Modern seasonality in Lake Challa (Kenya/Tanzania) and its sedimentary documentation in recent lake sediments

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Abstract

From November 2006 to January 2010, a sediment trap that was cleared monthly was deployed in Lake Challa, a deep stratified freshwater lake on the eastern slope of Mt. Kilimanjaro in southern Kenya. Geochemical data from sediment trap samples were compared with a broad range of limnological and meteorological parameters to characterize the effect of single parameters on productivity and sedimentation processes in the crater basin. During the southern hemisphere summer (November–March), when the water temperature is high and the lake is biologically productive (nondiatom algae), calcite predominated in the sediment trap samples. During the “long rain” season (March–May) a small amount of organic matter and lithogenic material caused by rainfall appeared. This was followed by the cool and windy months of the southern hemisphere winter (June–October) when diatoms were the main component, indicating a diatom bloom initiated by improvement of nutrient availability related to upwelling processes. The sediment trap data support the hypothesis that the light–dark lamination couplets, which are abundant in Lake Challa cores, reflect seasonal delivery to the sediments of diatom-rich particulates during the windy months and diatom-poor material during the wet season. However, interannual and spatial variability in upwelling and productivity patterns, as well as El Niño–Southern Oscillation (ENSO)-related rainfall and drought cycles, exert a strong influence on the magnitude and geochemical composition of particle export to the hypolimnion of Lake Challa.

The influence of global warming on freshwater availability in Africa is attracting increasing attention (Cullen et al. 2006; Mölg et al. 2008; Olaka et al. 2010). Annually laminated lacustrine sediments provide potential information about changes in the large-scale atmospheric circulation and associated moisture-balance variation (Gasse et al. 1989; Johnson et al. 2002; Wolff et al. 2011). Paleoclimatic reconstructions on the basis of lake sediments require understanding of the lake’s response to the seasonality of environmental parameters, such as temperature and precipitation (Mackay et al. 2005). Monitoring of the modern lake systems provides insight into sediment formation and preservation in the context of seasonal climatic variability (Pilskaln 2004; Bluszcz et al. 2008; Stockhecke et al. 2012). However, the majority of published records on seasonal aspects of lacustrine sediment formation are from temperate region lakes (Ojala et al. 2012). Microfacies and geochemical data extracted from sediment trap and water samples from Lake Challa, a tropical crater lake on the lower eastern flank of Mt. Kilimanjaro, were compared with a broad range of limnological and meteorological parameters to understand the forcing of external factors and the influence of lake internal processes on the formation of seasonal laminae in tropical lake sediment. Our aim is to understand the genesis of the sedimentary record of Lake Challa to provide a reliable mechanistic model for paleoclimatic interpretation of the lake’s sediment record and varve formation.

Study site and modern-day climate—Lake Challa (3°19’ S, 37°42’ E) is a crater lake with steep walls (up to 170 m) on the lower eastern slope of Mt. Kilimanjaro (Fig. 1). Maximum depths varied between 92 m and 98 m in 1999–2010. The lake surface area is approximately 4.5 km², and it is ~ 880 m above sea level. The drainage basin is composed of igneous rocks (predominantly trachy-basalts) of the tertiary Kilimanjaro complex (Bear 1955) covered by “calcareous tuffaceous grits,” a calcite-cemented tuffaceous breccia (Downie and Wilkinson 1972). This volcanic complex is underlain by metamorphic rocks (predominantly gneisses) that outcrop east and south of Lake Challa to the Indian Ocean coast (Petters 1991). A seismic survey detected a sediment infill of ~ 210 m on the bottom of the crater, which accumulated over ~ 250,000 yr (Moernaut et al. 2010).

Evaporation (1700 mm yr⁻¹) exceeds the annual rainfall onto the surface of the lake (600 mm; Payne 1970). Therefore, the water budget of the lake is mostly controlled by subsurface in- and outflow in which the groundwater input is probably fed by seepage of precipitation falling...
further uphill in the forest zone on the slopes of Mt. Kilimanjaro. Local climate is controlled by the seasonal passage of the Inter-Tropical Convergence Zone (ITCZ; Fig. 1), with rainfall occurring predominantly from October to December and March to May (Fig. 2; Nicholson 2000). These two rainy seasons are referred to as the “long rains” (March–May), usually the major rainfall of the year, and the “short rains” (October–December), which are more variable. Long rains and short rains presumably respond to changes in Indian Ocean surface-water temperatures and related forcing on atmospheric circulation (Black et al. 2003; Hastenrath et al. 2004). Furthermore, rainfall in East Africa is linked to the El Niño–Southern Oscillation (ENSO), with more rain during El Niño years and severe droughts in La Niña years (Rojelowiski and Halpert 1987; Nicholson 2000). Temperatures in the region south of the equator are high during northern hemisphere winter (November–March) and decline by about 4°C during the northern hemisphere summer (June–August). In contrast to temperature, seasonal wind speed variations show the

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Fig. 1. (a) Location of Lake Challa (3°19′S, 37°42′E) and (b) bathymetric map (depth contours at 10 m intervals (Moernaut et al. 2010). (a) Black outline indicates coverage area shown in (c). (c) Seasonal migration of the ITCZ over Eastern Africa with northeasterly (NE) and southeasterly (SE) trade winds. Monthly precipitation (shading, mm month$^{-1}$) based on satellite precipitation estimates over ocean areas and rain gauge data over land regions; contours are drawn at 50 mm intervals. Large-scale wind vectors for the 925 hectopascal pressure level indicate wind direction with wind speed proportional to the length of the vectors. Source: http://iridl.ldeo.columbia.edu.

