# Amino acid racemization in *Neogloboquadrina pachyderma* and *Cibicidoides wuellerstorfi* from the Arctic Ocean and its implications for age models

5 Gabriel West<sup>1,2</sup>, Darrell S. Kaufman<sup>3</sup>, Martin Jakobsson<sup>1,2</sup>, Matt O'Regan<sup>1,2</sup>

Correspondence to: Gabriel West (gabriel.west@geo.su.se)
Matt O'Regan (matt.oregan@geo.su.se)

#### 15 Abstract

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We report the results of amino acid racemization (AAR) analyses of aspartic and glutamic acids in the planktic foraminifera, Neogloboquadrina pachyderma, and the benthic species, Cibicidoides wuellerstorfi, collected from sediment cores from the Arctic Ocean. The cores were retrieved at various deep-sea sites of the Arctic, which cover a large geographical area from the Greenland and Iceland seas to the Alpha and Lomonosov Ridges in the central Arctic Ocean. Age models for the investigated sediments were developed by multiple dating and correlation techniques, including oxygen isotope stratigraphy, magnetostratigraphy, bio-, litho-, and cyclostratigraphy. The extent of racemization (D/L values) was determined on 95 samples (1028 subsamples) and shows a progressive increase downcore for both foraminifera species. Differences in the rates of racemization between the species were established by analysing specimens of both species from the same stratigraphic levels (n = 21). Aspartic acid and glutamic acid racemize on average  $16\pm2$  % and  $23\pm3$  % faster, respectively, in C. wuellerstorfi than in N. pachyderma. D/L values increase with sample age in nearly all cases, with a trend that follows a simple power function. Scatter around least square regression fits are larger for samples from the central Arctic Ocean than for those from the Nordic Seas. Calibrating the rate of racemization in C. wuellerstorfi using independently dated samples from the Greenland and Iceland seas for the past 400 ka enables estimation of sample ages from the central Arctic Ocean, where bottom water temperatures are presently relatively similar. The resulting ages are older than expected when considering the existing age models for the central Arctic Ocean cores. These results confirm that the differences are not due to taxonomic effects on AAR and further warrant a critical evaluation of existing Arctic Ocean age models. A better understanding of temperature histories at the investigated sites, and other environmental factors that may influence racemization rates in central Arctic Ocean sediments is also needed.

<sup>&</sup>lt;sup>1</sup> Department of Geological Sciences, Stockholm University, Stockholm, 10691, Sweden

<sup>&</sup>lt;sup>2</sup> Bolin Centre for Climate Research, Stockholm University, Stockholm, 10691, Sweden

<sup>&</sup>lt;sup>3</sup> School of Earth and Sustainability, Northern Arizona University, Flagstaff, AZ 86011, USA

#### 40 1. Introduction

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The first application of amino acid geochronology to Arctic Ocean sediments analysed the extent of epimerization in the protein amino acid, isoleucine, over time in samples of the planktic foraminifera, *Neogloboquadrina pachyderma* and the benthic species, *Cibicidoides wuellerstorfi* (Sejrup et al., 1984). Not only did this study provide some of the first amino acid racemization (AAR) data from a polar environment, but it also exposed crucial chronological issues associated with Arctic Ocean sediments. The results contradicted available age interpretations obtained from palaeomagnetic data (Sejrup et al., 1984; Backman, 2004). The problems of dating Pleistocene Arctic marine sediments continue to exist today and are well known (e.g. Alexanderson et al., 2014). Over the past few decades, amino acid geochronology received limited attention in the Arctic, but several studies provided promising results (Sejrup & Haugen, 1992; Kaufman et al., 2008, 2013) that highlighted its potential as a dating technique, and the need for its continued development in Arctic settings. This is particularly desirable, since theoretically, it could provide age control up to a few million years, using even limited amounts of calcium carbonate.

N. pachyderma and C. wuellerstorfi are commonly used in stable isotope stratigraphy and paleoceanographic reconstructions (e.g. Shackleton et al., 2003) and are associated with cold water masses. N. pachyderma is considered to be a primarily 'high-latitude' species (Darling et al., 2017), and C. wuellerstorfi is thought to show strong preference for bottom waters below ~5°C (Rasmussen and Thomsen, 2017). These characteristics and their frequent occurrence in sediment cores from the Arctic Ocean make them particularly useful for amino acid geochronology studies in this region.

The rates of racemization for aspartic and glutamic acids were previously calibrated in N. pachyderma from central Arctic Ocean samples by Kaufman et al. (2008) for the past 150 ka. The calibration relied on the established age models of sediments from the Lomonosov Ridge (O'Regan et al., 2008). Subsequently, however, the extent of racemization in these samples was shown to be higher than expected when compared with those of similar ages from other cold bottom water sites from the Atlantic and Pacific oceans (Kaufman et al., 2013), despite the cold bottom water in the central Arctic Ocean. The reason for this apparently higher extent of racemization in N. pachyderma from central Arctic Ocean samples is unclear, but not considered to be caused by taxonomic effects, since the rate of racemization observed in this species is lower than in other taxa (Kaufman et al., 2013). Either the established ages that were used to calibrate the rate of racemization in the central Arctic Ocean sediments are too young, such that units currently correlated with substages of marine oxygen isotope stage (MIS) 5 instead represent MIS 9, 7 and 5, or other undetermined processes influence protein degradation and preservation in central Arctic Ocean foraminifera. At the Yermak Plateau in the eastern Arctic Ocean (Fig. 1), racemization rates for N. pachyderma generally conform to the rates determined for other cold bottom water sites (West et al., 2019), further challenging the established ages previously used to calibrate the rate of AAR from the Lomonosov Ridge in the central Arctic Ocean. However, it is unknown how racemization progresses in other regions of the Arctic Ocean.

If the apparently higher extent of racemization in *N. pachyderma* from the central Arctic Ocean is not the result of taxonomic effects, a higher rate of racemization can also be anticipated in other taxa from the area, e.g. in *Cibicidoides wuellerstorfi*. However, little is known about racemization of amino acids in this species. The earliest studies involving *C. wuellerstorfi* investigated taxonomical applications (Haugen et al., 1989), or focused on the epimerization of isoleucine (Sejrup et al., 1984; Sejrup and Haugen, 1992) utilising HPLC ion exchange analysers. Since the publication of these seminal papers, analysis of amino acids has become significantly faster, with reduced sample mass requirements, due to improvements in analytical methods (Kaufman & Manley, 1998), yet no studies have addressed amino acid racemization in *C. wuellerstorfi* despite its palaeoceanographical importance (e.g. Yu & Elderfield, 2008; Wollenburg et al., 2015; Burkett et al., 2016; Raitzsch et al., 2020), and the relatively faster and easier sample processing offered by its larger tests (up to ~4-5 times the size) when compared to *N. pachyderma*.

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Here we report the results of aspartic acid and glutamic acid racemization analyses of *Neogloboquadrina* pachyderma and *Cibicidoides wuellerstorfi* obtained from well-dated Quaternary deep-sea sediment cores from the Greenland and Iceland seas, and from sediment cores from the central Arctic Ocean, where sediment ages continue to be debated (Purcell et al., 2022). The long-term rates of racemization in the two species are compared, and the relationship between the extent of racemization and sample age is investigated in both species.

