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FitzPatrick, M

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Meriel E.J. FitzPatrick, David A. Forber, Malcolm B. Hart

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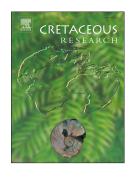
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1	Dinocyst stratigraphy and palaeoenvironmental interpretation of
2	the Cretaceous/Paleogene boundary at Stevns Klint, Denmark
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4	Meriel E.J. FitzPatrick ^{a*} , David A. Forber ^b , Malcolm B. Hart ^a
5	
6	^a School of Geography, Earth and Environmental Sciences, Plymouth University,
7	Drake Circus, Plymouth PL4 8AA, United Kingdom
8	^b College of Engineering, Mathematical & Physical Sciences, Harrison Building,
9	Streatham Campus, University of Exeter EX4 4QF, United Kingdom
10	
11	*Corresponding author
12	E-mail: me.fitzpatrick@plymouth.ac.uk (M.E.J. FitzPatrick)
13	
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15	ABSTRACT
16	A re-examination of the uppermost Maastrichtian chalks and an expanded section of
17	the Fish Clay (Fiskeler Member) undertaken at Stevns Klint, Denmark, one of the
18	classic outcrops of the Cretaceous/Palaeogene boundary, has identified diverse and
19	abundant dinoflagellate cyst assemblages, which are compared with earlier
20	dinoflagellate cyst (dinocyst) studies. The uppermost part of the white coccolith chalk
21	(Sigerslev Member) of the uppermost Maastrichtian is placed in the Palynodinium
22	grallator Zone. The overlying 'Grey Chalk' (Højerup Member) represents a shallower-
23	water marine succession that is characterised by a series of dune-like structures.
24	The last common occurrence of <i>P. grallator</i> is within the Højerup Member, confirming
25	the Danian age for the Fish Clay (Fiskeler Member) and a latest Maastrichtian age
26	for the P. grallator Zone. Within the Fiskeler Member, Damassadinium californicum
27	(previously known as Danea californica), a key biostratigraphical marker, is only
28	intermittently present, while the distributions of Senoniasphaera inornata and

29	Xenicodinium reticulatum provide useful biostratigraphic control. The well-preserved
30	dinocysts of the Fiskeler Member are replaced by reduced assemblages in the
31	Cerithium Limestone Member with only pollen grains recorded.
32 33	Keywords: K/Pg boundary, dinoflagellate cysts, biostratigraphy, Fiskeler Member
34	
35	1. Introduction
36	In an investigation of the Stevns Klint, Maastrichtian-Paleogene succession, Hart et
37	al. (2005) identify an expanded section of the boundary clay at Kulstirenden. As
38	earlier studies of the dinocyst assemblages had been undertaken on much thinner
39	successions of the boundary clay, this research aims to record the dinocyst changes
40	in more detail across the K/Pg boundary.
41	While the basal Paleocene 'Global Stratigraphic Section and Point' (GSSP) is

42 located in Tunisia (Molina et al., 2006), the Cretaceous/Paleogene (K/Pg) boundary at Stevns Klint, Denmark, remains an important reference point (Damholt and Surlyk, 43 44 2012). Stevns Klint includes the original site where the K/Pg boundary was first 45 described by Desor (1847), lying between the coccolith chalks of the Maastrichtian 46 and the bryozoan-rich chalks of the lower Danian. Desor (1847) also included the Faxe (Fakse) Quarry in his original definition of the Danian. While this working quarry 47 48 is located only 17 km to the west of Rødvig, and has exceptionally interesting 49 occurrences of bryozoan mounds and cool-water corals (Bernecker and Weidlich, 50 1990, 2005; Willumsen, 1995; Surlyk and Håkansson, 1999), other invertebrate 51 fossils (Graversen, 2001), gastroliths (Noe-Nygaard, 1975) and evidence of 52 coprolites (Milàn, 2010), the K/Pg boundary was not exposed there during our 53 fieldwork. The old quarries at Holtug and Boesdal contain similar features and were 54 also un-sampled.

55 The K/Pg transition is exposed almost continuously along the ~14 km-long cliff from Rødvig in the south to Bøgeskov in the north (Fig. 1). As reported by Hart et al. 56 (2004, 2005), the Fish Clay, now renamed the Fiskeler Member by Surlyk et al. 57 58 (2006), has been found to display considerable variation along the length of the cliffs at the eastern end of the Stevns Peninsula. At Kulstirenden the Fiskeler Member 59 attains its maximum thickness (~50 cm). It is worth noting that in the definition of the 60 GSSP the greater thickness of the boundary clay at El Kef, as compared to Stevns 61 Klint, was used (Molina et al., 2006, p. 265) as evidence of the greater continuity of 62 sedimentation in the Tunisian section. While this is certainly true at Rødvig and 63 Højerup, the Kulstirenden exposure is guite comparable in thickness to that at El Kef. 64 In their definition of the basal-Paleogene GSSP at El Kef, Molina et al. (2006, p. 265) 65 66 recorded that there was no 'mass extinction' of dinocysts across the boundary and Damassadinium californicum – while at the base of the boundary clay in Scandinavia 67 - appears 10 cm above the base of the boundary clay in the El Kef succession 68 69 (Brinkhuis and Leereveld, 1988; Brinkhuis and Zachariasse, 1988; Donze and Méon, 1997). Despite the relatively low numbers of *D. californicum* three of the voting 70 members of the Paleocene Working Group supported the FO of *D. californicum* as 71 72 marking the boundary while 11 voted for the base of the boundary clay and, therefore, the impact event as the marker for the GSSP. 73

In recent years there have been a number of papers on the Stevns Klint K/Pg
boundary succession, including a revision of the lithostratigraphy (Surlyk et al.,
2006). The micropalaeontology and stable isotope stratigraphy have been described,
in some detail, by Schmitz et al. (1992), Hart et al. (2004, 2005) and Rasmussen et
al. (2005). Missing from all these recent investigations is any re-assessment of the

organic-walled dinocyst stratigraphy which has, in the past, been somewhatproblematic.

There have been a number of dinocyst investigations at Stevns Klint, dating 81 82 back to the 1960s. These include the works of Morgenroth (1968), Wilson (1974), J.M. Hansen (1977) and H.J. Hansen et al. (1986). Unfortunately, these 83 84 investigations all used samples collected from the thinner sections of the Fiskeler Member. Hultberg (1985, 1986, 1987) did sample from slightly thicker parts of the 85 Fiskeler Member, but the representativeness of some of his results has been 86 87 questioned (Brinkhuis et al., 1998). In particular, two of Hultberg's (1986) Fiskeler Member samples contained a monospecific or near-monospecific assemblage of 88 89 Manumiella druggii, which Hultberg interpreted as indicative of a lagoonal or 90 brackish-water environment. J.M. Hansen (1977), H.J. Hansen et al. (1986) and Brinkhuis et al. (1998) all found more diverse assemblages at these locations. 91 92 Furthermore, despite sampling from the slightly thicker parts of the Fiskeler Member, 93 Hultberg only had eight samples within that member that actually contained dinocysts. Isotopic data from Stevns Klint presented by Hart et al. (2005) indicate 94 potential environmental changes on a scale that Hultberg's sampling resolution 95 would not be expected to achieve. 96

97 The primary objectives of our research are, therefore: 1) to increase the 98 sampling resolution through the Fiskeler Member, which will provide a more detailed 99 analysis of the dinocyst assemblages and species distributions across the K/Pg 100 boundary at Stevns Klint; and 2) to use this new information on dinocyst stratigraphy 101 to make comparisons with the distribution of these taxa from a number of other K/Pg 102 boundary sites.

103

104 **2.** Geological setting

105 Forming the edge of the Stevns Peninsula in eastern Sjaelland, Stevns Klint is 106 located ~40 km south of Copenhagen (Fig. 1). The cliff, which varies in height 107 between 20 and 41 m, hosts several guarries along its 14 km length (Surlyk, 1979; Hart et al., 2004, 2005; Surlyk et al., 2006; Damholt and Surlyk, 2012). One of the 108 109 first descriptions of Stevns Klint and the classic Cretaceous–Paleogene boundary sections, dates back to the eighteenth century (Abildgaard, 1759), but it was not until 110 111 a century later that the complicated stratigraphy of the boundary was appreciated. Desor (1847) was the first to recognise the K/Pg boundary, while Rosenkrantz 112 113 (1924) later described the palaeontology of the Maastrichtian and Danian strata. 114 Together with Faxe (Fakse) Quarry, Stevns Klint constitutes the classic type area of 115 the Danian Stage in Europe (Surlyk, 1979), but has been passed over as the Global Stratotype Section and Point for the base of the Danian because of its rather 116 complex stratigraphy and lack of many key species of planktic foraminifera (Hart et 117 al., 2004, 2005). The two localities used in this investigation are at Højerup, where 118 119 most previous field studies have taken place (Schmitz et al., 1992; Hart et al., 2004, 120 2005) and Kulstirenden, which is located approximately 8 km to the north of Højerup. 121 Although laterally variable over the ~14-km long cliff, the composite stratigraphical sequence at Stevns Klint can be divided into five major lithological 122 units (Hart et al., 2005; see Fig. 2 here) all of which have now been re-named by 123 124 Surlyk et al. (2006). Except where stated otherwise, the following lithological details are derived from Hart et al. (2004, 2005) and Surlyk et al. (2006). 125 Low, rounded chalk mounds constitute the lowermost 5 -10 m of the visible white 126 coccolith chalk which is now known as the Sigerslev Member of the Tor Formation. 127 The mounds grade upwards into ~20 m of Zoophycos-rich, horizontally-bedded 128

129 chalks, which host gently undulating lines of (generally) small black flints. Two 130 features are present near the top of this member. There is a prominent, laterally-131 continuous nodular flint layer approximately 3–4 m below the Fiskeler Member (Fig. 132 3) at Højerup. At Harvig there are two phosphatised 'hardgrounds', approximately 100 mm apart, that are located ~ 30 cm above this flint horizon. Along the section 133 134 northwards, towards Højerup, these merge into a single horizon. This line of flints (and the overlying hardground) can be followed along almost all of Stevns Klint and 135 appear to be a reliable datum level (Surlyk et al. 2006, plates 7,10). At Rødvig this 136 137 line of flints is just below beach level and rarely visible. On the coast to the southeast of Kulstirenden it is visible in the cliff just below the Fiskeler Member as the 138 139 overlying Højerup Member is locally absent. 140 The Højerup Member is <4 m thick at Højerup and consists of grey-coloured, bryozoan- and Thalassinoides-rich chalk that was deposited in low, elongate, 141 asymmetric mounds (3.5 to 4 m in height and with a basal dimension of ~25 m). Flint 142 143 bands populate parts of the more gently-dipping northern slopes (~5° compared to ~15° for the southern slopes). At Kulstirenden, the Højerup Member is very thin or 144 completely absent where the Fiskeler Member is at its thickest (Hart et al., 2005, fig. 145 7). At this location, the hardground at the boundary between the Sigerslev and 146

147 Højerup members is immediately below the Fiskeler Member, almost giving the

148 impression of a phosphatised sub–K/Pg boundary hiatus.