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Fig. 2. (a) Meteorological data from Voi, Kenya (~ 100 km east of Lake Challa). Black line, annual temperature trend; bars, monthly rainfall amount; dashed line, monthly wind speed data; source: monthly station data via the Global Climate Perspective System from the National Climatic Data Center by the National Oceanic and Atmospheric Administration (NOAA NCDC GCPS). (b) Comparison between the 13 months running mean of wind speed and negative precipitation anomalies (13 months running mean) indicates the inverse relationship between rainfall and wind speed on interannual timescales in the region around Lake Challa (4°S–3°S, 37°E–38°E); source: National Centers for Environmental Prediction and National Center for Atmospheric Research (NCEP-NCAR) reanalysis dataset (Kalnay et al. 1996).
Table 1. Hydrochemical water profiles from Lake Challa, 23 Aug 2007 (18:00 h) and 28 Nov 2007 (12:00 h). All units mg L\(^{-1}\).

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<th>Sr</th>
<th>Ba</th>
<th>F</th>
<th>DIC</th>
<th>SO(_4)</th>
<th>SRP</th>
<th>NPOC</th>
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NA, not analyzed; DIC, dissolved inorganic carbon; SRP, soluble reactive phosphorus; NPOC, nonpurgeable dissolved organic carbon; TDS, total dissolved solids; *, average of two water analyses.
Table 2. Chemical composition of local rocks and soils (all collected within 30 km around Lake Challa). Element ratios refer to mass ratios.

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<tr>
<th>Sample description</th>
<th>SiO₂</th>
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<th>Fe₂O₃</th>
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<td>Unweathered grits (Challa)</td>
<td>35.94</td>
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<td>9.70</td>
<td>14.21</td>
<td>0.21</td>
<td>8.53</td>
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<td>0.67</td>
<td>8.28</td>
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<tr>
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<td>17.32</td>
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<td>3.03</td>
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<td>0.21</td>
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</table>

* Ravine (~ 500 m long) enters Lake Challa in northwest corner of caldera and is, according to information from local people, only active in very wet years; fsp, feldspar; element ratios refer to mass ratios.

Methods

Temperature profiles were recorded at 2 h intervals between November 2006 and January 2010 by eight temperature loggers (Minilog, Vemco, Canada) placed (10, 20, 30, 40, 50, 60, 65, 85 m) at a mooring site close to the center of the lake. Water for chemical analyses was sampled along vertical profiles at various depths on 23 August 2007 and 28 November 2007, using a Hydro-Bios Plastic Water Sampler (Niskin type). Water samples (100 mL each) were filtered through 0.2 μm, 25 mm diameter Whatman GD/X polypropylene syringe filters, and 50 mL was acidified with concentrated HNO₃ for mass ratios.

For hydrogen and oxygen isotope composition, following the methods of Eggermont et al. (2007), water samples for stable isotope analyses of oxygen (δ¹⁸O) and deuterium (δD) were collected in August and November 2006 from various water depths (0 to 90 m in 10 m steps) and were stored frozen until further preparation in the GeoForschungsZentrum (GFZ) laboratories in Potsdam, where the solids were separated by filtration, washed with Merck (Suprapur) acids comprised the following steps: (1) dissolution in 2 mL HCl and filling to a final volume of 50 mL was acidified with concentrated HNO₃ for analyses of cations and was stored at 4°C until processing. Chemical analyses closely followed Eggermont et al. (2007). Profiles of temperature, pH, conductivity, and dissolved oxygen were measured using a Hydrolab Quanta conductivity, temperature, depth (CTD) profiler and a multiparameter water quality meter (Yellow Springs Instrument Company 600XL-B-D), respectively.

Water samples for stable isotope analyses of oxygen (δ¹⁸O) and deuterium (δD) were collected in August and November 2007 from various water depths (0 to 90 m in 10 m steps) and were stored in sealed air-tight polypropylene bottles until analysis at the Alfred Wegner Institute for Marine and Polar Research in Potsdam. A Finnigan Delta-S stable isotope ratio mass spectrometer equipped with two equilibration units was used for the determination of hydrogen and oxygen isotope composition, following Meyer et al. (2000). Analytical precision was better than 0.8% for δD and 0.1‰ for δ¹⁸O.