#### 2. Materials and methods

#### a) Investigated sediment cores

Foraminifera samples were taken from sediment cores from the Greenland and Iceland seas, the Lomonosov Ridge, and the Alpha Ridge (Fig. 1). The Greenland and Iceland seas, although part of the Arctic Ocean, are under the direct influence of Atlantic surface waters, and thus are predominantly characterised by open water conditions, unlike the sea ice covered areas of the Lomonosov and Alpha Ridges. The studied sediment cores (Table 1) were collected from deep water (811 – 2952 m) environments, which presently experience similar, very cold (<0 °C), and relatively stable bottom water temperatures.

Region	Core	Latitude (°)	Longitude (°)	Water depth (m)	Bottom water temperature est. (°C)	Age reference
Alpha Ridge	AO16-9-PC1	85.95570	-148.32580	2212	-0.4	Cronin et al. (2019)
Lomonosov Ridge	AO16-5-PC1	89.07800	-130.54700	1253	-0.3	ACEX age model*
Lomonosov Ridge	LOMROG07-PC04	86.70117	-53.76720	811	0.1	Hanslik et al. (2013)
Lomonosov Ridge	LOMROG12-PC03	87.72470	-54.42528	1607	-0.4	O'Regan et al. (2020)
Lomonosov Ridge	LOMROG12-PC07	88.19760	-55.68450	2952	-0.8	ACEX age model*
Lomonosov Ridge	LOMROG12-PC09	89.02672	-73.73444	1318	-0.3	ACEX age model*
Iceland Sea	ODP 151/907A	69.24982	12.69823	1801	-0.8	Jansen et al. (2000a, b)
Greenland Sea	PS17 / 1906-2	76.84630	-2.15050	2901	-1.0	Bauch (2002, 2013)

Table 1: Sediment cores investigated in this study. Current bottom water temperatures were approximated by using annual mean temperature observations from the nearest location from the World Ocean Atlas (Locarnini et al., 2018). \*The ACEX age model is based on Backman et al. (2008), Frank et al. (2008), and O'Regan et al. (2008).

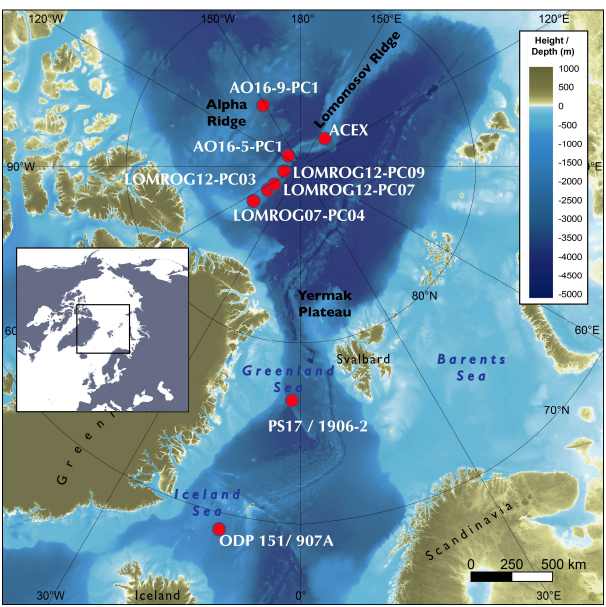


Figure 1: Location of sediment cores referred to in this study. Basemap: Jakobsson et al. (2020).

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Age-depth models for the investigated sediment cores have been developed using a variety of dating techniques. Cores from the Nordic Seas (ODP151/907A and PS17/1906-2) primarily relied on oxygen isotope stratigraphy, complemented by magnetostratigraphy in the case of ODP151/907A (Jansen et al., 2000a, b), and by carbon isotope stratigraphy for core PS17/1906-2 (Bauch, 2002, 2013). The age-depth model of the latter is less certain beyond marine isotope stage (MIS) 6 (Bauch, 2013), due to the large uncertainty associated with isotope stratigraphy, a characteristic issue of Arctic Ocean records.

The age-depth models of sediment cores from the central Arctic Ocean utilise a more diverse toolset, reflecting the difficulties of dating Arctic marine sediments, and heavily depend on a combination of bio- and lithostratigraphy (e.g. Cronin et al., 2019). The lithostratigraphy of the central Arctic Ocean cores investigated in this study can be correlated with that of the Integrated Ocean Drilling Program Expedition 302, the Arctic Coring Expedition (ACEX). This correlation is most apparent in bulk density profiles (Fig. 2), but other sedimentological properties including grain size and a variety of XRF-scanning properties also correlate coherently among cores (O'Regan et al., 2019). The currently accepted age model for the ACEX sedimentary sequence was developed using cyclostratigraphic analysis (O'Regan et al., 2008) and produced similar estimated Quaternary sedimentation rates as obtained by the decay of beryllium isotopes (Frank et al., 2008). The late Quaternary chronology (MIS 1 – 6) for ACEX included constraints from <sup>14</sup>C dating, the correlation with nearby records AO96/12-1PC (Jakobsson et al., 2001) and PS2185 (Spielhagen et al., 2004), where MIS 5 was identified based on the occurrence of the calcareous nannofossil *Emiliania huxleyi* (Jakobsson et al., 2001), and further supported by results from optically stimulated luminescence dating of quartz grains (Jakobsson et al., 2003). The age model of core LOMROG07-PC04 is based on correlation with PS2185 (Hanslik et al., 2013).

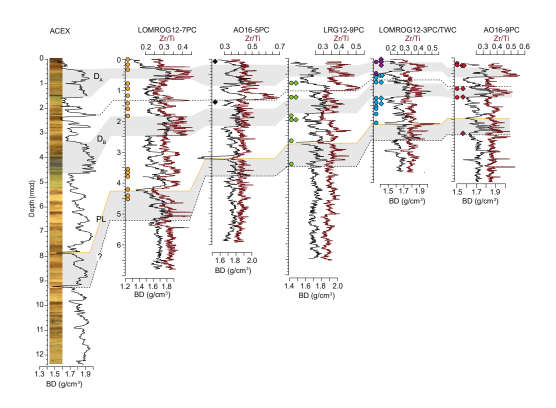


Figure 2: Bulk density (BD) (black line) and XRF-scanning Zr/Ti (red line) profiles of sediment cores from the central Arctic Ocean, with correlations among cores and with the ACEX composite section. D<sub>A</sub> and D<sub>B</sub> are prominent diamict units found in all the cores, while PL is a characteristic fine-grained peach coloured layer. Locations of *N. pachyderma* (circles) and *C. wuellerstorfi* (triangles) samples analysed in this study are shown for each core.

#### b) Analytical procedures

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Sediment samples were wet sieved (63  $\mu$ m), and air dried prior to picking foraminifera tests of *N. pachyderma* and *C. wuellerstorfi*. Initially, the >250  $\mu$ m fraction was targeted to isolate the largest and best-preserved tests. For some samples this was not possible, and the tests were collected from the 180–250  $\mu$ m fraction instead. The

tests were kept in glass vials, and stored in a refrigerator prior to racemization analyses. A total of 95 stratigraphic depths were sampled, with some depths containing both foraminifera species (n = 21). Each sample was further subsampled – on average with 9.6 N. pachyderma and 11.2 C. wuellerstorfi subsamples per sample, producing 1009 analysed subsamples. Each subsample comprised between 10 and 12 N. pachyderma, or 2 and 4 C. wuellerstorfi tests.

