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Research article

## The Eocene–Oligocene transition in the C-isotope record of the carbonate successions in the Central Mediterranean

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## ABSTRACT

The Eocene–Oligocene transition marks a fundamental step in the evolution of the modern climate. This climate change and the consequent major oceanic reorganisation affected the global carbon cycle, whose dynamics across this crucial interval are far from being clearly understood. In this work, the upper Eocene to lower Oligocene  $\delta^{13}\text{C}_{\text{carb}}$  and  $\delta^{13}\text{C}_{\text{TOC}}$  records of a shallow-water and a hemipelagic carbonate settings within the Central Mediterranean area have been studied and discussed. The shallow-water carbon isotope signal has been analysed in the northern portion of the Apula Platform, cropping out in the Majella Mountain, Central Apennines (Santo Spirito Formation). A coeval Umbria–Marche basinal succession has been investigated in the Massignano section (Conero area, Central Italy). The purposes of this work are: to discriminate between the global and the local (Mediterranean) signature of C-isotope record during the Oi-1 event, to correlate the regional C-isotope signal with the global record, and to evaluate the carbon cycle dynamics across the greenhouse–icehouse transition through the integration of complementary records (shallow-water vs pelagic settings,  $\delta^{13}\text{C}_{\text{carb}}$  vs  $\delta^{13}\text{C}_{\text{TOC}}$ ). The upper Eocene carbon isotope record of the analysed successions matches with the global signal. The overall trend shows a decrease of the  $\delta^{13}\text{C}_{\text{carb}}$  and a contemporary increase of the  $\delta^{13}\text{C}_{\text{TOC}}$ . The decoupling of the two curves is consistent with a reduced fractionation effect by primary producers that characterised the interval between the Middle Eocene Climatic Optimum and the onset of the Oi-1 event. However, regional factors superimposed the global signal. In fact, the upper Eocene basinal  $\delta^{13}\text{C}_{\text{TOC}}$  record is marked by short-term negative spikes, which possibly represent times of higher productivity triggered by the westward subtropical Eocene Neotethys current entering from the Arabian–Eurasian gateway. On the contrary, the shallow-water record does not display these short-term productivity pulses. A change in the carbonate factory is only recorded at the Eocene–Oligocene transition, marked by a reduction of the larger benthic foraminifera and the spread of seagrass and corals. Moreover, in the shallow-water record of the Santo Spirito Formation, no major carbon isotope shift related to the Oi-1 event is recorded due to the presence of extensive slumps that disrupt the bedding. These slumps are the main evidence of the sea-level drop that occurred concomitantly with the onset of the Antarctica ice-sheet, which caused the deepening of the storm wave base and increased the instability over the entire ramp.

## 1. Introduction

The Eocene–Oligocene transition (EOT) represents the most important step in the evolution of the modern icehouse climate. It marks the transition from the warm ‘greenhouse world’ when the atmospheric  $\text{pCO}_2$  was higher than 1000 ppm (Beerling and Royer, 2011; Pagani et al., 2014) and only transient ice-sheets could develop at low latitudes (Carter et al., 2017), to the modern ‘icehouse’ climate that is characterised by stable polar ice caps and lower  $\text{CO}_2$  concentrations (Pearson et al., 2009; Beerling and Royer, 2011). The EOT (Houben

et al., 2012) culminates with a  $\delta^{18}\text{O}$  peak recorded in the lower Oligocene (33.55 Ma) deep-sea successions (Miller et al., 1991). This positive oxygen isotope shift, named Oi-1 event (Miller et al., 1991; Zachos et al., 1996, 2001; Coxall and Pearson, 2007; Lear et al., 2008), marks the development of a permanent ice sheet in Antarctica as much as 50% in volume of the present one (DeConto and Pollard, 2003; Liu et al., 2009; Miller et al., 2009; Bohaty et al., 2012). Several studies testify for a very dynamic and unstable polar ice cap in Antarctica from the early Oligocene to the late Miocene (Pekar and DeConto, 2006; Cook et al., 2013; Liebrand et al., 2017; Sangiorgi et al., 2018). Pekar and

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Deconto (2006) report different records showing that the ice-volume fluctuated between 50% and 125% of the present Eastern Antarctic ice sheet, while Liebrand et al. (2017) state a minimum waxing and waning of 85% to 110% of the present-day volume. However, it is overall accepted that an ice sheet persisted in Antarctica since the Oi-1 event.

Both the causes and the effects of the glaciation in Antarctica are still under debate. Overall, the main theories approaching this problem rely on two different hypotheses e.g. the isolation of Antarctica due to plate reorganisation, or the decrease of the atmospheric CO<sub>2</sub> during late Eocene. The opening of the Tasmanian and Drake Passages during late Eocene was assumed as the main triggering cause of the glaciation since it led to the development of the Antarctic Circumpolar Current and therefore the thermal isolation of Antarctica (Kennet et al., 1975; Kennet, 1977). However, more recently, the decrease of pCO<sub>2</sub> is considered the main controlling factor on the ice-growth in Antarctica (DeConto and Pollard, 2003; Pagani et al., 2011; Egan et al., 2013; Goldner et al., 2014; Galeotti et al., 2016). Regardless of the debate on the triggering causes, both the opening of the Drake Passage (Scher and Martin, 2004) and the onset of an Antarctic ice sheet (Goldner et al., 2014; Wright et al., 2018) altered the ocean circulation resulting in a general global ocean reorganisation that affected the carbon cycle, as marked by a positive  $\delta^{13}\text{C}$  isotope excursion recorded in the deep-sea successions (Zachos et al., 1996, 2001; Cramer et al., 2009) associated with a major deepening of the Calcite Compensation Depth (CCD) (Coxall et al., 2005). Different hypotheses have been suggested to explain this positive carbon isotope excursion, most of which are related to changes in the oceanic trophic regime and the associated primary productivity induced by sea level changes or changes in the global oceanography (Salamy and Zachos, 1999; Zachos and Kump, 2005; Coxall et al., 2005; Coxall and Pearson, 2007; Miller et al., 2009; Coxall and Wilson, 2011; Plancq et al., 2014).

The carbon cycle perturbation related to the Oi-1 event has been clearly identified in the deep-sea record of all the oceans at different latitudes (Zachos et al., 1996, 2001; Cramer et al., 2009), whereas little is known about the response of the shallow-water carbonate systems' to this carbon cycle perturbation (Jaramillo-Vogel et al., 2013, 2016). Notwithstanding that, even if often discontinuous, shallow-water carbonates are significantly more sensitive to temperature and trophic changes than deep-sea successions. In fact, during the transition from the warm Eocene to the Oligocene 'icehouse', these major changes in climate, ice volume, and ocean circulation deeply influenced the composition and production of the carbonate factory (Nebelsick et al., 2005; Brandano et al., 2009, 2017a; Pomar et al., 2017).

The larger benthic foraminifera, which dominated the Eocene carbonate platforms (Racey, 2001, and references therein; Bassi, 2005; Beavington-Penney et al., 2005), experienced a major decline. In turn, zooxanthellate corals and coralline algae spread as main biota-producing sediment, and seagrass environments expanded strongly in the euphotic zone, influencing the facies association of the Cenozoic carbonate platforms (Nebelsick et al., 2005; Pomar and Kendall, 2008; Brandano et al., 2017a).

In this work, the carbon isotope record is studied in the EOT interval in two different contexts from the Central Mediterranean area. The first case study refers to sedimentary succession from the Apula Carbonate Platform (Central Apennines, Central Italy), in which the main lithofacies of the Santo Spirito Formation and  $\delta^{13}\text{C}$  of the whole-rock signal are analysed and discussed. This shallow-water carbon isotope signal is correlated with the  $\delta^{13}\text{C}$  record obtained in the Massignano section (Scaglia Variegata and Scaglia Cinera Fm, Conero area, Central Italy), which is the well-known Global Stratotype Section and Point (GSSP) of the Eocene-Oligocene boundary (Premoli Silva and Jenkins, 1993), and represents the Umbria-Marche basinal succession.

Through a precise correlation of the C-isotope records of the Mediterranean area with the global signal, we aim (i) at assessing the impact of C-cycle perturbations on shallow- and deep-water successions, and (ii) at evaluating the relationships between carbon cycle and

shallow carbonates across this major greenhouse-icehouse transition.

## 2. Geological setting

The Apennine fold-and-thrust belt is the result of the collision of the Adria plate with the southern margin of Europe (Alvarez et al., 1974; Bally et al., 1986; Dewey et al., 1989; Doglioni, 1991; Rosenbaum et al., 2002a; Devoti et al., 2008; Carminati et al., 2012). The Apennine subduction started in the late Eocene (Lustrino et al., 2009), and it continued through Neogene until today (Boccaletti et al., 1990; Rosenbaum et al., 2002b; Doglioni et al., 1998; Carminati et al., 2010, 2012). The westward-dipping Apennine slab led to an eastward migration of the deformation fronts and related foredeeps, and subsequent extensional tectonics in the back-arc area where several basins opened during the Oligocene-Miocene interval on thin continental (the València Through) and oceanic (Alborán, Provençal and Thyrrenian Basin) crust (Carminati et al., 2012 and references therein). The Central Apennine consists of Triassic to Miocene deposits that can be ascribed to three different paleogeographic domains: the Apennine carbonate platforms (Latium-Abruzzi and Apula platforms), the Umbria-Marche basin, and the Molisano basin (Colacicchi, 1967; Crescenti et al., 1969; Parotto and Praturlon, 1975; Bernoulli, 2001; Vezzani et al., 2010). In this study, the northern extension of the Apula platform, represented by the Majella Mountain and the Marche basinal successions, are investigated (Fig. 1a).

