1	Massive and permanent decline of symbiont bearing morozovellids and $\delta^{13}C$
2	perturbations across the Early Eocene Climatic Optimum at the Possagno
3	section (Southern Alps of northeastern Italy)
4	
4 5	Valaria Luciani ¹ Jan Paalman ² Eliana Farmaciari ³ Luca Ciucharti ³ Claudia Acuini ³ Daharta
6	Valeria Luciani , Jan Dackmani , Enana Fornaciari , Luca Giusberti , Ciautia Aginni , Roberta D'Onofrio ¹
7	
8	
9	¹ Department of Physics and Earth Sciences, Ferrara University, Polo Scientifico Tecnologico, via G. Saragat 1,
10	44100, Ferrara, Italy
11	² Department of Geological Sciences, Stockholm University, SE-10691 Stockholm, Sweden
12	³ Department of Geosciences, Padova University, via G. Gradenigo 6, 35131, Padova, Italy
13	
14	Correspondence to: V. Luciani (valeria.luciani@unife.it)
15	
16	Abstract - The Early Eocene Climatic Optimum (EECO) records the highest prolonged global
17	temperatures over the past 90 Ma. Understanding the causes and timing of Eocene climate change
18	remains a major challenge in Cenozoic paleoceanography, which includes the biotic response to
19	climate variability and the changes among planktic foraminiferal assemblages across the EECO.
20	The symbiont bearing and shallow dwelling genera Morozovella and Acarinina were important
21	calcifiers in the tropical-subtropical early Paleogene oceans but almost completely disappeared at
22	about 38 Ma, near the Bartonian/Priabonian boundary. We show here that morozovellids record a
23	first critical step across the EECO through a major permanent decline in relative abundance from
24	the Tethyan Possagno section and ODP Site 1051 in the western subtropical North Atlantic.
25	Possible causes may include increased eutrophication, weak water column stratification, changes in
26	ocean chemistry, loss of photosymbionts and possible complex interaction with other microfossil
27	groups. Relative abundances of planktic foraminiferal taxa at Possagno parallel negative shifts in
28	both $\delta^{13}C$ and $\delta^{18}O$ of bulk sediment from Chron C24r to basal Chron C20r. The post-EECO stable
29	isotopic excursions towards lighter values are of modest intensity. Significant though ephemeral
30	modifications in the planktic foraminiferal communities occur during these minor isotopic

31 excursions. These modifications are marked by pronounced increases in relative abundance of 32 acarininids, in a manner similar to their behaviour during pre-EECO hyperthermals in the Tethyan 33 settings, which suggest a pronounced biotic sensitivity to climate change of planktic foraminifera 34 even during the post-EECO interval.

35

36 1 Introduction

37

38 The Early Eocene Climatic Optimum (EECO) is the interval in which the Earth's climate attained its 39 warmest state of the past 90 Ma and it represents a major turning point in the Cenozoic climate, as it 40 was followed by a long term cooling throughout the remainder of the Eocene, which culminated with 41 the formation of permanent large ice sheets on Antarctica at the end of the Eocene (Zachos et al., 2001; 42 Coxall et al., 2005). Superimposed on the long term early Eocene climate trend, short lived (<200 kyr) 43 warm events occurred, named hyperthermals, the most extreme of which is the well known Paleocene 44 Eocene Thermal Maximum (PETM) (Agnini et al., 2009; Coccioni et al., 2012; Cramer et al., 2003; 45 Kennett and Stott, 1991; Lourens et al., 2005; Nicolò et al., 2007; Quillévéré et al., 2008; Zachos et al., 46 2008). Massive, rapid releases of isotopically light carbon are linked to hyperthermals and concomitant 47 deep-sea carbonate dissolution events (Dickens, 2011; Dickens et al., 1995, 1997; Zachos et al., 2005, 48 2008). The series of early Eocene hyperthermals (Littler et al., 2014; Slotnick et al., 2012; Zachos et 49 al., 2010) has been suggested to continue into the earliest middle Eocene, with an additional thirteen 50 brief (~40 kyr) and less pronounced events (Kirtland Turner et al., 2014; Sexton et al., 2011). 51 Although the EECO still lacks a formal definition in terms of absolute age and duration (Slotnick 52 et al., 2012), this interval is thought to represent a ca. 2-3 Ma long early Eocene interval of extreme 53 warmth between about 52 and 50 Ma (Zachos et al., 2001) or between about 53 Ma and 51 Ma (Zachos 54 et al., 2008). Current thought about the placement of the EECO thus encompasses the interval from 55 approximately 50 Ma to 53 Ma. Regardless of the exact duration and position on the chronometric 56 scale of the EECO, it is clearly much longer than any of the brief early Eocene hyperthermals (Zachos 57 et al., 2001; 2008; 2010). Exceptionally high and long lasting atmospheric pCO_2 conditions are 58 considered to have played a primary control on the EECO warmth (Fletcher et al., 2008; Hyland and 59 Sheldon, 2013; Komar et al., 2013; Lowenstein and Demicco, 2006; Pearson and Palmer, 2000; Smith 60 et al., 2010). Slow addition of depleted carbon dioxide from volcanism, the second emplacement phase

61 of the North Atlantic Igneous Province, and increased weathering of silicate rocks are often invoked as 62 the main trigger of the EECO warming and the subsequent long term cooling trend (Demicco, 2004; 63 Raymo and Ruddiman, 1992; Sinton and Duncan, 1998; Vogt, 1979; Zachos et al., 2008). The 64 influence of a major switch from continental to island arc volcanism around 50 Ma may also have 65 played a role in perturbing the carbon cycle change that helped end the warm EECO interval (Dickens 66 et al., 2014; Lee et al., 2013). 67 It is well established that major plant and mammal faunal evolutionary turnovers occurred during 68 the EECO (Falkowski et al., 2005; Figueirido et al., 2012; Wilf et al., 2003; Wing et al., 1991; 69 Woodbourne et al., 2009; Zonneveld et al., 2000). In the marine realm, changes in evolutionary trends

70 have also been observed, for example the inception to the modern structure among calcareous

nannofossil communities (Agnini et al., 2006; Shamrock and Watkins, 2012; Schneider et al., 2011)

72 and possibly in diatom lineages (Oreshkina, 2012; Sims et al., 2006). These observations both from

continents and the oceans support the hypothesis of a primary interaction between climate change andbiotic evolution.

The relationship between the EECO and the paleoecology and evolution of planktic foraminifera is insufficiently known. At the beginning of the Eocene, planktic foraminiferal history was far enough from the Cretaceous-Paleogene mass extinction to have originated several phylogenetic lines with taxa occupying different ecological niches in the upper water column. The Eocene is a crucial interval in evolution of planktic foraminifera that encompassed one of their major diversifications reaching a peak in the middle Eocene (Norris, 1991; Pearson et al., 2006).

Within this plankton group, the symbiont bearing and shallow dwelling morozovellids and
acarininids are of particular interest because they dominated the tropical and subtropical assemblages
of the early Paleogene oceans. These genera belong to the muricate group, from the muricae that form
conical pustules on the test wall. Among calcareous microplankton, the muricates were one of the
major calcifiers in the low latitude early Paleogene oceans and almost completely disappeared at about
38 Ma, near the Bartonian/Priabonian boundary (Agnini et al., 2011; Luciani et al., 2010; Wade, 2004;
Wade et al., 2012).

The hemipelagic Possagno sedimentary succession is located in the Venetian Prealps of
northeastern Italy (Fig. 1). This section represents continuous deposition of the early through early
middle Eocene interval (55-46 Ma) from a bathyal setting in the central-western Tethys. A robust

91	Eocene biomagnetostratigraphy was established by Agnini et al. (2006). The Possagno section thus
92	spans the EECO interval, here agreed as the interval from about 53 to 50 Ma (Slotnick et al., 2012).
93	According to Agnini et al. (2006), the Possagno section is 66 m thick extending from the
94	Paleocene/Eocene boundary at about 56 Ma to the early middle Eocene (lower Chron C20r) at 46 Ma
95	on the time scale of Cande and Kent (1995). The aim here is to investigate the response of the
96	muricates in terms of relative abundance throughout the early and early middle Eocene, including the
97	EECO interval, in the 56 Ma through 46 Ma interval at Possagno, encompassing biozones planktic
98	foraminiferal Zones E1 to lower E8 (Luciani and Giusberti, 2014). In addition, the planktic
99	foraminiferal changes recorded at Possagno are compared with those observed from ODP Site 1051 in
100	the western subtropical North Atlantic. Additional aims include (1) to document planktic foraminiferal
101	changes in the Possagno section in relation to stable carbon and oxygen isotopes for the purpose to
102	unravel paleoenviromental conditions of the upper water column and (2), to distinguish ephemeral
103	biotic modifications during brief peaks of warming from permanent evolutionary changes in the
104	Tethyan and North Atlantic realms.
105	
106	2 The Possagno section and Site 1051: setting and stratigraphy
100	
107	
107 108	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in
107 108 109	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the
107 108 109 110	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia
107 108 109 110 111	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975;
107 108 109 110 111 112	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975; Thomas, 1998) in the western part of the Belluno Basin, a Mesozoic–Cenozoic paleogeographic unit
 107 107 108 109 110 111 112 113 	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975; Thomas, 1998) in the western part of the Belluno Basin, a Mesozoic–Cenozoic paleogeographic unit of the Southern Alps characterized by Jurassic to Eocene pelagic sequences (Bosellini, 1989). The
 107 108 109 110 111 112 113 114 	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975; Thomas, 1998) in the western part of the Belluno Basin, a Mesozoic–Cenozoic paleogeographic unit of the Southern Alps characterized by Jurassic to Eocene pelagic sequences (Bosellini, 1989). The basal part of the Possagno section straddles a 30 cm thick interval of dark-red clayey marls (Arenillas
 107 108 109 110 111 112 113 114 115 	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975; Thomas, 1998) in the western part of the Belluno Basin, a Mesozoic–Cenozoic paleogeographic unit of the Southern Alps characterized by Jurassic to Eocene pelagic sequences (Bosellini, 1989). The basal part of the Possagno section straddles a 30 cm thick interval of dark-red clayey marls (Arenillas et al., 1999; Agnini et al., 2006), referred to as the Clay Marl Unit (CMU) by Giusberti et al. (2007).
107 108 109 110 111 112 113 114 115 116	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975; Thomas, 1998) in the western part of the Belluno Basin, a Mesozoic–Cenozoic paleogeographic unit of the Southern Alps characterized by Jurassic to Eocene pelagic sequences (Bosellini, 1989). The basal part of the Possagno section straddles a 30 cm thick interval of dark-red clayey marls (Arenillas et al., 1999; Agnini et al., 2006), referred to as the Clay Marl Unit (CMU) by Giusberti et al. (2007). The CMU is the lithological expression of the main carbon isotope excursion (CIE) of the Paleocene
107 108 109 110 111 112 113 114 115 116 117	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975; Thomas, 1998) in the western part of the Belluno Basin, a Mesozoic–Cenozoic paleogeographic unit of the Southern Alps characterized by Jurassic to Eocene pelagic sequences (Bosellini, 1989). The basal part of the Possagno section straddles a 30 cm thick interval of dark-red clayey marls (Arenillas et al., 1999; Agnini et al., 2006), referred to as the Clay Marl Unit (CMU) by Giusberti et al. (2007). The CMU is the lithological expression of the main carbon isotope excursion (CIE) of the Paleocene Eocene Thermal Maximum (PETM), the base of which defines the Paleocene-Eocene boundary
107 108 109 110 111 112 113 114 115 116 117 118	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975; Thomas, 1998) in the western part of the Belluno Basin, a Mesozoic–Cenozoic paleogeographic unit of the Southern Alps characterized by Jurassic to Eocene pelagic sequences (Bosellini, 1989). The basal part of the Possagno section straddles a 30 cm thick interval of dark-red clayey marls (Arenillas et al., 1999; Agnini et al., 2006), referred to as the Clay Marl Unit (CMU) by Giusberti et al. (2007). The CMU is the lithological expression of the main carbon isotope excursion (CIE) of the Paleocene Eocene Thermal Maximum (PETM), the base of which defines the Paleocene-Eocene boundary (Aubry et al., 2007). The bio- lithostratigraphic assignment of the Possagno sediments follows Luciani
107 108 109 110 111 112 113 114 115 116 117 118 119	An upper Cretaceous-Miocene succession crops out at the bottom of the Monte Grappa Massif in the Possagno area, about 60 km NW of Venice. The lower to middle Eocene interval, which is the focus of this study, is represented by pelagic to hemipelagic Scaglia facies sediments. The Scaglia beds at Possagno are considered to have been deposited at middle to lower bathyal depths (Cita, 1975; Thomas, 1998) in the western part of the Belluno Basin, a Mesozoic–Cenozoic paleogeographic unit of the Southern Alps characterized by Jurassic to Eocene pelagic sequences (Bosellini, 1989). The basal part of the Possagno section straddles a 30 cm thick interval of dark-red clayey marls (Arenillas et al., 1999; Agnini et al., 2006), referred to as the Clay Marl Unit (CMU) by Giusberti et al. (2007). The CMU is the lithological expression of the main carbon isotope excursion (CIE) of the Paleocene Eocene Thermal Maximum (PETM), the base of which defines the Paleocene-Eocene boundary (Aubry et al., 2007). The bio- lithostratigraphic assignment of the Possagno sediments follows Luciani and Giusberti (2014), and the magnetostratigraphy is from Agnini et al. (2006) (Figs. 2,3).

