Calcareous nannofossil biostratigraphy and paleoenvironmental variations across the upper Paleocene–lowermost Eocene at Gebel Nezzazat, West Central Sinai, Egypt

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Abstract
Detailed lithostratigraphic and calcareous nannofossil biostratigraphic studies were carried out across the Paleocene–Eocene (P-E) that outcrops at Gebel Nezzazat (G. Nezzazat) in West Central Sinai (Egypt). The study interval spans from the upper part of the Tarawan Formation to the lowermost Thebes Formation covering the whole Esna Formation in between them. The Esna Formation had been subdivided into four members: the Hanadi, Dababiya Quarry, Mahmiya, and Abu Had Members. Five calcareous nannofossil biozones (NP7/8, NP9 to NP12) and four subzones (NP9a and NP9b and NP10a and NP10b) were recognized. The lowest occurrences (LOs) of Fasciculithus alanii group, Neochiastozygus junctus, Sphenolithus radians, and Blackites herculesii as well as the highest occurrence of F. alanii group and the increased frequency of N. junctus are biostratigraphically significant. On contrast, the LOs of Discoaster binodosus, Discoaster mahmoudii, Discoaster diastypus, Zygrhablithus bijugatus, and Campylosphaera dela as well as the LOs of Fasciculithus tympaniformis are unreliable bioevents. Calcareous nannofossils increase in abundance close to the P-E transition. Ericsonia subpertusa suddenly increases above the base of Eocene, whereas the diversity of Fasciculithus drops close to this level. The P-E boundary at G. Nezzazat was placed at the base of the Dababiya Quarry Member in coincidence with the base of Subzone NP9b that was delineated by the LOs of Discoaster araneus, Rhomboaster cuspis, Rhomboaster calcitrapa, Rhomboaster spineus, and Rhomboaster bitrifida. A small gap was recorded across the P-E boundary as indicated by the lack of the four beds of the Dababiya Quarry Member. The changes in calcareous nannofossil assemblages reveal warm-water and oligotrophic conditions prevailed during the transition at G. Nezzazat.

Keywords Nezzazat · Calcareous nannofossils · Paleocene/Eocene · Rhomboaster · Sinai · Egypt

Introduction

The Paleocene–Eocene was distinguished by various climatic, sedimentologic, isotopic, and paleontologic variations (Dupuis et al. 2003). This interval was accompanied by a short-lived episode of global warming associated with negative shifts in the δ13C and δ18O values as well as carbonate...
contents (e.g., Cramer et al. 2003; Zachos et al. 2005; Westerholm et al. 2009; Frieling et al. 2018). Furthermore, the P-E transition was associated with changes in the biotic assemblages which include origins of new species, radiations, migration towards higher latitudes, and turnovers of some taxa (e.g., Bralower et al. 2002; Gingerich 2003; Bown et al. 2004; Tantawy 2006; Agnini et al. 2007a; Westerholm et al. 2009; Kasem et al. 2020a, b). As a result of the location of Egypt at the Tethys’ southern margin during the P-E transition, Egypt had complete P-E sequences, which have a perfect record of changes in sedimentology and biotic assemblages throughout this interval (Dupuis et al. 2003; Kasem et al. 2020b).

Thus, Egyptian sequences attracted several authors to track the biotic, isotopic, and climatic responses to climatic variations through the P-E interval (e.g., Youssef 2016; Obaidalla et al. 2017; Faris et al. 2018; Kasem et al. 2020a, b). The Global Boundary Stratotype Section and Point (GSSP) that hosts the base of the Eocene Series had been officially ratified at the Dababiya Quarry in southern Egypt (Dupuis et al. 2003). The onset of the Eocene Series at the GSSP coincides with a distinctive bed known as the Dababiya Quarry Beds (DQBs) that are easily distinguished based on their lithological, biotic, and isotopic features (Dupuis et al. 2003). Further subdivisions to the DQBs were achieved by several authors based on the variations in color, texture, and the phosphatic as well as carbonate contents (e.g., Aubry et al. 2007; Ouda et al. 2013). The DQBs had been recorded in many sections in Egypt (e.g., Tantawy 2006a, b; Ouda et al. 2016a, b; Faris et al. 2017; Kasem et al. 2020b).

Calcaneous nannofossils are highly sensitive to climatic changes and underwent considerable variations during the latest Paleocene–early Eocene. Therefore, they are a significant tool for tracking the biotic response across this transition (e.g., Perch-Nielsen 1985; Bralower et al. 2002; Dupuis et al. 2003; Tantawy 2006a, b; Agnini et al. 2006, Agnini et al. 2007a, b; Bernaola et al. 2007; Mutterlose et al. 2007; Raffi and Raffi and De Bernaola 2008; Bown and Pearson 2009; Raffi et al. 2009; Self-Trail et al. 2012; Kasem et al. 2020a, b). The calcaneous nannofossil assemblage’s changes

Table 1 Abundances of Paleocene–early Eocene calcaneous nannofossils at G. Nezzazat, WC Sinai, Egypt

| Sample | Formation | Species                | Abundance |
|--------|-----------|------------------------|-----------|
| NP10   |           |                        |           |
| NP9    |           |                        |           |
| NP7/8  |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
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| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
| NP12   |           |                        |           |
| NP11   |           |                        |           |
that include the origination and radiation of genus *Rhombobaster* and the origination of new species of genera *Discoaster* and *Fasciculithus* are important bioevents that mark the P-E boundary, and are useful in global biostratigraphic correlations (Table 1).

The main intents of this study are to describe the lithostratigraphy of the uppermost Paleocene–lowermost Eocene succession at Gebel Nezzazat, to provide detailed calcareous nannofossil biostratigraphy, to reveal the variations in calcareous nannofossil assemblages, and to discuss the biostratigraphic reliability of significant bioevents across the P-E interval, as well as to track the changes in water temperature and fertility through the P-E transition.

**Location, material, and methods**

Sixty-one rock samples covering the Paleocene–Eocene sequence at Gebel Nezzazat (Lat. 28° 47′ 45.4″N and Long. 33° 14′ 9.79″E), West Central Sinai, Egypt (Fig. 1), were investigated in this study. Smear slides from each sample were prepared for investigation of calcareous nannofossil assemblages according to Bown and Young (1998). The slides had been investigated by a Euromex iScope microscope at 1250× magnification at Damanhour University, Egypt. The state of preservation of the calcareous nannofossil assemblages is good, where little or no evidence of overgrowth and/or etching was noted. A quantitative analysis was carried out to calcareous nannofossil assemblages. Specimens had been counted in 50 fields of view to reveal the abundances of various species. Additional 50 fields of view were viewed just to reveal scarce species. To ensure even distribution of the sediments on the slide, about 0.1 g of raw sediments was transferred into 10 ml of distilled water in a test tube and they were dispersed; 0.25 mm of the suspension was pipetted onto 22 mm x 22 mm coverslips and dried on a hot plate. The coverslips were fixed onto glassy slides by using DPX Mountant. The materials used for this study are deposited at Faculty of Science, Geology Department, Damanhour University, Damanhour, Egypt.
Fig. 1 Location map shows the study section (Gebel Nezzazat, West Central Sinai, Egypt).

