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This is a pre print version of the following article:

*Original:*

Di Stefano, A., Baldassini, N., Maniscalco, R., Speranza, F., Maffione, M., Cascella, A., et al. (2015). New bio-magnetostratigraphic data on the Miocene Moria section (Northern Apennines, Italy): connections between the Mediterranean region and the North Atlantic Ocean. NEWSLETTERS ON STRATIGRAPHY, 48(2), 135-152 [10.1127/nos/2015/0057].

*Availability:*

This version is available <http://hdl.handle.net/11365/990263> since 2021-03-25T18:18:10Z

*Published:*

DOI:10.1127/nos/2015/0057

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# New bio-magnetostratigraphic data on the Miocene Moria section (Northern Apennines, Italy): connections between the Mediterranean region and the North Atlantic Ocean

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With 8 figures and 3 tables

**Abstract.** New bio-magnetostratigraphic data have been acquired from the Burdigalian part of the Moria section in the Umbria–Marche Apennine (Central Italy). The investigated sedimentary sequence is 55 meters thick and comprises the transition between the Bisciario and the Schlier formations (*Auctorum*), composed of five meters of indurated marly limestones, followed by about 40 meters of blue marly clays capped by 10 m of alternating clays and calcareous layers. According to existing literature, the sequence contains the so-called “Piero della Francesca Level” (*Auctorum*), a few-centimeters-thick biotite-rich level. The sequence has been sampled for paleomagnetic and calcareous plankton analyses. Four magnetic polarity zones have been documented in the middle and upper part of the section. The calcareous plankton content shows different degrees of preservation, from poor to good for calcareous nannofossils and poor to medium for planktonic foraminifers. Several useful biohorizons have been observed and chronologically constrained through calibration to the magnetostratigraphic data.

The studied section ranges from the top of Chron C5Dr.1n to Chron C5Cn.2r, covering the time interval from 17.7 to 16.4 Ma (according to the ATNTS04) almost continuously, with a small hiatus of at least 34 ka at the C5Dn/C5Cr transition. The integration of our results with those from previous studies of the same section and comparisons with data from other Mediterranean and North Atlantic sequences allows to evaluate the synchrony of the observed biohorizons with other parts of the Mediterranean and the Atlantic Ocean during a still relatively poorly known time interval.

**Key words.** Miocene, calcareous plankton biostratigraphy, magnetostratigraphy, Mediterranean, North Atlantic

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## 1. Introduction

The Moria section in the Umbro–Marche Apennine of Central Italy represents a well-known lower–middle Miocene sedimentary sequence in the Mediterranean region (Fig. 1). The section has been subject to detailed integrated stratigraphic studies over the past two decades because of the favorable exposure, the presence of abundant calcareous plankton and the presence of radiometrically datable layers (Fornaciari and Rio 1996, Fornaciari et al. 1996, Deino et al. 1997, Di Stefano et al. 2008). A detailed lithostratigraphy has been presented by Deino et al. (1997), who published an integrated bio-magnetostratigraphic study (Fig. 2) for the ca. 100 m thick succession. However, reliable paleomagnetic directions are limited and discontinuous throughout the studied section. In this study we re-examined the lower, 55-m-thick portion of the section coinciding with the lower part of the section reported by Deino et al. (1997) (Figs. 2, 3), namely the segment encompassing the Bisciario/Schlier transition.

The interpretation of the observed magnetozones was made through tie-points provided by main calcareous plankton horizons. Furthermore, a number of additional plankton events were detected and magnetostratigraphically constrained. The obtained data were integrated with those gathered previously by Di Stefano et al. (2008), and compared with other Mediterranean sections and with North Atlantic DSDP Hole

608 (Ruddiman et al. 1987, Miller et al. 1991, Olafsson 1991, Gartner 1992, Di Stefano et al. 2011a). This resulted in an improved stratigraphic resolution for the interval ranging from the base of Chron C6n (19.7 Ma) to Chron C5ADr (14.7 Ma), and the establishment of correlations between the Mediterranean region and the North Atlantic Ocean across the lower–middle Miocene boundary.

## 2. Geological setting

The Moria section (base: 43° 30.176' N, 12° 35.619' E; top: 43° 30.151' N, 12° 35.533' E) crops out in the Umbria–Marche Apennines of Central Italy, along the road connecting the villages of Palcano and Moria, on the west side of Mount Petrano in the Marche Region (Figs. 1, 3). The section includes the uppermost part of the Bisciario Formation (Fm) and most of the Schlier Fm, which represent the lower–middle Miocene deposits of the Umbria–Marche succession (Capuano 2009).

The Umbria–Marche succession was deposited in a pelagic environment from the early Jurassic to the early Cretaceous, within an extensional tectonic regime linked to the Tethyan rifting phase, followed by hemipelagic deposition within a sector that connected the flexured foreland and the foredeep. Thus, during the late Cretaceous to early Oligocene, the calcareous

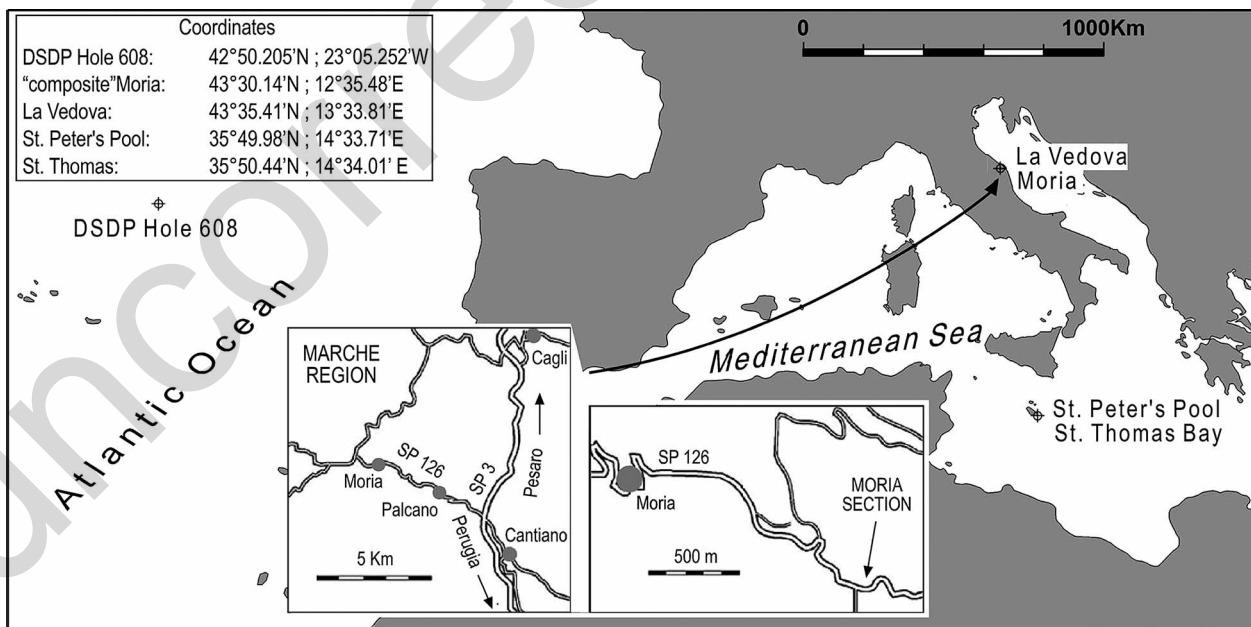


Fig. 1. Location map of the Moria section and of the other sections considered in the text.

Maiolica, Marne a Fucoidi, Scaglia Bianca, and Scaglia Rossa formations were deposited in the Umbria–Marche basin (Capuano 2009, Guerrera et al. 2012). After the late Oligocene, the complex geodynamic setting of the Apennine orogenesis was recorded in the deposition of the Scaglia Variegata and Scaglia Cinerea formations, and subsequently in Miocene sediments characterized by a progressive reduction of the carbonaceous fraction and the increase of fine clastic material (Guerrera et al. 2012), which gave rise to the Bisciario, Schlier and Marnoso-Arenacea formations.

The Bisciario Fm consists of marls and siliceous marly limestones that are finely layered, and gray and greenish-gray in color (ocher if altered), with a progressive upward increase of the terrigenous fraction. Blackish-gray cherty lists and nodules are locally present in the lower and middle part of the formation. The Bisciario Fm is characterized by frequent volcanic ashes, tuffites and bentonites resulting from syn-tectonic volcanic activity. The well-known “Raffaello Bed” (Coccioni and Montanari 1994) dated at about 22 Ma traditionally marks the Scaglia Cinerea/Bisciario lithostratigraphic boundary.