121 Escarpment in the western North Atlantic (Norris et al, 1998). ODP Site 1051 is located well above 122 the local lysocline and the carbonate compensation depth. The sediments studied here are from 452.24 123 to 353.10 meters below sea floor (mbsf) and consists of lower to middle Eocene carbonate ooze and 124 chalk (Norris et al., 1998). This part of the Eocene section shows good recovery except between 382 125 mbsf and 390 mbsf (Fig. 4) and contains abundant calcareous plankton. Magnetostratigraphy is from 126 Ogg and Bardot (2001). Paleodepth estimates from benthic foraminiferal assemblages indicate lower 127 bathyal depth (1000-2000 m) during late Paleocene-middle Eocene (Norris et al., 1998). Bohaty et al. 128 (2009) derived a paleodepth of about 2200 m for the interval around to 50 Ma through a standard 129 subsidence model.

130

131

132 3 Methods

133

134 Analyses of foraminifera and stable isotopes were performed from the identical sample set of the 135 Possagno section previously used for calcareous nannofossils (Agnini et al., 2006). Relative 136 abundances have been determined from about 300 specimens extracted from each of the 110 samples 137 investigated in the >63 μ m size fraction. A sampling interval of 2–5 cm was used in the basal 0.7 m of 138 the Possagno section, followed by 50 cm spacing for the 0.7-14 m interval, and 20 cm for the 14-66 m 139 interval. Washed residues were prepared following standard procedures, which varied with the 140 different lithologies. Foraminifera were successfully extracted from the indurated marly limestones and 141 limestones using the cold-acetolyse technique (Lirer, 2000; Luciani and Giusberti, 2014), a highly 142 successful method for disaggregating strongly lithified samples (Fornaciari et al., 2007; Luciani et al., 143 2007), otherwise analyzable only in thin section. The marly samples were disaggregated using 30 % 144 hydrogen peroxide and subsequently washed and sieved using a 63 µm sieve. In most cases, gentle 145 ultrasonic treatment improved the cleaning of the tests. In the Possagno section, foraminifera are 146 continuously present and diverse throughout the studied interval with a preservation varying from 147 moderate to fairly good, even though tests are recrystallized and essentially totally filled. 148 The weight percent of the $>63 \,\mu\text{m}$ size fraction relative to the weight of the bulk sample, typically 149 100 g/sample, for the 110 Possagno samples is referred to as the coarse fraction, following Hancock 150 and Dickens (2005). Investigation of fifty Eocene samples at Site 1051 (Hole 1051A) from 452.24 to

151	353.1 mbsf, corresponding to ~52-47 Ma, had a spacing varying from 2.0 m to 0.5 m. These were
152	prepared using disaggregation using distilled water and washing over 38 μm and 63 μm sieves. Washed
153	residues were dried at <50°C. Planktic foraminifera from Site 1051 are abundant and well preserved.
154	The degree of dissolution, expressed as fragmentation index (F index), has been evaluated by
155	counting the number of planktic foraminiferal fragments or partially dissolved tests versus entire tests
156	on 300 elements, following Hancock and Dickens (2005). These data are expressed in percentages.
157	Fragmented foraminifera include specimens showing missing chambers and substantial breakage.
158	Carbon and oxygen stable isotope data of bulk sediment samples were analysed using a Finnigan
159	MAT 252 mass spectrometer equipped with a Kiel device at Stockholm University. Precision is within
160	± 0.06 ‰ for carbon isotopes and ± 0.07 ‰ for oxygen isotopes. Stable isotopes values are calibrated to
161	the Vienna Pee Dee Belemnite standard (VPDB) and converted to conventional delta notation ($\delta^{13}C$
162	and δ^{18} O).
163	
164	
165	4 Results
166	
167	4.1 Foraminiferal fragmentation
168	
169	The F index recorded at Possagno (Fig. 3) displays a large variability throughout the interval
170	investigated. The highest values, up to 70 %, were observed in the 16-22 m interval. The maximum F
171	index values correspond to the minimum values in the δ^{13} C record. A number of F index peaks mimic
172	the δ^{13} C negative peaks below 16 m, showing values between 60 % and 70 %.
173	F index values at Site 1051 (Fig. 4) show less variability with respect to Possagno. A maximum
174	value of 60 % is reached in Zone E5, just below an interval of uncertain magnetostratigraphic
175	attribution (Norris et al., 1998), here referred to as Chron C23r. Relatively high F index values around
176	50 % occur the upper portions of Chrons C24n and C22r. The interval across the EECO, on the basis of
177	biomagnetostratigraphic correlation and here placed between ca 397 and 425 mbsf, displays F index
178	
	values (<20 %). The relatively lower F index values at Site 1051 are presumably caused by less
179	values (<20 %). The relatively lower <i>F</i> index values at Site 1051 are presumably caused by less carbonate dissolution at that site and, to some extent, the lower resolution of the investigated samples at

181	
182	4.2 Weight percent coarse fraction
183	
184	Carbonate dissolution generally causes the bulk sediment coarse fraction to decrease because of
185	fragmentation of foraminiferal tests (Hancock and Dickens, 2005). The coarse fraction and F index
186	data from Possagno (Fig. 3) do not show such an anti-phasing, especially in the post-EECO interval.
187	The coarse fraction at Possagno shows minor fluctuations with a mean value of 5.3 ± 1.5 % from the
188	base of the EECO and upsection, with pre-EECO values varying around 10 %.
189	
190	4.3 Carbon and oxygen isotopes
191	
192	The δ^{13} C data from Possagno show a negative shift of about 1.5 ‰ at the 0 m level, which
193	corresponds to the Paleocene-Eocene boundary (Agnini et al., 2009) There are nine additional negative
194	carbon isotope excursions above the Paleocene-Eocene boundary in the lower 21.4 m of the Possagno
195	section (Fig. 2, Tab. S1):
196	1. 0.9 ‰ at 21.4 m (C22r) within EECO
197	2. 1.1 ‰ at 20.8 m (C22r) within EECO
198	3. 0.6 ‰ at 19.8 m (C23n) within EECO
199	4. 0.8 ‰ at 18.0 m (C23n) within EECO
200	5. 0.9 ‰ at 16.8 m (C23n) within EECO
201	6. 0.4 ‰ at 14.8 m (C24n.1n) within EECO (X event)
202	7. 0.3 ‰ at 12.5 m (near C24n.2n/C24n.2r boundary; J event)
203	8. 0.3 ‰ at 10.5 m (mid C24n.3n; I event)
204	9. 0.3 ‰ at 8.0 m (upper C24r; ETM2/ELMO event)
205	The two oldest of these carbon isotope excursions are determined using 45-50 cm sample spacing,
206	implying that their true magnitudes are probably not fully captured. Their precise positions may also
207	change as higher resolution data become available from this relatively condensed part of the section
208	showing sedimentation rates <0.5 cm/kyr. The remaining five are determined using 20 cm sample
209	spacing. The number, magnitudes and stratigraphy of the above carbon isotope excursions are similar
210	to the results of other studies (Agnini et al., 2009; Slotnick et al., 2012; Zachos et al., 2010).

Above Chion C221, a series of additional minor negative carbon isotope excursions (CIES) at	211	Above Chron C22r, a series of additional minor negative carbon isotope excursions (CIEs) are
---------------------------------------------------------------------------------------------	-----	----------------------------------------------------------------------------------------------

- recorded in Chron C22n, Chron C21r and Chron C21n from the Possagno section. By combining the
- 213 chron identification with the number of CIEs starting at the old end of the chron, these CIEs are coined
- 214 C22n-CIE1, C22n-CIE2, etc., up to C21n-CIE4 (Fig. 2). We have tentatively named the isotope shifts
- 215 of small magnitude as events only when changes in isotopic composition are associated with evident
- 216 modifications in planktic foraminferal assemblages and/or in fragmentation index. This is because
- 217 increase in both fragmentation index and some foraminiferal taxa is similar to the record observed
- during early Eocene hyperthermals from the same geological setting (Luciani et al. 2007; Agnini et al.,
- 219 2009). As the Possagno section is measured from the base of the PETM (0 m) and upsection,
- 220 increasing distance from the PETM level yields increasing positive meter values:
- 221 1. C21n-CIE4 0.3% from 56.6 m to 57.0 m
- 222 2. C21n-CIE3 0.3 ‰ from 55.6 m to 56.2 m
- 223 3. C21n-CIE2 0.3% from 54.8 m to 55.0 m
- 224 4. C21n-CIE1 0.8 ‰ from 48.8 m to 49.4 m
- 225 5. C21r-CIE4 0.3 ‰ from 39.6 m to 39.8 m
- 226 6. C21r-CIE3 0.5 ‰ from 38.8 m to 39.2 m
- 227 7. C21r-CIE2 0.7 ‰ from 37.6 m to 38.2 m
- 228 8. C21r-CIE1 0.9 ‰ from 32.8 m to 33.2 m
- 229 9. C22n-CIE3 0.5 ‰ from 31.2 m to 31.4 m
- 230 10. C22n-CIE2 0.5 ‰ from 30.0 to 30.2 m
- 231 11. C22n-CIE1 0.6 ‰ from 27.2 m to 27.4 m

Oxygen isotopes of bulk rock measurements from indurate marly limestones and limestones may be affected by diagenetic overprints (Marshall, 1992), which presumably apply also to the rocks in the Possagno section. Despite of this preservation caveat, it is assumed that oscillations in oxygen isotopes chiefly represent temperature fluctuations during the ice-free early Eocene world. Lighter oxygen isotope values in the Possagno section indeed show a clear correspondence with lighter carbon isotope values (CIEs) and vice versa in the post-EECO interval (Fig. 2). Thus, despite the possibility of some diagenetic overprinting in several individual samples, especially in the lower part of our record, a three-

- 239 point moving average of oxygen isotope data should reveal early to early middle Eocene climate
- 240 variability in the Possagno section.

242 **4.4 Planktic foraminiferal quantitative analysis**

243

244 The planktic foraminiferal assemblages show significant modifications in the early to early 245 middle Eocene interval at Possagno (Fig. 3). The mean relative abundance of Acarinina is about 46 % 246 of the total assemblage throughout the section. Members of this genus show peak abundances of 60-70 247 % of the total assemblage during the early to early middle Eocene CIEs. Particularly prominent is the 248 increase to ~80 % during the EECO interval (Fig. 3). Acarininids clearly thrived and expanded in 249 abundance during the CIEs, including the EECO. 250 This increase of acarininids is counter balanced by a transient decrease in members of 251 subbotinids. This latter group recovers above the EECO interval and increases moderately from ~24 % 252 to \sim 36 % in terms of mean relative abundance of the total assemblage, up to the top of the section. The 253 North Atlantic Site 1051 also shows a slight increase of ca. 7 % in the mean value among the 254 subbotinids during the corresponding time interval. 255 A permanent reduction in the abundance of members of the genus Morozovella represents a

256 major change within the planktic foraminiferal assemblages within Zone E5. This group collapses from 257 a mean value of ~24 % in the 0-15 m interval to less than 6 % above 15 m. Qualitative examination of 258 species variability shows that, in the lower part of the Zone E5 where the greater morozovellids 259 abundance is recorded, no dominance of particular species is recognized, even though M. 260 marginodentata, M. subbotinae and M. lensiformis are relatively more common forms with respect to 261 M. aequa, M. aragonensis, M. formosa and M. crater. In the interval with the low abundance of 262 morozovellids within the EECO, an overturn is observed since M. aragonensis, M. formosa, M. crater 263 and, in the upper part of the E5 Zone, *M. caucasica* are the most common species.

264 Morozovellids never recover to their pre-EECO abundances, even if including the appearance of 265 the ecologically comparable genus *Morozovelloides* (Pearson et al., 2006) in samples above 36 m.

