



Last Glacial Maximum deep water masses in southwestern Gulf of Mexico: Clues from benthic foraminifera

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Abstract

The repercussions of climate change during the Last Glacial Maximum (LGM) associated with changes in the structure and chemistry of abyssal waters in the southwestern Gulf of Mexico were studied by numerical analysis of benthic foraminiferal data from three sediment cores (water depth 1988-2735 m). In this context, to better understand the distribution patterns of modern benthic foraminifera in bathyal and abyssal waters of the area, species data from twelve core-tops (water depth 960-3255 m) were also analyzed. Multivariate Q-mode factor analysis separated the modern foraminiferal faunas of the North Atlantic Deep Water (NADW, deeper than ~2000 m), dominated by *Nuttallides decorata*, *Alabaminella turgida*, *Ioanella tumidula*, and *Globocassidulina subglobosa*, from those of the shallower Subantarctic Intermediate Water (AAIW) and Caribbean Midwater (CMW), characterized by the association of *Bolivina lowmani*, *Bulimina aculeata*, *Alabaminella turgida*, *Globocassidulina subglobosa*, *Epistominella exigua*, and *E. vitrea*.

Although most of the species are found in the entire stratigraphic interval studied, detailed examination shows differences between the dominant LGM and Holocene assemblages, which can be traced to climate-related modifications in world ocean circulation in the LGM, particularly those of NADW. During LGM, *Alabaminella turgida* (a species sensitive to oxygen depletion) along with a group of dissolution-prone species such as *Biloculinella irregularis* and *Cornuloculina inconstans*, thrived in the deepest part of the study area, indicating well-oxygenated waters of glacial NADW (GNADW). As the production of GNADW ceased in the Holocene, the deep areas in the Gulf were occupied by the present NADW that contains less oxygen and more CO₂. This water mass is more corrosive and precluded the presence of dissolution-prone species during the LGM, except in the most calcium carbonate- and oxygen-rich areas next to the Yucatan platform. Conditions in this area allowed the proliferation of *Nuttallides decorata* during the Holocene, while the *N. decorata* and *Alabaminella turgida* populations decreased in the western Gulf.

Keywords: Benthic foraminifera, paleoceanography, Last Glacial Maximum, water masses, southwestern Gulf of Mexico.

Resumen

Se estudió la repercusión de los cambios climáticos ocurridos durante el Último Máximo Glacial (UMG) en la estructura y química del agua de fondo (1988-2735 m) del suroeste del Golfo de México por medio del análisis numérico de foraminíferos bentónicos en tres núcleos sedimentarios. En este contexto, para entender mejor la distribución de foraminíferos bentónicos modernos en las aguas batiales y abisales del área, se analizaron también los datos de especies en la parte superior de doce núcleos sedimentarios (960-3255 m de profundidad). Por medio de un análisis multivariado de factores modo Q, se delimitaron dos asociaciones principales:

la asociación dominada por *Nuttallides decorata*, *Alabaminella turgida*, *Ioanella tumidula* y *Globocassidulina subglobosa* en Agua Profunda Noratlántica (APNA, a profundidades mayores de ~2000 m) y la asociación de *Bolivina lowmani*, *Bulimina aculeata*, *Alabaminella turgida*, *Globocassidulina subglobosa*, *Epistominella exigua* y *E. vitrea*, característica de profundidades más someras del Agua Antártica Intermedia y el Agua del Caribe.

Aunque la mayoría de las especies se encuentran en todo el intervalo estratigráfico estudiado, el examen detallado de las faunas muestra claras diferencias entre las asociaciones dominantes del UMG y el Holoceno. Dichas diferencias se pueden relacionar con las modificaciones producidas por los cambios climáticos en el UMG en la circulación oceánica global, particularmente con aquellas del APNA. Durante el UMG, *Alabaminella turgida* (una especie altamente sensitiva a bajas concentraciones de oxígeno disuelto en el agua) y un grupo de especies susceptibles a la disolución tales como *Biloculinella irregularis* y *Cornuloculina inconstans*, prosperaron en la parte más profunda del área de estudio, indicando la presencia de las aguas bien oxigenadas de la APNA glacial. Al cesar la producción de esta masa de agua en el Holoceno, las áreas profundas del Golfo de México fueron ocupadas por el APNA actual, con menor contenido de oxígeno y mayor CO₂. Esta masa de agua de mayor corrosividad, no permitió la presencia del conjunto de especies poco tolerantes a la disolución presente durante el UMG, excepto en las zonas cercanas a la plataforma de Yucatán, donde se presenta un mayor contenido de oxígeno y carbonato de calcio. Las condiciones en esta área permitieron la proliferación de *Nuttallides decorata* durante el Holoceno, mientras que las poblaciones de *N. decorata* y *Alabaminella turgida* disminuyeron en la parte occidental del Golfo

Palabras clave: Foraminíferos bentónicos, paleoceanografía, Último Máximo Glacial, masas de agua, Golfo de México suroccidental.

1. Introduction

The oceans played a major role in climatic changes associated with the last-glacial period. The modification of the thermohaline circulation and, particularly, shifts in the production/distribution of North Atlantic Deep Water (NADW) changed the balance of heat and salt transport, affecting climate on a planetary scale. In the modern ocean, NADW forms in the Greenland, Iceland, Labrador and Norwegian seas, sinks and flows southward into the major basins of the world's oceans. However, due to the extension of the northern ice sheets during the Last Glacial Maximum (LGM), NADW formed southward, altering its structure, physical and chemical properties (*i.e.* temperature, salinity, oxygen content, nutrient inventory, CO₂).

The benthic foraminiferal faunas were affected by these changes in glacial NADW. An earlier investigation (Machain-Castillo *et al.*, 1998) reports that less corrosive, CO₂-depleted, LGM bathyal and abyssal waters of the Gulf of Mexico and Caribbean Sea supported populations of high-Mg-calcitic benthic foraminifera and aragonitic pteropods that are not present in modern sediments. In this paper we examine the entire benthic foraminiferal suite, and compare modern with LGM assemblages in order to assess deep-water variations related to late Pleistocene climatic changes.

Modern benthic foraminifera from the Gulf of Mexico are well known. There have been major studies on their geographic and bathymetric distributions from the 1950s to the 1980s (Phleger and Parker, 1951; Parker, 1954; Pflum and Frerichs, 1976; Culver and Buzas, 1981, 1983; Poag, 1981, 1984). More recently, their relationships with water masses in the northern Gulf have also been investigated (Denne and Sen Gupta, 1991, 1993; Jones and Sen Gupta, 1995; Osterman, 2003). Pleistocene benthic foraminiferal faunas of the Gulf are much less known in spite of several

significant studies (Dignes, 1979; Jones and Sen Gupta, 1996; Machain-Castillo *et al.*, 1998; Denne and Sen Gupta, 2003).

2. Material and methods

This study is based on (a) 12 core-top samples (960–3255 m water depth) to delineate the modern faunas in sediments underlying the intermediate and deep water masses, and (b) three gravity cores collected below sill depth (1988–2735 m) in the central southwestern Gulf of Mexico to compare LGM assemblages and assess deep-water variations related to Pleistocene climatic changes (Figure 1, Table 1). The JS cores were taken by the Universidad Nacional Autónoma de México from the O/V *Justo Sierra* and the K cores by the U.S. Naval Oceanographic Office from the USNS *Kane*. The foraminifera were obtained from 15 cc sediment samples collected at 10 cm stratigraphic intervals. The sediments were oven dried at 50 °C, weighed, washed through a 63 µm mesh, dried and weighed again. The washed residues were subdivided with the aid of an Otto microsplitter to obtain aliquots of approximately 1000 benthic foraminifers. All species were identified and counted. A Q-mode factor analysis was performed on the species-abundance data with the Statistica 6.0 program (see section 5.1 for detail).

