Late Pleistocene and Holocene Afromontane vegetation and headwater wetland dynamics within the Eastern Mau Forest, Kenya

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ABSTRACT: The Mau Forest Complex is Kenya’s largest fragment of Afromontane forest, providing critical ecosystem services, and has been subject to intense land use changes since colonial times. It forms the upper catchment of rivers that drain into major drainage networks, thus supporting the livelihoods of millions of Kenyans and providing important wildlife areas. We present the results of a sedimentological and palynological analysis of a Late Pleistocene–Holocene sediment record of Afromontane forest change from Nyabuyabi wetland in the Eastern Mau Forest, a highland region that has received limited geological characterization and palaeoecological study. Sedimentology, pollen, charcoal, X-ray fluorescence and radiocarbon data record environmental and ecosystem change over the last ~16 000 cal a BP. The pollen record suggests Afromontane forest characterized the end of the Late Pleistocene to the Holocene with dominant taxa changing from Apodytes, Cellis, Dracaena, Hagenia and Podocarpus to Cordia, Croton, Ficus, Juniperus and Olea. The Late Holocene is characterized by a more open Afromontane forest with increased grass and herbaceous cover. Continuous Poaceaee, Cyperaceae and Juncaceae vegetation currently cover the wetland and the water level has been decreasing over the recent past. Intensive agroforestry since the 1920s has reduced Afromontane forest cover as introduced taxa have increased (Pinus, Cupressus and Eucalyptus). © 2021 The Authors. Journal of Quaternary Science published by John Wiley & Sons Ltd on behalf of Quaternary Research Association

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Introduction

African wetlands are dynamic ecosystems experiencing substantial land use and increasing hydroclimatic variability and stresses to biodiversity (Chapman et al., 2001; MEMR, 2012). Pollen-based analyses that reconstruct changes in past vegetation assemblages and distributions across the highlands of eastern Africa are beginning to characterize the spatiotemporal complexity of montane forests (Livingstone, 1967; Olago et al., 1999; Rucina et al., 2009; Finch and Marchant, 2011; Schüler et al., 2012; Opiyo et al., 2019). Several mountains in Kenya support lake systems that preserve lacustrine sediment archives of palaeovegetation dynamics since the Late Pleistocene (Hamilt, 1982; Marchant et al., 2018; Gil-Romera et al., 2019). Palustrine ecosystems in montane environments have similarly been used to establish environmental histories in equatorial eastern Africa (Hamilt, 1982; Heckmann, 2014; Finch et al., 2017). Among the montane ecosystems of Central Kenya with elevations >3500 m above sea level (asl), such as the Aberdare Range and Mau Escarpment, permanent lake ecosystems are not frequently supported due to steep topography, hydroclimate and sediment infilling. Thus, palustrine, soil, cave and other terrestrial geochronologies are important sources useful for analyses of vegetation change in response to climate, anthropogenic and local-scale mechanisms of environmental change. Longer term insights on ecosystem change from these crucial landscapes that are rich in biodiversity and provide a wide range of ecosystem services can be useful to inform their contemporary management (Gilson and Marchant, 2014).

Kenyan highlands are the headwaters to several large catchments; their forests generate and capture orographic and occult precipitation forming crucial headwater sources for major river systems (Nkako et al., 2005; Cuní-Sanchez et al., 2016; Los et al., 2019). The Mau Forest Complex is one of the five key water towers in Kenya (Nkako et al., 2005; MEMR, 2012). As most of rural sub-Saharan population relies on rain-fed agriculture (Wolf, 2011), and ~70% of Kenyans live in rural areas (Pieterse et al., 2018), understanding the functioning of these headwaters can inform management on the historical evolution and variability of these ecosystems and the ecosystem linkages to the hydrology. Notwithstanding the important roles that mountain ecosystems play with impacts across their wider catchments, high-elevation wetlands receive less ecosystem protection than large lowland wetlands, but are important contributors to biodiversity, landscape diversity, habitat connectivity and social-ecological resilience. This contribution is not just within the highland areas but across the catchment; for example the Mara River flows some 500 km to the south-west through agricultural areas and the Maasai Mara–Serengeti ecosystems and into Lake Victoria and the Nile. The Mau Forest Complex is one of the remaining forest blocks of the western Rift Valley in Kenya, supporting indigenous forests and wildlife and several large communities. The forests have undergone significant change since the late 1800s, conspicuously with the development of industrial forestry during colonial government administration (Klopp, 2012) and ongoing land-use change
towards agriculture and continued industrial forestry of exotic tree species (Kenya Gazette Supplement 2012).

Mau forests are divided into seven forest blocks: Eastern Mau Forest is the smallest of these, making the small and isolated remnant indigenous forests the most susceptible to further anthropogenic land-use modifications, climate change effects, ecological disturbances and introduced species (Okeyo-Owuor, 2007; Kinyanjui, 2011; Were et al., 2015). The remaining patches of indigenous forest are protected by legislation for their environmental and ecosystem services and cultural use and heritage (Republic of Kenya, 2016). Much of the lower elevation forests have been converted to agriculture over the past decades (Olang and Musula, 2011; Swart, 2016) with ~25% of the forest converted from AD 1994 to 2009 due to encroachment and settlement encroachment into the forest (Mwangi et al., 2017). The Government of Kenya recently excised ~353 km² of the forest to resettle victims of ethnic clashes as well as members of the Ogiek community previously evicted from the forest. Recent politicking has led to increased illegal settlement, logging and charcoal burning in the forest as a source of income (Nkako et al., 2005; Were et al., 2013). These increased human population pressures on the forests have further fragmented wildlife populations, with subsequent erosion and water distribution issues having consequences for downstream ecosystems and populations (Gichana et al., 2015; Mwangi et al., 2017; Dutton et al., 2018; Mwanake et al., 2019).

Neighbouring Mau forest blocks have undergone varying degrees of anthropogenic modifications impacting vegetation biodiversity, soil geochemistry and topsoil seed bank; however, ecological restoration potential remains (Kinjanji et al., 2013).

The environmental history of the Eastern Mau Forest complex is relatively understood (Marchant et al., 2018). Early colonial maps described the Mau escarpment as forested but explorers did not penetrate the region before the 20th century. The colonial government established the Mau Forest Reserve (among others) as demands for timber increased with road and railway construction, and exports (Cranworth, 1912). By the AD 1920s, early forest delineations (Troup, 1932) already noted heavy modifications to forests although there was no investigation of the natural history until geological mapping in the 1980s and 1990s (Williams, 1991).