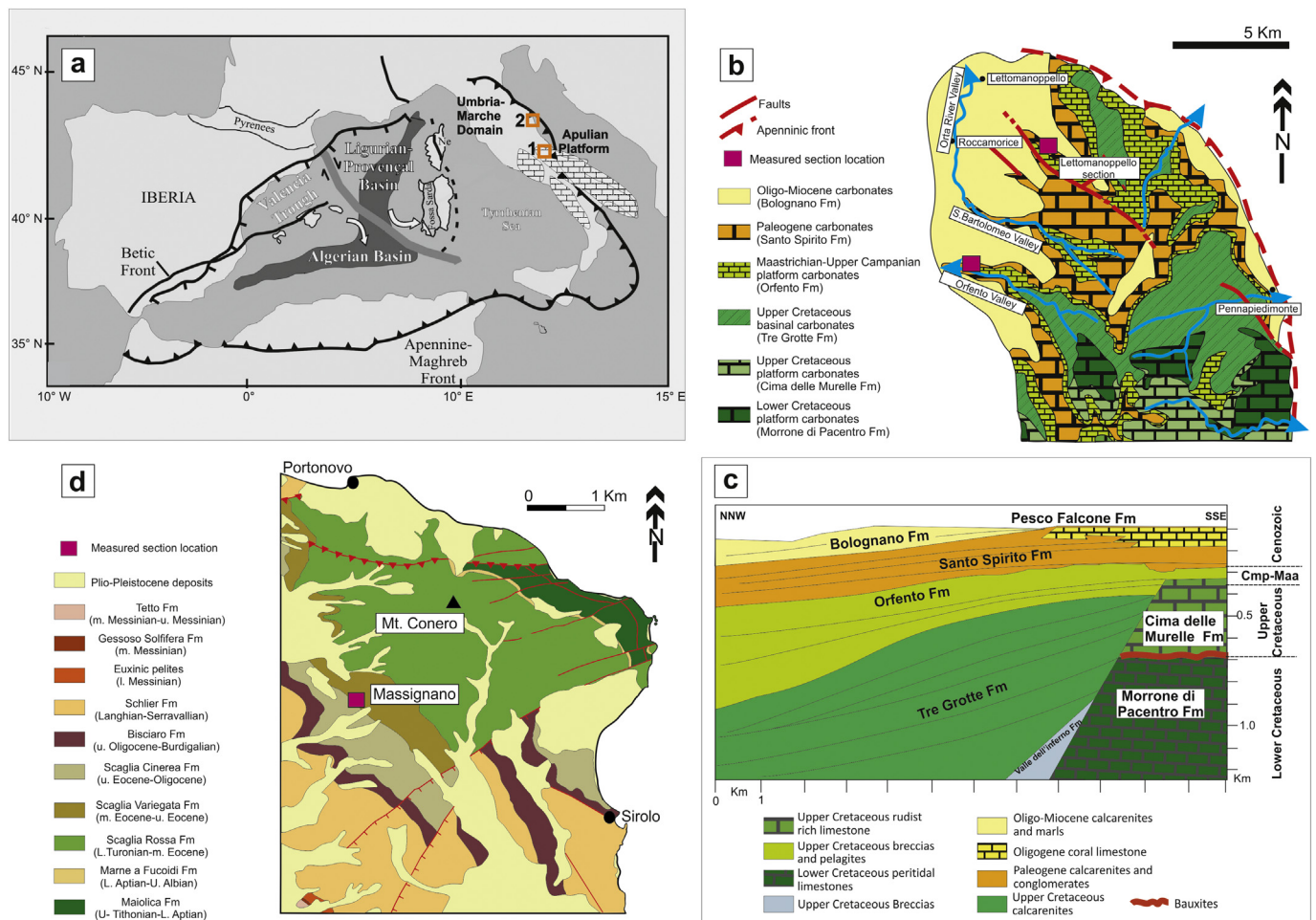
### 2.1. The Majella Mountain

The Majella Mountain (Fig. 1b) is a 35 km-long anticline that is convex towards the northeast, and plunges both northward and southward (Patacca et al., 2008). Its outcropping succession consists of Upper Jurassic to upper Miocene limestones and dolostones (Crescenti et al., 1969). During the Mesozoic, a steep erosional escarpment separated the platform top from the basin, which extended northward (Fig. 1c). By the late Campanian, the platform prograded over the basin, that was filled up by overlapping sediments (Vecsei et al., 1998). Thus, the Paleogene evolution of the Majella carbonate platform, corresponding to the Santo Spirito Formation, is represented by a continuous sedimentation along the platform margin and the slope, while the platform top shows long-term hiatuses and discontinuous deposits. In the upper Rupelian a discontinuity surface occurs, separating the Santo Spirito Formation from the Bolognano Formation (upper Rupelian-lower Messinian), which represents a carbonate ramp developed above the former shallow deposits of the platform (Mutti et al., 1997; Brandano et al., 2012, 2016a). The evolution of the Oligocene-Miocene ramp ended in the Messinian with the deposition of the *Turborotalita multiloba* Marls (Carnevale et al., 2011), followed by the Gessoso-Solfifera Formation (Crescenti et al., 1969). Lastly, during the early Pliocene, the Majella Mountain was involved into the foredeep system of the Apennine orogeny (Cosentino et al., 2010).

### 2.2. The Umbria-Marche Domain

The Umbria-Marche Domain consists of an Upper Triassic-Miocene sedimentary succession deposited in the northern margin of the Tethys (Fig. 1a and d). During the Late Triassic, the rifting of the Tethys led to a marine transgression, testified by the evaporitic deposition of the Anidridi di Burano and, subsequently, by the development of the Calcarea Massiccio carbonate platform (Pialli, 1971). During the Early Jurassic, the Calcarea Massiccio platform drowned due to the rift-related extensional tectonics, and the Umbria-Marche domain evolved into a wide basin characterised by intrabasinal structural highs (Centamore et al., 1971; Bernoulli and Jenkyns, 1974; Brandano et al., 2016b) until the Late Jurassic.

This heterogeneous paleogeography ended with the deposition of pelagic mudstones (the Maiolica Formation, Tortonian to Albian in age)



**Fig. 1.** a) Map of the Mediterranean area with the studied successions location. 1: Majella Mountain; 2: Conero area. Modified after Gueguen et al. (1998). b) Simplified geological map of the Majella Mountain (modified after Vecsei and Sanders, 1999) with section locations. c) Schematic stratigraphic architecture of the Majella carbonate platform, modified after Vecsei et al. (1998). d) Simplified geological map of the Conero area, with Massignano section location (modified after Coccioni et al., 1997).

that levelled the inherited topography (Alvarez, 1990). The evolution of the Umbria-Marche basin during the Cretaceous to Miocene interval resulted in the transition from pelagic to hemipelagic sedimentation, characterised by a constant increase of siliciclastic content, in a context of significant subsidence (Marchegiani et al., 1999; Guerrero et al., 2012). The last phase of orogeny is represented by a W-to-E migration of siliciclastic turbidites (Marnoso Arenacea Fm.) occurring in middle to late Miocene (Alvarez, 1999; Guerrero et al., 2012).