3. Water masses

3.1. Present-day water masses

The water masses entering the Gulf of Mexico through the Yucatan Channel are modified as they travel towards the west and mix with resident and coastal waters. According

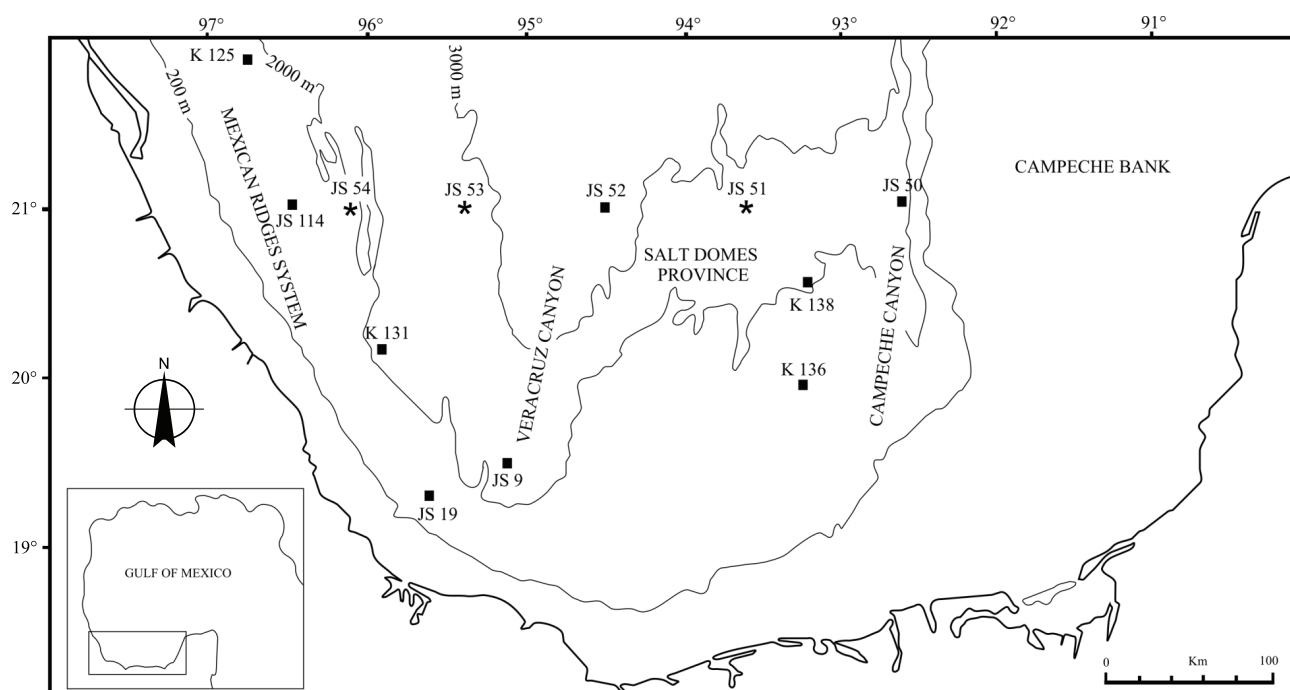


Figure 1. Study area. Squares mark locations of surface (core-top) samples; asterisks mark locations of cores that include Pleistocene sediments (Modified from Martínez-Trápaga, 1990).

Table 1. Locations and depths of core-top and down-core* samples.

Core No.	Latitude (N)	Longitude (W)	Water depth (m)	Core length (cm)
JS114	21°00.0'	96°37.4'	960	
JS19	19°21.0'	95°38.1'	980	
K136	19°58.9'	93°15.0'	1215	
K125	21°50.0'	96°38.0'	1245	
K138	20°31.0'	93°13.5'	1711	
JS54*	21°00.0'	96°06.1'	1988	130
K131	20°11.2'	95°59.3'	2001	
JS9	19°30.4'	95°18.3'	2117	
JS50	21°00.0'	92°42.0'	2240	
JS51*	20°59.9'	90°09.9'	2680	130
JS53*	20°59.7'	95°24.1'	2735	160
JS52	20°59.9'	94°30.2'	3255	

to Aldeco-Ramírez *et al.* (2009) three water masses are found in the upper 120 m: Gulf Common Water (GCW) with $T \sim 22.5^\circ\text{C}$, $S \sim 36.3\text{--}36.4$ (Vidal *et al.*, 1994), Caribbean Tropical Surface Water (CTSW) with $T > 28^\circ\text{C}$, $S > 36.4$, and the deeper Caribbean Subtropical Underwater (CSUW) with $T \sim 22.2\text{--}26^\circ\text{C}$, $S \sim 36.4\text{--}36.7$ (Schroeder *et al.*, 1974). In the eastern part of the Gulf, CSUW (also reported as Subtropical Underwater or Tropical Water) is present to depths of 150–250 m (salinity maximum ~ 36.7 at $T \sim 23^\circ\text{C}$), whereas in the western Gulf, the Gulf Water (GW, salinity 36.4–36.5) is found to ~ 250 m (Morrison *et al.*,

1983). Around the Yucatan Channel, Kinard *et al.* (1974) and Rivas *et al.* (2005) report sporadic occurrences of the Sargasso Sea Water (18°C , $S \sim 36.3$, oxygen maximum $\sim 3.4\text{ ml L}^{-1}$) between 200–400 m, while its presence in the loop current and adjacent anticyclonic rings is reported by Morrison and Nowlin (1977) and Morrison *et al.* (1983). The next underlying water mass, the Tropical Atlantic Central Water (TACW) with its associated oxygen minimum ($2.5\text{--}2.9\text{ ml L}^{-1}$), also called Oxygen Minimum Water (OMW), is found down to $\sim 600\text{--}700$ m (Nowlin, 1972; Vidal *et al.*, 1994; Rivas *et al.*, 2005). At $\sim 600\text{--}1000$ m, a remnant of the Antarctic Intermediate Water, or Subantarctic Intermediate Water, is found (AAIW, or SAIW) with salinity minimum ~ 34.8 , $T \sim 7^\circ\text{C}$ (Vidal *et al.*, 1994; Rivas *et al.*, 2005). At depths greater than about 1000 m, Upper NADW enters the Gulf of Mexico via the Caribbean basins. Sill depth of about 1900 m prevents deeper NADW from entering the Gulf. Within the Gulf, the upper part of this NADW is mixed with Caribbean Intermediate Water (CIW), also called Caribbean Midwater (CMW), which has a slightly lower salinity and higher silicate content than pure NADW (Morrison *et al.*, 1983; Metcalf, 1976; Vidal *et al.*, 1994). Below about 1100–1400 m, high-oxygen NADW (salinity < 35 , $\sim 4^\circ\text{C}$) or Gulf Basin Water (GBW) fills the entire basin. McLellan and Nowlin (1963) show that potential temperature and salinity distributions below 1500 m are uniform; however, dissolved oxygen content decreases away from the Yucatan Channel (Rivas *et al.*, 2005).

3.2. Last Glacial Maximum NADW

Today the deepest water mass entering the Gulf of Mexico is derived from the upper NADW that enters the Caribbean Sea (sill depth ~1900 m) and the Gulf of Mexico via the Yucatan Channel, with a sill depth of 2040 m (Rivas *et al.*, 2005). However, paleoceanographic reconstructions by several authors (with various proxies) indicate that during LGM two distinct water masses occurred in the depth range presently occupied by NADW, with the boundary at about 2–2.5 km depth. The shallower one contained fewer nutrients and less CO₂ than the present NADW, and the deeper one was nutrient- and CO₂-enriched (Curry *et al.*, 1988; Duplessy *et al.*, 1988; Haddad and Droxler, 1996; Boyle, 1997; Marchitto and Broecker, 2006; Lynch-Stieglitz *et al.*, 2007). Thus, the glacial bathyal and abyssal waters of the Gulf of Mexico were depleted in nutrients and CO₂ compared to the present ones.

4. Stratigraphic framework

Machain-Castillo *et al.* (1998) delineated the stratigraphic framework and boundaries for the cores in the present study, using the standard planktonic foraminiferal zonation and chronology for the Gulf of Mexico (Kennett and Huddlestun, 1972; Kennett *et al.*, 1985; Flower and Kennett, 1990), as well as the oxygen-isotope record of *Globigerinoides sacculifer* in core JS 54. Following Kennett and Huddlestun (1972), they recognized the upper part of the Y zone (Y2 and Y1) by high frequencies of *Globorotalia crassaformis* and *Globigerina falconensis*, and the virtual absence of *Pulleniatina obliquiloculata*. All cores reached the Pleistocene subzone Y2, which is correlated with Marine Isotope Stage 2 (Kennett and Huddlestun, 1972); thus the basal portion of the cores represents the LGM.

The Y2–Y1 boundary was recognized by the last consistent occurrence of *Globorotalia inflata*, the increase of *G. crassaformis* and *Globigerinoides ruber*, and the sporadic low frequencies of the *Globorotalia menardii* group. The Y1 subzone was recognized by high frequencies of *Neogloboquadrina dutertrei* and *G. ruber*, low frequencies of *G. crassaformis* and the virtual absence of *G. inflata*. This subzone represents the incursion of meltwater from the Laurentide Ice Sheet into the Gulf from 16 to 11.6 kys B.P. (Flower and Kennett, 1990).

The Pleistocene–Holocene boundary (Y–Z, 11 ka) and subzone Z2 were recognized by the consistent presence of *G. menardii* (Holocene) and an abrupt change in $\delta^{18}\text{O}$ values of *Globigerinoides ruber* (Figure 2). The Y/Z boundary is at about 55 cm in JS-51, 65 cm in JS-53, and 75 cm in JS-54. The Z zone is characterized by the presence of warm water forms (*G. menardii* and *P. obliquiloculata*) and the absence of cool water forms (*G. inflata* and *G. falconensis*), with the highest frequencies of *G. menardii* and *P. obliquiloculata* during the Z1 subzone, from 6 kys B.P. to Present (Kennett

et al., 1985).