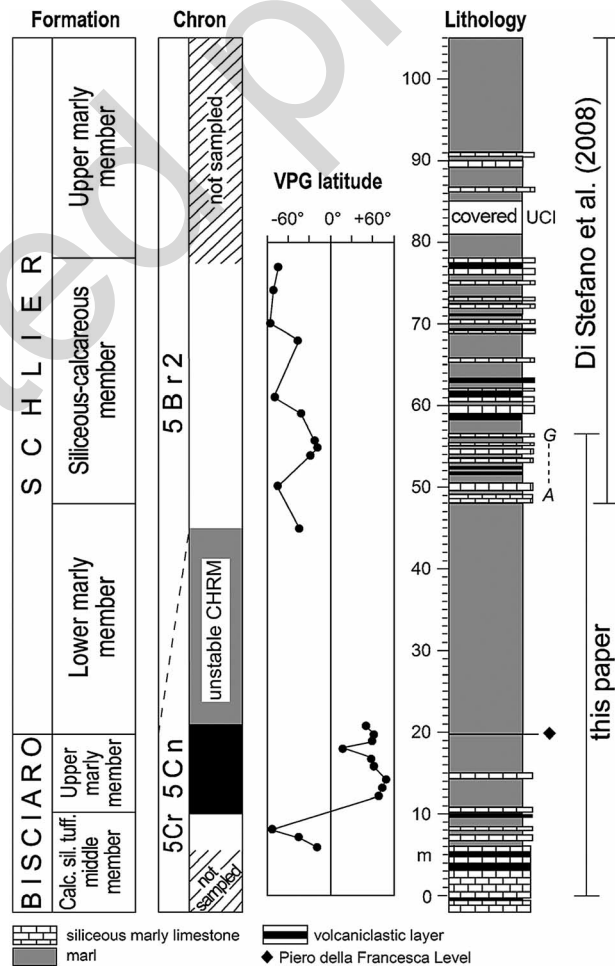
The Bisciario-Schlier transition is gradual and not always well defined. Coccioni and Montanari (1992) identified the boundary between the two formations in a 10–15 cm thick biotite-rich layer referred to as “Piero della Francesca Level”, radiometrically ( $^{40}\text{Ar}/^{39}\text{Ar}$ ) dated at  $17.1 \pm 0.16$  Ma (Deino et al. 1997). However, more recent interpretations place the boundary in correspondence with a significant reduction (or disappearance) of the calcareous and volcanoclastic layers (Capuano 2009, Guerrera et al. 2012).

The turbidite-type Schlier Fm is characterized by a sedimentation rate of ca. 40 m/Ma (Capuano 2009). Subdivided into a lower and an upper marly member, it consists mainly of marls and clayey marls. An intermediate member is composed of alternating marls/clayey marls and carbonate-rich layers. The upper boundary to the Marnoso Arenacea Fm is traditionally placed at the level of the first arenaceous horizon (Capuano 2009).

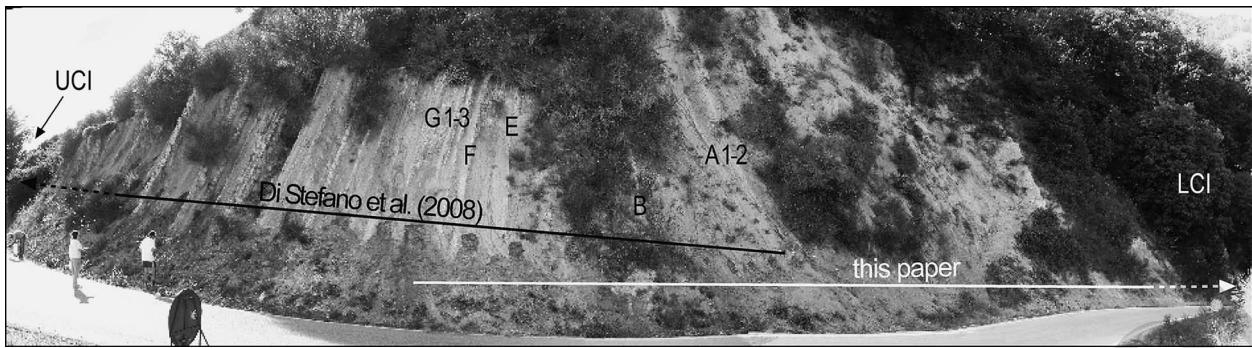
### 3. Previous studies and description of the Moria section

The stratigraphy of the Moria section has been described previously by Fornaciari et al. (1996), Fornaciari and Rio (1996), and Deino et al. (1997) (Figs. 2, 3). The lower part of the 100 m thick section consists

of few meters of indurated gray/beige limestones, a short interval of alternating soft marls and carbonates, and about 30 meters of gray/green marls. The upper part is represented by 40 m of alternating marls and clayey marls interlayered with whitish indurated calcareous layers, capped by marl. Deino et al. (1997) reported the “Piero della Francesca Level” at about 20 m above the base of the section. Biostratigraphically, the section falls within the planktonic foraminifer zones N6 (top) to N8 of Blow (1969) (mainly corresponding to the *G. trilobus* – *P. glomerosa* s.l. zones of Iaccarino and Salvatorini 1982 and Iaccarino 1985) and the calcareous nannofossil zones MNN3b–MNN5a (Fornaciari and Rio 1996, Fornaciari et al. 1996) straddling the Burdigalian-Langhian boundary.



**Fig. 2.** Lithostratigraphic log and paleomagnetic data of the Moria section, redrawn after Deino et al. (1997). The part considered in the present study and the part examined by Di Stefano et al. (2008) are indicated. The letters refer to the limestone layers as reported in Figs. 3 and 4. UCI = Upper Covered Interval.



**Fig. 3.** Panoramic view of the composite Moria section. The part considered in the present study and the one examined by Di Stefano et al. (2008) are indicated. LCI = Lower Covered Interval; UCI = Upper Covered Interval. The letters refer to the limestone layers as reported in Figs. 2 and 4.

The earliest magnetostratigraphic study could resolve magnetization directions in 23 samples (Deino et al. 1997) (Fig. 2), suggesting the presence of a 4-m-thick reversed polarity interval at the base of the section that was interpreted as Chron C5Cr based on the comparison with the Contessa section (Montanari et al. 1997). A normal polarity zone extending for about 11 meters follows, interpreted as Chron C5Cn (Deino et al. 1997). No reliable paleomagnetic data were acquired from the middle part of the section. An interval of reversed polarity was observed in the intermediate member of the Schlier Fm, interpreted to represent Chron C5Br2 (Deino et al. 1997). It must be noted that this interpretation is not consistent with the radiometric age of the “Piero della Francesca Level” reported by the same authors ( $17.1 \pm 0.16$  Ma), as the C5B/C5C boundary, which should be very close to it (Fig. 2), has an age of 15.97 Ma (Lourens et al. 2004).

A high-resolution calcareous plankton biostratigraphy of the upper part of the section (encompassing the uppermost layers of the lower marly, the siliceous-calcareous and the upper marly members of the Schlier Fm; Fig. 2) was recently published by Di Stefano et al. (2008). Unfortunately, no magnetostratigraphic age control could be provided for this interval. The calcareous plankton events recognized by Di Stefano et al. (2008) can now be chronologically constrained through the comparison with the subsequent study of Turco et al. (2011) who detected and dated the same bioevents at the La Vedova section, about 80 km east of the Moria section, along the Conero Riviera (Marche region, Central Italy; Fig. 1).

In this study, the lower 55 m of the Moria section have been re-sampled for bio-magnetostratigraphic analysis (Figs. 4, 5, 6), starting from the limestones of the Bisciario Fm and recognizing the “Piero della

Francesca Level”, 22 m from the base of the section. Minor differences with respect to the description reported by Deino et al. (1997), for instance the different position of the “Piero della Francesca” layer, are due to the presence of a 4-m-thick vegetation-covered interval in the basal part of the section (LCI in Fig. 3). The top of the studied section, characterized by thin calcareous layers within prevailing clays of the Schlier Fm, overlaps the basal part of the section studied by Di Stefano et al. (2008) (Figs. 2, 3).