Here we present the first investigation of long-term terrestrial ecosystem change in Eastern Mau documenting how the forest has changed since the Late Pleistocene. After the Last Glacial Maximum, glaciers retreated on the highest mountains of eastern Africa followed by changing elevation vegetation patterns on the mountains (Hamilton, 1982; Van Zinderen Bakker and Coetzee, 1988). As conditions warmed (Loomis et al., 2017), the African Humid Period generally brought higher moisture regimes to the regime from 14 000 to 6000–4000 years BP; with the timing of the transition to relatively drier conditions being time-transgressive and having high spatiotemporal complexity across Africa (Shananah et al., 2015; Phelps et al., 2020), including highland regions (Street-Perrott et al., 2007). East Africa experienced high precipitation variability characterized by high rainfall in the Early Holocene and increasing, yet highly variable aridity towards the present as evidenced by major drought events (Stager et al., 2003; Verschuren et al., 2005). The effect of increasing CO₂ through the Holocene and varying C₂/C₄ vegetation varied between high- and low-elevation environments (Urban et al., 2015), the last 10 000 yr from the Sacred Lake record on Mount Kenya (Olago et al., 1999) are summarized as dominated by C4 vegetation that reflect the increasing atmospheric CO₂, temperature and precipitation. By the end of the African Humid Period during the Late Holocene, moisture regimes became relatively drier but with high spatiotemporal variability. This was characterized by wetland and lake level variability in lowlands (Verschuren, 2001; Oberg et al., 2012; De Cort et al., 2018), changing montane moisture regimes (Barker et al., 2001, Street-Perrott et al., 2007) and forest pollen assemblages (Rucina et al., 2009; Githumbi et al., 2018a,b; van der Plas et al., 2019; Courtney Mustaphi et al., 2020). Pollen evidence of recent human land use and forest resource use varies between mountains and sites (Ryner et al., 2008; Heckmann et al., 2014; Iles, 2019).

For example, in Uganda (Hamilton et al., 1986; Jolly et al., 1998; Leju, 2009), forest resource use and conversion of land cover have increasingly occurred during in recent centuries (Troup, 1932; Petrusson et al., 2013; Gil-Romera et al., 2019; Courtney Mustaphi et al., 2020).

**Study region: Kiptunga Forest Block**

The Nyabuibiabubi wetland (2865 m asl) is located in the Kiptunga Forest Block of the Eastern Mau Forest Block (Fig. 1) that covers an area of 29 000 ha. The surficial geology consists of an extensive thick catena of Early Pleistocene Mau ashes with ferrous basal tuffs (Jennings, 1971; Williams, 1991). Soils are relatively young and productive Udands (Andisols) that contain volcanic tephra. The mantling and subsequent aeolian and hydrological erosion of these deposits have shaped much of the current topography of the mountain ridge and the basin and fluvial channels of the forest and wetland. Kiptunga Forest currently supports populations of birds, reptiles, antelopes, primates and hyenas. The Ogiek community, who historically practised predominantly hunter-gatherer livelihoods, inhabit the region, yet recently, the local populations have increasingly practised pastoralism with cattle, goats and sheep; much of the forested lower elevations have been converted to agriculture (Sang, 2001; Spruyt, 2011).

The Mau forests contain several headwater catchments that flow into the Rift Valley or towards Lake Victoria, most notably through the Mara River, which flows into Maasai Mara and Serengeti. As an orographic precipitation water tower, Mau provides water for rural and urban settlements, pastoral communities and wildlife. The high biodiversity of the Mau forest, the Maasai Mara National Reserve and Serengeti National Park have led to the designation of the three regions as Important Bird Areas. They are also habitat to high numbers of large game and host both indigenous and threatened animals such as the bongo and the yellow-backed duiker, the golden cat, the leopard and the African elephant (Nkako et al., 2005).

Scattered minor pockets of remnant indigenous broadleaf forests include Croton, Dombeya, Hagenia, Juniperus, Olea spp., Podocarpus and Prunus. The Maasai Mau block is entirely indigenous comprising a Juniperus–Podocarpus mosaic interspersed with indigenous vegetation glades (Nkako et al., 2005). The highly valued indigenous timber species are Albizia gummifera, Olea capensis, Juniperus procera, Polyscias kikuyuensis, Podocarpus spp., Pouteria spp., Prunus africana and Strombosia spp. The major land cover classes (forests–shrublands, grasslands, croplands, urban areas, barren land cover and open water) have undergone tremendous changes over the last few decades with a decrease in Afromontane vegetation cover accompanied by an increase in agricultural and fragmented land and modifications to wetlands (MEMR, 2012; Were et al., 2013). In addition to broad-scale land-use transition there is quite common selective harvesting of tree species, and collection of poles, non-timber forest products and firewood. Few patches of indigenous forest remain due to intensive tree replanting–harvesting cycles of introduced tree species partitioned into plantation plots (Sanya, 2008).
Agroforestry species include indigenous *Juniperus procera*, and introduced taxa include *Cupressus lusitanica*, *Eucalyptus maculata* (syn. *Corymbia maculata*) and *Pinus patula*.

**Study site: Nyabuiyabui wetland**

The Nyabuiyabui hydronym is an Ogiek word meaning ‘spongy’ (Spruyt, 2011) or ‘marshy’, and may support a floating vegetation mat during wetter intervals. Nyabuiyabui wetland (Fig. 1) covers an area of 122 ha within the Kiptunga Forest Block. The current waterlogged/open water area extends to about 6 ha although water levels vary in response to local hydroclimatic conditions. Nyabuiyabui has minor ephemeral inflows and a single outflow to the south-west that forms a tributary of the Mara River network. The wetland is shallow with observed water depths at the centre of the basin well below 50 cm during March–April 2014 and lower still (<20 cm) during April 2015. Ancedotal discussions with local people suggested that the water level has been decreasing during the remembered past and that there was open water during the early to mid-20th century. Cattle graze along the wetland margin and impact the hummocky ground and marshy morphology of the Poaceae–Cyperaceae–Juncaceae tussocks. The wetland is surrounded by small patches of indigenous forest and monoculture tree plots of varying ages with planting dates from AD 1935 to 2006 and predominantly during the 1960s (Sanya, 2008; Courtney Mustaphi et al., 2016). As of 2015, plantation plots of *Cupressus lusitanica* were the most predominant in the watershed surrounding the wetland. The closest archaeological records are human and animal remains from an excavated burial cave, north-east of the wetland but significant forestland use could not be determined from the finds. The finds were ascribed to the Late Stone Age due to their similarity to finds from that period (Faugust and Sutton, 1966; Merrick and Monaghan, 1984).

**Methods**

**Field methods**

In 2014, a suitable coring site was determined by probing the sediments with fibreglass rods along transects to locate the maximum accumulation. A 537-cm core was recovered from 0°26′11.28″S, 35°47′58.74″E, 2920 m asl near the wetland centre using a hand pushed hemicylindrical Russian corer 5 cm in diameter (Fig. 1). Sediment cores were collected in 50-cm drives with 10-cm overlapped sections from parallel coring holes. Cores were transferred to longitudinally split PVC tubes, wrapped in plastic wrap and aluminium foil, shipped to the University of York, UK, and refrigerated at 4 °C.