5. Benthic foraminifera

5.1. Separation into factors

The core-top sediments contain 70 species of benthic foraminifera, most of which occur in very low abundances. To delineate the main associations, and avoid the noise due to sporadic or reworked species, only the species that were present in at least three samples and constituted at least 1% of the assemblage in one sample were used in the Q-mode factor analysis (Appendix 1). We selected a two-factor solution (79% of total variance) because it better explains the data and separates the intermediate water faunas (shallower than ~2000 m) from the deeper ones (Tables 2, 3). Although a three-factor solution increased the total explained variance to 90%, the third factor only separated those samples from the NADW that contained reworked shallower water species (*i.e.* JS 9, JS 50, K138). Other authors using a cutoff similar to ours (~80% of total variance) are Nomura (1991), Jian and Wang (1997), Schmiedl *et al.* (1997) and Murgese and De Deckker (2007).

5.2. Factor 1

The core-top Factor 1 (F 1 c-t) assemblage is dominated by *Bolivina lowmani*, *Bulimina aculeata*, *Alabaminella turgida*, *Globocassidulina subglobosa*, *Epistominella exigua*, and *E. vitrea* (Figure 3, Table 3), and is represented in core-tops from 960–1711 m water depth, and also in cores JS 9 (2711 m) and JS 50 (2680 m). Core JS 9 was taken from the western flank of the Veracruz Canyon, and JS 50 from the west of the Campeche Canyon (Figure 1); the presence of shelf species (*e.g.*, *Elphidium* spp., Figure 3c) and species of Factor 1 assemblage in these cores indicates slumped material in the sediment. The bathymetric ranges and environments of these species in the Gulf of Mexico and Caribbean Sea have been reported by various authors. *Bolivina lowmani* seems to have a wide tolerance to environmental parameters; it is widely distributed and has a meroplanktonic stage that helps dispersal. Its reported depth range in the Gulf of Mexico is 0–3632 m (Sen Gupta *et al.*, 2009). Denne and Sen Gupta (1991, 1993) find it a common species in the 820–1341 m range, and Jones and Sen Gupta (1995) report it as one of the most abundant species in the 2700–3900 m range in the northwestern Gulf. *Bulimina aculeata* has been reported from 55 to 3200 m in the Gulf (Sen Gupta *et al.*, 2009), but it is most abundant in the lower bathyal region (Phleger and Parker, 1951; Pflum and Frerichs, 1976). Denne and Sen Gupta (1991) report it as one of the main components of their CMW assemblage. *Alabaminella turgida* also has a large depth range (1–3850 m; Sen Gupta *et al.*, 2009), but it is more abundant in the abyssal waters of the Gulf, especially in well-oxygenated

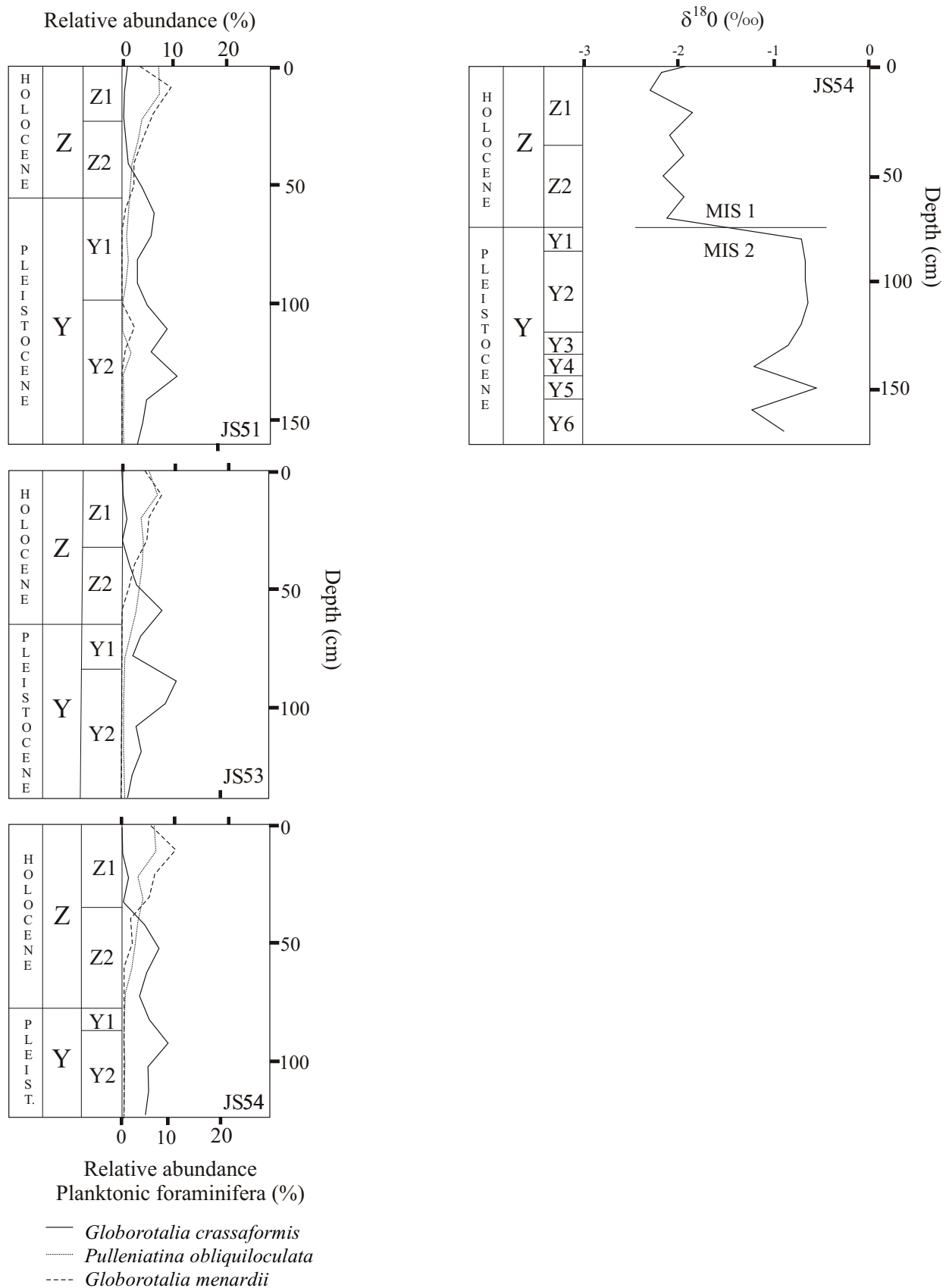


Figure 2. Left: Biostratigraphic zonation in cores JS51, JS53, and JS54. Right: $\delta^{18}\text{O}$ curve for *Globigerinoides ruber* in core JS54 (Modified from Machain-Castillo *et al.*, 1998).

Table 2. Eigenvalues and factor loadings for core-top samples.

	Eigenvalue	% Total variance	Cumulative eigenvalue	Cumulative %
1	7.45173	62.09775	7.45173	62.09775
2	2.047987	17.06656	9.499717	79.16431

Factor Loadings (Varimax normalized) Extraction: Principal components

	Factor 1	Factor 2
JS9	0.818327	0.372051
JS19	0.858388	0.039756
K131	0.370763	0.901973
K136	0.891491	0.119481
K125	0.700796	0.123143
K138	0.667599	0.621318
JS114	0.772131	0.441085
JS54	0.514242	0.84918
JS51	-0.177407	0.829354
JS50	0.711386	0.475975
JS53	0.454662	0.875396
JS52	0.149038	0.794799

waters, with frequencies as high as 20% (Phleger and Parker, 1951; Parker, 1954; Pflum and Frerichs, 1976). Denne and Sen Gupta (1993) report that it is associated with a group of species sensitive to low concentrations of dissolved oxygen and better represented away from the influence of the Mississippi River Delta. The species is common in the Caribbean but typically shows lower frequencies (Galluzzo et al., 1990). *Globocassidulina subglobosa* is regarded as a cosmopolitan and eurybathyal species, resistant to dissolution (Dignes, 1979; Denne and Sen Gupta, 1991). *Epistominella exigua* is widely distributed in the northern Gulf, except around the Mississippi River Delta. It has been reported from 2 to 3700 m (Sen Gupta et al., 2009) with the highest frequencies generally between 500-1000 m (Phleger and Parker, 1951; Parker, 1954; Pflum and Frerichs, 1976). Denne and Sen Gupta (1991, 1993) consider it to be a characteristic species in AAIW (650-1000 m). The depth range of *Epistominella vitrea* in the Gulf is 1-3632 m (Sen Gupta et al., 2009). Parker (1954) and Osterman (2003) find the species more abundant above 100 m and off the Mississippi River Delta. Besides the species with highest scores on Factor 1c-t, *Bolivina albatrossi*, *Bulimina alazanensis*, *B. mexicana*, *Gyroidinoides polius*, and *Osangularia culter* are more abundant in samples related to this factor than in deeper samples (Figures 3d, 3e). The wide-ranging *B. albatrossi* is most abundant in middle and lower bathyal environments (Phleger and Parker, 1951; Pflum and Frerichs, 1976). *Bulimina alazanensis* has been found in the Gulf from 42 to 3640 m (Sen Gupta et al., 2009), and is most abundant in middle bathyal environments (Phleger and Parker, 1951; Pflum and Frerichs, 1976). *Bulimina mexicana* is known to be associated with the

Table 3. Factor scores for core-top samples.