Fig. 2 Google earth map, shows the study section (Gebel Nezzazat, West Central Sinai, Egypt).
Lithostratigraphy

The uppermost Paleocene–lowermost Eocene at Gebel Nezzazat (G. Nezzazat) extends in three formations that are, from base to top, the Tarawan, Esna, as well as Thebes Formations (Fig. 3, Table 2).

The Tarawan Formation

Awad and Ghabrial (1965) had introduced this unit to describe a succession of chalk passing into chalky limestones and marly limestone at Gebel Tarawan, Kharga Oasis (Western Desert, Egypt). About 2.3 m from the upper part of the Tarawan Formation had been investigated in this study and consists of hard, yellow argillaceous limestone grading into white chalky limestone (Figs. 3 and 4A, B). It lies conformably above the Dakhla Formation and below the Esna Formation (Fig. 4A). Depending on the calcareous nannofossils, it was assigned to the (NP7/8 Zone) Thanetian Age (Figs. 3 and 5).

The Esna Formation

The Esna Shale Passage Beds was originally established by Beadnell (1905) to characterize a succession of 104-m-thick greenish gray shale intercalated with calcareous interbeds, grades upward into argillaceous limestone, and overlies the Cretaceous sequence and underlies the Eocene succession at Gebel Oweina, near Esna, southern Egypt. He named it Esna Shale. This unit was assigned to the Esna Formation by Said (1960). At G. Nezzazat, this unit is about 9.2 m of gray to dark gray shale intercalated with black shale and yellow, gray to brown calcareous shale, and upgrades to yellow marl and white chalky limestone (Figs. 3 and 4A, C, D). It conformably overlies the Tarawan Formation and is overlain by the Thebes Formation conformably (Fig. 4A, C, D). The Esna Formation in our study contains the following calcareous nannofossil biozones (the upper part of NP7/8, NP9, NP10, and NP11 and the lower part of NP12), so it is assigned to Thanetian–Ypresian ages (Figs. 3 and 5).

It was further subdivided into smaller units (e.g., Abdel Razik 1972; Dupuis et al. 2003; Tantawy 2006a, b; Ouda et al. 2016a, b; Kasem et al. 2020a, b) (Table 2). In this study, we followed the Esna Formation subdivision of Aubry et al. (2007), from base to top as follow.

The Hanadi Member

This unit was originally established to extend from the ground of the Esna Formation (Fm) to the top of the phosphatic bed (Abdel Razik 1972). Later, Aubry et al. (2007) delimit this unit to extend from the base of the Esna Fm to the base of the phosphatic bed. It consists mainly of gray, massive, calcareous shales. At G. Nezzazat, this member consists of 1.8-m-thick gray to dark gray shale (Fig. 3).
The Dababiya Quarry Member

It was originally established by Aubry et al. (2007) to distinguish the distinct beds that mark the base of the Eocene at the Dababiya Quarry area in Egypt. These marker beds reflect climatic changes in the southern Tethys during the P-E transition as indicated by their lithological, geochemical, and biotic characteristics (Aubry et al. 2007). The base of this unit in this study is coincident with the entry of Subzone NP9b (Fig. 3) (Dupuis et al. 2003; Kasem et al. 2020b).

Further subdivision of these beds was introduced by several authors depending on their color, texture, contents of carbonate and coprolite, and/or the composition and patterns of foraminifers (e.g., Aubry et al. 2007; Ouda et al. 2013). Moreover, Tantawy (Tantawy 2006a, b) subdivided the interval of the Esna Formation that covers the Dababiya Quarry Member of Aubry et al. (2007) into five units (DQBs 1–5). The DQB1, at section E, which lies east of the P-E GSSP is missed (Khozyem et al. 2015). At Naqb Assiut section, Kharga Oasis, Western Desert, Egypt, the Dababiya Quarry Member is only represented by the upper two beds (nos. 4 and 5) (El-Dawy et al. 2016), while at Gaga section, Baris Oasis, Western Desert, Egypt, the first bed (clay layer) of the Dababiya Quarry Member at the Paleocene/Eocene GSSP which represents the onset of the P-E boundary is missing, whereas the rest of the beds (nos. 2–4) are well represented (Metwally and Mahfouz 2018). In the present study, the Dababiya Quarry Member (DQM) is represented by only one sample of black shale, which may represent DQB1, of about 15 cm thick (Fig. 3).

![Fig. 3 Lithostratigraphic columnar section of the upper Paleocene-lower Eocene succession at G. Nezzazat, WC Sinai, Egypt.](image-url)
The Mahmiya Member

Aubry et al. (2007) established this member to cover the main part of the Esna Unit 2 of Dupuis et al. (2003) at the Dababiya Quarry section. It consists of monotonous, dark, clayey shales without marked bedding and has low calcium carbonate content (<50%) and clear cyclic color variations. The thickness of the Mahmiya Member is 65 m in the Dababiya Quarry section. Aubry et al. (2007) used the term El Quda Bed to describe the thin (10–30 m) calcarenite bed within the Mahmiya Member which consists generally of accumulations of cylindrical coprolites, phosphatic and shell clasts, and a variable amount of glauconite at the base (Table 2). It differs from the beds of the DQM by the presence of glauconite and the erosive and bioturbated contact with the underling lithology. In this study, the Mahmiya Member is ~3.4 m of gray to dark gray shales (Fig. 3).

The Abu Had Member

This member was originally established by Abdel Razik (1972) as the lower member of the Thebes Formation and covers the interbeds of limestones and shales at the transition between the shales of the Esna Formation and the massive carbonates of the Thebes Formation. Aubry et al. (2007) assigned this member to the Esna Formation at the Dababiya Quarry (Table 2). At the Dababiya Quarry, this member is 43.5 m thick of alternating shales and limestone. In the present study, the upper fraction of the Esna Formation at the G. Nezzazat section is assigned to the Abu Had Member and is about 4 m of yellow calcareous shale, gray shale, gray to brownish calcareous shale, dark gray shale, yellow to gray calcareous shale, yellow marl, white chalky limestone associated with chert nodules and bands, as well as brown calcareous shale (Fig. 3).

Fig. 4 A: Field photograph shows different formations covers the study interval at the study section; B: Field photograph shows the contact between Dakhla and Tarawan formations; C: Field photograph shows the lower part of the Esna Formation; D: Field photograph shows the contact between the Esna and Thebes formations.
The Thebes Formation

It was described by Said (1960) to cover a 290-m white to grayish white limestone with chert bands that superimpose the Esna Formation at the Gebel Gurnah, southern Egypt. About 2 m of hard yellow to white argillaceous limestone from the lowermost part of this formation was examined at G. Nezzazat and was assigned to the Ypresian Age based on the presence of the calcareous nannofossil biozone (NP12) (Figs. 3, 4A, D, and 5). It conformably underlain by the Esna Formation.