## 4. Methodology

Sixty-nine samples were considered for qualitative and semiquantitative analyses of planktonic foraminifers. Samples were washed through a 63  $\mu\text{m}$  sieve. Some encrusted specimens of biostratigraphically significant taxa were cleaned by ultrasonic bathing to allow better observation of diagnostic characters. Sixty-eight smear slides for calcareous nannofossil analyses were prepared and studied using standard methods (Haq and Lohman 1976, Rio et al. 1990), including analysis under a 1000x magnification polarized microscope (transmitted light and crossed nicols). Following Rio et al. (1990) and Fornaciari et al. (1996), the relative abundances of all species within the genera *Sphenolithus* and *Helicosphaera* were determined by counting 30 and 50 specimens, respectively.

Sixty paleomagnetic cores were drilled using a petrol-powered portable drill cooled by water, and oriented in situ using a magnetic compass corrected for the local magnetic declination for the year 2009 (ca.  $2^\circ$ ). Afterwards, the cores were cut into standard paleomagnetic specimens of 22 mm height. Paleomagnetic measurements were performed at the Istituto Nazio-

nale di Geofisica e Vulcanologia (INGV Rome, Italy). Natural remanent magnetization (NRM) of the specimens was measured in a magnetically shielded room with a DC-SQUID cryogenic magnetometer (2G Enterprises, USA). Demagnetization diagrams were obtained through a stepwise (25°C increments) thermal cleaning treatment carried out in a magnetically shielded oven up to complete demagnetization of the samples. Demagnetization data were then plotted on orthogonal demagnetization diagrams (Zijderveld 1967), and the magnetization components were isolated by principal component analysis (Kirschvink 1980) according to standard paleomagnetic procedures.

## 5. Magnetostratigraphy

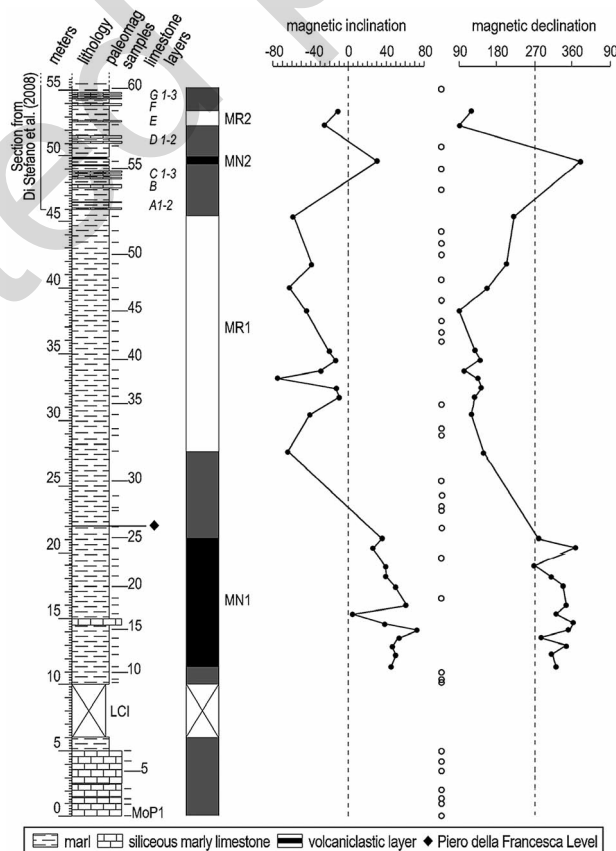
Magnetization of the analyzed samples is very weak and ranges between 17 and 293  $\mu\text{A}/\text{m}$  (mean value = 58  $\mu\text{A}/\text{m}$ ), denoting a very low concentration of the magnetic minerals. Being close to the instrumental sensitivity of the magnetometer ( $\sim 5 \mu\text{A}/\text{m}$ ), only 28 out of 60 samples yielded stable and interpretable demagnetization diagrams (Figs. 4, 5a). Most of the samples are characterized by two components of magnetization (Fig. 5a). The low-stability component, possibly resulting from a present-day geomagnetic field overprint, was removed at 100–120°C (denoting possible presence of goethite). The high-stability component was in general isolated between 120 and 300°C, pointing to iron sulphides (likely greigite) as the main magnetic carrier.

Demagnetization experiments were not continued above 300°C due to the occurrence of mineralogical transformation upon heating (i. e., iron sulphides transformed into magnetite) that resulted in a spurious increase of magnetic susceptibility and magnetization. Nevertheless, although this process inhibited complete removal of the high-stability components, in the demagnetization diagrams (Fig. 5a) they clearly decay towards the origin of the axes (typical feature of high-stability components of magnetization).

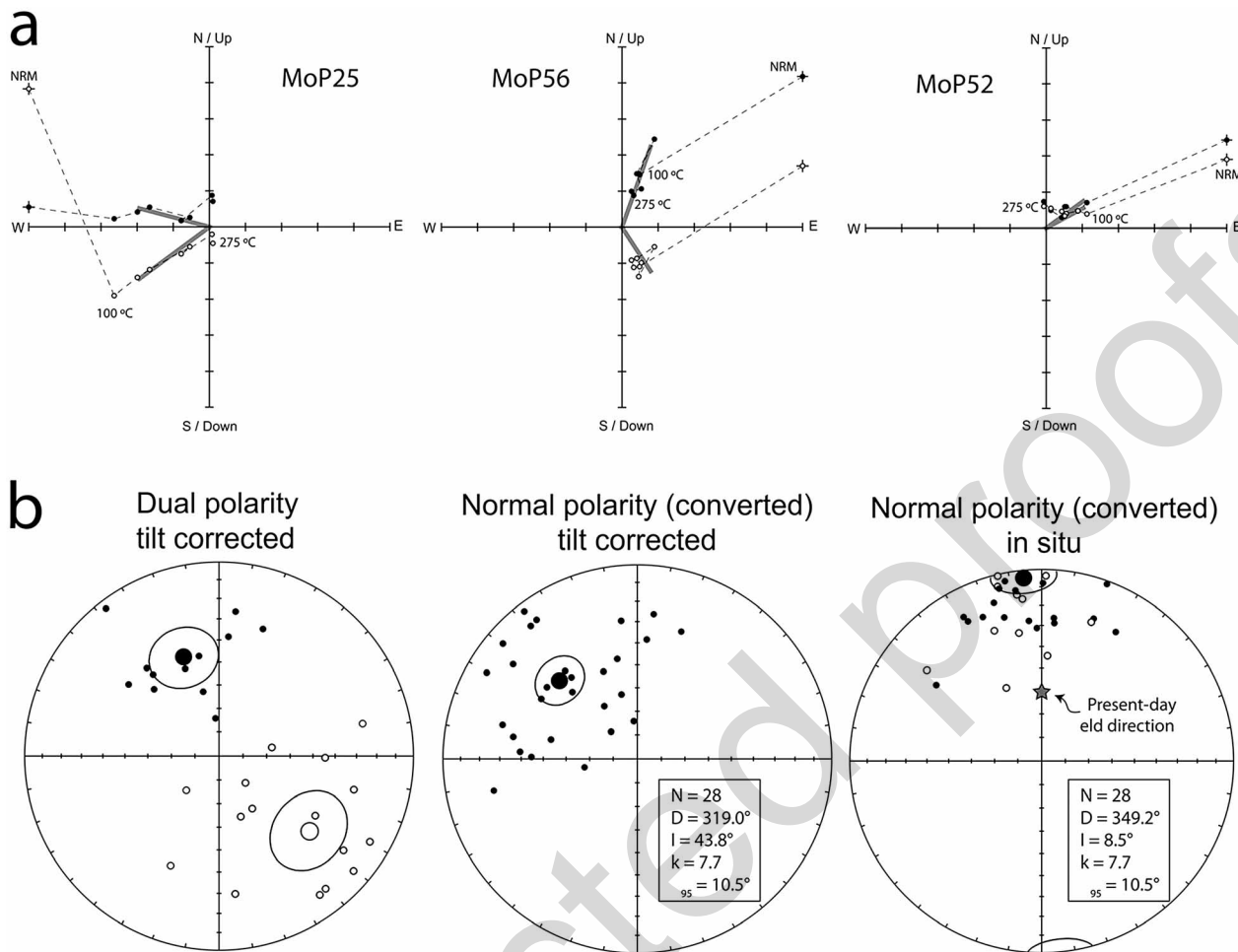
The most commonly used approaches to test the reliability of the isolated remanence directions in paleomagnetic studies (primary vs. secondary magnetization) are the reversal test (McFadden and McElhinny 1990), and the fold test (McFadden 1990). A positive result of the reversal test provides evidence that magnetic remanence was acquired over a longer time interval during which at least one geomagnetic reversal occurred, excluding in this way that a rapid remagne-

tization event occurred since the deposition of the sediment. Out of 28 isolated directions, 14 have a reversed polarity and 14 a normal polarity (Figs. 4, 5b). Their distribution is, at a first glance, somewhat antipodal, supporting the possibility of a positive reversal test. However, likely due to the low intensity of the magnetizations, which resulted into a larger scatter of the paleomagnetic directions, the performed reversal test provided an undetermined result (note this is not a negative reversal test). On the other hand, the entire section displays a quite homogenous bedding attitude of 214°/63° (dip direction/dip), implying that the statistical parameters associated with the distribution of the isolated high-stability components (i. e.,  $k$  and  $\alpha_{95}$ ) are identical before and after bedding correction, hence inhibiting any meaningful fold test.