**Laboratory analysis**

Six bulk sediment subsamples and three sieved and picked organic matter (plant material) samples were accelerator mass spectrometry (AMS) radiocarbon dated at Queen’s University Belfast 14CHRONO laboratory, UK; Scottish Universities Environmental Research Centre (SUERC), Glasgow, UK; or DirectAMS, Bothell, USA. The IntCal13 curve (Reimer et al., 2013) was used to calibrate the dates and an age-depth model was developed using a BACON R script with default settings (Blauuw, 2010; Blauuw and Christen, 2011; R Development Core Team, 2017). Several iterations of potential age–depth models were run to explore the potential ranges for bounds to
stratigraphic zonations and to extrapolate a basal date. More confidence was given to sieved plant material (mostly grass charcoal fragments) that frequently provide narrower dating uncertainties over bulk sediment AMS radiocarbon dates (Rey et al., 2019) because of the relatively short growth time of above-ground grassy fuel.

Loss-on-ignition (LOI) and particle-size distribution analyses were carried out to characterize the sediments: organic matter content, carbonate content and mean clastic particle size every 5 cm down core. LOI analysis involved weighing the wet samples and then again after drying at 105, 550 and 950 °C (for 24, 5 and 3 h, respectively) to calculate the dry weight, organic matter and carbonate contents, respectively (Heiri et al., 2001). Particle-size distribution analysis was carried out using the Malvern laser granulometer (MEH/MG1B9194). The procedure involved pretreating 1-cm³ wet sediment subsamples with 30% hydrogen peroxide in a hot water bath to digest organic matter and reduce particle aggregation (Syvitski, 1991). If the subsample contained <3.5% organic matter, the hydrogen peroxide treatment was skipped. At a pump speed of 1500, 1–2 g of sample was added until laser obscuration in the measurement column was 4%. The granulometer repeated three measurements and calculated an average result (Malvern Instruments Ltd, 2007).

The cores were scanned using a Cox Analytical Systems ITRAX core scanner at the Department of Geography and Earth Sciences, Aberystwyth University, UK. The ITRAX core scanner collected optical imagery of the core face using an RGB digital camera, and measured magnetic susceptibility using a Bartington M2S2E sensor at 1-cm intervals and air-corrected between measurements. Magnetic susceptibility is the degree of magnetization in response to a magnetic field measured in intensity values that are dimensionless units (X). The X-ray fluorescence (XRF) results represent a semiquantitative measurement of elemental composition in kcps (thousands of counts per second) of the sediment matrix. In total, 22 elements were examined at 0.05-cm intervals through XRF with a 3-kW water-cooled molybdenum anode X-ray tube (60 kV, 35 mA, 200-ms exposure). The results are influenced by potential X-ray absorption and/or scattering across the core due to variability in water content, particle size distributions, mineralogy and surface roughness of cleaned core face (Croudace et al., 2006).

Subsamples of 1 cm³ of sediment were extracted at 1-cm intervals from the wet core face for macroscopic charcoal analysis. This was soaked in sodium hexametaphosphate solution to disaggregate the organic material and clay particles (Bamber, 1982). A drop of hydrogen peroxide whitened non-charcoal organic matter (Schlachter and Horn, 2010; Whitlock et al., 2010). Samples were gently wet sieved through a 125-µm mesh, and the retained charcoal pieces were identified by visual inspection and probed with a metal needle (Hawthorne and Mitchell, 2016; Vachula, 2019) and tallied under a Zeiss Axio Zoom V16 microscope at magnifications of 10–40x. Counts were converted to charcoal concentration values (number of particles per unit volume, pieces cm⁻³).

A 1-cm³ subsample was obtained every 10 cm for pollen and spore analysis following sequential chemical digestion steps using 10% HCl, 10% KOH and acetylsalicylic (Aspirin and Iversen, 1950, 1989; Erdman, 1960). Heavy liquid separation using sodium polytungstate (3NaWO4·5WO3·H2O with relative density = 2) was carried out after acetylsalicylic (Neumann et al., 2010; Quick, 2013; Colomboari et al., 2018). One tablet of the acetylsalicylic exotic marker Lycopodium spores (n=9666, σ=212 spores per tablet; University of Lund batch no. 3682) was added before pollen preparation to aid in calculation of absolute concentrations (Stockmarr, 1971; Bonny, 1972).

The data were analysed using Rbaco package version 2.3.9.1 to develop the age–depth model using Bayesian approaches. Dates identified as outliers in the initial BACON run (default settings) were the bulk sediment samples. A study published in 2018 looking at the impact of model choice, dating density and quality on chronologies found that using BACON outliers were found to have little to no impact on model precision (Blaauw et al., 2018). Rioja version 0.9-21 (Juggins, 2020) was used to run the hierarchical constrained clustering – CONISS (ITRAX, charcoal and pollen), calculate the statistically significant number of assemblage zones (Bennett, 1996) and C2 for stratigraphic plots (Juggins, 2003). CONISS analysis was carried out on the complete pollen dataset but for ease of view we present the dominant taxa in each grouping in the text and the complete pollen diagram in the Supporting Information.

Data accessibility
All data generated from this study will be openly available via the African Pollen Database (Vincens et al., 2007), a constituent of the Neotoma Paleoecology Database and data repository (Grimm et al., 2018; Williams et al., 2018).

Results

General lithology description and geochronology
Six lithological units were identifiable from the 537-cm sediment core that was mainly dark organic-rich silt sediment. The top 8 cm comprised organic-rich detritus, mainly plant roots. This changed into a layer of dark grey clayey silt that extended until 124 cm (between 84 and 124 cm there is an increase in the coarse sand fraction). From 124 to 385 cm the sediment changes to a dark brown silt layer (>80%) with grey/black laminations between 224 and 274 cm. Starting with a thick black layer at 386 cm, the sediment becomes a darker brown sandy silt until 474 cm. From 475 cm to the bottom, the sediment changes back to a dark grey sandy silt with light coloured laminations and concretions at depths with increasing sand. The transitions in sediment colour and type along the core are rapid and distinct except in the bottom 1 m where laminations are visible.

Nine radiocarbon dates were used in BACON (Blaauw, 2010) to develop a plausible age–depth model (Fig. 2; Table 1). The Bayesian model recognizes four ages as outliers: 2449 ± 35 cal a BP at 50–51 cm, 10 721 ± 47 cal a BP at 100–101 cm and 14 424 ± 45 cal a BP at 384–853 cm are older than the dates below them, and 13 963 ± 60 cal a BP at 315–316 cm is younger than the dates above it. Due to the relatively low number of available dates to construct a robust age–depth model, we use the most parsimonious suite of dates based on macrofossils and the most likely date sequence (Fig. 2). In addition, the palaeoenvironmental changes are primarily discussed within depth boundaries and the ages discussed within broad time intervals/stratigraphic stages (Walker et al., 2019), i.e. Late Pleistocene to Early Holocene (538–240 cm), the early and Middle Holocene (240–100 cm) and the Late Holocene from 100 cm to the top (present).

