## Palynology of Early Cretaceous (Barremian to Aptian) hydrocarbon (methane) seep carbonates and associated mudstones, Wollaston Forland, Northeast Greenland

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ABSTRACT. Palynostratigraphic and palynofacies analysis have been performed on hydrocarbon seep carbonate, carbonate nodule and mudstone samples from the Early Cretaceous Kuhnpasset Beds in the Kuhnpasset area of Wollaston Forland, Northeast Greenland. Three informal palynostratigraphic zones have been defined based on dinoflagellate cyst occurrences. The zones range from ?early Barremian to early Aptian in age, and correlate with previously defined dinoflagellate cyst zones in Northeast Greenland. These zones indicate hydrocarbon seepage in Kuhnpasset spanned the ?early–late Barremian age range, but did not continue into the Aptian, and thus seep activity continued for as much as three million years. Palynofacies analysis is used to interpret the depositional environment of the Kuhnpasset Beds and indicate that the hydrocarbon seeps and associated mudstones from this sedimentary sequence were deposited in a proximal shelf setting.

KEYWORDS: Palynostratigraphy, dinoflagellate cysts, palynofacies, hydrocarbon seep carbonates, Early Cretaceous, Northeast Greenland

## INTRODUCTION

Hydrocarbon seeps are places on the continental margins where fluids rich in hydrocarbons, principally methane, flow on to the seafloor (Sibuet and Olu, 1998). First discovered in 1984, they are now found in all the World's oceans. Hydrocarbon seeps support highly unusual biotic communities, which, together with the similarly structured hydrothermal vent communities, have altered our view of life in the deep sea, in part because the primary energy source for these ecosystems is not solar, but geochemical (Levin, 2005). In the shallow subsurface at seep sites methane is utilized by a consortium of methanotrophic archaea and sulphate-reducing bacteria in the anaerobic oxidation of methane reaction (e.g. Hinrichs et al., 1999; Boetius et al., 2000; Reitner et al., 2005), leading to the supersaturation of pore fluids with respect to carbonate ions and resulting in the formation of distinctive authigenic carbonate deposits with multi-phase carbonate cements and very negative  $\delta^{13}$ C values (e.g. Ritger et al., 1987; Aloisi et al., 2000; Naehr et al., 2007; Haas et al., 2010). Seep carbonates

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**Figure 1. A.** Geological locality map with field locality indicated. Map modified from Bjerager et al. (2020). Insert **B**. Map of Greenland with area of map A highlighted in red. White shows the extent of the Greenland icecap

have a wide variety of morphologies (e.g. nodules, tubular/tabular concretions, cemented breccias and pavements) and sizes (e.g. Campbell, 2006), and because of rapid precipitation commonly contain well-preserved macrofossils and microfossils, including palynomorphs (e.g. Kiel et al., 2013; Dalseg et al., 2016a, b).

The Mesozoic is a crucial time in the evolutionary history of seep faunas, with a change from brachiopod dominated communities of the Triassic, Jurassic and Early Cretaceous to bivalve dominated communities of the Late Cretaceous (Sandy, 2010). Palaeobathymetric controls on Mesozoic seep faunas occurred in seep communities in the Cretaceous, as in modern seeps, with the number of obligate species decreasing from the slope and deep shelf to the shallow shelves (Kiel, 2010). Compared to the Cenozoic, the Mesozoic fossil record of hydrocarbon seep communities is relatively sparse and comes largely from the circum-Pacific area (e.g. Japan and Western USA) and sites that were in the Tethys Ocean (principally Southern to Eastern Europe; Kiel et al., 2008; Kaim et al., 2013). Four areas of Jurassic to Cretaceous aged hydrocarbon seepage are known from the present-day Arctic region, including two sites in Northeast Greenland: one Barremian (Kuhnpasset, Wollaston Forland; Kelly et al., 2000)

and one Campanian (Leitch Bjerg, Geographical Society Ø); Svalbard (Jurassic-Cretaceous boundary; Hammer et al., 2011); Arctic Canada (Albian of Prince Patrick Island and Ellef Ringnes Island; Beauchamp and Savard, 1992; Williscroft et al., 2017) and Novaya Zemlya (three ages of seepage: late Oxfordian-early Kimmeridgian, late Tithonian and latest Berriasian-early Valanginian; Hryniewicz et al., 2015). In the Mesozoic, the latter three areas were part of the Boreal Ocean, which was a relatively isolated sea with limited marine connections with the Tethys and ancient Pacific (Panthalassic) Oceans (Zakharov et al., 2002). The Northeast Greenland seeps occurred in a more southerly position in a narrow seaway that linked the Boreal Ocean to the Tethys, through Northwest Europe.

The seeps in Kuhnpasset have been described by Kelly et al. (2000) and comprise over 30 exposed carbonate mounds (27 individually numbered) on the eastern side of Kuhnpasset, below the western flank of Aucellabjerget Mountain (Fig. 1). All but one of these are concentrated along a ridge in an area of  $\sim 1 \text{ km}^2$  (Kelly et al., 2000: fig. 4). The individual seep carbonates are subcircular to suboval in plan view, with relatively flat bases, where these can be seen. They vary in size from 1 to 3 m in diameter and are up to 1.8 m in height. The host rocks to the carbonate bodies are silty to sandy mudstones, and subordinate thin sandstone beds (Fig. 2). Kelly et al. (2000) erected the name Kuhnpasset Beds for this sequence and suggested a late Barremian age for it, based on the occurrence of the ammonite genera Audouliceras, Epicheloniceras and Sanmartinoceras, and the presence of dinoflagellate cysts indicative of the Pseudoceratium toveae Subzone (Nøhr-Hansen, 1993). Subsequently, Bjerager et al. (2020) incorporated the Kuhnpasset Beds into their Stratumbjerg Formation of the Brorson Halvø Group. Bjerager et al. (2020) suggested that the Stratumbjerg Formation was predominantly deposited in slope and basin-floor settings, with local passive infill of submarine gullies and shallow marine deposition. The latter facies is represented in the Kuhnpasset-Aucellabjerg area by the cross-bedded sandstones that overly the seep bearing mudstone sequence (Fig. 2). Bjerager et al. (2020) named these sandstones the Aucellabjerg Member, and this is thought