In November 2006, a Uwitec (double-funneled 86 mm diameter) sediment trap was installed below the thermalcline at 35 m water depth close to the center of the lake (core position Fig. 1) to record seasonal changes in primary production, carbonate precipitation, and deposition of silicilastics. The trap was emptied and redeployed every ~ 4 weeks, providing 34 monthly samples between December 2006 and January 2010. After settling for a minimum of 24 h and the decanting of superfusive water, samples were stored frozen until further preparation in the GeoForschungsZentrum (GFZ) laboratories in Potsdam, where the solids were separated by filtration, washed with Merck (Suprapur) acids comprised the following steps: (1) carbonate dissolution and wet oxidation of organic material (HNO₃ 30%, 3 h, 160°C), (2) oxidation of refractory organic compounds (HClO₄ 70%, 5 h, 160°C), (3) silicate dissolution (HF 40%, 2 days, 70°C), (4) evaporation of HF and Si-F compounds (HClO₄ 70%, 2 h, 210°C), and (5) dissolving in 2 mL HCl and filling to a final volume of 14 mL. Four multielement standards in an acid matrix similar to that of the digestion solutions were prepared for external calibration on the basis of commercial single-element standards of Spex and Merck, respectively. The total carbon (TC) and total organic carbon (TOC) analyses were performed using an elemental analyzer (NC2500 Carlo Erba Instruments, Milan, Italy).

Meyer et al. (2000).
Erba). The results were referenced against lab-internal soil standards and a reference sample (urea). Total inorganic carbon (TIC) was determined as the difference between TC and TOC. Saturation indices for calcite, aragonite, and amorphous silica in the water column were calculated using PHREEQC software (Version 2, database: Phreeqc; Parkhurst and Appelo 1999).

Sediment investigations were carried out on a 49 cm long gravity core, taken from the center of the lake. Sedimentary structures were studied under the microscope on thin sections from 10 cm long resin-impregnated blocks (Merkt 1971; Brauer et al. 1999). Sediment components were studied in smear slides, which were prepared with Merck Kaiser's glycerol gelatin for microscopy (melts at 40°C). Scanning electron microscope (SEM) images were performed using a Carl Zeiss Ultra 55 Plus system. Changes in sediment composition and intensities of chemical elements in the top 10 cm of the gravity core, last 100 yr, were studied by micro X-ray fluorescence (μXRF) scanning of the sediment, imbedded in a resin-block. Scanning was performed using an EAGLE III XL μXRF spectrometer (Röntgenanalytik) at GFZ, applying 40 kV tube voltage, 300 μA tube current, 123 μm spot size, and 100 μm step size.

Samples of powdered catchment rocks and soils were mixed and fused with Li tetraborate–metaborate flux (FLUXANA, FX-X65). The prepared glass discs were analyzed using wavelength-dispersive XRF spectrometry (Panalytical Axios Advanced XRF spectrometer). Quantification of element concentrations was achieved using calibration curves generated from multiple certified reference samples.

Results

**Physical and chemical lake water characteristics**—Lake Challa is a tropical hard-water lake with HCO₃ as the dominant anionic component, followed by Cl and SO₄ (Table 1). According to Talling and Talling (1965), this is the dominant (prevailing) type of lake water in Africa. A substantial proportion of the high dissolved inorganic carbon (DIC) inventory of the lake may originate from chemical dissolution of the local calcite-bearing tuffaceous breccia (cf. Table 2; CO₂ contents of unweathered and weathered grit). The DIC : Ca equivalent ratio of the lake water is approximately three times higher than that of fluids derived from carbonate dissolution. This relatively high DIC : Ca may have been generated by inflow of fluids from CO₂-forced chemical silicate weathering and aerobe mineralization of organic matter over time. There is no evidence for active CO₂-rich sublacustrine springs within the lake basin.

Sulfate concentrations in Lake Challa are generally low, and they decrease in the anoxic bottom water (Fig. 3f). Soluble reactive phosphorus (SRP) was not detectable in the upper 40 m of the water column (Fig. 3d,j); it is obviously limiting for phytoplankton in Lake Challa, in particular for diatom blooms, which would be supported by high Si concentration of the lake (Barker et al. 2011). High concentrations of SRP and NH₄ in deep water (> 40 m water depth; August 2007) probably reflect the release of PO₄ and NH₄ from sedimenting particles and the diffusive reflux of both components across the sediment–water interface under anoxic conditions. In November 2007, seasonal depletion of both ions was due to oxygenation of the deep water (Fig. 3).