Results

Calcareous nannofossil biostratigraphy

Martini’s (1971) zonation scheme of the Paleogene was adopted in this study, and Romein’s (1979) recommendation of combining Zones NP7 and NP8 was followed. In addition, Aubry et al.’s (1999) proposition of subdividing Zone NP9 into Subzones NP9a and NP9b was applied. Furthermore, the subdivision of Zone NP10 suggested by Kasem et al. (2020a) into Subzones NP10a and NP10b was followed. The stratigraphic importance of calcareous nannofossil bioevents that associate the P-E transition was discussed in this study. Abbreviations that are applied in this study are LO for the lowest occurrence, HO for the highest occurrence, LCO for the lowest common occurrence, and HCO for the highest common occurrence (Kasem et al. 2017b). Table 1 shows the counts and stratigraphic distributions of calcareous nannofossil assemblages. Microphotographs of the recorded species are shown in Plates 1 and 2. Five calcareous nannofossil zones and four subzones were recognized and discussed as follows.

Discoaster mohleri Zone (NP7/8)

This combined zone spans from the LO of taxon Discoaster mohleri to the LO of Discoaster multiradiatus (Romein 1979). It is comparable to Zones CP6 and CP7 of Okada and Bukry (1980) and Zones CNP9 and CNP10 of Agnini et al. (2014). Originally, it was introduced by Hay (1964), yet the
marker species *Heliolithus riedelii* was not recorded from several sections besides having inconsistent stratigraphic ranges (Perch-Nielsen 1985; Romain 1979; Agnini et al. 2007b). Accordingly, Romain (1979) excluded this bioevent from being a zonal marker and lumped Zones NP7 and NP8 together into the *D. mohleri* Zone (NP7/8). In this study, this zone is ~2.8 m thick (samples 1 to 11) and covers almost the whole Tarawan Formation and extends to the basal part of the Esna Formation (Table 1, Fig. 2). It was assigned to Thanetian.

**Discoaster multiradiatus Zone (NP9)**

This zone had been established by Bramlette and Sullivan (1961) and then had been emended by Martini (1971) to the biostratigraphic interval between the LOs of *Discoaster multiradiatus* and *Tribrachiatus bramlettei*. It is comparable to Zone CN8 of Okada and Bukry (1980), Subzone NTp16b to Zone NTp20 of Varol (1989), and Zone CNP11 plus the lower part of Zone CNE1 of Agnini et al. (2014). Zone NP9 was documented from several Paleocene–Eocene successions in Egypt (Kasem et al. 2020a, b). At G. Nezzazat, this zone extends from sample 11 to sample 22 and is ~2.4 m thick in the basal portion of the Esna Formation and dates Thanetian–Ypresian (Table 1, Fig. 3).

Most *Discoaster* species that mark the Paleocene first occur in this zone, and genera *Fasciculithus* and *Rhomboaster* radiate in this zone (Table 1). Furthermore, many species belonging to the genus *Fasciculithus* disappear in the upper part of this zone (Table 1), among them are, e.g., *Fasciculithus lillianiae*, *Fasciculithus alanii*, *Fasciculithus thomasi*, and *Fasciculithus clinitus*. Further subdivisions of Martini’s (1971) Zone NP9 or equivalent interval had been recommended by several authors (Bukry 1973; Varol 1989; Kasem et al. 2020a, b).

Bukry (1973) subdivided this interval (Zone CP8) based on the LO of *Campylosphaera eodela*, *Rhomboaster cuspis*, and related taxa in shallow-ocean areas, and by *C. eodela* in deep-ocean areas (Bukry and Percival 1971). This subdivision is consistent with the occurrences documented by many authors (e.g., Tantawy 2006a, b). However, later studies confirmed that *Campylosphaera eodela* has taxonomic disputes and inconsistent stratigraphic ranges and, thus, is an unreliable zonal marker (Agnini et al. 2007b; Kasem et al. 2020a, b).

Bybell and Self-Trail (1997) further subdivided Zone NP9 based on the HOs of fasciculiths species (e.g., *F. clinatus*, *F. lillianae*, *Fasciculithus hayi*, *Fasciculithus bobii*, *F. alanii*, and *Fasciculithus mirenus*). Yet Khoozem et al. (2013) noted that these species disappear within Zone NP9. Furthermore, Kasem et al. (2020b) recorded *Fasciculithus involutus* plus *Fasciculithus tympaniformis* in the basal portion of Zone NP10. Aubry et al. (1999) suggested subdividing Zone NP9 into two subzones (NP9a and NP9b), relying on the LOs of genus *Rhomboaster*, *Discoaster araneus*, plus/or *Discoaster anartios*. Several studies confirmed the applicability of this suggestion (e.g., Dupuis et al. 2003; Agnini et al. 2007a, b; Faris et al. 2015; Kasem et al. 2020a, b). In this study, Zone NP9 has been subdivided into two subzones as follows.

**NP9a Subzone**

It covers the biostratigraphic interval between the LOs of *Discoaster multiradiatus* and *Rhomboaster* spp., *D. araneus*, plus/or *D. anartios* (Aubry et al. 1999). It extends from sample 11 to sample 18 at G. Nezzazat (Table 1) and covers about 1.5 m in the lower portion of the Esna Formation (Fig. 3). This subzone belongs to the Thanetian Age. Several new species come in this interval; among them are *Fasciculithus lillianiae*, *Neochiastozygus pusillus*, *Ellipsolithus distichus*, *Discoaster barbadensis*, *Discoaster falcatus*, *Neo-chiastozygus dientus*, and *Lanternithus simplex* (Table 1).

**NP9b Subzone**

The base of this subzone was located by the LOs of *Rhomboaster* spp., *D. araneus*, and/or *D. anartios*, and its upper limit was delineated by the LO of taxon *Tribrachiatus bramlettei* (Aubry et al. 1999). It is 0.9 m thick in the Esna Fm at G. Nezzazat (Fig. 3), covering from sample 19 (black shale) to sample 22, and belongs to the Ypresian Age (Table 1). Many new species first occur in this interval; among them are *Rhomboaster cuspis*, *Rhomboaster spinus*, *Rhomboaster bitritida*, *Rhomboaster calcitrapa*, *Chiasmolithus nitidus*, *Discoaster mahmoudii*, *D. araneus*, *D. anartios*, *Calcidiscus* sp., *Blackites herculesii*, *Discoaster diastypus*, *Pontosphaera plana*, and *Pontosphaera distincta* (Table 1). The HO of *Fasciculithus alanii* as well as the LOs of *Rhomboaster calcitrapa*, *Rhomboaster intermedia*, and *Blackites herculesii* were used to delineate the lower limit of Subzone NP9b (Faris and Salem 2007). Some authors documented that *Fasciculithus alanii* go extinct in coincidence with the ground of Subzone NP9b (Faris and Salem 2007), yet it has its HO earlier at the Dababiya Quarry (Dupuis et al. 2003), and extend up to the lowermost Eocene in some sections (Kasem et al. 2020b). In this study, *F. alanii* occurs in Subzone NP9b (Table 1).