The Fiskeler Member lies within the troughs formed by the mounds in the underlying chalk. At some locations along Stevns Klint it can be missing as a result of the erosion at the top of the Cerithium Limestone Member. At Højerup, the maximum thickness of the clay is 7 cm (Christensen et al., 1973). At Kulstirenden the clay attains a maximum thickness of 45–50 cm in the centre of a large trough but

154 thins away from this centre so that at some locations it is absent from the succession (Hart et al., 2005). Where the clay is thickest, at Kulstirenden, the Højerup Member 155 is absent and the clay lies just above the hardground that marks the boundary 156 157 between the Sigerslev and Højerup members. Christensen et al. (1973) divided the Fiskeler Member into four units: 1) the 158 159 lowermost grey, laminated marl; 2) black marl, with pyrite concretions, that is often stained reddish-yellow in weathered outcrops; 3) black, laminated marl; 4) light grey 160 marl also containing pyrite concretions. Figure 4 shows the Fiskeler Member at 161 162 Kulstirenden where most of the samples for this study were collected. The Fiskeler Member is observed to grade into the overlying Cerithium 163 164 Limestone Member, with small lenses of chalk near the transition (making the 165 boundary imprecise). Schmitz et al. (1992) reported the presence of a hiatus marked by an erosion surface but this has not been confirmed, especially in the expanded 166 succession at Kulstirenden. The Cerithium Limestone Member is formed of <1 m of 167 168 yellowish-white hard chalk with two hardgrounds at its uppermost boundary at Kulstirenden, which converge into a single hardground at other locations, including 169

Højerup and Rødvig. This surface is regionally important as a sequence boundary 170 (Rasmussen et al., 2005; Hart et al., 2011) and can be seen cross-cutting parts of 171 the succession along the length of Stevns Klint. The overlying, bryozoan-rich chalks 172 (Korsnæb Member) are cream-coloured, bryozoan-rich packstones and wackestones 173 that occur as large asymmetric mounds. These mounds are on a much larger scale 174 than those recorded from the Højerup Member and are best seen east of Rødvig, in 175 176 the old quarries of Boesdal and immediately north of the old church at Højerup (where they were previously quarried). 177

178

179 **3.** Material and methods

Over the last few years several suites of samples have been collected from various locations along Stevns Klint by Malcolm Hart, Sean Feist, Eckhart Håkansson and Claus Heinberg. Of these, thirty-one were selected for palynological processing and analysis, with two slides prepared for each sample. For each slide 300 dinocyst specimens were counted; bisaccate pollen, spores and foraminiferal test linings were noted but did not form part of this count. The Gonyaulacoid/Peridinioid (G/P) and Protoperidinioid /Gonyaulacoid (P/G) ratios were calculated.

The G/P ratio calculated here follows Harland (1973), whereby G/P = nG/nP; n=number of species (identified as G/P ^{species}). The G/P ratio was also calculated using G/P = nG/nP; n=number of specimens (identified as G/P ^{specimens}) in order to allow comparison with some other authors. The P/G ratio follows Versteegh (1994), whereby P/G = nP/(nP+nG); n = number of specimens.

Twenty-seven of the thirty-one samples selected for our investigation were 192 193 collected from Kulstirenden. The K8 samples came from the white coccolith chalk of 194 the Sigerslev Member immediately below the prominent flint horizon, only two of which were sent for processing (K8/7 and K8/5). The KUL/I and KUL/K samples 195 196 were collected between the prominent flint horizon and the hardground(s) that usually mark the boundary between the Sigerslev and Højerup members. To 197 198 complete the Maastrichtian succession, four samples (L5 series) were selected from 199 the Højerup Member at Højerup, because this member is missing in the accessible section at Kulstirenden. Sixteen, mostly contiguous samples (KU14-29), represent 200 the Fiskeler Member at Kulstirenden. Samples KU45–50 (excluding KU48, which 201 was not sent for processing) were collected from the Cerithium Limestone Member. 202 By using this selection of samples there has been a complete coverage of the K/Pg 203

transition (Fig. 5). The sampling of the Fiskeler Member is particularly detailed and
this has allowed a more complete analysis of the dinocyst stratigraphy than has
previously been the case.

207

208 4. Palynological processing

209 The full processing procedure is, perhaps unusually, described here as it was the sample preparation methods adopted by Hultberg (1985, 1986, 1987) that concerned 210 Brinkhuis et al. (1998). All samples were processed in the Palynology Laboratory of 211 the Natural History Museum (London) in February 2005 by Jonah Chitolie. Samples 212 213 were processed using standard (see Batten, 1999) palynological techniques 214 (hydrochloric acid followed by hydrofluoric acid for demineralisation). Oxidation was 215 not deemed necessary, however, because of the well-preserved nature of the palynomorphs. 216

The chalk samples all weighed between 119 and 166 g, while the clay 217 218 samples weighed 9–13 g. All the samples were cleaned of surface contamination 219 either by scraping the outer layer of the sediment with a stainless steel scalpel (for hard chalks), or by brushing with a dry nylon toothbrush (for clays, marls and soft 220 221 chalks). The clean samples were oven-dried at 100° for approximately ten hours. After drying and weighing (chalk samples weighed 50–118 g; clay samples weighed 222 5-8 g), the samples were placed in glass beakers: 2-litre beakers for the chalk and 1 223 224 litre for the clay, 100 ml of distilled water was added to each sample, subsequently followed by a small amount of 50% hydrochloric acid, with any reaction being 225 allowed to subside before the addition of more acid. The chalk samples foamed due 226 to their high reactivity. 227

228 After being washed through a 15 µm sieve with filtered tap water, the residues 229 were returned to the glass beakers where concentrated hydrochloric acid was added to eliminate un-dissolved carbonates. After further washing, the samples were 230 231 placed in 250 ml polypropylene beakers where 50% hydrofluoric acid was added and left overnight. Samples were then washed through a 15 µm polyester sieve. 232 The recovered residues, in almost all of the samples, contained well-233 234 preserved palynomorphs, some of which were transparent. All residues were 235 examined before mounting and any that were found to contain calcium fluoride precipitate were treated with 50% hydrochloric acid and heated at 80°C for 236 approximately one hour. The residues were divided in two, with one part stained 237 238 using Bismark brown R.

For each sample two glass coverslips were cleaned and placed on a warm plate. A small amount of residue was placed in a vial and mixed with polyvinyl acetate. The residue was pipetted onto the coverslip and left to dry. A small amount of Elvacite was used as the mounting medium and was placed on each slide and then inverted onto the coverslip. Elvacite is made by dissolving 20 g of Elvacite 2044 resin (a trade name for methacrylate resin) in 35 ml of xylene. The slides were kept in a fume hood over night and then placed in a drying oven at 80°C for two hours.

246

247

5. Dinocyst distribution

A range of well-known taxa have been identified in this study, all of which are listed and counts given in Appendix 1, while the ranges of selected taxa are shown in Table 1 and images of selected taxa are shown in Figure 6. Taxonomic classification follows Fensome et al. (2008), although, it is accepted that some aspects of the taxonomy may be in need of further investigation and revision.

253	
254	5.1. Kulstirenden – Sigerslev Member (K8 samples)
255	Sample K8/7 was dominated by <i>P. grallator</i> , which recorded ~30% (112 specimens)
256	of the count. The Achomosphaera-Spiniferites complex of dinocysts, together
257	accounted for 80 specimens or ~ 25%, with A. ramulifera, S. ramosus multibrevis
258	and S. ramosus reticulates being particularly common. The Achomosphaera-
259	Spiniferites complex of dinocysts is here defined as a grouping of all species and
260	subspecies belonging to these two genera. Palaeoperidinium pyrophorum showed its
261	highest abundance at this level with 15 specimens counted. This produces a G/P
262	^{species} ratio of 25; G/P ^{specimens} ratio of 293 and a P/G ratio of 0.05 (see Fig. 7).
263	Sample K8/5 was similar to K8/7 with P. grallator comprising 180 specimens or 30%
264	of the cysts counted, and the Achomosphaera-Spiniferites complex of dinocysts
265	accounting for 79 specimens, c. 25% of the count, S. ramosus multibrevis, however,
266	was not quite as abundant as in K8/5, while <i>P. pyrophorum</i> was only observed once.
267	Species diversity was 25 and 21 for K8/7 and K8/5, respectively, similar to the L5
268	series of samples from Højerup. This produces the following ratios: G/P ^{species} =21;
269	$G/P^{specimens} = 305$ and $P/G = 0$.

270

271 5.2. Kulstirenden – Højerup Member (KUL samples)

KUL/K produced a near-monospecific assemblage of *P. grallator* (235 specimens;
87.7%). Of the other taxa only *Oligosphaeridium complex* and *S. ramosus gracilis*,
were found in abundances greater than one. *Impagidinium cristatum* was relatively
common (32 specimens). There were no peridiniacean cysts resulting in a G/P ^{species}
ratio of 0. Only 17 species and sub-species were observed in this sample, the lowest

diversity of dinocysts encountered (see Fig.7). KUL/I was barren and did not containany dinocysts.

279

280 5.3. Højerup – Højerup Member (L5 samples)

These four samples showed a dinocyst assemblage almost completely dominated by 281 282 gonyaulacacean cysts. Species diversity varied between 27–30 (Fig. 7). The only peridiniacean cyst observed in these samples was Palaeoperidinium pyrophorum, 283 284 which was found in samples L5/60-65 and L5/290-295, but was absent from the other two. Where it did occur, abundance was low: 3 specimens in the former 285 sample and 8 specimens in the latter. Consequently, the G/P^{species} ratio increases 286 287 and decreases from 26 to 0 and from 29 to 0 from the lowest to uppermost samples (see Fig. 7). Similarly the G/P^{specimens} ratio varied between 0–103. The P/G ratio 288 varied from 0.01 to 0 and 0.03 to 0 from the lowest to the uppermost samples (see 289 Fig. 7). Collectively, the Achomosphaera-Spiniferites complex cysts are the most 290 291 abundant species (78–143 specimens) in the Højerup Member samples, comprising between 24.6–45 % of the count. Within this complex, A. ramulifera, A. sagena 292 sagena, and several subspecies of Spiniferites ramosus were identified (see Fig. 6). 293 294 Tanyosphaeridium xanthiopyxides is a common species throughout all of the samples (14–37 specimens). Impagidinium cristatum, is also common; its 295 abundance is consistently higher in the older samples (16/12 specimens in L5/60-65 296 297 and L5/125-130 respectively). Of the other taxa, only P. grallator, Spiniferites ramosus ramosus, Spiniferites ramosus gracilis and Hystrichosphaeridium tubiferum 298 299 appear in any significant numbers. In addition, abundant algae and foraminiferal 300 linings were noted in all of the samples, particularly L5/125–130.