When converted to normal polarity state, the isolated directions ( $N = 28$ ) after bedding correction display



**Fig. 4.** Lithological log of the succession investigated in the present study; sample positions are indicated on the right. The black, white and gray intervals in the polarity record represent normal, reversal and uncertain polarity intervals, respectively. The empty circles represent samples that were not interpreted. LCI = Lower Covered Interval.



**Fig. 5.** **a** – Zijdeveld diagrams (Zijdeveld 1967) of three representative samples showing normal (samples MoP25 and MoP56) and reverse (sample MoP52) polarity of magnetisation (tilt-corrected coordinates). Solid and open dots indicate projections on the horizontal and vertical planes, respectively. Demagnetization step values in °C are shown, together with the interval that was used to isolate the characteristic remanent magnetisation (thick grey line). **b** – Magnetic component analysis and graphic output were carried out with REMASOFT 3.0 software [Chadima and Hroudá, 2006].

a well-clustered distribution with declination ( $D$ ) =  $319.0^\circ$ , inclination ( $I$ ) =  $43.8^\circ$ , and statistical Fisherian (Fisher 1953) parameters  $k = 7.7$ , and  $\alpha_{95} = 10.5^\circ$  (Fig. 4). Before bedding correction (in situ), the mean value ( $D = 349.2^\circ$ ,  $I = 8.5^\circ$ ) is clearly different from the direction of the present-day geomagnetic field (Fig. 5b), hence a recent magnetic overprint can be firmly excluded. Furthermore, the computed mean direction for the Moria section is consistent with the moderate (few tens of degrees) counterclockwise rotations characterizing coeval rocks from adjacent sectors of the northern Apennines (Speranza et al. 1997). This provides strong evidence for the primary origin of the isolated directions. Accordingly, the obtained paleomagnetic data are reliable and will be used in the following magnetostratigraphic reconstruction.

The distribution of the magnetic polarities within the Moria section resulting from this study (Fig. 4, Table 1) indicates the presence of four magnetozones, from bottom to top: MN1 = normal polarity interval (samples MoP11–MoP25), MR1 = reversed polarity interval (samples MoP31–MoP53), MN2 = normal polarity interval (sample MoP56), and MR2 = reversed polarity interval (samples MoP58–MoP59).

## 6. Biostratigraphy

### 6.1 Planktonic foraminifers

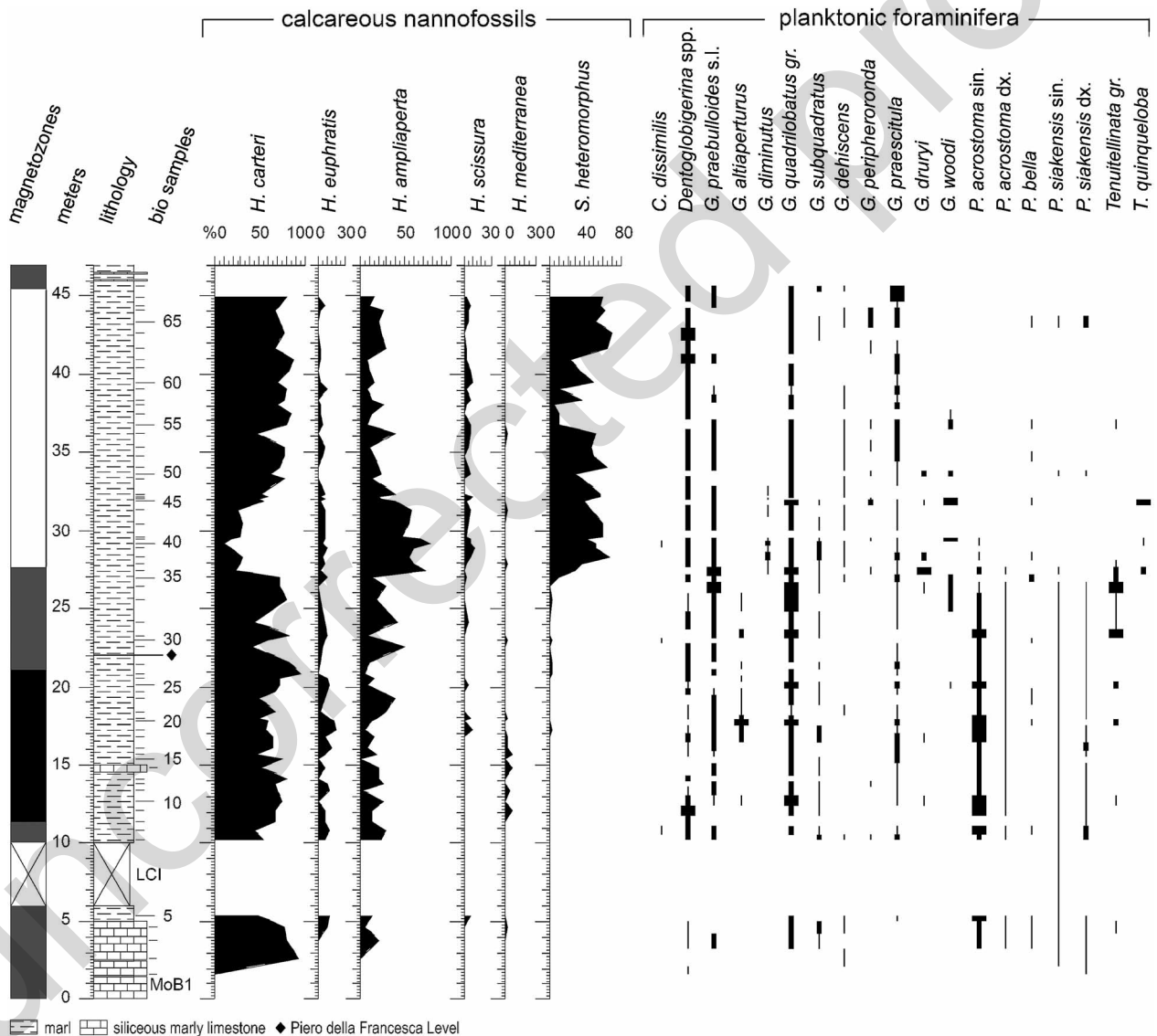
Planktonic foraminifers are generally abundant throughout the investigated part of the Moria section, but poorly to medium preserved in samples from the calcare-

ous strata of the Bisciario Fm. The poor preservation due to encrustation and carbonate dissolution increases the portion of dissolution-resistant taxa within the assemblages, thus altering the original abundance pattern. The preservation of planktonic foraminifers improves upwards in the marls of the Schlier Fm. However, recrystallized, broken and mechanically deformed tests are common throughout the section.

The assemblages are dominated by *Dentoglobigerina* spp., *Globigerina praebulloides*, *Globigerinoides quadrilobatus* group, *Globorotalia praescitula*, and *Paragloborotalia acrostoma* (Fig. 6). Less abundant taxa

include *Globigerinoides altiapertura*, *G. diminutus*, *G. subquadratus*, *Globoquadrina dehiscons*, *Globoturborotalita druryi*, *G. woodi*, and the *Tenuitellinata angustiumbilitata* group. Finally, *Catapsydrax dissimilis*, *Paragloborotalia bella*, *P. siakensis*, and *Turborotalita quinqueloba* show rare and scattered occurrences.

The generally poor preservation of the assemblages hampered the application of quantitative approaches, and only semiquantitative distributions of selected taxa are reported (Fig. 6). The ranges and occurrence characteristics of the most significant taxa are discussed in the following.



**Fig. 6.** Magnetostratigraphy and quantitative (calcareous nannofossil) and semi-quantitative (planktonic foraminifer) distribution patterns from the lower part of the Moria section. The black, white and gray areas indicate normal, reversal and uncertain polarity, respectively. LCI = Lower Covered Interval.