**Tribrachiatus contortus Zone (NP10)**

Hay (1964) reported that this biozone covers from the LO of *Tribrachiatus bramlettei* to the HO of *T. contortus*. This zone is comparable to Subzone CP9a of Okada and Bukry
The start of Zone NP10 was delineated variously by different authors because of the taxonomic disputes among them concerning whether *T. bramlettei* has a different structure from *Rhomboaster* or not (Kasem et al. 2020a). Faris et al. (2015) proposed to locate the base of Zone NP10 at the LCO of *T. bramlettei*. At G. Nezzazat, *T. bramlettei* first occurs in sample number 23 with common occurrence (Table 1) and has been adopted to mark the base of Zone NP10. Abu Shama et al. (2007) suggested approximating the base of Zone NP10 at the increased frequency of *Neoehiazoxygus junctus* to delineate the lower limit of Zone NP10. At G. Nezzazat, *N. junctus* first occurs in sample 19 and shows increased frequency in sample 21 slightly below the base of Zone NP10 (sample 23) (Table 1).

The genus *Fasciculithus* commonly disappears in the basal part of Zone NP10 (Romein 1979); thus, it can help in the approximation of the lower limit of this zone in case of the scarcity or absence of the zonal marker (Perch-Nielsen 1985). However, Kasem et al. (2020a) noted that *F. tympaniformis* occurs with common occurrence up to the middle part of Zone NP10. Representatives of genus *Tribrachiatus* are important in biostratigraphic correlation and further subdivision of Zone NP10. The co-occurrences of *Tribrachiatus bramlettei* and *T. contortus* in Zone NP10 from one side and *T. contortus–Tribrachiatus orthostylus* from another side are a good indicator to the completeness of this interval (e.g., Agnini et al. 2006; Kasem et al. 2020a, b).

At G. Nezzazat, an overlap between the stratigraphic ranges *T. bramlettei* and *T. contortus* which were documented extends from sample 26 to sample 33, and *T. contortus* and *T. orthostylus* have a stratigraphic overlap extending from sample 26 to sample 34 (Table 1). Some authors suggested further subdivisions for Martini’s (1971) Zone NP10 (e.g., Aubry 1996; Kasem et al. 2020a).

Aubry (1996) suggested subdividing Zone NP10 into four subzones: NP10a to NP10d, based on LOs and HOs of *Tribrachiatus digitalis* and *T. contortus*. The validity of this suggestion was confirmed by various studies (e.g., Dupuis et al. 2003; Al Wosabi 2015). Yet the reliability of *T. digitalis* as a marker is questionable because of the disputes concerning its taxonomic position and biostratigraphic range (Raffi et al. 2005). Consequently, Kasem et al. (2020a, b) recommended to exclude *T. digitalis* from being a reliable subzonal marker and used the LO of *T. contortus* to subdivide Zone NP10 into Subzone NP10a plus NP10b. This suggestion is followed in this study.

**Subzone NP10a**

Kasem et al. (2020a) introduced Subzone NP10a to extend between the subsequent LOs of *Tribrachiatus bramlettei* and *T. contortus*. This subzone is comparable to Aubry’s (1996) Subzones NP10a to NP10c and Tantawy’s (1998) Subzones NP10a and NP10b. At G. Nezzazat, this subzone extends from sample 23 to sample 25 (~40 cm thick) in the Esna Formation (Fig. 3, Table 1) and is assigned to the Ypresian Age.

**Subzone NP10b**

Aubry (1996) introduced Subzone NP10d to cover the total range of *Tribrachiatus contortus*. Later, Kasem et al. (2020a) assigned this Subzone NP10b. This subzone is comparable to Tantawy’s (1998) Subzone NP10c. It covers from sample 26 to sample 34 (~1.9 m thick) in the Esna Formation (Table 1, Fig. 3) and is assigned to Ypresian. The incoming species that appear in this interval include *Tribrachiatus digitalis*, *T. orthostylus*, *Zygrhablithus bijugatus*, and *Neoehiazoxygus macilentus* (Table 1).

**Discoaster binodosus Zone (NP11)**

This zone covers from the HO of *Tribrachiatus contortus* to the LO of *Discoaster lodoensis* as delineated by Hay and Mohler (1967). It is comparable to Subzone CP9b of Okada and Bukry (1980) and is equivalent to Zone CNE3 of Agnini et al. (2014). At G. Nezzazat, this zone is ~3.9 m thick in the Esna Formation (Table 1, Fig. 3) and is assigned to the Ypresian Age.
Calcareous nannofossil bioevents

Considerable changes in calcareous nannofossils are associating the Paleocene–Eocene Thermal Maximum (PETM) and are useful for the placement of the P-E boundary (Romein 1979; Dupuis et al. 2003; Raffi et al. 2005; Agnini et al. 2007a, b). These changes include incoming of new species, radiations, migration towards higher latitudes, and turnovers of some taxa (Kasem et al. 2020a, b).

Several authors tracked the variations in calcareous nannofossil assemblages that associated the PETM in Egypt (e.g., Dupuis et al. 2003; Tantawy 2006a, b; Abu Shama et al. 2007; Abu 2020; Faris and Abu Shama 2007; Berggren et al. 2012; Youssef 2015; Faris and Farouk 2015; Faris et al. 2017; Faris et al. 2014; Faris et al. 2018; Youssef 2015, 2016; Kasem et al. 2020a, b). According to these studies, the P-E boundary was delineated either in Zone NP9, at the base of Zone NP10 or within Zone NP10 (Kasem et al. 2020a, b; Table 3). The biostratigraphic importance of calcareous nannofossil bioevents that predate, coincide, and postdate the P-E transition is discussed below (Fig. 5).

The LO and HO of Discoaster multiradiatus

Discoaster multiradiatus had been regarded as a reliable Paleocene zonal marker by various authors (e.g., Varol 1989; Agnini et al. 2017). Moreover, this species is supposed to go extinct in the lower portion of Zone NP11 (Perch-Nielsen 1985); however, Kasem et al. (2020a) noted that it disappears in Subzone NP10b. At Nezzazat, it disappears within the upper part of Zone NP11 (Table 1). Thus, the HO of D. multiradiatus seems to be inconsistent and unreliable for biostratigraphic correlations.

