RESEARCH ARTICLE

Clonal growth strategy, diversity and structure: A spatiotemporal response to sedimentation in tropical *Cyperus papyrus* swamps

Addisie Geremew¹*, Iris Stiers¹, Tim Sierens¹, Alemayehu Kefalew², Ludwig Triest¹

¹ Laboratory of Plant Biology and Nature Management (APNA), Department of Biology, Vrije Universiteit Brussel (VUB), Brussels, Belgium, ² Department of Plant Biology and Biodiversity Management, College of Natural Sciences, Addis Ababa University, Addis Ababa, Ethiopia

* addisie2009@yahoo.com, ageremew@vub.ac.be

Abstract

Land degradation and soil erosion in the upper catchments of tropical lakes fringed by papyrus vegetation can result in a sediment load gradient from land to lakeward. Understanding the dynamics of clonal modules (ramets and genets) and growth strategies of plants on such a gradient in both space and time is critical for exploring a species adaptation and processes regulating population structure and differentiation. We assessed the spatial and temporal dynamics in clonal growth, diversity, and structure of an emergent macrophyte, *Cyperus papyrus* (papyrus), in response to two contrasting sedimentation regimes by combining morphological traits and genotype data using 20 microsatellite markers. A total of 636 ramets from six permanent plots (18 x 30 m) in three Ethiopian papyrus swamps, each with discrete sedimentation regimes (high vs. low) were sampled for two years. We found that ramets under the high sedimentation regime (HSR) were significantly clumped and denser than the sparse and spreading ramets under the low sedimentation regime (LSR). The HSR resulted in significantly different ramets with short culm height and girth diameter as compared to the LSR. These results indicated that *C. papyrus* ameliorates the effect of sedimentation by shifting clonal growth strategy from guerrilla (in LSR) to phalanx (in HSR). Clonal richness, size, dominance, and clonal subrange differed significantly between sediment regimes and studied time periods. Each swamp under HSR revealed a significantly high clonal richness ($R = 0.80$) as compared to the LSR ($R = 0.48$). Such discrepancy in clonal richness reflected the occurrence of initial and repeated seedling recruitment strategies as a response to different sedimentation regimes. Overall, our spatial and short-term temporal observations highlighted that HSR enhances clonal richness and decreases clonal subrange owing to repeated seedling recruitment and genets turnover.
Introduction

Clonality is a widespread life history trait in angiosperms that maximizes plant fitness [1, 2] and is significant for population persistence under a heterogeneous and stochastic environment [3, 4, 5]. However, the downside of persistent clonal reproduction is a complete loss of recombination that could lead to extinction [6]. Although the reaction norm is species dependent, clonal plants have developed different adaptive strategies to contrasting environmental gradients [5, 6, 7]. Shifting clonal growth strategy is an adaptive mechanism crucial for colonization, establishment, and distribution of clonal plants [6, 8, 9, 10]. The two clonal growth strategies, guerrilla and phalanx, reveal endpoints of ramet arrangement in space [5, 11]. The guerrilla strategy produces loosely organized spreading growth modules that enable plants to escape adverse environmental conditions and competition [5, 11]. In contrast, the phalanx strategy results in clumped ramets occupying suitable patches to use profuse resources [5, 12]. Under heterogeneous environments, ramets expand and produce intermingled distribution among genets [13, 14]. Recent evidences show that the change in allele frequencies in response to an environmental gradient acts as an indicator of the effect of selective processes that modulate population genetic structure [15, 16]. Similarly, the shift in clonal growth strategy and genet frequencies may reflect how clonal plants are resilient to dynamic environmental factors that finally lead to clonal structuring and genetic divergence. The dynamics of the ramets and genets of clonal plants in both space and time is critical to explore species adaptation and processes regulating population differentiation [14]. In addition to the clonal growth strategy, clonal plants have the capacity to reproduce both sexually and vegetatively, and the balance between these reproduction modes regulates how genets colonize space and recruitment [17]. Depending on the turnover of genets, two recruitment strategies have been known for clonal plants [18]: (i) the 'Initial Seedling Recruitment' (ISR) strategy; and (ii) the 'Repeated Seedling Recruitment' (RSR) strategy. Under the ISR strategy, the development of a population includes selective elimination of genotypes, as a result the population ends up with a small number of large and evenly aged clones. In contrast, RSR strategy enables survival of small clones of different age and size to coexist and leads to high genetic and clonal diversity within a population [19]. Differential intensity between these recruitment strategies dictates clonal richness [19, 20] and contributes to patch growth, clone coexistence and evolution, and disturbance recovery [18, 21]. Therefore, in examining the populations of a species occurring in different environments, it is important to explore how each reproduction mode and recruitment strategy contributes to genets distribution and dynamic.