Nitrate is very low in the top 20 m (August 2007) to 40 m (November 2007), respectively, where it is probably consumed by primary producers. The presence of NO₃ in 20 m to 30 m and 50 m to 70 m water depth in August 2007 (Fig. 3e; Table 1) and at ~ 60 m in November 2007 reflects microbe-mediated organic matter degradation and nitrification under oxic conditions (Buckles et al. 2013). Oxygen

### Table 2. Extended.

<table>
<thead>
<tr>
<th></th>
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<td>0.00</td>
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<td>0.01</td>
<td>0.07</td>
<td>0.02</td>
<td>1.48</td>
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concentrations in water profiles indicate permanent anoxic conditions in the deep water. Moreover, the density contrast along the water column between 40 m and deep water inhibits mixing of the entire water column during weakening of thermal stratification during southern hemisphere winter (Fig. 4a,b,d).

The presence of NO₃ in deep water from August 2007 indicates previous deep-water oxygenation and related nitrifying activity. Furthermore, the combined increase of SO₄ and decrease of total dissolved solids (TDS) between August and November 2007 indicate a short-term overturn event of the water column. Alternatively, the presence of NO₃ and SRP at 60 m in November 2007 could be an indication of groundwater inflow. However, the profiles of inert ions (Na, K, Mg, Cl) do not support this interpretation (Table 1). Notably, the water profile from 23 August 2007 shows increased TDS and electric conductivity (EC) in deep water, which correlates with an increase in Ca and DIC (Table 1). The increase of both components most likely reflects the dissolution of autochthonous calcite and an increase in DIC due to microbe-mediated mineralization of organic matter. Undersaturation of calcite in deep water supports this interpretation.

In general, Si concentrations are high in Lake Challa (> 20 mg L⁻¹, Table 1). Under stratified conditions in August 2007, higher Si concentrations in the hypolimnion (Fig. 3) reflect diatom dissolution in the water column and Si reflux from the sediments.

The profiles of Ca : Sr and DIC during seasonal stratification (Fig. 3) reflect biogenically induced precipitation of
low-Sr CaCO3, driven by biogenic uptake of CO2 in the euphotic zone, dissolution of settling autochthonous carbonate particles in water depths > 40 m, and Ca reflux across the sediment–water interface. An oversaturation increase of calcite in the upper 40 m of the water column was recorded from 23 August to 28 November 2007 (Table 3).

Very low Fe and Mn concentrations, down to 40 m water depth in August 2007 and in the whole water column (except at ~ 60 m and 95 m) in November 2007 (Fig. 3c,i), reflect the precipitation of Fe(Mn)OOH in oxygenated water. Increasing Fe and Mn concentrations downward from the upper hypolimnion in August 2007 document the diffusive reflux of Fe and Mn from the surface sediments, which is sustained by the microbe-mediated reduction of reactive Fe(Mn)OOH and Fe(Mn)-release into the pore water of the sediments. The greater increase of Mn in comparison to Fe (Mn : Fe maximum at ~ 50 m water depth; Fig. 3b) indicates that prevailing pH–Eh conditions favored the precipitation of dissolved Fe or the release of Mn2+ from settling Fe(Mn)OOH precipitates and from the sediments (excluding subsurface inflow with high Mn:Fe ratios) or both.

Compared to other African stratified lakes (Talling 1966; Davison 1993), Fe concentrations in the anoxic hypolimnion of Lake Challa (in August 2007) are relatively low. Profiles from 28 November 2007 document the chemical stratification shortly after overturn. The deep-water concentrations

### Table 3. Saturation indices of selected minerals with pH and temperature values during an ongoing mixing event (23 Aug 2007) and during stratified conditions (28 Nov 2007).

<table>
<thead>
<tr>
<th>Depth (m)</th>
<th>Temp(°C)</th>
<th>pH</th>
<th>Calcite</th>
<th>Aragonite</th>
<th>Si(a)</th>
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<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>0</td>
<td>23</td>
<td>8.5</td>
<td>0.58</td>
<td>0.43</td>
<td>−0.42</td>
</tr>
<tr>
<td>10</td>
<td>23.1</td>
<td>8.5</td>
<td>0.59</td>
<td>0.44</td>
<td>−0.42</td>
</tr>
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<tr>
<td>30</td>
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<td>0.32</td>
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<tr>
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<tr>
<td>50</td>
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<tr>
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<td>7.12*</td>
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<tr>
<td>80</td>
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<td>7</td>
<td>−0.47</td>
<td>−0.61</td>
<td>−0.26</td>
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<tr>
<td><strong>28 Nov 2007</strong></td>
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<tr>
<td>0</td>
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<td>9.1</td>
<td>1.15</td>
<td>1.01</td>
<td>−0.5</td>
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<tr>
<td>10</td>
<td>26.4</td>
<td>9.1</td>
<td>1.15</td>
<td>1</td>
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<tr>
<td>20</td>
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<tr>
<td>40</td>
<td>23.8</td>
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</tr>
<tr>
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<td>7.5</td>
<td>−0.01</td>
<td>−0.22</td>
<td>−0.31</td>
</tr>
<tr>
<td>80</td>
<td>23.4*</td>
<td>7.5*</td>
<td>−0.3</td>
<td>−0.44</td>
<td>−0.31</td>
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<tr>
<td>90†</td>
<td>23.2*</td>
<td>7.5*</td>
<td>−0.29</td>
<td>−0.44</td>
<td>−0.41</td>
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<tr>
<td>95</td>
<td>23</td>
<td>7.5</td>
<td>−0.3</td>
<td>−0.44</td>
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* pH and temperature values interpolated.
† Average of two water analyses, Si(a) amorphous silica, hydrochemical data for Phreeq-calculations of saturation indices (SI) see Table 1.
of Fe$^{2+}$ and Mn$^{2+}$ certainly increased during the stratification period until their seasonal precipitation.