Sedimentology

The sedimentology is highly variable along the core (Fig. 3). LOI results show that the sediment bulk density ranged between 0.1 and 3.2 g cm⁻³ with an average of ~1 g cm⁻³ and standard deviation (SD) of ~0.5, organic matter ranged

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from 0 to 100% with an average of ~21% and SD of ~18, while the carbonate content ranged between 0 and 40% with an average of ~6% and SD of ~7.

The sediment is composed of organic material mixed with varying amounts of silt, sand and clay (Fig. 3). The silt content is highest with an average of 72.44 ± 1.13% (ranging from ~34 to 86%), followed by sand with an average of 17.84 ± 1.13% (ranging between 2 and ~63%) and finally clay with an average of 9.63 ± 0.39% (ranging between ~1.6 and 19%). The sand component is most variable throughout the sediment core. A stratigraphically constrained cluster analysis highlights three distinct zones. From the bottom of the core to 370 cm covering the end of the Pleistocene, the clay content ranges between ~3 and 15%, silt between 52 and 81%, and highest variance is in the sand content between 10 and 55%. Clay content is the lowest, averaging ~7% followed by ~67% silt and ~26% sand; this zone can be described as a sandy silt. The second zone from 365 to 120 cm, covering the end of the Pleistocene and Early Holocene, shows an increase in clay and silt percentages to ~11 and ~80%, respectively, and a decrease in the sand content to ~9%. The clay content ranges between 5 and ~18%, silt ~57 to ~86%, and sand 2 to ~33%; this zone can be described as a clayey silt. The top of the record from 115 cm to the top of the core consists of a sandy silt-like bottom section where the average clay is ~11%, silt is ~65% and sand is ~26%. The clay content remains at ~11% while the silt decreases from 80 to 65% and the sand increases from ~9 to ~26%.

Bulk density is higher in the bottom half of the core (530 to ~260 cm) during what would be the Late Pleistocene. It gradually decreases, and between 155 and 75 cm (Early and Middle Holocene) it averages 0.5 g cm<sup>−3</sup>. From 70 cm to the top of the core (Late Holocene), average bulk density increases back up to ~1 g cm<sup>−3</sup>. The organic matter and carbonate content are lowest in the lower half of the record (Late Pleistocene) at ~12 and 4%, respectively. This increases to 16 and 10% between 260 and 160 cm around the Late Pleistocene/Early Holocene interval. The section from 155 to 75 cm (Early to Late Holocene period) has the highest increase in organic matter at an average of 40%, while the carbonate content decreases to 6%. The top of the record (from 100 to 0 cm), representing the Late Holocene, has organic matter and carbonate content of ~35 and ~6%, respectively.

Table 1. Radiocarbon dating results from the 537-cm-long core collected in 2014.

| Depth (cm) | Age (14C a BP) | pMC (%) | Material | Laboratory code |
|-----------|----------------|---------|----------|-----------------|
| 0         | 201 ± 23       | 97.57 ± 0.28 | Plant remains | Top of core |
| 30–31     | 2449 ± 35      | 73.72 ± 0.32 | Bulk sediment | SUERC-57340 |
| 74–76     | 1865 ± 37      | 79.37 ± 0.13 | >63 μm charcoal | UBA-27553 |
| 100–101   | 10 721 ± 47    | 26.33 ± 0.15 | Bulk sediment | UBA-26117 |
| 128–129   | 7616 ± 33      | 38.75 ± 0.16 | >63 μm plant remains | D-AMS-009663 |
| 230–231   | 9837 ± 42      | 29.39 ± 0.15 | Bulk sediment | UBA-26118 |
| 315–316   | 13 963 ± 60    | 17.58 ± 13 | Bulk sediment | UBA-27554 |
| 384–385   | 14 424 ± 45    | 16.602 ± 0.094 | Bulk sediment | D-AMS 011823 |
| 536–537   | 13 692 ± 83    | 18.19 ± 0.19 | Bulk sediment | UBA-26116 |

Radiocarbon ages are presented with 1σ measurement error. pMC, per cent modern carbon with 1σ measurement error.

*Dates identified as outliers in the age–depth model.
Elemental profile and magnetic susceptibility ($\chi$)

The ITRAX core scanner was set to detect counts for 22 elements (Fig. 4; Table 2); Ni and I were not included in further data analyses because both occurred at very low concentrations and had a limited stratigraphic pattern (Githumbi, 2017). Magnetic susceptibility readings expressed as $\chi$, or volume susceptibility, represent the ratio of magnetization in samples (per unit volume) to the magnetic field created by the sensor, and are dimensionless with a scale of $10^{-5}$ SI units (Burrows et al., 2016).

Elemental composition was dominated by Fe (average 85.25 cps), Zr (6.5 cps) and Y (2.6 cps). Rb (1.3 cps) and Ti (1.2 cps) are the only other elements with average cps values $>1$ through the sediment core. Magnetic susceptibility ($\chi$) varied through the sediment record between $-4.14$ and $70.89 \times 10^{-5}$ SI with an average of 2.97. A stratigraphically constrained cluster analysis divided the elemental composition record into three significant zones labelled ITRAX1, ITRAX2 and ITRAX3 (Fig. 4).

ITRAX1 extends from 484 to 260 cm covering the Late Pleistocene to the Early Holocene. Fe (ranging from 82.46 to 92.82 cps), Zr (2.67 to 7.14 cps), Y (1.03 to 3.24 cps), Rb (0.91 to 2.33 cps), Mn (0.23 to 7.69, with an average of 0.9 cps) and Ti (0.79 to 1.59, with an average of 1.2 cps) had the highest average counts in this zone. Magnetic susceptibility averaged 4.47. In the next zone, ITRAX2 (from 260 to 127 cm), covering part of the early to mid-Holocene, the same elements still dominated with a slight increase in Zr, Y and Rb, while Ti and Mn decreased. Mn decreases from $-1.6$ to 0.6 cps. The top zone, ITRAX3, extends from 127 cm to the core top and includes the rest of the Holocene to the present. Between 113 and 100 cm, all the elements experience a spike except Ti, K and Fe, which decrease. In this zone, all the elements exhibit an increased trend in counts towards the top except K, Rb and Sr (Fig. 4). Pb shows the greatest increase from 0.022 (ITRAX3), 0.031 (ITRAX2) to 0.21 in ITRAX1, almost a 10-fold increase. Hg shows a similarly large increase in the uppermost sediment samples.