Factor Scores (Varimax normalized) Extraction: Principal components		
Species	Factor 1	Factor 2
<i>Alabaminella turgida</i>	1.54681	2.29915
<i>Bolivina albatrossi</i>	0.92765	-0.8929
<i>B. lowmani</i>	4.3341	2.8112
<i>B. ordinaria</i>	0.26811	-0.4704
<i>Bulimina aculeata</i>	1.70276	-1.12526
<i>B. alazanensis</i>	0.43242	-0.60811
<i>B. mexicana</i>	0.78418	-0.75732
<i>Globocassidulina subglobosa</i>	1.35106	-0.1366
<i>Cassidulina neocarinata</i>	-0.56123	-0.12679
<i>C. nocrossi australis</i>	-0.43871	-0.17515
<i>G. cf. subglobosa</i>	-0.28406	-0.08476
<i>Cibicides wuellerstorfi</i>	-0.47839	0.06455
<i>Cibicidoides</i> sp	-0.44291	-0.03866
<i>C pachydermus</i>	-0.01445	-0.43852
<i>Coryphostoma subspincensis</i>	-0.43539	-0.0989
<i>Eggerella bradyi</i>	-0.48734	-0.05847
<i>Elphidium discoideale</i>	-0.50019	-0.16772
<i>E. sp. 1</i>	-0.44391	-0.16104
<i>Epistominella exigua</i>	1.14976	-0.87946
<i>E. vitrea</i>	1.03069	-0.58951
<i>Gavelinopsis translucens</i>	-0.2906	-0.26801
<i>Glomospira charoides</i>	0.40678	-0.27906
<i>Gyroidina altiformis</i>	-0.52845	-0.12233
<i>G. polius</i>	-0.18983	-0.22452
<i>G. regularis</i>	-0.31505	-0.28685
<i>G. sp 1</i>	-0.55477	-0.12455
<i>G. sp 2</i>	-0.17726	-0.21015
<i>Hoeglundina elegans</i>	-0.54949	0.08174
<i>Ioanella tumidula</i>	-0.97852	1.12121
<i>Neocorbina</i> sp	-0.50165	-0.14997
<i>Neocrosbya minuta</i>	-0.54587	-0.20941
<i>Nonionella</i> sp	-0.43061	-0.16485
<i>Nuttallides decorata</i>	-2.07369	4.6269
<i>Oridorsalis</i> sp	-0.48362	-0.10688
<i>Osangularia culter</i>	0.51185	-0.4993
<i>Planulina exorna</i>	-0.51518	-0.17688
<i>Pullenia subspherica</i>	-0.36245	-0.11168
<i>P. sp</i>	-0.38538	-0.20578
<i>Quinqueloculina venusta</i>	-0.71844	0.17673
<i>Q. sp 7</i>	-0.63882	-0.03695
<i>Sphaeroidina bulloides</i>	-0.25687	-0.34214
<i>Uvigerina lavéis</i>	-0.54501	-0.16009
<i>U. peregrina</i>	0.68198	-0.69247

OMZ (Oxygen Minimum Zone; 350-550 m; Denne and Sen Gupta, 1991). Parker (1954) records the maximum abundance of *Gyroidinoides polius* (range 99-3850 m; Sen Gupta et al., 2009) at 2788 m, and Denne and Sen Gupta (1991, 1993) consider it a species associated with CMW. *Osangularia culter* is found in much of the northern Gulf, except around the Mississippi River Delta, at depths greater than 400 m, with higher frequencies above 2000 m (Phleger and Parker, 1951; Parker, 1954; Pflum and Frerichs, 1976).

Overall, the assemblage associated with Factor 1

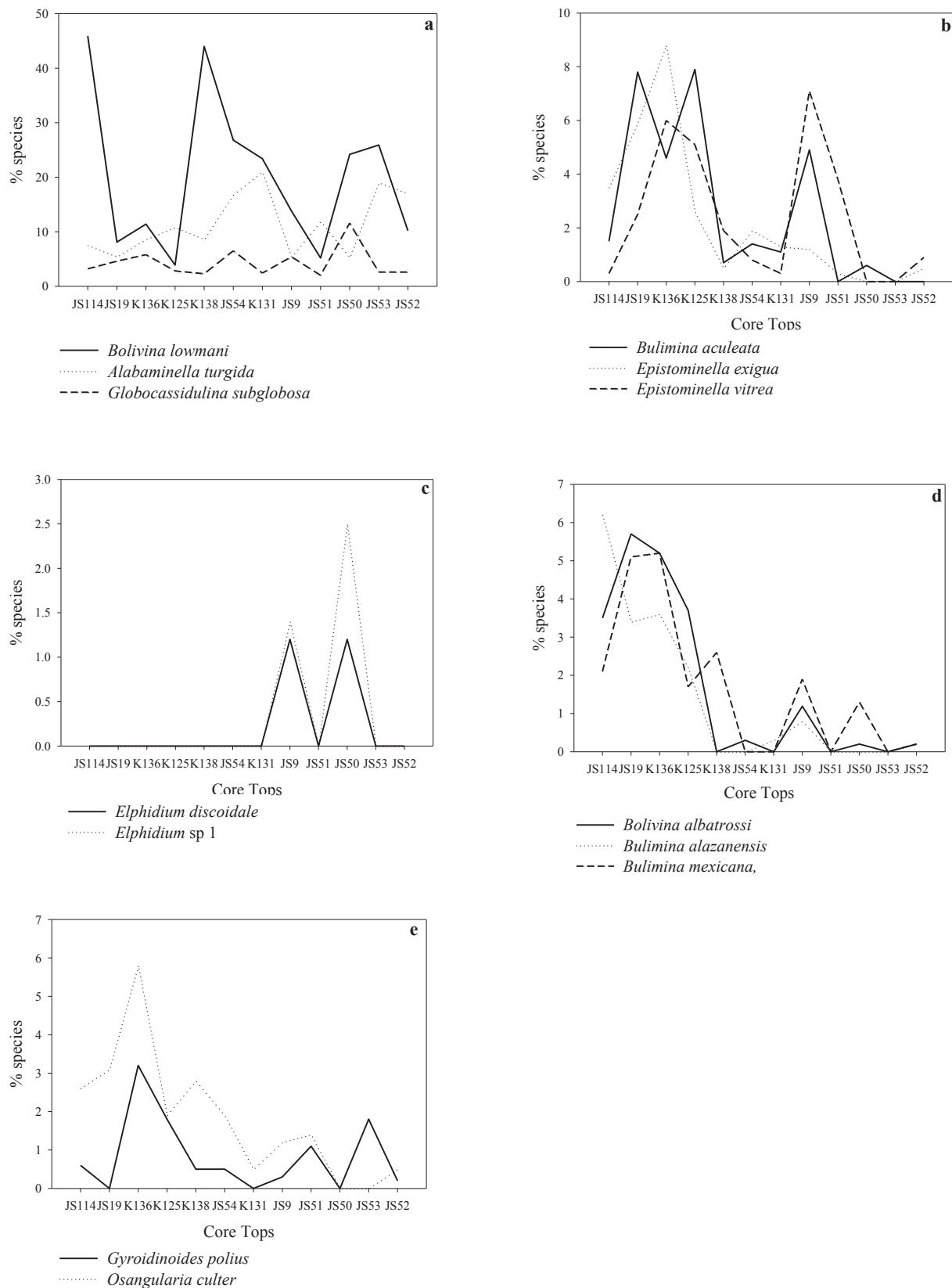


Figure 3. Core-top distribution of Factor 1 assemblage. a) *Bolivina lowmani*, *Alabaminella turgida*, *Globocassidulina subglobosa*; b) *Bulimina aculeata*, *Epistominella exigua*, *Epistominella vitrea*; c) *Elphidium discoidale* and *Elphidium* sp 1; d) *Bolivina albatrossi*, *Bulimina alazanensis*, *Bulimina mexicana*; e) *Gyroidinoides polius*, *Osangularia culter*.

c-t is similar to the northwestern Gulf assemblage that straddles the AAIW-CMW boundary (Table 4). Our inability to differentiate the AAIW and CMW foraminiferal assemblages may be partly related to the small number of core-top samples, but the distinction is unclear even in the larger data sets of Culver and Buzas (1983), Denne and Sen Gupta (1991), and Osterman (2003). They suggest that the difference between these water masses may not be sharp, because the Gulf AAIW has been modified in its trajectory and it is only a remnant of the original water mass.