301

302 5.4. Kulstirenden – Fiskeler Member (KU samples)

Of the species that were recorded below the boundary, *I. cristatum* was absent and 303 304 *P. grallator* disappears above KU15. The *Achomosphaera–Spiniferites* complex 305 cysts were still present in large numbers (between 60-114 specimens in samples KU14-22). Tanyosphaeridium xanthiopyxides continued its predominance from the 306 307 L5 samples (20–81 specimens) forming between 5 and 26% of the total species count. Species that were not present below the K/Pg boundary but were present in 308 309 samples KU14–KU29 include Fibrocysta capitata. Senoniasphaera inornata as well 310 as Cribroperidinium spp. and Thalassiphora pelagica. These species were present in 311 low numbers (1-8 specimens) and not seen in every KU sample. A cribroperidinioid 312 cyst attributed here to Cribroperidinium sp. A of Brinkhuis and Schiøler (1996), has 313 been recorded by Brinkhuis et al. (1998), Slimani et al. (2010) and Vellekoop et al. (2014). Cribroperidinium sp. A of Brinkhuis and Schiøler (1996) is considered a 314 315 transitional form with Carpatella cornuta sensu stricto (Damassa, 1988). It has a 316 short apical horn and spongy appearance but lacks the definite antapical protrusion of C.cornuta s.s. This taxon only occurs at Stevns Klint above the K/Pg boundary in 317 318 the Fiskeler Member. This is consistent with other studies (Brinkhuis et al., 1998; Schiøler et al., 1997; Vellekoop et al., 2014) which have described transitional forms 319 320 such as this occurring in the lowermost Danian, below the first appearance of 321 *C.cornuta* s.s., however, Vellekoop et al. (2015) and Slimani et al. (2010) record its 322 first appearance just below the boundary. The acritarch Raphidodinium sp. cf. armatum was always present in these samples and occurred in significant numbers 323 (23-59 specimens) from samples KU19 to KU23 and KU27-29. Xenicodinium 324 325 reticulatum was observed in low numbers (2-5 specimens) in two of the uppermost

326	samples (KU23 andKU29). In addition, Damassadinium californicum was only
327	observed intermittently and as single to rare occurrences.
328	Peridinioid cysts were found to be more common and diverse than below the
329	boundary, increasing in number up through the succession. There were more
330	peridiniacean specimens recorded in the higher, more productive KU samples
331	(KU24-29). Palaeoperidinium pyrophorum continued to be intermittently present and
332	Cerodinium diebelii was often common (8-47 specimens) with rare Deflandrea spp.
333	and Palaeocystodinium australinum. Manumiella seelandica (formerly, M. druggii)
334	was only observed intermittently and in variable numbers (1-14 specimens).
335	Trithyrodinium evittii was very common, becoming abundant (163 and 214
336	specimens) in the highest samples (KU28 and KU29, respectively), where it
337	accounted for 50-65% of the assemblages. Species diversity (31-41) remained high
338	in the KU samples, similar to the L5 and K8 samples throughout the clay, with an
339	overall increase through the upper samples (see Fig. 7).
340	No dinocysts were observed in samples KU48–KU50 (Cerithium Limestone
341	Member), although bisaccate pollen grains were recorded.

342

343

6. Biostratigraphy

Wilson (1974), using a relatively limited number of samples, was the first to propose
a dinocyst zonation for Denmark. Subsequently, J.M. Hansen and co-workers (J.M.
Hansen, 1977, 1979, Kjellström and Hansen, 1981) and Hultberg (1986), working
across a range of sites in southern Scandinavia (including Stevns Klint), generated a
zonation for the uppermost Maastrichtian and lowermost Danian. More recently,
Schiøler and Wilson (1993) refined Wilson's (1974) dinocyst zonation for the whole
of the Maastrichtian. The problem with many of these zonations is a lack of precise

information on how the sampling of the variable lithostratigraphical units wasaccomplished, especially in Denmark.

In the discussion of the K/Pg boundary at El Kef (Molina et al., 2006) the stratigraphical value of *Danea californica* (= *Damassadinium californicum*) was regarded as important: it was one of the potential criteria for the definition of the GSSP.

Sample KUL/I contained no dinocysts but was collected immediately below
the hardground surface at the top of the Sigerslev Member. This suggests that
hardground formation (which also characterises the upper levels of the Cerithium
Limestone Formation) inhibits the preservation of dinocysts. This has previously
been noted by FitzPatrick (1992) in the Turonian chalk succession of the United
Kingdom.

Palynodinium grallator occurs, often in large numbers (e.g., 110-235 363 specimens) in the upper parts of the Sigerslev Member, aside from sample KUL/I 364 365 adjacent to the hardground and continues in reduced numbers (0-38) throughout the Højerup Member to the very top of the Maastrichtian. In the Højerup Member, P. 366 grallator occurs in reduced numbers while Achomosphaera ramulifera and 367 Impagidinium cristatum are present in reasonable numbers (6-20 and 2-16, 368 respectively) before the latter species disappears at the K/Pg boundary. Records of 369 rare P. grallator (1-5 specimens) within the Fiskeler Member are possibly the result 370 of inclusion of minute clasts of chalk within the samples processed. Vellekoop et al. 371 (2015) record the first appearance of *P.grallator* above the K/Pg boundary in Tunisia 372 and Slimani et al. (2010) describe a similar occurrence from Morocco. This 373 374 occurrence of *P.grallator* was, in both cases, interpreted as the southward migration of a cooler water species in response to cooling during the early Danian. 375

Palynodinium minus occurs throughout the Fiskeler Member, from KU15–KU28, but
in low numbers (1–5 specimens). This species has reduced lateroventral protrusions
similar to that described as *P*. cf. grallator by Machalski et al. (2016) which was
noted from the Maastrichtian of Lechówka, south east Poland. The sporadic
occurrence of *P.grallator* and *P.minus* above the K/Pg boundary is, therefore, not
unusual.

382 *Damassadinium californicum* does occur in the Stevns Klint material (see 383 Table 1), but is very rare. It is present only in the Fiskeler Member, although the 384 environmental changes recorded higher in the succession probably were a limiting 385 control on its reported range.

386 The occurrence of Cribroperidinium sp. A of Brinkhuis and Schiøler (1996), 387 although rare within the Fiskeler Member is guite characteristic, appearing in the sample below *D. californicum* which is only millimetres above the K/Pg boundary as 388 defined by Molina et al. (2006). A number of other taxa (e.g., Fibrocysta capitata, 389 390 *Cribroperidinium* spp.) also occur throughout the Fiskeler Member, which contains a diverse dinocyst assemblage throughout (31–41 species per sample). This result is 391 different to the near-monospecific association described by Hultberg (1985, 1986, 392 1987). 393

Appearing in KU23 is *Xenicodinium reticulatum*, another marker species to have a first appearance within the Fiskeler Member, including *Trithyrodinium evitii*, *Senoniasphaera inornata* and *Manumiella seelandica*. We suggest, therefore, that the upper *X. reticulatum* Sub-zone *of the D. californicum* Zone, of the lowermost Danian can be recognised here.

399

400 6.1 Comparison with biostratigraphic zonations of Scandinavia

In the following paragraphs the results from this study are discussed in relation to
previously published distributions of stratigraphically significant dinocysts from
southern Scandanavia.

404

405 6.1.1 Palynodinium grallator Zone (new definition)

The uppermost part of the upper Maastrichtian is characterised by the P. 406 grallator Zone (Wilson, 1974; Schiøler and Wilson, 1993). This zone, is here defined 407 by the first and last common occurrences of *P. grallator*, (K8/7–KUL-K) respectively 408 and is different to Wilson's (1974) Zone Vb, as the top of this zone was defined by 409 410 the last occurrence of *P.grallator*. The uppermost part of this zone has been divided 411 into the Tanyosphaeridium magdalium (= T. xanthiopyxides) and Thalassiphora pelagica Subzones by Schiøler and Wilson (1993). In previous studies at Stevns 412 Klint, and throughout the Danish region (e.g., Hultberg, 1986; Schiøler and Wilson, 413 1993), T. xanthiopyxides has always been referred to as T. magdalium. Lentin & 414 Williams (1993), however, noted that T. magdalium was almost certainly a junior 415 synonym of T. xanthiopyxides. 416

Kjellström and Hansen (1981) also included the Chiropteridium inornatum-417 418 Palynodinium grallator Concurrent Range Zone in their zonation, with its upper boundary corresponding to the Maastrichtian–Danian (K/Pg boundary). This 419 420 Concurrent Range Zone was later renamed the Senoniasphaera inornata-421 Palynodinium grallator Concurrent Range Zone by Hultberg (1985). The common occurrence of *T.xanthiopyxides*, and the consistent occurrence of *Thalassiphora* 422 pelagica, in this study, through the Højerup Member and into the Fiskeler Member, 423 prevents the recognition of these subzones as defined by Hansen (1977) and 424 Kjellström and Hansen (1981). 425

426	
427	6.1.2 Damassadinium californicum (Danea californica) Zone
428	Originally named after its taxonomic junior synonym Danea mutabilis (Hansen, 1977)
429	this zone begins with the first occurrence of <i>D. californicum</i> (Kjellström and Hansen,
430	1981; Hultberg, 1985). The zone extends over much of the lower Danian and has
431	previously been divided into two subzones (Chiropteridium inornatum and
432	Hafniasphaera cyrptovesiculata Subzones), of which only the lower one (the
433	Senoniasphaera inornata Subzone, formerly the Chiropteridium inornatum Subzone)
434	has been observed at Stevns Klint (e.g., J.M. Hansen, 1977). Hultberg (1985, 1986)
435	did not record D. californicum at Stevns Klint but this is unsurprising as it is relatively
436	rare in our samples (1–3 specimens in some, but not all samples of the Fiskeler
437	Member). The rare but consistent occurrence of S. inornata here, with its first
438	appearance just below that of D. californicum, within the Fiskeler Member, supports
439	this zonation. The Senoniasphaera inornata Subzone was divided into three zonules:
440	Carpatella cornuta, Xenicodinium rugulatum, Xenicodinium reticulatum Zonules, by
441	Hansen (1977) and Kjellström and Hansen (1981), but Hultberg (1986) added a
442	fourth: the Fibrocysta axialis Zonule. These zonules, as defined by Hansen (1977)
443	and Kjellström and Hansen (1981), were not all recognised here in the samples from
444	Stevns Klint.
445	

446 6.1.3 Carpatella cornuta Sub-zone (new definition)

This subzone should contain *C. cornuta* (Kjellström and Hansen, 1981), however, it
is not recognised here. The transitional form, *Cribroperidinium* sp. A of Brinkhuis and
Schiøler (1996) present here, is known to occur in lowermost Danian samples
(Brinkhuis et al. 1998; Slimani et al., 2010; Schiøler et al.,1997; Vellekoop et al.,

451 2015). Xenicodinium reticulatum is not present. Kjellström and Hansen (1981) indicated that Cordosphaeridium inodes longipes was not found at this level and this 452 is consistent with our results. Cribroperidinium sp. A of Brinkhuis and Schiøler (1996) 453 454 is, therefore, interpreted as a marker species for the basal Danian from samples KU14–KU20. 455 456 6.1.4 Xenicodinium reticulatum Sub-zone 457 458 This subzone is only tentatively identified here within the Fiskeler Member as samples from all over southern Scandinavia (as attempted by Kjellström and 459 Hansen, 1981) have not been investigated. Xenicodinium reticulatum was observed 460 461 in samples KU23-KU29 and so is the basis for the identification of this subzone. No

462 dinocysts were observed in samples KU45–KU50, although bisaccate pollen grains
463 were recorded.