The analytical procedures followed that of previous analyses as described in detail by Kaufman et al. (2013) and West et al. (2019), and were performed at the Amino Acid Geochronology Laboratory (AAGL), Northern Arizona University. The foraminifera tests were first sonicated (1-30 s) to remove any loose sediment particles, treated with 1 mL hydrogen peroxide (3%) to remove surficial organic matter, and then rinsed three times with reagent grade (grade I) water. Multiple tests were picked into micro-reaction hydrolysis vials (defining one subsample), and dissolved in 8 μL hydrochloric acid (6 M). The vials were then sealed with nitrogen gas, and the subsamples hydrolysed for 6 hours at 110°C. After the hydrolysis was complete, the subsamples were evaporated in a vacuum desiccator, and then rehydrated in 4 μL of 0.01 M HCl spiked with 10 μM L-homoarginine. Each subsample was injected onto a high-performance liquid chromatograph (HPLC) with a fully automated, reversed phase procedure (Kaufman & Manley, 1998) to separate pairs of D- and L-amino acids. The peak-area ratio of D and L stereoisomers of eight amino acids (aspartic acid, glutamic acid, serine, alanine, valine, phenylalanine, isoleucine and leucine) were analysed to determine the extent of racemization. This study only utilised the racemization results of aspartic acid (Asp) and glutamic (Glu) acid, which are among the two most abundant and chronographically well-resolved amino acids. Asp and Glu reported in this study also include any Asn and Gln present in the biomineral.

#### 3. Results

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## a) Data screening

Initial data screening was based on the procedure of Kosnik & Kaufman (2008). First, subsamples with L-Ser / L-Asp ≥ 0.8 – an indicator of potential contamination by modern amino acids – were excluded. The D/L values of Asp and Glu positively covary in fossil proteins. Subsamples that did not adhere to this expected trend were omitted. Finally, subsamples with D/L Asp or Glu values beyond ±2σ of the sample mean were also removed (Supplementary Fig. S1). As a result of this screening process, 17.9 % of all subsamples were rejected
 (Supplementary Table S1).

Following the subsample screening process, sample means, and related standard deviation values were calculated for Asp and Glu for all samples. Stratigraphically reversed samples (mean D/L values lower than expected for their stratigraphic depths with no overlap within  $1\sigma$  with the sample from shallower depth) were identified within each core (Table 2). These include five samples of *N. pachyderma* and three *C. wuellerstorfi*. Of these, two samples contained sufficient tests of both species to analyse AAR, but only *C. wuellerstorfi* were stratigraphically reversed.

Lab ID (UAL)	Core	Core depth (m)	Age (ka)	nª	Excluded	Excl. ratio (%)	Asp D/L	1 σ	Glu D/L	1 σ
Neoglob	oquadrina pachyderm	ıa		1				<u> </u>		
	Alpha Ridge									
17336	AO16-9-PC1	0.190	16	10	1	10	0.141	0.014	0.069	0.011
17337	AO16-9-PC1	0.210	22	10	1	10	0.210	0.010	0.104	0.005
17338	AO16-9-PC1	0.970	127	8	3	38	0.419	0.020	0.242	0.029
17339	AO16-9-PC1	1.260	188	10	5	50	0.445	0.015	0.254	0.022
	Lomonosov Ridge	200	100				01110	0.0.0	0.20.	0.022
21406	LOMROG12-TWC03	0.095	38	7	1	14	0.193	0.017	0.077	0.010
21407	LOMROG12-TWC03	0.495	60	7	1	14	0.332	0.017	0.168	0.022
21408	LOMROG12-PC03	0.505	61	8	2	25	0.340	0.032	0.163	0.026
21409	LOMROG12-PC03	0.610	76	8	1	13	0.395	0.021	0.206	0.015
21410	LOMROG12-PC03	0.761	108	8	3	38	0.415	0.040	0.212	0.025
21411	LOMROG12-PC03	1.291	204	8	2	25	0.468	0.015	0.240	0.010
21412	LOMROG12-PC03	1.401	236	6	2	33	0.443	0.013	0.239	0.016
21413	LOMROG12-PC03	1.551		6	2	33	0.443	0.021	0.239	0.020
21413	LOMROG12-PC03	1.601	290	8	3	33	0.400	0.066	0.242	0.070
21414	LOMROG12-PC03	1.781	311	5	0	0	0.417	0.032	0.185	0.026
21415	LOMROG12-PC03	2.101	391	8	5	63	0.377	0.060	0.191	0.048
	LOMROG12-PC03		489							
22759		0.025	MIS 1-3? (~19)	17	4	24	0.102	0.018	0.044	0.011
21587	LOMROG07-PC04	0.040	MIS 1-3? (~39)	8	3	38	0.097	0.006	0.036	0.004
21588	LOMROG07-PC04	0.140	MIS 5.1? (~77)	8	3	38	0.278	0.034	0.114	0.021
22760	LOMROG07-PC04	0.225	MIS 5.1? (~82)	11	1	9	0.278	0.008	0.123	0.009
22761	LOMROG07-PC04	1.025	MIS 5.5? (~115)	13	2	15	0.320	0.011	0.151	0.012
21589	LOMROG07-PC04	1.110	MIS 5.5? (~123)	8	4	50	0.347	0.029	0.146	0.022
22762	LOMROG07-PC04	1.385	> MIS 7? (~405)	13	2	15	0.387	0.029	0.195	0.032
<sup>b</sup> 21590	LOMROG07-PC04	1.410	MIS 1-3? (~12.8)	8	3	38	0.318	0.035	0.137	0.035
21591	LOMROG07-PC04	1.720	MIS 1-3? (~40)	8	3	38	0.391	0.036	0.175	0.043
22763	LOMROG07-PC04	1.785	MIS 5.1? (~74)	12	4	33	0.386	0.023	0.187	0.016
21592	LOMROG07-PC04	2.010	MIS 5.1? (~78)	8	7	88	0.357		0.140	
22764	LOMROG07-PC04	2.025	MIS 5.5? (~115)	9	2	22	0.417	0.069	0.230	0.066
<sup>b</sup> 21594	LOMROG07-PC04	3.000	MIS 5.5? (~123)	7	4	57	0.270	0.099	0.136	0.057
15866	LOMROG12-PC07	0.020	9	9	2	22	0.081	0.004	0.034	0.002
15867	LOMROG12-PC07	0.165	31	10	1	10	0.205	0.036	0.090	0.022
15868	LOMROG12-PC07	0.315	40	10	2	20	0.272	0.025	0.124	0.013
15869	LOMROG12-PC07	0.780	52	9	1	11	0.294	0.026	0.122	0.019
15870	LOMROG12-PC07	0.980	63	6	1	17	0.354	0.024	0.163	0.022
15871	LOMROG12-PC07	1.170	79	9	3	33	0.373	0.029	0.189	0.028
15872	LOMROG12-PC07	1.425	96	9	3	33	0.399	0.035	0.183	0.027
15873	LOMROG12-PC07	1.580	111	9	2	22	0.368	0.030	0.168	0.024
15874	LOMROG12-PC07	1.720	124	9	3	33	0.385	0.024	0.173	0.025
b15875	LOMROG12-PC07	3.537	350	9	3	33	0.305	0.036	0.133	0.018
15876	LOMROG12-PC07	3.657	374	9	0	0	0.354	0.048	0.185	0.031
15877	LOMROG12-PC07	3.787	402	4	3	75	0.348		0.174	
15878	LOMROG12-PC07	4.210	485	6	3	50	0.402	0.012	0.193	0.011
<sup>b</sup> 15879	LOMROG12-PC07	4.405	502	7	2	29	0.335	0.014	0.155	0.012
15880	LOMROG12-PC07	4.510	510	5	3	60	0.307	0.029	0.145	0.034
22765	LOMROG12-PC09	0.800	55	17	2	12	0.379	0.012	0.197	0.010
22766	LOMROG12-PC09	1.250	129	10	0	0	0.400	0.029	0.209	0.030
22767	LOMROG12-PC09	1.840	205	12	2	17	0.416	0.023	0.207	0.027
22768	LOMROG12-PC09	1.980	233	17	3	18	0.435	0.019	0.232	0.024
22769	LOMROG12-PC09	2.160	276	13	1	8	0.475	0.053	0.278	0.040
b22770	LOMROG12-PC09	3.400	603	11	4	36	0.407	0.005	0.203	0.006
	Iceland Sea									
22745	ODP 151/ 907A	1.050	42	14	2	14	0.200	0.008	0.079	0.006
				40		15		0.005		0.003
22746	ODP 151/ 907A	1.870	107	13	2	15	0.281	0.005	0.114	0.003

