Northern Hemisphere control of deglacial vegetation
changes in the Rufiji uplands (Tanzania)

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Abstract

In tropical Eastern Africa, vegetation distribution is largely controlled by regional hydrology which has varied over the past 20,000 years. Therefore, accurate reconstructions of past vegetation and hydrological changes are crucial to better understand climate variability in the tropical southeastern African region. We present high-resolution pollen records from a marine sediment core recovered offshore the Rufiji River. Our data document significant shifts in pollen assemblages during the last deglaciation identifying, through respective changes in both upland and lowland vegetation, specific responses of plant communities to atmospheric (precipitation) and coastal (coastal dynamics/sea level changes) alterations. Specifically, arid conditions reflected by maximum pollen representation of dry and open vegetation occurred during the Northern Hemisphere cold Heinrich event 1 (H1) suggesting the expansion of drier upland vegetation to be synchronous with cold northern hemisphere conditions. This arid period is followed by an interval in which forest and humid woodlands expanded, indicating a hydrologic shift towards more humid conditions. Droughts during H1 and the shift to humid conditions around 14.8 kyr BP in the uplands are consistent with latitudinal shifts of the Intertropical Convergence Zone (ITCZ) driven by high-latitude Northern
Hemisphere climatic fluctuations. Additionally, our results show that the lowland vegetation, consisting of well developed salt marshes and mangroves in a successional pattern typical for vegetation occurring in intertidal habitats, has responded mainly to local coastal dynamics related to marine inundation frequencies and soil salinity in the Rufiji Delta as well as the local moisture availability. Lowland vegetation shows a substantial expansion of mangrove trees after ~14.8 kyr BP suggesting an increased moisture availability and river runoff in the coastal area. The results of this study highlight the de-coupled climatic and environmental processes to which the vegetation in the uplands and the Rufiji Delta has responded during the last deglaciation.

1. Introduction

The African tropics, a region of major importance for the global hydrologic cycle, have experienced large-scale changes in hydroclimate and rainfall over the last deglaciation and the Holocene (e.g. Street-Perrot and Perrot, 1990; Lézine et al., 1995; Gasse, 2000; Gasse et al., 2008; Johnson et al., 2002; Vincens et al., 2005; Castañeda et al., 2007; Tierney et al., 2008; Schefuß et al., 2011; Stager et al., 2011; Bouimetarhan et al., 2009, 2012, 2013; Ivory et al., 2012). While millennial-scale hydroclimatic variations in Northwest Africa are commonly linked to atmospheric processes involving latitudinal migrations of the Intertropical Convergence Zone (ITCZ) related to North Atlantic climate anomalies (Dahl et al., 2005; Stouffer et al., 2006; Tjallingii et al., 2008; Mulitza et al., 2008; Itambi et al., 2009, Penaud et al., 2010; Bouimetarhan et al., 2012; Kageyama et al., 2013), the mechanisms responsible for tropical southeastern African climate fluctuations remain a matter of debate. Whereas Indian Ocean sea surface temperatures (SST) have been suggested to influence East African rainfall variability on longer timescales (Tierney et al., 2008, 2013; Tierney and deMenocal, 2013; Stager et al., 2011), other studies suggest that East African rainfall variations were atmospherically linked to North Atlantic climate fluctuations through a southward shift of the ITCZ (Johnson et al., 2002; Broccoli et al., 2006; Brown et al., 2007; Castañeda et al., 2007; Schefuß et al., 2011; Chiang and Friedman, 2012; Mohtadi et al., 2014).
On interannual timescales, the Indian Ocean Dipole (IOD) has been shown to influence modern East African rainfall variability (Saji et al., 1999; Saji and Yamagata, 2003). The El Niño-Southern Oscillation (ENSO) has also been invoked to explain extreme rainfall variability over modern East Africa (e.g. Nicholson, 1996; Plisnier et al., 2000; Indeje et al., 2000; Kijazi & Reason, 2005). As the distribution of tropical African vegetation is largely controlled by regional hydrology, past climate changes are commonly associated with reorganizations of biomes (Gasse et al., 2008; Dupont, 2011). Therefore, understanding the response of vegetation to climate change is crucial for a meaningful assessment of possible forcing mechanisms. Today, most evidence of tropical Eastern African vegetation changes during the last 25,000 years derives from pollen records with the majority reconstructed from continental archives (Gasse, 2000; Vincens et al., 2005; Garcin et al., 2006, 2007; Ivory et al., 2012). These archives have provided explicit evidences of environmental and vegetation changes. However, it appears that the response of southeast African tropical ecosystems to climatic fluctuations during the last deglaciation varied geographically and no definitive consensus has been reached on defining which climatic pattern was causing tropical southeast African vegetation changes. While terrestrial records register, in most cases, a local signal of continental climate conditions through changes in vegetation cover, marine pollen records might, given they have sufficient temporal resolution to resolve millennial-scale climate oscillations, provide a signal integrating a much larger region. Complementary to terrestrial paleorecords from the region, we present new palynological evidence from a marine core offshore the Rufiji River that provides detailed vegetation reconstructions in the Rufiji catchment (Southern Tanzania, SE Africa) during the last deglaciation and more insights into the timing of arid and humid phases in a regional context and their connection to global climate. Furthermore, except for few studies that investigated Holocene mangrove ecosystems in the Tanzanian coast (Punwong et al., 2013 a, b, c), this is the first study from the marine realm that emphasizes the ecological implications of intertidal tropical ecosystems in this area, which are known to be very sensitive to environmental changes at the sea-continent interface. We present detailed information on the development of intertidal plant communities, through a high resolution reconstruction of sensitive salt marsh and mangrove communities during the last deglaciation. We link
them to the intertidal conditions in the Rufiji Delta, such as river runoff and soil salinity, which are influenced by marine inundation frequencies, sea level changes, and coastal moisture. The present study allows to discern, specific responses of plant communities to oceanic (marine inundations/sea level changes) alterations in the Rufiji Delta and to atmospheric (rainfall) changes in the uplands underlying the local and regional mechanisms which control the observed patterns of tropical southeast African vegetation.