266 Genera and species with low abundances show minor changes throughout the interval studied at267 Possagno (Fig. S1).

268 The major change in planktic foraminiferal assemblages at Site 1051 includes a distinct decrease of

269 *Morozovella*, from mean values around 40 % to 10 % in the middle part of Zone E5 (Fig. 4). Similarly

to Possagno, the lower part of Zone E5 with the higher percentages of morozovellids does not record

271	the dominance of selected species, but at Site 1051 M. aragonensis and M. formosa besides M.
272	subbotinae are relatively common whereas M. marginodentata is less frequent. Within the interval of
273	low morozovellid abundances, M. aragonensis and M. formosa are the most common taxa. The general
274	decline of morozovellids does not appear therefore related, both at Possagno and Site 1051, to the
275	extinction or local disappearance of a dominant species.
276	The differences in the relative abundance of morozovellids at Possagno and Site 1051 indicate a
277	morozovellid preference for open ocean settings of low-latitudes, as suggested also by other authors
278	(Berggren, 1978; Boersma et al., 1987; Premoli Silva and Boersma, 1988). Like at Possagno,
279	morozovellids never recover at Site 1051 in the Zone E5 through E8 interval. The abundance of
280	subbotinids shows little variations around mean values of 20 % at Site 1051. Acarinina displays an
281	increase in mean relative abundance from 35 % (base to ca. 450 mbsf) to around 50 % (ca. 430 mbsf),
282	with maximum values of about 60 %. The relatively low resolution used here does not permit
283	comparison between the early Eocene CIEs at Site 1051 (Cramer et al., 2003) and how the relative
284	abundance of planktic foraminiferal genera varies with respect to CIEs.
285	
286	4.5 Radiolarian abundance
287	
288	Radiolarians are rare to absent in the Possagno section. Brief temporary occurrences of this
289	group have been observed in coincidence with some of the most negative $\delta^{13}C$ excursions. Specifically,
290	they reach a maximum relative abundance of 28 % in the lower part of the major $\delta^{13}C$ perturbation
291	recorded in the lower to middle part of C23n, of ~10 % at 27.5 m and of 15 % at 31.4 m (Fig. 3). At
292	Site 1051, radiolarians fluctuate in abundance from 0 to 78 % throughout the studied interval.
293	
294	
295	
	5. Discussion
296	5. Discussion
296 297	 5. Discussion 5.1 The δ¹⁸O and δ¹³C stratigraphies at Possagno
296 297 298	5. Discussion 5.1 The $\delta^{18}O$ and $\delta^{13}C$ stratigraphies at Possagno
296 297 298 299	5. Discussion 5.1 The δ^{18} O and δ^{13} C stratigraphies at Possagno The δ^{18} O and δ^{13} C records from the Possagno section display both details and trends (Fig. 2) that

301 stratigraphies (Cramer et al., 2009; Schmitz et al., 1997; Shackleton et al., 1985; Slotnick et al., 2012; 302 Zachos et al., 2001; 2008). For example, the major CIE (-1.5 ‰) close to the base of the Possagno 303 section represents the PETM, followed upsection by hyperthermals ETM2, I, J, and ETM3 (Agnini et 304 al., 2009). The initiation and termination of the EECO are not well constrained in any single 305 sedimentary record (Slotnick et al., 2012), nor so at Possagno, yet the interval between 16 m and 22.5 306 m in Possagno shows the lightest δ^{18} O values during the entire post-PETM interval of the early Eocene 307 and early middle Eocene (Fig. 2) and is considered to represent at least part of the EECO. In terms of 308 oxygen isotope stratigraphy, both the EECO and post-EECO intervals are characterized by a series of 309 rapid oscillations. The oxygen isotope amplitude range shows up to 1.5 ‰ differences between 310 adjacent samples, which possibly may reflect potential diagenetic overprint. By running a 3-point 311 running mean of the oxygen isotope data, single overprint outliers are dampened. Even so, these data 312 show rapid amplitude changes of up to $0.5 \,$ %, suggesting correspondingly rapid temperature changes 313 in the western Tethys on the order of 2°C during the ice-free early and early middle Eocene world. The 314 underlying cause of these distinct and rapid temperature changes may be sought in the stable carbon 315 isotope data. 316 Several lines of evidence suggest that high CO₂ concentrations were driving the EECO global 317 warmth as well as the hyperthermal events of the early Eocene (Fletcher et al., 2008; Hyland and 318 Sheldon, 2013; Komar et al., 2013; Lowenstein and Demicco, 2006; Lunt et al., 2011; Pearson and 319 Palmer, 2000; Royer et al., 2007; Smith et al., 2010). A series of CIEs occur within and above the 320 EECO interval at Possagno. A number of CIEs have been observed from ODP Site 1258 in the western 321 tropical Atlantic (Kirtland Turner et al., 2014; Sexton et al., 2006; 2011), which they interpreted as 322 minor hyperthermals and referred to as 21 numbered H-events (H for Hyperthermal) in Chrons C23r 323 through C21r. Sedimentation rates at Possagno are 2-6 times lower than those at Site 1258 in the 324 indentical time interval. The number of CIEs within individual magntochron zones at Possagno and 325 Site 1258 differ slightly, presumably because of differences in sedimentation rates and sample 326 resolution. Kirtland Turner et al. (2014) listed three CIEs/H-events in Chron C23r, none of which are 327 evident in the Possagno record at the present sample resolution, probably due to strongly condensed 328 sedimentation or presence of a hiatus. At Possagno, three events are recorded in C23n, compared with 329 the two CIEs/H-events from Demerara Rise, whereas only two of the six CIEs/H-events in Chron C22r 330 are distinguishable in Possagno. Five CIEs/H-events are listed in Chron C22n from Demerara Rise,

three of which are evident in the Possagno record. Finally, five CIEs/H-events are listed in Chron C21r,

four of which are evident in the Possagno record. It follows that the lower sample resolution at

333 Possagno likely has blurred both the number and true magnitudes of the isotopic amplitude changes. It

- remains uncertain if the CIEs and H-events at Site 1258 and the CIEs in the Possagno section are in
- 335 synchrony (Tab. 1).

Regardless of the potential synchrony between Possagno and Site 1258 at the Demerara Rise,

both regions clearly demonstrate that after the EECO, from ca. 50 Ma to ca. 46-47 Ma during a trend of

338 cooling (2-2.4°C) climate, about 18 brief negative CIEs coincide with hyperthermal-like brief episodes

339 of warming. These brief CIEs induced environmental perturbations that are expressed in the planktic

340 for a primary relationship between increased

- 341 CO₂ concentration and warmth during early through early middle Eocene times (Dickens et al., 2005;
- 342 Quillévéré et al., 2008; Zachos et al., 2005; 2008;).
- 343

5.2 The EECO interval and hyperthermals at Possagno: Acarinina dominated or dissolutioncontrolled assemblages?

346

347 One of the most prominent changes in the planktic foraminiferal assemblages is the dominance of 348 acarininids across the EECO interval, resulting in a corresponding decrease in relative abundance of 349 morozovellids and subbotinids (Fig. 3). These changes parallel relatively high values of the F index and coincide with the most negative parts in the δ^{13} C record. The increase in shell fragmentation 350 351 suggests some carbonate dissolution. The high pCO_2 atmospheric concentration during the EECO may 352 have induced carbonate dissolution at the deep-water Possagno setting, resulting from deep-water 353 acidification and a rise of the lysocline, similar to patterns observed during the main hyperthermal 354 events. This interpretation for the Possagno section should however imply a considerable shallowing of 355 the CCD/lysocline, assuming that the section was deposited in a middle to lower bathyal setting. 356 Further studies on the Tethyan CCD in the Eocene interval will help investigate the hypothesis that the 357 CCD was shallower there, with respect to the open ocean CCD. Intensified water column 358 remineralization of organic matter, forced by augmented metabolic rates at elevated temperatures, may 359 have caused pH to decrease in the uppermost water column, inducing dissolution of calcitic tests 360 (Brown et al., 2004; John et al., 2013, 2014; O'Connor et al., 2009; Olivarez Lyle and Lyle, 2006).

361 Questions arise on how to estimate the possible dissolution artefacts from the primary ecological 362 signal. Nguyen et al. (2011) and Petrizzo et al. (2008) studied Pacific Ocean assemblages of latest 363 Paleocene to initial Eocene age, and suggested that subbotinids are more dissolution susceptible than 364 morozovellids and acarininds, which previously were thought to be the most dissolution prone forms 365 (Berggren and Norris, 1997; Boersma and Premoli Silva, 1983). Paleogene assemblages affected by 366 extensive dissolution could be expected to be impoverished with respect to the more dissolution 367 susceptible subbotinids. These results have been challenged by other analyses that document a 368 dominance of subbotinids within intervals affected by a high F index and enhanced carbonate 369 dissolution (Luciani et al., 2010), who suggested that dissolution has affected the planktic assemblages 370 rather equally. The degreee of dissolution of planktic foraminifera appears to have varied during 371 different time intervals, being species related rather than exclusively associated with different genera. 372 However, since data on dissolution susceptibility on different genera are so far lacking for early and 373 early middle Eocene times, we cannot exclude that dissolution may have changed the planktic 374 foraminiferal assemblages.

When assuming that dissolution has affected assemblages, it follows that the dominance of acarininids during the EECO and hypertermal events may represent a taphonomic artifact. This assumption appears yet to conflict with the results from the upper part of Possagno in the Chron C21n interval, where significant decreases of subbotinids, associated with distinct acarininid increases, correspond to negative shifts in δ^{13} C values in the absence of carbonate dissolution, as expressed in low *F* index values (Fig. 3).

381 The similarity in the major planktic foraminiferal modifications throughout the EECO at Site 382 1051(Fig. 4), which appears only marginally affected by dissolution, suggests that the Possagno 383 assemblages represent a reasonably genuine paleoecological response rather than assemblages 384 primarily modified by carbonate dissolution.

385 The decrease of CF values (Fig. 3) in the EECO interval might indicate loss of carbonate shells
386 due to carbonate dissolution. Similarly, relatively low CF-values with only minor fluctuations are

387 recorded to the top of the section, independently from changes in F index values. The CF curve

388 parallels the EECO/post-EECO trend of the morozovellid abundance thus suggesting a relationship

389 with the morozovellid decline rather than carbonate dissolution.

390

5.3 The Possagno and Site 1051 records: planktic foraminiferal response to the EECO

392

393	The planktic foraminiferal assemblages show significant variations in the Possagno material that
394	correlate with the pronounced δ^{13} C perturbations in the EECO interval (Fig. 3). When the warm
395	preferring acarininds become dominant during the EECO, this results in a reduction in relative
396	abundance of the warm preferring morozovellids. This feature is recurring in planktic foraminiferal
397	assemblages across some hyperthermals (PETM and X events), as recorded from a number of Tethyan
398	successions of northeastern Italy and it has been interpreted as a result of relatively enhanced
399	eutrophication of surface waters in a near continental setting (Agnini et al., 2009; Arenillas et al., 1999;
400	Luciani et al., 2007; Molina et al., 1999).
401	Planktic foraminiferal analyses of the pre-EECO hyperthermals ETM2, I, J and ETM3 at Possagno
402	show PETM-like responses, consisting of strongly increasing relative abundances of acarininids, as in
403	the PETM interval of the nearby Forada section (Luciani et al., 2007). The multi-proxy analyses of the
404	X-event at the nearby Farra section (Agnini et al., 2009) corroborate at higher resolution the record
405	from Possagno. Increased surface water eutrophication has been proposed to favour acarininids, in
406	being able to temporarily colonize deeper waters that previously were occupied by subbotinids and in
407	being able to tolerate relatively high eutrophic conditions that suppressed the abundances of
408	morozovellids (Agnini et al., 2009; Luciani et al., 2007). Slight differences in paleobiology between
409	morozovellids and acarininids are documented in several cases by minor variations in stable isotopes
410	that commonly indicate a more surface habitat for the former group (Boersma et al., 1987; Pearson et
411	al., 1993; 2001).
412	The increased surface water eutrophication during hyperthermals was forced by strengthening of
413	the hydrological cycle and increased weathering as a consequence to enhanced greenhouse conditions.
414	The effects improved the nutrient availability in this near continental, pelagic setting of the western
415	Tethys (Agnini et al., 2009: Giusberti et al., 2007; Luciani et al., 2007). The hypothesis of increased
416	nutrient availability in the lower part of the EECO interval at Possagno is supported by the entry of
417	relatively high concentration of radiolarians, considered as eutrophic indices (Hallock, 1987).
418	The decline of morozovellids across the EECO at Possagno and Site 1051 is irreversible and

cannot be explained by brief perturbations, as during the pre-EECO hyperthermals. The morozovellidcrisis is coupled with the gradual disappearances of several species, including *M. aequa, M. gracilis,*

M. lensiformis, M. marginodentata, and *M. subbotinae*, and it is not counterbalanced by the appearance
of species of *Morozovelloides*, a minor component of middle Eocene assemblages. The latter genus
appeared at Possagno around the Ypresian/Lutetian boundary (Luciani and Giusberti, 2014) and it is
morphologically highly convergent with *Morozovella* although probably did evolve from *Acarinina*(Pearson et al., 2006).

426 The similar behaviour across the EECO of morozovellids in the Tethyan Possagno section and 427 Site 1051 in the western subtropical North Atlantic supports the hypothesis of a geographically wide 428 spread morozovellid crisis that is caused by climate change. This change must be a consequence of the 429 major modifications across the EECO, both in terms of temperature and pCO_2 , which in turn must have 430 induced water column reorganizations leading to a reduction of the morozovellid habitat. Because 431 morozovellids exhibit transient reduction in abundance during pre-EECO hyperthermals, and due to the 432 imprecise definition of this event, it is not possible to precisely pinpoint the exact turning point of the 433 morozovellid decline, i.e. whether it began just at the onset, within, or at the termination of the EECO 434 event. Current data from Possagno and Site 1051 however record that their massive drop in abundance 435 began across the C24n1n-C23r transition. The decrease apparently started at the top of C24n1n at 436 Possagno, but it has not been possible to determine whether or not this decrease is transitory because a 437 potential recovery may be hidden by condensation/hiatus across the C24n1n-C23r interval. Present data 438 from Site 1051 record the decline in lower C23r, even though some uncertainties are caused by the low 439 resolution in the foraminiferal analysis and magnetostratigraphic attribution.