Figure 2. Lithological log with sample levels indicated (red boxes). Lithostratigraphy from Bjerager et al. (2020)

to be late early Aptian in age, based on dinoflagellates (Nøhr-Hansen et al., 2020) and the occurrence of the ammonite genus *Deshayesites* (Kelly et al., 2000).

In this paper we present a detailed investigation of the palynofloras from the seepbearing sedimentary sequence from Kuhnpasset, including, for the first time, from the seep carbonates themselves. Our aims are to better refine the age of the seep carbonates and the duration of hydrocarbon seepage in the area by comparing local palynostratigraphic zones with established Cretaceous dinoflagellate cyst zonations from Greenland and other areas from the Boreal realm. In addition, we perform a palynofacies analysis to interpret the depositional environment during the formation of the seeps in Kuhnpasset.

## MATERIALS AND METHODS

The material for this study was collected during fieldwork to Kuhnpasset between the 2<sup>nd</sup> and 12<sup>th</sup> August 2019 by Crispin T.S. Little, Hans Arne Nakrem, and Simon R.A. Kelly under Prospecting License No. 2017/15 from the Ministry of Industry, Energy and Research, Government of Greenland. A sedimentary log was made from seep carbonate number 27 of Kelly et al. (2000) at the base, to the sandstones of the Aucellabjerg Member at the top of the section (Fig. 2). We selected 13 mudstone samples from this section, one carbonate nodule and six seep carbonates for subsequent analysis (Supplementary File 1<sup>1</sup>).

Between 700 and 2000 g of carbonate nodule and seep carbonate samples were treated with 10-15% acetic acid at the University of Oslo. After five days, the acid had neutralized and the resulting residues were sieved. The  $<63 \mu m$  fraction was retained and sent with the mudstone samples (approximately 100 g per sample) to Applied Petroleum Technology A/S (APT), Oslo for standard processing onto palynological slides. These slides were studied by Emil Bang at the University of Oslo using a Leica DMPL microscope. Photomicrographs of palynomorphs were taken with a Leica camera MC170HD and Leica Application Suite v. 3.4.0 - LAS software. The images were stacked in Helicon Focus v.7.0 and edited in ACDsee Ultimate 8 and Adobe Photoshop CS6 software. The dinoflagellate cysts in the slides were identified based on shape, location of archaeopyle, and shape of processes (if present) using works by Davey (1982), Heilmann-Clausen and Birkelund (1987), Nøhr-Hansen (1993), Nøhr-Hansen et al. (2020) and Śliwińska et al. (2020).

Dinoflagellate cyst abundance was calculated as individuals of each taxon per total individuals in each sample. The abundances were sorted into four abundance groups: Rare (0-0.99%), Common (1.0-9.9%), Abundant (10.0-24.9%) and Dominant (25.0-100%). For each sample 300 specimens were counted. For dinoflagellate cyst diversity indices Simpson 1-D index, Shannon index (Shannon-Wiener index) and Fisher- $\alpha$  index were calculated. The calculations were performed using PAST software ver. 4.06 (Hammer et al., 2001; Hammer, 2021). For the palynofacies analysis, approximately 300 organic particles were counted for each sample. Ideally 500 organic particles should be counted (Tyson, 1995), but many of our slides contained fewer than this, therefore we consider 300 organic particles to be adequate for our analyses. The counts were divided into the following groups: structured organic matter (spores, pollen grains, dinoflagellate cysts, cuticles, coaly phytoclasts, woody phytoclasts, other plant debris) and structureless organic matter (amorphous organic matter; AOM). Cuticles, coaly phytoclasts, woody phytoclasts and other plant debris are furthermore grouped in phytoclasts.

Figured specimens (Pl. 1) are curated in the Natural History Museum, University of Oslo, abbreviated PMO (former Paleontologisk Museum Oslo). The taxonomy used here follows that of Williams et al. (2017).

### RESULTS

The preservation of palynomorphs in the samples was very good throughout all samples, with seemingly no difference between the seep carbonate and mudstone samples. However, palynomorphs in the uppermost two samples were less well preserved than those in the other samples. This material also had a darker color (TAI 2/2+) than the other samples, which generally had a bright yellow coloration (TAI 1/1+). In total, 99 different dinoflagellate cyst taxa were identified from the carbonate and mudstone samples (Pl. 1, Supplementary File  $2^2$ ). This included 41 genera, 54 species, one affinity (aff.) to a species, four confers (cf.) to a species and five confers to a genus; four were primarily identified to genus level (Gonyaulacysta sp., Kiokansium sp., Spiniferites sp., Subtilisphaera sp.). From the mudstone samples a total of 84 taxa were identified, including 48 which were identified to species level. The remaining taxa were placed in open nomenclature. For the carbonate samples a total of 69 taxa were identified, 39 to species level, 30 in open nomenclature. The terrestrial palynomorphs observed in the samples were not identified more specifically than general groups of spores or pollen. In addition, some freshwater algae were observed. Only two observed palynomorphs were identified as being reworked. These were a Jurassic dinoflagellate cysts of the genus *Rigaudella* Below 1982 (sensu Nøhr-Hansen, 1993) in Seep 3A (90.8 m level), and the Permian bisaccate pollen Protohaploxypinus sp. in the carbonate nodule sample (78.3 m level).