2. Regional setting and background

The Rufiji River, formed by the convergence of three principal tributaries, Kilombero, Luwegu and the Great Ruaha located in the high elevations (750 to 1900 m) of the East African Rift (Temple and Sundborg, 1972; Sokile et al., 2003), lies entirely within Tanzania (Fig. 1). With a mean annual discharge of ~ 30 x 10^9 m^3 and a catchment basin area of ~ 174,846 km^2, the Rufiji forms the second largest delta in eastern Africa after the Zambezi (Temple and Sundborg, 1972). The north-south extent of the Rufiji Delta along the eastern Tanzanian coast is ~65 km and comprises largely undisturbed saline swamps, tidal marshes and woodlands (Temple and Sundborg, 1972). The delta contains the largest estuarine mangrove forest in East Africa with a total area of 53,000 ha (Masalu, 2003) found along shorelines and tidal channels that are protected from high-energy wave action and periodically flooded by seawater. Typical mangrove species in the delta include *Avicenna marina*, *Ceriops tagal* and *Rhizophora mucronata* (Masalu, 2003).

The climate of Tanzania is tropical and particularly sensitive to the seasonal migration of the ITCZ. As such, the northern part experiences a bimodal rainfall regime with a long rainy season from March to May and a short rainy season from October to December (e.g. Nicholson, 1996, 2000; Indeje et al., 2000). In contrast, the southern regions of Tanzania (8-12°S), that contain the major part of the Rufiji catchment and the southern uplands, experience tropical summer rainfall with a single well defined rainy season that lasts from November to April (Temple and Sundborg, 1972; Kijazi and Reason, 2005). The dry season occurs during May-October and is dominated by the southeasterly trade winds (Fig. 2) (Walter and Lieth, 1960-1967; Griffiths, 1972; Nicholson et al., 1988).
This seasonality results in strong precipitation gradients that have a clear influence on plant distribution.

The vegetation distribution of tropical Africa is controlled mainly by rainfall and its seasonality although temperature is also an important controlling factor at high altitudes (White, 1983; Hély et al., 2006). In Southeast Africa, the vegetation is very diverse, representing different communities ranging from Somali-Masai deciduous bushland/wooded grassland to Zambezan woodlands and includes closed forest, dry scrubland, alpine open grassland and semi-evergreen lowland forest (Fig. 1) (White, 1983). The Somali-Masai semi-desert grassland and shrublands are dominated by Acacia, Boscia, Asteraceae, Artemisia, Euphorbia, Indigofera and Tamarindus. The Zambezan humid woodland dominated by Uapaca, Brachystegia, and Isoberlina, is mainly well developed in the low to mid-altitudes. These woodlands are replaced by Afromontane communities above 1800-2000 m altitude and vary from montane forests to montane grasslands depending on rainfall. In the lowlands, flooded grasslands host an important community of Cyperaceae and Typha. Many species of fern and halophytes are common along rivers and streams. Halophytes grow on saline soils in intertidal areas, lagoons and depressions as well as salt-lake shores. They are frequently found in arid and semi-arid regions where rainfall is insufficient to remove salt from soils. Halophytic plant communities in SE Africa are mainly dominated by Amaranthaceae, grasses and some species of Cyperaceae (Kindt et al., 2011).

3. Material and methods

3.1. Gravity core GeoB12624-1

We studied marine sediment core GeoB12624-1 (8°14.05’S, 39°45.16’E), recovered off the Rufiji Delta in the Western Indian Ocean at ~655 m water depth during R/V Meteor cruise M75-2 (Savoye et al., 2013). The 600 cm-long core consists of dark olive-gray mud. Generally, the regional wind system is dominated by northeasterly and southeasterly trade winds, which are not favorable for transporting palynomorphs from the continent to the Indian Ocean. Therefore, since the core location is close to the coast
and the mouth of the Rufiji River, we expect the pollen and spores to be mostly delivered
by fluvial transport.

### 3.2. Radiocarbon dating

The GeoB12624-1 age model is based on 7 accelerator mass spectrometry (AMS)
radiocarbon ages, measured on mixed samples of planktonic foraminifera at the Poznań
Radiocarbon Laboratory (Poland) and the National Ocean Sciences AMS Facility in
Woods Hole (USA). Conventional radiocarbon ages were converted to calendar ages with
CALIB 6.11 software, using 1σ age ranges (Stuiver and Reimer, 1993) and the marine 09
calibration (Reimer et al., 2009) with a constant reservoir correction of 140 years (±25 yr)
(Southon et al., 2002). Sediment ages between dated core depths were estimated by linear
interpolation.