### 3. Materials and methods

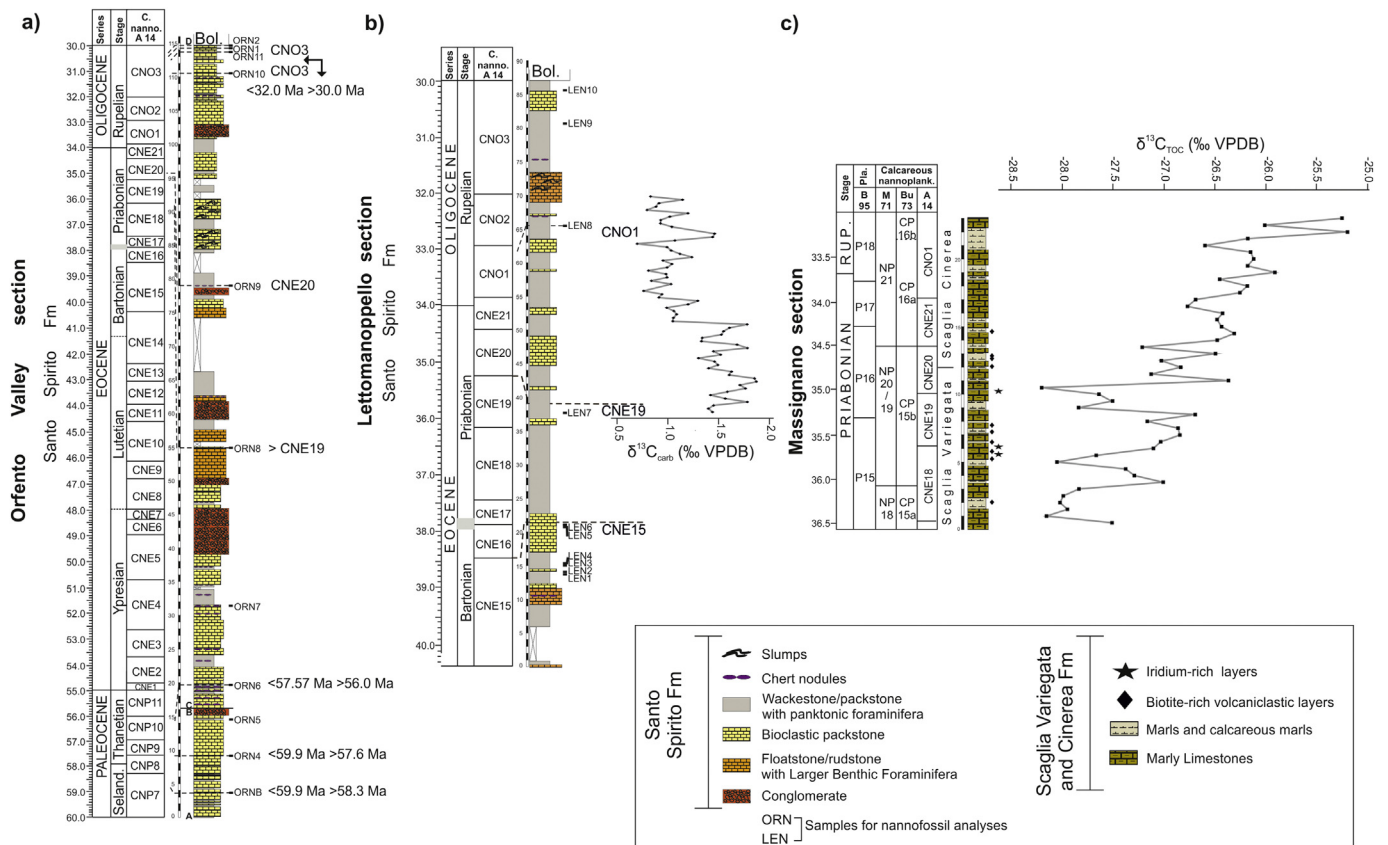
The studied materials originate from three different stratigraphic sections: the Lettomanoppello and the Orfento Valley sections (Majella Mountain, Central Italy), and the Massignano section (Conero area, Central Italy) (Fig. 2). Eighty-five thin sections of the Santo Spirito Formation, belonging to both the sections, were observed at the petrographic microscope for textural characterisation and skeletal components identification.

Previously published and newly generated data on calcareous nanofossil assemblages were considered for biostratigraphic characterisation. Eleven samples from the Orfento Valley section, collected from marly layers and chalk inclusions within cherty nodules if present, were examined. Smear-slides for nanofossils analysis were obtained following the procedure defined by Bown and Young (1998) and observed under a polarising microscope ( $\times 1200$  magnification). The nanofossil biostratigraphic data of the Lettomanoppello section are from Raffi

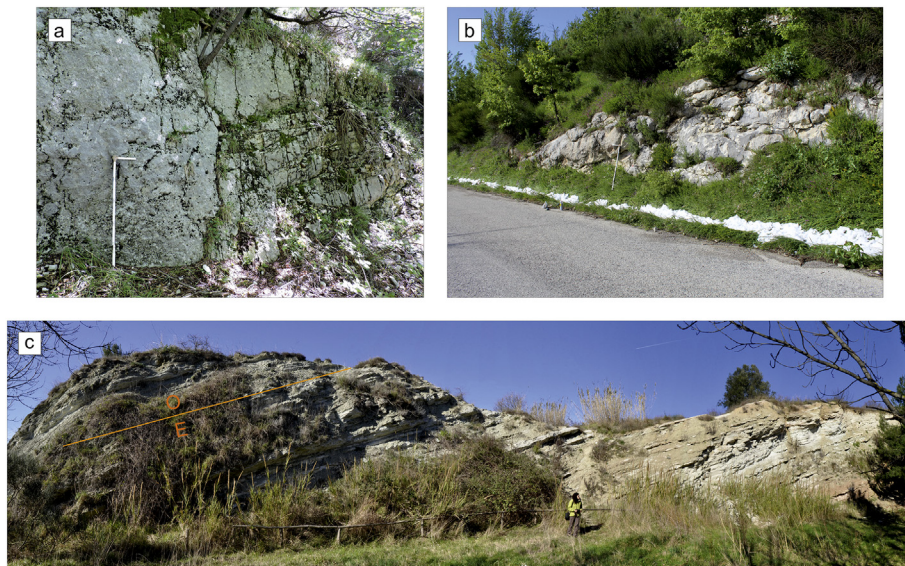
et al. (2016). For biozones and biochronology we refer to the Paleogene Zonation of Agnini et al. (2014).

The Lettomanoppello and the Massignano sections were sampled for carbon isotope stratigraphy, and a 0.5 m sampling interval was chosen. Stable isotope analyses were carried out at the Isotope Geochemistry Laboratory of the Istituto di Geologia Ambientale e Geoingegneria (IGAG-CNR) of Rome, and carbon and oxygen stable isotope ratios were measured on 63 bulk samples of the Lettomanoppello section. The whole-rock analyses were performed with a Finnigan Delta Plus Mass spectrometer coupled with a gas chromatography-based Gas Bench II, and stable isotope ratios were calibrated with the international NBS19 carbonate standard. All the results are expressed in the Vienna Pee Dee Belemnite (VPDB) scale. The analytical error was  $\pm 0.1\text{‰}$  (SD) based on replicate standards.

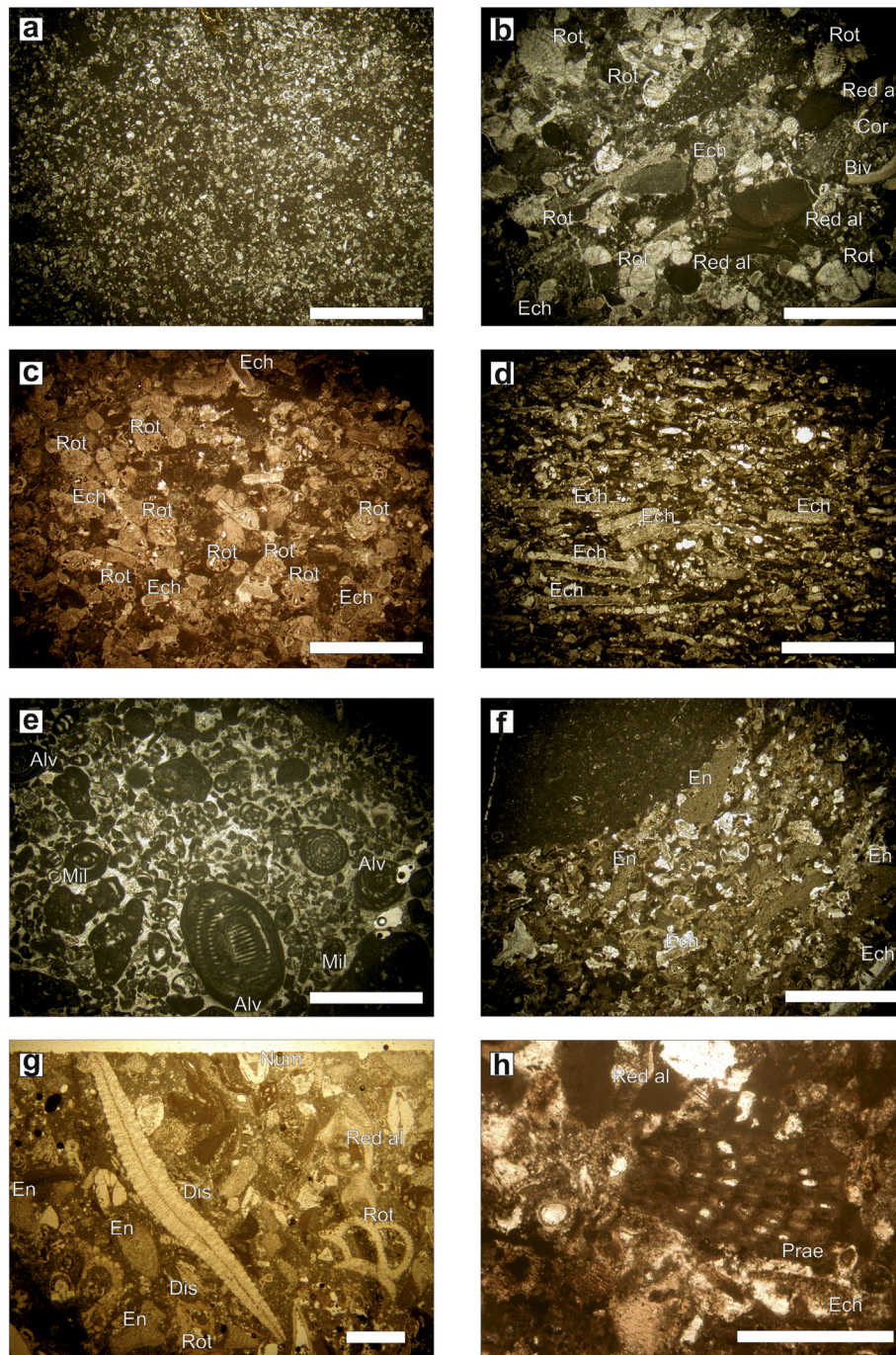
$\delta^{13}\text{C}_{\text{TOC}}$  was measured on 46 samples belonging to the Massignano section. Rock samples were hand-crushed in an agate mortar. To remove the carbonate fraction, 0.5 g of powder per sample was treated with 40 ml of an HCl 1 M solution and shaken with a magnetic stirrer for 20 min. After 20 min, the pH of the solution was  $\sim 2$ , indicating that the entire carbonate fraction was consumed and that the reaction was complete. The residual powder was rinsed with deionised water and oven-dried at  $50\text{ }^\circ\text{C}$ .  $\delta^{13}\text{C}_{\text{TOC}}$  analyses were performed with a Finnigan Delta Plus Mass spectrometer coupled with a Flash-1112 Thermo Elemental Analyzer. Organic matter carbon isotope ratios were calibrated with the IAEA-CH-6 and IAEA-CH-7 international standards. The analytical error was  $\pm 0.3\text{‰}$  based on replicate standards.