#### PALYNOSTRATIGRAPHIC ANALYSIS

The stratigraphic distribution of identified dinoflagellate cyst taxa is presented in Figs 3 and 4. Using these stratigraphic ranges we divide our Kuhnpasset section into three informal zones based on first occurrences (FO) and last occurrences (LO) of selected dinoflagellate cyst species (Fig. 4). An informal zoning is preferred as the materials in this study were not collected with regular sampling intervals; rather, sampling was adjusted to the seep

<sup>&</sup>lt;sup>1</sup> Supplementary File 1: Sample list and weights

 $<sup>^2\;</sup>$  Supplementary File 2: Dinoflagellate cyst taxa counts per sample.



Plate 1. Representative dinoflagellate cyst species from the Kuhnpasset samples. 1. Vesperopsis longicornis (Batten and Lister, 1988) Harding 1990a, 85.80 m level, PMO 222.088C, EF W23 (EF: England finder coordinate); 2. Batioladinium exiguum (Alberti, 1961) Brideaux, 1975, 85.80 m level, PMO 222.088, EF J39; 3. Sirmiodinium grossii Alberti, 1961 emend. Warren, 1973, carbonate nodule, 78.30 m level, PMO 222.091, EF R33; 4. Subtilisphaera perlucida (Alberti, 1959) Jain and Millepied, 1973 (Jain and Millepied, 1973), 101 m level, PMO 222.091, EF D23; 5. Hystrichosphaerina schindewolfii Alberti, 1961, Seep 26, PMO 222.107, EF E37; 6. Muderongia sp., 85.80 m level, PMO 222.089, EF J31; 7. Odontochitina nuda (Gocht, 1957) Dörhöfer and Davies, 1980, 86.60 m level, PMO 222.086C, EF C31; 8. Pseudoceratium toveae Nøhr-Hansen, 1993, carbonate nodule, 78.30 m level, PMO 222.107, EF P21; 10. Hystrichosphaeridium arborispinum Davey and Williams, 1966, 85.80 m level, PMO 222.088, EF U37; 11. Gardodinium traberculosum (Gocht, 1959) Davey, 1978 emend. Harding, 1996, Seep 8, PMO 222.106, EF T23; 12. Pseudoceratium anaphrissum (Sarjeant, 1966) Bint, 1986 emend. Harding, 1990b, Seep 4, PMO 222.105, EF C35; 13. Batioladinium longicornutum (Alberti, 1961) Brideaux, 1975, 86.70 m level, PMO 222.085, EF K29; 14. ?Endoscrinium sp., Seep 27, PMO 222.110, EF M35. Scale bar 20 µm in 1–6, 8–12 and 14; scale bar 50 µm in 7 and 13



Figure 3. Dinoflagellate cyst generic range chart, with abundance data and informal zones 1–3

carbonate and adjacent mudstone intervals. Our informal zones are defined as follows, from base upwards:

**Zone 1.** This encompasses all samples from Seep 27 up to Seep 26, at 97.6 m in the section. The upper zone boundary is defined at the LO of *Hystrichosphaerina schindewolfii* and the FO of *Palaeoperidinium cretaceum*. The lower boundary of the zone is left open. *Batioladinium exiguum*, *Endoscrinium campanula* and *Hystrichosphaeridium arborispinum* have their first and last occurrences within this zone.

**Zone 2.** This spans the interval from Seep 26 to the uppermost mudstone sample above New Seep 2019, at 104.6 m in the section. The zone is transitional with stepwise LOs of different species through the zone. The lower boundary is placed at the LO of *Hystrichosphaerina* schindewolfii, and the upper boundary at the LO of *Pseudoceratium anaphrissum*, *Chlamydophorella nyei* and *Gardodinium trabeculosum*. The zone boundaries coincide with the FO and LO of *Palaeoperidinium cretaceum*.

**Zone 3.** This comprises the two uppermost samples in the section (115.0 and 116.3 m levels). The lower boundary of the zone is defined by the LO of *Chlamydophorella nyei*, *Gardodinium trabeculosum* and *Palaeoperidinium cretaceum*. No upper boundary is defined here.

Our informal zones 1-3 can be roughly correlated with the Northeast Greenland dinoflagellate zones defined by Nøhr-Hansen (1993) and Nøhr-Hansen et al. (2020). The oldest possible age for zone 1 is early Barremian as Pseudoceratium anaphrissum is present throughout the entire zone. According to Nøhr-Hansen (1993), P. anaphrissum has its uppermost occurrence in the early Barremian Pseudoceratium anaphrissum Subzone. In our study Pseudoceratium toveae occurs as a dominant species at 86.6 m. This may correlate with the acme of this species recorded in the Pseudoceratium toveae Subzone by Nøhr-Hansen (1993). However, this subzone was defined above the LO of P. anaphrissum and dated as late Barremian (Nøhr-Hansen, 1993). If so, it suggests that our