### 3.3. X-ray fluorescence (XRF) scanning

XRF Core Scanner II (AVAATECH Serial No. 2) data were collected from the surface of
the archive half of core GeoB12624-1 at the MARUM - University of Bremen (Germany)
every 2 cm down core over a 1.2 cm² area with 10 mm down core slit size, generator
settings of 10 kV, a current of 350 µA, and a sampling time of 30 seconds. The split core
surface was covered with a 4 µm SPEXCert Prep Ultralene1 foil to avoid XRF scanner
contamination and desiccation of the sediment. The reported data were acquired with a
Canberra X-PIPS Detector (SDD; Model SXP 5C-200-1500) with 200eV X-ray
resolution, the Canberra Digital Spectrum Analyzer DAS 1000, and an Oxford
Instruments 50W XTF5011 X-Ray tube with rhodium (Rh) target material. Raw data
spectra were processed by the analysis of X-ray spectra by Iterative Least square software
(WIN AXIL) package from Canberra Eurisys.

The elements Fe, Al, Ba and Ca were measured, but only concentrations of Al and Ca
were used for this study. Ca mainly reflects the marine biogenic carbonate content
whereas Al is related to siliciclastic sedimentary components and varies directly with the
terrigenous fraction of the sediment (e.g. Govin et al., 2014). The Al/Ca ratio therefore
serves as an indicator of the ratio between terrigenous and marine material. High Al/Ca ratios correspond to increased terrigenous input.

3.4. Palynological analysis

In total, 54 sediment samples were prepared for palynological analysis using standard laboratory procedures (Faegri and Iversen, 1989). Sediment (4 cm³) was decalcified with diluted HCl (10%), and then treated with HF (40%) to remove silicates. One tablet of exotic *Lycopodium* spores (18,583±1708 spores/tablet) was added to the samples during the decalcification process in order to calculate palynomorph concentrations per volume of sediment and accumulation rates. After chemical treatment, samples were sieved over an 8 µm nylon mesh screen using an ultrasonic bath (maximum 60 seconds) to disaggregate organic matter. An aliquot (40-60 µl) was mounted on a permanent glass slide using glycerin. One to four slides per sample were counted under a Zeiss Axioskope light microscope at 400x and 1000x magnification. Pollen grains were identified following Bonnefille and Riollet (1980), the African Pollen Database (APD) (Vincens et al., 2007a) and the reference collection of the Department of Palynology and Climate Dynamics at the University of Göttingen (Germany). 32 pollen taxa were identified and listed in Table 1. Other microfossils such as fern spores and fresh water algae (*Botryococcus, Cosmarium, Pediastrum, Scenedesmus* and *Staurastrum*) were also counted. Pollen relative abundances are expressed as percentages of total pollen including herbs, shrubs, trees and aquatics throughout the whole manuscript. However, in order to solely identify the signal of taxa from the upland vegetation, pollen of Cyperaceae, Amaranthaceae mangrove and *Typha* have been excluded from the total pollen sum in Fig. 8.

4. Results

4.1. Age model and sedimentation rates

Radiocarbon dates from 7 samples ranging between 2 and 596 cm core depth are presented in Table 2. The time period represented by core GeoB12624-1 ranges from
~19.3 to 2.3 kyr BP (Fig. 3). High sedimentation rates are recorded, with maximum values of 90 cm/kyr between ~11.6-10.2 kyr BP. Minimum values (18 cm/kyr) are seen later during the Holocene (Fig. 3). The upper 8 samples show very low pollen counts and were excluded from the interpretation. Thus, this study focuses on the interval ~19-10 kyr BP.

4.2. Palynomorph concentrations and Al/Ca ratios
Plotting the concentrations of pollen and other palynomorphs shows significant changes of the terrestrial content in the marine sediment (Fig. 4). Pollen concentrations are relatively high throughout the studied sequence with an average of ~24 x 10^2 grains cm^{-3}, varying between ~5 x 10^2 and ~58 x 10^2 grains cm^{-3}. High values are recorded after ~14.8 kyr BP, while low values are recorded mainly between ~16.8-14.8 kyr BP and in the youngest part after ~10.6 kyr BP. Parallel to the increase in pollen concentrations, the Al/Ca ratios increase after ~14.8 kyr BP with a prominent peak between ~11.6-10.6 kyr BP (Fig. 4). Maxima in Al/Ca ratios and pollen concentrations are coeval with higher sedimentation rates and high fresh water algae concentrations.

4.3. Pollen assemblages
The interval between ~19-14.8 kyr BP was marked by the presence of afromontane taxa, such as Podocarpus, Celtis, Olea, and Artemisia, exhibiting higher values at the beginning of the interval, but decreased around ~16.6 kyr BP (Fig. 5). This interval was also characterized by the dominance of Poaceae pollen (up to ~30%) at the beginning. Poaceae pollen maxima were followed by a dominance of Cyperaceae (~60%), which, in turn declined around 16.6 kyr BP when Amaranthaceae pollen increased rapidly up to ~16% along with Asteraceae, Boscia and Acacia. Around 14.8 kyr BP, values of Rhizophora increased rapidly to their maximum of ~30%. This occurred right after the Amaranthaceae pollen maxima and simultaneously with the increase in Al/Ca ratios. In parallel, Uapaca pollen increased remarkably reaching up to ~15% of the assemblage along with other taxa from the forest and humid woodlands, such as Berlinia/Isoberlina,
Sterospermum, Ziziphus and Borreria. Abundances of pollen of the aquatic taxon Typha and fern spores also increased after ~14.8 kyr BP, while pollen percentages of Poaceae and taxa from dry woods and shrubs declined steadily. Afrotropical taxa were still present albeit with lower values than in the older part of the record (Fig. 5).