22748	ODP 151/ 907A	3.060	160	15	0	0	0.355	0.011	0.171	0.009
22749	ODP 151/ 907A	5.060	168	17	2	12	0.335	0.011	0.171	0.009
22749	ODP 151/ 907A	6.810	309	12	1	8	0.393	0.018	0.192	0.014
22750	ODP 151/ 907A	15.210	398	10	3	30	0.388	0.013	0.163	0.016
22/31	Greenland Sea	13.210	781	10	3	30	0.462	0.030	0.230	0.040
22732	PS17 / 1906-2	0.150	40	10	2	OF.	0.129	0.003	0.067	0.004
			10	12 17	3	25				
22733	PS17 / 1906-2	1.805	85		0	0	0.263	0.008	0.111	0.008
22734	PS17 / 1906-2	2.005	111	12	0	0	0.306	0.006	0.138	0.005
22735	PS17 / 1906-2	2.105	117	12	4	33	0.312	0.010	0.141	0.009
22736	PS17 / 1906-2	2.205	122	12	3	25	0.347	0.011	0.169	0.012
22737	PS17 / 1906-2	3.290	207	8	1	13	0.343	0.011	0.171	0.008
22738	PS17 / 1906-2	3.605	225	12	2	17	0.355	0.010	0.178	0.010
22739	PS17 / 1906-2	5.505	398	5	1	20	0.439	0.030	0.227	0.036
Cibicido	ides wuellerstorfi						1	Г	ı	
1=011	Alpha Ridge	0.400		4.0			0.00=	2 2 2 4		0.040
17341	AO16-9-PC1	0.190	16	10	0	0	0.235	0.061	0.099	0.040
17342	AO16-9-PC1	0.210	22	10	1	10	0.326	0.027	0.158	0.020
17343	AO16-9-PC1	0.970	127	10	0	0	0.502	0.005	0.311	0.010
17344	AO16-9-PC1	1.260	188	10	0	0	0.522	0.007	0.321	0.008
17346	AO16-9-PC1	2.415	599	10	2	20	0.523	0.101	0.302	0.094
	Lomonosov Ridge									
17331	AO16-5-PC1	0.090	30	10	1	10	0.342	0.012	0.155	0.006
17333	AO16-5-PC1	1.370	105	7	1	14	0.525	0.020	0.339	0.032
22752	LOMROG12-TWC03	0.025	16	13	1	8	0.144	0.012	0.047	0.006
22753	LOMROG12-TWC03	0.095	38	18	3	17	0.157	0.014	0.053	0.006
17327	LOMROG12-TWC03	0.215	44	8	0	0	0.341	0.006	0.161	0.006
22754	LOMROG12-PC03	0.520	66	13	3	23	0.412	0.014	0.208	0.013
17328	LOMROG12-PC03	0.540	98	8	0	0	0.433	0.009	0.234	0.009
17329	LOMROG12-PC03	0.741	198	8	1	13	0.485	0.008	0.284	0.008
17330	LOMROG12-PC03	1.271	63	8	1	13	0.541	0.010	0.328	0.019
<sup>b</sup> 22755	LOMROG12-PC03	1.441	247	9	1	11	0.491	0.017	0.267	0.035
22756	LOMROG12-PC09	0.800	73	12	1	8	0.433	0.011	0.236	0.011
22757	LOMROG12-PC09	1.250	140	22	2	9	0.476	0.007	0.266	0.011
22758	LOMROG12-PC09	1.980	224	5	0	0	0.508	0.015	0.281	0.015
	Iceland Sea									
22740	ODP 151/ 907A	1.050	42	9	3	33	0.166	0.026	0.059	0.013
22741	ODP 151/ 907A	1.870	107	21	1	5	0.337	0.015	0.151	0.010
<sup>b</sup> 22742	ODP 151/ 907A	2.350	131	15	6	40	0.160	0.013	0.064	0.006
22743	ODP 151/ 907A	3.060	168	10	1	10	0.388	0.025	0.198	0.026
22744	ODP 151/ 907A	15.210	781	18	1	6	0.549	0.016	0.345	0.032
	Greenland Sea									
22724	PS17 / 1906-2	0.150	10	15	0	0	0.175	0.009	0.059	0.004
22725	PS17 / 1906-2	1.805	85	18	3	17	0.328	0.010	0.144	0.007
22726	PS17 / 1906-2	2.005	111	20	1	5	0.351	0.016	0.160	0.007
22727		2.105	117	18	1	6	0.357	0.009	0.171	0.007
	PS17 / 1906-2	2.100								
22728	PS17 / 1906-2 PS17 / 1906-2	2.205	122	21	1	5	0.379	0.016	0.191	0.015
			122 207	21 6	1	5 17	0.379	0.016 0.050	0.191 0.258	0.015
22728	PS17 / 1906-2	2.205								

Table 2: Extent of racemization (D/L) for aspartic acid (Asp) and glutamic acid (Glu) in samples of *Neogloboquadrina* pachyderma and *Cibicidoides wuellerstorfi* from Arctic Ocean sediment cores. Reported ages are from published age models (Table 1). 190

 $<sup>^{\</sup>rm a}$  Number of subsamples used to calculate the mean and standard deviation  $^{\rm b}$  Stratigraphically reversed sample

## b) Inter-species comparison

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The overall subsample rejection rate was higher for *N. pachyderma* (22.7 %) than for *C. wuellerstorfi* (9.9 %) (Table 3). Only one *C. wuellerstorfi* subsample was rejected due to high serine content (L-Ser / L-Asp  $\geq$  0.8), significantly fewer than those for *N. pachyderma*, implying that secondary amino acids (contamination) were not introduced during core storage or laboratory analysis because both species were treated similarly.