464

465 **7. Discussion**

466 7.1 K/Pg boundary sections

K/Pg boundary sections world-wide, fall into three categories; those proximal to the
Chixculub impact site (e.g., Texas, Alabama, Georgia, Gulf of Mexico), distal sites
(e.g., Denmark, Tunisia, Italy, Austria, etc.) and those that are intermediate in
character e.g. Demerara Rise (MacLeod et al., 2007) and New Jersey (Olsson et al.,
1997).

472 Stevns Klint is an important distal K/Pg boundary site, with others located in
473 Southern Spain, Morocco, Tunisia, Egypt, Italy, Austria, Czech Republic, Poland,
474 etc. These sites extend across a wide range of latitudes from 'Tethyan' to 'Boreal'
475 locations and dinocyst studies from these areas have produced varying results in

476	terms of species diversity and abundance (Molina et al., 2006; Donze and Méon,
477	1997; Brinkhuis and Leereveld, 1988; Brinkhuis and Zachariasse,1988; Brinkhuis et
478	al.,1998; M'Hamdi et al. 2015; Vellekoop et al., 2015).
479	Our information on the dinocyst stratigraphy of Stevns Klint can be compared
480	to many of these areas with some reliability, including successions from a wide
481	range of different environments and water depths.
482	
483	7.2 Dinocyst occurrences and biostratigraphic markers
484	In addition to the publications already mentioned, there is extensive literature on
485	K/Pg boundary successions that include dinocyst studies. Many of these studies
486	describe diverse and abundant dinocyst assemblages, characteristic of the
487	uppermost Maastrichtian and lowermost Danian (Wilson,1974; J.M. Hansen, 1979;
488	De Coninck and Smit,1982; Firth,1987, 1993; Hultberg and Malmgren, 1987; Habib
489	et al.,1992; Moshkovitz and Habib, 1993; Schiøler and Wilson, 1993; Olsson et al.,
490	1997; Schiøler et al.,1997; Nøhr-Hansen and Dam,1997; Mohamed and Wagreich,
491	2013; Vellekoop et al., 2016). Others, often the thinner successions, are usually not
492	very productive and yield poor or barren dinocyst assemblages (Gedl, 2004;
493	Mohamed et al., 2013; Beiranvand et al., 2014; Machalski et al., 2016).
494	Conclusions relevant to this study that can be made from the literature:
495	1. <i>P. grallator</i> often, and quite widely, typifies the uppermost Maastrichtian
496	(Wilson, 1974; Schiøler and Wilson, 1993) with C. cornuta, D. californicum and
497	S. inornata frequently characterising the lowermost Danian, (Kjellström and
498	Hansen, 1981) though the exact details of their ranges are dependent on the
499	individual site, the sampling interval used and issues relating to re-working of
500	fine sediment across the boundary.

501	2.	An increase in the abundance of peridinioid species, demonstrated by the G/P
502		and P/G ratios across the K/Pg boundary into the lower Danian, has been
503		recognised previously (Brinkhuis et al., 1998; Prauss, 2009). More specifically,
504		an increase in the abundance of Trithyrodinium evittii is noted by Brinkhuis et
505		al. (1998) and Slimani et al. (2010). The predominance of <i>T.evittii</i> has also
506		been recognised in central and northern Europe (Schiøler et al., 1997) and the
507		north-eastern USA (Firth, 1987, 1993).
508	3.	Vellekoop et al. (2015) describes the K/Pg boundary succession at Elles (75 km
509		south of EI Kef) and shows very similar results to those from the K/Pg
510		stratotype section. Diverse and abundant dinocyst assemblages were
511		recovered, and an increase in the abundance of Cribroperidinium sp. A of
512		Brinkhuis and Schiøler (1996) in the lowermost Danian is noted.
513		
514	7.3	Interpretation of results from Stevns Klint
515	In t	he present study, only four dinocyst taxa are found to be present throughout the
516	pro	ductive samples. These are A. ramulifera, S. ramosus ramosus, S. ramosus
517	gra	cilis and T. xanthiopyxides. These are all cosmopolitan species (Brinkhuis et al.,
5 10	400	(Cluiia

518 1998) and are considered characteristic of normal, marine shelf environments (Sluijs519 et al., 2005).

520 The K8/7 and K8/5 samples of the coccolith-rich chalk both contain *P.* 521 *grallator* (112 and 110 specimens, respectively) and *T. xanthiopyxides* (9 and 4 522 specimens, respectively) and, therefore, confirm previous research on the uppermost 523 Maastrichtian (Wilson, 1974; Hultberg, 1986; Schiøler and Wilson, 1993). The 524 predominance of *P. grallator* and, to a lesser extent, species of *Achomosphaera* (29 525 and 32 specimens, respectively) and subspecies of *Spiniferites ramosus* (51 and 49

526 specimens, respectively), is consistent with other findings at Stevns Klint (e.g., J.M. 527 Hansen, 1977; Hultberg, 1986; Brinkhuis et al., 1998). Palaeoperidinium pyrophorum is the only peridinioid cyst encountered in the Maastrichtian samples. Its highest 528 529 abundance (15 specimens) occurs in sample K8/7. 530 Of the other Kulstirenden samples within the coccolith-rich chalk, KUL/I was barren but KUL/K yielded assemblages very similar to K8/7 and K/8/5 samples, with 531 a predominance of *P.grallator*, *I. cristatum* and the associated *Achomosphaera* 532 /Spiniferites complex of species. Consequently, the G/P species ratio for the 533 Maastrichtian is also at its highest here (based on one peridinioid species), although 534 it still remains low (K8/7 G/P ratio = 25). Samples with a G/P species ratio of 18 are 535 536 considered by Harland (1973) as being indicative of a marine environment. Hultberg 537 (1986) is the only other author to have calculated the G/P ratio at Stevns Klint showing values of 1–300, using the number of specimens. Our findings here are 538 539 consistent with Hultberg (1986) for the Maastrichtian, which also shows gonyaulacacean-dominated assemblages for the upper Maastrichtian. The G/P^{species} 540 ratio calculated here varies between 0 and 29 and G/P^{specimens} ratio varies from 0 to 541 304, both showing a decreasing trend through the Maastrichtian and into the Danian. 542 Hultberg's (1986) G/P ratio varied between 75 and 300 in the Maastrichtian chalks, 543 decreasing to <50 at the base of the Fiskeler Member. Palynodinium grallator and H. 544 545 tubiferum tubiferum are both considered to be indicative of high-latitude climates (Brinkhuis et al., 1998), with the latter "Hystrichosphaeridium-type" being noted for its 546 ability to tolerate stressed environments (Schiøler et al., 1997). Their combined 547 548 presence and the high abundances of *P. grallator* are, therefore, indicative of relatively cool climatic conditions at this time. 549

550 Within the Højerup (L5) samples of the 'Grey Chalk', the Achomosphaera-Spiniferites complex cysts dominate the flora, replacing *P. grallator* as the major 551 taxon, although T. xanthiopyxides becomes increasingly abundant (14-37 552 553 specimens) and is one of the most common species in the three uppermost samples. Spiniferites ramosus ramosus and S.ramosus gracilis are also very common in these 554 samples. While P. grallator remains present, its abundance is relatively low 555 compared to the K8 samples. In contrast, Impagidinium spp. becomes a major 556 component of the assemblage, their highest abundances occurring in the lowermost 557 558 samples.

559 The biostratigraphy shows that *P. grallator* was observed rarely at the base of 560 the Fiskeler Member. This agrees with Brinkhuis et al. (1998) who found this species in the bottom layer of the Fiskeler Member. Moreover, Hultberg (1986) had 561 previously noted P. grallator, in parts of Stevns Klint, in Layer 3 (Christensen et al., 562 1973) of the Fiskeler Member; while in other parts of the peninsula its range was 563 564 found to extend only to Layer 1. Hultberg drew two main conclusions from these findings: (i) that the Fiskeler Member is Maastrichtian in age, and (ii) had been 565 deposited diachronously over Denmark. This was in marked contrast to other fossil 566 and geochemical evidence, including the large negative shift in the δ^{13} C record (e.g., 567 Hart et al., 2005, fig. 10) record from Stevns Klint. The top of the Højerup Member 568 and therefore, the end of the Maastrichtian in higher latitudes can be defined by the 569 570 last common occurrence of *P. grallator*. Our data clearly support a Danian age for the Fiskeler Member at this location. The 'diachronism' reported by Rasmussen et al. 571 (2005) for the planktic foraminiferal zonation is considered an artefact of the variation 572 in thickness of the Fiskeler Member and the position of the facies change to the 573 Cerithium Limestone Member. The iridium spike that is recorded within K/Pg 574

575 boundary sediments from a variety of locations (Alvarez et al., 1980), however, 576 precludes diachronous deposition as does the global correlation of the δ^{13} C data 577 (Hart et al., 2005; Hart et al., 2014).