440 In contrast to the deterioration of the morozovellid habitat, relatively favourable conditions for 441 thermocline dwellers such as subbotinids and parasubbotinids are suggested by the new species 442 appearing progressively during the post-EECO interval at Possagno (Luciani and Giusberti, 2014), in 443 good agreement with the low latitude data presented by Pearson et al. (2006). Most of the new species 444 will characterize the thermocline of the middle and late Eocene oceans: Subbotina corpulenta, S. 445 eocena, S. hagni, S. senni, S. yeguanesis, Parasubbotina griffinae, and P. pseudowilsoni. The 446 appearance of the radially chambered *Parasubbotina eoclava*, which is considered to be the precursor 447 of the truly clavate chambered *Clavigerinella* (Coxall et al., 2003; Pearson and Coxall, 2014) occurs at 448 19.8 m (Luciani and Giusberti, 2014). Clavigerinella is the ancestor of the genus Hantkenina that 449 successfully inhabited the sub-surface middle through late Eocene oceans. Prior to the evolution of 450 genuine Clavigerinella, P. eoclava made several aborted attempts to evolve towards the genus

451 Clavigerinella. This is reflected in presence of rare and scattered specimens that are morphologically

452 close to *Clavigerinella* within the EECO interval even though true representative of genus

453 Clavigerinella were not observed in the Possagno section.

454 The EECO and post-EECO intervals indeed proved to be crucial in Eocene planktic foraminferal 455 evolution.

- 456
- 457

458

5.4 Possible causes of morozovellids decline across the EECO 459 The data from Possagno and Site 1051 demonstrate that the early Paleogene planktic 460 foraminiferal symbiont bearing groups were strongly affected by a habitat deterioration across the 461 EECO. The early Eocene crisis was followed by a second step, involving the large sized acarininids 462 and Morozovelloides, documented through their reduction in abundance as well as test size during the 463 Middle Eocene Climate Optimum (MECO) at ca. 40 Ma in Tethyan (Fig. 5), Southern Ocean, and 464 northwest Atlantic settings (Edgar et al., 2012; Luciani et al., 2010). Furthermore, the muricate crisis 465 culminates near the Bartonian/Priabonian boundary with a major demise in the Acarinina lineage and 466 the extinction of *Morozovelloides*. Only small (<125 µm) and relatively rare acarininids survived into 467 the late Eocene and Oligocene (Agnini et al., 2011; Berggren et al., 2006; Wade, 2004; Wade et al., 468 2012). An episode with loss of symbiosis resulting in bleaching caused by global warming has been 469 proposed to explain the second muricate crisis (Edgar et al., 2012). If the MECO warmth has been the 470 main cause of bleaching of acarininids, we would expect that this phenomenon also involved 471 morozovellids during the EECO, as this warm interval records the highest temperatures of the 472 Paleogene. Considering the importance of photosymbiosis in extant species for foraminiferal test 473 calcification and ecology (Bé, 1982; Bé et al., 1982; Hemleben et al., 1989), we may assume similar 474 requirements for fossil taxa. The algal-symbiotic relationship is considered one of the most successful 475 strategies adopted by muricates during the earliest Paleogene (Norris, 1996; Quillévéré et al., 2001). A 476 crisis in that relationship may represent one possible hypothesis to explain the decline of early Eocene 477 morozovellids. Further studies that include stable isotope analyses, including δ^{13} C gradients, on 478 morozovellid tests are needed to further elucidate this scenario. There is however scarce documentation

- 479 on mechanisms responsible for bleaching and besides elevated sea surface temperature, a number of
- 480 other factors, for example high ultraviolet radiation, decrease in pH decrease, increase in CO₂, changes

481 in salinity and nutrient availability, may have been involved (Douglas et al., 2003; Wade et al., 2008)). 482 The protracted exceptional warming of the EECO is expected to have increased metabolic rates 483 particularly in heterotrophs that are more sensitive to temperature than rates of primary production 484 (Brown et al., 2004; O'Connor et al., 2009; Olivarez Lyle and Lyle, 2006). This effect may have 485 increased water column remineralization of organic matter and caused pH to decrease in the uppermost 486 water column (John et al., 2013; 2014; O'Connor et al., 2009; Olivarez Lyle and Lyle, 2006). The 487 potential lowering of pH in surface waters, which could have been further enhanced by the huge EECO 488 CO₂ pressure, may have affected calcification (De Moel et al., 2009; Moy et al., 2009; Zeebe et al., 489 2008) of most surface dwellers such as the morozovellids, also by reducing or completely halting their 490 symbiont relationships. 491 Complex interactions with other microfossil groups, such as radiolarians, diatoms or

dinoflagellates, may have contributed to the morozovellid crisis across the EECO, for example by
competing for the use of the same algal-symbionts in the case of radiolarians, or symbiont-providers.
Detailed comparisons of trends in other fossil groups are necessary to investigate this hypothesis.

495 Seawater chemistry influences the biomineralization of organisms producing CaCO₃ skeletons, 496 especially for many algae and invertebrates that have less control over the chemical composition of 497 their mineralized parts (Stanley, 2006; 2008). High magnesium/calcium ratios are known to have 498 favoured aragonitic and high-Mg calcite skeletons throughout the Phanerozoic. This insight is 499 corroborated by experiments with living organisms, confirming, for example, population growth 500 among the calcitic coccolithophores in conditions of low concentration of Mg and high concentration 501 of calcium in seawater (Stanley et al., 2005). A strong reduction in Ca concentration occurred during 502 the Cenozoic, following the 'calcitic' Cretaceous ocean, possibly driven by changes in rates of deep-503 sea igneous activity (Hardie, 1996). We cannot exclude that a decrease of Ca concentration in seawater 504 chemistry may have affected morozovellid calcification. Planktonic foraminifera have not been widely 505 employed to study the effects of the Mg/Ca ratio of the seawater on calcification, however, they have 506 been found to produce heavier skeleton when the saturation state of the ambient seawater with respect 507 to calcite is elevated. It would be interesting to compare flux data of calcareous nannofossils before and 508 after the major evolutionary change recorded across the EECO (Agnini et al., 2006; Schneider et al., 509 2011) to test a potential reduction in their overall productivity.

510 Extended time intervals of weak water column stratification and increased eutrophication are

511 known to provide hostile ecological conditions for the highly specialized oligotrophic morozovellids 512 (Boersma et al., 1987; Bralower et al., 1995; Pearson et al., 2006; Premoli Silva and Boersma, 1989). 513 Such conditions are documented in several ocean sites by the recorded decline in surface-to-benthic 514 δ^{13} C gradients (Hilting et al., 2008) and have been considered linked to evolutionary turnovers among 515 calcareous nannofossil assemblages (Schneider et al., 2011). Weakened thermal stratification with 516 increased vertical mixing is predicted for many, although not all, oceanic areas during hyperthermals. 517 The fact that the permanent morozovellid collapse occurs during the EECO implies that a threshold 518 was surpassed, not previously experienced by the morozovellid communities.

519 Available data indicate that the protracted conditions of extreme warmth and high CO₂ pressure 520 during the EECO may have been the key element inducing a permanent impact in the marine surface 521 water ecosystem that became detrimental for the morozovellids. Even the peculiar PETM, that records 522 the most dramatic changes among the hyperthermals both in terms of temperature increase and carbon 523 cycle perturbation, did not adversely affected the morozovellid habitat in a permanent way. On the 524 contrary, morozovellids increased in abundance in open oceanic settings (Kelly et al., 1996; 1998, 525 2002; Lu and Keller, 1993; Petrizzo, 2007), and only a transient decrease in abundance is recorded in 526 pelagic Tethyan near continental settings (Luciani et al., 2007). It is intriguing to note that the second 527 main crisis of the muricate symbiont bearing forms occurred during the MECO (Fig. 5), that is also a 528 warming event of much longer duration (about 400-500 kyr) than the early Paleogene hyperthermals 529 (Bohaty et al., 2009; Westerhold and Röhl, 2013).

530

531 5.5 Planktic foraminiferal changes during post-EECO stable-isotope perturbations at Possagno532

533 The small δ^{13} C excursions recorded in the post-EECO interval at Possagno, from C22n to C21n,

induced perturbations on the planktic foraminiferal assemblages that mirror those recorded in the pre-

535 EECO interval (Fig. 3). These perturbations are expressed as marked increases of acarininids,

536 representing warmer thriving taxa that were tolerant to relatively enhanced surface water eutrophic

537 conditions. Peaks in surface water eutrophication could have been reached during the C22n-CIE1 and

538 C22n-CIE3 events, as indicated by the relatively high production of radiolarians.

539 The post-EECO CIEs are concomitant with δ^{18} O excursions and coupled to distinct modifications 540 in the planktic foraminiferal assemblages comparable to those recorded at other early Eocene brief

541	warming events in Tethyan settings. These data make it tenable to refer to these events as
542	hyperthermals, although of less intensity and magnitude compared to some of the pre-EECO
543	hyperthermals. A number of these δ^{13} C and δ^{18} O excursions, of much smaller magnitude and intensity
544	with respect to the PETM, probably correlate with the recently discovered late early Eocene through
545	early middle Eocene post-EECO hyperthermals in the Atlantic and Pacific (Kirtland Turner et al.,
546	2014; Sexton et al., 2006; 2011; Tab.1). According to Sexton et al. (2011) and Kirtland Turner et al.
547	(2014) their shorter duration and more rapid recovery phases, with respect to the PETM, imply a
548	different forcing and feedback mechanism involving redistribution of carbon among the ocean
549	reservoirs rather than release of greenhouse gases from buried sediments. This mechanism was active
550	also during the EECO interval and was similar to the orbital forcing of the carbon cycle operating
551	during the Oligocene and Miocene.
552	
553	
554	6 Summary and conclusions
555	
556	The investigation of planktic foraminifera from the Possagno section in northern Italy represents a
557	first case history recording changes in relative abundance among planktic foraminiferal assemblages
558	across the EECO warm interval and post-EECO climatic variability in the Tethys realm from about 55
559	Ma to 46 Ma.
560	The most crucial change emerging from the Possagno and ODP Site 1051 data is the irreversible
561	decline across the EECO of the symbiont bearing genus Morozovella, one of the most important
562	calcifiers of the early Paleogene tropical and subtropical oceans. The Possagno data indicate that the
563	EECO event had a permanent impact on the planktic foraminiferal communities, representing a critical
564	phase in the reorganization of Eocene pelagic ecosystems. Possible causes for this reorganization, that
565	deserve further investigations, include increased eutrophication, changes in ocean chemistry, weak
566	water column stratification, loss of symbionts, complex interaction with other microplankton groups
567	such us radiolarians, diatoms or dinoflagellates that represented possible competitors in the use of
568	symbionts or as symbiont providers. A critical threshold was reached across the EECO, evidently never
569	reached before, which induced unfavourable habitats for continued morozovellid diversification and
570	proliferation but not harsh enough to cause their extinction. This threshold appears to be related to the

571 duration of extreme conditions characterizing the extended warmth during the EECO.

572 Even though several questions still remain to be answered, the data presented from Possagno add 573 significant information about the complex evolution of the muricate planktic foraminifera and stimulate 574 additional investigations across the EECO from different paleoceanographic settings.

The post-EECO interval at Possagno is punctuated by relatively small, negative δ^{13} C shifts. 575 576 interpreted as potential minor hyperhthermals in the light of available oxygen isotope data, a number of 577 which probably are in synchrony with those recorded in the tropical Atlantic by Sexton et al. (2011) 578 and Kirtland Turner et al. (2014). These modest post-EECO δ^{13} C and δ^{18} O perturbations are associated 579 with significant, though ephemeral, modifications in the planktic foraminiferal communities showing 580 distinct increases of the warm acarininids in a manner similar to their behaviour during pre-EECO 581 hyperthermals in Tethyan settings. Changes in planktic foraminiferal assemblages occur during 582 environmental perturbations associated with minor negative carbon and oxygen isotope excursions, 583 which suggest a pronounced biotic sensitivity to climate change of planktic foraminifera even during 584 the post-EECO interval.

585

586 Acknowledgements. Funding for this research was provided by MIUR/PRIN COFIN 2010-2011,

587 coordinated by D. Rio. V. Luciani was financially supported by FAR from Ferrara University, and L.