Species	Total number	L-Ser / L-Asp	Non-covarying D/L Asp	D/L Asp or D/L Glu not within		
	of subsamples	≥ 0.8	and D/L Glu	±2σ of sample mean		
N. pachyderma	635	81	20	43		
C. wuellerstorfi	374	1	16	20		

Table 3: Number of rejected subsamples per rejection criterion.

The proportion of samples with high intra-sample variability (coefficient of variation for D/L Asp and Glu > 10 %, following data screening) was approximately twice as high for *N. pachyderma* as for *C. wuellerstorfi*.

The rate of amino acid racemization varies between different foraminifera species (King & Neville, 1977). As some samples (n = 21) from certain stratigraphic intervals contained specimens of both N. pachyderma and C. wuellerstorfi, this allowed a direct comparison of the rates of racemization in the two species (Fig. 3). One stratigraphically reversed sample was omitted prior to analysis. The slope of the least-squares regression fit to the D/L versus D/L data indicates that Asp racemized, on average,  $15\pm2$  % (n = 19) faster in C. wuellerstorfi than in N. pachyderma, and similarly, Glu also racemized faster ( $23\pm2$  %) in C. wuellerstorfi than in N. pachyderma.

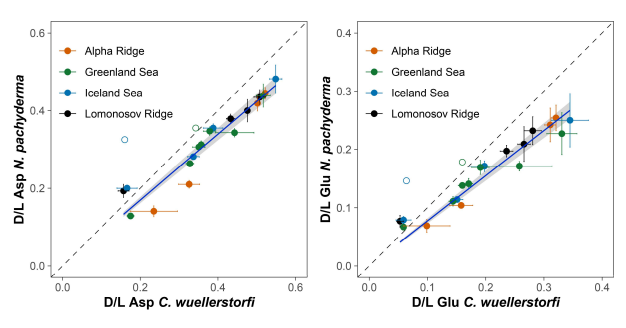


Figure 3: Extent of racemization for aspartic acid (Asp) and glutamic acid (Glu) in *Neogloboquadrina pachyderma* and *Cibicidoides wuellerstorfi* from samples from the same stratigraphic depths collected from different regions of the Arctic Ocean. Error bars represent  $\pm 1\sigma$  intra-sample variability, solid line is least squares regression with 95%

## c) Relationship between D/L values and age / stratigraphic depth

The extent of racemization (D/L values) for Asp and Glu show a systematic increasing trend with increasing age of samples of both species from the Greenland and Iceland seas, and follow a simple power function (Fig. 4). A simple power law was used to model the forward rate of racemization in this study, as this is typical for racemization of amino acids in biominerals held under isothermal conditions (e.g. Kaufman, 2006; Clarke and Murray-Wallace, 2006).

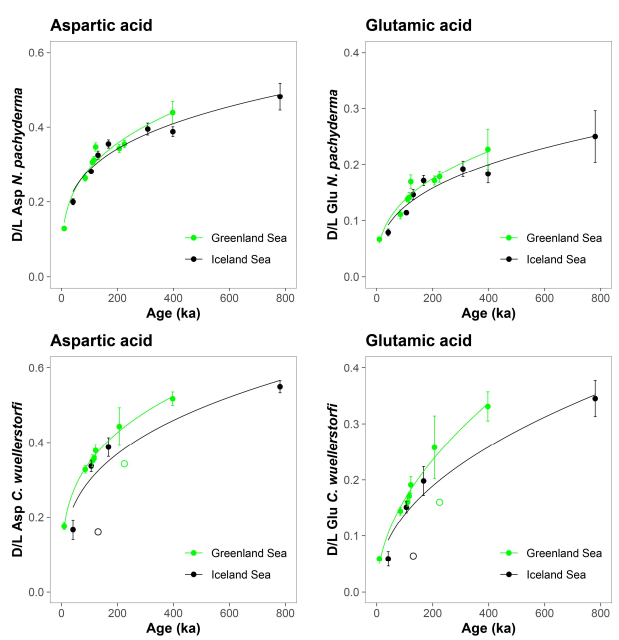


Figure 4: Extent of racemization for aspartic acid (Asp) and glutamic acid (Glu) in samples of *N. pachyderma* and *C. wuellerstorfi* from sediment cores PS17/1906-2 (Greenland Sea) and ODP 151/907A (Iceland Sea). Error bars represent  $\pm 1\sigma$  intra-sample variability and curves are power functions. Stratigraphically reversed samples are marked with unfilled symbols and were excluded to improve the goodness of fit.

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D/L values obtained from sediment cores from the central Arctic Ocean, where age-depth models are less certain, are displayed on the ACEX composite depth scale based on correlations using bulk density profiles (Fig. 5). The extent of racemization follows a simple power function (stratigraphically reversed samples excluded), as also observed in samples from the Nordic Seas, and D/L values generally overlap between correlative samples from multiple cores. This directly supports the accuracy of the lithostratigraphic correlations established among the sites. D/L values from *N. pachyderma* appear to reach a plateau at ~0.4 for Asp and ~0.2 for Glu (~6 m depth in the ACEX record). Plateauing of D/L Asp values was previously documented in other Arctic Ocean cores as well (Kaufman et al., 2008). It is unclear whether plateauing is present in *C. wuellerstorfi* samples, as only a few samples were analysed in the corresponding age range, and the oldest sample has the largest intrasample variability.

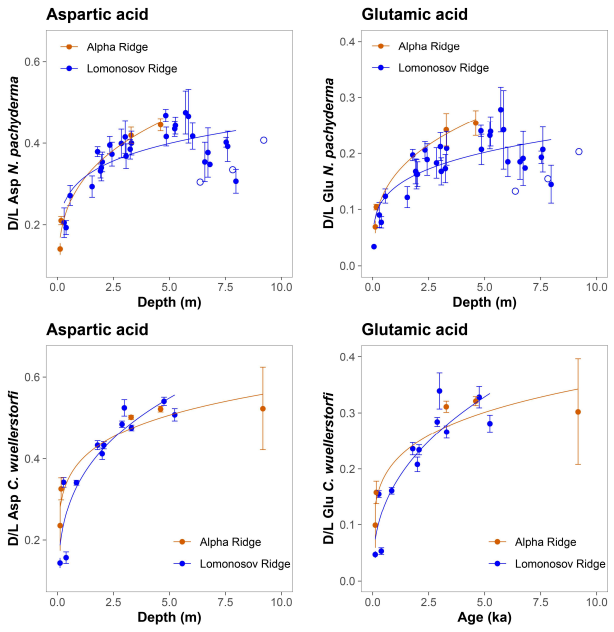


Figure 5: Extent of racemization for aspartic acid (Asp) and glutamic acid (Glu) in *N. pachyderma* and *C. wuellerstorfi* from the central Arctic Ocean displayed on the ACEX composite depth. Sediment cores investigated in

this study were grouped based on their geographical location (Alpha Ridge – red, Lomonosov Ridge – blue). Error bars are  $\pm$  1 $\sigma$  intra-sample variability, curves are power functions and unfilled symbols mark stratigraphically reversed samples.