**Fig. 2.** Stratigraphic sections with carbon isotope records plotted against stratigraphic depth. a) Orfento Valley stratigraphic section; b) Lettomanoppello section; c) Massignano section. Age constraints for the Santo Spirito Formation (Orfento Valley and Lettomanoppello sections) are provided by calcareous nannofossil biostratigraphy (this work; Raffi et al., 2016) and refer to the Paleogene Zonation of Agnini et al. (2014). Age constraints for the Massignano section are referred to in Coccioni et al. (1988) and Premoli Silva and Jenkins (1993). A 14 = calcareous nannofossil zonation of Agnini et al. (2014). B 95 = planktonic foraminifera zonation of Berggren et al. (1995). Bu 73 = calcareous nannofossil zonation of Bukry (1973). M 71 = calcareous nannofossil biozones of Martini (1971).



**Fig. 3.** a) Conglomerate interval interrupting the normal outer ramp sedimentation in the Orfento Valley section. b) Slumps within the lower Oligocene portion of the Lettomanoppello section. c) Massignano section, the Eocene-Oligocene boundary is marked with the orange line.



**Fig. 4.** Microfacies of the Santo Spirito Formation. a) Highly bioturbated packstone with planktonic foraminifera. b) Bioclastic packstone. c) Packstone with rotalids. d) Packstone with echinoid fragments. e) Grainstone with *Alveolina* and miliolids. f) Packstone with encrusting foraminifera and intraclasts. g) Floatstone with LBF. h) Floatstone with LBF. Alv: *Alveolina*. Biv: Bivalve fragments. Cor: coral fragments. Dis: *Discocyclina*. Ech: echinoid fragments. En: encrusting foraminifera. Mil: miliolids. Num: nummulitids. Prae: *Praerhapydionina* cf. *40elicate*. Red al: red algae crusts and fragments. Rot: rotalids. Scale bar = 1 mm.

## 4. Results

### 4.1.1. The Orfento Valley section

The Orfento Valley section (Figs. 2a and 3a) consists of the upper Paleocene-lower Oligocene portion of the Santo Spirito Formation. The composite section is 114.6 m thick, the first section (A-B), is 16.2 m thick, the second (C-D) is 98.4 m thick (Fig. 2a). The first 15.1 m of the Orfento A-B section are represented by wackestones to packstones rich in planktonic foraminifera (Fig. 4a) alternated with laminated

bioclastic packstones. The main components of the bioclastic packstones are highly fragmented larger benthic foraminifera (LBF), such as *Nummulites*, *Discocyclina*, rotalids, and red algae crusts, together with echinoids and bivalve fragments (Fig. 4b, c, and d). Thin marly layers alternate with this calcareous facies of the Orfento A-B section, which is interrupted by a 1.1 m-thick conglomerate characterised by an erosional basal surface with LBF and intraclasts. The base of the Orfento C-D section consists of 22.2 m of coarse packstones with LBF alternated with mudstones/wackestones rich in planktonic foraminifera. Cherty nodules are very abundant in this interval.

At m 39.0, a 7 m-thick conglomerate crops out. This conglomerate is

chaotic and poorly sorted. LBF and coral fragments were identified together with intraclasts. The section continues with a 16.8 m-thick interval dominated by bioclastic packstones with LBF, such as *Nummulites* and *Alveolina*, alternated with floatstones with *Alveolina* in a grainstone matrix dominated by abundant miliolids and peneroplids, and with packstones containing encrusting foraminifera, such as *Gypsina moussaviani*, red algae crusts, and nodules (Fig. 4e and f). These facies are interrupted by two conglomerate intervals 0.6 m- (m 49.5) and 2.8 m-thick (m 59). This portion of the Orfento C-D section ends with a 3.50 m-thick interval of wackestones with planktonic foraminifera. Cherty nodules are common.

At m 74.2, after an 8 m covered interval, the upper portion of the Orfento C-D section crops out. It consists of an alternation of bioclastic packstones with LBF and wackestones with planktonic foraminifera. A 1.0 m-thick conglomerate interrupts the normal sedimentation at 77.5 m. From 84.5 m to 92.0 m, an interval of bioclastic packstones with LBF characterised by extensive slumps occurs. The uppermost conglomerate, which is 1.9 m-thick, crops out at 101 m. The 13.6 m interval at the top of the section consists of alternating very fine bioclastic packstones and planktonic-rich wackestones. Chert occurs in nodules and lists up to the top of the section (Fig. 2a).

**4.1.1.1. Biostratigraphy and age constraints.** Although the observed nannofossil assemblages in the sample from the Orfento Valley composite section are generally scarce and show moderate to poor preservation, it was possible to obtain some biostratigraphic and biochronologic constraints based on the presence of some marker species. In the few samples from the Orfento A-B section, and in a marly interlayer cropping out at m 3.4 of the Orfento C-D section (Fig. 2a), the presence of specimens belonging to the genus *Fasciculithus* (*F. ulii* group, *F. pileatus*, *F. tympaniformis*) indicate a biostratigraphic position included in the interval between Zones CNP7 and CNP9 of Agnini et al. (2014) (Plate 1–10). Therefore, the lower part of the composite Orfento section corresponds to the interval Selandian-lower Thanetian (< 59.9 Ma and > 57.0 Ma). In the interval from m 20 to m 55 in the composite section, the abundance of calcarenites and resedimented material limited the availability of samples suitable for biostratigraphy (Fig. 2a). The nannofossil assemblage observed in a marly interlayer at 55 m, and in the chalk found in a cherty nodule at 79 m, allows us to ascribe this interval to the Priabonian (CNE 19 and CNE20 of Agnini et al., 2014) due to the presence of very rare *Discoaster saipanensis* and *D. barbadiensis* and *Ericsonia formosa* (Plate 1 11–14). In the uppermost portion of the section (m 110.5 and m 113.5), the few samples show a scarce, but typical, lower Oligocene nannofossil assemblage, which places the top of the section in the CNO3 Zone (Agnini et al., 2014).

#### 4.1.2. The Lettomanoppello section

The Lettomanoppello section represents the middle Eocene-lower Oligocene portion of the Santo Spirito Formation, and the section is 87 m thick. The first interval consists of 9 m of highly bioturbated wackestones/packstones rich in planktonic foraminifera. The foraminiferal assemblage is dominated by globigerinids, such as *Globigerina*, *Globigerinatheka*, and *Turborotalia cerroazulensis*, and minor components are sponge spicules, echinoid and mollusc fragments. The following interval is represented by a 2.4 m thick interval of LBF floatstones to rudstones. The main components are LBF, such as *Nummulites*, *Assilina*, *Discocyclusina*, and rotalids (Figs. 2b and 4f). Other components are encrusting foraminifera, such as *Gypsina* and *Acervulina*, *Textularia*, *Nodosaria*, red algae, and echinoid and mollusc fragments (Fig. 4g).