578 The D. californicum Zone is representative of the Danian. In contrast to data presented by Kjellström and Hansen (1981), however, the expanded section of the 579 Fiskeler Member studied here documents only an intermittent presence of D. 580 californicum and in reduced numbers. Hultberg (1986) and Brinkhuis et al. (1998) did 581 not record *D. californicum* at Stevns Klint, confirming its rareness. This suggests 582 that, although the *D. californicum* Zone has been applied relatively successfully to 583 584 the rest of southern Scandinavia, this zonation is difficult to apply at Stevns Klint, or 585 at least a part of it. The rare occurrence of S.inornata, a species that is supposed to 586 be coincident with *D. californicum* throughout the lower parts of the Danian, supports this zonation. With the mass extinction of calcareous nannofossils and almost all 587 planktic foraminifera at the K/Pg boundary, there is a dramatic loss of sediment 588 589 supply leaving only a thin clay-rich succession and an apparent concentration of organic-walled microfossils. Even our small samples, therefore, represent a 590 significant interval of time as indicated by the precessional cyclicity in the δ^{13} C data 591 592 (Hart et al., 2005; Leighton et al., 2017). If this is true in the expanded succession seen at Kulstirenden then it is particularly so in the thinner successions (e.g., at 593 Højerup). If greater condensation of sediments has occurred in these thinner 594 595 sections, then each sample represents a longer period of time. Consequently, species that were only intermittently present might appear to be present throughout 596 the whole time represented by the sediment. For this location, therefore, it is 597 proposed that the lowermost Danian is defined by the appearance of 598 Cribroperidinium sp. A of Brinkhuis and Schiøler (1996) together with the 599

600 appearance of S. inornata and D.californicum because C.cornuta s.s. is not present. In the middle of the Fiskeler Member (sample KU23), X. reticulatum appears and 601 602 characterises an X. reticulatum Sub-zone. With all dinocyst taxa disappearing in the 603 Cerithium Limestone Member and the overlying Korsnæb Member the vertical extension of this sub-zone is not currently known. 604 Another feature of the KU (14–50) samples of the Fiskeler Member is the 605 predominance of *T. xanthiopyxides*, where it commonly represents approximately 606 one-third of the assemblage. Previous studies have shown this species to be present 607 608 in all stratigraphic levels at Stevns Klint, but no one has commented on its 609 predominance within this stratigraphic level. Indeed, both H.J. Hansen et al. (1986) 610 and Brinkhuis et al. (1998) show T. evittii to be the dominant species through layers 611 2 and 3 of the Fiskeler Member. Trithyrodinium evittii is also found to be common to very common (19–48 specimens in samples KU18–27) within the Fiskeler Member in 612 our study and becomes abundant in the topmost samples (KU28 and KU29, 613 614 recording 163-214 specimens, respectively). Another peridinioid, Manumiella seelandica (as M.druggii) has previously been reported to be abundant in these 615 sediments at the base of the Danian (Hultberg, 1986) but these data are not 616 confirmed here. In the current study occasional specimens of *M. seelandica* are 617 618 found at the base of the Fiskeler Member, becoming more common (5-14 619 specimens) in the topmost samples (KU26–KU29). Instead of abundant M. seelandica, large numbers of *T.evittii* are recorded here and this confirms the results 620 of Brinkhuis et al. (1998). 621 The lack of dinocysts, and the presence of bisaccate pollen, which have a 622

hinterland provenance (Schiøler et al., 1997), could be due to variable preservation

and/or complex diagenesis of the Cerithium Limestone Member (Rasmussen et al.2005).

626

627 7.4 Palaeoenvironmental interpretation

The Gonyaulacoid/Peridinioid (G/P) and Protoperidinioid /Gonyaulacoid (P/G) ratios were calculated here based on the data collected; the former after Harland (1973), and the latter follows Versteegh (1994). Note that these methods produce different results to those obtained by Hultberg (1986,1987) for samples from Stevns Klint but the overall decreasing trend in the G/P^{species} and G/P ^{specimens} ratios is similar . The method of calculation for the G/P ratio used in Hultberg (1986, 1987) is not clear but probably involved using the number of specimens.

The G/P ratio, based on specimen percentage, was calculated by Prauss (2009) for the K/Pg boundary at Brazos River, Texas. An overall increase in peridinioid cysts was recorded across the K/Pg boundary. This is interpreted as reflecting an increase in palaeoproductivity through the succession, into the Danian. Brinkhuis et al. (1998) found that their P/G ratios did not correlate with the drop in δ^{13} C values across the K/Pg boundary at El Kef and interpreted this as reflecting changes in temperature as well as productivity.

Both G/P^{species} and G/P ^{specimens} ratios calculated here display an overall decreasing trend from the uppermost Maastrichtian into the Danian, while the P/G ratio increases. The interpretation here is of a marine environment with increasing nutrients and productivity.

Small numbers of *Impagidinium cristatum* (10 specimens) occur in the
Sigerslev Member (K8/7) with an increased abundance occurring in the
stratigraphically younger K8/5 (20 specimens). *I. cristatum* is considered to be

indicative of more oceanic rather than neritic conditions (Schiøler et al., 1997), and so its presence indicates that there was still an open marine influence to the environment in the Danish Basin. This is recognised by Sheldon et al. (2010) as an anti-clockwise circulatory system in the Danish Basin with possible warmer water influences coming from the southeast. The decrease in abundance of *I. cristatum* and other *Impagidinium* spp. implies a shallowing-upward sequence.

Bromley (1979) and Hart et al. (2004, 2005) suggest that the succession of the Højerup Member represented a shallower environment compared to the white coccolith chalk below (Sigerslev Member). The stable isotope data derived from benthic foraminifera (Schmitz et al., 1992; Hart et al., 2005) provide evidence that, although shallower, the environment was still open marine.

This shallowing upwards is confirmed by the occurrence of *Senoniaspharea inornata* in the Fiskeler Member (KU14-29). This species belongs to the *Areoligera* group of Gv cysts (Fensome et al., 2008) which has been interpreted as reflecting high energy, shallower environments (Sluijs et al., 2005). The occurrence of this species at Stevns Klint, reflects the shallower conditions both above, and near to the Rynkøbing-Fyn High in comparison to the more basinal area of northern Jutland (Jylland).

Brinkhuis et al. (1998, p.77) indicate that the differences between their results and those of Hultberg (1986) were the result of the processing techniques employed and then conclude (Brinkhuis et al. 1998, p.80) that there were warm, stable conditions in the latest Maastrichtian, and cooler but fluctuating conditions in the earliest Danian, with water masses controlling the migration and distribution of taxa. In the past, *T. evittii* has been used as a warmer-water, low latitude indicator species and as a peridinioid, it also indicates increased primary productivity (Brinkhuis et al.,

1998, fig.6). The abundance of *T.evittii* in the lower Danian at Stevns Klint agrees
with observations made by Brinkhuis et al. (1998) and others (e.g., Aydin, 2013;
Pross and Brinkhuis, 2005).

677

678 8 Conclusions

679 The evidence obtained from this investigation of the dinocysts provides data that helps to document the changes across the Maastrichtian/Danian (K/Pg) boundary. 680 Our results do not replicate those of earlier studies, although, this may be due to the 681 682 limited number of samples analysed in previous investigations, the sample interval, sampling from thinner sections of the Fiskeler Member or the processing techniques 683 684 adopted (e.g. Hultberg, 1986). Also previous authors have rarely identified their 685 precise sample locations on Stevns Klint, so lateral variations, as indicated by the variable lithology may account for some of these differences. 686

The top of the Højerup Member and, therefore, the top of the Maastrichtian is 687 688 characterised by the last common appearance of *P. grallator*. Within the Fiskeler Member, the intermittent presence of *D. californicum* precludes its use as a primary 689 biostratigraphical indicator although, when used together with other 690 691 biostratigraphically useful species such as S. inornata, X. reticulatum and T. evittii, the lowermost Danian can be characterised. Conditions throughout the latest 692 693 Maastrichtian and the early Danian, as indicated by the G/P ratio, were changing and 694 becoming dominated by peridinioid dinocysts at the base of the Danian, reflecting an increase in primary productivity (Sluijs et al. 2005). 695

Although our sampling was at a much higher resolution than earlier
 investigations and from an expanded succession of Fiskeler Member, there is still
 the potential for further dinocyst work at Stevns Klint and additional refinement of the

699	biostratigraphy, especially if coupled with data from successions in North Jutland
700	(Jylland) and off-shore.
701	
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710	
711	References
712	Abildgaard, S., 1759. Beskrivelse over Stevns Klint og dens naturlige
713	Mærkværdigheder. Svare, København, 50pp.
714	Alvarez, L.W., Alvarez, W., Asaro, F., Michel, H.V., 1980. Extraterrestrial cause for
715	the Cretaceous-Tertiary extinction. Science 208, 1095–1108.
716	Aydin, T., 2013. Dinoflagellate cyst biostratigraphy, palynofacies and
717	palaeoenvironmental analysis of the Maastrichtian and basal Danian, Brazos
718	River, Texas. Unpublished MSc Thesis, Texas A&M University, College Station,
719	Texas, 121pp.
720	Batten, D.J., 1999. Small palynomorphs. In: Jones, T.P., Rowes, N.P. (eds.), Fossil
721	plants and spores: modern techniques, Geological Society, London, pp.15-
722	19.
723	Beiranvand, B., Zaghbib-Turki, D., Ghasemi-Nejad, E., 2014. Integrated
724	biostratigraphy based on planktic foraminifera and dinoflagellates across the
725	Cretaceous/Paleogene (K/Pg) transition at the Izeh section (SW Iran). Comptes
726	Rendus Palevol 13, 235–258.