5.3. Factor 2

The Factor 2 c-t assemblage is typical of the 1988-3255 m core-top samples and cores JS 9 and JS 50, all collected from below the sill depth. The species with highest scores (> 1) on this factor are: *Nuttallides decorata*, *Bolivina lowmani*, *Alabaminella turgida* and *Ioanella tumidula* (Figure 4). *Nuttallides decorata*, *A. turgida*, *I. tumidula*, and *Globocassidulina subglobosa* have been reported as predominant taxa in the deep Gulf of Mexico (Phleger and Parker, 1951; Phleger, 1960; Dignes, 1979) and the Caribbean (Gaby and Sen Gupta, 1985; Bertrand, 1986; Galluzzo et al., 1990). *Nuttallides decorata* has been reported from 155 to 3850 m in the Gulf of Mexico (Sen Gupta et al., 2009); it is generally considered as typical of bathyal or abyssal waters (Phleger and Parker, 1951; Phleger, 1960; Pflum and Frerichs, 1976). Dignes (1979) suggests its distribution could be partly controlled by calcite dissolution, since its relative abundances (particularly down-core) increase in areas of low-carbonate dissolution. Denne and Sen Gupta (1993) report the occurrence of *N. decorata* from 1204 to 1361 m (their deepest sample) and consider it characteristic of the CMW (1000-1500 m). The reported depth range of *Ioanella tumidula* in the Gulf is 68-

3850 m (Sen Gupta et al., 2009). Phleger and Parker (1951) record its highest abundances below 2000 m, and Pflum and Frerichs (1976) find an abundance increase with depth ($> 10\%$ at > 3300 m in the central and western Gulf). In the study area, its frequencies are $< 2\%$ above 2700 m, 6% at 2735 m, and 17% at 3255 m. In the Grenada, Venezuela, and Colombia basins, *I. tumidula* is known from 1000-4000 m, increasing in abundance with water depth; a maximum value of 13% is recorded at 4000 m in the Colombia Basin (Gaby and Sen Gupta, 1985; Bertrand, 1986; Galluzzo et al., 1990). *Hoeglundina elegans* and *Cibicides wuellerstorfi* also show maximum abundances in the Factor 2 c-t assemblage (Figure 4b). Phleger (1960) considers them as characteristic of the lower continental slope and deep sea, along with *N. decorata*, *I. tumidula*, and *A. turgida*. Denne and Sen Gupta (1991, 1993) recognize *C. wuellerstorfi* as a characteristic species of their deepest assemblage (CMW, 1000-1500 m) in the northwestern Gulf.

5.4. Down-core distributions

To delineate changes in foraminiferal assemblages in waters below sill depth during the late Pleistocene (LGM and deglaciation), we looked at the down-core distribution of factors and species in 3 cores along 21°N (JS 54, JS 53, JS 51; 1988-2735 m; Appendix 2). A core that contains reworked material (JS 50) and one that did not reach the Pleistocene (JS52) were not used.

Factor analysis of down-core foraminiferal data produces 3 factors that explain 97.8% of the total variance (Tables 5, 6, 7). Down-core Factor 1 (F 1 d-c) accounts for 82.4% of the total variance; only two species show positive scores: *Nuttallides decorata*, with 6.08 and *Oridorsalis* sp. 1 with 0.09 (Figure 5). The species with strongest negative scores are *Bolivina lowmani* (-0.98), *Alabaminella turgida*

Table 4. Comparison of benthic foraminiferal assemblages for the slope and abyssal water masses in the Gulf of Mexico (mwd = meters water depth).

Culver and Buzas, 1983	Osterman, 2003	Denne and Sen Gupta, 2003	This paper
Bathyal and abyssal	Bathyal	Subantactic Intermediate Water (SAIW)	AAIW + CMW
	640-1020 (mwd)	671-1000 (mwd)	960-1700 (mwd)
<i>Laticarinina pauperata</i>	<i>Laticarinina pauperata</i>	<i>Epistominella exigua</i>	<i>Bolivina lowmani</i>
<i>Cyclamina cancellata</i>	<i>Bulimina aculeata</i>	<i>Osangularia rugosa</i>	<i>Bulimina aculeata</i>
<i>Osangularia culter</i>	<i>Cyclamina cancellata</i>	<i>Osangularia culter</i>	<i>Alabaminella turgida</i>
	<i>Osangularia culter</i>	<i>Bulimina alazanensis</i>	<i>Globocassidulina subglobosa</i>
	<i>Gyroidina neosoldanii</i>	<i>Uvigerina peregrina dirupta</i>	<i>Epistominella exigua</i>
			<i>Epistominella vitrea</i>
		Caribbean Midwater (CMW)	NADW (GBW)
		1000-1500 (mwd)	1988-3255 (mwd)
		<i>Gyroidinoides laevis</i>	<i>Nuttallides decorata</i>
		<i>Bulimina aculeata</i>	<i>Bolivina lowmani</i>
		<i>Nuttallides decorata</i>	<i>Alabaminella turgida</i>

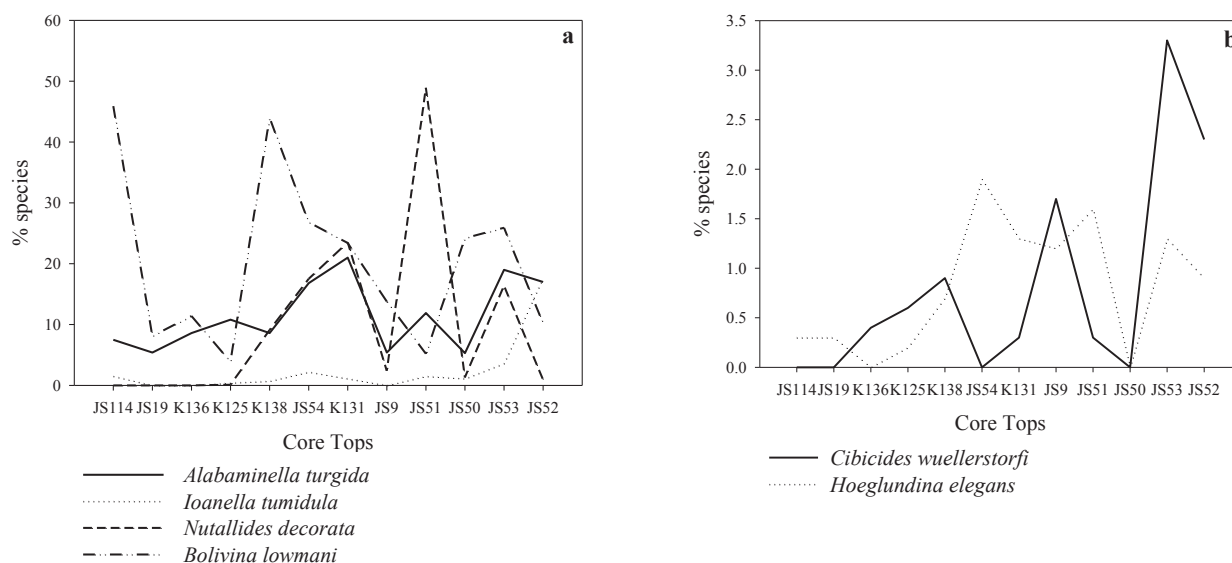


Figure 4. Core-top distribution of Factor 2 assemblage. a) Species with higher scores: *Alabaminella turgida*, *Ioanella tumidula*, *Nuttallides decorata*, *Bolivina lowmani*, b) Other species abundant in this factor: *Cibicides wuellerstorfi*, *Hoeglundina elegans*.

Table 5. Eigenvalues and factor loadings for down-core samples.

	Eigenvalue	% Total variance	Cumulative eigenvalue	Cumulative %
1	35.41598	82.36275	35.41598	82.36275
2	3.66996	8.53479	39.08594	90.89754
3	2.97063	6.90843	42.05657	97.80597

(-0.52) and *Ioanella tumidula* (-0.49). As explained above, *N. decorata* has been considered as characteristic of CMW and the abyssal Gulf. In our core-tops, this species has its upper depth limit at 1245 m (similar to the one found by Denne and Sen Gupta, 1993, in the northwestern Gulf), and it is the second most abundant species in our samples, after *B. lowmani*.