From m 10.6 to m 69, the section consists of the alternation of bioclastic packstones and highly bioturbated wackestones rich in planktonic foraminifera (Fig. 2b). The main components of the bioclastic packstones are encrusting and epiphytic foraminifera, such as *Gypsina moussaviani* usually hooked, *Planorbulina*, *Lobatula lobatula* and

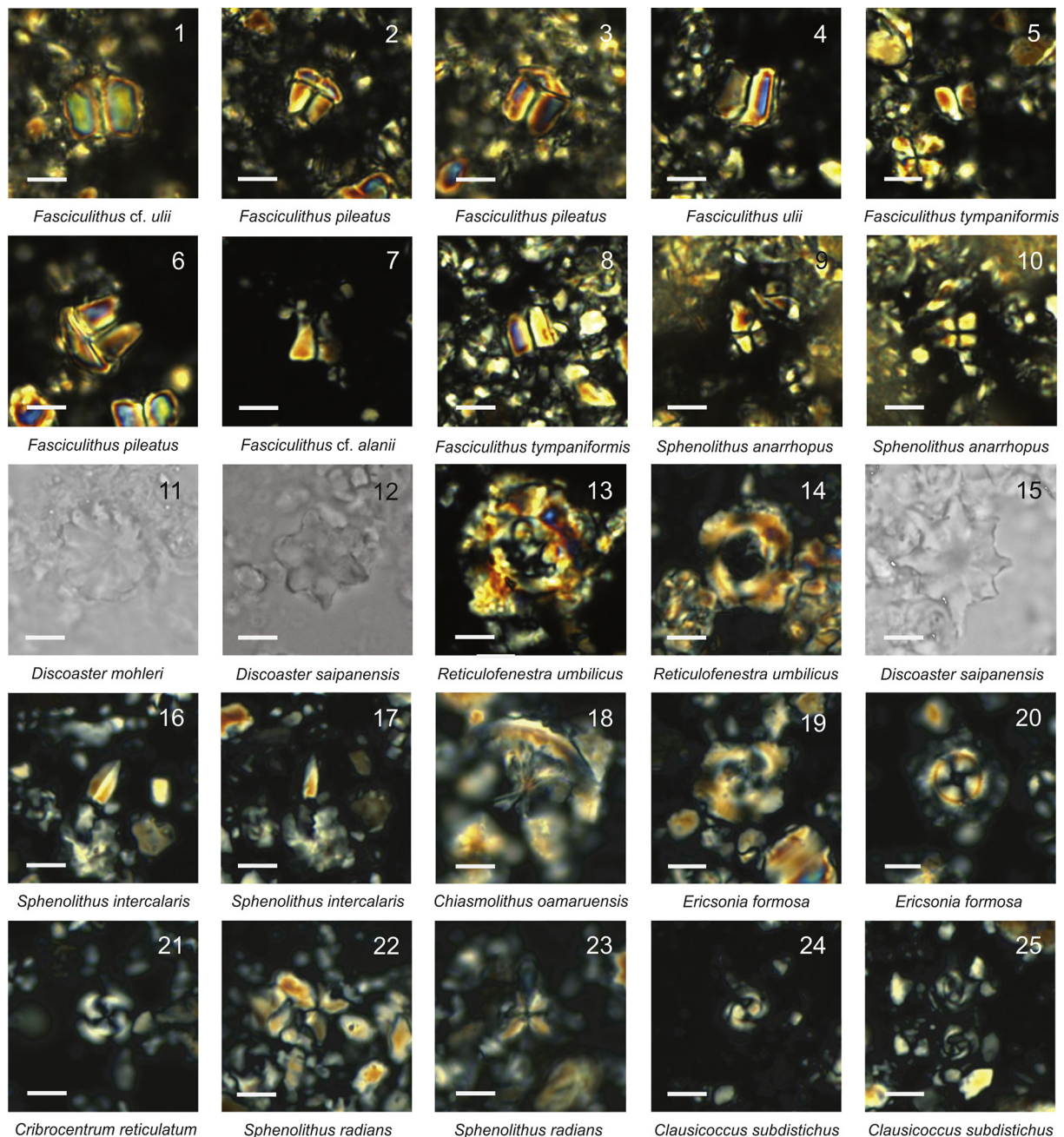
nubecularids. Both articulated branches and non-articulated crusts of red algae, sometimes hook-shaped, among which *Sporolithon* and *Subterraniphylum*, are frequent. Echinoid and mollusc, bryozoan, and *Ditrupea* fragments are common, together with small benthic foraminifera (SBF), such as miliolids, small rotalids, *Lenticulina*, textularids, and buliminacea, such as *Bolivina*. At 69 m, a 4.50 m-thick interval of floatstones/rudstones with LBF occurs. Among the LBF tests, *Phraerhapydionina cf. delicata* has been identified (Fig. 4h). This interval is characterised by extensive slumps. The last 13.6 m of the section consist of bioturbated wackestones rich in planktonic foraminifera, interrupted at 82.5 m by 3 m of bioclastic packstones. Cherty nodules occur at 75 m. A discontinuity surface marks the passage to the overlying Bolognano Formation at m 87.

**4.1.2.1. Biostratigraphy and age constraints.** Previously obtained nannofossil data from the Lettomanoppello section (Raffi et al., 2016) permitted us to obtain a detailed biostratigraphic classification of the section referring to Agnini et al. (2014) Zonation. The lower portion (up to 21.2 m) corresponds to the mid-Eocene (Bartonian) Zone CNE15 based on the presence of *Criboecium reticulatum*, *Sphenolithus obtusus*, *Sphenolithus spiniger*, and *Dictyococcites bisectus* (Plate 1 20–23). Above, in a sample from a marly layer at 37.8 m, the observed nannofossil assemblage indicates Zone CNE19 (*Discoaster saipanensis* TZ), thus ascribing the central portion of the section to the upper Eocene (Priabonian) (Plate 1 15–19). The biostratigraphic results from the upper portion of the section indicate the transition to the lower Oligocene. In fact, a marly layer at m 64.50 contains a nannofossil assemblage belonging to Zone CNO1 (*Ericsonia formosa* CRZ, age between 34 and 33 Ma), followed above (just below the boundary with the Bolognano Formation) by sediments of Zone CNO3 (*Dictyococcites bisectus* PRZ, age < 32 Ma) with *D. bisectus*, *Sphenolithus predistentus*, and *Sphenolithus celsus* (Plate 1 24–25).

**4.1.2.2. The C isotope record.** The upper Eocene-lower Oligocene  $\delta^{13}\text{C}_{\text{Carb}}$  values of the Lettomanoppello section range between +0.7‰ and +1.9‰ (Fig. 2b). The lower portion of the carbon isotope curve, from m 37.8 to m 50.8 in the section, shows a stationary trend with the values fluctuating between +1.4‰ and +1.9‰. A sharp negative shift of 0.7‰ amplitude is recorded starting from 50.8 m with values falling first to +1.1‰ at 51.8 m, and then falling further to +0.7‰ at 62.8 m in correspondence to the base of the Oligocene. This overall negative shift of 1.1‰ amplitude is the most significant in the analysed record. A 0.8‰ amplitude positive shift is recorded above (from 62.8 to 64.3 m), with values rising to +1.5‰. Whereas, the upper portion of the curve shows a stationary trend, with the  $\delta^{13}\text{C}_{\text{Carb}}$  values fluctuating around +1.0‰ up to the end of the measured interval in the section (at 69.8 m).

#### 4.1.3. The Massignano section

The Massignano section represents the upper Eocene-lower Oligocene interval of the Umbria-Marche basin succession. It is 23.0 m thick, and comprises the Scaglia Variegata and Scaglia Cinerea Formations (Figs. 2c and 3c). The Scaglia Variegata Formation consists of alternating reddish and greenish marls, calcareous marls, and marly limestones. This interval is characterised by horizontal and homogeneous beds, ranging in thickness between few and 50 cm centimetres. Bioturbation is common. The Scaglia Cinerea consists of homogeneous hemipelagic grey marls and marly limestones. Beds are tabular and range between few and 80 cm. Bioturbation is present in all the lithologies. Ten biotite-rich layers related to volcanic events characterise the section: six in the Scaglia Variegata, and four in the Scaglia Cinerea (Jovane et al., 2009). Lastly, three iridium-rich layers, cropping out at 5.61 m, 6.17 m, and 10.28 m, are interpreted as being linked to impactoclastic events (Bodiselsch et al., 2004) (Fig. 2c). The boundary between the Scaglia Variegata and the Scaglia Cinerea falls at 12.0 m (Coccioni et al., 1988).



**Plate 1.** Microphotographs of selected calcareous nannofossils in samples from the Orfento (ORN) and Lettomannoppello (LEN) sections. Scale bar 5  $\mu\text{m}$ .

1. *Fasciculithus cf. ulii* Perch-Nielsen, 1971; crossed nicols. Sample ORN6. 2, 3, 6. *Fasciculithus pileatus* Bukry; crossed nicols. 2) Sample ORN 2; 3, 6) Sample ORN4. 4. *Fasciculithus ulii* Perch-Nielsen, 1971; crossed nicols. Sample ORN4. 5, 8. *Fasciculithus tympaniformis* Hay and Mohler in Hay et al., 1967; crossed nicols. Sample ORN4. 7. *Fasciculithus cf. alanii* Perch-Nielsen, 1971; crossed nicols. Sample ORN4. 9, 10. *Sphenolithus anarrhopus* Bukry and Bramlette, 1969; 9) crossed nicols 45°, 10) crossed nicols 0°. Sample ORN6. 11. *Discoaster mohleri* Bukry and Percival Jr, 1971; parallel nicols. Sample ORN6. 12. *Discoaster saipanensis* Bramlette and Riedel, 1954; parallel nicols. Sample ORN8. 13, 14. *Reticulofenestra umbilicus* (Levin, 1965) Martini and Ritzkowski, 1968; crossed nicols. 13) Sample ORN8; 14) Sample LEN7. 15. *Discoaster saipanensis* Bramlette and Riedel, 1954; parallel nicols. Sample LEN7. 16, 17. *Sphenolithus intercalaris* Martini, 1971; 16) crossed nicols 0°, 17) crossed nicols 45°. Sample LEN7. 18. *Chiasmolithus oamaruensis* (Deflandre and Fert, 1954) Hay et al., 1966; broken specimen, crossed nicols. Sample LEN7. 19, 20. *Ericsonia formosa* (Kamptner 1963) Haq, 1971; crossed nicols. 19) Sample LEN7; 20) Sample LEN5. 21. *Cribrocentrum reticulatum* (Gartner Jr. and Smith, 1967) Perch-Nielsen, 1971; crossed nicols. Sample LEN5. 22, 23. *Sphenolithus radians* Deflandre, 1952; 22) crossed nicols 45°, 23) crossed nicols 0°. Sample LEN5. 24, 25. *Clausicoccus subdistichus* Roth and Hay in Hay et al., 1967 Prins; crossed nicols. Sample LEN8.