The LO of the Fasciculithus alanii group

The last radiation of Fasciculithus occurs during the latest Paleocene (Romein 1979). Previous studies revealed that F. alanii first occurs in Subzone NP9a and disappears in this subzone close to the CIE (Dupuis et al. 2003; Faris and Farouk 2015). Thus, it can distinguish the Paleocene portion of Zone NP9 from the Eocene part (Aubry and Salem 2013a). However, Faris and Abu Shama (2007) recorded F. alanii shortly above the base of Subzone NP9b.

Agnini et al. (2007b) lumped Fasciculithus richardi, F. mitreus, F. hayi, and Fasciculithus schaubii within the F. richardi group. Similarly, Agnini et al. (2014, 2017) lumped Fasciculithus species that first appear in Zone NP9 into the F. richardi group. Later, Kasem et al. (2020a) recommended assigning this group to the F. alanii group, where the occurrence and disappearance of F. alanii were the most important bioevents in this interval (Agnini et al. 2007a, b; Aubry and Salem 2013a).
At G. Nezzazat, *F. richardii* appeared in sample 10 just below the base of Zone NP9, whereas *F. thomasii* appears in sample 10 in coincidence with the base of Zone NP9 and followed by the appearance of *F. alanii* in sample 12 then *F. lillianiae* in sample 13 (Table 1). These taxa were grouped in the *F. alanii* group and seem to be useful in the approximation of the base of Zone NP9 when its zonal marker is absent or poorly preserved.

The LOs and HOs of the calcareous nannofossil excursion taxa

Calcareous nannofossil excursion taxa (CNET) or the *Rhomboaster–Discoaster* (RD) assemblage includes *Discoaster araneus* and *D. anartios*, as well as *Rhomboaster* spp. They appear suddenly and dominate the nannofossil assemblages of the lowermost Eocene (Dupuis et al. 2003; Kahn and Aubry 2004; Bown and Pearson 2009). Kahn and Aubry (2004) included *D. falcatus*, *Bomolithus supremus*, *Coccolithus bownii*, and *Toweius serotinus* in the CNET.

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**Table 3** Compilation of calcareous nannofossil bioevents used in delineation of the P-E boundary at various localities in Egypt and the present study

| Section            | Location             | Author                        | P-E boundary   | Calcareous nannofossil bioevents                                      |
|--------------------|----------------------|-------------------------------|----------------|-----------------------------------------------------------------------|
| Gebel Nezzazat     | WC Sinai             | Present study                 | NP9a/NP9b      | LOs of *D. araneus* and *Rhomboaster* spp.                            |
| Farafra-Bahariya asphaltic road | Central Sinai    | Kasem et al. (2020b)          | NP9a/NP9b      | LOs of *D. araneus*, *D. anartios*, and *Rhomboaster* spp.           |
| G. Misheiti        | Western Desert       | Kasem et al. (2020b)          | NP9a/NP9b      | LOs of *D. araneus*, *D. anartios*, and *Rhomboaster* spp.           |
| G. Naqp Assiut and G. Umm El Ghanayim | Kharga Oasis  | Faris et al. (2017)           | NP9a/NP9b      | LOs of *D. araneus* and *Rhomboaster* spp.                            |
| Taramsa            | West Qena            | Youssef (2016)                | NP9a/NP9b      | LOs of *D. araneus*, *D. anartios*, *Discoaster aegyptiacus*, *Rhomboaster* spp., and *C. eodelalda* |
| Duwi               | Quseir               | Wadi Qena                     |                |                                                                       |
| Qeryia             | Safaga               | Faris and Farouk (2015)       | NP9a/NP9b      | LOs of *D. araneus*, *D. anartios*, and *Rhomboaster* spp.           |
| Arras              | Sinai                | Faris and Farouk (2015)       | NP9a/NP9b      | LOs of *D. araneus*, *D. anartios*, and *Rhomboaster* spp.           |
| G. Matulla         | Eastern Desert       | Al Wosabi (2015)              | NP9a/NP9b      | LOs of *Rhomboaster* spp.                                            |
| G. Markha          | Farafra Oasis        | Youssef (2015)                | NP9a/NP9b      | LOs of *D. araneus* and *Rhomboaster* spp.                            |
| G. Duwi            | Central Sinai        | Khalil and Al Sawy (2014)     | NP9a/NP9b      | LOs of *Rhomboaster* spp.                                            |
| G. El-Qus Abu Said | Upper Nile Valley    | Faris and Abu Shama et al. (2007) | NP9a/NP9b     | LOs of *Rhomboaster* spp.                                            |
| W. Nukhul          | Kharga Oasis         | Faris and Abu Shama et al. (2007) | NP9a/NP9b     | LOs of *Rhomboaster* spp.                                            |
| Nag El-Quda        | Central Sinai        | Faris (2007)                  | NP9a/NP9b      | LOs of *Rhomboaster* spp.                                            |
| G. Ghanima         | Central Sinai        | Faris and Abu Shama et al. (2007) | NP9a/NP9b     | LOs of *Rhomboaster* spp.                                            |
| G. Matulla         | Themed area, Sinai   | Faris and Abu Shama (2007)    | NP9a/NP9b      | The LOs of *D. araneus*, *Rhomboaster cuspis*, *R. calcitrapa*, and *R. bitri-fida* |
| G. El Kheeb        | Central Sinai        | Faris and Salem (2007)        | NP9a/NP9b      | LOs of *Rhomboaster intermedia* and *R. calcitrapa*                   |
| G. Nukhul          | El Serai             | Tantawy (Tantawy 2006a, b)    | NP9a/NP9b      | LOs of *C. eodelalda* and *D. araneus*                                |
| W. Feiran          | East Qena            | Tantawy (Tantawy 2006a, b)    | NP9a/NP9b      | Abrupt decrease of *Fasciculithus* spp.                               |
| Taramsa            | West Qena            | Dupuis et al. (2003), and Aubry et al. (2007) | NP9a/NP9b | Barren interval followed by the LOs of *D. araneus* and *Rhomboaster* spp. |
| Dababiya Quarry (GSSP) | Nile Valley     | Faris et al. (1999)           | Within NP9    | HOs of *Fasciculithus* spp.                                          |
| Teir/Tarawan       | Kharga Oasis         | Faris et al. (1999)           | Within NP9    | LOs of *T. branlettei* and *D. binodosus*                             |
| Um El-Ghanayim     | Nile Valley          | Dupuis et al. (2003), and Aubry et al. (2007) | NP9a/NP9b | Barren interval followed by the LOs of *D. araneus* and *Rhomboaster* spp. |
| El Sheikh Eissa    | East Qena            | Tantawy (Tantawy 2006a, b)    | Within NP9    | LO of *T. branlettei*                                                |
| El-Homra El-Shanka | Esna                 | Tantawy (Tantawy 2006a, b)    | Within NP9    | LO of *T. branlettei*                                                |
| G. El-Shaghab      | Quseir               | Tantawy (1998)                | NP9a/NP9b      | LO of *C. eodelalda*                                                 |
| Duwi               | Quseir               | Tantawy (1998)                | NP9a/NP9b      | LO of *C. eodelalda*                                                 |
| Oweina             | Nile Valley          | Tantawy (1998)                | NP9a/NP9b      | LO of *C. eodelalda*                                                 |
| Abu Had            | Eastern Desert       | Tantawy (1998)                | NP9a/NP9b      | LO of *C. eodelalda*                                                 |
| Wadi Tarfa         | East Qena            | Tantawy (1998)                | NP9a/NP9b      | LO of *C. eodelalda*                                                 |
The CNET which had been considered the most remarkable change in calcareous nannofossils marks the P-E boundary and points to unusual conditions prevailed during this interval (Mutterlose et al. 2007; Table 3).