Sedimentation affects plant growth and distribution in lacustrine wetlands by altering moisture level, aeration, temperature and other factors of substrate-plant-microenvironment conditions [22, 23]. Despite contradictory conclusions, a shift in clonal growth strategy (from phalanx to guerrilla and vice versa) has been reported as an adaptive mechanism to sediment burial by a few wetland and coastal dune plants such as Suaeda salsa [24], Phalaris arundinacea [25] and Carex brevicuspis [11]. Arnaud-Haond et al. [26] found no significant correlation between sedimentation rate and plasticity of clonal traits in a seagrass but rather, with the demographic parameter, mortality. More specifically, severe land degradation and soil erosion in the upper catchment of tropical lakes bordered by Cyperus papyrus L. dominated vegetation result in a sediment load gradient from land to lake ward [27]. Investigating the shift in clonal growth strategy, exploring the genotypic diversity and structure metrics along such a sediment gradient may shed light on the resilience of C. papyrus to sediment accretion and help to forecast the evolutionary potential of a population [28]. A high level of genotypic diversity is related to increased resistance [29] and resilience [30]. It is possible that the spatial variability of sediment loads can create microenvironments that are significant for the persistence of
multiple genotypes through seedling recruitment, and at the same time favor directional selection and increase genotypic diversity within a population [31].

Biologically productive papyrus swamps, dominated by the keystone species paper reed (Cyperus papyrus L) have been widely recognized for their valuable regulatory and provisional ecosystem services [32, 33, 34, 35, 36, 37, 38, 39]. More explicitly, these swamps control sediment load into the adjoining waterbodies [40, 41, 42]. Recent quantitative studies have shown that catchments with disturbed papyrus swamps resulted in sediment yield three times greater than those with pristine papyrus vegetation [43, 44]. Since most of the papyrus swamps are threatened by anthropogenic pressures [27, 37], there is an imperative need for restoration and management measures to reduce sediment accretion in adjacent lakes and wetlands, supported by serious study of the population dynamics and adaptive response of the dominant and keystone species, C. papyrus. The effect of sedimentation on the growth, distribution and diversity of clonal plants in coastal ecosystems is well known [7, 45, 46]. In contrast, to our knowledge, there is less information on the effect of sedimentation on the responsive survival strategy, i.e. dynamics in seedling recruitment strategy, clonal growth strategy, structure and diversity of emergent macrophytes at the temporal and spatial scales.

In this study, we combined morphological traits and genotype data using 20 novel microsatellite markers to assess the spatial and temporal dynamics in clonal growth, diversity and structure of the paper reed (C. papyrus), as a response to sedimentation regimes. We hypothesized that a higher level of sedimentation would lead to: (1) increased occurrence of phalanx strategy over guerrilla strategy; (2) increased ramet density and genet size; and (3) higher genotypic diversity through seedling recruitment. Our findings contribute to the understanding of population dynamics, resilience and survival strategy of clonal plants, especially of the key- stone species C. papyrus in the context of sedimentation. It will also provide an important guideline for designing conservation and restoration plans of C. papyrus and protection of lakes from over-sedimentation.

Materials and methods
Species description
The paper reed, Cyperus papyrus (Cyperaceae), is a perennial rhizomatous keystone emergent species native to Africa that occurs particularly along freshwater lakes, rivers and stream margins. In the Lake Tana region, papyrus forms monocultures and co-occurs with other emergent macrophytes, such as Typha latifolia, Polygonum senegalense, Phragmites australis, Vossia cuspidata, Echinocloa stagnina and E. pyramidalis [47]. The species occupied the center of the lake and its basin between 16,700 and 15,100 BP following an intermittent flooding event after draw down [48]. The species produces inter-meshed horizontal rhizomes that shield the lake and adjacent wetlands from over-sedimentation and pollutants [40]. Reproduction is primarily sexual via very small seeds and asexual through extension of horizontally creeping rhizomes. In the Lake Tana basin, it flowers and fruits from June to August during the rainy season. Due to over-exploitation and ongoing habitat conversion [27], C. papyrus is locally on the brink of extinction. The short term threats come from direct habitat loss and damage from over-collection [49].