727	Bernecker, M., Weidlich, O. 1990. The Danian (Paleocene) Coral Limestone of
728	Fakse, Denmark: A model for ancient aphotic, azooxanthellate coral mounds.
729	Facies 22, 103–138.
730	Bernecker, M., Weidlich, O. 2005. Azooxanthellate corals in the Late Maastrichtian -
731	Early Paleocene of the Danish Basin: Bryozoan and coral mounds in a boreal
732	shelf setting. In: Freiwald, A., Roberts, J.M. (eds.), Cold-water corals and
733	ecosystems, Springer Verlag, Heidelberg, 3–25.
734	Brinkhuis, H., Leereveld, H. 1988. Dinoflagellate cysts from the Cretaceous/Tertiary
735	boundary sequence of El Kef, Northwest Tunisia. Revue of Paleobotany and
736	Palynology 56, 5–19.
737	Brinkhuis, H., Schiøler, P., 1996. Palynology of the Geulhemmerberg
738	Cretaceous/Tertiary boundary section (Limburg, SE Netherlands). Geologie en
739	Mijnbouw 75, 193-213
740	Brinkhuis, H., Zachariasse, W.J. 1988. Dinoflagellate cysts, sea level changes and
741	planktonic foraminifers across the Cretaceous/Tertiary boundary at El Haria,
742	northwest Tunisia. Marine Micropaleontology 13, 153–191.
743	Brinkhuis, H., Bujak, J.P., Smit, J., Versteegh, G.J.M., Visscher, H., 1998.
744	Dinoflagellate-based sea surface temperature reconstructions across the
745	Cretaceous-Tertiary boundary. Palaeogeography, Palaeoclimatology,
746	Palaeoecology 141, 67–83.
747	Bromley, R.G., 1979. Chalk and bryozoan limestone: facies, sediments and
748	depositional environments. In: Birkelund, T., Bromley, R.G. (eds.), Cretaceous-
749	Tertiary Boundary Events, I. The Maastrichtian and Danian of Denmark, pp16-
750	32. University of Copenhagen.
751	Christensen, L. E., Fregerslev, S., Simonsen, A., Thiede, J., 1973. Sedimentology
752	and depositional environments of Lower Danian Fish Clay from Stevns Klint,
753	Denmark. Bulletin of the Geological Society of Denmark 22, 193–212.
754	Damassa, S.P. 1988. <i>Carpatella cornuta</i> Grigorovich 1969 (Dinophyceae) – a
755	member of the Aptiana-Venticosum complex. Palynology 12, 167–177.
756	Damholt, T., Surlyk, F., 2012. Nomination of Stevns Klint for inclusion in the World
757	Heritage List. Østsjællands Museum, Store Heddinge, Denmark, 160 pp.
758	De Coninck, J., Smit, J. 1982. Marine organic-walled microfossils at the Cretaceous-
759	Tertiary boundary in the Barranco del Gredero. Geologie en Mijnbouw 61, 173-
760	178.

761	Desor, É., 1847. Sur le terrain danien, nouvel étage de la craie. Bulletin de la Société
762	géologique de France 4, 179–182.
763	Donze, P., Méon, H. 1997. Les extinctions à la limite Crétacé-Tertiare: disparité
764	entre le domain marin et le domaine contental révelé par la coupe
765	stratigraphique du Kef (nord-ouest de la Tunisie). Comptes Rendus de
766	l'Academie des Sciences, Paris 325, 961–966.
767	Fensome, R.A., MacRae, R.A. and Williams, G.L., 2008. DINOFLAJ2, Version 1.
768	American Association of Stratigraphic Palynologists, Data Series no. 1.
769	Available at: http://dinoflaj.smu.ca/wiki/Main_Page. Accessed on 08/06/2017
770	Firth, J.V. 1987. Dinoflagellate biostratigraphy of the Maastrichtian to Danian interval
771	in the U.S. Geological Survey Albany core, Georgia, U.S.A. Palynology 11,
772	199–216.
773	Firth, J.V. 1993. Dinoflagellate assemblages and sea-level fluctuations in the
774	Maastrichtian of southwest Georgia. Review of Palaeobotany and Palynology
775	79, 179–204.
776	FitzPatrick, M.E.J., 1992.Turonian dinoflagellate cyst assemblages from Southern
777	England. Unpublished PhD Thesis, University of Plymouth, 373pp.
778	Gedl, P. 2004. Dinoflagellate cyst record of the deep-sea Cretaceous-Tertiary
779	boundary at Uzgruň, Carpathian Mountains, Czech Republic. In: Beaudoin,
780	A.B., Head, M.J. (eds.), The palynology and micropalaeontology of boundaries,
781	Geological Society, London, Special Publications 230, 257–273.
782	Graversen, P. 2001. Den geologiske udforskning af Fakse Kalkbrud frå midten af
783	1700-tallet til nu. Geologisk Tidsskrift 2, 1–40.
784	Habib, D., Moshkovitz, S., Kramer, C. 1992. Dinoflagellate and calcareous
785	nannofossil response to sea-level change in Cretaceous-Tertiary boundary
786	sections. Geology 20, 165–168.
787	Hansen, H.J., Gwozdz, R., Hansen, J.M., Bromley, R.G., Rasmussen, K.L., 1986.
788	The diachronous K/T plankton extinction in the Danish Basin, pp. 381–384. In:
789	Walliser. O. (Ed.), Global Bioevents, [Lecture Notes on Earth Sciences, 8],
790	Springer, Berlin/Heidelberg.
791	Hansen, J.M. 1977. Dinoflagellate stratigraphy and echinoid distribution in Upper
792	Maastrichtian and Danian deposits from Denmark. Bulletin of the Geological
793	Society of Denmark 26, 1–26.

794	Hansen, J.M., 1979. Dinoflagellate zonation around the boundary. In: Birkelund, T.,
795	Bromley, R. G. (eds.), Cretaceous-Tertiary Boundary Events Symposium: I.
796	The Maastrichtian and Danian of Denmark, pp. 136–141, University of
797	Copenhagen, Copenhagen.
798	Harland, R., 1973. Dinoflagellate cysts and acritarchs from the Bearpaw Formation
799	(Upper Campanian) of southern Alberta, Canada. Palaeontology 16, 665–706.
800	Hart, M.B., Feist, S.E., Håkansson, E., Heinberg, C., Price, G.D., Leng, M.J.,
801	Watkinson, M. P., 2005. The Cretaceous-Palaeogene boundary succession at
802	Stevns Klint, Denmark: foraminifers and stable isotope stratigraphy.
803	Palaeogeography, Palaeoclimatology, Palaeoecology 224, 6–26.
804	Hart, M.B., Feist, S.E., Price, G.D., Leng, M.J., 2004. Reappraisal of the K-T
805	boundary succession at Stevns Klint, Denmark. Journal of the Geological
806	Society, London 161, 885–892.
807	Hart, M.B., Searle, S., Feist, S.E., Leighton, A.D., Price, G.D., Smart, C.W.,
808	Twitchett, R.J., 2011. The distribution of benthic foraminifera across the
809	Cretaceous-Paleogene boundary in Texas (Brazos River) and Denmark
810	(Stevns Klint). In: Keller, G., Adatte, T. (eds.), The end-Cretaceous mass
811	extinction and the Chicxulub Impact in Texas, Society for Sedimentary Geology
812	(SEPM), Special Publication 100, 179–196.
813	Hart, M.B., Leighton, A.D., Smart, C.W., Pettit, L.R., Medina-Sanchez, A.N., Harries,
814	P.J., Cardenas, A.L. Hall-Spencer, J.M. and Prol-Ledesma, R.M. 2014. Ocean
815	acidification in modern seas and its recognition in the geological record: the
816	Cretaceous/Paleogene boundary in Texas and Alabama. Transactions of the
817	Gulf Coast Association of Geological Societies 64, 193-213.
818	Hultberg, S.U., 1985. Dinoflagellate studies of the Upper Maastrichtian and Danian in
819	southern Scandinavia, Unpublished PhD Thesis, University of Stockholm,
820	Stockholm, 189pp.
821	Hultberg, S.U., 1986. Danian dinoflagellate zonation, the C-T boundary and the
822	stratigraphical position of the Fish Clay in southern Scandinavia. Journal of
823	Micropalaeontology 5, 37–47.
824	Hultberg, S.U., 1987. Palynological evidence for a diachronous low-salinity event in
825	the C-T boundary clay at Stevns Klint, Denmark. Journal of Micropalaeontology
826	6, 35–40.

827	Hultberg, S.U., Malmgren, B.A. 1987. Quantitative biostratigraphy based on Late
828	Maastrichtian dinoflagellates and planktonic foraminifera from South
829	Scandinavia. Cretaceous Research 8, 211–228.
830	Kjellström, G.J., Hansen, J.M., 1981. Dinoflagellate biostratigraphy of the
831	Cretaceous-Tertiary boundary in southern Scandinavia. Geologiska
832	Föreningens i Stockholm Förhandlingar 103, 271–278.
833	Leighton, A.D., Hart, M.B., Smart, C.W., Leng, M.J., Hampton, M., 2017. Timing
834	recovery after the Cretaceous/Paleogene boundary: Evidence from the Brazos
835	River, Texas, USA. Journal of Foraminiferal research, 47,(3), 229-238.
836	Machalski, M., Vellekoop, J., Dubicka, Z., Peryt, D., Harasimiuk, M., 2016. Late
837	Maastrichtian cephalopods, dinoflagellate cysts and foraminifera from the
838	Cretaceous-Paleogene succession at Lechówa, southeast Poland:
839	Stratigraphic and environmental implications. Cretaceous Research 57, 208–
840	227.
841	MacLeod, K.G., Whitney, D.L., Huber, B.T., Koeberl, C. 2007. Impact and extinction
842	in remarkably complete Cretaceous-Tertiary boundary sections from Demerara
843	rise, tropical western North Atlantic. Geological Society of America, Bulletin
844	119, 101–115.
845	M'Hamdi, A., Slimani, H., Louwye, S., Soussi, M., Ismail-Lattrache, K.B., Ben Ali, W.,
846	2015. Dinoflagellate cysts and palynofacies from Mastrichtian-Danian transition
847	at the stratotype EI kef (Tunisia). Comptes Rendus Palevol, 14, 167-180.
848	Milàn, J. 2010. Coprolites from the Danian Limestone (Lower Paleocene) of Faxe
849	Quarry, Denmark. In: Milàn, J.,Lucas, S.G., Lockley, M.G., Spielmann, J.A.
850	(eds.), Crocodyle tracks and traces, New Mexico Museum of Natural History
851	and Science, Bulletin 51, 215–218.
852	Mohamed, O., Wagreich, M. 2013. Organic-walled dinoflagellates cyst
853	biostratigraphy of the Well Höflein 6 in the Cretaceous-Paleogene
854	Rhenodanubian Flysch Zone (Vienna Basin, Austria). Geologica Carpathica
855	64(3), 209–230.
856	Mohamed, O., Piller, W.E., Egger, H. 2013. Dinoflagellate cysts and palynofacies
857	across the Cretaceous/Paleogene boundary at the Neritic Waidach section
858	(Eastern Alps, Austria). Review of Palaeobotany and Palynology 190, 85–103.
859	Molina, E., Alegret, L., Arenillas, I., Arz, J.A., Gallala, N., Hardenbol, J., von Salis,
860	K., Steurbaut, E., Vandenberghe, N., Zaghbib-Turki, D., 2006. The Global