The LOs of the CNET were useful for further subdivision of Zone NP9 (Aubry and Sanfilippo 1999; Agnini et al. 2007a, b; Kasem et al. 2020a, b). Yet, the lowermost Eocene (~73 cm) is marked by a drop in carbonate content and, subsequently, the dissolution of calcareous plankton at the GSSP (Dupuis et al. 2003), which make it hard to either recognize the exact LOs of the CNET or track the precise changes in calcareous nannofossil assemblages throughout the P-E transition (Raffi et al. 2005).

At G. Nezzazat, the LOs of Rhomboaster calcitrapa, R. cuspis, Rhomboaster spinosa, and Rhomboaster bitrifida Discoaster araneus were used to differentiate the Paleocene Subzone NP9a from the Eocene Subzone NP9b (Table 1, Fig. 3). Furthermore, it was suggested to assign the interval from the HO of CNET and the top of Zone NP9 to Subzone NP9c (Aubry and Salem 2013a). However, in the present study, the CNET extends up the upper part of Zone NP10 (Table 1).

The LO of Discoaster mahmoudii

Previous studies revealed that Discoaster mahmoudii appears in Zone NP9 (Monechi et al. 2000; Aubry and Salem 2013b). Aubry and Salem (2013b) pointed to the restriction of the LO of D. mahmoudii to the top section of Zone NP9 (Subzone NP9c). However, it appears to be ~3.5 m above the base of Eocene at the Dababiya Quarry (Dupuis et al. 2003). Moreover, it was recorded within Zone NP10 at ~7 m above the CIE at G. Nezzi (Monechi et al. 2000).

At G. Nezzazat, the LO of D. mahmoudii is coincident with the base of Subzone NP9b (Table 1). Thus, the biostratigraphic significance of this event is weakened (Agnini et al. 2007a; Kasem et al. 2020a).

The LO of Blackites herculesii

This species had been introduced as Rhabdopsphaera herculea (Stradner 1969) then emended to Rhabdolithus solus (Perch-Nielsen 1971) and later revised to Blackites herculesii (Bybell and Self-Trail 1997). It had been documented in Zone NP9 (Kasem et al. 2020a, b) and was suggested to be restrictive to the topmost of this zone (Aubry and Salem 2013a). However, it was recorded in Zone NP10 at the Misheiti section in Sinai, Egypt (Kasem et al. 2020a).

At G. Nezzazat, it appears in the topmost portion of Zone NP9b (sample 21) (Table 1). Thus, it can approximate the top of Zone NP9.

The LO of Campylosphaera dela

It is hard to distinguish between Campylosphaera dela and C. eodela (Bramlette and Sullivan 1961; Hay and Mohler 1967; Bukry and Percival 1971). Therefore, they were regarded as synonyms (Kasem et al. 2020a, b).

Bukry (1973) used the LO of C. dela in subdividing the interval equivalent to Martinis’s (1971) Zone NP9. However, Kasem et al. (2020a) noted this species in Zone NP7/8. Thus, the stratigraphic range of this taxon is inconsistent (Agnini et al. 2007b).

In this study, C. dela and C. eodela were viewed as a single taxon and first occurred in the topmost of NP9a (sample 18) just below the base of Subzone NP9b (Table 1). As a result of the inconsistent ranges and taxonomic disputes, C. dela was excluded from being a reliable biostratigraphic event (Kasem et al. 2020a).

The LO of Discoaster binodosus

It had been proposed to use the LO of Discoaster binodosus in the approximation of the top of Zone NP9 when its zonal marker, Tribrachiatus bramlettei, is scarce or absent (Perch-Nielsen 1985); however, the range of this taxon is inconsistent where it was recorded in the lower part of Zone NP9 (Faris and Salem 2007) and within Zone NP10 (Kasem et al. 2020a, b). Thus, the biostratigraphic significance of these bioevent is weakened (Faris and Salem 2007; Kasem et al. 2020a, b).

In this study, D. binodosus appears within Subzone NP9b, but it is sporadic in the early stage of its range and its LCO is coincident with the base of Subzone NP10b (Table 1).

The drop in the frequency and diversity of Fasciculithus species

Several studies documented a severe decrease in the diversity of Fasciculithus close to the upper limit of Paleocene (e.g., Agnini et al. 2007a, b; Kasem et al. 2020a).

In this study, the diversity of Fasciculithus decreases from 10 species in sample 17 slightly below the ground of Subzone NP9b into 5 and 3 just below and above the P-E boundary at G. Nezzazat (Table 1, Fig. 7). Furthermore, the abundance of Fasciculithus/Lithoptychius drops from 179 specimens per 50 fields of view (S/50 FOV) in sample 17 into 51 and 11S/50 FOV specimens just below and above the base of the Eocene (Table 1, Fig. 7).

The HOs of Fasciculithus tympaniformis and F. alanii group

Fasciculithus taxa diversified in the uppermost Paleocene and go extinct shortly above this interval (Martini 1971;
Raffi et al. 2005; Tantawy 2006a, b). Thus, it can approximately locate the top of Zone NP9 when the zonal marker *Tribrachiatus bramletti* is poorly preserved or absent (Perch-Nielsen 1985).

Agnini et al. (2014, 2017) used the HO of *F. tympaniformis* to mark the upper limit of the earliest Eocene Zone CNE1. However, several *Fasciculithus* species were recorded up to Zone NP10 (Romein 1979; Bralower and Mutterlose 1995; Aubry 1996; Raffi et al. 2005; Agnini et al. 2007b).

Kasem et al. (2020a) noted that *F. tympaniformis* extends with common occurrences up to the topmost section of Zone NP10. In this study, *F. clinatus, F. involutus, Fasciculithus billii, Fasciculithus pileatus, Fasciculithus janii, and Fasciculithus bitectus* disappear in Zone NP9 (Table 1). However, *F. tympaniformis* occurs with common occurrences up to Zone NP9 and extends with rare and discontinuous occurrences up to Zone NP11 (Table 1), which indicates reworking of these specimens and supports the biostratigraphic reliability of its HCO rather than the HO that is an inconsistent bioevent, possibly due to the diachronous nature of the HO of this taxon or discrepancies concerning the taxonomic position of *T. bramletti* (Tantawy 2006a, b; Kasem et al. 2020a).

