 1992). The small differences between δ^{18} O of *N. pachyderma* and *G. ruber-ruber* show that the two taxa share 323 324 a similar habitat indicating more homogenous conditions during this interval between surface and 325 intermediate waters (Fig. 4).

The climatic conditions detected during early MIS 12 in core KC01^B are coeval with the relative wintertime warmer sea surface temperatures (SST) documented in the nearshore waters off Portugal and in the western Mediterranean basin by Voelker et al. (2010) and associated with the increased heat transport by the Azores Current across the Atlantic. Moreover, the warming detected during the lower half of MIS 10 displays the same warm temperature anomaly in the SST record_-of Site 980 (North Atlantic, Feni Drift) (McManus et al., 1999) and site 1089 (South Atlantic Subtropical Front) (Cortese et al., 2007) during full glacial MIS 10 centered at ca. 350 ka <u>and</u>-linked by the authors to a stronger than usual Agulhas Current influence. These similarities can be explained by the quick response of sub-surficial Mediterranean waters to atmospheric processes during both glacial stages. The warming occurring during the glacial half of both glacial MIS 10 and MIS 12 implies_-an extra-regional connection between the Mediterranean sea and the northern and southern hemisphere.

338 The subsequent increase in abundances of cold-water indicators T. guingueloba, N. pachyderma 339 (dex) and G. bulloides document the establishment of full glacial conditions during MIS 12 and MIS 340 10 at about 455 ka and 360 ka, respectively (Fig._4).-). Well---known environmental preferences of 341 these species for nutrient-rich environments (Be and Tolderlund, 1971; Fairbanks and Wiebe, 342 1980; Reynolds and Thunell, 1986; Pujol and Vergnaud-Grazzini, 1995; Sierro et al., 2003) 343 suggest productive sea surface waters as also supported by the highest values of the 344 paleoproductivity curve (Fig. 4). At this time the fertilization can be triggered by the concurrence of 345 different factors. One is represented by eolian input due to enhanced North-African dust deposition 346 in the eastern Mediterranean (Roberts et al., 2011) -during the upper part of the last five glacial 347 stages. However, the eolian dust in general does not seem able to provide an adequate and/or 348 continuous source of nutrients to enhance primary production (Krom et al., 2005; Incarbona et al., 349 2008). Another important factor may be the higher buoyancy gradient -due to the reduced Atlantic 350 surface-waters inflow that has altered the equilibrium of vertical mixing in the water column.

The glacial sea-level lowstand at <u>the_time of MIS 12 and MIS 10, leads_to</u> a reduced Atlantic surface-water influx and thus_-to_-shoaling of the density gradient (pycnocline) between intermediate and surface waters within the Mediterranean. <u>This shoaling is</u> similarly to what has been suggested for other glacial sea-level lowstands (Rohling and Gieskes, 1989; Rohling, 1991; Rohling and Bryden, 1994; Myers et al., 1998).

This factor is more evident_-during MIS12 when sea level was about 125 m below present (Rohling et al., 2009)_-(Fig._4). MIS 12 is generally dominated by higher abundances (<u>respect_compared to</u> MIS 10) of *N. pachyderma* (dex) and *N. dutertrei* reaching up to 60–% of the total assemblage. These taxa are indicative <u>of_for_the</u> intensity of the deep chlorophyll maximum (DCM), and <u>occursoccurring</u> when the upper part of the water column is isothermal and cold (Rohling and Gieskes, 1989; Pujol and Vergnaud Grazzini, 1995). Such DCM may develop during periods of reduced deep mixing._-A-similarSimilar evidence comes from the offset observed between *N. pachyderma* and *G. ruber* oxygen isotope values (Fig._4) that document a stratificationstratification between the surficial and the lower part of the photic zone. This is consistently with a different density gradient in the water column due to_-the large decrease in Atlantic surface inflow -inflow.