Figure 4. Dinoflagellate cyst species range chart, with abundance data and informal zones 1-3

zone 1 is of early-late Barremian age. It is also possible that *P. anaphrissum* is reworked, but the common occurrence of this species at the top of subzone 1 makes this unlikely. Our zones 1 and 2 could represent the transition from the P. anaphrissum Subzone to the P. toveae Subzone, but a more detailed investigation is needed to address this question. Within our zone 2, Batioladinium longicornutum has its LO, suggesting that this zone has a latest Barremian age. This is based on correlation with the Batioladinium longicornutum zone redefined by Nøhr-Hansen et al. (2020), which is recognized on the basis of the FO and LO of Batioladinium longicornutum. With Odontochitina nuda present in our zone 3, and the LO of Batioladinium longicornutum in our zone 2, zone 3 is likely of early Aptian age. This is based on correlation with the Odontochitina nuda Zone of early Aptian age in the work of Nøhr-Hansen (1993), the lower boundary of which is defined on the basis of the LO of Batioladinium longicornutum and the upper boundary at the LO of Odontochitina nuda. Thus, seepage in Kuhnpasset spanned the early-late Barremian age range, but did not continue into the Aptian. This extends the duration for seepage suggested by Kelly et al. (2000) from the late to the early Barremian, and indicates that there was seep activity over some extended period of time, possibly as much as three million years, beginning with the formation of Seep 27 and ending at New Seep 2019. Saying that, the majority of seepage would have occurred over a shorter period of time, perhaps confined to the late Barremian, as the majority of seep deposits are confined to 20 m of the Kuhnpasset section (Fig. 2).

In addition to the key species listed above, the occurrence of Pseudoceratium anaphrissum (Late Hauterivian–late early Barremian) and Vesperopsis longicornis (late Barremianearly Albian) in our material is also of biostratigraphic significance. In our study we found Batioladinium longicornutum had an LO before that of Pseudoceratium anaphrissum. This contrasts with the Northeast Greenland dinoflagellate cyst biostratigraphy of Nøhr-Hansen (1993) and Nøhr-Hansen et al. (2020), where the late Barremian Pseudoceratium toveae Subzone is defined from the LO of Pseudoceratium anaphrissum to the LO of Batioladinium longicornutum. An explanation for this difference could be that the LO

of *Batioladinium longicornutum* is within the upper part of the section, from which we did not collect samples for palynomorph study (between 104.8 and 115.0 m), or because of reworking, as we found reworked Permian and Jurassic palynomorphs in some of our samples.

Although not the subject of this study, the Kuhnpasset Beds have a rich ammonite fauna, dominated by lytoceratids, but also including aconeceratids, cheloniceratids and heteromorphs. These are currently undescribed, but based on preliminary investigation, compare with some of Rosenkrantz's (1934) material from Kuhn Ø and Frebold's (1935) from Storre Koldewey, both of stated Aptian age. However, the earliest reliably dated ammonite which we discovered in 2019 at Kuhnpasset was a specimen of Deshayesites boegvadi Rosenkrantz (1934), from about 2 m above the base of the Aucellabjerg Member (122 m on Fig. 2). This ammonite was taken as an indicator for the mid early Aptian in Hold with Hope (Kelly and Whitham, 1999) and confirms an early Aptian age for the strata above the Kuhnpasset seeps. Further work to reconcile ammonite and dinoflagellate cyst biostratigraphy in Kuhnpasset specifically and Northeast Greenland in general would be beneficial.

## DINOFLAGELLATE CYST DIVERSITY

Within our zone 1, the number of taxa per sample varies between 27 and 46, in zone 2, between 29 and 37. In zone 3, the number varies between 19 and 20. Our diversity data corresponds well with the observations of Nøhr-Hansen (1993) who reported similar diversities in the Barremian samples, and a lower diversity in the late early Aptian – early Albian samples. Note that Nøhr-Hansen (1993) recorded diversity as the number of species, whereas here diversity is recorded as the total number of taxa, therefore a direct comparison of the actual number of species is not possible.

Our statistical analyses indicate that in the mudstone samples the Simpson 1-D index averages at 0.9 and varies between 0.9 and 1.0. In the carbonates the Simpson 1-D also averages at 0.9, with no variation. The Shannon H index in the carbonate samples has the lowest value at 2.9 and highest at 3.2, averaging at 3.0. This is slightly lower than the mudstone samples averaging at 3.1. The



**Figure 5.** Variation in diversity indices (**A**. Fisher- $\alpha$ ; **B**. Shannon-H; **C**. Simpson-1-D) for three sample material types (mudstones, seep carbonates, and carbonate nodule)

mudstone samples also have larger outliers, with lowest H at 2.4 and highest at 3.5. The Fisher- $\alpha$  averages at 10.8 in the mudstone samples and 9.8 in the carbonates. The lowest value for Fisher- $\alpha$  in the mudstones is 5.8 and 8.9 for the carbonates. The highest value for the latter is 11.1 while for the mudstones it is 15.0. The variation between sample types and diversity indices is presented in Fig. 5. In general, our recorded diversity indices show a relatively stable Simpson 1-D value for all samples, whereas the Fisher- $\alpha$  and Shannon H values have larger variation. According to this, it seems that the dinoflagellate cyst diversity in general is quite high, but with a slightly lower diversity in the seep carbonates. This could be the result of a changing environment when hydrocarbon seepage took place, which impacted the water column around the area of seepage, or due to differential preservation in the sediment due to the anaerobic oxidation of methane producing chemical gradients not conducive to the preservation of organic matter. There is also a decreasing trend in diversity in the two uppermost mudstone samples. This may be partly due to the poorer preservation observed in these samples.