## 4. Discussion

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Aspartic acid and glutamic acid racemization analyses of multiple foraminifera taxa obtained from cold bottom water sites across the globe showed that sample ages can be confidently estimated by calibrated age equations, which relate the extent of racemization of the amino acids to independently determined sample age (Kaufman et al., 2013). These globally derived age equations are based on both planktic and benthic foraminifera species with *N. pachyderma* contributing up to ~21% of all samples, but *C. wuellerstorfi* is not represented. Ages derived using the global age equations generally agree with independently derived ages at the Yermak Plateau (West et al., 2019), but do not agree with previously published age models from the central Arctic Ocean, beyond about 40 ka. If existing age models in the central Arctic Ocean are correct, the extent of racemization for *N. pachyderma* is higher than expected when compared with other oceans (Kaufman et al., 2008, 2013).

The results of AAR presented here show that Asp and Glu racemize in a predictable manner in both *N. pachyderma* and *C. wuellerstorfi* samples from the Nordic Seas (Fig. 4), although the rates differ systematically between the species (Fig. 3). Both D/L Asp and D/L Glu values obtained from *N. pachyderma* samples from the Nordic Seas clearly follow a trend previously observed at the Yermak Plateau (Fig. 6), implying that racemization kinetics are indistinguishable in these areas. These data from independently dated cores from coldwater sites provide further support for the integrity of the AAR technique, and for the globally calibrated age equations of Kaufman et al. (2013).

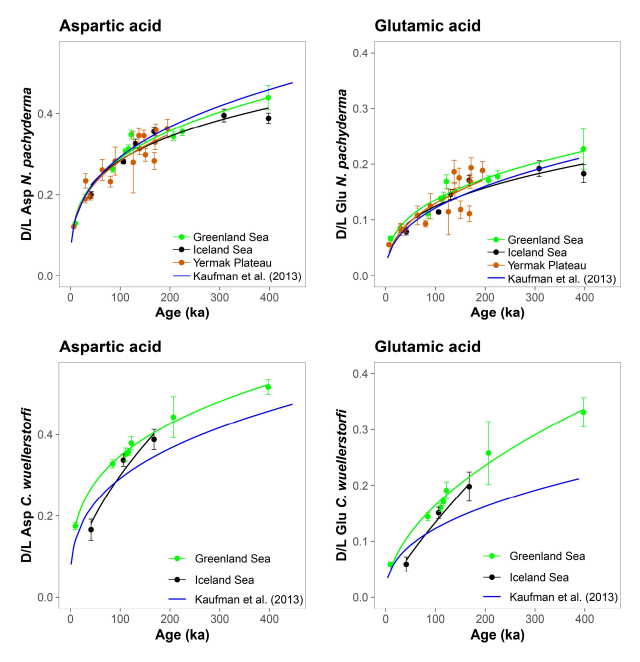


Figure 6: Extent of racemization in aspartic acid (Asp) and glutamic acid (Glu) in *Neogloboquadrina pachyderma* and *Cibicidoides wuellerstorfi* from the Greenland and Iceland seas and the Yermak Plateau. Black, green and red curves are power functions fit to the data. Blue curves are the globally calibrated age equations of Kaufman et al. (2013). Data for the Yermak Plateau from West et al. (2019). Error bars represent  $\pm 1\sigma$  intra-sample variability.

The inter-species comparisons show that Asp racemizes approximately 16 % faster in *C. wuellerstorfi* than in *N. pachyderma* (Fig. 3). Differences of similar magnitude were previously observed between *N. pachyderma* and *Pulleniatina obliquiloculata*, in which Asp racemizes 12-16 % faster than in *N. pachyderma* (Kaufman et al., 2013). Quantifying the relative differences of the racemization rates between species can facilitate future AAR studies. When a taxon is unavailable at a certain stratigraphic depth, the extent of racemization in one species could be adjusted to that of another based on the difference in the observed rate of racemization. *C. wuellerstorfi* was not represented in the globally calibrated age equations and the higher rate of AAR in this species suggests that D/L values should not be used in combination with the equations without adjustment (Fig. 6).

The purportedly higher rates of racemization in *N. pachyderma* from the central Arctic Ocean were argued to be caused by factors other than taxonomic effects (Kaufman et al., 2013). Either racemization generally progresses at a higher rate here due to an unknown reason, or existing age models used to constrain the rate of AAR from the central Arctic Ocean underestimate the true age of deposition. Given the existing age models, higher-than-expected rates of racemization are also observed in this study in both *N. pachyderma* and *C. wuellerstorfi* samples from the central Arctic Ocean (Fig. 7). Differences between ages predicted by the global calibrated age equation and currently available sample ages based on the ACEX age model are larger for *C. wuellerstorfi*, as expected, since racemization proceeds faster in this taxon than in *N. pachyderma*.



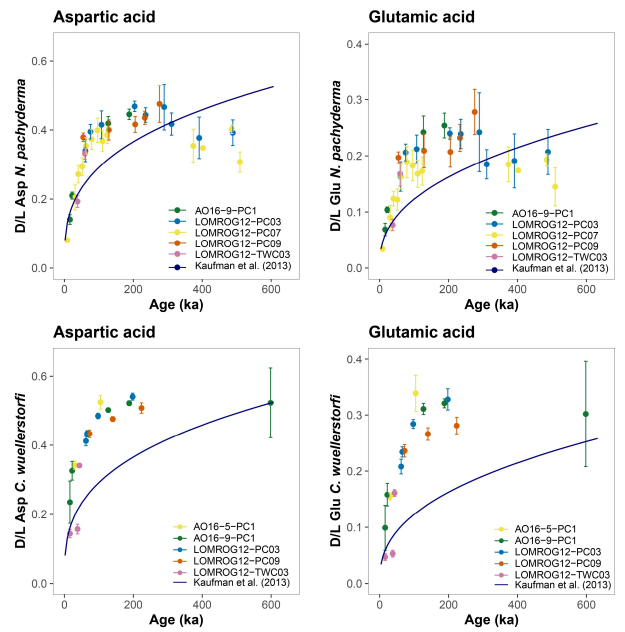


Figure 7: Extent of racemization for aspartic acid (Asp) and glutamic acid (Glu) in *N. pachyderma* and *C. wuellerstorfi* from the central Arctic Ocean. D/L values are displayed against the ACEX age model. D/L values versus

age trend defined by the globally calibrated age equations (Kaufman et al., 2013) is shown in blue. Error bars represent  $\pm 1\sigma$  intra-sample variability.

The AAR trends in *C. wuellerstorfi* samples from the Nordic Seas, where age models are better constrained, can also be compared with those in the central Arctic Ocean samples. The racemization of Asp and Glu in *C. wuellerstorfi* samples from both the Greenland and Iceland seas (GIS) follow the same trend (Fig. 4), which can be approximated with regression analysis. Following the removal of stratigraphically reversed samples (n = 3) and samples with high analytical uncertainty (coefficient of variation > 10 %, n = 5), simple power functions (Eq. 1 and 2)

$$t = 3827.6 * (D/L Asp)^{3.395}$$
 (1)

$$t = 4979.1 * (D/L Glu)^{2.138}$$
 (2)

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(where t = age in ka) fit the D/L Asp and D/L Glu versus age relationship well for the past 400 ka (the period for which these models are robust). Applying these equations to D/L values in *C. wuellerstorfi* from the central Arctic Ocean reveals that sample ages are younger than predicted by the model based on the Nordic Sea samples (Fig. 8). The higher-than-expected D/L values observed for *C. wuellerstorfi* from the central Arctic Ocean confirm earlier findings, which argued that higher racemization rates in the central Arctic Ocean are not the result of taxonomic effects (Kaufman et al., 2013).