Charcoal record (125 µm)

Charcoal concentration fluctuated from the Late Pleistocene to present at 0–1198 pieces cm$^{-3}$ with a mean of 217 pieces cm$^{-3}$. Charcoal varies throughout the record and the CONISS analysis identifies three significant zones. The bottom of the record to 307 cm is the first zone, CHAR1, and covers the Late Pleistocene period, the second zone is from 306 to 105 cm and is divided into two subzones CHAR2B and CHAR2A, and the top zone starts at 104 cm and covers the Late Holocene.

CHAR3, covering the Late Pleistocene and Early Holocene transition, experiences fluctuations in charcoal concentration; minimum, maximum and mean charcoal concentration values are 25, 863 and 230 pieces cm$^{-3}$. The mean concentration is higher than the mean throughout the whole record. These values are lower in the next zone, CHAR2B, to 14, 727 and 205 pieces cm$^{-3}$, respectively. In CHAR2A there is a significant increase in the charcoal concentration, with minimum, maximum and mean values of 16, 1198 and 238 pieces cm$^{-3}$, respectively. The topmost zone is CHAR1 where charcoal concentration decreases significantly. Minimum, maximum and mean values here are 0, 513 and 121 pieces cm$^{-3}$, respectively.

Pollen analysis

Pollen taxon diversity varied down the Nyabuiyabui sediment core with $>$70 pollen types (Supporting Information, Fig. S1; Table 3) observed and enumerated. The sample with the highest diversity had 68 pollen taxa identified while the sample with the lowest had 18 pollen taxa identified. To aid interpretation and discussion, only the most common pollen types or those types that consistently contributed $>2\%$ to the pollen sum for any level are presented, although the full spectra were used for cluster analysis and were grouped into Afromontane, trees, shrubs, herbs and aquatic taxa (Fig. 5; Table 3). Aquatic taxa included Ludwigia, Nymphaeae,
Cyperaceae and *Typha*. *Cupressus* and *Pinus* are Neophytes that appear in the record during the last ~200 years. Table 3 contains the list of taxa that comprise each grouping.

Throughout the sediment core, Afromontane pollen accounted for ~24% of the taxa (ranging from 12 to 44%), tree pollen accounted for ~15% (0 to 27%), shrub pollen accounted for 8% (0 to 15%), herb pollen accounted for 34% (21 to 37%), aquatics accounted for ~13% (2 to 35%), Poaceae accounted for 7% (0 to 18%) while unknown pollen accounted for ~0.4% (0 to 1%) (Fig. 5).

The pollen diagram was divided into three significant pollen zones using a broken-stick and clustered hierarchical analysis (Bennett, 1996) (Fig. 5). The lowermost pollen zone NBPOLL1 extended from the bottom of the core to 340 cm (the end of the Pleistocene). Afromontane taxa accounted for ~28%, with *Podocarpus*, *Cordia* and *Dracaena* dominating at ~3% each. Tree pollen accounted for ~20%, shrubs for ~11%, herbs for ~32% and unknown pollen for ~0.4% (Supporting Information, Fig. S1). In this zone, arboreal pollen (Afromontane, trees and shrubs) accounted for ~59% and non-arboreal taxa for ~41% (Fig. 5).

The second pollen zone, NBPOLL2, extended from 330 to 100 cm (end of the Pleistocene to the mid-Holocene) and was divided into sub-zone NBPOLL2A from 330 to 220 cm (end of the Pleistocene and Younger Dryas) and sub-zone NBPOLL2B from 210 to 100 cm (Early and Middle Holocene). There was a sharp decrease in the NBPOLL2A pollen counts across all the vegetation types (average sample count of ~970 in zone NBPOLL1 fell to ~390 in zone NBPOLL2A). The Afromontane taxa remained at ~29%, with *Olea*, *Podocarpus* and *Juniperus* each dominating at ~6%. However, the tree and shrub taxa decreased significantly dropped from 20 to 7% and from 11 to 5%, respectively. This was accompanied by an increase in herbs and aquatic taxa from 32 to 34%.

In zone NBPOLL2B, the average for Afromontane taxa fell significantly to ~18% from 29% in the previous zone (Fig. 5). Afromontane genera dominating this zone were *Cordia* (~4%), *Podocarpus* (~4%), *Croton* (~2%) and *Olea* (~2%). Most of the tree, shrub and herbaceous taxa that had disappeared in NBPOLL2A reappear (*Alangium*, *Commiphora*, *Lannea*, *Maytenus*, *Polyscias*, *Syzygium*, *Abutilon*, *Fagonia* and *Rumex*). Tree taxa increased from 7 to 13% while shrub taxa increased

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**Figure 4.** A stratigraphic plot of the sediment elemental characterization using ITRAX (expressed as relative abundance of cps, counts per second) alongside the CONISS dendrogram (Bennett, 1996) which divided the record into three significant zones. The element iodine (I) was also measured, but did not have values above the analytical detection limit of the ITRAX scanner. Radiocarbon dates, ‘+’ symbols on the left. [Color figure can be viewed at wileyonlinelibrary.com]
from ~5 to ~7%. Herbaceous and aquatic taxa increased to 40 and 11%, respectively.

The third pollen zone (NBPOLL3) extends from 90 cm to the top (Late Holocene to present), where the average Afromontane pollen taxa reduced to 16%, tree pollen remained at ~12%, and shrubs remained ~7%. Herbs fell significantly to ~24% from 40% while the aquatic taxa and Poaceae increased to ~24 and ~13%, respectively. The aquatics and Poaceae in this zone were at their highest recorded level. Podocarpus (~5%), Cordia (~3%), Ficus (~3%), Commiphora (~2%) and Rhus (~2%) dominate this zone. Introduced tree taxa (Neophytes), namely Cupressus and Pinus, appear towards the top of the sediment core (20 cm) at high abundance as they were introduced taxa at commercial timbering scales and frequently used on residential and industrial properties.