**4.1.3.1. Biostratigraphy and age constraints.** Based on planktonic foraminifera biostratigraphy, the Massignano section spans from Priabonian to Rupelian (Coccioni et al., 1988; Premoli Silva and Jenkins, 1993). The base of the section is placed in the upper P15 Zone of Blow (1969) and Berggren et al. (1995), or the top of the nannofossil biozone CNE17 of Agnini et al. (2014). The top of the section falls into the P18 Zone (Blow, 1969; Berggren et al., 1995),

which corresponds to the CNO1 Zone of Agnini et al. (2014). The Eocene/Oligocene boundary, marked by the extinction of the *Hantkeninidae*, is placed at 19 m (Coccioni et al., 1988). The calcareous nannofossils commonly used as markers for the late Eocene-early Oligocene are *Discoaster saipanensis* (last occurrence at 34.44 Ma; Pälke et al., 2006) and *Clausicoccus subdistichus*, whose increase (equivalent to 'Acme *Ericsonia obruta*') approximates the

Eocene/Oligocene boundary at 33.88 Ma (Pälike et al., 2006). Both biohorizons are recognised in the Massignano section (Premoli Silva and Jenkins, 1993).

**4.1.3.2. The C isotope record.** The upper Eocene-lower Oligocene  $\delta^{13}\text{C}_{\text{TOC}}$  record at Massignano ranges between  $-28.2\text{‰}$  and  $-25.2\text{‰}$  (Fig. 2c). An overall increasing trend is evident throughout the entire carbon isotope curve. The lowest portion of the curve (from 0 to 8.5 m) shows a stationary trend, with the values fluctuating around  $-27.5\text{‰}$ . A  $1.5\text{‰}$  abrupt negative shift occurs between 8.5 and 10.5 m, with the values decreasing from  $-26.7\text{‰}$  down to  $-28.2\text{‰}$ , that represents the lowest value recorded in the section. The distinct positive trend ( $> 2\text{‰}$  in amplitude) recorded from 10.5 to 19.0 m, with the values rising to  $-26.0\text{‰}$ , corresponds to the Eocene-Oligocene boundary interval. The carbon isotope record in the lowermost Oligocene shows the values fluctuating between  $-26.7\text{‰}$  and  $-26.0\text{‰}$  from 19.0 to 21.5 m, and rising to  $-25.2\text{‰}$  at the top of the section.

## 5. Discussion

### 5.1. Facies association and depositional model of the Santo Spirito formation

The Santo Spirito Formation represents a carbonate platform developed in the Majella Mountain during the Paleogene, and it can be referred to as the standard model for Cenozoic carbonate ramps of Buxton and Pedley (1989). The recognised lithofacies can be ascribed to an outer ramp and to the adjacent basin environments. In particular, the facies with highly bioturbated wackestones to packstones and planktonic foraminifera is interpreted as deposited within the aphotic zone and below the storm wave base of the outer ramp where only photo-independent organisms accumulate. The facies typical of an outer ramp environment gradually pass into facies corresponding to basinal settings, where marly wackestones to mudstones with planktonic foraminifera occur. In this lower portion of the ramp, accumulation of abundant bioclastic turbidites and debris flow deposits occurs, which constitute a complex fan system and interrupt the normal outer ramp (cf. Payros et al., 2007).

Analysis of the skeletal components of the bioclastic packstones, the floatstones/rudstones lithofacies, and the clasts of the conglomerates, permits the inference of different facies' belts that constitute the inner portions of the platform. The major components identified in the bioclastic packstones, such as encrusting foraminifera and hooked-shaped red algae crusts, together with miliolids and alveolinids, indicate the presence of a vegetated environment developed within the euphotic zone as in the inner ramp (Beavington-Penney et al., 2004; Tomassetti et al., 2016).

The seagrass meadows hosted an important carbonate factory of the Cenozoic inner ramps, where encrusting, epiphytic, and miliolids foraminifera lived together with red algae and molluscs (Buxton and Pedley, 1989; Beavington-Penney et al., 2004; Brandano et al., 2016c; Tomassetti et al., 2016; Tomás et al., 2016). The middle ramp was dominated by nummulitids in the upper portion, and distally by orthofragminids, as evident by the occurrence of common *Discocyclina* tests in the LBF-rich floatstones/rudstones. During the late Eocene, zooxanthellate corals and red algae also occurred, as testified by the presence of these components in the conglomerates and in the LBF floatstones to rudstones lithofacies observed in the Orfento Valley section.

### 5.2. Diagenetic overprint over $\delta^{13}\text{C}_{\text{Carb}}$ and $\delta^{13}\text{C}_{\text{TOC}}$ records

Precipitation of carbonates is associated with only little carbon isotopic fractionation relative to the dissolved inorganic carbon (DIC). Therefore, the  $\delta^{13}\text{C}$  of both inorganic and biologically precipitated

carbonate must be very close to the value of dissolved inorganic carbon in the ocean (Saltzman and Thomas, 2012), thus making the  $\delta^{13}\text{C}_{\text{Carb}}$  a very good proxy of the isotopic composition of the DIC at the moment of the carbonate precipitation. Furthermore, both  $\delta^{13}\text{C}_{\text{Carb}}$  and  $\delta^{13}\text{C}_{\text{TOC}}$  are almost insensitive to temperature changes (Grossman and Ku, 1986; Hayes et al., 1999; Maslin and Swann, 2005; Galimov, 2006; Burla et al., 2008). Pelagic carbonates are considered the most reliable record to study carbon isotope stratigraphy as they are not affected by meteoric diagenesis, and they are more continuous and better age-constrained (Weissert et al., 2008 and references therein).

Several recently published papers have demonstrated that shallow-water carbonates can preserve the original carbon isotope signature (John et al., 2003; Mutti et al., 2005; Franceschi et al., 2014; Brandano et al., 2015; Frijia et al., 2015), and show higher amplitude carbon isotope shifts in comparison to the coeval pelagic successions (Brandano et al., 2017b). Within the shallow-water skeletal associations, the heterozoan carbonates have greater potential to preserve the original marine isotopic ratios compared to the photozoan skeletal assemblages (Mutti et al., 2006) due to their mineralogical composition that is low-Mg calcite-dominated, which makes them more resistant to diagenetic alteration.

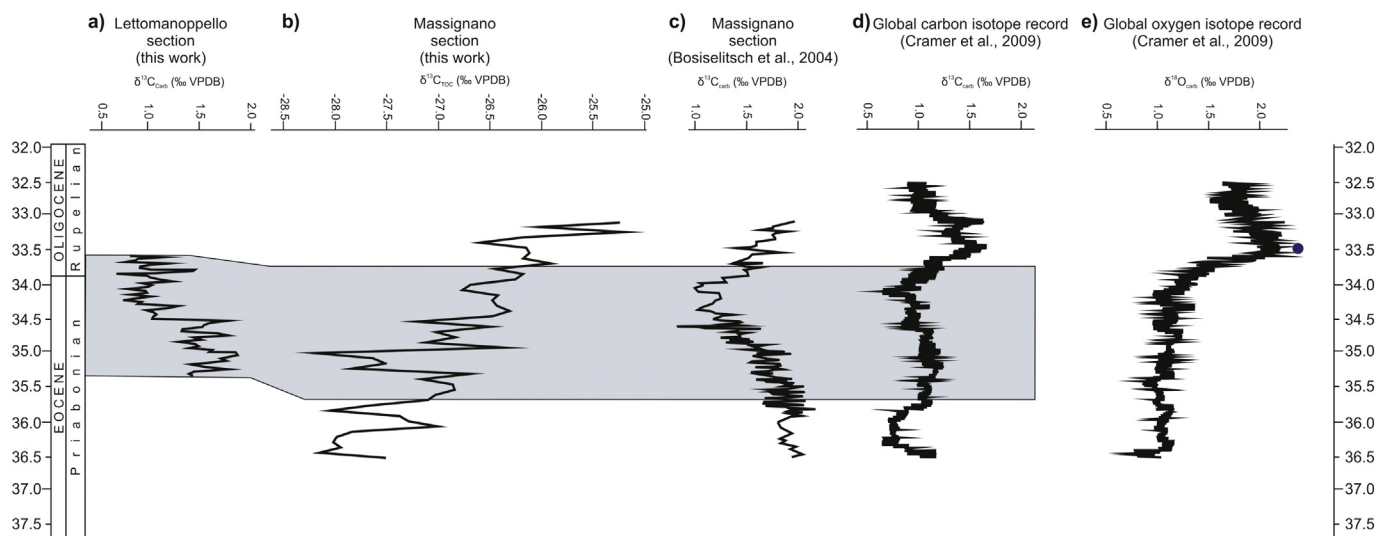
The microfacies analyses of the samples used for carbon isotope measurements show that the skeletal assemblages are dominated by low-Mg calcite shells, coupled with occasionally common H-Mg calcite organisms such as echinoids or red algae, whereas no aragonitic organisms are present. Only a minor growth of calcite cements is present, testifying for a diagenesis in a normal marine environment (Flügel, 2004). There isn't any evidence of deep burial diagenesis, neither of subaerial exposure, nor of diagenesis in a meteoric environment. Furthermore, the global compilation of  $\delta^{13}\text{C}_{\text{Carb}}$  values from benthic foraminifera records for the late Eocene-early Oligocene shows a range varying between  $0\text{‰}$  and  $+2\text{‰}$  (Cramer et al., 2009). Therefore, the Santo Spirito Formation carbon isotope record falls precisely within the coeval carbonate  $\delta^{13}\text{C}_{\text{Carb}}$  range allowing us to rule out any major diagenetic overprint that could have affected the investigated succession (Fig. 2b). Likewise, the  $\delta^{13}\text{C}_{\text{TOC}}$  record of the Massignano section is consistent with the  $\delta^{13}\text{C}$  of marine phytoplankton, with a significantly minor portion of terrestrial organic matter. Overall, the marine phytoplankton  $\delta^{13}\text{C}$  ranges nowadays between  $-10\text{‰}$  and  $-32\text{‰}$  (Sarmiento and Gruber, 2006). However, Hayes et al. (1999) state that during Eocene the marine organic matter  $\delta^{13}\text{C}$  was between  $-26\text{‰}$  and  $-28\text{‰}$  (Fig. 2c). Whereas, plants characterised by a C3 photosynthesis pathway, that were the most abundant type of plants during Eocene, show a  $\delta^{13}\text{C}$  ranging between  $-23\text{‰}$  and  $-33\text{‰}$  (mean value  $\sim -26\text{‰}$ , Maslin and Thomas, 2003).