588 Giusberti and E. Fornaciari received financial support from Padova University (Progetto di Ateneo

589 GIUSPRAT10 CPDA108242/10). J. Backman acknowledges support from the Swedish Research

590 Council. We are grateful to Domenico Rio who promoted the researches on the "Paleogene Veneto"

and for the fruitful discussion. Members of the "Possagno net", Simone Galeotti, Dennis Kent,

592 Giovanni Muttoni, who sampled the section in 2003, are gratefully acknowledged. We warmly

593 acknowledge the Cementi Rossi s.p.a. and Mr. Silvano Da Roit for fruitful collaboration during

samplings at the Carcoselle Quarry (Possagno, TV). This research used samples and data provided by

the Ocean Drilling Program (ODP). ODP is sponsored by the U.S. National Science Foundation (NSF)

and participating countries under management of Joint Oceanographic Institution (JOI) Inc. We

597 specially thank from the ODP Bremen Core Repository. Finally, we are grateful to the reviewers,

598 B.Wade and R. Speijer, who helped to strengthen the manuscript.

599

eferences

603

604 Possagno, Italy: the calcareous nannofossils response to climate variability, Earth Planet. Sci. Lett., 605 241, 815-830, 2006. 606 607 Agnini, C., Macrì, P., Backman, J., Brinkhuis, H., Fornaciari, E., Giusberti, L., Luciani, V., Rio, D., 608 Sluijs, A., Speranza, F.: An early Eocene carbon cycle perturbation at 52.5 Ma in the Southern Alps: 609 chronology and biotic response, Paleoceanogr., 24, PA2209. doi:10.1029/2008PA001649, 2009 610 611 Agnini, C., Fornaciari, E., Giusberti, L., Grandesso, P., Lanci, L., Luciani, V., Muttoni, G., Plike, H., 612 Rio, D., Spofforth, D. J. A., Stefani, C.: Integrated biomagnetostratigraphy of the Alano section (NE 613 Italy): a proposal for defining the middle-late Eocene boundary, Geol. Soc. Am. Bull., 123, 841-872, 614 2011. 615 616 Arenillas, I., Molina, E., Schmitz, B.: Planktic foraminiferal and δ^{13} C isotopic changes across the 617 Paleocene/Eocene boundary at Possagno (Italy), Int. J. Earth Sc., 88, 352-364, 1999. 618 619 Aubry, M.-P., Ouda, K., Dupuis, C., Berggren, W.A., Van Couvering, J.A., and the Members of the 620 Working Group on the Paleocene-Eocene Boundary .: The Global Standard Stratotype-Section and 621 Point (GSSP) for the base of the Eocene Series in the Dababiya section (Egypt)., Episodes 30, 271– 622 286, 2007. 623 624 Be', A.W.H.: Biology of planktonic foraminifera, in: Foraminifera: notes for a short course, Broadhead 625 T., Stud. Geol., 6, Univ. Knoxville, Tenn., 51-92, 1982. 626

Agnini, C., Muttoni, G., Kent, D.V., Rio, D.: Eocene biostratigraphy and magnetic stratigraphy from

- Be', A.W.H., Spero, H.J., Anderson O.R.: Effects of symbiont elimination and reinfection on the life
 processes of the planktonic foraminifer *Globigerinoides sacculifer*, Marine Biol. 70, 73–86, 1982.
- 629
- 630 Berggren, W.A.: Rates of evolution in some Cenozoic planktonic foraminifera, Micropaleont. 15, 337-

631 370, 1978.

632

- Berggren, W.A., Norris, R.D.: Biostratigraphy, phylogeny and systematics of Paleocene trochospiral
 planktic foraminifera, Micropaleont., 43 (Suppl. 1), 1–116, 1997.
- 635
- 636 Berggren, W. A., Pearson, P. N., Huber, B.T., Wade, B. S.: Taxonomy, Biostratigraphy and Phylogeny
- 637 of Eocene Acarinina, In: Atlas of Eocene Planktonic Foraminifera, Cushman Foundation Spec. Publ.
- 638 41, Pearson, P. N., Olsson, R. K., Huber, B.T., Hemleben, C., Berggren, W. A., 257–326, 2006.
- 639
- 640 Boersma, A., Premoli Silva, I.: Paleocene planktonic foraminiferal biogeography and the
- 641 paleoceanography of the Atlantic-Ocean, Micropaleont., 29, 4, 355–381, 1983.
- 642
- 643 Boersma, A., Premoli Silva, I., Shackleton, N.: Atlantic Eocene planktonic foraminiferal biogeography
- and stable isotopic paleoceanography, Paleoceanogr., 2, 287–331, 1987.
- 645
- 646 Bohaty, S.M., Zachos, J.C., Florindo, F., Delaney, M.L.: Coupled greenhouse warming and deep-sea
- acidification in the Middle Eocene, Paleoceanogr., 24, PA2207, doi:10.1029/2008PA001676, 2009.
- 648
- 649 Bosellini, A.: Dynamics of Tethyan carbonate platform, in: Controls on Carbonate Platform and Basin
- 650 Platform, Crevello, P.D., Wilson, J.L., Sarg, J.F., Read, J.F., SEPM Spec. Publ., 44, 3–13, 1989.
- 651
- Bralower, T.J., Zachos, J.C., Thomas, E., Parrow, M., Paull, C.K., Kelly, D.C., Premoli Silva, I., Sliter,
- 653 W.V., Lohmann, K.C.: Late Paleocene to Eocene paleoceanography of the equatorial Pacific Ocean:
- 654 Stable isotopes recorded at Ocean Drilling Program Site 865, Allison Guyot, Paleoceanogr., 10, 841–
- 655 865, 1995.

- Brown, J.H., Gillooly, J.F., Allen, A.P., Savage, V.M., and West, G.B.: Toward a metabolic theory of
 ecology, Ecology, 85(7), 1771–1789, 2004.
- 659
- 660 Cande, S. C., Kent, D. V.: Revised calibration of the geomagnetic polarity timescale for the Late

661 Cretaceous and Cenozoic, J. Geophys. Res., 100, 6093–6095, 1995.

- 663 Cita, M. B.: Stratigrafia della Sezione di Possano. In Bolli, H. M. (Ed.), Monografia
 664 Micropaleontologica sul Paleocene e l'Eocene di Possagno, Provincia di Treviso, Italia, Schweiz.
 665 Palaeontol. Abhandl., 97, pp.9–33, 1975.
- 666
- 667 Coccioni, R., Bancalà, G., Catanzariti, R., Fornaciari, E., Frontalini, F., Giusberti, L., Jovane, L.,
- 668 Luciani, V., Savian, J., Sprovieri, M.: An integrated stratigraphic record of the Palaeocene-lower
- Eocene at Gubbio (Italy): new insights into the early Palaeogene hyperthermals and carbon isotope
- 670 excursions, Terra Nova, 24, 380–386, 2012.
- 671
- 672 Coxall, H. K., Wilson, P.A; Pälike, H.; Lear, C. H; Backman, J.: Rapid stepwise onset of Antarctic
 673 glaciation and deeper calcite compensation in the Pacific Ocean. Nature, 433, 53-57,
 674 doi:10.1038/nature03135, 2005.
- 675
- 676 Cramer, B.S., Wright, J.D., Kent, D.V., Aubry, M.-P.: Orbital climate forcing of δ^{13} C excursions in the 677 late Paleocene–early Eocene (chrons C24n–C25n), Paleoceanogr., 18, 21-1. 678 doi:10.1029/2003PA000909, 2003.
- 679
- Cramer, B.S., Toggweiler, J.R., Wright, M.E., Katz, J.D., Miller, K.G.: Ocean overturning since the
 Late Cretaceous: Inferences from a new benthic foraminiferal isotope compilation, Paleoceanogr., 24,
 PA4216, doi:10.1029/2008PA001683, 2009.
- 683
- 684 Coxall, H.K., Huber, B.T., Pearson, P.N.: Origin ad morphology of the Eocene planktic foraminifera
 685 *Hantkenina*, J. Foram. Res., 33, 237-261, 2003.
- 686
- 687 Coxall, H. K., Wilson, P.A; Pälike, H.; Lear, C. H; Backman, J.: Rapid stepwise onset of Antarctic
 688 glaciation and deeper calcite compensation in the Pacific Ocean. Nature, 433, 53-57,
 689 doi:10.1038/nature03135, 2005.
- 690

- 691 De Moel, H., Ganssen G.M., Peeters F.J.C., Jung, S.J.A., Kroon, D., Brummer, G.J.A., Zeebe, R.E.:
- 692 Planktic foraminiferal shell thinning in the Arabian Sea due to anthropogenic ocean acidification?
- 693 Biogeosciences, 6, 1917-1925, 2009.
- 694
- 695 Demicco, R.V.: Modeling seafloor-spreading rates through time, Geology 32, 485-488, 2004.
- 696
- 697 Dickens, G.R.: Methane release from gas hydrate systems during the Paleocene–Eocene Thermal
- Maximum and other past hyperthermal events: setting appropriate parameters for discussion, Clim.
 Past., 7(2), 1139–1174. doi:10.5194/cpd-7-1139-2011, 2011.
- 700
- Dickens, G.R., O'Neil, J.R., Rea, D.K., Owen, R.M.: Dissociation of oceanic methane hydrate as a
 cause of the carbon isotope excursion at the end of the Paleocene, Paleoceanogr., 10, 965–971,
 doi:10.1029/95PA02087, 1995.
- 704
- 705 Dickens, G.R., Castillo, M.M., Walker, J.C.G.: A blast of gas in the latest Paleocene: simulating first-
- roder effects of massive dissociation of oceanic methane hydrate, Geology, 25, 259–262, 1997.
- 707
- 708 Dickens G.R., Lee C.A., CIA Operatives: Continental island arc fluctuations through time and the
- Eocene transition from a greenhouse to an icehouse world, Rend. Online Soc. Geol. It., 31, 62-63. doi:
- 710 10.3301/ROL.2014.46, 2014.
- 711
- 712 Douglas, A.E.: Coral bleaching- how and why? Mar. Pollut. Bull., 46, 385–392, 2003.
- 713
- 714 Edgar, K.M., Bohaty, S.M., Gibbs, S.J., Sexton, P.F., Norris, R.D., Wilson, P.A.: Symbiont 'bleaching'
- in planktic foraminifera during the Middle Eocene Climatic Optimum, Geology, 41, 15-18,
 doi:10.1130/G33388.1, 2012.
- 717
- 718 Falkowski, P.G., Katz, M.E., Milligan, A.J., Fennel, K., Cramer, B.S., Aubry, M.P., Berner, R.A.,
- 719 Novacek, M.J., Zapol, W.M.: Evolution: The rise of oxygen over the past 205 million years and the
- revolution of large placental mammals, Science, 309 (5744), 2202-2204, 2005.

70	1
1.1	
1 /.	

721	
722	Figueirido, B., Janis, C.M., Pérez-Claros, J.A., De Renzi, M., Palmqvist, P.: Cenozoic climate change
723	influences mammalian evolutionary dynamics, Proc. Natl. Acad. Sci. USA, 109(3), 722-727, 2012.
724	
725	Fletcher, B.J., Brentnall, S.J., Anderson, C.W., Berner, R.A., Beerling, D.J.: Atmospheric carbon
726	dioxide linked with Mesozoic and early Cenozoic climate change, Nat. Geosci., 1, 43-48, 2008.
727	
728	Fornaciari, E., Giusberti, L., Luciani, V., Tateo, F., Agnini, C., Backman, J., Oddone, M., Rio, D.: An
729	expanded Cretaceous-Tertiary transition in a pelagic setting of the Southern Alps (central-western
730	Tethys), Palaeogeogr. Palaeoclimatol. Palaeoecol., 255, 98–131, 2007.
731	
732	Giusberti, L., Rio, D., Agnini, C., Backman, J., Fornaciari, E., Tateo, E., Oddone, M.: Mode and tempo
733	of the Paleocene–Eocene thermal maximum in an expanded section from the Venetian pre-Alps, Geol.
734	Soc. Am. Bull., 119, 391–412, 2007.
735	
736	Hallock, P.: Fluctuations in the trophic resource continuum: a factor in global diversity cycles?
737	Paleoceanogr., 2, 457–471, 1987.
738	
739	Hancock, H.J.L., Dickens, G.R.: Carbonate dissolution episodes in Paleocene and Eocene sediment,
740	Shatsky Rise, west-central Pacific, in: Bralower, T.J., Premoli Silva, I., Malone, M.J., Proc. ODP, Sci.
741	Results 198. http://www-odp.tamu.edu/publications/198_SR/116/116.htm, 2005.
742	
743	Hemleben, C, Spindler, M., Anderson, O.R.: Modern planktonic foraminifera, in: Springer-Verlag,
744	New York, 1-363, ISBN-13: 9780387968155, 1989.
745	
746	Hilting, A.K., Kump, L.R., Bralower, T.J.: Variations in the oceanic vertical carbon isotope gradient
747	and their implications for the Paleocene-Eocene biological pump, Paleoceanogr., 23 (3) (PA3222).
748	DOI: 10.1029/2007PA001458, 2008.
749	
750	Hyland, E.G., Sheldon, N.D.: Coupled CO2-climate response during the Early Eocene Climatic