### PALYNOFACIES ANALYSIS

Results of the palynofacies analyses are presented in Figs 6 and 7 which shows the relative percentage of AOM, phytoclasts and palynomorphs (both marine and terrestrial) in each sample as single points in Tyson's Ternary Diagram (Tyson, 1995; Miles, 2021; Supplementary Files 3 and  $4^3$ ). The dinoflagellate cyst contribution in the carbonates is generally low, varying between 0.3% and 3.2%. This is also the case with cuticles (0.3-2.9%) and AOM (0.9-4.2%). For spores the general trend lies between ~10% and ~20%, and for pollen grains it is  $\sim 3-15\%$ . Both spores and pollen grains have a similar outlier in Seep 8, where they comprise only 1.0% of the total counts. The Seep 8 sample also differs from the other carbonate samples in containing 47.7% coaly particles counting in the total assemblage. In the remaining carbonate samples coaly particles comprise between  $\sim 18\%$  and  $\sim 33\%$ . In the Seep 4 sample, the plant debris constitutes 51.0% of the palynomorph assemblage. For the other carbonate samples this value is between ~15% and ~27%. Woody particles in the carbonate samples vary between  $\sim 15-30\%$ , apart from a lower outlier in Seep 4 with only 9.4%.

In the mudstone samples (85.8-87.6 m interval) below Seep 4, spores contribute between ~ 6% and ~18%, pollen grains ~5–10% with a higher value at level 86.7 m with 18.4%. Dinoflagellate cysts contribute between  $\sim 1\%$ and  $\sim 3\%$ , with an outlier at level 86.3 m with 4.5%. The proportion of cuticles is  $\sim 1.7\%$  in these samples, with a higher contribution at level 86.3 m, where they contribute 4.2%. Most of the samples have a larger portion of woody particles than coaly particles, except for the sample level 86.9 m, where the coaly material counts for 26.1% and woody particles only for 21.7%. The palynomorph group that makes up the most of these mudstone samples is plant debris, varying between ~25% and ~35%. AOM is generally equal in the samples,  $\sim 3.7\%$ .

At sample level 86.9 m, the palynomorph groups that dominate the palynofacies assemblage are coal (20.8-25.9%), woody particles (20.8-23.3%) and plant debris (30.6-33.7%). AOM, cuticles and dinoflagellate cysts each make up less than 5% of the total assemblage.

<sup>&</sup>lt;sup>3</sup> Supplementary File 3: Palynology similarity indices; Supplementary File 4: Palynofacies organic particle data



Figure 6. Tyson palynofacies ternary diagram for the Kuhnpasset samples, grouped into four types. AOM – amorphous organic matter; I – highly proximal shelf or basin; II – marginal dysoxic-anoxic basin; III – heterolithic oxic shelf (distal shelf); IVa – shelf to basin transition (dysoxic); IVb – shelf to basin transition (anoxic); V – mud dominated oxic shelf (distal shelf); VI – proximal suboxic-anoxic shelf; VII – distal dysoxic-anoxic shelf; VIII – distal dysoxic-suboxic shelf; IX – distal suboxic-anoxic shelf

Spore contents vary between 10.6% at 103.3 m and 9.1% at 104.8 m. In the upper of the two mudstone samples in our zone 3 (at 116.3 m), coaly particles make up the largest part with 37.6%, while plant debris and woody particles make up 27.4% and 20.2% respectively. The same groups dominate in the lower sample



Figure 7. Contents of organic particles through the Kuhnpasset samples. Stippled pattern indicates non-sampled interval

from zone 3 (at 115.0 m), but in this plant debris makes up the largest fraction with 24.9%. Woody and coaly particles both make up ~20% of this sample. The dinoflagellate cyst fraction is very low in the zone 3 samples, with 0.6% in the upper sample, while in the lower sample the value reaches 3.4%. AOM is close to the same value in both samples (4.4% and 4.8%) and cuticles in the lower sample constitute 4.0% while being only 1.9% in the upper sample. Spores and pollen make up a larger part in the lower sample, with 9.3% and 12.8% respectively, than in the upper sample where they are 2.9% and 4.8%.

We use the results of the palynofacies analyses to interpret the possible depositional environment of the carbonate seeps and mudstones in our section. Palynofacies alone is not a definitive way to determine a depositional environment, but it is useful as a supplement to other paleontological and sedimentological observations (Batten and Stead, 2005). The carbonate samples and the mudstone samples all plot within palynofacies fields I and III (Fig. 6), which correspond to "highly proximal shelf or basin" and "heterolithic oxic shelf (proximal shelf)" of Tyson (1995) respectively. The carbonate samples have a slightly greater spread in the diagram. With respect to kerogen type, the palynofacies field I represents kerogen type III which is gas prone, and palynofacies field III represents kerogen type III or IV, both are gas prone kerogen. Based on the position of the samples in the ternary diagram and variations in terrestrial vs. marine palynomorphs, we interpret that the seep containing sedimentary sequence at Kuhnpasset was deposited in a proximal shelf environment, shallower than the majority of the Stratumbjerg Formation in Northeast Greenland, which Bjerager et al. (2020) suggested was predominantly deposited in a basin-floor and slope environment.

## CONCLUSIONS

From the palynostratigraphic and palynofacies analyses performed on six hydrocarbon seep carbonates, one carbonate nodule and 13 associated mudstone samples from the Kuhnpasset Beds, Wollaston Forland, Northeast Greenland, we draw the following conclusions: - We propose three informal dinoflagel-

late cyst zones from the samples. Based on

stratigraphically important species, our lower zone 1 is assigned an ?early–late Barremian age, our middle zone 2 is late Barremian and our upper zone 3 is early Aptian in age.

- Based on dinoflagellate cyst biostratigraphy, hydrocarbon seepage in the Kuhnpasset area spanned the early – late Barremian age range, but did not continue into the Aptian. This indicates seep activity continued for as much as three million years, although the majority of seepage would have occurred over a shorter period of time, perhaps confined to the late Barremian.