With TDS concentrations of approximately 350 mg L$^{-1}$ (Table 1), the water of the mixed Lake Challa is moderately mineralized (Talling and Talling 1965). The salinity of the lake was fairly constant between September 1999 and November 2007 (Fig. 4d). The maximum EC variation in the upper 45 m of the water column ($\sim 25 \mu S cm^{-1}$) may largely depend on interannual variations in the lake water overturn and calcite precipitation. The current TDS of

![Diagram of sediment trap materials composition changes](image_url)

Fig. 5. Selected data for changes in the composition of sediment trap materials in the context of seasonal meteorology and changes in water-column stratification. (a) Mn:Fe ratios and Mn contents; shaded bars mark the seasonal overturn. (b) Seasonal changes in water-column stratification on temperature records received from thermistor data. (c) Ti contents and Ti:Al ratios of sediment trap materials; increase in Ti:Al marks higher proportions of siliciclastic input of local provenance. (d) Carbonate contents of sediment trap materials calculated as CaCO$_3$ on the basis of TIC analyses (solid line) and monthly rainfall data from the meteorological station in Voi (Kenya) (bars).
Lake Challa may have been built up over a long period by evaporative enrichment.

**Thermal and chemical stratification**—Limnological surveys conducted between September 1999 and November 2007, in combination with meteorological monitoring data from Voi (Kenya) (Fig. 2; ~100 km to the east of Lake Challa), were used to interpret seasonal changes in the thermal and chemical stratification of Lake Challa. The seasonal formation of a well-defined thermocline is related to warming during the southern hemisphere winter (October–May) and generally low wind stress during this season. The thermal stratification of the water column is not sustained year-round, due to cooling and intensified wind during the dry season. Our monitoring data imply a yearly minimum turnover of the upper 45 m of the lake (Fig. 5b), with reduced vertical temperature gradients during the southern hemisphere winter (Figs. 4a, 5b). Temperature and oxygen profiles document distinct interannual variations in the intensity of seasonal mixing events, depending on cooling and wind stress during the year (Figs. 4b, 5b). There is, however, no simple correlation between water-column mixing and the penetration depth of O2. The advective and diffusive downward flux of O2 is additionally determined by the diurnal production and consumption of O2 by phytoplankton and by microbe-mediated organic matter degradation.

The increase in EC and TDS between ~45 m and 60 m of the water profile from 23 August 2007 suggests that seasonal mixing was limited to a water depth of ~50 m (Fig. 4d; Table 1). The undisturbed finely laminated pattern of the sediments (Fig. 6) indicates that the lake bottom was seasonally anoxic over longer periods, which is supported by the oxygen profiles (Fig. 4b).

The O2 profile in November 2006 exhibits maximum O2 concentrations at ~10 to 15 m, whereas in all other profiles O2 concentrations are constant (or decrease with depth) within the upper ~15 m (Fig. 4). Such maxima commonly
develop in stratified nutrient-limited lakes and reflect enhanced primary productivity around the thermocline (Wetzel 1983). Transparency of surface waters in the stratified Lake Challa is high (Secchi depth of 5 to 7 m; Damsté et al. 2009).

Stable isotope profiles for August and November 2007 provide information about the water budget and seasonal mixing of the water column (Fig. 4e). The August 2007 data show that seasonal mixing did not include the entire water column. The heavier isotope composition of the epilimnion vs. deep water in November 2007 indicates seasonal evaporative enrichment of Lake Challa. The δ$_{18}^{0}$O data are strongly linked to precipitation and evaporation, with lighter δ$_{18}^{0}$O caused by precipitation and heavier δ$_{18}^{0}$O values by an increase in evaporation. The overall heavier isotope composition in Lake Challa vs. precipitation (cf. precipitation Entebbe [Uganda]: average annual δ$_{18}^{0}$O = −2.26‰; average annual deuterium = −5.8‰; International Atomic Energy Agency 1992) has been adjusted over longer timescales. At multiannual scales, paleo-δ$_{18}^{0}$O data for Lake Challa were inferred from a diatom isotope study and reflect the aggregated balance between the amount of precipitation and the intensity and duration of dry-season evaporation (Barker et al. 2011).