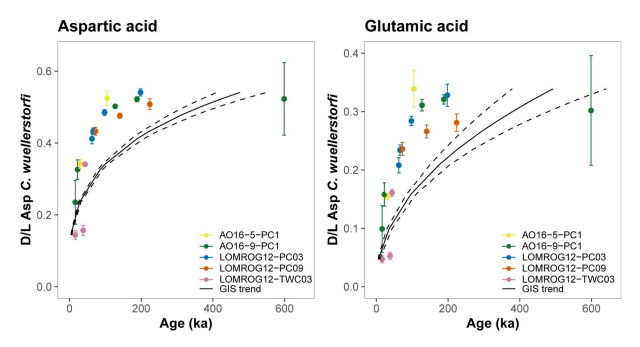


Figure 8. Aspartic acid (Asp) and glutamic acid (Glu) racemization versus sample age in *C. wuellerstorfi* from the central Arctic Ocean. Black curves are the age equations determined by samples from the Greenland and Iceland seas (GIS trend); black dashed curves mark 95% confidence intervals. Error bars represent ± 1σ intra-sample variability.

The discrepancy between the globally (*N. pachyderma*) and GIS (*C. wuellerstorfi*) calibrated AAR ages and the established chronology for central Arctic Ocean sediments is illustrated in Figure 9 and Table 4. The calibrated

AAR ages suggest that intervals previously interpreted as substages in MIS 5 are instead separate interglacial periods extending back to MIS 9. This interpretation would indicate that the diamict unit previously assigned to MIS 6, and representing the onset of a fundamentally different depositional regime in the Arctic characterised by recurrent coarse-grained facies (O'Regan et al., 2010), is likely older than MIS 8 and possibly as old as MIS 12. Exact age determinations are difficult due to a paucity of data from these lower depths, and increased downcore scatter in the stacked AAR results (*discussed below*).

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It cannot be excluded that the established chronologies of the central Arctic Ocean sediments underestimate their true ages, but significant shifts (representing multiple glacial cycles) in sediment ages are required. It is important to recognise that this results in a number of fundamental inconsistencies when compared with other geochronological data. For example, recent work (O'Regan et al., 2020) placed the first occurrence of the coccolithophore, Emiliania huxleyi, in core LOMROG12-PC03 at 1.39±0.02 mbsf in MIS 7 (~220 ka). The AAR age equations (1) and (2), defined by the trend in C. wuellerstorfi samples from the Greenland and Iceland seas, yield estimated ages of 475±12 ka (D/L Asp) or 459±21 ka (D/L Glu) (MIS 12/13) based on the mean D/L ±1SE for a sample from 1.27 m in this core. Similarly, the globally calibrated age equation (Kaufman et al., 2013) returns an MIS 12 age of 427±20 ka for N. pachyderma samples from 1.29 m in LOMROG12-PC03. These ages are substantially older than the evolutionary occurrence of E. huxleyi during MIS 8 (Thierstein et al., 1977; Anthonissen and Ogg, 2012). Furthermore, E. huxleyi is abundant in the 0.58-0.82 mbsf core interval of LOMROG12-PC03, which was assigned an MIS 5 age (O'Regan et al., 2020). In contrast, the AAR trend from the Greenland and Iceland seas assigns ages of 189±6 to 328±7 ka (MIS 6-9) for the 0.52-0.74 m core depths (Table 4). This could only be reconciled with the existence of E. huxleyi in this interval, if it had entered the Arctic shortly after its first evolutionary occurrence. However, the MIS 6-9 age is not consistent with results from optically stimulated luminescence dating in stratigraphically coeval sediments from another core on the Lomonosov Ridge, which support the MIS 5 age assignment (Jakobsson et al., 2003).

Stratigraphic depth (m)	ACEX equivalent depth (m)	Marine Isotope Stage (MIS) correlation						
		ACEX age model (Backman et al., 2008; O'Regan et al., 2008) and <i>E. huxleyii</i> occurrences (O'Regan et al., 2020)		AAR geochronology – aspartic acid in C. wuellerstorfi (this study)				
0.52-0.74	1.96-3.35	MIS 5	MIS 6-9	MIS 6-9				
1.27-1.39	4.62-5.44	MIS 7	MIS 9-12	MIS 9-12				

Table 4: Age estimates for selected stratigraphic intervals in core LOMROG12-PC03 based on the ACEX age model and occurrences of *E. huxleyi*, and AAR geochronology.

On the other hand, sedimentation rates estimated using the AAR global and GIS calibration equations are consistent with radiocarbon derived estimates over the past 40 ka, but beyond this point remain intermediate between the proposed age model for the ACEX record (O'Regan et al., 2008) and those recently inferred from <sup>230</sup>Th and <sup>231</sup>Pa extinction ages in sedimentary sequences from this region of the Lomonosov Ridge (Hillaire-Marcel et al., 2017; Purcell et al., 2022). Additional work is required to understand the origin and resolve the differences between these geochronological approaches. However, a critical observation is that even the older estimated ages from the global and GIS calibration equations remain considerably younger than the long-

overturned paleomagnetic-derived age model that would place the Bruhnes-Matuyama boundary (~780 ka) at approximately 5 m depth in the ACEX record (O'Regan et al., 2008) (Fig. 9).

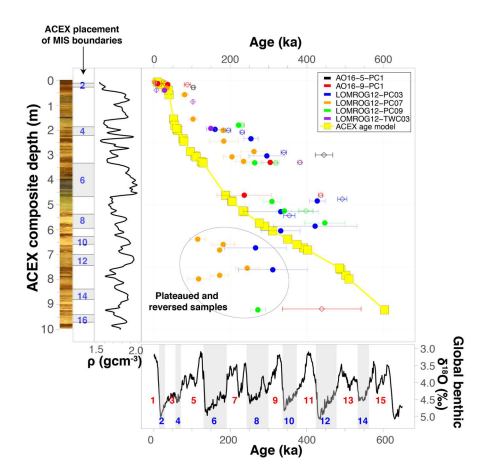


Figure 9: Alternative age–depth relationships in sediment cores from the central Arctic Ocean. Circles and diamonds mark ages estimated by globally calibrated rates of racemization of aspartic acid in N. pachyderma (Kaufman et al., 2013) and GIS-calibrated rates of racemization of aspartic acid in N. wuellerstorfi (this study), respectively. Error bars represent 95 % confidence intervals. The ACEX age model is based on Backman et al. (2008), Frank et al. (2008), and O'Regan et al. (2008). Also shown are ACEX digital core image, marine oxygen isotope stage (MIS) boundaries based on the ACEX age model, bulk density ( $\rho$ ) profile of the ACEX core, and the global benthic  $\delta^{18}$ O record and corresponding MIS 1-15 based on Lisiecki & Raymo (2005).