The colder intervals of the investigated glacials are characterized by peaks in abundances of *N. pachyderma* (sin) (with values of 18%) at times of extremely negative, cold values of the climate curve (Fig._5). *N. pachyderma* (sin) is known not only to prefer polar-subpolar waters (Hemleben et al., 1989; Bé and Tolderlund, 1971; Reynolds and Thunell, 1986; Dieckmann et al., 1991; Johannessen et al., 1994), but also to be the dominant planktonic foraminiferal species during Heinrich events recorded in the North Atlantic <u>oceanOcean</u> (Heinrich, 1988; Bond et al., 1992).

373 Very rare specimens of N. pachyderma (sin) have been towed at the end of summer in the Ionian 374 Sea (Pujol and Vergnaud-Grazzini, 1995); however this taxon is generally uncommon -(<5%) in the 375 Mediterranean during the Quaternary (Thunell, 1978; Rohling and Gieskes, 1989; Rohling et al., 376 1993; Hayes et al., 1999; Sprovieri et al., 2003; Hayes et al., 2005; Triantaphyllou et al., 2009; 377 Siani et al., 2010-; Sprovieri et al., 2012). -Significant percentages of N. pachyderma (sin) in the 378 western Mediterranean Sea during the last glacial period have been interpreted as the result of 379 polar water intrusions into the Mediterranean via the Strait of Gibraltar at_-the time of the Atlantic 380 Heinrich events (Cacho et al., 1999; Pérez-Folgado et al., 2003; Sierro et al., 2005). The increases 381 in the relative abundance of N. pachyderma (sin), coeval with a SST drop comparable for 382 distribution and amplitude to the Heinrich-type events, have also been documented during MIS5 383 (Combourieu Neobout et al., 2002; Martrat et al., 2004), throughout MIS 15-9 (Girone et al., 2013) 384 and during MIS 100 (Becker et al., 2006) in the western and eastern basin.

At the investigated site (Fig. 5), we observe that the distribution of *N. pachyderma* (sin) is positively correlated with that of neogloboquadrinids and that, despite the different time-resolution of the data, the significant increase in abundance of *N. pachyderma* (sin) correlates well with that observed in the western Mediterranean (ODP Site 975) (Girone et al., 2013) and North Atlantic 389 Heinrich-type events (Stein et al., 2009; Rodrigues et al., 2011). In the Mediterranean Sea high 390 abundances of neogloboquadrinids are often associated with low-density water masses as 391 observed in sapropel deposition during colder climatic phases (Capotondi and Vigliotti, 1999; Negri 392 et al., 1999; Stefanelli et al., 2005). We can speculate that the cooling of Atlantic inflowing waters 393 due to iceberg melting, already documented in the western Mediterranean (Sierro et al., 2005; 394 Frigola et al., 2008), could also have affected the hydrological setting of the Ionian Sea. On the 395 other hand, this may have occurred because of the intense freshwater discharges into the 396 Mediterranean through the Black Sea from meltwater of lakes dammed by the Scandinavian ice 397 sheet (Sprovieri et al., 2012) and/or from NW African (Atlas) mountain glaciers (Rogerson et al., 398 2008). At the present, we do not have clear evidences for of a local sources of water; and further 399 high-resolution studies are necessary to understand the occurrence and distribution of N. 400 pachyderma (sin) in the Mediterranean Sea.

401

402 7.2 Interglacials

403 From the paleoclimatic curve_During MIS 11_typical full interglacial conditions can for the first time
 404 be recognized during MIS 11 at around ~420 ka-on the paleoclimatic curve.