Between ~12.8-11.6 kyr BP, percentages of Amaranthaceae and Poaceae increased simultaneously with Asteraceae and Boscia representatives of dry woods and shrubs. The decrease in representation of Cyperaceae pollen, Rhizophora, Typha, fern spores, afrotropical and taxa from the forest and humid woodlands occurred during this time interval along with a slight decrease in Al/Ca ratios. Around 11.6 kyr BP, the record was marked by a rapid increase in percentages for Rhizophora, Typha and fern spores followed by a dominance of Cyperaceae pollen which were in turn replaced by percentage maxima of Poaceae and Amaranthaceae by the end of the record (Fig. 5). These changes were concordant with the increase of Al/Ca ratios that peak ~11 kyr BP, only to decrease again at the end of the record.

The terrestrial palynomorph content presented in this study shows that the most abundant pollen are from Poaceae (grasses), Cyperaceae (e.g. sedges), Rhizophora (mangrove tree), and Amaranthaceae (herbs including many species growing in salt marshes and on salty soils) followed by pollen of Podocarpus (yellow wood). The development of these plant communities interacts differently with inherent environmental variability such as soils, topography, and climate. Therefore, our site received an integrated contribution from both the lowland and upland vegetation.

5. Expansion of the salt marshes and mangrove: deglacial ecological implications for lowland vegetation and coastal processes

The pollen record indicates a directional alternation of three pollen families, between ~19 to 14.8 kyr BP, in the following order: Poaceae, Cyperaceae and Amaranthaceae, followed by an increase in mangrove around 14.8 kyr BP (Fig. 6, steps 1 to 4). The former pollen taxa belong to plant families that host the most common representatives of halophytic vegetation in tropical SE Africa (White, 1983; Kindt et al., 2011). Although they inhabit a wide range of environments, their development in this sequence in addition
to the following expansion of mangrove around 14.8 kyr BP suggests a gradational pattern typical of salt marshes occurring in intertidal habitats (between mean sea level and high water spring level) in coastal areas. Therefore, they are considered, due to their proximity to the shoreline, to be affected by marine inundation frequencies and sea level changes and thus to reflect the coastal dynamics in the Rufiji Delta (Blasco et al., 1996; Hogarth et al., 1999). The East African coast located in the southwestern Indian Ocean lies in a “far-field” location (Woodroffe and Horton, 2005) considered to be situated at significant distances from ice sheet melting. This implies that isostatic effects from large ice sheets are considered to be minimal in this area (Punwong et al., 2013a). Therefore, it is justified to compare our high-resolution pollen record with general sea-level reconstructions (Waelbroeck et al., 2002; Rohling et al., 2009). This comparison shows that when sea level was ~80-120 m lower relative to today, the exposed shelf allowed the grass (Poaceae) and sedges (Cyperaceae) to expand (Fig. 6, Fig 7e). The coastline was also substantially closer to the core site when sea level was low (Fig. 1). During the subsequent sea-level rise, only pioneer species from the Amaranthaceae tolerating highly saline environments with a permanent tidal influence and having high colonizing abilities could expand under these stressful conditions. The development of mangrove at ~14.8 kyr BP might reflect either the expansion of mangrove vegetation along the Rufiji Delta or the erosion of mangrove peat during sea-level rise (Hooghiemstra and Agwu 1986; Dupont and Agwu, 1991; Lézine et al., 1995; Lézine, 1996; Dupont, 1999; Kim et al., 2005; Scourse et al., 2005). Mangroves are most common in wetter habitats and swamps where brackish water accumulates. They are known to be very sensitive to sea-level fluctuations and runoff variability (Hooghiemstra and Agwu, 1986; Dupont and Agwu, 1991; Lézine et al., 1995; Lézine, 1996; Woodroffe, 1999). Their development would suggest a permanent marine influence, but also less saline coastal environments as they do not survive in hypersaline soils due to the rapid sea-level increase (Woodroffe, 1999). Consequently, the expansion of mangrove vegetation along the Rufiji Delta in our record, during the period of global sea-level rise (Waelbroeck et al., 2002; Rohling et al., 2009) (Fig. 6), is likely the result of changes in local hydrologic conditions through an increased river runoff promoted by higher moisture availability in the coast after ~14.8 kyr BP. By this means, higher freshwater input and increased sedimentation rates may dominate over
local sea-level rise, suppressing the intrusion of sea water and allowing complex plant
communities to develop on the delta and mangroves to expand landward in response to
increased rainfall over the Rufiji Delta. Our results corroborate previous findings in the
Rufiji Delta and the coast of Zanzibar where dynamics of Holocene mangrove systems
were related to past sea level changes and local moisture availability (Punwong et al.,
2013a, b, and c). Furthermore, the development of Suwayh mangrove near the littoral of
the Indian Ocean in Oman clearly records the influence of enhanced tropical summer
precipitation (Lézine et al., 2010). Increasing both freshwater supply and sediment load
would also fit the development of aquatic taxa such as Typha, which is represented
parallel to the Rhizophora pollen maximum reflecting wetter coastal conditions and
continuous input of freshwater. Therefore, the erosion of mangrove peat during sea-level
rise is less likely because this would imply reduced freshwater flow to the coast and dry
climatic conditions.