Pollen taxa, charcoal concentration, organic matter content, clay, sand and some elements/elemental ratios are plotted alongside the lithology and radiocarbon dates. Summary aspects

Table 3. Pollen taxa identified from the analysis of the 537-cm core collected from Nyabuiyabui

| Afrotomtane | Trees       | Shrubs          | Herbs              | Aquatics          |
|-------------|-------------|-----------------|--------------------|-------------------|
| Apodytes    | Allophylus  | Ruhiaceae       | Laportea           | Ludwigia          |
| Celtis      | Annona      | Salvadoria      | Malvaceae          | Nympheaceae       |
| Cordia      | Carissa     | Scheffleria     | Ormocarpum         | Cyperaceae        |
| Croton      | Combretaceae| Syzygium        | Phyllanthus        | Typha             |
| Cupressus * | Commiphora  | Tachonanthes    | Plantago           |                   |
| Dracaena    | Lannea      | Abutilon        | Rumex              |                   |
| Ficus       | Maera       | Acanthaceae     | Sansevieria        |                   |
| Hagenia     | Maesa       | Aphyranthes     | Solanaceae         |                   |
| Juniperus   | Maytenus    | Amaranthaceae/Chenopodiaceae | Umbelliferae |
| Olea        | Monotes     | Asteraceae      | Urtica             |                   |
| Podocarpus  | Morella (Myrica) | Brassicaceae   | Vernonia           |                   |
| Prunus      | Myrsine     | Blaeria         | Capparidaceae      |                   |
| Alangium    | Pinus *     | Cadaba          | Cissus             |                   |
|             | Polysicas   | Fabaceae        | Commelina          |                   |
|             | Psidium     | Cyathula        | Fagonia            |                   |
|             | Rhus        | Erica           | Gomphrena          |                   |
|             |             | Euphorbia       | Heliotropium       |                   |
|             |             | Hypericum       | Hypoestes          |                   |
|             |             | Impatien        | Liliaceae          |                   |
|             |             | Ipomea          | Narcissus          |                   |
|             |             | Justicia        | Polygonum          |                   |
|             |             |                 | Rubia              |                   |
|             |             |                 | Trianthema         |                   |
|             |             |                 | Poaceae            |                   |

*Pinus and *Cupressus are agroforestry neophytes at the study site.

Figure 5. Relative pollen abundances of the terrestrial pollen sum from the Nyabuiyabui sediment core, radiocarbon dates (‘+’ symbols on the left, black font for dates used in the suggested age-depth model in Fig. 2) and charcoal (>125 μm) for comparison. Aquatic taxa are shown as relative abundances of the total aquatic pollen sum (Table 2).
from the different proxies are plotted in a single stratigraphic plot (Fig. 6) to ease comparison across datasets. Significant changes in each of the proxies occur at around the same time, implying that the changes noted have the same or similar drivers.

Discussion
The Nyabuiyabui wetland sediment record covers the interval from the end of the Late Pleistocene (~17 000 cal aBP) to the present and provides insight into the dynamics of the wetland as well as the wider Mau Forest ecosystem. Given its geographical location, insights from the Nyabuiyabui sediment record have relevance for areas downstream within the Sondu and the Mara River catchment. The results also provide another comparison point of long-term environmental change in the East African highlands.

Although lacustrine sedimentary archives are often preferred due to their high time resolution and reliable dating (Ojala et al., 2012), the lack of undisturbed lakes in the Mau Forest necessitates investigations of wetland palustrine sediments to generate knowledge of past ecosystem and environmental changes. The temporal resolution of wetland sediment records in eastern Africa is variable and these shallow water ecosystems frequently experience higher levels of physical and bioturbation due to their small size and volume compared to lakes (Rucina et al., 2010; Githumbi, 2017). Sedimentary hiatuses have frequently been observed during the Late Pleistocene to Early Holocene from lacustrine and palustrine sediment records across eastern Africa, such as the Rukiga Highlands (Taylor, 1990), Laikipia Plateau (Taylor et al., 2005), Munsa in Uganda (Leju et al., 2005), Mount Kenya (Street-Perrott et al., 2007; Rucina et al., 2009), Eastern Arc Mountains in Tanzania (Mumbi et al., 2008; Finch et al., 2009, 2014), Pare Mountains in Tanzania (Heckmann, 2014), the Ruliji Delta (Punwong et al., 2013) and Virunga (McGlynn et al., 2013).

The distribution of radiocarbon dates collected from Nyabuiyabui is complex and reveals some age–depth reversals, which has been observed in several palustrine sediment studies in the region (Hamilton, 1982; Courtney Mustaphi and Marchant, 2016). This has been observed in lowland wetlands (Awuor, 2008; Öberg et al., 2012; Githumbi et al., 2017a,b; Gorman et al., 2020) and montane wetlands (Bonnefille and Riollet, 1988; Mumbi et al., 2008; Finch et al., 2009). The radiocarbon dates from Nyabuiyabui group the sediment stratigraphy into a basal pre-Holocene (Late Pleistocene), Early to Mid-Holocene, and Late Holocene section. By focusing our age–depth model on the dates that are derived from macrofossils, we are able to construct a coherent age–depth relationship that is very useful for assessing broad patterns of sedimentological and vegetation change at Nyabuiyabui (Fig. 6). Similar to several other palaeoenvironmental records established from palustrine sediments, the radiocarbon date uncertainties limit the precision of exploring the specific timing of events or rates of change. However, the vegetation change is explored and discussed with reference to depth to acknowledge this uncertainty. We describe the long-term variability in forest composition and fire during the broad periods of the Late Pleistocene, Early to Mid Holocene and Late Holocene.

Changes in pollen composition and abundance throughout the core indicate the persistence of an upland forest dominated by Podocarpus, Cordia, juniperus and Olea with varying degrees of openness possibly caused by ecological turnover and a variable fire regime. Podocarpus, Cordia, juniperus and Olea appear in all the samples, Afrotropical taxa in general appear in all pollen zones. Pollen zone NBPO2L2A (Late Pleistocene to Early Holocene transition period) experienced a loss of montane forest taxa (Abutil, Alangium, Commiphora, Fagonia, Lannea, Maytenus, Polyscias, Psidium, Rumex and Syzygium). The disappearance as well as the general decrease in tree and shrub taxa is accompanied by an increase in Poaceae and aquatic taxa. This suggests a drying environment with a contraction in the open water area providing more extensive shallow-water littoral areas for taxa such as Typha and Cyperaceae to become established. The general trend towards the top of the core is an increase in herbaceous taxa compared to the woody taxa.

Figure 6. A stratigraphic plot of radiocarbon dates, lithology and pollen and ITRAX CONISS zonations, and select palaeoecological and sedimentological measurements. Pollen percentages are grouped into Afrotropical and herbs, Poaceae, and aquatic pollen counts, as well as charcoal (>125 μm), organic, clay and sand contents. [Color figure can be viewed at wileyonlinelibrary.com]
Late Pleistocene to Early Holocene development of Eastern Mau: 538–240 cm