- 751 Optimum, Palaeogeogr. Palaeoclimatol. Palaeoecol., 369, 125-135, 2013.
- 752
- John E.H., Pearson P.N., Coxall H.K., Birch H., Wade B.S., Foster G.L.: Warm ocean processes and carbon cycling in the Eocene, Phil. Trans. R. Soc., A, 371, 20130099, 2013.
- 755
- John E.H., Wilson J.D., Pearson P.N. Ridgwell A.: Temperature-dependent remineralization and
 carbon cycling in the warm Eocene oceans, Palaeogeogr. Palaeoclimatol. Palaeoecol., 413, 158-166,
 2014
- 759
- Kelly, D.C., Bralower, T.J., Zachos, J.C., Premoli Silva, I., Thomas, E.: Rapid diversification of
 planktonic foraminifera in the tropical Pacific (ODP Site 865) during the late Paleocene thermal
 maximum, Geology 24, 423–426, 1996.
- 763
- Kelly, D.C., Bralower, T.J., Zachos, J.C.: Evolutionary consequences of the latest Paleocene thermal
 maximum for tropical planktonic foraminifera, Palaeogeogr., Palaeoclimatol., Palaeoecol., 141, 139–
 161, 1998.
- 767
- Kennett, J.P., Stott, L.D.: Abrupt deep-sea warming, palaeoceanographic changes and benthic
 extinctions at the end of the Palaeocene, Nature 353, 225–229, 1991.
- 770
- Kirtland Turner S., Sexton P.F., Charled C.D., Norris R.D.: Persistence of carbon release events
 through the peak of early Eocene global warmth, Nat. Geosc., 7, 748-751, DOI: 10.1038/NGEO2240,
 2014.
- 774
- 775 Komar, N., Zeebe, R.E., Dickens, G.R.: Understanding long-term carbon cycle trends: the Late
- Paleocene through the Early Eocene, Paleoceanog., 28, 650-662, doi: 10.1002/palo.20060, 2013.
- 777
- 778 Lee C.T., Shen B., Slotnick B.S., Liao K., Dickens G.R., Yokoyama Y., Lenardic A., Dasgupta R.,
- Jellinek M., Lackey J.S., Schneider T., Tice M.M.: Continental arc-island arc fluctuations, growth of
- rustal carbonates, and long-term climate change, Geosphere, 9, 21-36, 2013.

782	Lirer, F.: A new technique for retrieving calcareous microfossils from lithified lime deposits.
783	Micropaleontol., 46, 365–369, 2000.
784	
785	Littler, K., Röhl, U., Westerhold, T., Zachos, J.C.: A high-resolution benthic stable-isotope for the
786	South Atlantic: implications for orbital-scale changes in Late Paleocene-early Eocene climate and
787	carbon cycling, Earth Planet. Sci. Lett., 401, 18-30. <u>http://dx.doi.org/10.1016/j.epsl.2014.05.054</u> , 2014.
788	
789	Lourens, L.J., Sluijs, A., Kroon, D., Zachos, J.C., Thomas, E., Röhl, U., Bowles, J., Raffi, I.:
790	Astronomical pacing of late Palaeocene to early Eocene global warming events, Nature, 7045, 1083-
791	1087, 2005.
792	
793	Lowestein, T.K. Demicco R.V.: Elevated Eocene Atmospheric CO2 and Its Subsequent Decline,
794	Science, 313 (5795), 1928. DOI:10.1126/science.1129555, 2006.
795	
796	Lu, G., Keller, G.: The Paleocene-Eocene transition in the Antarctic Indian Ocean: inference from
797	planktic foraminifera, Mar. Micropaleontol., 21, 101–142, 1993.
798	
799	Luciani, V., Giusberti, L., Agnini, C., Backman, J., Fornaciari, E., Rio., D.: The Paleocene-Eocene
800	Thermal Maximum as recorded by Tethyan planktonic foraminifera in the Forada section (northern
801	Italy), Mar. Micropaleont., 64, 189–214, 2007.
802	
803	Luciani, V, Giusberti L., Agnini C, Fornaciari E, Rio D, Spofforth D.J.A., Pälike H: Ecological and
804	evolutionary response of Tethyan planktonic foraminifera to the middle Eocene climatic optimum
805	(MECO) from the Alano section (NE Italy), Palaeogeogr. Palaeoclimatol. Palaeoecol., 292, 82-95, doi:
806	10.1016/j.palaeo.2010.03.029, 2010.
807	
808	Luciani, V., Giusberti L.: Reassessment of the early-middle Eocene planktic foraminiferal
809	biomagnetochronology: new evidence from the Tethyan Possagno section (NE Italy) and Western
810	North Atlantic Ocean ODP Site 1051., J. Foram. Res., 44, 2, 187-201, 2014.

812	Lunt, D.J., Ridgwell, A., Sluijs, A., Zachos, J., Hunter, S., Haywood A.: A model for orbital pacing of
813	methane hydrate destabilization during the Palaeogene, Nat. Geosc. Lett., 4, 775-778, DOI:
814	10.1038/NGEO1266, 2011.
815	
816	Marshall, J.D.: Climatic and oceanographic isotopic signals from the carbonate rock records and their
817	preservation, Geol. Mag., 129, 143-160, 1992.
818	
819	Molina, E., Arenillas, I., Pardo, A.: High resolution planktic foraminiferal biostratigraphy and
820	correlation across the Palaeocene Palaeocene/Eocene boundary in the Tethys, Bull. Soc. Géol. France,
821	170, 521–530, 1999.
822	
823	Moy, A.D., Howard, W.R., Bray, S.G., Trull, T.W.: Reduced calcification in modern Southern Ocean
824	planktonic foraminifera, Nat. Geosc., 2, 276-280.doi:10.1038/NGEO460, 2009.
825	
826	Müller-Merz, E., Oberhänsli, H.: Eocene bathyal and abyssal benthic foraminifera from a South
827	Atlantic transect at 20-30° S, Palaeogeogr. Palaeoclimatol. Palaeoecol., 83, 117-171, 1991.
828	
829	Nguyen, T.M.P., Petrizzo, M.R., Stassen, P., Speijer, R.P.: Dissolution susceptibility of Paleocene-
830	Eocene planktic foraminifera: Implications for palaeoceanographic reconstructions, Mar.
831	Micropaleont., 81, 1-21, 2011.
832	
833	Nicolò, M. J., Dickens, G. R., Hollis, C. J., Zachos, J. C.: Multiple early Eocene hyperthermals: their
834	sedimentary expression on the New Zealand continental margin and in the deep sea, Geology, 35, 699-
835	702, 2007.
836	
837	Norris, R.D.: Biased extinction and evolutionary trends, Paleobiology, 17 (4), 388-399, 1991.
838	
839	Norris, R.: Symbiosis as an evolutionary innovation in the radiation of Paleocene planktic foraminifera,
840	Paleobiology, 22, 461–480, 1996.

841	
842	Norris, R. D., Kroon, D., Klaus, A.: Proceedings of the Ocean Drilling Program, Initial Reports, 171B,
843	Ocean Drilling Program, College Station, TX, p. 1–749, 1998.
844	
845	O'Connor, M., Piehler, M.F., Leech, D.M., Anton, A., Bruno, J.F.: Warming and resource availability
846	shift food web structure and metabolism, Plos Biol., 7(8), 1-6. doi: 10.1371/journal.pbio.1000178,
847	2009.
848	
849	Ogg, J. G., Bardot, L.: Aptian through Eocene magnetostratigraphic correlation of the Blake Nose
850	Transect (Leg 171B), Florida continental margin, in: Proc. Ocean Drill. Progr., Scientific Results,
851	171B: Ocean Drilling Program, College Station, TX, Kroon, D. et al., 1–58,
852	www.odp.tamu.edu/publications/171B-SR/VOLUME/CHAPTERS/SR171B09, 2001.
853	
854	Olivarez Lyle, A., and Lyle, M.W.: Missing organic carbon in Eocene marine sediments: Is metabolism
855	the biological feedback that maintains end-member climates? Paleoceanogr., 21, PA2007,
856	doi:10.1029/2005PA001230, 2006.
857	
858	Oreshkina, T.V.: Evidence of Late Paleocene - Early Eocene hyperthermal events in biosiliceous
859	sediments of Western Siberia and adjacent areas, Austrian J. Earth Sci., 105, 145-153, 2012.
860	
861	Pearson, P.N., Palmer, M.R.: Atmospheric carbon dioxide concentrations over the past 60 million
862	years, Nature, 406, 695-699, doi:10.1038/35021000, 2000.
863	
864	Pearson P.N., Coxall H.K.: Origin of the Eocene planktonic foraminifer Hantkenina by gradual
865	evolution, Palaeontology, 57, 243-267, 2014.
866	
867	Pearson, P.N., Shackleton, N.J., Hall, M.A.: Stable isotope paleoecology of middle Eocene planttonic
868	foraminifera and multi-species isotope stratigraphy, DSDP Site 523, south Atlantic, J. Foram. Res. 23,
869	123-140, 1993.
870	

- 871 Pearson, P.N., Ditchfield, P.W., Singano, J., Harcourt-Brown, K.G., Nicholas, C.J., Olsson, R.K.,
- 872 Shackleton, N.J., Hall, M.A.: Warm tropical sea surface temperatures in the Late Cretaceous and
- 873 Eocene epochs, Nature 413, 481-487, 2001. doi:10.1038/35097000

- 875 Pearson, P.N., Olsson, R.K., Huber, B.T., Hemleben, C., Berggren, W.A.: Atlas of Eocene planktonic
- foraminifera, in: Cushman Found. Spec. Publ. 1, 1-514, 2006.
- 877
- Petrizzo, M.R.: The onset of the Paleocene–Eocene Thermal Maximum (PETM) at Sites 1209 and
 1210 (Shatsky Rise, Pacific Ocean) as recorded by planktonic foraminifera, Mar. Micropaleont., 63,
 187–200, 2007.
- 881
- Petrizzo, M.R., Leoni, G., Speijer, R.P., De Bernardi, B., Felletti, F.: Dissolution susceptibility of some
 Paleogene planktonic foraminifera from ODP Site 1209 (Shatsky Rise, Pacific Ocean), J. Foram. Res.
 38, 357–371, 2008.
- 885
- 886 Premoli Silva, I., Boersma, A.: Atlantic Eocene planktonic foraminiferal historical biogeography and
- 887 paleohygrographic indices, Palaeogeogr. Palaeoclimatol. Palaeoecol. 67, 315-356, 1988.
- 888
- 889 Premoli Silva, I., Boersma, A.: Atlantic Paleogene planktonic foraminiferal bioprovincial indices, Mar.
- 890 Micropaleont., 14, 357–371, 1989.
- 891
- 892 Quillévéré, F., Norris, R.D., Moussa, I., Berggren, W.A.: Role of photosymbiosis and biogeography in
- the diversification of early Paleogene acarininids (planktonic foraminifera), Paleobiology, 27, 311–326,
- 894 2001.
- 895
- Quillévéré, F., Norris, R.D., Kroon, D., Wilson, P.A.: Transient ocean warming and shift in carbon
 reservoir during the early Danian, Earth Planet. Sci. Lett., 265, 600–615, 2008.
- 898
- Raymo, M.E., Ruddiman W.F.: Tectonic forcing of late Cenozoic climate, Nature, 359, 117–122, 1992.

901 Schmitz, B., Pujalte, V.: Abrupt increase in seasonal extreme precipitation at the Paleocene-Eocene

902 boundary, Geology, 35(3), 215–218, doi:10.1130/G23261A.1, 2007.

903

- 904 Schneider, L.J. Bralower, T.J., Kump, L.J.: Response of nannoplankton to early Eocene ocean
- 905 destratification, Palaeogeogr. Palaeoclimatol. Palaeoecol., 310, 152-162, 2011.
- 906
- 907 Sexton, P.F., Wilson, P.A., Norris, R.D.: Testing the Cenozoic multisite composite DDO and
- 909 Drilling Program Leg 207), Paleoceanogr., 21, PA2019, 2006.
- 910
- 911 Sexton, P.F., Norris R.D., Wilson, P.A., Pälike, H., Westerhold, T., Röhl, U., Bolton, C.T., Gibbs, S.:
- 912 Eocene global warming events driven by ventilation of oceanic dissolved organic carbon, Nature 471,
- 913 349-353, doi:10.1038/nature09826, 2011.
- 914
- 915 Shackleton, N.J., Corfield, R.M., Hall, M.A.: Stable isotope data and the ontogeny of Paleocene 916 planktonic foraminifera, J. Foraminifer. Res., 15, 321–376, 1985.
- 917
- 918 Shamrock, J.L., Watkins, D.K., Johnston, K.W.: Eocene bio-geochronology of ODP Leg 122 Hole

919 762C, Exmouth Plateau (northwest Australian Shelf), Stratigraphy, 9, 55-76, 2012.