Factor 2 d-c (8.5% of total variance) is represented by *Alabaminella turgida* (score 5.9) and, to a minor extent, by *Ioanella tumidula* (0.7), *Biloculinella irregularis* (0.3), *Cornuloculina inconstans* (0.3), *Nuttallides decorata* (0.2), *Globocassidulina subglobosa* (0.06), *Francesita advena* (0.02), *Pyrgo nasuta* (0.02), *Cibicides wuellerstorfi* (0.02), and *Oridorsalis umbonatus* (0.005). *Alabaminella turgida*, the main species of this factor, is one of the most abundant species in the deepest Gulf, and is characteristic of well-oxygenated waters (Phleger and Parker, 1951; Parker, 1954; Pflum and Frerichs, 1976; Denne and Sen Gupta, 1991, 1993, 2003). The rest of the species with positive scores on Factor 2 d-c are mainly found in the Pleistocene section of the cores (Figure 6), and represent a mixture of porcelaneous, high-Mg-calcitic species (*B. irregularis*, *C. inconstans*, *P. nasuta*) and deep-water hyaline species (*C. wuellerstorfi*, *O. umbonatus*, *N. decorata*). *Bolivina lowmani*

shows a strong negative score (-1.48).

Factor 3d-c shows positive scores (Figure 7) for 6 species (*B. lowmani*, *A. turgida*, *N. decorata*, *G. subglobosa*, *H. elegans*, and *I. tumidula*) and the strongest negative scores: *C. inconstans* (-0.55), *B. irregularis* (-0.53), and *F. advena* (-0.40).

Bolivina lowmani, *A. turgida*, *N. decorata*, and *G. subglobosa* are the four most abundant species at depths greater than ~2000 m in the present southwestern Gulf of Mexico, as well as in the northwestern Gulf (Phleger and Parker, 1951, Phleger, 1960; Dignes, 1979) and the Caribbean (Gaby and Sen Gupta, 1985; Bertrand, 1986; Galluzzo *et al.*, 1990). The species with negative scores are the LGM dissolution-prone fauna of Factor 2 d-c.

The distribution of these three factors is different in each core (Table 6). Core JS 51 (Figure 2) contains a sequence of 160 cm reaching up to the Y2 subzone. All of the sequence is characterized by high loadings of Factor 1 d-c, implying there is no change in the foraminiferal assemblage (*Nuttallides decorata*). The Y and earliest Z2 subzones of core JS 53 show high loadings of Factor 2 d-c, indicating that during this time interval the central waters of the Gulf, around 2700 m depth, were dominated by the *Alabaminella turgida* assemblage while the rest of the Holocene is characterized by the *Bolivina lowmani* assemblage (Factor 3 d-c). The Pleistocene to early Holocene part of core JS 54 (Z2 subzone) is characterized by Factor 1 d-c while the late Holocene part (Z1 subzone) is dominated by Factor 3 d-c.

6. Discussion

Two modern assemblages of benthic foraminifera are clearly recognized in deep-bathyal and abyssal sediments

Table 6. Factor loadings for down-core samples H (Holocene), P (Pleistocene). Factor Loadings (Varimax normalized) Extraction: Principal components.

	Factor 1	Factor 2	Factor 3
H	51-0	0.9231	0.22432
	51-10	0.8737	0.2477
	51-20	0.80369	0.1387
	51-30	0.78297	0.20926
	51-40	0.74035	0.20746
	51-50	0.90841	0.21669
	51-60	0.87874	0.34457
	51-70	0.93555	0.25534
	51-80	0.83535	0.45333
	51-90	0.8674	0.34951
P	51-100	0.73579	0.22142
	51-110	0.73277	0.30082
	51-130	0.68069	0.27902
	51-140	0.82915	0.16496
	51-150	0.6857	0.32571
	51-160	0.7112	0.37767
	53-0	0.28341	0.36409
	53-9	0.28214	0.57505
	53-20	0.37231	0.32941
	53-30	0.38978	0.3377
H	53-40	0.42991	0.50014
	53-50	0.26435	0.63278
	53-60	0.44867	0.7964
	53-70	0.24843	0.91807
	53-80	0.37139	0.85569
	53-90	0.26028	0.90469
	53-100	0.47335	0.78436
	53-110	0.3397	0.86524
	53-120	0.34184	0.86517
	53-130	0.21011	0.9345
H	54-0	0.31957	0.27756
	54-10	0.45472	0.46989
	54-20	0.46182	0.18233
	54-30	0.3023	0.28707
	54-40	0.68429	0.24249
	54-50	0.71178	0.49289
	54-60	0.80466	0.44438
	54-70	0.81045	0.47125
	54-80	0.79011	0.50655
	54-90	0.8386	0.48014
P	54-100	0.82936	0.46951
	54-110	0.86875	0.41812

Table 7. Factor scores for down-core samples.

Factor Scores (Varimax normalized) Extraction: Principal components

Factor 1	Factor 2	Factor 3	Species
-0.520715	5.87532	1.272981	<i>Alabaminella turgida</i>
-0.136135	-0.12706	-0.322489	<i>Alliatina</i> sp
-0.103293	-0.20135	-0.349119	<i>Bolivina</i> spp
-0.976955	-1.48219	5.80151	<i>B. lowmani</i>
-0.167808	-0.30136	-0.201467	<i>Bulimina aculeata</i>
-0.114377	-0.28414	-0.284841	<i>B. alazanensis</i>
-0.110745	-0.29777	-0.250602	<i>B. mexicana</i>
-0.004404	0.06282	0.284103	<i>Globocassidulina subglobosa</i>
-0.141741	-0.18462	-0.179735	<i>G.</i> sp
-0.1584	0.01881	-0.121419	<i>Cibicides wuellerstorfi</i>
-0.192581	-0.08462	-0.019906	<i>Cibicoides</i> sp
-0.13144	-0.26214	-0.228562	<i>C. mundulus</i>
-0.104661	0.33212	-0.546921	<i>Cornuloculina inconstans</i>
-0.091557	-0.3095	-0.269348	<i>Coryphostoma subspiniensis</i>
-0.10642	-0.02422	-0.268427	<i>Eggerella bradyi</i>
-0.10052	-0.31492	-0.119329	<i>Epistominella exigua</i>
-0.048002	-0.29412	-0.279927	<i>E. vitrea</i>
-0.187141	0.01971	-0.404333	<i>Francesita advena</i>
-0.205615	-0.26626	-0.062128	<i>Glomospira charoides</i>
-0.13581	-0.30892	-0.013805	<i>Gyroidinoides polius</i>
-0.1874	-0.24545	-0.186902	<i>G.</i> sp 1
-0.017026	-0.25941	-0.265178	<i>G.</i> sp 2
-0.100343	-0.37994	0.194361	<i>Hoeglundina elegans</i>
-0.491116	0.67247	0.1921	<i>Ioanella tumidula</i>
-0.162841	-0.18693	-0.307176	<i>Melonis pompilioides</i>
-0.002824	0.33451	-0.532868	<i>Biloculinella irregularis</i>
6.077148	0.21256	0.910969	<i>Nuttalides decorata</i>
-0.105409	-0.37949	-0.092758	<i>Osangularia culter</i>
-0.112729	0.00475	-0.343378	<i>Oridorsalis umbonatus</i>
0.089207	-0.08044	-0.271356	<i>O.</i> sp 1
-0.116026	-0.23512	-0.127233	<i>Pullenia subspherica</i>
-0.144682	-0.19971	-0.252358	<i>P.</i> sp
-0.195534	0.01924	-0.313539	<i>Pyrgo nasuta</i>
-0.15952	-0.1263	-0.277673	<i>Quinqueloculina bosciana</i>
-0.178826	-0.04046	-0.12443	<i>Q. venusta</i>
-0.15523	-0.08407	-0.173469	<i>Q.</i> sp7
-0.084813	-0.06898	-0.377979	<i>Q.</i> sp 8
-0.069897	-0.15336	-0.342098	<i>Q.</i> spp
-0.06656	-0.11371	-0.390179	<i>Robertinoides bradyi</i>
-0.077259	-0.25577	-0.355092	<i>Uvigerina peregrina dirupta</i>

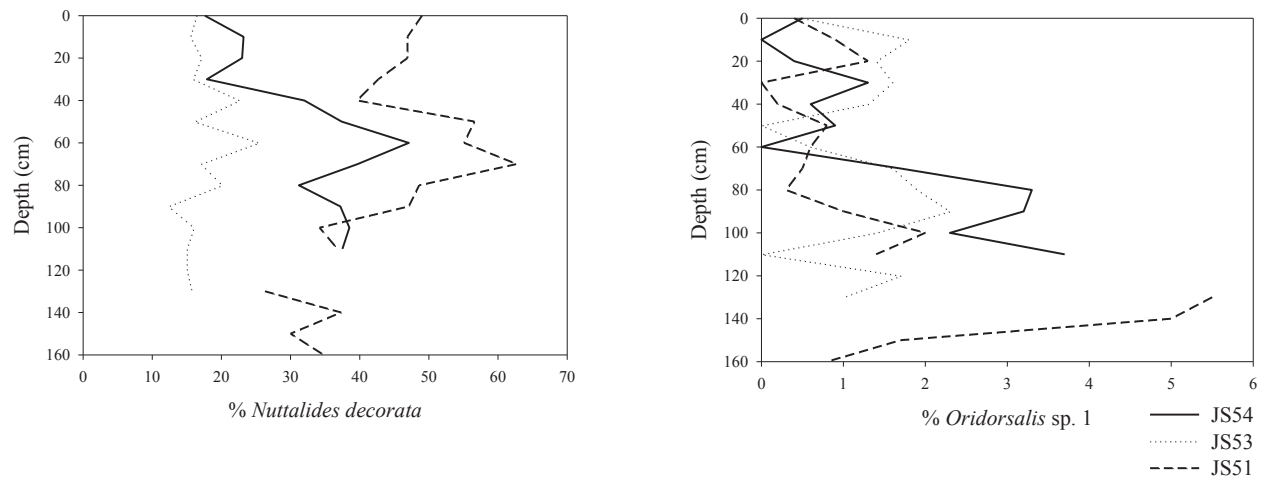


Figure 5. Down-core distribution of characteristic species of Factor 1: *Nuttalides decorata*, *Oridorsalis* sp. 1.