- Statistical analysis shows a generally high dinoflagellate cyst diversity with a slightly lower diversity in the carbonate samples compared to the mudstone samples. The two uppermost mudstone samples differ from the rest of the studied material by yielding the lowest diversity and poorest preservation of dinoflagellate cysts.

- The dinoflagellate cyst assemblages recorded in this study show close similarities with other assemblages of similar age recorded in the Arctic/Boreal region.

- Based on the observed palynofacies material and relative proportions of marine and terrestrial palynomorphs, the carbonates and mudstones analyzed in this study were deposited in a proximal shelf setting.

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#### REFERENCES

- Alberti, G., 1959. Zur Kenntnis der Gattung Deflandrea Eisenack (Dinoflag.) in der Kreide und im Alttertiär Nord-und Mitteldeutschlands. Mitteilungen aus dem Geologischen Staatsinstitut in Hamburg 28(9), 3-105.
- Alberti, G., 1961. Zur Kenntnis mesozoischer und alttertiärer Dinoflagellaten und Hystrichosphaerideen von Nord- und Mitteldeutschland sowie einigen anderen europäischen Gebieten. Palaeontographica, Abt. A 116(1-4), 1-58.

- Aloisi, G., Pierre, C., Rouchy, J.-M., Foucher, J.-P., Woodside, J., 2000. Methane-related authigenic carbonates of eastern Mediterranean Sea mud volcanoes and their possible relation to gas hydrate destabilization. Earth and Planetary Science Letters 184, 321–338. https://doi.org/10.1016/S0012-821X(00)00322-8
- Batten, D., Lister, J., 1988. Early Cretaceous dinoflagellate cysts and chlorococcalean algae from freshwater and low salinity palynofacies in the English Wealden. Cretaceous Research 9(4), 337–367. https://doi.org/10.1016/0195-6671(88)90007-9
- Batten, D.J., Stead, D.T., 2005. Palynofacies analysis and its stratigraphic application. In: Koutsoukos, E.A.M. (ed.), Applied stratigraphy. Springer, Dordrecht, pp. 203–226. https://doi.org/10.1007/1-4020-2763-X\_10
- Beauchamp, B., Savard, M., 1992. Cretaceous chemosynthetic carbonate mounds in the Canadian Arctic. PALAIOS 7(4), 434–450. https://doi. org/10.2307/3514828
- Below, R., 1982. Scolochorate Zysten der Gonyaulacaceae (Dinophyceae) aus der Unterkreide Marokkos. Palaeontographica, Abt. B Paläophytologie 182(1–3), 1–51.
- Bint, A.N., 1986. Fossil Ceratiaceae: A restudy and new taxa from the mid-Cretaceous of the western interior, USA. Palynology 10(1), 135–180. https:// doi.org/10.1080/01916122.1986.9989307
- Bjerager, M., Alsen, P., Bojesen-Koefoed, J., Fyhn, M.B., Hovikoski, J., Ineson, J.R., Nøhr-Hansen, H., Nielsen, L.H., Piasecki, S., Vosgerau, H., 2020. Cretaceous lithostratigraphy of North-East Greenland. Bulletin of the Geological Society of Denmark 68, 37–93. https://doi.org/10.37570/bgsd-2020-68-04
- Boetius, A., Ravenschlag, K., Schubert, C.J., Rickert, D., Widdel, F., Gieseke, A., Amann, R., Jorgensen, B.B., Witte, U., Pfannkuche, O., 2000. A marine microbial consortium apparently mediating anaerobic oxidation of methane. Nature 407(6804), 623–626. https:// doi.org/10.1038/35036572
- Brideaux, W.W., 1975. Taxonomic note: redefinition of the genus *Broomea* and its relationship to *Batioladinium* gen. nov. (Cretaceous). Canadian Journal of Botany 53(12), 1239–1243. https://doi. org/10.1139/b75-148
- Campbell, K.A., 2006. Hydrocarbon seep and hydrothermal vent paleoenvironments and paleontology: Past developments and future research directions. Palaeogeography, Palaeoclimatology, Palaeoecology 232, 362–407. https://doi. org/10.1016/j.palaeo.2005.06.018
- Dalseg, T.S., Nakrem, H.A., Smelror, M., 2016a. Dinoflagellate cyst biostratigraphy, palynofacies, depositional environment and sequence stratigraphy of the Agardhfjellet Formation (Upper Jurassic-Lower Cretaceous) in central Spitsbergen (Arctic Norway). Norwegian Journal of Geology 96(2), 119–133. http://dx.doi.org/10.17850/njg96-2-04
- Dalseg, T.S., Nakrem, H.A., Smelror, M., 2016b. Organic-walled microfossils and palynodebris in

cold seep carbonate deposits: The Upper Jurassic– Lower Cretaceous Agardhfjellet Formation on Svalbard (Arctic Norway). Norwegian Journal of Geology 96(2), 135–146. http://dx.doi.org/10.17850/njg96-2-01