### 5.3. The Central Mediterranean carbon isotopes shifts at the Eocene-Oligocene transition

#### 5.3.1. Comparison of the Mediterranean C-isotope signal with the global trend

The upper Eocene-lower Oligocene  $\delta^{13}\text{C}_{\text{Carb}}$  record of the Santo Spirito ramp and the coeval  $\delta^{13}\text{C}_{\text{TOC}}$  record of the Massignano hemipelagic succession matches the global signal (cf. Cramer et al., 2009, Fig. 5). The  $\delta^{13}\text{C}_{\text{Carb}}$  curves of Lettomannoppello and Massignano sections show a clear negative trend during the late Priabonian (Fig. 5a, c). This trend reflects the interval between the end of the C-cycle perturbation associated with the MECO and the onset of the carbon isotope anomaly contemporary to the Oi-1 event, as marked by a cooling trend associated with a decrease of primary production. This decrease is well recorded by the Eocene  $\delta^{13}\text{C}_{\text{TOC}}$  of the Massignano section, which shows an overall positive trend. Variations in the  $\delta^{13}\text{C}_{\text{TOC}}$  might be due to the exogenic carbon pool or related to changes in the source of the organic matter (terrestrial vs marine) (Sluijs and Dickens, 2012). Further studies would be needed to entirely exclude biases related to the different sources of organic matter, such as the analysis of the





**Fig. 5.** Comparison of the carbon isotope record of the studied sections with the global carbon and oxygen isotope record. All the curves have been calibrated with the Geological Time Scale 2004.

palynofacies, or the integration of the N/C ratio across the investigated section, to quantify precisely the variation of the amount of the terrestrial organic matter. These studies, besides useful, go beyond the scope of this work. Furthermore, the site and the paleodepth of the Massignano section already guarantee that the potential biases related to the influence of the terrestrial organic matter, even if possible, must have been minor or unlikely. In fact, the paleodepth of the Massignano section is estimated between 1000 and 1500 m (Coccioni and Galeotti, 2003). At this depth the marine phytoplankton derived organic matter is always significantly more abundant than the terrestrial organic matter (Bernier, 1989). Secondly, during late Eocene, the Apennine orogeny front was located far more west than today, and the absence of coarse-grained sediments within the Massignano succession provides an evidence that the continent was not close. Lastly, in oxic deposits the terrestrial organic matter is strongly degraded due to heterotrophs feeding and bioturbation, and usually appears as sparse amorphous particles and pellets (Lallier-Vergès and Albéric, 1990). For these reasons, even if we cannot rule out completely the influence of the terrestrial organic matter on the  $\delta^{13}\text{C}_{\text{TOC}}$  record at Massignano, we can definitely assume that it would have been unlikely and scarce.

The c-cycle perturbation that occurred during the Oi-1 has long been debated. The different interpretations vary from feedbacks among atmospheric  $\text{pCO}_2$  and ice sheet coverage and silicate weathering rates, to increased marine organic carbon burial/cycling. Different hypotheses suggest a global switch in the ecology of plankton that favoured siliceous organisms over calcareous ones, a shift in global carbonate sedimentation from the shelf to the deep ocean, changes in riverine chemical inputs, a heavier isotopic composition of the terrestrial biosphere, or an increase in the rain ratio of the inorganic/organic carbon from surface to deep ocean, to a reduced ocean acidity linked to an increased ocean overturning (Coxall and Wilson, 2011; Plancq et al., 2014).

According to Goldner et al. (2014), the decrease of atmospheric  $\text{pCO}_2$  during the EOT produced the cooling phase that promoted the growth of the Antarctica ice cap, which in turn enhanced the ocean thermal gradient and invigorated its circulation. The increased wind-driven upwelling around Antarctica drove an increase in ocean productivity, potentially drawing down carbon. Furthermore, it strengthened the cooling across the EOT, pointing towards a C-cycle perturbation related to the enhanced burial rates of organic carbon in marine sediments, coupled with increased surface water productivity (Salamy and Zachos, 1999; Zachos and Kump, 2005; Dunkley Jones et al., 2008).

### 5.3.2. Decoupling between $^{13}\text{C}_{\text{Carb}}$ and $^{13}\text{C}_{\text{TOC}}$

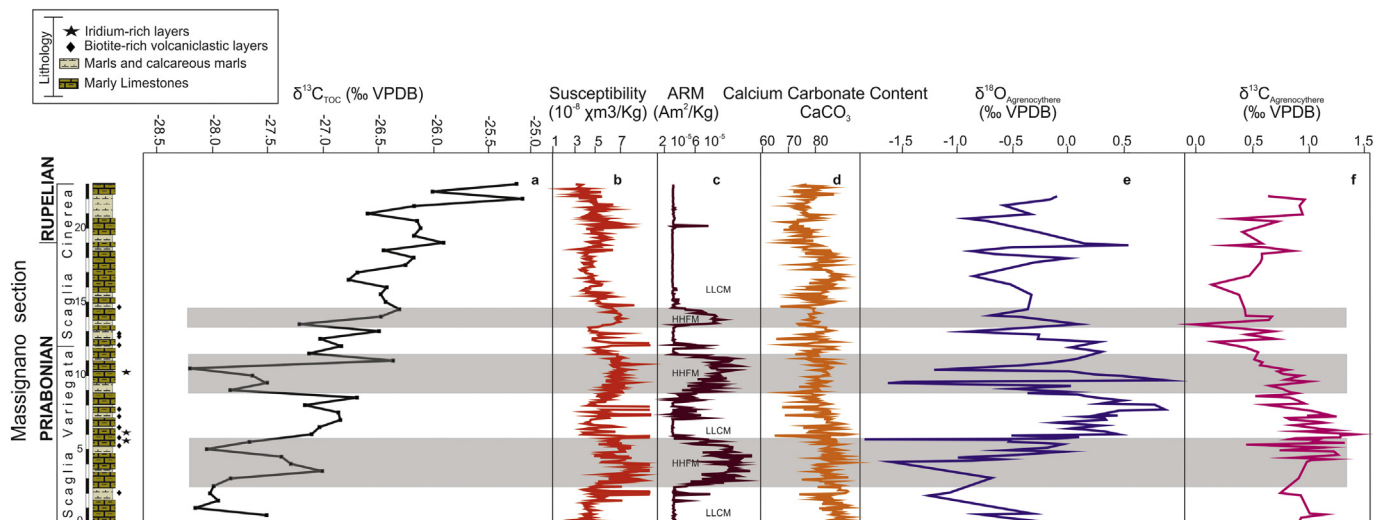
The upper Eocene  $\delta^{13}\text{C}_{\text{Carb}}$  and the  $\delta^{13}\text{C}_{\text{TOC}}$  curves of the analysed successions show opposite trends. This decoupling during the late Priabonian is consistent with the decreasing values of  $\text{CO}_2$  that characterise the late Eocene (Beerling and Royer, 2011; Foster et al., 2017) (Fig. 5a, b). The net isotopic fractionation between the total organic carbon (TOC) and sedimentary carbonates depends on four processes: the fractionation associated with primary producers, the fractionation between Dissolved Inorganic Carbon (DIC) and dissolved  $\text{CO}_2$ , the fractionation between DIC and carbonate minerals, and the fractionation associated with secondary biological processes (see Hayes et al., 1999 for a review). Among these processes, the first one is the most effective, but it has not been constant throughout time. In fact, Hayes et al. (1999) point out how the isotopic effect associated with primary production decreases as dissolved  $\text{CO}_2$  lowers, affecting the TOC sequestered in sediments, thus increasing the difference between the TOC and the DIC.