Agnini et al. (2014, 2017) documented the HO of *F. richardii* group below the LOS of *Rhomboaster* spp. and suggested delineating the base of the entry of the Eocene (base of Zone CNE1) based on the HO of this group. Abu Shama et al. (2007) recorded the HO of *F. alanii* in coincidence with the base of Subzone NP9b. Thus, the HO of *F. alanii* group can approximate the lower limit of Subzone NP9b when its subzonal markers are absent or poorly preserved.

In this study, the LOS of *F. lillianiae* and *F. thomasi* are coincident with the base of Subzone NP9b, whereas *F. alanii* extends within Subzone NP9b (Table 1).

The acme of Ericsonia subpertusa

An increased frequency of *Ericsonia subpertusa* was documented across the P-E interval (Dupuis et al. 2003; Abu Shama et al. 2007; Kasem et al. 2020a).

At G. Nezzazat, *E. subpertusa* dominates the latest Paleocene–earliest Eocene nannofossil assemblages; nevertheless, it suddenly increases from 121 S/50 FOV in samples 17 to 270 and 446 S/50 FOV just below and above the P-E boundary (Table 1).

The LO of Discoaster diastypus

The appearances of both *Tribrachiatus contortus* plus *Discoaster diastypus* had been regarded synchronous (Raffi et al. 2005). Therefore, Bukry (1973) used their LOS to limit the base of the *Discoaster diastypus* Zone of Bukry (1973) that is correlative to Zone CP9 of Okada and Bukry (1980). The co-occurrences of *T. bramletti* and *D. diastypus* were recorded in some areas in Egypt (e.g., Abu Shama et al. 2007), yet their LOS were not concurrent in other sections (Tantawy 2006a, b; Kasem et al. 2020a).

At G. Nezzazat, the LO of *D. diastypus* precedes the appearance of *T. bramletti* (Table 1) that supports the diachronous nature of *D. diastypus* and/or *T. bramletti*.

The LO of Sphenolithus radians

The appearance of *Sphenolithus radians* and the disappearance of species *Tribrachiatus contortus* were very associated; therefore, the LO of *S. radians* can approximate the entry of Zone NP11 when *T. contortus* is absent (Perch-Nielsen 1985). Kasem et al. (2020a) noted that the LOS of *S. radians* and *T. contortus* are coincident at Misheiti in Sinai, Egypt. However, *S. radians* were recorded up to the topmost of Zone NP11 (Abu Shama et al. 2007; Al Wosabi 2015).

At G. Nezzazat, the LO of *S. radians* and the HO of *T. contortus* are coincident (Table 1), which supports the reliability of this bioevents in approximation of the base of Zone NP11.

The LO of Zygrhablithus bijugatus

The LO of *Zygrhablithus bijugatus* at the base of Subzone NP9b was recorded (Kasem et al. 2020b). In this study, it occurs at the base of Subzone NP10b (Table 1). This indicates inconsistent LO of this taxon.

The LO and increased frequency of Neochiastozygus junctus

Kasem et al. (2020b) noted that *Neochiastozygus junctus* first occurs in Subzone NP9b and blooms just above the top of Zone NP9. The increased frequency of this taxon was used to limit the basis of Zone NP10 (Abu Shama et al. 2007).

At G. Nezzazat, the LO of *N. junctus* is coincident with the ground of Zone NP9b and increases in frequency in the upper section of Zone NP9 and in the lower part of Zone NP10 in this study (Table 1). Thus, this bioevent can approximate this interval in case of the absence of the zonal markers (Table 1).

The LOs and HOs of Tribrachiatus species

Genus *Tribrachiatus* is useful in biostratigraphic zonation (Martini 1971). Previous investigations documented that *T. bramletti* appears slightly above the base of the Eocene (Agnini et al. 2007a, b). Yet its earliest occurrences are rare to very rare and sporadic (Agnini et al. 2007a).

Thus, Faris et al. (2015) suggested depending on the LCO of this taxon to delineate the lower limit of Zone NP10.
Furthermore, the LO of *T. bramlettei* is inconsistent (Agnini et al. 2007b), probably due to the diachronous nature of this bioevent and/or dissolution that affects the lowermost Eocene sediments (Agnini et al. 2007b). In addition, *T. bramlettei* and *Rhomboaster cuspis* were considered synonyms by some authors (e.g., Von Salis et al. 2000); whereas others count them two different species (e.g., Raffi et al. 2005; Kasem et al. 2020a, b; the present study).

Furthermore, several authors reported that *Tribrachiatus bramlettei* disappears in the uppermost of Zone NP10, shortly after the HO of *T. contortus* and below the LO of *T. orthostylus* (e.g., Aubry and Sanfilippo 1999). Nevertheless, *T. bramlettei* extends up to the top of Zone NP10 at Galala Mountains, Eastern Desert, Egypt (Marzouk and Scheibner 2003).

In this study, the HO of *T. bramlettei* occurs in sample 33 at a level somewhat below the end of Zone NP10 and slightly below the LCtO of *T. orthostylus* (Table 1).

The taxonomic position of *Tribrachiatus digitalis* underwent criticism (Raffi et al. 2005), and consequently, its biostratigraphic significance is weakened (Kasem et al. 2020a, b).
On the other hand, *T. contortus* is significant in biostratigraphic studies (Martini 1971), where its HO delineates the top of Zone NP10, and its LO is useful in subdividing Zone NP10 into Subzones NP10a and NP10b (Kasem et al. 2020a, b, and the present study).

The appearance of *T. orthostylus* can approximate the topmost of Zone NP10 when its zonal marker is absent (Perch-Nielsen 1985). In this study, *T. orthostylus* appears just below the base of Subzone NP10b (Table 1).

**The calcareous nannofossil species richness and abundance**

Calcareous nannofossils reached their maximum diversity in the Paleogene interval within Zone NP9 (Perch-Nielsen 1985). This had been confirmed in this study, where several incoming species occur in Zone NP9 such as *Discoaster falcatus*, *D. binodosus*, *Discoaster medius*, *D. mahmoudii*, *D. araneus*, *D. anartios*, *Discoaster lenticularis*, *D. diasty pus*, *D. barbadiensis*, *Rhomboaster cuspid*, *R. calcitrapa*, *R. sp.", *R. bitriffida*, *Fasciculithus lillianiae*, *Neochiasmatozygus distentus*, *N. pusillus*, *Lanternithus simplex*, *Pontosphaera distincta*, *P. plana*, *Chiasmolithus nitidus*, *Calcidiscus sp.*, *Blackites herculies*, and *Ellipsolithus distichus* (Table 1). Nevertheless, the abundance of calcareous nannofossil assemblages shows increased frequency close to the P-E boundary (Table 1, Fig. 6).