The end of the Late Pleistocene (540–220 cm) exhibits the highest arboreal pollen diversity (number of taxa identified) and abundance (total counts in each sample) while the herbs and grasses are at minimal abundances. This diversity is in part due to turnover at 330 cm to an ecosystem dominated by Cordia, Croton, Ficus, Juniperus and Olea from one dominated by Apodytes, Celtis, Dracaena, Hagenia and Podocarpus. The turnover signifies a change from Afromontane taxa that prefer a cooler, dry environment to one characterized by more mesic conditions. This ecosystem transition is similar to that documented by pollen records from the Rukiga Highlands (Taylor, 1990), Ruwenzori (Livingstone, 1967), Burundi highlands (Bonnefille and Riollet, 1988), Lake Albert (Beuning et al., 1997), Mount Elgon (Hamilton, 1987) and Mount Kenya (Street-Perrott et al., 2007; Rucina et al., 2009). This interval is also characterized by the highest biomass burning with high macroscopic charcoal concentrations implying a continuous connected source of fuel. Juniperus, a pioneer species colonizing gaps after a fire, is a dominant taxon in this forest transition, and more broadleaved species such as Olea are also established. The successional role of Juniperus in forests with recurring fire events is observed on Mount Kenya (Rucina et al., 2009) and the southern Aberdare Range (Bussmann, 2001). In the Nyabuiyabui pollen record, peaks in Juniperus lag behind those in Hagenia and may signal a sequence of ecological response to fire. The increase in Afromontane and tree taxa corresponds to increases in magnetic susceptibility, silt and clay as well as peaks in detrital elements. The detrital elements, silicon (Si), titanium (Ti), iron (Fe), rubidium (Rb) and strontium (Sr), show increased terrigenous input indicating higher erosional inputs to the basin through surface runoff conditions after episodes of heavy or continuous rainfall.

Several East African Pleistocene palaeoenvironmental records are dominated by signals inferring cooler, drier conditions such as forest compositional changes to semi-deciduous forest and lower lake levels (Van Zinderen Bakker and Coetzee, 1988; Sonzogni et al., 1998; Olago, 2001; Chalié and Gasse, 2002). Late Pleistocene sediments from the Rumii Swamp from Mount Kenya or the Lake Emakaat record show stratigraphic changes as well as increases in wetland fringe taxa (Cyperaceae, Poaceae and Typha), indicating lower water levels (Rymer et al., 2006; Rucina et al., 2009). However, the Lake Challa record indicates increased precipitation due to the intensification of the south-easterly Indian Ocean Monsoon from ~16 500 cal a BP to the Early Holocene that was interrupted during the Younger Dryas between ~13 300 and 11 700 cal a BP (Verschuren et al., 2009).

The varied responses among palaeo-vegetation data from different highland areas of equatorial eastern Africa suggest heterogeneity in hydroclimate-vegetation interactions since the Late Pleistocene to present day. This is not surprising given the local topographic-climate system feedback or position of the mountain in bioclimatic space being important controls on the response of the ecosystem through time (Hamilton, 1982; Loomis et al., 2017; Los et al., 2019). The drivers responsible for changes through time recorded by multiple palaeoenvironmental proxies across different mountains are uncertain; even for those records that are characterized by similar climate change, there are localized factors such as topography, hydrology and soil development to take into account. Given the differential response across the East African Mountains, a comparison of available studies across mountain ranges will improve our understanding of coherent responses across different mountain ecosystems and lead to an understanding of typologies of ecosystem response to rapid climate transitions.

The Early and Middle Holocene: 240–100 cm

At Nyabuiyabui, the low Ti, Fe and Mn concentrations in the sediments from 256 to 213 cm could indicate increasing dryness (Burrows et al., 2016). There is also a marked decrease in the number of pollen taxa identified and abundance until 190 cm, while the contribution of Poaceae pollen increases, implying a relatively warm and dry interval. The forest becomes increasingly open after 190 cm (the Early Holocene), and the subalpine forest association of Hagenia–Juniperus is interpreted as an early successional phase following fire-induced disturbance (Bussmann, 2001). There is an increase in Apodytes, Celtis, Olea, Podocarpus and Erica representative of montane forest. On Kilimanjaro, a warm and wet climate enabled the development and expansion of Afromontane forest around this period (Schüler et al., 2014). The Early Holocene experienced the highest increase in organic matter and decrease in carbonate content, accompanied by an increased silt and clay content as well as increased bulk density, implying high sedimentation. The increased organic matter content peaking with sulphur content could indicate that the increased sedimentation was anoxic (Tierney and Russell, 2007).

Compared to the Late Pleistocene, the climate of equatorial eastern Africa was generally warm and wet (Bonnefille and Riollet, 1988) and may signal a sequence of ecological response to fire. The increase in Afromontane and tree taxa corresponds to increases in magnetic susceptibility, silt and clay as well as peaks in detrital elements. The detrital elements, silicon (Si), titanium (Ti), iron (Fe), rubidium (Rb) and strontium (Sr), show increased terrigenous input indicating higher erosional inputs to the basin through surface runoff conditions after episodes of heavy or continuous rainfall.

The Late Holocene: 100 cm to the top

Increasingly arid conditions are recorded in the Nyabuiyabui record with continued replacement of Afromontane taxa with...
began in Kenya ~4000 cal aBP, i.e. pastoralism at ~4000 cal aBP, Neolithic hunters and herders. The spread of agropastoralism~500 cal aBP, and foraging declines ~250 cal aBP, altering landscapes through grazing and cultivation (Archibald et al., 2005; García et al., 2005). Global transformations of the landscape are hypothesized to have reached significant levels globally ~3000 cal a BP (Ellis et al., 2020). Data from eastern Africa indicate a shift in the adoption of livelihood patterns, for example as indicated by Wright (2005) while exploring resource exploitation among Neolithic hunters and herders. The spread of agropastoralism in Africa is believed to have occurred from 4000 cal a BP, altering landscapes through grazing and cultivation (Archibald et al., 2012; Phillips et al., 2020). Different land-use activities began in Kenya ~4000 cal a BP, i.e. pastoralism at ~4000 cal a BP, extensive agriculture at ~1000 cal a BP and intensive agriculture ~500 cal a BP, and foraging declines ~250 cal a BP (Stephens et al., 2019). Archaeological studies in this landscape would improve our understanding of resource use in the catchment; unfortunately, the two closest archaeological sites (burial caves) have not unearthed information about forest and wetland resource use (Faugust and Sutton, 1966; Merrick and Monaghan, 1984).

Several severe arid events are recorded within the Late Holocene; a diatom-chironomid record from Lake Naivasha using salinity to infer lake levels over the last 11000 years shows several intervals more arid than any recorded in the 20th century (Verschuren et al., 2000). Sediments from Lake Tanganyika and Lake Kivu record salinity peaks with Lake Kivu having salinity levels three times higher than the modern lake (Haberyan and Hecky, 1987). Within the Nyabuiyabui record, geochemical analysis shows a drop in all element concentrations except Cl, Ar, Cu, Hg and Pb. Increased Cl often results from precipitated chloride of NaCl that occurs during dry conditions (Kristen, 2009). The Ti/Rb ratio is useful to understand sediment source (Arnaud et al., 2014); this peaks twice in the Late Holocene (Fig. 6) at the same time as the Ti/Zn ratio and increases in sand and could indicate sedimentation of eroded material in the area. Lake Bogoria records a decline in high-altitude forest taxa (Kiage and Liu, 2006), whereas there is an increase in drought-tolerant taxa on Mounts Kenya and Elgon (Hamilton, 1982; Vincens, 1986), which imply the establishment of more arid conditions. This loss in high-altitude taxa could also be a signal of widespread clearance recorded in several montane forest sites (Bessems et al., 2008; Kiage and Liu, 2009; Rucina et al., 2009; Gelorini et al., 2012) attributed to land use activities across eastern and central Africa.