Within the overall positive  $\delta^{13}\text{C}_{\text{TOC}}$  trend, negative spikes are present in the interval between 36.5 and 36.0 Ma of the Massignano record (Figs. 2 and 6a). These spikes may be related to the environmental changes produced by the alternating activity of the westward subtropical Eocene Neo-tethys (STENT) current, as proposed by Jovane et al., 2007; Jovane et al., 2009. According to the authors, the paleoceanography of the Neo-tethys was characterised by alternating periods of low productivity, represented by intervals of deposition driven by local terrigenous material, and expressed by the authors as coarser magnetite and periods of high productivity and freshening, displayed by abundant fine-grained magnetite plus hematite layers.

The high productivity and freshening periods represent times when the westward subtropical Eocene Neo-tethys current entered through the Arabia-Eurasian gateway, bringing large amounts of fresh and deep-sea waters rich in dissolved iron. The negative spikes of  $\delta^{13}\text{C}_{\text{TOC}}$  of the Massignano section show a good correspondence with the fine hematite-rich sediments interval recognised by Jovane et al., 2007, and confirm the coexistence of the high biogenic productivity phases (Fig. 6). Lastly, in the Massignano section, the onset of the positive carbon isotope excursion of the early Oligocene is recorded (Bodiselsch et al., 2004), although in the section the entire perturbation is not displayed (Fig. 5c).

### 5.3.3. C-isotope trend and sea-level changes

The Priabonian  $\delta^{13}\text{C}_{\text{Carb}}$  curve of the Santo Spirito ramp at Lettomanoppello section shows the end of upper Eocene carbon isotope



**Fig. 6.** a) Comparison of the  $\delta^{13}\text{C}_{\text{TOC}}$  record of the Massignano (this work). b) Low-field magnetic susceptibility of the Massignano section. c) Anhysteretic Remanent Magnetization (ARM). d) Calcium Carbonate Content ( $\text{CaCO}_3\%$ ), e)  $\delta^{18}\text{O}$ , and f)  $\delta^{13}\text{C}$  isotope record measured on valves of the *Agrenocythere* ostracod genera throughout the Massignano section. (b), (c), (d), and (f) are modified from Jovane et al. (2009). HHFM = high concentration, high coercivity, and fine-grained magnetic; LLCM = low concentration, low coercivity, and coarse-grained magnetic (modified from Jovane et al., 2004).

negative trend, which is linked to the decrease of primary production at the end of the Eocene (Fig. 5a). The prominent  $\delta^{13}\text{C}$  excursion occurring in the early Oligocene associated with the onset of Antarctic glaciation is not recorded because the regular bedding is interrupted by the occurrence of extensive slumps. These slumps are indeed interpreted as the major consequence of the sea-level drop related to the cooling corresponding to the Oi-1 event. Slumps in the lower Oligocene (within Zone CNO1 of Agnini et al., 2014) have been identified not only in the Lettomannoppello and in the Orfento Valley sections (Figs. 2a, b, and 3b), but also in the eastern sector of the Santo Spirito ramp, namely in the Pennapedimonte stratigraphic section (Raffi et al., 2016), and indicate an enhanced instability over the entire ramp.

Houben et al. (2012), in a study of the Eocene-Oligocene succession of the Berici Mountain (Northern Italy), estimated a  $\sim 20$  m sea level drop in correspondence with the Eocene-Oligocene boundary related to the EOT, and then a further  $\sim 60$  m drop related to the Oi-1 event and the glaciation of Antarctica. Similarly, Miller et al. (2009) suggested a sea level drop related to the Oi-1 event of  $80 \pm 25$  m. In this framework, the occurrence of extensive slumps indicates the deepening of the storm weather wave base. It is widely accepted that slumps and soft-sediment deformation structures can be triggered by storm waves (Molina et al., 1998). The intensification and the more frequent occurrence of storm events on the Santo Spirito ramp could have also been enhanced by the onset of the new oceanographic conditions and strong wind-driven currents related to the stronger latitudinal gradient (Coxall and Pearson, 2007; Miller et al., 2009).

#### 5.3.4. Productivity conditions and carbonate factory response

The high biogenic productivity phases recorded during the late Eocene in the pelagic realm, testified by the negative  $\delta^{13}\text{C}_{\text{TOC}}$  spikes at Massignano section, do not seem to be equally recorded in the platform domain. Apparently, no significant changes in the carbonate factory of the Santo Spirito ramp occurred concomitantly with the EOT. On the other hand, since the depositional environment of Santo Spirito Formation identifies an outer ramp setting, the carbonate factory was consequently dominated by aphotic biota, such as small benthic foraminifera, mollusc and echinoid fragments, coupled with the pelagic components represented by the planktonic foraminifera accumulating in the outer ramp. Relevant compositional changes are observed only in the resedimented interval (floatstone to rudstone lithofacies) marked by the disappearance of *Discocyclina* specimens, the reduced occurrence of

*Nummulites*, and the spreading of red algae and corals.

A drastic change of the carbonate factory is recorded in the western margin of the Lessini shelf (Jaramillo-Vogel et al., 2013, 2016). While red algae and LBF dominate in the lower portion of the succession, its uppermost part is characterised by bryozoan-rich beds. According to Jaramillo-Vogel et al. (2013, 2016), the disappearance of *Discocyclina* and *Asterocyclina*, and the replacement of phototrophic organisms by heterotrophic ones could be due to a significant change in the environmental conditions. This change corresponds to a sharp positive shift in  $\delta^{13}\text{C}$  that starts just below the bryozoan beds and is interpreted as the main C-cycle perturbation occurring at the EOT.

This perturbation is associated with a major change in trophic resources as indicated in the Lessini Shelf by a rise in the total phosphorus content, which triggered an increased primary productivity and thus, a shallowing of the photic zone (Jaramillo-Vogel et al., 2016). Decreased water transparency would have limited light-dependent biota and left bryozoans as dominant benthic calcifying organisms.

The coeval deposition of bryozoan beds reported from other several localities in northern Italy is interpreted to represent the regional response of a carbonate platform depositional system to global oceanographic changes related to the beginning of the Antarctic glaciation. In the Southern Tethys platforms (Hyblea, southern Apulia, Malta), the progressive global cooling at the EOT produced a major decline of larger benthic foraminiferal assemblages. The *Discocyclina* genus disappeared as well as most of the *Nummulites* species, whereas coral bioconstructions underwent renewed development. Lastly, coralline algae became a dominant sediment-producing biota, and seagrass environments expanded, causing a change in the facies association of the Cenozoic carbonate platforms (Pedley, 1998; Nebelsick et al., 2005; Brandano et al., 2009, 2017a).

## 6. Conclusions

The data here presented testify that the oceanic carbon isotope trends have been recorded in the Neo-tethys shallow-water (Santo Spirito Formation, Majella Mountain, northern Apulia Platform) and basinal successions (Massignano section) since they overall match the deep-sea signal. The shallow-water  $\delta^{13}\text{C}_{\text{Carb}}$  record of the upper Eocene shows an overall negative trend, while the contemporary  $\delta^{13}\text{C}_{\text{TOC}}$  records a positive one. This general trend is related to reduced fractionation by primary producers during the late Eocene due to the

decreasing pCO<sub>2</sub> after the C-cycle perturbation of the MECO, and before the onset of the carbon anomaly linked to the Oi-1 event. However, this study proves that regional factors, superimposed to the global forcing, influenced the Neo-thetys carbon isotope record, as suggested by the sharp transient negative spikes that mark the δ<sup>13</sup>C<sub>TOC</sub> record of the Massignano section. The Neothetys paleoceanography and current patterns mostly controlled these short-term anomalies. They are indeed interpreted as short times of higher productivity linked to enhanced nutrient availability and triggered by the westward subtropical Eocene Neo-thetys current entering from the Arabian-Eurasian gateway.

In contrast, the shallow-water carbon isotope signal does not record these short-term pulses of productivity. In fact, no significant changes in the carbonate factory of the Santo Spirito Formation are recorded. This fact does not mean a minor sensitivity of shallow-water successions, but it is related to the investigated depositional environments that range from the outer ramp to a pelagic setting where only photo-independent organisms occur and accumulate. However, the compositional characteristics of the resedimented sediments from the photic zone of the ramp throughout the EOT indicate a change in the composition of the carbonate factory during the Oligocene, with the disappearance of orthophragminids, the reduction of *Nummulites*, and the expansion of biota associated with the seagrass together with the spread of corals and red algae. Furthermore, the Santo Spirito Formation does not record the major carbon isotope shift concomitantly with the Oi-1 event as in the lowermost Oligocene the regular bedding of the sedimentary succession is interrupted by the occurrence of extensive slumps. The slumps represent evidence of the sea-level drop that occurred at the onset of the Antarctica ice-sheet, which caused the deepening of the storm wave base, and eventually increased instability over the whole ramp. This discovery is even more important if the rarity of continuous shallow-water successions across the Eocene-Oligocene transition, due to the major sea level drop, is taken into consideration.

Finally, this study proves how different evidences may characterise the shallow-water systems response to Eocene-Oligocene transition in comparison to the pelagic settings, attesting the importance of integrated studies to understand the complex dynamics related to the greenhouse-icehouse climate transition.

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