405 The warm conditions culminate between 415 to 395 ka when the microfauna is characterized by 406 the presence of the-tropical---subtropical species G. sacculifer that prefers warmer and shallower 407 waters than G. ruber (Kuroyanagi and -- Kawahata, 2004). At present, this species lives in the 408 western Mediterranean at the end of summer (Pujol and Vergnaud-Grazzini, 1995) and its 409 distribution in surface sediments from west to east reflects its strong temperature dependence 410 (Thunel, 1978). The increase in abundance of G. sacculifer in the Mediterranean has been 411 reported to be associated to "Climatic Optima" during the late Holocene intervals e.g. Medieval 412 Warm Period, the Roman Age, the late Bronze Age and the Copper Age (Piva et al., 2008; Lirer et 413 al., 2013; 2014). According, we interpret the occurrence of this species in core KC01B during 414 interglacial MIS 11 and MIS 9 as indicative of the "Climatic Optimum interval/Thermal maximum". 415 This interpretation concurs with the paleoclimate reconstruction of the same period and for the 416 same core, based on calcareous nannofossil assemblages (Maiorano et al., 2013). - The presence

of an extended "Climatic Optimum", lasting ~30 ka, has been documented in SST records across the world ocean (McManus et al., 1999; Hodell et al., 2000; De Abreu et al., 2005; Kandiano and Bauch, 2007; Dickson et al., 2009; Stein et al., 2009; Voelker et al., 2010), from Antarctic ice cores (Petit et al., 1999; Jouzel et al., 2007) and in terrestrial pollen records (Tzedakis, 2010)...). It occurs after a prolonged deglacial warming_-(Termination \mp V) that extends into MIS 11 with global and regional climate variability (Milker et al., 2013 and references therein).

The present_-high-resolution database provides new climatostratigraphic insights for the "Climatic Optimum" of the central Mediterranean region. It is interesting to note that, during MIS 11, *G. sacculifer* shows three different_-peaks of abundance concomitant with the decrease/absence of *G. truncatulinoides_*-(Fig. 4). *G. sacculifer* reaches its maximum quantity in the Eastern Basin, where surface waters remain relatively warm, and low nutrient contents prevail throughout the year due to the a relatively stable_-pycnocline at depth (Pujol and Vergnaud Grazzini, 1995).

Higher values of living *G. truncatulinoides* have been observed in the Mediterranean in_-areas of intense water mixing during winter, i.e. the North Western Basin (Pujol and Vergnaud Grazzini, 1995), while in the more stratified waters of the Eastern Basin it is rarely found (Core top database of Kallel et al. (1997).

Distributional trends of these taxa document that the "Climatic Optimum" of MIS 11 was not uniform but characterized by three distinct intervals with a year-around water column stratification and/or a weaker winter mixing with a consequent limited food supply. At the same time, higher values of the paleoclimate curve imply warmer superficial conditions_-or elevated winter sea surface temperatures (i.e. increase of the warm water taxa). The three warmer intervals present an estimated age of 415-410 ka, 408-403 ka, 400and 400-395 ka, respectively.

An interesting remark is that the peak from 400 to 395 ka is coeval with and shows the same length of the warm interval (400-395 ka Fig._4) recorded in marine paleoclimate data off Iberia at the end of the MIS_11c (Koutsodendris and Zahn, 2014)._-Since that interval corresponds to an insolation minimum, this implies that orbital insolation forcing is not the warming cause.

443 Koutsodendris and Zahn_–(2014) motivate the North-Atlantic warmth as an interhemispheric 444 teleconnection between strong leakage in the South Atlantic and Atlantic Meridional Overturning Circulation (AMOC)-driven warmth in the North Atlantic maintaining temperate conditions off Iberia and the continental Europe during the MIS11c. Accordingly, we speculate that the warmer intervals identified in our record are the result of an ocean-climate teleconnection between the high- and low- latitudes. Nevertheless–_additional sites are necessary in–_order to understand the exact mechanisms and extent of such climatic variability at time of the "Climatic Optimum".

The observed post-glacial temperature increase after Termination V does not match the Holocene SST trend when the "Climatic Optimum" already began after Termination I coincident with Sapropel S₁ deposition (Rohling and De Rijk, 1999;__Capotondi et al., 1999;__Cacho et al., 2001; Triantaphyllou et al., 2009). However a direct comparison between MIS 11 and MIS 1 is difficult to make because of a different phase in the orbital parameters. In fact, the present interglacial (Holocene) spans a–_single summer insolation maximum (summer at 65º_N), while MIS 11 interglacial optimum spans two (weak) insolation maxima (Laskar et al., 2004).