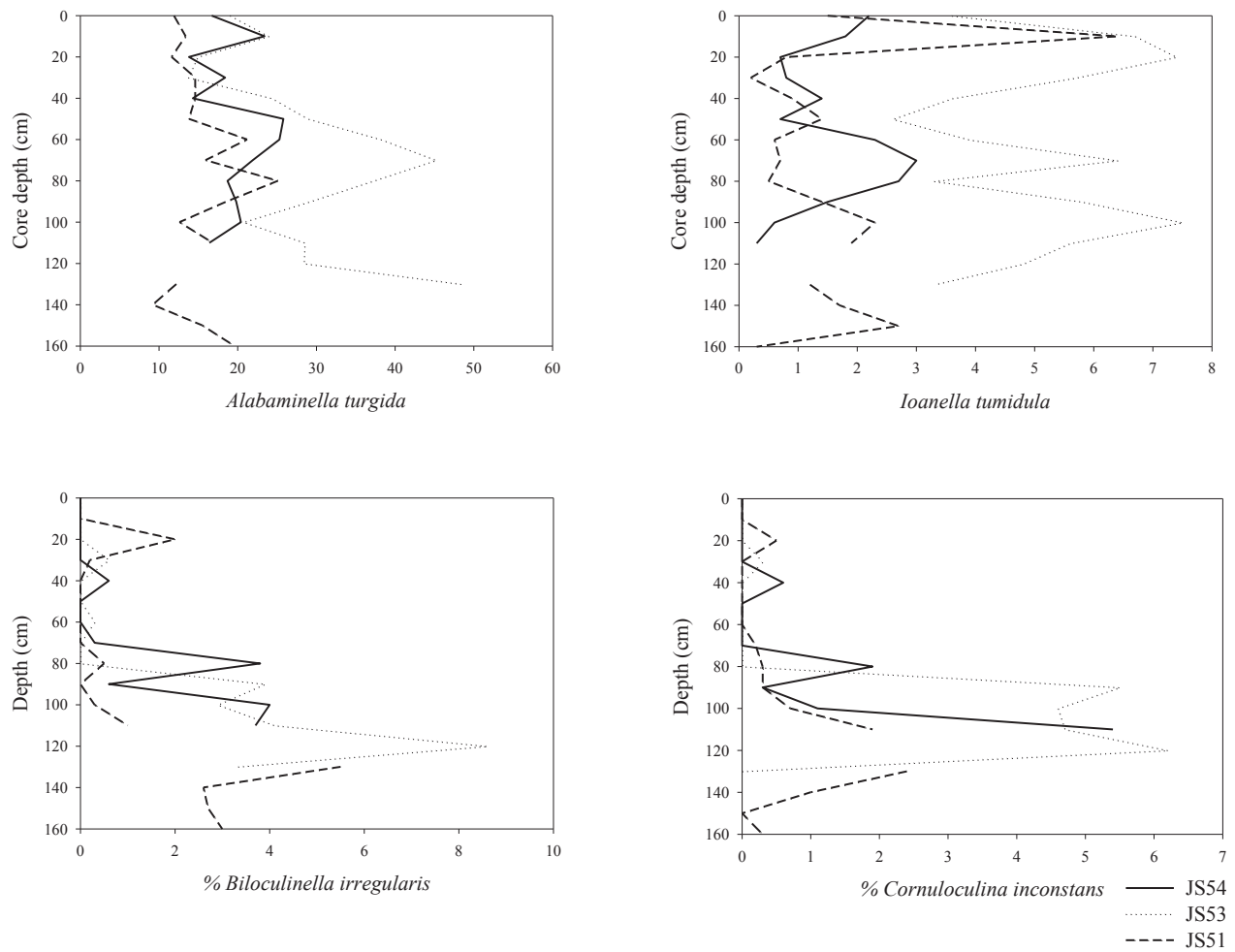


Figure 6. Down-core distribution of characteristic species of Factor 2: *Alabaminella turgida*, *Ioanella tumidula*, *Biloculinella irregularis*, *Cornuloculina inconstans*.

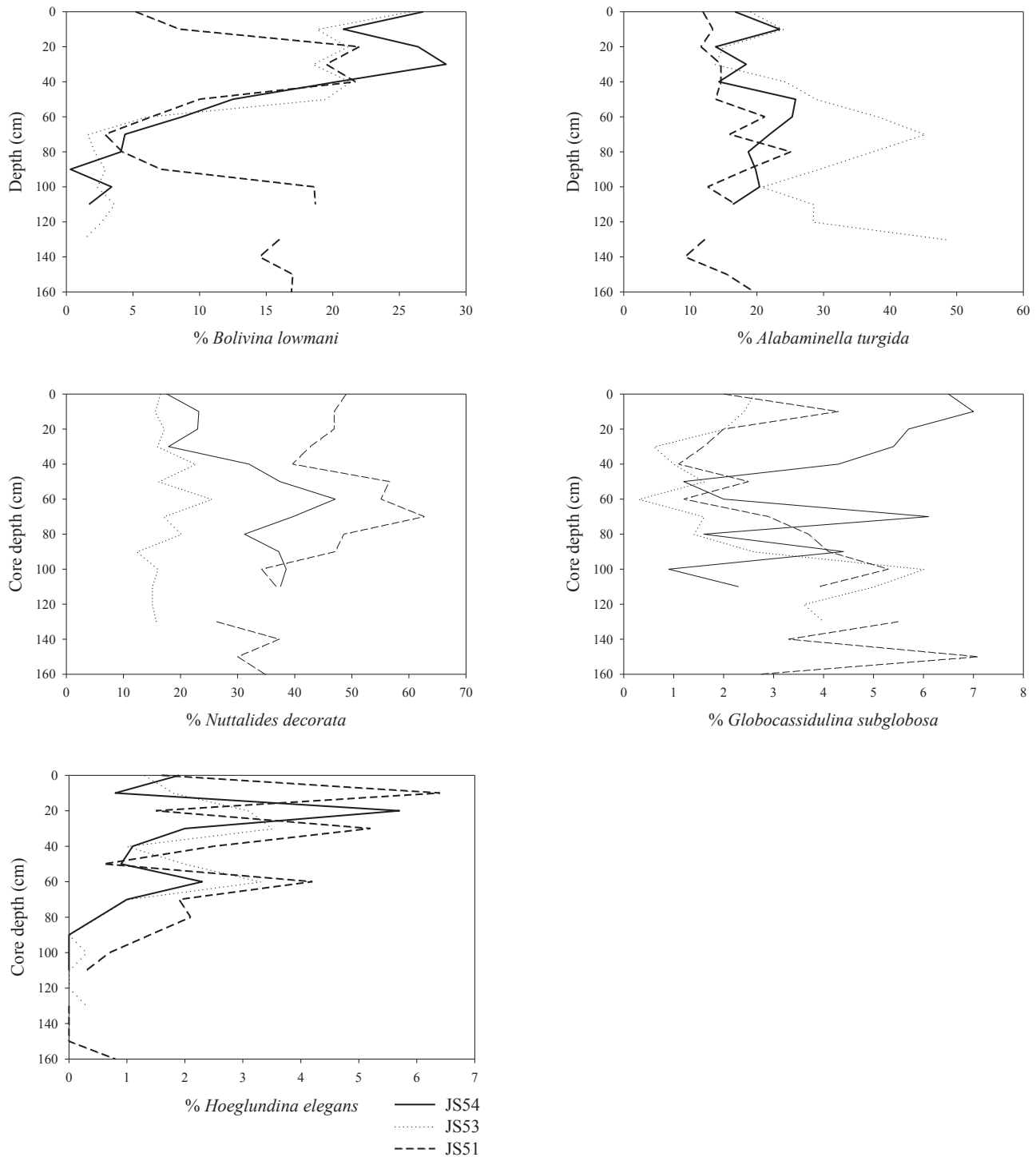


Figure 7. Down-core distribution of characteristic species of Factor 3: *Bolivina lowmani*, *Alabaminella turgida*, *Nuttalides decorata*, *Globocassidulina subglobosa*, *Hoeglundina elegans*.

of the southwestern Gulf of Mexico. One (shallower than 1700 m) contains species related to AAIW and CMW; the other is a deeper fauna related to NADW. Most of the species found in the modern NADW assemblage were also present in the late Pleistocene. However, important differences can be recognized. The results of factor analysis indicate that

during the late Pleistocene (LGM and deglaciation), the benthic foraminiferal fauna between ~2000 and 2250 m was dominated by the *N. decorata* assemblage, while the assemblages at depths ~2750 m were characterized by the *A. turgida* assemblage.