The detrital allochthonous sediment fraction—The siliciclastic sediment fraction comprises the input of local soil particles by surface run-off from a restricted area and aeolian influx of a wider source region. Table 2 compiles the bulk chemical composition of various representative materials from around Lake Challa. Local soils and their basic precursor rocks have much higher Fe, Mg, Ti, Ca,
and P than acidic gneisses of the basement, which outcrops at some distance from the lake (Table 2). Dust influx of remote provenance, if contributing in substantial amounts to the bulk siliciclastic influx, may be characterized by geochemical signatures similar to mean Upper Continental Crust (UCC) composition (Rudnick and Gao 2003).

Sediment trap samples—Optical and electron microscopy of sediment trap samples reveal three major components: (1) autochthonous calcite, predominantly precipitated during southern hemisphere summer, (2) amorphous organic and fine-grained (clay-sized) siliciclastic matter settling during the long rainy season, and (3) diatoms mainly deposited during the southern hemisphere winter period (Table 4). The percentage of siliciclastic matter in the trap materials is estimated on the basis of ICP-AES analyses, assuming a constant Al$_2$O$_3$ concentration of the inorganic detrital influx (15.6 wt%, mean continental crust composition; Taylor 1964). On this basis the siliciclastic matter in the traps ranged between 20.3 wt% and 49.6 wt% (average: 26.7 wt%, SD 7.0). Values above 30 wt% are only recorded during the rainy seasons between November and March. The Al : Ti mass ratio of most samples is rather low (minimum 0.7, average 14) as expected for input of local rock particles of basaltic composition (average basalt 11.0; Taylor 1964). However, six trap samples show distinctly higher Al : Ti (29–97) and lower Fe : Al ratios (0.07–0.17), probably representing a second siliciclastic component. Siliciclastic matter of remote provenance is delivered by aeolian input and contributes to the detrital sedimentary component. The highest Al : Ti ratio was determined for material collected in August 2008 during the dry and windy season (Table 4).

Seasonal changes between oxic and anoxic conditions lead to enrichment and depletion of dissolved Fe and Mn in the hypolimnion (Balistrieri et al. 1994; Schettler and...
The hydrochemical composition of the groundwater input (no data available), the diffusive reflux of Fe and Mn from the sediments, and the pH–Eh conditions in the hypolimnion cause seasonal enrichment of Mn$^{2+}$ vs. Fe$^{2+}$ in the deep water of Lake Challa (Fig. 5). Seasonal overturn events oxygenate the deep water, causing precipitation of Fe:MnOOH and, consequently, high Mn and Mn : Fe ratios in sediment trap materials (Fig. 5).

Monthly Ca varies substantially, from 0.2 wt% to 32.2 wt%, reflecting distinct seasonal variation in the precipitation of autochthonous calcite (cf. microscopic component analyses, Table 4). Highest CaCO$_3$ occurred between November and March in the short rainy season. A positive correlation between CaCO$_3$ and Ca : Sr ($R^2 = 0.78$; $n = 33$) indicates that autochthonous CaCO$_3$ is a low-Sr component (cf. Ca : Sr water profile; Fig. 3).

Total P in the sediment trap samples comprises (1) organically bound P of planktonic matter, (2) dissolved PO$_4$ removed by chemical (co-)precipitation or sorption on settling inorganic particles, and (3) P associated with the allochthonous detrital organic and inorganic input. The monthly molar TOC : P ratios of the trap material vary between 9.0 and 250.1 (average 119), showing major positive and negative deviations from the Redfield ratio (C : P = 106 : 1; Redfield 1958). In our interpretation, TOC : P values distinctly below 106 indicate substantial P contributions from abiotic nonplanktonic components. P and TOC positively correlate ($R^2 = 0.51$) with excess P, if samples with TOC : P $<$ 60 are not considered (Fig. 7).

The same trap samples also show a positive correlation with sulfur ($R^2 = 0.80$), indicating that most sulfur is bound to organic matter (Fig. 7). Ca shows no positive correlation with P. Therefore, sorption of PO$_4$ onto settling calcite crystals can be rejected as a significant controlling factor for the abiotic removal of dissolved P from the lake water.

Sediments—In 2005, a 20.81 m long continuous core spanning the past 25,000 yr was recovered from the center of Lake Challa. The sediment profile is dominated by sequences of fine light–dark couplets, the same as those found in the uppermost 10 cm of the short gravity core (CH05-1G). The chronology of the long sediment profile
is based on 168 AMS \(^{14}\)C dates and \(^{210}\)Pb dating (Verschuren et al. 2009; Blaauw et al. 2011). Radiometric dating and analysis of the monthly collected sediment trap material confirmed that the light–dark couplets record annual deposition (supplement in Wolff et al. 2011). Based on microscopic inspection, the dark layers are mainly composed of amorphous organic matter and fine-grained siliciclastics, with single silt-sized calcite crystals imbedded in this matrix. Diatom skeletons (\textit{Gomphocymbella} sp. and \textit{Nitzschia} sp.) are the dominant constituents of the light layers (Fig. 8; Barker et al. 2011; Barker et al. 2013).