457 From 392 ka onward, the MIS11 in core KC01B is characterized by oligotrophic surface waters 458 during summer (Fig. 4) and eutrophic during winter, when deep-water masses -are well-ventilated, 459 as testified by the presence of G. inflata and G. truncatulinoides. These conditions are comparable 460 to those in the present-day Ionian Sea. At present, G. inflata and G. truncatulinoides dominate the 461 fauna of the winter assemblages in the Ionian Sea (Pujol and Verganud Grazzini, 1995). 462 Distribution -of planktonic foraminifera during MIS 9 document similar general trends as described 463 for MIS 11, with a thermal maximum between 326 and 320 ka (Fig.4). However, MIS9 is 464 characterized by lower percentages of warm-water taxa compared to MIS 11, thus suggesting that 465 SSTs were lower during MIS9 than during MIS11.

466

467 7.3 Sea surface changes during transition T-V and T-IV

468

Termination V in core KC01B_-is marked by a decrease of δ^{18} O of *N. pachyderma* from +2.5‰ at 430 ka to +0.8–‰ at 420 ka (Fig._6).–_During this transition the planktonic foraminifera content shows significant changes related to different environmental conditions (Fig._6) and provides new 472 data on how the transitions evolved from glacial to interglacial. 473 The high abundance of the herbivorous species G. bulloides and T. guingueloba from ~430 to 474 ~425 ka (Fig._6)-_suggests enhanced nutrient supply in the sub-surface water masses. As these 475 species reach the highest concentrations in upwelling regions or in areas of vigorous vertical 476 mixing in the water column (Reynolds and Thunell, 1986), where high phytoplankton productivity 477 prevails, we can hypothesize the presence of an upwelling regime or continental coastal input at 478 this time, probably induced by the presence or intensification of the gyre similar to the one 479 observed at present in the Ionian Sea (Civitarese et al., 2010, Gacic et al., 2010). The absence of 480 G. inflata inflata -(Fig. 4) supports this oceanographic scenario, as this species prefers temperate 481 and high nutrient waters not affected by upwelling processes (Giraudeau 1993. The increase in 482 primary productivity is also documented by the -calcareous nannofossil data (Maiorano et al., 483 2013). This -phase is coeval with -a -peak of iron--rich terrigenous dust at ODP Site 958 (Helmke et 484 al., 2008), related to the strengthening of trade winds over northwestern Africa. Probably, this 485 strengthening of westerlies is also associated with the strengthening of the Atlantic Ionian Stream 486 jet leading to a dynamic activity of mesoscale features such as meanders and eddies. The latter 487 results in mixing thus in enhanced nutrient supply, and leads to increased enhanced primary 488 production. Several oceanographic studies performed in different areas of the Mediterranean 489 documented that the variability in mesoscale hydrographic features leads to an increase of 490 biological productivity (Estrada, 1996; Christaki et al., 2011).

The relatively low abundance of *G. bulloides* between 424 and 420 ka, together with the increasing percentages of *N. pachyderma* (dex) and *N. dutertrei_*-(Fig._6). documents the transition to stratified water conditions and the development of a DCM. The co-occurrence of *N. incompta<u>, a species</u>* that prefers shallower and warmer waters than *N. pachyderma* (Kuroyanagi and Kawahata, 2004), also suggests amelioration of climate (Fig. 6).

We interpret this microfaunal assemblage as the result of fresh-water input/land-derived nutrients associated with the climatic transition from glacial conditions of MIS 12 to the interglacial conditions of MIS 11._-Further evidence is provided by the steadily decreasing δ^{18} O of *N. pachyderma* during the same period (Fig. 6), which might be explained by a SST increase or a salinity decrease due to surface water freshening during this phase. The increase in *G. ruber* abundance at around 420 ka together with <u>ligherlighter</u> oxygen isotopic values in *N. pachyderma*_-reflects the influence of the African humid phase in the Ionian sea that culminates with sapropel layer S11 deposition at 418.9 ka (Konijnendijk et al., 2014) (Fig. 6).