Table 1 Magnetozones recognized in the Moria section.

POLARITY	THICKNESS (m)	INTERVAL		MEAN (m)	ERROR +/- (m)	MAGNETO ZONE	
		Sample	Meters				
Und. interval	1.70						
Reverse	1.10	T	MoP59	53.50	54.35	0.85	MR2
		B	MoP58	52.40	51.00	1.40	
Und. interval	2.80						
Normal	0.85	T	MoP56	49.60	51.00	1.40	MN2
		B	MoP56	49.60	47.50	2.10	
Und. interval	4.20						
Reverse	17.80	T	MoP53	45.40	47.50	2.10	MR1
		B	MoP31	27.60	24.40	3.20	
Und. interval	6.40						
Normal	9.80	T	MoP25	21.20	24.40	3.20	MN1
		B	MoP11	11.40	5.90	5.50	
Und. interval	11.00						

### *Catapsydrax dissimilis*

The Last Occurrence (LO) of the genus *Catapsydrax* is considered a first-order bioevent and has been widely used for tropical open-ocean settings (Blow 1969, Berggren et al. 1995) and the Mediterranean region (Iaccarino 1985). The LO of *C. dissimilis* s.l. (Blow 1969, Kennett and Srinivasan 1983), or of *C. dissimilis/unicavus* according to Berggren et al. (1995), marks the top of the *Globigerinatella insueta/Catapsydrax dissimilis* zone (N6 and M3 biozones of Blow 1969 and Berggren et al. 1995, respectively).

In the examined samples, *Catapsydrax* is rare, and the poor preservation makes its distinction from *Globigerina venezuelana* and dentoglobigerinids sometimes difficult. The latter two taxa tend to form an aberrant small last chamber partially covering the umbilicus (Bolli and Saunders 1985), developing an element similar to the bulla in the genus *Catapsydrax*. *Catapsydrax dissimilis* is present up to sample MoB39 (29.0 m), close to the abundance increase of *Sphenolites heteromorphus* (Fig. 6).

### *Globigerinoides altiapertura*

*Globigerinoides altiapertura* has been used as a zonal marker in the lower Miocene of the Mediterranean region (Iaccarino 1985, Mancin et al. 2003, Iaccarino et al. 2007). Specimens with the typical high-arc apertures are present continuously in the middle part of the investigated section (samples MoB18–MoB31, 17.6–23.8 m). The absence of the taxon in the lower part is probably due to poor preservation. The highest presence of *G. altiapertura* in sample MoB33 (Fig. 6) may represent the true LO of the species, in agreement with Iaccarino (1985) who reports a late Burdigalian age for this event.

### *Globigerinoides diminutus*

*Globigerinoides diminutus* is morphologically similar to *G. ruber* and *G. subquadratus*, but has a smaller and more compact test as well as a higher and narrower aperture (Bolli and Saunders 1985). The species appears in the late Burdigalian N7 zone (Blow 1969, Postuma 1971, Salvatorini and Cita 1979, Kennett and Srinivasan 1983, Bolli and Saunders 1985, Iaccarino 1985).

According to Foresi et al. (2014), this species appears in the upper part of their *P. acrostoma/C. dissimilis* (MMi2c) subzone, which corresponds to Blow's (1969) N6 zone. The first occurrence (FO) of typical specimens is in sample MoB36 (28.0 m), just below the last common occurrence (LCO) of *P. acrostoma* and the LO of *C. dissimilis* (Fig. 6).

### *Globigerinoides quadrilobatus* group

The group comprises *Globigerinoides quadrilobatus*, *G. sacculifer* and *G. trilobus*, which Blow (1969) considered to be subspecies of *G. quadrilobatus*. Specimens of this group are common and continuously present throughout the investigated section (Fig. 6), with *G. trilobus* and *G. quadrilobatus* being dominant.

Typical *G. trilobus* are accompanied by small and medium-sized specimens with ovoidal, sub-spherical or bi-spherical outline of the test, showing some affinity with the *Globigerinoides-Praeorbulina* group (Turco et al. 2011b). There are consistently two apertures at the base of the last chamber, and the wall texture is cancellate. Unfortunately, the poor preservation often hampers the identification of the secondary apertures. No convincing specimens with three apertures at the base of the last chamber were found; this implies that *G. sicanus* is absent from the investigated samples, which is consistent with the FO of the species located at a stratigraphically higher level in the Moria section (Di Stefano et al. 2008).

### *Globigerinoides subquadratus*

*Globigerinoides subquadratus* is an important biostratigraphic marker for the Miocene. The FO of the taxon has been reported in the middle early Miocene (Blow 1969, Bizon and Bizon 1972, Kennett and Srinivasan 1983, Bolli and Saunders 1985, Iaccarino 1985, Spezzaferri 1994, Coccioni et al. 1997, Hakyemez and Toker 2010, Foresi et al. 2014). Typical specimens of *G. subquadratus*, with two supplementary apertures, occur from sample MoB3 (3.80 m) upsection, but this observation may be biased by diagenetic effects due to the poor preservation in the lower part of

the section (Fig. 6). The species shows a discontinuous distribution up to sample MoB41 (29.5 m), becoming further scattered upwards. The *G. subquadratus* concept includes very small specimens as they are mainly present in the basal part of the section; these specimens resemble *Globigerinoides diminutus*, from which they differ by the large secondary apertures, the incised intercameral sutures and the less box-shaped form of the test.

#### *Globorotalia peripheroronda*

The FO and LO of *G. peripheroronda* have been reported at the base of the Aquitanian (Kennett and Srinivasan 1983, Spezzaferri 1994) and in the Serravallian (Foresi et al. 2002, Hilgen et al. 2009), respectively. In the Mediterranean region, *G. peripheroronda* occurs from the late Burdigalian *Globigerinoides trilobus* zone onwards (Iaccarino 1985), but it is generally rare in the lower part of its range (Foresi et al. 2001, Lirer and Iaccarino 2005, Di Stefano et al. 2011b).

In the investigated samples, *G. peripheroronda* shows rare occurrences in two samples from the lower part of the succession, and rare to few discontinuous occurrences from sample MoB36 (27.4 m) upwards, together with *Turborotalita quinqueloba* (Fig. 6). This distribution pattern of the two taxa agrees with data from the St. Thomas section (Foresi et al. 2014), which crops out along the southeastern coast of Malta Island (Fig. 1).

#### *Globorotalia praescitula*

According to Iaccarino (1985), *G. praescitula* first occurs in the Burdigalian *G. trilobus* zone and becomes common in the Mediterranean region during the Langhian (Foresi et al. 2001). In the Moria section, the first specimens of *G. praescitula* were found in sample MoB5 (5.40 m). The species is rare to common and has a discontinuous occurrence upwards, which confirms the scattered distribution reported by Deino et al. (1997). Considering the poor preservation of foraminifers in the lower part of the section, it cannot be excluded that this species also occurs below sample MoB5.

#### *Paragloborotalia acrostoma*

The stratigraphic distribution of *P. acrostoma* is reported from the latest Oligocene to early Langhian by Spezzaferri (1994, 1996), but according to Foresi et al. (2001) the species is typical of the early-early middle Miocene. In the Moria section, the first specimens of *P. acrostoma* occur in sample MoB3 (3.80 m). The

species shows a prevailing sinistral coiling and is common to abundant up to sample MoB36 (27.4 m). Its LCO is close to the LO of *C. dissimilis*. These two bio-events basically occur at the same level in other Mediterranean sections as well, such as at Colle Amarena (Lirer et al. 2007) and Pianosa (Cornamusini et al. 2014). In the uppermost part of its distribution range, *P. acrostoma* is rare. Its highest occurrence is in sample MoB41 (29.5 m).

#### *Paragloborotalia siakensis*

The stratigraphic distribution of *P. siakensis* ranges from the Oligocene to the Tortonian; it has been studied in detail as it is considered extremely useful for correlation of middle Miocene sections on the basis of alternating acme and paracme intervals (Dall'Antonia et al. 2001, Sprovieri et al. 2002, Di Stefano et al. 2008, 2011b). *Paragloborotalia siakensis* occurs rarely, but continuously from the base of the Moria section up to sample MoB36 (27.4 m), whereas it is scattered from this level upsection. The species shows random coiling patterns along its entire range in the Moria section.