- Davey, R.J., 1970. Non-calcareous microplankton from the Cenomanian of England, northern France and North America, part II. British Museum (Natural History) Geology Bulletin 18(8), 333–397.
- Davey, R.J., 1978. Marine Cretaceous palynology of Site 361, D.S.D.P. Leg 40, off southwestern Africa. Initial Reports of the Deep Sea Drilling Project, Volume XL, Washington, 883–913. http://dx.doi. org/10.2973/DSDP.PROC.40.125.1978
- Davey, R.J., 1982. Dinocyst stratigraphy of the latest Jurassic to Early Cretaceous of the Haldager No 1 borehole, Denmark. Danmarks Geologiske Undersøgelse Serie B(6), 57 pp. https://doi.org/10.34194/ serieb.v6.7061
- Davey, R.J., Williams, G.L., 1966. V. The genus Hystrichosphaeridium and its allies. In: Davey, R.J., Downie, C., Sarjeant, W.A.S., Williams, G.L. (eds), Studies on Mesozoic and Cainozoic dinoflagellate cysts. British Museum (Natural History) Geology, Bulletin, Supplement 3, 53–106.
- Dörhöfer, G., Davies, E.H., 1980. Evolution of archeopyle and tabulation in rhaetogonyaulacinean dinoflagellate cysts. Miscellaneous Publication, 91 pp., Royal Ontario Museum, Life Sciences Division, Toronto.
- Frebold, H., 1935. Marines Aptien von der Köldewey Insel (Nördliches Ostgrönland). Meddelelser om Grønland 95(4), 1-112, pls 1-8.
- Gocht, H., 1957. Mikroplankton aus dem nordwestdeutschen Neokom (Teil I). Paläontologische Zeitschrift 31(3-4), 163-185.
- Gocht, H., 1959. Mikroplankton aus dem nordwestdeutschen Neokom (Teil II). Paläontologische Zeitschrift 33(1-2), 50-89.
- Haas, A., Peckmann, J., Elvert, M., Sahling, H., Bohrmann, G., 2010. Patterns of carbonate authigenesis at the Kouilou pockmarks on the Congo deepsea fan. Marine Geology 268, 129–136. https://doi. org/10.1016/j.margeo.2009.10.027
- Hammer, Ø., 2021. PAST PAleontological STatistics Version 4.06. Available from: https://www.nhm.uio. no/english/research/infrastructure/past/. Accessed April 2021.
- Hammer, Ø., Harper, D.A.T., Ryan, P.D., 2001. PAST: Paleontological statistics software package for education and data analysis. Palaeontologia Electronica 4(1), 9. https://palaeo-electronica.org/2001\_1/ past/past.pdf
- Hammer, Ø., Nakrem, H.A., Little, C.T.S., Hryniewicz, K., Sandy, M.R., Hurum, J.H., Druckenmiller, P., Knutsen, E.M., Høyberget, M., 2011. Hydrocarbon seeps from close to the Jurassic– Cretaceous boundary, Svalbard. Palaeogeography, Palaeoclimatology, Palaeoecology 306(1-2), 15-26. https://doi.org/10.1016/j.palaeo.2011.03.019
- Harding, I.C., 1990a. Palaeoperidinium cretaceum: a brackish-water peridiniinean dinoflagellate from the Early Cretaceous. Palaeontology 33(1), 35–48.

- Harding, I.C., 1990b. A dinocyst calibration of the european Boreal Barremian. Part I: introduction, stratigraphy, systematics. Part II: systematics, biostratigraphy. Palaeontographica, Abt. B 218(1-3), 1-76.
- Harding, I.C., 1996. Taxonomic stabilisation of dinoflagellate cyst taxa, as exemplified by two morphologically complex Early Cretaceous species. Review of Palaeobotany and Palynology 92(3–4), 351–366. https://doi.org/10.1016/0034-6667(95)00114-X
- Heilmann-Clausen, C., Birkelund, T., 1987. Lower Cretaceous dinoflagellate biostratigraphy in the Danish Central Trough. Danmarks Geologiske Undersøgelse Serie A (17), 1–89. https://doi.org/10.34194/ seriea.v17.7036
- Hinrichs, K.-U., Hayes, J.M., Sylva, S.P., Brewer, P.G., DeLong, E.F., 1999. Methane-consuming archaebacteria in marine sediments. Nature 398(6730), 802–805.
- Hryniewicz, K., Hagström, J., Hammer, Ø., Kaim, A., Little, C.T.S., Nakrem, H.A., 2015. Late Jurassic– Early Cretaceous hydrocarbon seep boulders from Novaya Zemlya and their faunas. Palaeogeography, Palaeoclimatology, Palaeoecology 436, 231– 244. https://doi.org/10.1016/j.palaeo.2015.06.036
- Jain, K.P., Millepied, P., 1973. Cretaceous microplankton from Senegal basin, NW Africa. 1 Some new genera, species and combinations of dinoflagellates. The Palaeobotanist 20, 22–32.
- Kaim, A., Skupien, P., Jenkins, R.G., 2013. A new Lower Cretaceous hydrocarbon seep locality from the Czech Carpathians and its fauna. Palaeogeography, Palaeoclimatology, Palaeoecology 390, 42–51. https://doi.org/10.1016/j.palaeo.2013.03.010
- Kelly, S.R.A., Blanc, E., Price, S.P., Whitham, A.G., 2000. Early Cretaceous giant bivalves from seeprelated limestone mounds, Wollaston Forland, Northeast Greenland. Geological Society, London, Special Publications 177(1), 227–246. https://doi. org/10.1144/GSL.SP.2000.177.01.13
- Kelly, S.R.A., Whitham, A.G., 1999. Deshayesitid ammonites from the lower Aptian (Lower Cretaceous) of North-East Greenland. Scripta Geologica, Special Issue, 3, 83–95.
- Kiel, S., 2010. On the potential generality of depthrelated ecologic structure in cold-seep communities: Evidence from Cenozoic and Mesozoic examples. Palaeogeography, Palaeoclimatology, Palaeoecology 295, 245-257. https://doi.org/10.1016/j.palaeo.2010.05.042
- Kiel, S., Campbell, K.A., Elder, W.P., Little, C.T.S., 2008. Jurassic and Cretaceous gastropods from hydrocarbon-seeps in forearc basin and accretionary prism settings, California. Acta Palaeontologica Polonica 53, 679–703. http://dx.doi.org/10.4202/ app.2008.0412
- Kiel, S., Birgel, D., Campbell, K.A., Crampton, J.S., Schiøler, P., Peckmann, J., 2013. Cretaceous methane-seep deposits from New Zealand and their fauna. Palaeogeography, Palaeoclimatology, Palaeoecology 390, 17–34. https://doi.org/10.1016/j. palaeo.2012.10.033