This is suggested by the <u>coherent</u>-timing <u>that is coherent</u> with the onset of the wet phase over North-West Africa (Helmeke et al., 2008), when the enhanced influence of the West African Monsoon system on the Saharan-Sahel region led <u>to</u>higher fresh-water input into the Mediterranean.__During the early MIS 11_<u>-the</u> African monsoon system intensification is__also documented_-in marine records from the North Atlantic and western Mediterranean <u>seaSea</u> and it has been related to the northward-moving of the Intertropical Convergence Zone (ITCZ) (Kandiano et al., 2012).

In general, changes in the planktonic foraminiferal distribution observed during transition T_-IV lead to <u>a</u> similar climatic reconstruction as outlined for T_-V (Fig. 6). The dominance of *G. bulloides* and *T. quinqueloba*_-from 342_to_- 337 ka_-points to enhanced nutrient content and mixing/upwelling during the first part of <u>the</u> transition from MIS 10 to MIS9. The subsequent replacement (~ 336-333 ka) of *N. pachyderma* and *N. dutertrei* suggests stratified conditions with <u>a</u> shallow mixed layer-low surficial water masses. Then, Sapropel S10 occurs at 332 ka (Konijnendijk et al. 2014) (Fig. 6).

517 Based on our data, the dynamics of the sea surface property during Termination V are due to the 518 deglaciation and wind system variability. The timing and modalities of climate dynamics during the 519 Terminations in different regions are as yet not fully understood. Recent investigations focus on the 520 relative position of the Heinrich cold events that characterize all last five Terminations at a global 521 scale (Cortese et al., 2007; Cheng et al., 2009; Barker et al., 2011; Marino et al., 2015). During -the 522 most recent Terminations I (MIS -2/MIS -1) and II (MIS 6/MIS 5) their positions show strong 523 differences in these two deglaciations and evidence that point to a bipolar seesaw controls the 524 mechanism of for termination Termination II (Marino et al., 2015).

525 Observing the distributional trend of *N. pachyderma* (sin) in core KC01B, a "Heinrich-like" cold 526 event (see paragraph above) is clearly detected within both the Transitions transitions T_zV and T_z 527 IV (Fig. 6) in correspondence with a small drop in the oxygen isotope record –(Fig. 6). Based on 528 paleoceanographic inferences provided by planktonic foraminifera, these cold episodes occur

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approximately in the mid-point of the deglaciations, _-coincident with the increase in trade wind
intensity off NW Africa (Fig._6). This suggests that they are probably linked to the wind influence
and so-thus to the atmospheric conditions.

This interpretation is also consistent with reported results from other ocean basins, indicating that Heinrich-like events are associated_-with stronger winds (e.g. Wang et al., 2001; Moreno et al., 2002; Itambi et al., 2009; Roberts et al., 2011). These are-and probably induced by southward shifts of the Inter Tropical Convergence Zone (Jullien at al., 2007).

Accordingly, we conclude that the T-V and T-IV observed in the Mediterranean are not only regional events but are associated with a dynamic reorganization of global atmospheric conditions. Our environmental scenarios are consistent with the sequence of major events documented in the last four Terminations that link the displacements of the ITCZ, the AMOC and the North Atlantic cooling (Cheng et al., 2009; Schneider et al., 2014; Marino et al., 2015).