#### Other taxa

Other members of the planktonic foraminiferal assemblages are of secondary biostratigraphic interest. They include the following taxa:

*Dentoglobigerina* spp. and *Globigerina praebulloides* s.l., are common to abundant and occur continuously throughout the Moria section. *Dentoglobigerina* spp. comprise predominantly *D. altispira* and subordinately *D. baroemoenensis*. According to Blow (1959, 1969) and Blow and Banner (1966), the early Miocene *G. praebulloides* comprises several morphotypes and subspecies. In the samples from the Moria section, this taxon is referred to as *G. praebulloides* s.l., which comprises *G. praebulloides praebulloides*, *G. praebulloides oclusa*, and *G. aff. falconensis*.

*Paragloborotalia bella* is present in the Moria Section, but occurs rarely and discontinuously. The *Tenuitellinata* group comprises typical specimens of *T. angustiumbilitata* and other five-chambered globigerinids. Some of these resemble *G. ciperoensis* due to a wide umbilical area, but have a smooth rather than cancellate text surface. The *Tenuitellinata* group is discontinuously present in the lower part of the Moria section, where it is abundant from samples MoB31 to MoB36 (23.2–27.4 m), and virtually absent in the uppermost part.

*Globoturborotalita woodi* and *G. druryi* only occur in the middle part of the Moria section. *Globoquadri-*

*na dehiscens* is consistently rare and shows discontinuous occurrences throughout the section. Finally, *Turbotalita quinqueloba*, represented also by specimens showing the typical, slightly elongate last chamber covering the umbilicus, is present only in the interval from samples MoB35 to MoB45 (27.0–31.9 m). The comparison with the Foresi et al. (2014) St. Thomas section allows us to consider the lowest presence of the taxon in the Moria section as its stratigraphical FO.

## 6.2 Calcareous nannofossils

The calcareous nannofossil assemblages are characterized by poor preservation in the basal calcareous strata of the Bisciario Fm (Fig. 6). Preservation is medium to good in the remainder of the section. The nannofossil associations consist of placoliths, helicoliths and sphenoliths; discoasterids are rare, occur sporadically and are almost exclusively represented by *Discoaster deflandrei*. Within the placoliths, estimated through a qualitative analyses, the most representative are *Coccolithus pelagicus*, *Cyclicargolithus floridanus*, *Dicthyococcites* spp., and “small” and “medium” *Reticulofenestra*.

Following the most recent literature on Miocene nannofossil biostratigraphy in the Mediterranean region (Di Stefano et al. 2008, 2011b, Foresi et al. 2011, 2014, Iaccarino et al. 2011, Mourik et al. 2011, Turco et al. 2011a), helicoliths and sphenoliths were selected for quantitative analyses (Fig. 6) in order to determine positions of biostratigraphic horizons.

### *Helicosphaera carteri* and *H. euphratis*

Among the helicoliths, *H. carteri* is the most common (average percentages > 60%) taxon. It is continuously present in most of the analyzed samples, with the exception of the central part of the succession (28–32 m) where it is replaced by *H. ampliaperta*. *Helicosphaera euphratis* occurs in all samples, even if it only sporadically exceeds percentages of 10%. Fornaciari and Rio (1996) and more recently Backman et al. (2012) have highlighted the close inverse abundance relationship of *H. carteri* and *H. euphratis*. They used the sharp quantitative decrease of *H. euphratis* and the concomitant increase of *H. carteri* (reported as “*H. euphratis/H. carteri* cross-over”) to define the boundary between the MNN1/MNN2a (Fornaciari and Rio 1996) and the CNM3/CNM4 (Backman et al. 2012) zones in the late Aquitanian. The high abundance of *H. carteri* together with the low abundance of *H. euphratis* suggests that the base of the section is younger

than the *H. euphratis/H. carteri* cross-over dated at 20.9 Ma (Backman et al. 2012).

### *Helicosphaera ampliaperta* and *H. scissura*

*Helicosphaera ampliaperta* is almost continuously present across the Moria section, becoming abundant between 28 and 30 meters (samples MoB36 to MoB44) where it accounts for more than 50% of the total helicoliths. The distribution of *H. ampliaperta* spans from the Aquitanian (Young 1998) to the middle Langhian (Di Stefano et al. 2008). In Lourens et al. (2004), its FO is adopted to provisionally approximate the Aquitanian/Burdigalian boundary at 20.43 Ma. The basal part of its range is discontinuous and characterized by temporary absence intervals (Gartner 1992, Fornaciari and Rio 1996, Foresi et al. 2014). The LCO of the species appears sharp and synchronous, while its LO seems to be diachronous among different sections in the Mediterranean region (Di Stefano 1995, Fornaciari et al. 1996, Di Stefano et al. 2008, 2011b, Foresi et al. 2011, Iaccarino et al. 2011, Turco et al. 2011a).

Data from the Central Paratethys suggest that *H. ampliaperta* may have evolved from *H. scissura*, whose FO is recorded in the early Aquitanian (Marunteanu 1999, Rögl and Nagymarosy 2004, Garecka 2005, Bartol 2009, Makrodimitras et al. 2010, Chira et al. 2011, Holkova 2013). For open-ocean settings, the FO of *H. scissura* is reported in the middle Burdigalian, at about 18.28 Ma, thus following that of *H. ampliaperta* (Young 1998). In Mediterranean sequences the two species generally occur together, and it is difficult to establish which one represents the ancestor of the other.

In the Moria samples, *H. scissura* exhibits low abundances (< 10%) and a discontinuous distribution, but seems to match the distribution pattern of *H. ampliaperta*. *Helicosphaera scissura* may hence represent a morphotype within the *H. ampliaperta* population.

### *Helicosphaera mediterranea*

*Helicosphaera mediterranea* is a lower–middle Miocene species, spanning from the early Aquitanian to the late Langhian (Theodoridis 1984). Marunteanu (1999) recognizes a basal Miocene age (NN1 zone of Martini 1971) for the FO of this taxon in the Carpathian region, in contrast to the early Burdigalian age (MNN2a zone) proposed by Fornaciari and Rio (1996) for the Mediterranean region. Furthermore, in some Mediterranean sections, the FO of *H. mediterranea* precedes that of *H. ampliaperta*, whereas in others this event seems to be younger (Fornaciari and Rio 1996). Thus, the relation between *H. mediterranea* and *H. ampliaperta* is still

controversial. However, the Moria section, where the taxon shows a low and discontinuous occurrence from the basal levels onwards, is not suitable to solve this conundrum.

*Sphenolithus heteromorphus*

*Sphenolithus heteromorphus* is absent or rare in the lower part of the investigated succession. Sporadic specimens are recorded in a few samples from about 18 to 28 meters (Fig. 6). From this level upwards, the species becomes continuously present, with average percentages of about 30% and a maximum of about 70% with regard to the total sphenolith assemblages.

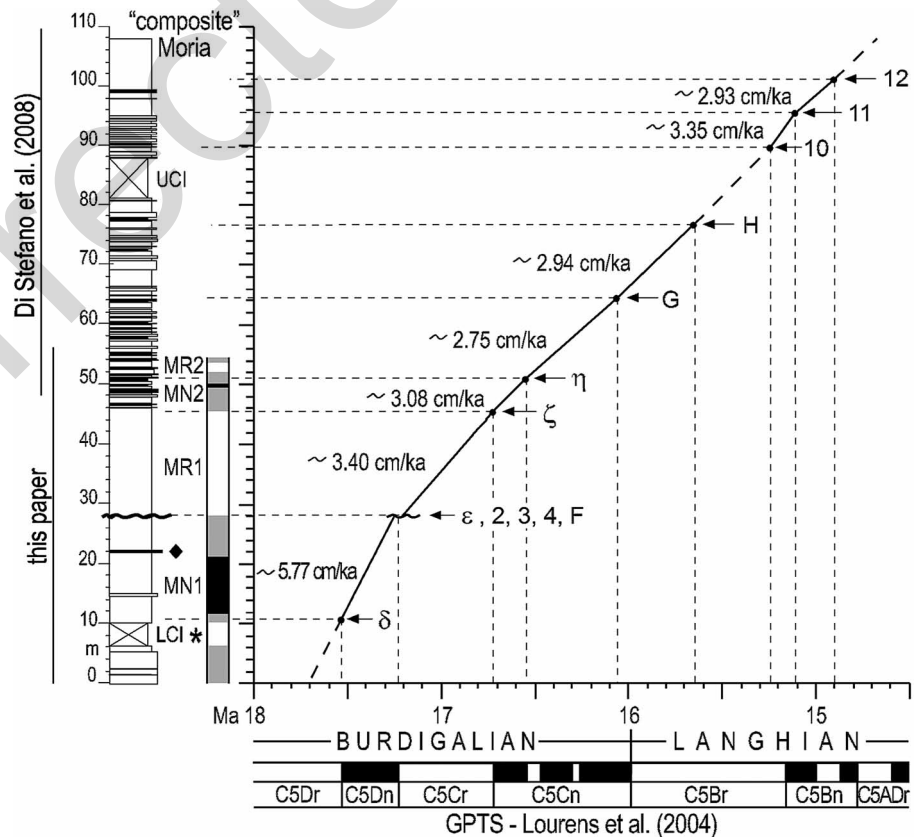
The FO of *S.heteromorphus* has been used by Okada and Bukry (1980) to define the base of their CN3 zone and by Theodoridis (1984) to recognize the boundary between its *Sphenolithus belemnos* subzone and *Helicosphaera ampliapertura* zone. Moreover, Fornaciari and Rio (1996) and Backman et al. (2012) used the FCO of the taxon to define the base of their MNN3b and CNM6 zones, respectively. In its initial range *S.heteromorphus* is generally accompanied by discontinuous, low-percentage occurrences of *S.belemnos*, whose LCO is below the FO of *S.heteromor-*