Taken together, the succession of salt marshes and mangrove reflects the response of
coastal plant communities to changes in intertidal environments (soil development and
salinity gradient) and coastal dynamics in the Rufiji Delta influenced by sea-level
changes as suggested by González and Dupont (2009). These results add to the scarce
knowledge on the East African coastal vegetation, a major biodiversity hotspot in the area
(Myers, 2000), and provide an independent evidence on the close relationship between
sea level changes and coastal community dynamics. In this context, our new
palynological record has great ecological implications as it deals with sensitive
ecosystems that are poorly documented on longer timescales.

6. Paleoclimate and controlling mechanisms in the uplands during H1

The total pollen assemblage is dominated by afromontane forest taxa in the earliest part
of the record until ~16.6 kyr BP (Fig. 7c). Afromontane forest mainly developed in
mountains favoured by cold and humid conditions (White, 1983, Kindt et al., 2011).
Their presence in the pollen record would thus be expected if the afromontane forest had
spread to lower altitudes than currently found and its pollen did not need to be transported
over long distances. Therefore, the high pollen abundances of the afromontane forest in
the marine pollen record corroborates previous pollen records that suggest the development of afromontane taxa at a lower elevation (Vincens et al., 2007b, Ivory et al., 2012) due to freezing conditions at higher altitudes, cooler conditions at lower altitudes, and lower $\rho CO_2$ (Street-Perott et al., 1997; Wu et al., 2007). During the decline of the afromontane taxa, the pollen representatives of dry wood and shrub vegetation increase significantly between ~16.6-14.8 kyr BP (Fig. 7b). This transition suggests a change towards drier conditions compared to the previous period and coincides with the timing of the North Atlantic H1 (Hemming, 2004; Stanford et al., 2011 (H1 sensu stricto)).

Around 14.8 kyr BP, the vegetation cover became denser. The decline of elements from dry woods and shrubs and the drastic decrease in afromontane forest was followed by an increase in pollen from forest and humid woodlands (Fig. 7a). A similar vegetation trend has been recorded in several pollen records from Lakes Malawi, Tanganyika, Rukwa and Masoko, indicating the retreat of the afromontane vegetation to higher altitudes due to progressive warming after H1 and the expansion of moist forest due to enhanced rainfall (Vincens, 1993; Vincens et al., 2005; 2007b; Ivory et al., 2012).

Between ~12.8-11.6 kyr BP, the presence of elements from both the forest and humid woodland vegetation and from dry woods and shrubs (Figs. 7a, b) suggests that vegetation was more heterogeneous. In contrast to other records from most of the African tropics (Gasse, 2000; Barker et al., 2007; Mulitza et al., 2008; Tierney et al., 2008; Junginger et al., 2014) where indicators of aridity have been observed during this time interval coincident with the YD (YD, 12.8 – 11.5 kyr BP) (Alley, 2000; Muscheler et al., 2008), our records do not show a clear climatic trend.

Around 11.6 kyr BP, sharply rising Al/Ca ratios and high sedimentation rates along with the presence of pollen from forest and humid woodlands would indicate increased precipitation. However, the decline of nearly all the pollen taxa percentages, Al/Ca ratios and sedimentation rates at the end of the record, around 10.6 kyr BP, reflects either a return to drier conditions or the end of active terrestrial input.

In sum, our data show that during H1 upland vegetation changed from afromontane forest to dry woods and shrubs (Fig. 7b and c). Forest and humid woodlands developed after ~14.8 kyr BP and continued to expand through the YD (Fig. 7a).
If we exclude the dominant pollen taxa (salt marshes and mangrove) from the total sum, dry woods and shrubs still show a substantial expansion during H1 as we can see in Fig. 8b. This, together with the sharply reduced Al/Ca ratios indicate increased aridity in the uplands during H1. The direct comparison of our record with terrestrial studies, shows that the signal of decreased precipitation coincides with lowered lake levels of Sacred Sacred Lake in Kenya (Street-Perrot et al., 1997), Lake Challa, Tanzania (Verschuren et al., 2009), Lake Rukwa, Tanzania (Vincens et al., 2005) and Lake Tanganyika (Burnett et al., 2011). Dry H1 conditions are also suggested by isotope records of the Tanganyika basin (Tierney et al., 2008) and Lake Malawi (Johnson et al., 2002; Brown et al., 2007; Castañeda et al., 2007). The expansion of forest and humid woodlands (Fig. 8c) along with higher Al/Ca ratios and sedimentation rates after H1 suggests a significant change in the hydrological regime towards enhanced rainfall and increased terrigenous discharge. We thus infer a shift towards more humid conditions. Significant increase in moisture after ~ 14.8 kyr BP has been reported from vegetation records in continental archives (Vincens, 1993; Vincens et al., 2005; 2007b; Ivory et al., 2012) as well as from lake records (Gasse, 2000; Junginger et al., 2014). Taken together, upland aridity during H1 and the increased humidity around 14.8 kyr BP as reconstructed from our records correlate (within age model uncertainties) with changes inferred from continental archives that show a similar pattern in most of the tropical eastern and south-eastern African lakes and are in agreement with northwest tropical African records (e.g. Hooghiemstra, 1988; Zhao et al., 2000; Mulitza et al., 2008; Itambi et al., 2009; Niedermeyer et al., 2009; Bouimetarhan et al., 2012, 2013).