- 541 During glacial-interglacial transitions T-IV and T-V the climate/ocean interaction was probably 542 related to strong feedback processes: the weakening or shutdown of the Atlantic Meridional 543 Overturning Circulation (AMOC) due to the <u>enhanced</u> freshwater input to the North Atlantic 544 resulted in an increase in sea surface temperature within the tropics as well as in cooling of the 545 North Atlantic and in the geographical shift of the wind system over North Africa.
- 546 Clearly, more data with good age control are needed from a wider area so to substantiate and 547 evaluate extent and intensity of these events. This is not only needed to better understand 548 mechanisms of paleo-climate change but is also relevant for our abilities to forecast potential future 549 climate change processes.
- 550

551 8. Conclusions

A detailed study on planktonic foraminiferal assemblages from sediment core (KC01B) collected in the Ionian Basin (central Mediterranean Sea) allowed us to reconstruct the climate variability during glacial- interglacial periods between 500 and 300 ka (MIS 13_-_MIS9). The main results can be summarized as follows:

20

- The early part of MIS 12 and MIS 10 is characterized by relatively "warm conditions" with a deepening of the summer thermocline_-derived from the quantitative distribution of *G. ruber* s.l. with respect to *G. ruber* s.s. Glacial conditions and eutrophic regimes are established in the upper half of the interval evidenced by significant increase of *T. quinqueloba*, N. *pachyderma* (dex) and *G. bulloides*. The colder intervals are interrupted by peaks in abundance of *N. pachyderma* (sin) coeval with north-Atlantic Heinrich-type cold events suggesting the close association of Central Mediterranean climate and North-Atlantic millennial-scale climate instability.

- Interglacials MIS 11 and MIS 9 have a prolonged "Climatic Optimum" lasting ~20 and 6 Ka
respectively, as documented_by the increase of the warm species *G. sacculifer*._Here for the first
time we document that the extended warmth during the MIS 11c is characterized by three intervals
with elevated -winter sea surface temperatures -and a weaker winter mixing.

567 - Complex paleoceanographic changes occurred during the glacial - interglacial transitions (T-V

and T-IV) consistent with the sequence of major events documented in the last four Terminations

that link the displacements of the ITCZ, the AMOC and the North Atlantic cooling.

570 The high-resolution investigations allow_-us to provide the timing of these changes occurring in the

571 Mediterranean region<u>and to link these to global climate events</u>.

572

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Responses to the reviewers

Reviewer #1

- *I would like to highlight a list of typos throughout the manuscript (clean version): lines 50, 51, 54, 55, 58, 59, 63, 68, 70, 77, 79, 81, 83, 85, 86, 95, 100, 101, 106, 122, 125, 126, 128, 129, 137-139, 141-144, 147, 148, 153, 154, 175, 185, 189, 222-225, 228, 229, 238, 292, 297, 298, 304-306, 308, 314, 319, 321-325, 327, 330, 335, 339, 341, 344, 347, 351-353, 356, 359, 360, 371, 373, 400, 401, 410, 414, 415, 417, 419, 420, 422, 423, 430, 435, 439-442, 446, 449, 452, 456, 463-465, 467, 468, 474, 476-478, 485-488, 491, 495, 496, 499, 500, 505, 513, 521, 525, 535, 549, 556-558 and 562. OK. Done.*

Typing errors highlighted throughout the manuscript probably are due to the translation from the "file.word" to the "file.pdf" as I checked throughout the version".word" and they are not present. However, I did not understand why it happened. I think this is probably due to the fact that I have submitted the manuscript as a file "word.docx" and not "word.doc". Probably the system does not convert well the version "word.docx".

- *remove AMOC, the acronym was already cited.* OK. Done.

Reviewer #2

- I suggest that a native english speaker should get a last and fresh read over the manuscript

OK. Done.

Tie-point	Depth (mcd)	Age kyr	Reference
sapropel S'	15,74	287,5	Konijnendijk et al. (2014)
sapropel S10	16,84	332,3	Konijnendijk et al. (2014)
base MIS 9	17,15	337	Lisiecki and Raymo (2005)
sapropel S11	19,27	418,9	Konijnendijk et al. (2014)
base MIS 12	20,83	474	Lisiecki and Raymo (2005)
sapropel S12	21,74	504,5	Konijnendijk et al. (2014)

Tab. 1 - Tie-points used in this work for the construction of the age model

Figure1 Click here to download high resolution image