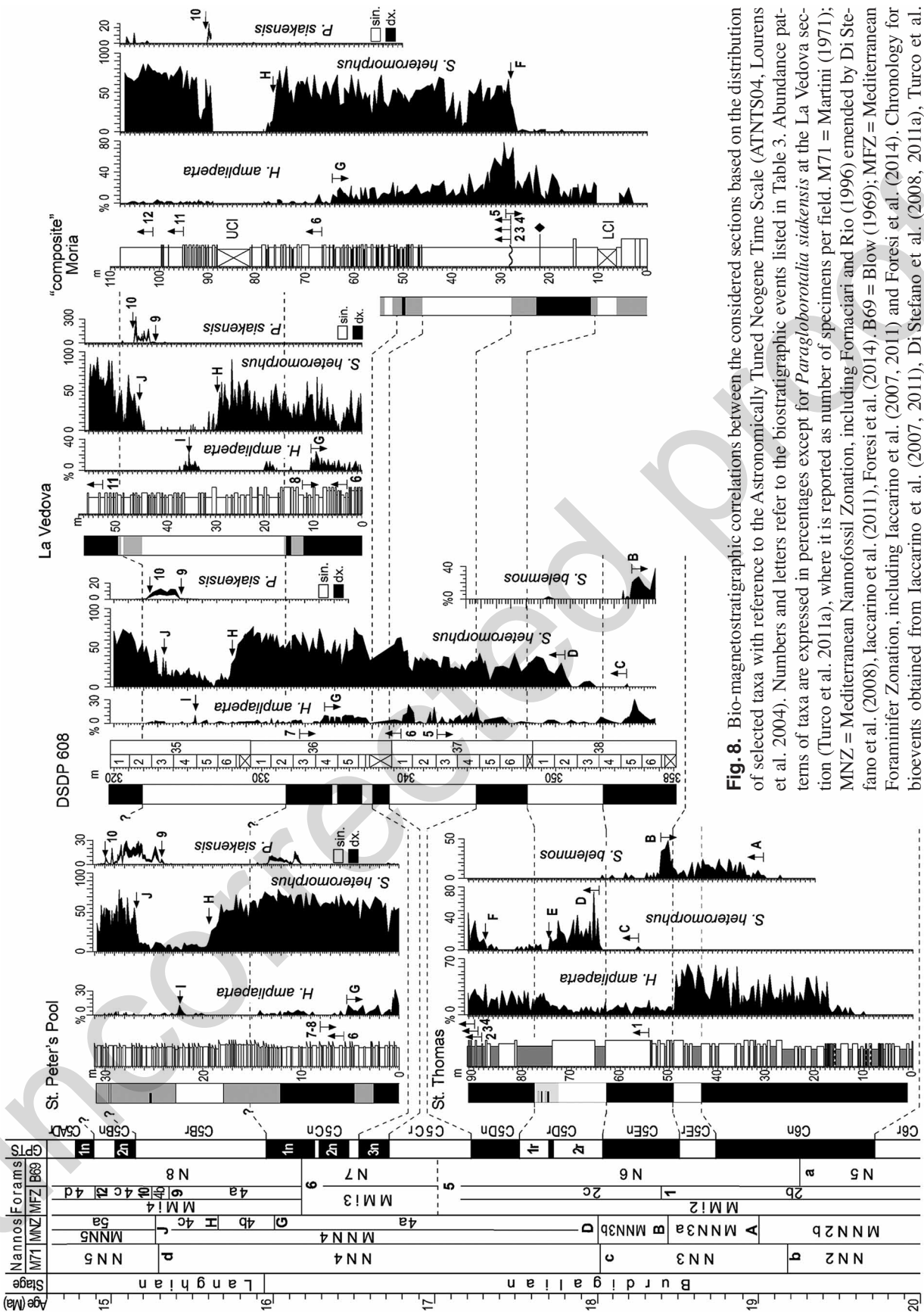
*phus* (Fornaciari and Rio 1996, Raffi et al. 2006). In the Moria section, *S.belemnos* is absent, which may suggest that the first specimens of *S.heteromorphus* recorded at the base of the section do not represent its true biostratigraphical FO.

### 7. Magnetostratigraphic interpretation and age model

The age-model herein proposed refers to the entire Moria Section (sensu Deino et al. 1997; composite Moria Section in Figs. 1, 7, and 8) and has been obtained through the integration of the data from the present study and those of Di Stefano et al. (2008). The two portions of the section were correlated using the calcareous marker horizons in the middle member of the Schlier Fm (Figs. 2, 3). This allows to define biostratigraphic events for the entire succession and to extend the age-model upsection where paleomagnetic data are absent. The data from the entire Moria section have been compared to data from other Mediterranean sections and from North Atlantic DSDP Site 608 (Figs. 1, 8; Tables 2, 3).

**Fig. 7.** Age-depth diagram and estimated sedimentation rates for the composite Moria section. Numbers and greek letters associated with arrows refer to the bio- and magnetostratigraphic events listed in Table 3; (\*) in Deino et al. 1997. The part considered in the present study and the part examined by Di Stefano et al. (2008) are indicated. LCI = Lower Covered Interval; UCI = Upper Covered Interval.





**Fig. 8.** Bio-magnetostratigraphic correlations between the considered sections based on the distribution of selected taxa with reference to the Astronomically Tuned Neogene Time Scale (ATNTS04, Lourens et al. 2004). Numbers and letters refer to the biostratigraphic events listed in Table 3. Abundance patterns of taxa are expressed in percentages except for *Paragloborotalia stakensis* at the La Vedova section (Turco et al. 2011a), where it is reported as number of specimens per field. M71 = Martini (1971); MNZ = Mediterranean Nannofossil Zonation, including Fornaciari and Rio (1996) emended by Di Stefano et al. (2008), Iaccarino et al. (2011), Foresi et al. (2007, 2011); MFZ = Mediterranean Foraminifer Zonation, including Iaccarino et al. (2007, 2011) and Foresi et al. (2008, 2011a), Turco et al. (2011b), and Foresi et al. (2014). LCI = Lower Covered Interval; UCI = Upper Covered Interval.



Chron C5Dn (17.533–17.235 Ma, Lourens et al. 2004) (Figs. 7, 8).

In the Moria section (this study), the FOs of *Globigerinoides diminutus*, *Turborotalita quinqueloba* and *Globorotalia peripheroronda*, and the end of the lower paracme of *Spennolithus heteromorphus* (P<sub>0</sub>E) (see events 2, 3, 4 and F, respectively, in Table 3) occur at the same stratigraphic level (28 m), close to the base of the MR1 magnetozone. In the St. Thomas section, events 2, 3 and 4 occur at different stratigraphic levels in upper Chron C5Dn. There, the *S. heteromorphus* P<sub>0</sub>E also occurs within Chron C5Dn, slightly preceding the abovementioned foraminiferal events. This comparison allows to draw the following conclusions regarding the Moria section: i) the interval characterized by absence of paleomagnetic signals from 23 to 28 m has been deposited during Chron C5Dn time; ii) MR1 can be referred to Chron C5Cr; iii) a short hiatus involving the C5Dn/C5Cr boundary is present at about 28 m; this can be estimated to have lasted at least 34 ka, which represents the difference of the ages between the *S. heteromorphus* P<sub>0</sub>E and the FO of *G. peripheroronda* as determined by Foresi et al. (2014), as 17.311 Ma and 17.277 Ma, respectively. Consequently, MN2 has been interpreted as Chron C5Cn.3n, MR2 as Chron C5Cn.2r, and the reverse magnetozone recognized by Deino et al. (1997) at the base of the section (Fig. 2) as Chron C5Dr.