For the tropical eastern African region where different processes can affect rainfall, several mechanisms have been proposed. Today, the IOD influences East African precipitation at the interannual timescale (Saji and Yamagata, 2003). However, recent hydrological records from the eastern equatorial Indian Ocean (Mohtadi et al., 2014) suggest similarly dry conditions during H1 and YD, ruling out a zonal IOD-like dipole structure between Indonesia and the eastern African lakes that was suggested earlier by Tierney et al. (2008). Many studies have proposed ENSO as an important driver of extreme rainfall anomalies over East Africa (e.g. Nicholson, 1996; Plisnier et al., 2000; Indeje et al., 2000). However, evidence for an El Niño- or La Niña-biased mean climate
state during H1 is ambiguous (Leduc et al., 2009; Prange et al., 2010). Moreover, it has recently been shown that the impact of the tropical Pacific on East African rainfall disappears on multidecadal and perhaps longer timescales (Tierney et al., 2013). We therefore suggest that an ENSO-like impact over southern Tanzania and hence the major portion of the Rufiji catchment area was not the main mechanism for the H1 drought. Results from climate model studies suggest a north-south anti-phase relation in African annual precipitation in response to North Atlantic cooling, consistent with latitudinal migrations of the ITCZ’s annual mean position (e.g., Lewis et al., 2010; Kageyama et al., 2013). In line with this hypothesis, the arid phase recorded in our data during H1 has (within age model uncertainties) a pronounced wet counterpart in the Zambezi region (Schefuß et al., 2011; Otto-Bliesner et al., 2014). Therefore, we suggest the observed H1 dry conditions in the uplands to be part of a north-south dipole rainfall anomaly over East Africa and the Indian Ocean corroborating the see-saw hypothesis supported by further climate model studies (Claussen et al., 2003) and which is consistent with a southward shift of the ITCZ annual mean position in response to Northern Hemisphere cooling (Mohtadi et al., 2014). The ITCZ shift is part of a reorganization of the annual mean Hadley circulation driven by Northern Hemisphere climatic fluctuations (Broccoli et al., 2006; Kang et al., 2009; Chiang and Friedman, 2012; Frierson et al., 2013) and is supported by several studies in the Indian Ocean realm (Johnson et al., 2002; Brown et al., 2007; Castañeda et al., 2007; Schefuß et al., 2011; Mohtadi et al., 2014). We suggest that the reorganization of the Hadley circulation and the associated southward ITCZ shift resulted in anomalous descent of air over the Rufiji region in the annual mean (and hence less rainfall), and anomalous ascent (and hence more rainfall) to the south. The modern seasonality of East African rainfall (Fig. 2) indicates that a southward shift of the ITCZ-related rainbelt (by a few degrees) would lead to significantly drier conditions associated with stronger surface northeasterlies in the Rufiji region, only during the austral summer season (DJF).

Alternatively, Indian Ocean sea surface temperatures (SSTs) might also play a role in influencing SE African hydrology and vegetation. Cooler SSTs during millennial-scale stadials would have reduced moisture transport from the Indian Ocean implying a reduction of monsoonal precipitation. Therefore, dry conditions during cold stadials have
been suggested to relate to low Indian Ocean SSTs (Tierney et al., 2008; Stager et al., 2011). Lower SSTs in the Indian Ocean have been proposed as a potential mechanism for extreme droughts in SE Africa during H1 as they would tend to reduce the evaporative moisture content of the ITCZ (Stager et al., 2011). However, Mg/Ca reconstructed SSTs from the nearby core GeoB12615-4 (7°08.30’S, 39°50.45’) in the western Indian Ocean show warming during H1 (Romahn et al., 2014), such that we rule out a dominant effect of Indian Ocean SST forcing on H1 aridity in the southern uplands of Tanzania.

7. Environmental changes during the YD

The prominent decrease in precipitation that we infer for H1 is however not recorded during YD. The vegetation reconstructions in our record show an alternation between humid and dry taxa during YD (Figs. 8b and 8c). This pattern reflects no clear climatic trend, while most records from the African tropics suggest drier conditions during YD (Gasse, 2000; Barker et al., 2007; Mulitza et al., 2008; Tierney et al., 2008; Junginger et al., 2014). In addition, marine records from the northern Indian Ocean realm have also shown dry conditions during YD as a response to a southward shift of the ITCZ (Mohtadi et al., 2014). However, two vegetation records from adjacent locations in tropical East Africa highlight different regional responses during the YD. Lake Masoko, a small lake within the Lake Malawi watershed, recorded an expansion of tropical seasonal forest during YD reflecting humid conditions (Garcin et al., 2006, 2007). In contrast, a record from Lake Malawi shows YD to occur in two phases progressing in a dry-to-wet pattern (Ivory et al., 2012) reflecting a more southerly ITCZ associated with an increase in rainfall seasonality (Ivory et al., 2012). Those differences in environmental responses to the YD are consistent with the heterogeneous vegetation observed in our record suggesting that the YD signal from this area is ambiguous which corroborates previous findings in the Indo-Pacific Warm Pool (Denniston et al., 2013; Dubois et al., 2014) where the YD is not well defined either. Therefore, our data suggest that H1 had a greater influence on East African hydrologic conditions than the YD, another North Atlantic cold event that likely, due to its shorter duration and weaker Northern Hemisphere cooling compared to H1, did not displace the annual mean ITCZ as far south as H1, thus causing these ambiguous signals. In addition, it has recently been suggested that gradually increasing greenhouse-gas forcing through the last glacial termination resulted in
increasingly wetter conditions in tropical Africa (Otto-Bliesner et al., 2014), leading to generally higher precipitation in the Rufiji region during the later stages of the deglaciation compared to H1.