- Lentin, J.K., Williams, G.L., 1975. A monograph of fossil peridinioid dinoflagellate cysts. Canada Bedford Institute Report BI. 1975-16, 237 pp.
- Levin, L.A., 2005. Ecology of cold seep sediments: interactions of fauna with flow, chemistry and microbes. Oceanography and Marine Biology. CRC Press, 11–56.
- Miles, N., 2021. Palynofacies A guide to the "Dark Side" of Palynolgy [Online]. Petrostrat – applied stratigraphy. Available from: https://www.petrostrat. com/educational-articles/palynofacies-a-guide-to-thedark-side-of-palynolgy/. Accessed July 2021.
- Naehr, T.H., Eichhubl, P., Orphan, V.J., Hovland, M., Paull, C.K., Ussler III, W., Lorenson, T.D., Greene, H.G., 2007. Authigenic carbonate formation at hydrocarbon seeps in continental margin sediments: a comparative study. Deep-Sea Research II 54, 1268– 1291. https://doi.org/10.1016/j.dsr2.2007.04.010
- Nøhr-Hansen, H., 1993. Dinoflagellate cyst stratigraphy of the Barremian to Albian, Lower Cretaceous, North-East Greenland. Grønlands Geologiske Undersøgelse Bulletin 166, 1–171. https://doi. org/10.34194/bullggu.v166.6722
- Nøhr-Hansen, H., Piasecki, S., Alsen, P., 2020. A Cretaceous dinoflagellate cyst zonation for NE Greenland. Geological Magazine 157, 1658–1692. https:// doi.org/10.1017/S0016756819001043
- Pocock, S.A.J., 1962. Microfloral analysis and age determination of strata at the Jurassic-Cretaceous boundary in the western Canada plains. Palaeontographica, Abt. B 111, 1–95.
- Reitner, J., Peckmann, J., Blumenberg, M., Michaelis, W., Reimer, A., Thiel, V., 2005. Concretionary methane-seep carbonates and associated microbial communities in Black Sea sediments. Palaeogeography, Palaeoclimatology, Palaeoecology 227, 18–30. https://doi.org/10.1016/j.palaeo.2005.04.033
- Ritger, S., Carson, B., Suess, E., 1987. Methane-derived authigenic carbonates formed by subductioninduced pore-water expulsion along the Oregon/ Washington margin. Geological Society of America Bulletin 98, 147–156. https://doi.org/10.1130/0016-7606(1987)98<147:MACFBS>2.0.CO;2
- Rosenkrantz, A., 1934. Beschreibung der von R. Bøgvad wahrend der Jahre 1929–30 in Ostgrönland gesammelten unterkretacischen Fossilien und die sich aus diesen ergebenden stratigraphischen Schlussfolgerungen. Meddelelser om Grønland 93(1), 17–28, pls 1–5.

- Sandy, M.R., 2010. Brachiopods from ancient hydrocarbon seeps and hydrothermal vents. In: Kiel, S. (ed.), The vent and seep biota, Topics in Geobiology 33, Springer, Berlin, pp. 279–314.
- Sarjeant, W.A.S., 1966. Further dinoflagellate cysts from the Speeton Clay. In: Davey, R.J., Downie, C., Sarjeant, W.A.S., Williams, G.L. (eds), Studies on Mesozoic and Cenozoic dinoflagellate cysts. British Museum (Natural History) Geology, Bulletin, Supplement 3, 199–214.
- Sibuet, M., Olu, K., 1998. Biogeography, biodiversity and fluid dependence of deep-sea cold-seep communities at active and passive margins. Deep Sea Research Part II. Topical Studies in Oceanography 45(1), 517–567.
- Śliwińska, K.K., Jelby, M.E., Grundvåg, S.-A., Nøhr-Hansen, H., Alsen, P., Olaussen, S., 2020. Dinocyst stratigraphy of the Valanginian-Aptian Rurikfjellet and Helvetiafjellet formations on Spitsbergen, Arctic Norway. Geological Magazine 157, 1693– 1714. https://doi.org/10.1017/S0016756819001249
- Tyson, R.V., 1995. Sedimentary Organic Matter: Organic facies and palynofacies. Springer, Dordrecht. https://doi.org/10.1007/978-94-011-0739-6
- Warren, J.S., 1973. Form and variation of the dinoflagellate Sirmiodinium grossi Alberti, from the Upper Jurassic and Lower Cretaceous of California. Journal of Paleontology 47(1), 101–114.
- Williams, G.L., Fensome, R.A., MacRae, R.A., 2017. DINOFLAJ3. American Association of Stratigraphic Palynologists, Data Series no. 2. Available at http://dinoflaj.smu.ca/dinoflaj3
- Williscroft, K., Grasby, S.E., Beauchamp, B., Little, C.T.S., Dewing, K., Birgel, D. Poulton, T., Hryniewicz, K., 2017. Extensive Cretaceous (Albian) methane seepage on Ellef Ringnes Island, Canadian High Arctic. Geological Society of America Bulletin 129, 788–805. https://doi.org/10.1130/B31601.1
- Zakharov, V.A., Shurygin, B.N., Kurushin, N.I., Meledina, S.V., Nikitenko, B.L., 2002. A Mesozoic ocean in the Arctic: Paleontological Evidence. Russian Geology and Geophysics 43, 143–170.