861	Boundary Stratotype Section and Point for the base of the Danian Stage
862	(Paleocene, Palaeogene, "Tertiary", Cenozoic) at El Kef, Tunisia – Original
863	definition and revision. Episodes 29, 263–273.
864	Morgenroth, P.,1968. Zue Kenntnis der Dinoflagellaten und Hystrichosphaeridien des
865	Danien. Geologisches Jahrbuch 86, 533–578.
866	Moshkovitz, S., Habib, D. 1993. Calcareous nannofossil and dinoflagellates
867	stratigraphy of the Cretaceous-Tertiary boundary, Alabama and Georgia.
868	Micropaleontology 39, 167–191.
869	Nøhr-Hansen, H., Dam, G. 1997. Palynology and sedimentology across a new
870	marine Cretaceous-Tertiary boundary section on Nuussuaq, West Greenland.
871	Geology 25(9), 851–854.
872	Noe-Nygaard, A.1975. Erratics from the Danish Maastrichtian and Danian
873	limestones. Bulletin of the Geological Society of Denmark 24, 75–81.
874	Olsson, R.K., Miller, K.G., Browning, J.V., Habib, D., Sugarman, P.J. 1997. Ejecta
875	layer at the Cretaceous-Tertiary boundary, Bass River, New Jersey (Ocean
876	Drilling Program Leg 174AX). Geology 25(8), 759–762.
877	Prauss, M.L., 2009.The K/Pg boundary at Brazos-River, Texas, USA – An approach
878	by marine palynology. Palaeogeography, Palaeoclimatology, Palaeoecology
879	283, 195–215.
880	Pross, J., Brinkhuis, H., 2005. Organic-walled dinoflagellate cysts as
881	palaeoenvironmental indicators in the Paleogene; a synopsis of concepts.
882	Paläontologische Zeitschrift 79/1, 53–59
883	Rasmussen, J.A., Heinberg, A., Håkansson, E., 2005. Planktic foraminifera from the
884	lowermost type Danian of Stevns Klint, Denmark. Bulletin of the Geological
885	Society of Denmark 52, 113–131.
886	Rosenkrantz, A.,1924. Nye lagttagelser over Cerithiumkalken i Stevns Klint med
887	Bemærtkninger om Grænsen mellem Kridt og Tertiær. Meddelelser fra Dansk
888	Geologisk Forening 6, 28–31.
889	Sheldon, E., Ineson, J., Bown, P., 2010. Late Maastrichtian warming in the Boreal
890	realm: Calcareous nannofossil evidence from Denmark. Palaeogeography,
891	Palaeoclimatology, Palaeoecology 295 , 55–75.
892	Schiøler, P., Brinkhuis, H., Roncaglia, L., Wilson, G. J., 1997. Dinoflagellate
893	biostratigraphy and sequence stratigraphy of the Type Maastrichtian (Upper

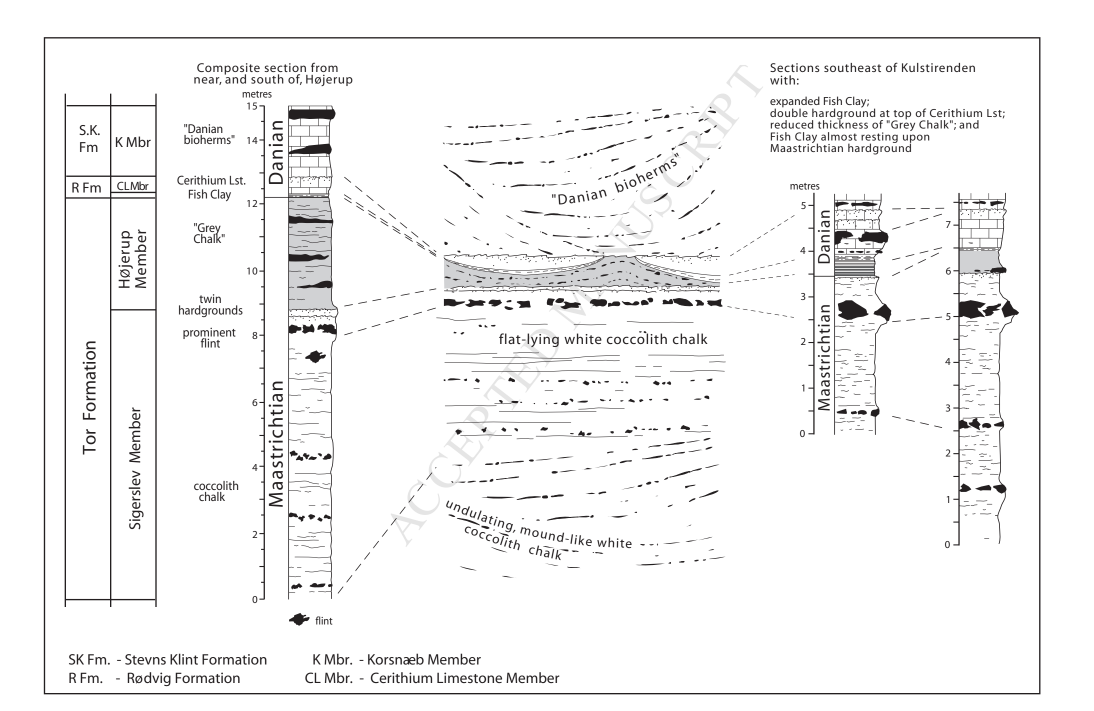
894	Cretaceous), ENCI Quarry, The Netherlands. Marine Micropaleontology 31,
895	65–95.
896	Schiøler, P., Wilson, G.J., 1993. Maastrichtian dinoflagellate zonation in the Dan
897	Field, Danish North Sea. Review of Palaeobotany and Palynology 78, 321–351.
898	Schmitz, B., Keller, G., Stenvall, O., 1992. Stable isotope and foraminiferal changes
899	across the Cretaceous-Tertiary boundary at Stevns Klint, Denmark: arguments
900	for long-term oceanic instability before and after bolide-impact event.
901	Palaeogeography, Palaeoclimatology, Palaeoecology 96, 233–260.
902	Slimani, H., Louwye, S., Toufiq, A. 2010. Dinoflagellate cysts from the Cretaceous-
903	Paleogene boundary at Ouled Haddou, southeastern Rif, Morocco:
904	biostratigraphy, palaeoenvironments and paleobiogeography. Palynology 34,
905	90–124.
906	Sluijs, A. Pross, J., Brinkhuis, H., 2005. From greenhouse to icehouse; organic-
907	walled dinoflagellate cysts as palaeoenvironmental indicators in the Paleogene.
908	Earth-Science Reviews 68, 281–315.
909	Surlyk, F.,1979. Guide to Stevns Klint. In: Birkelund, T., Bromley, R. G. (Eds.),
910	Cretaceous-Tertiary Boundary Events Symposium: I. The Maastrichtian and
911	Danian of Denmark,pp. 164-170. University of Copenhagen, Copenhagen.
912	Surlyk, F., Håkansson, E. 1999. Maastrichtian and Danian strata in the southeastern
913	part of the Danish Basin. In: Pedersen, G.K., Clemmesen, L. (eds.), Field Trip
914	Guidebook, 19 th Regional European Meeting of Sedimentology, 29–58.
915	Surlyk , F., Damholt, T., Bjerager, M., 2006. Stevns Klint, Denmark: Uppermost
916	Maastrichtian Chalk, Cretaceous-Tertiary boundary and lower Danian bryozoan
917	mound complex. Bulletin of the Geological Society of Denmark 54, 1-46.
918	Vellekoop, J., Sluijs, A., Schouten, S., Weijers, J.W.H., Sinninghe Damsté, J.S.,
919	Brinkhuis, H., 2014. Rapid short-term cooling following the Chicxulub impact at
920	the Cretaceous-Paleogene boundary. Proceedings of the National Academy of
921	Sciences, 111(21), 7537-7541
922	Vellekoop, J., Smit, J., Bas van de Schootbrugge, Weijers, J.W.H., Galeotti, S.,
923	Sinninghe Damsté, J.S., Brinkhuis, H., 2015. Palynological evidence for the
924	prolonged cooling along the Tunisian continental shelf following the K-Pg
925	boundary impact. Palaeogeography, Palaeoclimatology, Palaeoecology 426,
926	216–228.

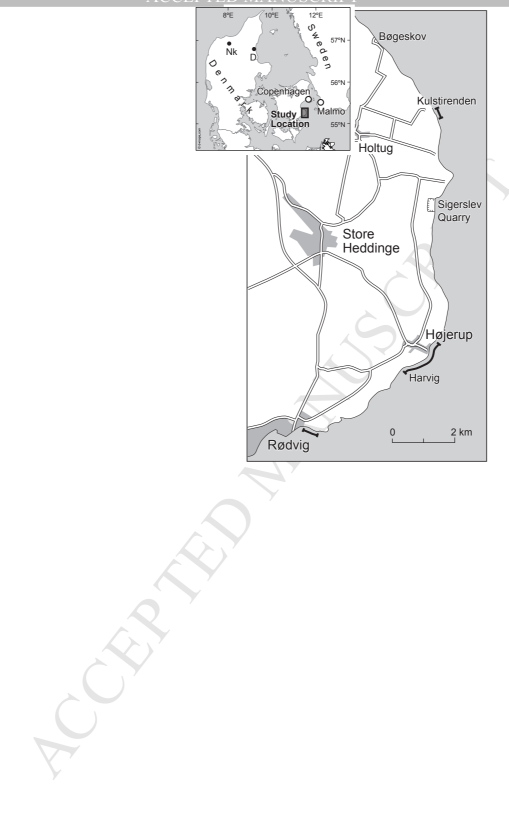
927	Vellekoop., J., Esmeray-Senlet, S., Miller, K.G., Browning, J.V., Sluijs, A., van de
928	Schootbrugge, B., Sinninghe Damsté, J.S., Brinkhuis, H., 2016 Evidence for
929	Cretaceous-Paleogene boundary bolide "impact winter" conditions from New
930	Jersey, USA. Geology 44 (8), 619–622
931	Versteegh, G.J.M., 1994. Recognition of cyclic and non-cyclic environmental
932	changes in the Mediterranean Pliocene: a palynological approach. Marine
933	Micropaleontology 23, 417–183.
934	Willumsen, M. 1995. Early lithification in Danian azooxanthellate scleractinian
935	lithotherm, Faxe Quarry, Denmark. Beiträge zur Paläontologie 20, 123–131.
936	Wilson, G. J., 1974. Upper Campanian and Maastrichtian Dinoflagellate cysts from
937	the Maastricht Region and Denmark, Unpublished PhD Thesis, University of
938	Nottingham, 601pp.
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942	Figures, Table and Plate Captions
943	
944	Fig. 1. Locality map of Stevns Peninsula, eastern Denmark and its location in
945	Southern Scandinavia.
946	
947	Fig. 2. Revised lithostratigraphy of the succession exposed on Stevns Peninsula,
948	following Hart et al. (2005) and Surlyk et al. (2006).
949	
950	Fig. 3. Photograph of the beach section north of Højerup church, showing the
951	horizontal chalk in the lower part of the cliff, the prominent flint line (above which is
952	the incipient hardground) and the Fiskeler Member deposited in troughs picked out
953	by the dune-like structures in the Højerup Member.
954	
955	Fig. 4. (a) Photograph of the Fiskeler Member at Kulstirenden prior to sampling. The
956	distinctive Fe-stained horizon is just below the scale card, at the base of the Fiskeler
957	Member; (b) the marked thinning of the Fiskeler Member to the north of the sampling
958	location.
959	
960	Fig. 5. Composite log of the Cretaceous/Palaeogene boundary succession.