- 920
- Sims, P.A., Mann, D.G., Medlin, L.K.: Evolution of the diatoms: insights from fossil, biological and
 molecular data, Phycologia, 45, 361-402, 2006.
- 923
- 924 Sinton, C. W., Duncan R. A.: 40Ar/39Ar ages of lavas from the southeast Greenland margin, ODP Leg
- 925 152, and the Rockall Plateau, DSDP Leg 81, in: Sc. Res. Oc. Drill. Progr. 152, College Station, TX,
- 926 Larsen, H.C., Saunders, A.P., Clift, P.D., 387–402, 1998.
- 927
- 928 Slotnik, B.S., Dickens, G.R., Nicolo, M.J., Hollis, C.J., Crampton, J.S., Zachos, J.C., Sluijs, A.: Large-
- 929 amplitude variations in carbon cycling and terrestrial weathering during the latest Paleocene and
- earliest Eocene: The Record at Mead Stream, New Zealand, J. Geol., 120, 487–505, 2012.

\mathbf{O}	2	1
9	Э	T

932	Smith, R.Y., Greenwood, D.R., Basinger, J. F.: Estimating paleoatmospheric pCO ₂ during the Early		
933	Eocene Climatic Optimum from stomatal frequency of Ginkgo, Okanagan Highlands, British		
934	Columbia, Canada, Palaeogeogr. Palaeoclimatol. Palaeoecol., 293, 120–131, 2010.		
935			
936	Stanley, S.M.: Influence of seawater chemistry on biomineralization throughout Phanerozoic time:		
937	paleontological and experimental evidence, Palaeogeogr. Palaeoclimatol. Palaeoecol., 232, 214-236.		
938	doi: 10.1016/j.palaeo.2005.12.010, 2006.		
939			
940	Stanley, S.M.: Effect of Global Seawater Chemistry on Biomineralization: Past, Present, and Future,		
941	Chem. Rew., 108, 4483-4498. Doi: 10.1021/cr800233u, 2008.		
942			
943	Stanley, S.M., Ries, J.B., Hardie, L.A.: Seawater chemistry, coccolithophore population growth, and		
944	the origin of Cretaceous chalk, Geology 33, 593–596, 2005.		
945			
946	Thomas, E.: Biogeography of the late Paleocene benthic foraminiferal extinction, in Aubry, MP., et		
947	al. (eds.), Late Paleocene-Early Eocene Climatic and Biotic Events in the Marine and Terrestrial		
948	records: Columbia University Press, New York, p. 214–243, 1998.		
949			
950	Vandenberghe N., Hilgen F.J., Speijer R.P., Ogg J.G., Gradstein F.M., Hammer 0., Hollis C.J., Hooker		
951	J.J., In: The Paleogene Period, In: Gradstein, F., Ogg, J.G., Schmitz, M.D., Ogg, G.M., The Geologic		
952	Time Scale 2012, 855-921, Elsevier, Amsterdam, 2012.		
953			
954	Vogt, P.R.: Global magmatic episodes: New evidence and implications for the steady state mid-oceanic		
955	ridge, Geology, 7, 93–98, 1979.		
956			
957	Wade, B. S.: Planktonic foraminiferal biostratigraphy and mechanisms in the extinction of Morozovella		
958	in the late Middle Eocene, Mar. Micropaleont., 51, 23–38, 2004.		
959			

- 960 Wade, B.S., Al-Sabouni, N., Hemleben, C., Kroon, D.: Symbiont bleaching in fossil planktonic
- 961 foraminifera, Evol. Ecol., 22, 253-265. doi:10.1007/s10682-007-9176-6, 2008.
- 962

963	Wade, B.S., Pearson, P.N., Berggren, W.A., Pälike, H.: Review and revision of Cenozoic tropical
964	planktonic foraminiferal biostratigraphy and calibration to the geomagnetic polarity and astronomical
965	time scale, Earth Sci. Rev., 104, pp.111-142. doi:10.1016/j.earscirev.2010.09.003, 2011.
966	
967	Wade, B.S., Fucek, V.P., Kamikuri, SI., Bartol, M., Luciani, V., Pearson, P.N.: Successive extinctions
968	of muricate planktonic foraminifera (Morozovelloides and Acarinina) as a candidate for marking the
969	base Priabonian. Newsletters on Stratigraphy, 45 (3) 245-262, 2012.
970	
971	Westerhold, T., Röhl, U.: Orbital pacing of Eocene climate during the Middle Eocene Climate
972	Optimum and the chron C19r event-missing link found in the tropical western Atlantic, Geochem.,
973	Geophys., Geosyst., 14, 4811-4825, doi:10.1002/2013GC004960, 2013.
974	
975	Wilf, P., Cúneo, R.N., Johnson, K.R., Hicks, J.F., Wing, S.L., Obradovich, J.D.: High plant diversity in
976	Eocene South America: evidence from Patagonia, Science, 300, 122-125, 2003.
977	
978	Wing, S.L., Bown, T.M., Obradovich, J.D.: Early Eocene biotic and climatic change in interior western
979	North America, Geology 19, 1189-1192, 1991.
980	

- Woodbourne, M.O., Gunnell, G.F., Stucky, R.K.: Climate directly influences Eocene mammal faunal
 dynamics in North America, PNAS, 106, 13399-13403, 2009.
- 983
- 284 Zachos, J.C., Pagani, M., Sloan, L., Thomas, E., Billups, K.: Trends, rhythms, and aberrations in global

985 climate 65 Ma to Present, Science, 292, 686–693, 2001.

- 986
- 287 Zachos, J.C., Röhl, U., Schellenberg, S.A., Sluijs, A., Hodell, D.A., Kelly, D.C., Thomas, E., Nicolo,
- 988 M., Raffi, I., Lourens, L.J., McCarren, H., Kroon, D.: Rapid acidification of the ocean during the
- 989 Paleocene–Eocene thermal maximum, Science, 308, 1611–161, 2005.

991	Zachos, J.C., S. Schouten, S. Bohaty, T., Quattlebaum, Sluijs, A., Brinkhuis, H., Gibbs, S.J., Bralower,
992	T.J.: Extreme warming of midlatitude coastal ocean during the Paleocene-Eocene thermal maximum:
993	Inferences from TEX86 and isotope data, Geology, 34(9), 737–740. doi:10.1130/G22522.1, 2006.
994	
995	Zachos, J.C., Dickens, G. R., Zeebe, R.E.: An early Cenozoic perspective on greenhouse warming and
996	carbon-cycle dynamics, Nature, 451, 279–283, 2008.
997	
998	Zachos, J.C., McCarren, H., Murphy, B., Röhl, U., Westerhold, T.: Tempo and scale of late Paleocene
999	and early Eocene carbon isotope cycles: Implications for the origin of hyperthermals, Earth Planet. Sci.
1000	Lett., 299, 242-249, doi:10.1016/j.epsl.2010.09.004, 2010.
1001	
1002	Zeebe, R.E., Zachos, J.C., Caldeira K., Tyrrel T.: Carbon emissions and acidification, Science, 321, 51-
1003	52, 2008.
1004	
1005	Zonneveld, J.P., Gunnell, G.F., Bartels, W.S.: Early Eocene fossil vertebrates from the southwestern
1006	Green River Basin, Lincoln and Uinta Counties, Wyoming, J. Vert. Paleontol., 20, 369-386, 2000.
1007	
1008	
1009	
1010	Figure Captions
1011	
1012	Figure 1. Left: Paleogeographic location of the Possagno section (star) in the Belluno Basin, a
1013	Mesozoic-Cenozoic pelagic sequence, delimited during the Eocene by the shallow water carbonates of
1014	the Lessini Shelf to the west (Modified from Bosellini, 1989). 1 - deep water mudstones of the Jurassic
1015	basins; 2 - Paleogene shallow water limestones, lagoons, and shelf-edge reefs of the Lessini Shelf; 3 -
1016	Paleogene deep water pelagic claystones and marlstones of the Belluno and Lombardian Basin. Right:
1017	Blake Nose map and location of Site 1051 in the western North Atlantic, modified from Norris et al.
1018	(1998).

1021 lithology and planktic foraminiferal E-Zonations from Wade et al. (2011), modified by Luciani and 1022 Giusberti (2014). Magnetostratigraphy is from Agnini et al. (2006). Thinner lines: original data; thicker 1023 lines average 3-points. The average 3-point curve is utilized to dampen some of the potential diagenetic 1024 overprint on the ¹⁸O data. The red line is referred to the stable carbon isotopes and the blue line to the 1025 oxygen data. Pre-EECO CIEs are labelled according to current literature; the EECO and post-EECO 1026 CIEs are labelled according to Kirtland Turner et al. (2014) and Sexton et al. (2011) by substituting H 1027 (hyperthermals) with CIE (carbon isotope excursion). The yellow band highlights the interval 1028 tentatively referred to the EECO. We have tentatively named the post-EECO isotope shifts of small 1029 magnitude as events only when changes in isotopic composition are associated with sharp variations in 1030 planktic foraminferal assemblages and/or fragmentation index. This is because increase in 1031 fragmentation index as well as increase in some taxa have been observed during the pre-EECO 1032 hyperthermals from the same geological setting (Luciani et al. 2007; Agnini et al., 2009). However, 1033 since the identification of some post-EECO minor shifts as hyperthermals is tentative, they are 1034 indicated with a question mark. Filled circles show occurrences of abundant radiolarians. 1035 1036 1037 Fig. 3. The Possagno δ^{13} C record and relative abundance of main planktic foraminifera across the early 1038 and basal middle Eocene interval, plotted against lithology, fragmentation index (F index) and coarse 1039 fraction (CF) data. The subbotinids includes the genera Subbotina and Parasubbotina. 1040 Magnetostratigraphy is from Agnini et al. (2006). The biozonal scheme is from Wade et al. (2011), 1041 modified by Luciani and Giusberti (2014). The yellow bands highlight the interval tentatively referred 1042 to the EECO, the pre-EECO hyperthermals and post-EECO stable isotope excursions that are 1043 considered to represent hyperthermals. Pre-EECO CIEs are labelled according to current literature; the 1044 EECO and post-EECO CIEs are labelled according to Kirtland Turner et al. (2014) and Sexton et al. 1045 (2011) by substituting H (hyperthermals) with CIE (carbon isotope excursion). We have tentatively 1046 named the post-EECO isotope shifts of small magnitude as events only when changes in isotopic 1047 composition are associated with sharp variations in planktic foraminferal assemblages and/or 1048 fragmentation index. This is because increase in fragmentation index as well as increase in acarininid 1049 abundance have been observed during the pre-EECO hyperthermals from the same geological setting

Figure 2. Carbon and oxygen isotopes of bulk sediment from the Possagno section plotted against

1020

- 1050 (Luciani et al. 2007; Agnini et al., 2009). However, since the identification of some post-EECO minor 1051 shifts as hyperthermals is tentative, they are indicated with a question mark. Filled circles show

1052 occurrences of abundant radiolarians.

- 1053
- 1054 Fig. 4. Relative abundance of the main planktic foraminiferal genera from ODP Site 1051, plotted
- 1055 against the biozones of Wade et al. (2011), partly modified by Luciani and Giusberti (2014), and F
- 1056 index data. The subbotinid group includes the genera Subbotina and Parasubbotina.
- 1057 Magnetostratigraphy is from Ogg and Bardot (2001); the gray bands are intervals of uncertainty in
- 1058 magnetostratigraphic boundaries. The striped band is an interval of non-recovery. The yellow band
- 1059 highlights the interval tentatively referred to the EECO.
- 1060
- 1061 Figure 5. The record of warm-indices muricates morozovellids and large acarininids (>200 micron) in
- 1062 the western Tethyan setting from the Possagno (below, this paper) and Alano sections (above, from
- 1063 Luciani et al., 2010) plotted against the generalized oxygen and carbon isotopic curves based on
- 1064 benthic foraminiferal record, slightly modified, shown in Vandenberghe et al. (2012, Fig. 28.11). The
- 1065 original oxygen and carbon isotopic values from Cramer et al. (2009) are recalibrated to GTS2012
- 1066 (Vandenberghe et al., 2012). The Tethyan record shows that the long-lasting EECO and MECO
- 1067 intervals mark two main steps in the decline of relative abundance within this group of important early
- 1068 Paleogene calcifiers. E-Zones follow Wade et al. (2011), partly modified by Luciani and Giusberti
- 1069 (2014). B-P=Bartonian-Priabonian.
- 1070
- Table 1. Position with respect to magnetochrons of nine early and lower-middle Eocene $\delta^{13}C$ shifts 1071
- 1072 (CIE) at the Possagno section. The analogous magnetostratigraphic position of the hyperthermals (H)
- 1073 recognized at Site 1258, Demerara Rise (Kirtland Turner et al, 2014; Sexton et al., 2011) suggests a
- 1074 possible correspondence of these events.
- 1075
- 1076 Appendix A. Supplementary material
- 1077
- Table S1. Possagno δ^{13} C (‰) and δ^{18} O (‰) values against thickness (meters). 1078
- 1079