The chronology for the upper 10 cm of core CH05-1G was established by counting the light–dark couplets representing the past 100 \pm 2 yr. This floating chronology was linked to the absolute timescale of the existing 25,000 yr chronology of the long core, using distinct marker layers. In lacustrine environments such varve types are classified as carbonaceous organic varves (Zolitschka 2007). Varved sequences potentially provide information about past environmental changes by variations in structure, composition, and thickness (Ojala et al. 2012).

Investigation of annually laminated lake sediments with \(\mu\)XRF scanning provides additional information on seasonal signals and long-term changes (Yancheva et al. 2007; Brauer et al. 2008). The \(\mu\)XRF scans, applied on a 10 cm resin block of core CH05-1G, show Ti and Fe maxima and Si:Al minima for the dark laminae, and the light diatom-rich layers show Si:Al maxima and Fe and Ti minima (Fig. 6). Ca:Al maxima occurring within the dark layers document seasonally enhanced deposition of autochthonous calcite. Light layers, documenting seasonal diatom blooms (Si:Al maxima), coincide with Mn, Mn:Fe, and Mn:Ti maxima that indicate chemical precipitation of Mn during overturn events (Figs. 5, 6).

In local rocks and soils, the concentration of Fe is approximately 70 times higher than the Mn concentration. Therefore, variations in the Fe:Ti and Fe:Al ratios document the chemical precipitation of dissolved Fe (with a low sensitivity compared to Mn ratios) or uncertainty due to the different chemical composition of detrital influx of local and remote provenance. Count ratios of Al:Ti calculated from \(\mu\)XRF scans display peaks in the light laminae. Al:Ti maxima indicate the influx of Al-rich siliciclastic matter of remote provenance during the windy and dry season. Total sulfur is clearly enriched in the dark layers (Fig. 6) and is related to organically bound S and/or Fe-sulfide.

The use of Ti for normalization probably overestimates the portion of chemically precipitated Mn for sediment...
calcite precipitation in the previous season, (2) temperature
precipitation is associated with a number of factors: (1)
rich layers. The decline or even absence of seasonal calcite
seasonal presence of light-colored noncalcareous diatom-
completely absent in the monthly trap material (Table 1).
calcite contents were distinctly lower or were even
contrast to stratified conditions in November 2007
period in August 2007 show undersaturation of calcite in
euphotic zone. Hydrochemical data during the overturn
nutrient cycle supports seasonal diatom blooms, particu-
during the dry season (May–October). The internal
related to an increase of local wind stress and cooling
layers in the varves (Figs. 6, 8). The darker layers
represent the two rainy seasons (November to December
and March to May) and the brief intervening dry season
with amorphous organic matter derived from phytoplank-
ton and calcite precipitation.
The influx of local siliciclastics is dominated by local
basaltic debris during the combined rainy seasons (No-
ember–April) washed into the lake by surface runoff. The
lake probably also receives dissolved nutrients from surface
run-off. Nutrient influx by groundwater inflow depends on
rainfall seepage and may also increase as a result of lake-
level lowering during the dry season. The influx of N and P
is hardly detectable in lake water because it is immediately
consumed by phytoplankton. During the rainy season,
calcite oversaturation increased in the upper 20 m of the
water column (Table 3). Seasonal biogenically induced
precipitation of low-Sr calcite crystals is well documented
by the water profiles of Ca and DIC, as well as Ca : Sr
ratios of monthly trap data and the composition of
calcareous organic-rich layers in the sediment.
Diatom growth is strongly limited by SRP in the
stratified lake. Partial overturn of the water column is related
to an increase of local wind stress and cooling
during the dry season (May–October). The internal
nutrient cycle supports seasonal diatom blooms, particu-
larly by vertical advective transport of SRP into the
euphotic zone. Hydrochemical data during the overturn
period in August 2007 show undersaturation of calcite in
contrast to stratified conditions in November 2007
(Table 3). During the dry seasons between 2006 and 2009,
calcite contents were distinctly lower or were even
completely absent in the monthly trap material (Table 1).
The 100-yr sediment record documents the regular
seasonal presence of light-colored noncalcareous diatom-
rich layers. The decline or even absence of seasonal calcite
precipitation is associated with a number of factors: (1)
calcite precipitation in the previous season, (2) temperature
decrease, (3) mixing with CO$_2$-rich deep water buffering the
pH increase associated with the CO$_2$ uptake by phyto-
plankton, and (4) possible kinetic inhibition of crystal
growth by PO$_4$ (Dittrich and Koschel 2002) or the
composition of Mg–calcite cements influenced by the
Mg : Ca ratio of the solution from which they precipitated
(Mucci and Morse 1983; Morse and Mackenzie 1990).