Study area and sampling sites
The field study was carried out in three papyrus wetlands situated in the Lake Tana basin (Fig 1), which is perched within a large dome of the Ethiopian highland. This large freshwater body (area 3,200 km²; mean depth 9 m) along the Blue Nile River is adjoined by seasonally and permanently flooded wetlands and also fed by a dendritic network of rivers [50]. The climate in
the basin is a tropical monsoon with a seasonal distribution of precipitation gauged by a shift in the inter-tropical convergence zone. Comprehensive accounts referred to the hydrology, climate, and lake level fluctuations [43, 51, 52]. The islands, peninsulas, adjacent wetlands, and flat depression areas are characterized by several soil types (e.g., Vertisols, Chromic Luvisols, Eutric Luvisols, and Lithic Leptosols). Because of severe land degradation and soil erosion in the adjoining landscape [53], alluvial sand deposition around the lake is amplified [43, 44, 54].
Three papyrus swamps, namely Gilgel Abay, Robit, and Sebatamye, were selected by considering differences in depth, large coefficients of runoff, and sediment deposition gradients from the shore of the lake (Fig 1; Table 2). The Robit swamp has inundation-drawdown cycles and agricultural encroachment in the surrounding landscape. Three perennial rivers from the east Tana catchment, the Gumara, Rib, and Gelda deposit a large proportion of the sediment load to Robit and other adjacent swamps resulting in sharp sedimentation gradients. The papyrus swamp located on the Gilgel Abay represents the stretches of the floodplain in the upper Blue Nile River in west Tana. The Gilgel Abay river delivers the highest amount (29%) of sediment load into Lake Tana and forms a visible delta with a clear gradient. The upper catchment of this area is also heavily disturbed and characterized by little vegetation cover. Unlike the Gilgel Abay and Robit swamps, the Sebatamye swamp is characterized by sediment depositions that were exported out of the lake via the Blue Nile River. This swamp is also typified by steep slopes and rocky substratum ahead of alluvial sediment deposition along the bank of the Blue Nile River and is upstream from the 40m Tis Isat fall.

**Sampling design and sample collection**

The field study was performed in 2014 and 2016. In 2013, a preliminary survey on the sediment gradient and vegetation zonation was conducted at the beginning of the rainy season. To verify sedimentation level and its effect on the depth of the wetland and the lake, wooden sticks of 10 m long were placed randomly until the next dry season. At each swamp, a transect was established landward to lakeward along the sediment gradients. Two rectangular plots, each 18 m x 30 m separated by 500 m, were established on two distinct sediment regimes: a high sediment level corresponded to the most shallow section (0.9 m sediment level) and a very low sediment regime represented a relatively deeper section of the lake (< 0.2 m sediment level). To characterize the clonal traits of papyrus, such as girth diameter, culm height, ramet density, spacer length (i.e. space between individual stands (shoots)), and biomass, ten 3 m x 3 m quadrats were delineated with cords in each plot. We used a Vernier caliper to measure girth diameter. Ramet density in a plot was estimated as the mean of the number of shoots per quadrat. Above the substrate, papyrus plants (n = 9 per quadrat) were clipped and sorted into leaves and culm. Subsequently, these parts were oven-dried at 70°C for 48 hours and weighed. Absolute aboveground biomass was determined as the sum of dried weights of leaves and culm. To monitor the temporal dynamics of ramets and genets and to avoid resampling (DNA extractions) of the same genet in 2016, all aerial shoots observed in 2014 within each plot were tagged using small, colored plastic strings.

To examine the pattern and dynamics of clonal diversity and structure, a total of 322 and 314 ramets shoots (40 to 60 per plot) were collected in 2014 and 2016, respectively. In order to minimize sampling of duplicates, individuals were sampled at an interval of three meters. The location of each sampling plot was recorded using a handheld GPS (Garmin 45XL, USA). Samples were dried with silica gel pending DNA extraction.

**DNA extraction and microsatellite generation**

Since our preliminary test using 11 microsatellite primers showed low polymorphism in *C. papyrus* populations of the Ethiopian highlands, additional polymorphic markers were required for an in-depth study of population connectivity, clonality, and seedling recruitment. To do so, genomic DNA was extracted from lyophilized green umbel rays (20 mg) of *C. papyrus* collected from Lake Tana (Ethiopia, 11°37.336’N, 37°20.542’) using the E.Z.N.A. SP plant DNA Mini Kit (Omega Bio-tek). The quantity of the DNA was determined by NanoDrop One Spectrophotometer (Thermo Scientific). Also, the quality of the DNA was evaluated following library DNA QC criteria. Twenty-five ng of DNA was subjected to Illumina sequencing on a
HiSeq, 100 bp paired-end library (Macrogen, Seoul, Republic of Korea). A total of 43.83 million reads of at least 80 bp, without a tag and adapter, were generated in FASTQ format, and we converted the FASTQ files into FASTA files using NET Bio v.1.1 (https://bio.codeplex.com). Both unassembled reads and contigs were applied to identify microsatellites and design primers using QDD2 v.2 [59] integrated with Primer3 [60]. After discarding the loci that showed compound SSR stretches, 5,510 target microsatellites displayed at least five pure repeats, a size of 100 bp PCR product and lowest penalty values. Subsequently, 70 primer pairs were selected for synthesis (Sigma Aldrich, Germany) and tested for amplification on eight papyrus samples.