1080	Figure S1. The Possagno δ^{13} C data and relative abundance of minor planktic foraminiferal genera and
1081	selected species plotted against lithology, fragmentation index (F index) data. Magnetostratigraphy is
1082	from Agnini et al. (2006). The biozonal scheme is from Wade et al. (2011), modified by Luciani and
1083	Giusberti (2014). The yellow bands highlight the interval tentatively referred to the EECO, the pre-
1084	EECO hyperthermals and post-EECO stable isotope excursions that are considered to represent
1085	hyperthermals. Pre-EECO CIEs are labelled according to current literature; the EECO and post-EECO
1086	CIEs are labelled according to Sexton et al. (2011) by substituting H (hyperthermals) with CIE (carbon
1087	isotope excursion). We have tentatively named the post-EECO isotope shifts of small magnitude as
1088	events only when changes in isotopic composition are associated with sharp variations in planktic
1089	foraminferal assemblages and/or fragmentation index. This is because increase in fragmentation index
1090	as well as increase in acarininid abundance have been observed during the pre-EECO hyperthermals
1091	from the same geological setting (Luciani et al. 2007; Agnini et al., 2009). However, since the
1092	identification of some post-EECO minor shifts as hyperthermals is tentative, they are indicated with a
1093	question mark. Filled circles show occurrences of abundant radiolarians.
1094	
1095	
1096	Appendix B. Taxonomic list of species cited in text and figures
1097	
1098	Globanomalina australiformis (Jenkins, 1965)
1099	Morozovella aequa (Cushman and Renz, 1942)
1100	Morozovella gracilis (Bolli, 1957)
1101	Morozovella lensiformis (Subbotina, 1953),
1102	Morozovella marginodentata (Subbotina, 1953)
1103	Morozovella subbotinae (Morozova, 1939)
1104	Parasubbotina eoclava Coxall, Huber and Pearson, 2003
1105	Parasubbotina griffinae (Blow, 1979)
1106	Parasubbotina pseudowilsoni Olsson and Pearson, 2006
1107	Subbotina corpulenta (Subbotina, 1953)
1108	Subbotina eocena (Guembel, 1868)
1109	Subbotina hagni (Gohrbandt, 1967)

- 1110 Subbotina senni (Beckmann, 1953)
- 1111 Subbotina yeguanesis (Weinzierl and Applin, 1929)
- 1112 Planoglobanomalina pseudoalgeriana Olsson & Hemleben, 2006











Site 1051

Fig. 5



Table 1.

MAGNETOCHRON	POSSAGNO (NORTHEASTERN ITALY)	DEMERARA RISE (ODP SITE 1258, WESTERN TROPICAL ATLANTIC)
	STABLE CARBON ISOTOPE SHIFTS	HYPERTHERMAL EVENTS
	CIE4	H4
C21r	CIE3	НЗ
6211	CIE2	H2
	CIE1	H1
	CIE3	НЗ
C22n	CIE2	НЕ
	CIE1	H1
C22r	CIE2	H2
	CIE1	H1

Supplementary material



Possagno section

Fig. S1

· · · · ·	Ŭ Ŭ	
Thickness (meters)	d13Cndh %-	d180ndb %
66	1 32/15/03865	-1 55/131513
65.7	1,324303803	-1,354131313
65.4	1,301202021	1 205765522
65,4	1,500550409	-1,265705555
65,1	1,289525470	-1,334902053
05,1	1,288019147	-1,301227875
64,75	1,337699738	-1,365512853
64,5	1,2/1493282	-1,449946289
64,35	1,048026588	-1,843804126
64	1,15/592263	-1,664853101
63,75	1,242287722	-1,654166107
63,5	1,369396415	-1,354916697
63,25	1,258468714	-1,405387406
63	1,328439322	-1,377815013
62,75	1,277403321	-1,585307364
62,5	1,255599346	-1,755604391
62,25	1,28272387	-1,411073928
62	1,268756475	-1,226212864
61,75	1,209940758	-1,47335947
61,5	1,178018179	-1,715059881
61,25	1,095738735	-1,870218997
61	1,313557115	-1,399695163
60,75	1,277940478	-1,35860539
60,5	1,291423645	-1,430644636
60,25	1,200942841	-1,498587494
60	1,250952568	-1,417450314
59,75	1,218475547	-1,393167024
59,5	1,225717988	-1,324035769
59,25	1,194266763	-1,51954699
59	1,080495861	-1,482955818
58.8	1.075773802	-1.522460465
58.6	1.11857554	-1.22201932
58.4	1.028116908	-1.323055034
58.2	0.92447057	-1.608414416
58	1.028708603	-1.348841195
57.8	1.07809129	-1.064679508
57,6	0.87	-1.86
57,0	0.892621305	-1 3938696/13
57,4	0,032021305	-1 8/83/981/
57	0.858608028	-1 97897778/
56.8	1 10/15652	-1 260/6008/
56,6	1 1/6252027	_1 /5152727/
56.4	1 086170407	-1,4J1J0/324
50,4	1,0001/0402	-1,2/140000
50,2	0.059074020	-1,/01198400
	0,9589/4626	
55,8	1,131002012	-1,33444/985
55,6	1,19	-1,25
55,4	1,18	-1,29

Table S1. Isotopic data of Possagno section

55,2	1,15	-1,45
55	0,74	-1,99
54,8	1,07	-1,5
54,6	0,96	-1,58
54,4	1,08	-1,25
54,2	1,05	-1,29
54	1,15	-1,11
53,8	1,1	-1,29
53,6	1,18	-1,14
53,4	1,03	-1,36
53,2	0,93	-1,8
53	0,96	-2,02
52,8	1,3	-1,26
52,6	1,25	-1,36
52,4	1,3	-1,48
52,2	1,26	-1,37
51,8	1,17	-1,7
51,6	1,14	-1,69
51,4	1,14	-1,69
51,2	1,21	-1,71
51	1,32	-1,25
50,8	1,14	-1,63
50,6	1,13	-1,61
50,4	1,13	-1,44
50,2	1,16	-1,28
50	0,99	-1,59
49,8	1,01	-1,48
49,6	0,9	-1,57
49,4	0,5	-2,55
49,2	0,82	-2,21
49	1,24	-1,43
48,8	1,33	-1,13
48,6	1,24	-1,82
48,4	1,53	-1,03
48,2	1,14	-1,78
48	1,26	-1,7
47,8	1,29	-1,56
47,6	1,23	-1,73
47,4	1,18	-1,65
47,2	1,19	-1,67
47	1,16	-1,58
46,8	1,2	-1,89
46,6	1,34	-1,42
46,4	1,31	-1,43
46,2	1,4	-1,64
46	1,35	-1,59
45,8	1,43	-1,55
45,6	1,44	-1,46
45,2	1,36	-1,71
45	1,34	-1,66

44,8	1,29	-2,02
44,6	1,59	-1,27
44,4	1,54	-1,41
44,2	1,47	-1,61
44	1,32	-1,84
43,8	1,48	-1,73
43,6	1,59	-1,27
43,4	1,55	-1,41
43,2	1,24	-2,19
43	1,56	-1,42
42,8	1,4	-1,67
42,6	1,4	-1,66
42,4	1,38	-1,65
42,2	1,49	-1,43
42	1,47	-1,44
41,8	1,38	-1,69
41,6	1,4	-1,52
41,4	1,3	-1,98
41,2	1,46	-1,65
41	1,51	-1,63
40,8	1,24	-1,89
40,6	1,18	-1,78
40,4	1,1	-1,71
40,2	1,36	-1,91
40	1,5	-1,55
39,8	1,01	-1,98
39,6	1,33	-1,94
39,4	1,18	-2,19
39,2	1,01	-2,09
39	1,32	-1,75
38,8	1,49	-1,66
38,6	1,3	-1,71
38,4	1,31	-1,46
38,2	1,09	-1,83
38	1,24	-2,66
37,8	1,52	-1,65
37,6	1,82	-0,92
37,4	1,44	-1,86
37,2	1,54	-1,72
37	1,62	-1,43
36,8	1,54	-1,6
36,6	1,37	-1,74
36,4	1,39	-2,15
36,2	1,47	-1,69
36	1,72	-1,19
35,8	1,34	-1,72
35,6	1,28	-1,79
35,4	1,49	-1,77
35,2	1,36	-1,64
35	1,31	-1,87

34,8	1,48	-1,16
34,6	1,41	-1,63
34,4	1,36	-1,84
34,2	1,38	-1,73
34	1,45	-1,49
33,8	1,35	-1,63
33,6	1,36	-1,2
33,4	1	-2,02
33,2	0,75	-2,58
33	1,15	-2,29
32,8	1,63	-1,12
32,6	1,35	-1,84
32,4	1,39	-1,79
32,2	1,58	-1,7
32	1,62	-1,47
31,8	1,37	-2,18
31,6	1,29	-1,65
31,4	0,78	-3,1
31,2	1,27	-1,87
31	1,16	-1,94
30,8	1,2	-1,81
30,6	1,26	-1,66
30,4	1,12	-1,67
30,2	0,66	-2,4
30	1,12	-1,81
29,8	1,06	-2,23
29,6	1,24	-1,74
29,4	1,24	-1,79
29,2	1,2	-2,16
29	1,38	-1,92
28,8	1,09	-2,23
28,6	1,25	-1,62
28,4	1,25	-1,8
28,2	1,15	-1,74
28	1,24	-1,7
27,8	1,18	-1,62
27,6	0,83	-2,15
27,4	0,71	-2,45
27,2	1,35	-1,41
27	1,25	-1,66
26,8	1,43	-1,25
26,6	1,54	-1,11
26,4	1,18	-1,74
26,2	1,27	-1,38
26	1,3	-1,74
25,8	1,3	-1,92
25,6	1,11	-1,91
25,4	1,42	-1,5
25,2	1,4	-1,51
25	1,27	-1,81

24.7	1 38	_1 13
24,7	1,38	-1,13
24,4	1,20	-1,94
24,2	1,20	1.01
24	1,19	-1,91
23,8	1,44	-1,75
23,0	1,45	-1,09
23,4	1,47	-1,83
23,2	1,41	-1,76
23	1,57	-1,19
22,8	1,28	-1,72
22,6	1,47	-1,64
22,4	1,25	-1,99
22,2	1,13	-2,11
22	1,31	-1,96
21,8	1,2	-2,07
21,4	0,78	-2,72
21,2	1,68	-0,78
21	1,05	-1,87
20,8	0,61	-1,79
20,6	0,17	-2,27
20,4	0,97	-1,69
20,2	1,32	-1,91
20	0,83	-1,64
19,8	0,33	-2,31
19,6	0,88	-1,89
19,4	1,19	-1,6
19,2	1,38	-0,81
19	1,4	-1,49
18,8	0,67	-1,73
18,6	0,72	-2,59
18,4	1,25	-1,59
18,2	0,85	-1,9
18	-0,22	-3,64
17,8	0,57	-2,43
17,6	0,87	-1,63
17,4	0,38	-2,11
17,2	1	-1,66
17	0,31	-2,16
16,8	-0,07	-3,25
16,6	0,87	-1,79
16,4	0,76	-2,08
16,2	0,62	-2,16
16	0,76	-1,7
15,8	0,47	-1,97
15,6	0,7	-1,41
15,4	0,86	-1,79
15,2	0,94	-1,58
15	0,94	-1,51
14,8	1,02	-1,42
14,6	0,67	-1,74

14.4	0.87	_1 73
14,4	0,76	-1,25
13.5	0,61	-1.79
13	0.53	-1.99
12.55	0.55	-1.95
12	0.8	-1.66
11.5	0.73	-1.87
10	0.91	-1.39
9.5	0.860407617	-1.745868284
9	0.847698043	-1.6249864
8.5	0.704323168	-1.620825392
8	0.697123094	-1.606233422
7.5	0.971220119	-1.70465462
7	0.960200544	-1.804953966
6.5	1.03282727	-1.798588563
6	1.093247595	-1.655217488
5.5	1.182569921	-1.685192373
5	1.147446246	-1.536443685
4.5	1.136631571	-1.751460814
4	1.131549897	-2.060952576
3.5	1.321208222	-1.972329429
3	1,210211548	-2,357527232
2,5	1,095834873	-2,265978774
2	1,205609198	-2,180916572
0,975	1,193025524	-2,39513048
0,825	1,169705849	-2,128674243
0,725	1,150951175	-2,193656665
0,625	1,187252381	-2,071204006
0,525	1,072769571	-1,900119279
0,425	0,996418462	-2,185873828
0,325	0,855189052	-1,957644402
0,275	0,325183243	-2,571328228
0,225	0,363733114	-2,577693458
0,175	0,395223833	-2,442193443
0,125	0,261679624	-2,517606276
0,075	0,056425765	-2,600329725
0,025	0,046204045	-2,697877737
0,01	0,715394386	-2,627456411
-0,025	0,825253976	-2,496448436
-0,075	1,512990667	-2,286158599
-0,125	1,549820857	-2,208348633
-0,175	1,458455048	-2,554977403
-0,225	1,393471238	-2,195702997
-0,325	1,366631429	-2,565834405
-0,425	1,545594619	-2,359719334
-0,525	1,67926181	-2,339723417
-0,725	1,762957082	-2,045519714
-0,825	1,772335157	-1,765052919