**Paleoecology**

The changes in temperature and nutrient availability affect the distribution of calcareous nannofossil; therefore, they can be used for tracking climatic changes (e.g., Bralower 2002; Bernaola et al. 2007). Oligotrophic conditions with warm waters that characterize low- to middle-latitude areas are sufficient for flourishing modern calcareous nannofossil assemblages (e.g., Watkins 1989; Bralower 2002; Bornemann 2003; Fuqua et al. 2008).

On the other hand, eutrophic forms are more common in cold nutrient-rich waters at high-latitude areas (Bralower 2002). Several investigations revealed the paleoecological preferences of several calcareous nannofossil taxa (e.g., Wei and Wise Jr 1990; Fuqua et al. 2008, etc.).

In this study, the variations in paleotemperature and paleofertility were tracked depending on the changes in abundances of calcareous nannofossil species that have certain ecological preferences (Fig. 7). The taxa that had been regarded in the previously mentioned references as warm water taxa include *Ericsonia subpertusa*, *Coccolithus pelagicus*, *Fasciculithus spp.*, *Lithoptychius spp.*, *Discoaster spp.*, *Heliolithus kleinpelli*, *Rhomboaster spp.*, *Tribrachiatus spp.*, *Bomolithus spp.*, *Zygrhabliithus bijugatus*, *Pontosphaera spp.*, *Sphenolithus spp.*, *Thoracosphaera operculata*, and *Thoracosphaera saxea*.
Based on calcareous nannofossils, the P-E boundary had been placed either in Zone NP9, at the NP9/10 zonal limit, or in Zone NP10 (Martini 1971; Bukry 1973; Bolle et al. 2000). At the Dababiya Quarry section in Egypt that was ratified as the GSSP for the base of Eocene, the P-E boundary was placed at the ground of Subzone NP9b (Dupuis et al. 2003). This boundary is marked by an abrupt decrease in carbon isotope and is coincident with the base of a distinctive bed of the Dababiya Quarry Member (Aubry et al. 2007).

At G. Nezzazat, this boundary had been placed at the NP9a/NP9b subzonal boundary that is denoted by the appearances of Discoaster araneus and/or Rhomboaster spp. (Table 1). This boundary is marked by the LOs of Rhomboaster cuspidis, R. calcitrapa, R. spineus, R. bitrifida, D. araneus, and Chiasmolithus nitidus as well as the HOs of Fasciculithus thomasi and F. lillianiae (Table 1). In addition, this level is coincident with the base of a black shale bed assigned to the DQM (correlative to DQB 4 of Tantawy 2006, ) in the lower portion of the Esna Formation (Fig. 3).

This reveals the presence of a gap represented by the missing of DQB2 to DQB5, but not detected by the calcareous nannofossils. Similar results were noted in different sections in Sinai and others in Egypt (see Aubry and Salem 2013b; Obaidalla et al. 2017; Kasem et al. 2020a, b for more discussion). Variations in calcareous nannofossil assemblages reveal warming and oligotrophic conditions prevailed during the P-E transition at G. Nezzazat in agreement with results from the GSSP as well as sections from various areas in the world (Dupuis et al. 2003; Bernaola et al. 2007; Mutterlose et al. 2007; Raffi et al. 2009; Kasem et al. 2020a, b).

Conclusions

Lithostratigraphic and calcareous nannofossil biostratigraphic investigations were carried out in this study at the Gebel Nezzazat section in Central Sinai, Egypt. The interval investigated extends throughout the Tarawan, Esna, and Thebes Formations. The Esna Formation has been partitioned into the Hanadi, Dababiya Quarry, Mahmiya, and Abu Had Members.

Five calcareous nannofossil biozones were recognized (NP7/8 through NP12). Zone NP9 had been subdivided into Subzones NP9a and NP9b depending on the LOs of Rhomboaster spp. and/or Discoaster araneus. Moreover, the LO of T. contortus had been used to partition Zone NP10 into Subzones NP10a and NP10b.

Exclusion of Tribrachiatus digitalis from being a reliable marker was recommended in this study. Stratigraphic overlaps between T. branlettei and T. contortus as well as between T. contortus and T. orthostylus were noted, indicating the completeness of this interval. Most Fasciculithus taxa (F. clinatus, F. involutus, F. billii, F. pileatus, F. janii,

Remarks

The Paleocene–Eocene transition was marked by geochemical, lithological, and biotic variations (Dupuis et al. 2003). Based on calcareous nannofossils, the P-E boundary had been placed either in Zone NP9, at the NP9/10 zonal limit, or in Zone NP10 (Martini 1971; Bukry 1973; Bolle et al. 2000).
and *F. bitectus*) disappear within Zone NP9. *Fasciculithus tympaniformis* occurs with common occurrences up to the top of Zone NP9 and extends with sporadic occurrences up to Zone NP11 (Table 1), supporting the insignificance of the HO of *F. tympaniformis* in biostratigraphy.

Subsequent appearances of *F. richardii*, *F. thomasii*, *F. alanii*, and *F. lillianiae* are closely related to the base of Zone NP9. These species were grouped in the *F. alanii* group, in which the LO and HO can approximate the entry of Subzone NP9a and Subzone NP9b, respectively, when their zonal markers are absent or poorly preserved.

Furthermore, *Blackites herculaesi* can approximate the top of Zone NP9. The LOs of *D. mahmoudii*, *D. binodosus*, *D. diastypus*, *Zygrrhabilithus bijugatus*, and *Campyllosphaera dela* are not reliable bioevents in biostratigraphy. On the other hand, the LO and increased frequency of *Neochias-totyzus junctus* can approximate the upper part of Zone NP9 and the lower part of Zone NP10.

Moreover, the LO of *S. radians* is a reliable bioevent for approximating the base of Zone NP11. *Triradiatothecus ortho-stylus* appears just below the base of Subzone NP10b. The abundance of calcareous nannofossils shows an increased frequency close to the P-E boundary. *Ericsonia subpertusa* dominates the calcareous nannofossil assemblages of the examined interval and suddenly increases above the P-E boundary. On contrast, a drop in the diversity of *Fasiculithus* is closely related to the P-E transition.

The P-E boundary at G. Nezzazat was placed at the base of Subzone NP9b that is denoted by the LOs of *Discocaster araneus*, *D. anartios*, and/or *Rhomboaster* spp. This boundary is coincident with the base of a black shale bed assigned to the DQM in the lower portion of the Esna Formation (Fig. 3). This reveals the presence of a small gap represented by the missing of DQB2 to DQB5. The variations in calcareous nannofossils reveal warming and oligotrophic conditions prevailed through the deposition of the upper Paleocene–lowermost Eocene at the study section.

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Declarations

Conflict of interest The authors declare that they have no competing interests.

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