961 Fig. 6. A) Palynodinium grallator, B) Thalassiphora pelagica; C) Palaeoperidinium 962 963 pyrophorum; D) Hystrichosphaeridium tubiferum; E) Spiniferites ramosus reticulatus; F) Spiniferites ramosus; G) Impagidinium cristatum; H) Tanyosphaeridium 964 965 xanthiopyxides; I) Damassadinium californicum; J) Cribroperidinium sp. A of Brinkhuis and Schiøler (1996); K) Xenicodinium reticulatum; L) Manumiella 966 967 seelandica; M) Cerodinium diebelii; N) Trithyrodinium evitti. Scale bar 10µm in all 968 images. 969 Fig. 7. Gonyaulacacean/Peridiniacean (G/P) ratio calculated using species and 970 971 specimen numbers. Peridiniacean/Gonyaulacacean (P/G) ratio ((Versteegh, 1994). 972 Total number of species counted in each sample. 973 974 **Table 1**. Occurrences of selected taxa. \circ < 5 specimens; • >5 specimens. 975 CLM - Cerithium Limestone Member; SM - Sigerslev Member 976 977 Appendix 1. Raw data of dinocyst counts. * marks taxa which require further 978 taxonomic investigation.

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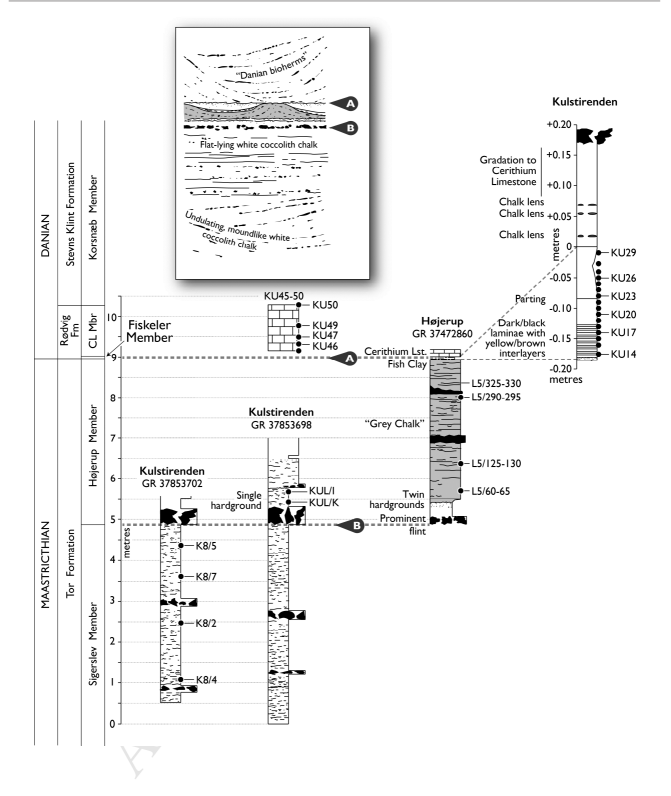
Lithostratigraphy		Sample number	Achomosphaera spp.	Hystrichosphaeridium spp.	Impagidinium spp.	Oligosphaeridium spp.	Palaeoperidinium pyrophorum	Palynodinium grallator	Spiniferites spp.	Tanyosphaeridium xanthiopyxides	Thalassiphora pelagica	Cribroperidinium sp. A	Cerodinium diebelii	Manumiella seelandica	Senoniasphaera inornata	Damassadinium californicum	Deflandrea spp.	Membranilarnacia tenella	Palynodinium minus	Trithyrodinium evittii	Palaeocystodinium australinum	Xenicodinium reticulatuum	
		CLM	KU45-50										Bar	ren									
			KU29	•	•		•	•		•	•	•		•	•	•		٠	•	•	•	•	•
	Rødvig Formation		KU28	•	•	0	•	•		•	•	•		•	•			•	•	•	•	•	
		Fiskeler Member	KU27	•	•	0	•	•		•	•	•		•	•	•		•	•	•	•	•	•
			KU26	•	•		•	•		•	•	•	0	•	•	•	0	•	•	•	•	•	•
			KU25	•	•		•	•		•	•	•	0	•	•	•	0		•	•	•	•	•
DANIAN			KU24	•	•	0	•	•		٠	•	•	0	•	•	•		٠	•	•	•	•	•
			KU23	•	•	0	•	•		•	•	•		•	•	•	0	•	•	•	•		•
			KU22	•	•		•			•	•	•		•	•	•			•	•	•	•	
			KU21	•	•		•	•		•	•	•		•	•	•			•	•	•	•	
			KU20	•	•	0	•	•		•	•	•	0	•	•	•	0	•	•	•	•		
			KU19	•	•		•	•		•	•	•	0	•	•	•	0	•	•	•	•		
			KU18	•	•	0	•	•		•	•	•	0	•		•	0		•	•	•		
			KU17	•	•	0	•	•		•	•	•	0	•	•	•	0		•	•	•		
			KU16	•	•	0	•	•		•	•	•	0	•	•	•	0		•	•	•		
			KU15	•	•	0	•	•	•	•	•	•	0	•	•	•	0	•	•	•	•		
			KU14	•	•	0	•	•	•	•	•	•	0	•	•	•							
	MAASTRICHTIAN Tor Formtion	Højerup Member	L5/325-330	•	•	•	•		•	•	•	•											
MAASTRICHTIAN			L5/290-295	•	•	•	•	•	•	•	•												
			L5/125-130	•	•	•	•		•	•	•	•											
			L5/60-65	•	•	•	•	•	•	٠	•	٠											
			KUL-I										Bar	ren			1						
			KUL-K	•	•	•	•		•	•	•	•											
		SM	K8/5	•	•	•	•	•	•	•	•												
			K8/7	•	•	•	•	•	•	•	•												

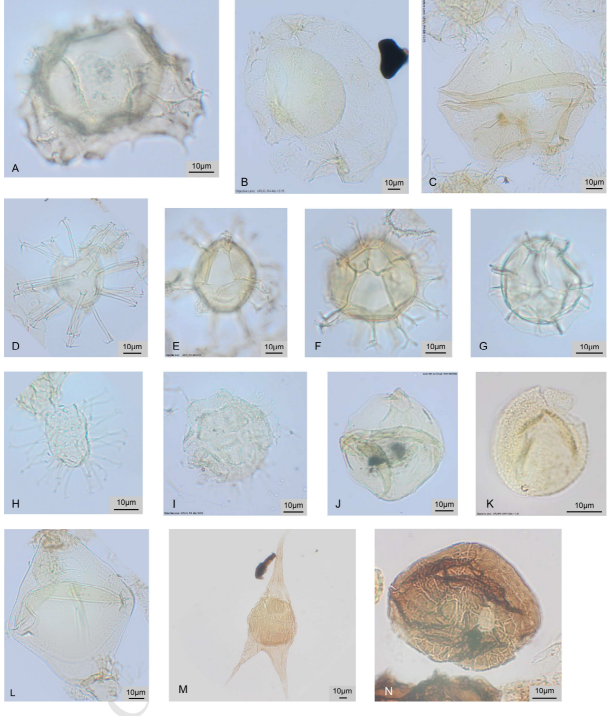


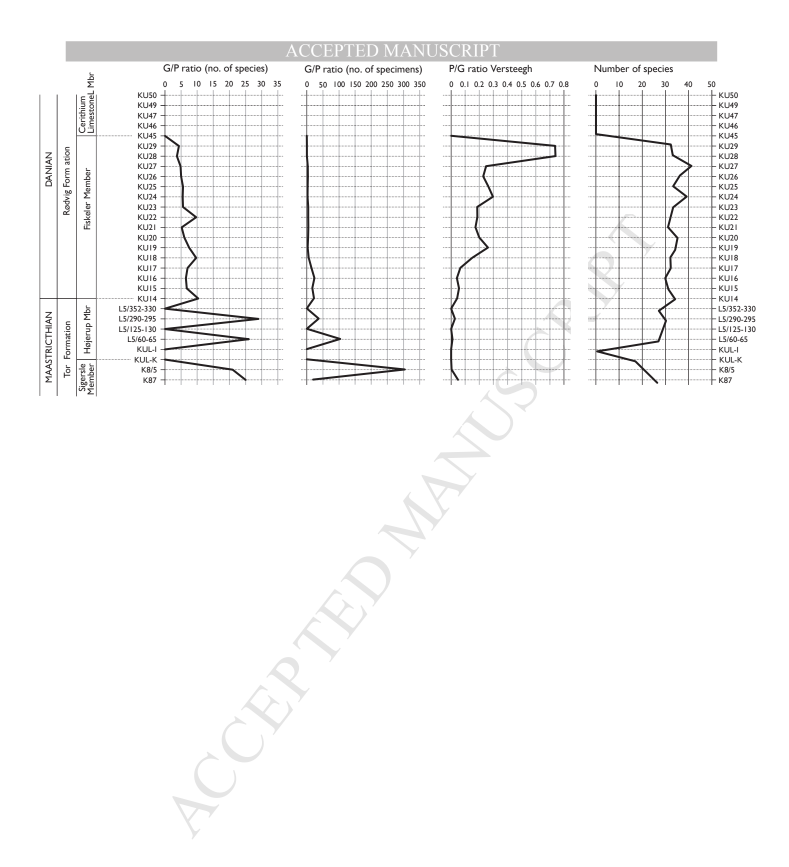












Highlights – Fitzpatrick et al.

The Cretaceous-Paleogene boundary succession at Stevns Klint is world famous and been a reference point for stratigraphy since the mid-nineteenth century. It is now a World Heritage Site and one of the most important of the distal K/Pg sites. Previous dinocyst research on the Stevns Klint succession has caused controversy, with some strange assemblages being described that may have been the result of preparation techniques. Unfortunately most of the sampling across the boundary has been limited to only a few sample points, often where the boundary clay (Fiskeler Member) is at its thinnest. Our research, using a large number of samples from an expanded succession of the boundary clay, has resolved many of the earlier problems and allowed us to identify a dinocyst stratigraphy across the boundary. This dinocyst stratigraphy can be used to compare the Stevns Klint succession with other boundary sections in Tunisia and Texas. One of the earlier suggestions, that the boundary clay is diachronous (despite containing the iridium anomaly), has been refuted.

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