Significant changes to forest compositions have been inferred from the Eastern Arc Mountains during the past 2000 cal a BP (Heckmann, 2014; Finch et al., 2017) that potentially relate to intensified mountain forest resource use (Illes et al., 2018). A sediment record from the Mara basin records a significant increase in sedimentation as well as increasing Hg levels from the late 1700s (Dutton et al., 2019) that resonates with our insights from the sedimentary record. In the Nyabuiyabui record, there are high levels of introduced Cupressus and Pinus taxa, and increased levels of Asteraceae, Acanthaceae and Verbenaceae. Cupressus and Pinus provide markers for the onset of colonial forestry operations in the early part of the 20th century (Finch et al., 2014). There is an increasing trend in element concentrations, with spikes in Cu, S, Hg, Ar and Pb towards the top of the sediment record (~25 cm). The onset of the increase corresponds to the advent of industrialization and thus in deposition of heavy elements while the later changes would be due to increased human presence and mechanized agroforestry (Troup, 1932). The Upper Mara was identified as a major source of sediment in the Mara wetland (Dutton et al., 2019) and so increased sedimentation would correspond to increased erosion and runoff in the upper Mara. In recent history (~200 years), eastern African montane forests were either reserved, industrially logged or maintained locally as culturally valued spaces during colonial government rule. Many Afrormontane forests were converted to agroforests by the 1930s (Troup, 1932; Wood, 1965) with the Kenya Forest Service managing the higher elevation forests for timber production in public-private partnerships following independence.

Recent land-cover change within the Mau Forest (Landsat data from 1986) indicates that forest cover has consistently shrunk in areal extent with an increase in cropland and grassland cover. These changes are correlated strongly with rapid population increase (Odawa and Seo, 2019). Increased population pressure leading to deforestation and illegal logging, together with land cover conversion, mainly to agriculture and settlement, in the Mau escarpment have been identified as contributing to the overall degradation of the Mara basin and changed hydrological and sedimentation regime (Defersha and Melesse, 2012). The upper catchment of the Mara River drains south-east of Mau, with 65% of the catchment area located in Kenya and the rest in Tanzania (Defersha and Melesse, 2012; Mwangi et al., 2016a) covering both the Maasai-Mara and Serengeti wildlife areas and important wetland areas in estuaries with Lake Victoria, such as Lake Masiro (Dutton et al., 2019).

The two main Mau land cover types are forest and grassland, in which rapid conversion to cropland and bare land is taking place. Controlling for slope, organic matter content and extreme rainfall events, a study combining field and modelling data to understand the effects of the land-cover change in the Mara basin identified deforestation as a significant cause of change in water quality and watershed degradation, particularly in the upper sections of the Amala and Nyangores tributaries into the Mara River (Defersha and Melesse, 2012). A study by Mwangi et al. (2016b), using the Soil and Water Assessment Tool (SWAT) to understand the effects of agroforestry on the Mara hydrology, concluded that agroforestry impacts cannot be generalized across a catchment without considering climate variability within the watershed. Increased deforestation and conversion to agriculture in the Mau Highlands increases the variability of catchment water flows. As observed in a paired study in the Kapchorwa catchment, peak discharge flow increased significantly after the deforestation of the Nanadi/Kakamega tropical rainforest due to the loss of ground cover that regulates surface runoff (Mwangi et al., 2016a,b). However, areas under current...
agricultural production when converted to agroforestry lead to a decrease in runoff while increasing groundwater uptake – this is often because of the choice of tree species such as Eucalyptus that has a massive water demand (Hubbard et al., 2020). Land use, predominantly deforestation and transition to agriculture, was found to influence nitrous oxide levels in the lower elevations of the Mara River network (Mwanake et al., 2019). The effect of forest cover loss leading to reduced capacity to act as a catchment during heavy rainfall and mediate streamflow was identified as a major disruptor to economic activities at the south-west of Mau (Otouma et al., 2012). Various streams connecting into the Sondu River catchment experience steep increases (floods) after a heavy downpour followed by long intervals of very low stream flow. This has adverse effects on tea plantations and the Sondu hydroelectric power station, which cannot adequately predict waterflow needed for planning (Otouma et al., 2012).

Despite the net loss in forest cover, some cropland and grassland areas have converted back to forest (Odua and Seo, 2019). Natural regeneration coupled with reforestation grassland areas have converted back to forest (Odawa and (Otuoma et al., 2012). Despite the net loss in forest cover, some cropland and grassland areas have converted back to forest (Odua and Seo, 2019). Natural regeneration coupled with reforestation grassland areas have converted back to forest (Odawa and (Otuoma et al., 2012). Despite the net loss in forest cover, some cropland and grassland areas have converted back to forest (Odua and Seo, 2019).

Conclusions

Nyabuiyabui is a unique montane forest wetland record providing insights into long-term forest dynamics due primarily to climate change as well as long-term wetland development for the largest last remaining closed-canopy forest in eastern Africa, with significant impacts across the wider landscape including the Mara river system. The Nyabuiyabui catchment has undergone significant changes in forest composition from a highly diverse Afromontane to a more open forest ecosystem indicative of increased aridity in the region.

Precipitation change is an important driver of vegetation composition over time. The additional effects of fires have been an important component of Mau forests, glades and vegetated wetlands. At Nyabuiyabui, the significant increase in elements such as Pb and Hg over the last 200 cal a BP combined with the rise of exotic taxa are clear indicators of human forestry activity at industrial scale that would impact forest ecosystems and associated service delivery across the catchment. Further focused studies on other wetlands located along the Mau Forest complex would greatly improve our understanding of the recent past and interpret how these changes on forest cover impact on hydrology and downstream impacts.

Managing and ensuring an intact and functioning forest-hydrological system is vital for the Mau Highlands and the wider lowland savanna ecosystems and livelihoods. Long-term data on vegetation change of highland watersheds provide useful context for current climate and land use change debates and how these insights can support management decisions for remediation efforts and future restoration outcomes.

Supporting information

Additional supporting information may be found in the online version of this article at the publisher’s web-site.

**Figure S1.** A stratigraphic plot of the complete pollen taxa identified and the associated CONISS pollen assemblage zonation.

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**Abbreviations.** asl, above sea level; LOI, loss-on-ignition; XRF, X-ray fluorescence.

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