8. Conclusions

The marine pollen record off the Rufiji River provides new information on the deglacial vegetation history and hydrologic variability in SE Africa. The upland versus lowland vegetation records allow to discern ecosystem responses to different environmental changes related to oceanic (coastal dynamics) and atmospheric (precipitation) alterations. The upland vegetation shows drier conditions during the Northern Hemisphere cold H1, with a shift to more humid conditions around 14.8 kyr BP inferred from the expansion of forest and humid woodlands. The lowland (coastal) vegetation shows a well-established salt marsh vegetation and mangroves along the Rufiji Delta throughout the whole record with a substantial expansion of mangroves after ~14.8 kyr BP as a positive reaction to higher moisture availability in the coastal area.

The observed H1 aridity in the uplands is consistent with a southward displacement of the annual mean ITCZ driven by high-latitude climate changes in the Northern Hemisphere. This finding suggests that the extension and composition of plant assemblages in the upland during H1 is primarily controlled by Northern Hemisphere climatic fluctuations corroborating previous studies from SE Africa and the Indian Ocean realm that evidenced the response of the regional hydrologic system to millennial-scale North Atlantic cold periods. Additionally, the coastal dynamics in the Rufiji Delta related to fluctuations in the sea level and available local moisture have played a major role in modulating the local coastal plant community by favoring/reducing the expansion of salt marsh vegetation and mangroves. Our new palynological record has a great ecological significance, as much as it deals with intertidal ecosystems that have not been intensively studied. It offers an important complement to previously published paleorecords from the region and highlights the contrasting processes to which upland and lowland vegetation have responded.
Acknowledgments

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Marine and terrestrial biomarker records for the last 35,000 years at ODP site 658C off
Table 1: List of identified pollen taxa in marine core GeoB12624-1. Taxa are grouped according to their phytogeographical assignment.

<table>
<thead>
<tr>
<th>Pollen type</th>
<th>Family</th>
</tr>
</thead>
<tbody>
<tr>
<td>Poaceae</td>
<td></td>
</tr>
<tr>
<td>Cyperaceae</td>
<td></td>
</tr>
<tr>
<td>Amaranthaceae (includes Chenopodiaceae)</td>
<td></td>
</tr>
</tbody>
</table>

**Dry woodlands and shrubs**

_Acacia_ Fabaceae-Mimosoideae

_Mimosa-type_ Fabaceae-Mimosoideae

_Boscia_ Capparaceae

_Asteroideae species_ Asteraceae

_Combretaceae_ Combretaceae

_Indigofera-type_ Fabaceae-Faboideae

_Caryophyllaceae_ Caryophyllaceae

_Plantago_ Plantaginaceae

_Tamarindus-type_ Fabaceae

_Artemisia_ Asteraceae

_Afromontane_

_Podocarpus_ Podocarpaceae

_Olea_ Oleaceae

_Celtis_ Cannabaceae

**Forest and humid woodlands**

_Uapaca_ Phyllanthaceae

_Psydrax type subcordatum_ Rubiaceae
<table>
<thead>
<tr>
<th>Species</th>
<th>Family</th>
</tr>
</thead>
<tbody>
<tr>
<td>Berlinia/Isoberlina</td>
<td>Fabaceae</td>
</tr>
<tr>
<td>Stereospermum-type</td>
<td>Bignoniaceae</td>
</tr>
<tr>
<td>Ziziphus-type</td>
<td>Rhamnaceae</td>
</tr>
<tr>
<td>Vernonia</td>
<td>Asteraceae</td>
</tr>
<tr>
<td>Alchornea</td>
<td>Euphorbiaceae</td>
</tr>
<tr>
<td>Cassia-type</td>
<td>Fabaceae</td>
</tr>
<tr>
<td>Cleome</td>
<td>Capparaceae</td>
</tr>
<tr>
<td>Borreria (=Spermacoce)</td>
<td>Rubiaceae</td>
</tr>
<tr>
<td>Pterocarpus-type</td>
<td>Fabaceae-Faboideae</td>
</tr>
<tr>
<td>Piliostigma</td>
<td>Fabaceae</td>
</tr>
<tr>
<td>Rhus-type</td>
<td>Anacardiaceae</td>
</tr>
</tbody>
</table>

**Mangrove trees**

<table>
<thead>
<tr>
<th>Species</th>
<th>Family</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rhizophora</td>
<td>Rhizophoraceae</td>
</tr>
</tbody>
</table>

**Bog vegetation and swamp plants**

<table>
<thead>
<tr>
<th>Species</th>
<th>Family</th>
</tr>
</thead>
<tbody>
<tr>
<td>Typha</td>
<td>Typhaceae</td>
</tr>
</tbody>
</table>

**Other elements**

<table>
<thead>
<tr>
<th>Species</th>
<th>Family</th>
</tr>
</thead>
<tbody>
<tr>
<td>Euphorbia</td>
<td>Euphorbiaceae</td>
</tr>
</tbody>
</table>
**Table 2**: Conventional radiocarbon age and mode values of calibrated dates for marine core GeoB12624-1. For reservoir corrections a constant ΔR of 140 ± 25 yrs has been applied to all dates (Southon et al., 2002).