The *A. turgida* assemblage (Factor 2 d-c) is composed

of the most oxygen-sensitive species and a group of dissolution-prone species. Among the latter, *B. irregularis* has been particularly recognized as an indicator of a less-corrosive, well-oxygenated water mass present in the Caribbean and the Gulf of Mexico during the Pleistocene (Machain-Castillo *et al.*, 1998; Denne and Sen Gupta, 2003). Such a dissolution-prone fauna would thrive if a less corrosive glacial NADW (GNADW) entered the Gulf of Mexico via the Caribbean Sea and filled the deeper parts of the basin, thus modifying the GNADW. This modified GNADW permitted the proliferation and preservation of the *A. turgida* assemblage in the central Gulf below 2500 m. At shallower depths (~2000–2250 m), the *N. decorata* fauna was dominant, but still included the dissolution-prone species (*B. irregularis*, *C. inconstans*). *Nuttallides decorata* has also been associated with periods of low-carbonate dissolution (Dignes, 1979), and is probably intolerant of low-oxygen concentrations, since its abundance decreases near river deltas (Pflum and Frerichs, 1976); this is one of the species with positive scores on Factor 2 d-c.

As the upper GNADW shut down and modern NADW started flowing, less-oxygenated waters entered the Gulf and the most dissolution-prone species (*B. irregularis*, *C. inconstans*, and *F. advena*) retreated, first from the deepest areas (JS 53), probably due to the lack of water exchange below the sill depth. *Alabaminella turgida* and *N. decorata* persisted in the Holocene NADW, but there are noticeable differences in their history. The abundance of *A. turgida* decreased in the less-oxygenated Holocene waters of the Gulf, while *N. decorata* increased in abundance, particularly in the eastern Gulf. The eastern Gulf is bathed by NADW, with decreasing oxygen levels away from the Yucatan Channel. Thus, the most oxygenated waters in the Gulf below sill depth are the ones closest to the Yucatan carbonate platform. For our three cores, the JS 51 site is the closest to the Yucatan Channel, and should have the highest dissolved-oxygen concentration (Rivas *et al.*, 2005). Dissolved oxygen in bottom water was not measured at these locations. However, Martínez-Trápaga (1990) measured CaCO_3 content in the sediments, and found a slight increase in carbonate in core JS 51 (27.4%) compared to the other two (JS 53–26%; JS 54–23.8%). As NADW moves to the west, it mixes with local water and loses oxygen (McLellan and Nowlin, 1963; Rivas *et al.*, 2005), and sets the scene for the settlement of species such as *Bolivina* spp., which are characteristic of lower-oxygen environments in the central and western Gulf.

7. Conclusions

Modern benthic foraminiferal faunas in the bathyal and abyssal southwestern Gulf of Mexico can be separated into an AAIW-CMW assemblage and a NADW assemblage at about 1700 m water depth. This AAIW-CMW assemblage is similar to that found in the northern Gulf, and is

characterized by the association of *B. lowmani*, *B. aculeata*, *A. turgida*, *G. subglobosa*, *E. exigua*, and *E. vitrea*, with lesser abundances of *B. albatrossi*, *B. alazanensis*, *B. mexicana*, *G. polius*, and *O. culter*. North Atlantic Deep Water hosts an assemblage dominated by *N. decorata*, *B. lowmani*, *A. turgida* and *I. tumidula*, species that are also predominant in the deep northern Gulf of Mexico and the Caribbean.

The stratigraphic distribution of benthic foraminiferal assemblages in the central southwestern Gulf of Mexico reflects the oceanographic changes since LGM.

Concomitant with changes in the oxygen and CO_2 content of the upper GNADW, benthic foraminiferal faunas during the late Pleistocene were dominated by species with preference for well-oxygenated waters and substrates with relatively high carbonate content. During the Holocene, however, the deeper Gulf of Mexico became filled with a more corrosive, less-oxygenated NADW, which caused the virtual disappearance of dissolution-prone species (*B. irregularis*, *C. inconstans*) and the restriction of the *N. decorata* assemblage to the more carbonated and oxygen-rich areas of the eastern Gulf, closer to the Yucatan Channel.

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Appendix 1. List of species used in core-tops factor analysis.

Alabaminella turgida (Phleger and Parker, 1951)
Bolivina albatrossi Cushman, 1922
Bolivina lowmani Phleger and Parker, 1951
Bolivina ordinaria Phleger and Parker, 1951
Bulimina aculeata d'Orbigny, 1826
Bulimina alazanensis Cushman, 1927
Bulimina mexicana Cushman, 1922
Cassidulina neocarinata Thalman, 1950
Cassidulina nocrassi australis Phleger and Parker, 1951
Cibicides wuellerstorfi (Schwager, 1866)
Cibicidoides pachydermus (Rzehak, 1886)
Cibicidoides sp
Coryphostoma subspinescens (Cushman, 1922)
Eggerella bradyi (Cushman, 1911)
Elphidium discoidale (d'Orbigny, 1839)
Elphidium sp 1
Epistominella exigua (Brady, 1884)
Epistominella vitrea Parker, Phleger and Pearson, 1953
Gavelinopsis translucens (Phleger and Parker, 1951)
Globocassidulina subglobosa (Brady, 1881)
Globocassidulina cf. *C. subglobosa*
Glomospira charoides (Jones and Parker, 1860)
Gyroidinoides altiformis (Stewart and Stewart, 1930)
Gyroidinoides polius (Phleger and Parker, 1951)
Gyroidinoides regularis (Phleger and Parker, 1951)
Gyroidinoides sp 1
Gyroidinoides sp 2
Hoeglundina elegans (d'Orbigny, 1826)
Ioanella tumidula (Brady, 1884)
Neocorbina sp
Neocrosbyia minuta (Parker, 1954)
Nonionella sp
Nuttalides decorata (Phleger and Parker, 1951)
Oridorsalis sp 1
Osangularia culter (Parker and Jones, 1865)
Planulina exorna Phleger and Parker, 1951
Pullenia subspherica Parr, 1950
Pullenia sp
Quinqueloculina venusta Karrer, 1868
Quinqueloculina sp 7
Sphaeroidina bulloides d'Orbigny, 1826
Uvigerina laevis Goës, 1896
Uvigerina peregrina Cushman, 1923

Appendix 2. List of species used in down-core factor analysis.

Alabaminella turgida (Phleger and Parker, 1951)
Alliatina sp 1
Biloculinella irregularis (d'Orbigny, 1839)
Bolivina lowmani Phleger and Parker, 1951
Bolivina spp
Bulimina aculeata d'Orbigny, 1826
Bulimina alazanensis Cushman, 1927
Bulimina mexicana Cushman, 1922
Cibicides wuellerstorfi (Schwager, 1866)
Cibicidoides mundulus (Brady, Parker and Jones, 1888)
Cibicidoides sp
Cornuloculina inconstans (Brady, 1879)
Coryphostoma subspinescens (Cushman, 1922)
Eggerella bradyi (Cushman, 1911)
Epistominella exigua (Brady, 1884)
Epistominella vitrea Parker, Phleger and Pearson, 1953
Francesita advena (Cushman, 1922)
Globocassidulina subglobosa (Brady, 1881)
Globocassidulina cf. *C. subglobosa*
Glomospira charoides (Jones and Parker, 1860)
Gyroidinoides polius (Phleger and Parker, 1951)
Gyroidinoides sp 1
Gyroidinoides sp 2
Hoeglundina elegans (d'Orbigny, 1826)
Ioanella tumidula (Brady, 1884)
Melonis pompilioides (Fichtel and Moll, 1798)
Nuttalides decorata (Phleger and Parker, 1951)
Oridorsalis umbonatus (Reuss, 1885)
Oridorsalis sp 1
Osangularia culter (Parker and Jones, 1865)
Pullenia subspherica, Parr, 1950
Pullenia sp
Pyrgo nasuta Cushman, 1935
Quinqueloculina bosciana d'Orbigny, 1839
Quinqueloculina venusta Karrer, 1868
Quinqueloculina sp 7
Quinqueloculina sp 8
Quinqueloculina spp
Robertinoides bradyi (Cushman and Parker, 1936)
Uvigerina peregrina dirupta Todd, 1948