The intensity, duration, and timing of the onset of
vertical mixing of the water column are annually variable.
During the period of seasonal stratification, dissolved Fe
and Mn are enriched in the anoxic deep water and
precipitate by oxygenation of the deep water during
overturn events. The net accumulation of chemically
precipitated Mn in the sediment distinctly exceeds that of
Fe due to hydrochemical characteristics of the groundwater
inflow and prevailing pH–Eh conditions or both. The
chemical precipitation of Mn is sensitively recorded in
μXRF scans of the sediments by Mn : Ti(Al) and Mn : Fe
maxima that coincide with light diatom-rich laminae
deposited during overturn events in the dry season.
Chemical characteristics of monthly trap materials demon-
strate that the siliciclastic influx comprises a local (basaltic)
component and a remote component characterized by high
Al : Ti and by a La : Y mass ratio close to 1. The ratio of
both end members in the depositional flux shows seasonal
variability. Siliciclastic matter deposited during the dry and
windy season occurring in the light layer is clearly enriched
in the high Al : Ti component and reflects aeolian input.

The climate archive of Lake Challa—Lake hydrology
reconstructions of the past 25,000 yr show that the lowermost
water column remained stratified and created anoxic
conditions at the sediment–water interface, which promotes
preservation of the seasonal laminae (Moernaut et al. 2010;
Barker et al. 2013). Biogenic silica resulting from diatom
blooms in the dry season is a major sediment component of
the Lake Challa sediments over the last 25,000 yr (biogenic
silica range 20–70%; Barker et al. 2011, 2013). These blooms
are associated with overturn events, which positively
correlate with the internal P cycle during overturn, and are
quantifiable by the thickness of the light diatom layers.
Seasonal upwelling of the deep-water SRP reservoir strongly
depends on wind strength and temperature decrease during
the dry season; therefore, the net accumulation of biogenic
opal (varve thickness and percentages of biogenic silica) can
be used as a proxy to reconstruct paleo wind variations
during the dry season (Wolf et al. 2011; Barker et al. 2013).
Assuming that present-day associations between El Niño
dynamics (wetter [drier during La Niña] and decreased
[increased during La Niña] wind speeds) in East Africa
existed throughout the period recorded in our long core, we
can interpret the Lake Challa sediments as indicating that
thick varves represent windier La Niña conditions and that
thinner varves are El Niño indicators. Variations in varve
thickness (between 0.08 and 7 mm yr$^{-1}$) thus show evidence
for interannual to centennial-scale changes in ENSO-related
rainfall variability over the past 25,000 yr (Wolf et al. 2011).
Photosynthetic production in the lake is strongly limited by
SRP availability. Accumulation of SRP in the deep
water depends on (1) external influx (mainly groundwater
inflow), (2) SRP removal during previous overturn events,
and (3) SRP flux from the surface sediments across the
sediment–water interface. The latter has to be considered for
paleoclimatic interpretations.

Al : Ti and La : Y may be additional suitable climate
proxies for local wind strength and mixing of the water
column. Increase of Al : Ti and decrease of La : Y potentially
record enhanced aeolian influx of remote provenance related to changes in the wind regime and the dryness in the source region of the dust, or both. Wind-induced mixing leads to higher Mn:Fe (Mn:Ti) and Si:Al ratios. The similar trend of Al:Ti illustrates the link between the external signal via dust and the internal lake response via mixing. Both ratios combined produce a reliable proxy for wind strength.

Limnological surveys conducted between September 1999 and November 2007 document seasonal variability in the sediment formation of Lake Challa, which is in agreement with meteorological observations in this region. During the generally warmer southern hemisphere summer months (October–May), stable thermal water stratification develops, and nutrient inflow during these rainy months supports peak photosynthetic activity in the euphotic zone. During southern hemisphere winter months (June–September), the vertical temperature gradient decreases strongly and south-easterly trade winds cause a pronounced thickening of the mixed layer, which varies from year to year. Deep and continuous mixing permits enhanced blooming of diatoms.

Based on lake water analyses, results from sediment trap analyses, and geochemical investigations on laminated sediments, we developed a model for the formation of laminated sediments in Lake Challa that is in good agreement with the observed seasonal changes in limnology and regional meteorological conditions. We show that diatom blooms are related to deeper mixing in the lake, which is documented by high Mn:Fe ratios within the light diatom-rich sediment laminae and in the sediment trap samples. Simultaneously, high Al:Ti ratios document enhanced aeolian influx of remote provenance. Dark laminae are characterized by higher detrital influx and, therefore, high Fe, Ti, and S concentrations. Low Mn:Fe ratios and observations of Fe-sulfides in dark laminae suggest strongly reducing conditions at the lake bottom during well-stratified periods. Sedimentological and geochemical investigations of recent sediments document a distinct seasonality in the record, and results from ongoing lake monitoring will probably allow even finer differentiation of these signals. These results from the Lake Challa sediment archive identify the site as a highly valuable target for high-resolution paleoclimate studies in East Africa.

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