<table>
<thead>
<tr>
<th>Core depth (cm)</th>
<th>Lab Code</th>
<th>$^{14}$C age ± age error (yr BP)</th>
<th>1σ calendar age ranges (yr BP)</th>
<th>Calibrated age (cal. yr BP)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Poz-30420</td>
<td>2810 ± 35</td>
<td>2308 - 2419</td>
<td>2340 (+79/-32)</td>
</tr>
<tr>
<td>124</td>
<td>Poz-47931</td>
<td>8680 ± 50</td>
<td>9091 - 9265</td>
<td>9178 (+87/-87)</td>
</tr>
<tr>
<td>210</td>
<td>OS-79104</td>
<td>9540 ± 65</td>
<td>10172 - 10332</td>
<td>10223 (+109/-51)</td>
</tr>
<tr>
<td>300</td>
<td>Poz-47932</td>
<td>10410 ± 60</td>
<td>11184 - 11312</td>
<td>11212 (+100/-28)</td>
</tr>
<tr>
<td>398</td>
<td>Poz-47933</td>
<td>11240 ± 60</td>
<td>12564 - 12664</td>
<td>12610 (+54/-46)</td>
</tr>
<tr>
<td>512</td>
<td>Poz-47934</td>
<td>13200 ± 70</td>
<td>14781 - 15116</td>
<td>15040 (+126/-259)</td>
</tr>
<tr>
<td>596</td>
<td>Poz-30421</td>
<td>16630 ± 80</td>
<td>19244 - 19417</td>
<td>19380 (+37/-136)</td>
</tr>
</tbody>
</table>
Figure 1. (a): Map of Southern Africa showing the location of marine sediment core GeoB12624-1, simplified phytogeography and modern vegetation after White (1983) and approximate position of the ITCZ during austral summer (December, January, February). Indicated are: the main course of Rufiji River, Zambezi River, and Limpopo River (blue lines), major lakes in the area and the outline of the Rufiji catchment in white. Other cores discussed in the text are also illustrated: GeoB9307-3 (Schefuß et al., 2011), GeoB12615-4 (Romahn et al., 2014). (b): Bathymetric map of the study area showing the location of marine sediment core GeoB12624-1 and the Rufiji Delta.
Figure 2. Modern atmospheric circulations over Africa: surface winds (m/s) (Kalnay et al., 1996) and precipitation (cm/month) (Adler et al., 2003) are illustrated during austral summer (DJF: December, January, February), autumn (MAM: March, April, May), winter (JJA: June, July, August) and spring (SON: September, October, November). The red dot denotes the location of marine sediment core GeoB12624-1.
**Figure 3.** Calibrated age-depth relation for core GeoB12624-1 (bars indicate the 1σ error range (yr BP)) and sedimentation rates (cm/kyr) (orange line).
**Figure 4.** Downcore variations of pollen concentrations, freshwater algae concentrations, Al/Ca ratios and sedimentation rate estimates during the interval 19-10 kyr BP. Shading indicates time intervals of Heinrich event 1 (H1) and the Younger Dryas (YD).
Figure 5. Palynological data from marine sediment core GeoB12624-1 showing relative abundances (%) of selected pollen taxa, percentages of fern spores and the total pollen and spores counts. Note scale changes on x-axes. Shading indicates time intervals of Heinrich event 1 (H1) and the Younger Dryas (YD). Triangles indicate age control points.
Figure 6. Comparison of the pollen record from marine core GeoB12624-1 with sea-level reconstructions: dark blue from Waelbroeck et al. (2002) and light blue from Rohling et al. (2009). Pollen percentages of Poaceae, Cyperaceae, Amaranthaceae indicates the succession of salt marshes (steps 1 to 3) and the mangrove forest (step 4) along the Rufiji Delta. Dashed lines denote the four steps of the directional alternation of those families.
Figure 7. Palynological data showing relative abundances of major pollen groups based on the total sum of pollen and spores. (a): pollen percentages of forest and humid woodlands, (b): pollen percentages of dry woods and shrubs, (c): afromontane taxa percentages pollen, (d): percentages of salt marshes (Cyperceae and Amaranthaceae), (e): Mangrove-pollen percentages. Shadings indicate the 95% confidence interval. Dashed lines denote time intervals of Heinrich event 1 (H1) and the Younger Dryas (YD). Triangles indicate age control points.
Figure 8. Palynological data showing relative abundances of (a): Grass-pollen percentages, (b): pollen percentages of dry woods and shrubs and (c): pollen percentages of forest and humid woodlands based on the sum of pollen and spores excluding Cyperaceae, Amaranthaceae, mangrove and Typha (aquatic pollen). Shadings indicate the 95% confidence interval. Dashed lines denote time intervals of Heinrich event 1 (H1) and the Younger Dryas (YD). Triangles indicate age control points.