For the Moria section, the age model presented in Fig. 7 takes into account the bio-magnetostratigraphic data of the present study and the position of the bioevents determined by Di Stefano et al. (2008), with ages derived from the neighbour La Vedova section (Table 3, Iaccarino et al. 2011, Turco et al. 2011a) and, for the lower portion, from the Maltese St. Thomas section (Table 3, Foresi et al. 2014). From 28 m upwards, the distribution of the bio-magnetostratigraphic events suggests near-constant sedimentation rates, with an average of about 3 cm/ka. In the lower part, the sedimentation rate slightly exceeds 5 cm/ka, in spite of the presence of a short hiatus. The average sedimentation rate, calculated from the top of the lower vegetation-covered interval (10 meters above the base of the section and marked as LCI in Figs. 3, 7) to the top of the composite section (ca. 108 meters from the base; Fig. 7) is 3.85 cm/ka, which is in close agreement with the sedimentation rate of ca. 4.0 cm/ka reported by Capuano (2009) for the Schlier Fm in the Urbino area (northern sector of the Marche region).

The proposed age model for the composite Moria section suggests deposition between about 17.7 and

14.7 Ma, corresponding to a late Burdigalian–Langhian age (Lourens et al. 2004); this is in agreement with the radiometric age of the “Piero della Francesca Level” reported by Deino et al. (1997). From a biostratigraphic perspective, the Moria section represents the interval between nannofossil biozones MNN4a and MNN5a, and between planktonic foraminifer biozones MMi2c and MMi4d (Di Stefano et al. 2008, Iaccarino et al. 2011, Foresi et al. 2014).

## 8. Mediterranean-Atlantic correlation

The lower part of the Moria section (sensu Deino et al. 1997) here investigated in detail is instrumental for generating a bio-magnetostratigraphy of the early to middle Miocene interval of the Mediterranean region. To achieve this goal, the composite Moria section has been correlated with other Mediterranean successions (Figs. 1, 8; Tables 2, 3). A series of magnetostratigraphically constrained calcareous plankton bioevents has been established that covers the time interval from about 19.7 Ma (base of the St. Thomas section) to about 14.7 Ma (top of the composite Moria section). Furthermore, the comparison of the results of Mediterranean sections to a reference succession in the Atlantic Ocean (DSDP Site 608) (Miller et al. 1991, Olafsson 1991, Gartner 1992, Krijgsman and Kent 2004, Di Stefano et al. 2011a, Wade et al. 2011) allowed us to discriminate between the events with a both Mediterranean and extra-Mediterranean biostratigraphical significance and those with more regional importance.

Besides the well-known, yet debated bioevents (such as the LCO and LO of *H. ampliaptera*) horizons based on abundance fluctuations of selected species merit special attention. These biohorizons, although seemingly less trustworthy than proper FOs and LOs, provide useful and reliable biostratigraphic constraints for the considered time interval. The best-known example is probably the *Spennolithus heteromorphus* “Paracme Interval” (Di Stefano 1995, Fornaciari et al. 1996) (H and J in Fig. 7 and Table 3), which is recorded almost synchronously in numerous Mediterranean sections (Iaccarino et al. 2011) and at Site DSDP 608 (Di Stefano et al. 2011a). In the same way, the *Paragloborotalia siakensis* “Acme a Interval” (Dall’Antonia et al. 2001, Di Stefano et al. 2008) (events 9 and 10 in Fig. 8 and Table 3) is a sharp interval clearly detectable both in the Mediterranean and North Atlantic

successions. Conversely, the older *S.heteromorphus* paracme (“Paracme 0” in Foresi et al. 2014) detected in the Mediterranean region at the top of the St. Thomas section (events E and F in Fig. 8 and Table 3) and at the base of the Moria Section, but missing in the Atlantic DSDP 608 succession, may be only of regional value as correlation tool. Similarly, other abundance peaks, such as the one of *P.siakensis* recognized in the intermediate part of the St. Peter’s Pool section (“Acme<sub>0</sub>” in Foresi et al. 2011, absent in the same levels of the La Vedova section of Turco et al. 2011), which crops out along the southeastern coast of Malta Island (Fig. 1) can also be considered of regional value and are possibly related to regional environmental conditions (Iaccarino et al. 2011).

The comparison between the St. Thomas section and the basal part of DSDP Site 608 clarifies the relationship between the final distribution range of *S.belemnos* (above its LCO) and the FO of *S.heteromorphus* in the frame of tight magnetostratigraphic control. The LCO of *S.belemnos* represents a well-defined event and is recorded within Chron C5En both in the St. Thomas section and at DSDP Site 608, while the LO of the species occurs at higher stratigraphic levels within Chron C5Dr. The stratigraphic positions of the FO and FCO of *S.heteromorphus* have been repeatedly debated in the literature (Martini 1971, Okada and Bukry 1980, Olafsson 1991, Gartner 1992, Fornaciari and Rio 1996, Backman et al. 2012). At DSDP Site 608, the FO of *S.heteromorphus* occurs between the LCO and the LO of *S.belemnos*, within Chron C5En. This is in good agreement with the St. Thomas section, where the FO of *S.heteromorphus* occurs in the same chron, within the final distribution range of *S.belemnos*. The FCO of *S.heteromorphus* occurs within Chron C5Dr in both the sections.

The LO of *C.dissimilis* is another discussed bio-event, reported to be within Chron C5Dn by Berggren et al. (1983), at 17.5 Ma by Shackleton et al. (1999) (C5Dr/C5Dn transition in Figs. 2 and 3 of Wade et al. 2011), and within Chron C5Cr by Di Stefano et al. (2011a). The occurrence of the last *C.dissimilis* specimens at the base of Chron C5Cr in the Moria section agrees with its position at DSDP Site 608 (Di Stefano et al. 2011a). Yet, this event does not seem to represent a reliable correlation tool within the Mediterranean area because in some cases, for example the St. Thomas section, the species is completely absent, probably due to environmental factors (Foresi et al. 2014).

## 9. Conclusions

Results are presented from an integrated bio-magnetostratigraphic study of the lower segment of the Moria section (sensu Deino et al. 1997), cropping out in the Umbria–Marche Apennine (Central Italy). According to our reconstruction, this portion of the section ranges from the top of Chron C5Dr to Chron C5Cn.2r, almost continuously spanning from about 17.7 to 16.4 Ma (according to the ATNTS04), with a small hiatus, estimated in at least 34 ka, at the C5Dn/C5Cr transition. This interpretation differs from that of Deino et al. (1997), but is consistent with the radiometric age of  $17.1 \pm 0.16$  Ma reported by the same authors for the “Piero della Francesca Level” in the section.

An age model is proposed for the composite Moria Section, based on the bio-magnetostratigraphic data of the present study and the biostratigraphic results of Di Stefano et al. (2008). It is chronologically constrained through a comparison with the La Vedova section (Turco et al. 2011). According to our reconstruction, the composite Moria section covers about 3 Ma (17.7–14.7 Ma), thus straddling the Burdigalian/Langhian boundary (Lourens et al. 2004, Iaccarino et al. 2011).

Calcareous plankton data from the composite Moria section are compared with previously published datasets from other Mediterranean sections. This yields a magnetostratigraphically framed calcareous plankton dataset that covers a time interval of about 5 Ma from the base of the St. Thomas section at 19.7 Ma to the top of the composite Moria Section at 14.7 Ma.

The presence/absence of biostratigraphical events as deduced from a comparison of datasets from the Mediterranean region and North Atlantic DSDP Site 608 enabled us to discriminate between bioevents exportable in extra-Mediterranean areas from those showing a bioprovincial or local stratigraphic significance.

**Acknowledgements.** Fabrizio Lirer and Federica Zangara are thanked for their help during the sampling. We are indebted to three anonymous reviewers for their constructive suggestions. The study has been conducted within the project “PRIN2012” entitled “The Burdigalian GSSP (Global Stratigraphic Section and Point): the missing tile to complete the Neogene interval of the Geological Time scale”, financed by the Italian MIUR and coordinated by Agata Di Stefano.



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