An alternative to central Arctic Ocean age models underestimating the true age of sediments is that the rate of AAR in foraminifera from the central Arctic Ocean is faster than in the eastern Arctic Ocean (West et al., 2019), the Nordic Seas (this study), and other globally distributed cold water sites (Kaufman et al., 2013). A fundamental premise of this study is that the investigated central Arctic Ocean sites have experienced similar temperatures histories to those other cold water sites. Modern bottom water temperatures near the coring sites are very similar (Table 1), but they may have differed in the past. Cronin et al. (2012) showed that the central Arctic basin waters, at ~1000-2500 m depth interval, were 1-2°C warmer during the past 50–11 ka than today. While this might account for part of the apparently higher rate of AAR in the central Arctic Ocean, West et al. (2019) showed that more substantial differences (~ >4 °C) in effective diagenetic temperatures would be required to achieve the differences observed between D/L values of equivalent age samples from sediment cores from the central Arctic Ocean and those from the Nordic Seas. Available heat flow data (Shephard et al., 2018)

show that geothermal flux is not unusually high in the central Arctic Ocean when compared to the Yermak Plateau, and thus cannot explain the apparently higher rates of racemization. Therefore, while offsets in D/L values in stratigraphically coeval sections of different central Arctic Ocean cores in this study might, in part, be attributed to differences in bottom water temperatures, such differences are unlikely able to explain the overall higher inferred rates of racemization compared to other global sites.

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D/L values in many samples of *N. pachyderma* from the central Arctic Ocean cores exhibit a distinct plateauing below the ~5.5–6 m composite ACEX depth (Fig. 5 and 9). Some of these samples are stratigraphically reversed (i.e. their D/L values are lower than of those from shallower depths). Such plateauing of D/L values was previously observed in cores from other areas of the Arctic Ocean (Kaufman et al., 2008), but its cause remains unclear. While sediment mass movements, glacial erosion and subsequent re-deposition are known to occur across the Arctic Ocean (e.g. Jakobsson & O'Regan, 2016; Boggild et al., 2020; Pérez et al., 2020; Schlager et al., 2021), this explanation would require that almost all of the material above 5.5–6 m composite depth was reworked similarly across multiple cores, which seems highly unlikely.

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Local differences in sedimentation rates might also account for some of the observed scatter in AAR results. For example, there are pronounced regional differences in the thickness of correlative units in the studied cores (Fig. 2), implying highly variable sedimentation rates between the sites. In cores with lower sedimentation rates, the influence of bioturbation could conceivably introduce significant scatter though mixing of individuals of different ages.

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Overall, sedimentation rates in the central Arctic Ocean are much lower in marginal areas closer to the shelves (Backman et al., 2004), and can be greatly reduced during glacials in some regions of the Arctic Ocean (Jakobsson et al., 2014). Conversely, thick diamict units suggest rapid influxes of ice rafted material during some glacials or glacial terminations. These punctuated episodes of sedimentation can not only introduce hiatuses, but also impact the length of time biocarbonates are exposed to microbial activity on the seafloor (Sejrup and Haugen, 1994), which in turn can speed up or slow down organic diagenetic processes that influence the rate of AAR. For example, where sedimentation rates are low, greater microbial activity could increase organic diagenesis and lead to higher apparent rates of AAR. On the other hand, continuous turnover and reworking of microbial necromass within the foraminifera test could instead lower the apparent rate of racemization via regeneration of L-amino acids, and alter the apparent AAR rates, as has been documented for bulk organic matter in marine sediments (Braun et al., 2017). The composition of the microbial community may also have an impact. Kubota et al. (2016) isolated alphaproteobacteria from deep-sea sediments from the Sagami Bay (Japan), which exclusively utilised D amino acids as a carbon and nitrogen source. If such microbes were also present within foraminifera tests of the central Arctic Ocean, they could potentially alter the progress of racemization. Apparent plateauing of D/L values (e.g. Fig. 5, 7, 9) could be associated with these processes, but would require a distinctively different microbial sedimentary environment in the central Arctic Ocean than elsewhere (such as the Yermak Plateau or Norwegian-Greenland Sea, where the global age-equation appears applicable). This is not inconceivable, as recently Yu et al. (2020) suggested that different marine environments could be characterised by distinct bacterial groups that utilise D-amino acids.

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In the future, pre-treating the foraminifera tests with bleach to isolate the intra-crystalline fraction, which approximates a closed system during diagenesis (Penkman et al., 2008; Wheeler et al., 2021), could minimise the influence of bacterial activity on racemization rates. However, recent work (Millman et al., 2022) showed that bleaching does not necessarily improve the quality of AAR results in foraminifera, thus the current study used the standard weak oxidative pre-treatment. Future work should investigate how bleaching impacts AAR results from central Arctic Ocean cores.

If the rate of AAR in foraminifera is indeed higher in the central Arctic Ocean, its exact cause remains unknown. The data reported in this study confirm that it is not caused by taxonomic effects. It is observed not only in the planktic *N. pachyderma* but also in the benthic species *C. wuellerstorfi*. We have highlighted the discrepancies that arise in central Arctic Ocean age models if the global (*N. pachyderma*) and GIS (*C. wuellerstorfi*) AAR age equations are applied. This alternate geochronological interpretation should be considered in future attempts to reconcile the currently disparate results from different dating techniques applied to Arctic Ocean sediments.

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#### 5. Conclusions

- Aspartic acid and glutamic acid racemize faster in *C. wuellerstorfi* than in *N. pachyderma*, and the extent of racemization for these amino acids increases progressively with sample age in both species from multiple sediment cores. Their trends conform to a simple power function.
- *C. wuellerstorfi* samples are characterised by lower intra-sample variability than those of *N. pachyderma*, and this, coupled with a reduced subsample rejection rate and faster sample processing offered by its larger tests make it an appealing target of future AAR studies.

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 Ages of N. pachyderma samples from the Greenland and Iceland seas agree with ages predicted by globally calibrated age equations for aspartic acid and glutamic acids (Kaufman et al., 2013), and confirm their applicability in these polar regions.

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• The rate of racemization for *C. wuellerstorfi* was calibrated for the past 400 ka using samples from the Greenland and Iceland seas. Applying this calibration to the *C. wuellerstorfi* samples from the central Arctic Ocean indicates that they are older than their currently accepted ages. This confirms that higher-than-expected D/L values in *N. pachyderma* from the central Arctic Ocean are not the result of taxonomic effects.

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• We cannot find a clear reason why age calibration equations that work globally in the eastern Arctic Ocean, and in the Nordic Seas cannot be applied in the central Arctic Ocean. As such the older ages predicted by these equations for central Arctic Ocean sediments remain a viable option when comparing and assessing results from different approaches to dating Arctic marine sediments.

 Regardless of the age equation applied, there remains substantial scatter in the AAR age estimates for stratigraphically coeval intervals in central Arctic Ocean cores. This likely reflects a combination of enhanced mixing in low sedimentation rate environments, differences in effective diagenetic burial temperatures, and potentially the poorly defined role of microbial activity.

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## Acknowledgements

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# **Supplementary Material**

Supplementary material associated with this article can be found in the online version.

# 490 Data availability

The results of amino acid analyses of all 1009 subsamples included in this study will be archived at the World Data Service for Paleoclimatology (https://www.ncdc.noaa).

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