**Primer testing and multiplexing**

Simplex PCRs were conducted in a total volume of 20 μL per reaction and contained: 0.2 mM of each dNTP (Promega), 2 mM MgCl₂, 0.5 μM of each forward and reverse primer (Sigma), 0.1 mM BSA, 1 U Taq DNA polymerase (Promega), and 3 μl (25 ng) template DNA. The PCRs were carried on a Bio-Rad Thermal Cycler using the profile of initial denaturation at 95°C for 3 min, followed by 35 cycles of 95°C for 0.4 min, annealing at 54°C for 1 min, extension at 72°C for 2 min and a final extension at 72°C for 5 min. To separate the PCR products we used capillary electrophoresis (QIAxcel Advanced System). The QX alignment marker of 15 bp/1600 bp (1.5 μl) and the QX size marker of 25 bp to 500 bp (5 μl) were employed. Fragments or allele size were visualized and established with QIAxcel Screen gel software that provides both electropherogram and gel images of DNA separation. Out of the 70 microsatellite loci tested, 30 produced scorable amplified products. A final subset of nine novel loci that appeared polymorphic and heterozygous, were fluorescently labeled (forward primers, Life Technologies, USA) and multiplexed in two sets, including the eleven primers developed by Triest et al. [58] using Multiplex Manager [61].

Multiplex PCRs were carried out for 420 samples from the three swamps within the Lake Tana basin in a 12.5 μL mix containing 6.25 μL 1x QIAGEN Master Mix (Valencia, USA), 2.5 μL of RNase free water, 2.5 μL of 25 ng DNA, and 1.25 μL of primers mix (see Triest et al. [58] for the reaction profile). The PCR products were resolved using ABI 3730xl Genetic Analyzer and fragment sizes were determined with an internal size standard GeneScan500 LIZ in Macrogen (Seoul, Republic of Korea). Allele sizes for each microsatellite locus per individual were scored with GeneMarker v2.4.1 (SoftGenetics, USA).

**Data analysis**

GenAlEx v. 6.5 [62] was employed to compute the number of alleles (Nₐ), expected (Hₑ) and observed (Hₒ) heterozygosity, and to assess departure from the Hardy–Weinberg equilibrium (HWE) at locus level. We also analyzed inbreeding coefficient (Fᵢₛ) and linkage disequilibrium (LD) among the 20 microsatellite loci implemented in FSTAT version 2.9.3.2 [63]. The test for presence of null alleles, large allele dropout, and genotyping errors was done with MICRO-CHECKER version 2.2.3 [64].

Clonal diversity was measured by different parameters using GENCLONE 2.0 [65]. Clonal richness (R) was estimated from the number of distinct multilocus genotypes (MLGs) (G) and the number of ramets (N) [66]. To estimate a comparable R across populations, we standardized the number of sample units collected (N) by the proportion of sample units collected at the site with higher density (Dₘₐₓ):  

\[ Nₛ = \frac{N}{Dₘₐₓ} \]  

(1)
We also applied the Pareto distribution index ($\beta$) as a proxy for clonal diversity [67] and the dispersion of ramets among genets. Spatial clone architecture and clonal structure were estimated among the six sampling plots (two plots per site) in three swamps by calculating the aggregation index ($Ac$) at plot level and clonal dominance index ($Dc$) for each genet using GENCLONE under 10,000 permutations. $Ac$ was computed as:

$$Ac = \frac{(P_{sg} - P_{sp})}{P_{sp}}$$

where $P_{sg}$ is the mean likelihood of clonal identity of all sample unit pairs, and $P_{sp}$ is the average probability of clonal identity among pairwise nearest neighbors. To determine the degree of intermingling among genets, $Dc$ was computed for each genet as:

$$Dc = \frac{(NR - 1)}{(NT - 1)}$$

where $NR$ is the genet size (i.e. genet size is the number of ramets per genet) and $NT$ is the total number of ramets sampled within the genet range[68]. A $Dc$ index of one points out that the spatial range of a genet is occupied fully by ramets of the identical genet. Only genets signifying more than two ramets were accounted for the calculation of $Dc$. Despite the edge effect was minimized by sampling across the center of a plot, we tested for its effect on dispersion of distinct or rare MLGs that overvalue clonal diversity following Arnaud-Haond et al. [67]. To estimate the minimum spatial extent of the largest clone in each plot, we determined the clonal subrange (CR) which corresponds to the maximum distance between two identical MLGs belonging to the same clone in meters. In other words, it is measured as the distance for which the probability of clonal identity becomes null [69]. Temporal and spatial differences in clonal growth, genotypic diversity, and structure measures were analyzed with a non-parametric Mann–Whitney U-test using SPSS XII (SPSS, Chicago). Spearman’s correlation coefficients were used to examine relationships between estimates of genotypic diversity, structure, and clonal growth traits. Furthermore a linear regression was carried out to assess the relationship between spacer length and ramet density.

**Results**

**Characteristics of the isolated microsatellites**

The 20 microsatellite loci were highly variable, having 3 (Cypap 62) to 12 (Cypap 23) alleles per locus over 282 ramets yielding 141 individuals (three populations) with distinct multilocus genotypes (MLGs) (Table 1). Levels of expected and observed heterozygosities per locus varied from 0.41 to 0.86 and from 0.25 to 0.87, respectively (Table 1). There was no evidence for large allele dropout or genotyping error. Putative null alleles were detected at only one locus (Cypap 28) in one population, with a very low frequency because of excess homozygotes ($P < 0.05$). The microsatellite loci developed and multiplexed showed a wide range of inbreeding coefficient ($F_{IS}$) of -0.046 (Cypap 10) to 0.611 (Cypap 15). The within-population tests revealed that there was no significant departure from the HWE ($P < 0.001$) nor LD between loci in the populations and thus, further analyses of clonal diversity and structures were carried out using these 20 loci.

**Clonal growth metrics**

All clonal growth metrics (culm height, ramet density, girth diameter, spacer length, and biomass) of *C. papyrus* were significantly different between the high sedimentation regime (HSR)
Mean culm height and girth diameter were about twice higher in the LSR compared with the HSR which contained dense ramets with short culms. Ramet density also varied by almost three orders of magnitude (Table 2), between the sedimentation regimes. The maximum density of *C. papyrus* ramets was 57.6 m$^{-2}$ at Robit

Table 1. Genetic diversity measures and characteristics of twenty microsatellite loci from *C. papyrus* genets (Total N = 141, per population = 47) of three populations of Lake Tana (Ethiopia).

<table>
<thead>
<tr>
<th>Loci</th>
<th>Primer sequences (5'-3')</th>
<th>Repeat motif</th>
<th>Accession No.</th>
<th>$T_{ar}$</th>
<th>Allele size</th>
<th>$A_T$</th>
<th>$A_E$</th>
<th>$H_O$</th>
<th>$H_E$</th>
<th>$F_{IS}$ (HWE)</th>
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<tbody>
<tr>
<td>Cypap22**</td>
<td>F: PET-TGGAGCTTACACGCA ATACGAGTTTCAACG &lt;br&gt; R: CACGGTAAATGCTAGAAAATTTT &lt;br&gt; (AAG) 14</td>
<td>KT873448</td>
<td>57</td>
<td>90–115</td>
<td>7</td>
<td>4.7</td>
<td>0.504</td>
<td>0.439</td>
<td>-0.148**</td>
<td></td>
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<tr>
<td>Cypap23**</td>
<td>F: NED-TGGTTCTTAATGTTTGGTATGCTT &lt;br&gt; R: TGGAGAGAGGAAGAATGTTTCAACG &lt;br&gt; (AG) 9</td>
<td>KT873449</td>
<td>57</td>
<td>98–135</td>
<td>12</td>
<td>7.6</td>
<td>0.871</td>
<td>0.591</td>
<td>-0.476**</td>
<td></td>
</tr>
<tr>
<td>Cypap27**</td>
<td>F: VIC-CATGCGCTCCCGGTTAATACTT &lt;br&gt; R: CAGAGTAGTGCCTCGAAGCTTTTC &lt;br&gt; (AT) 8</td>
<td>KT873450</td>
<td>57</td>
<td>82–124</td>
<td>6</td>
<td>4.3</td>
<td>0.393</td>
<td>0.762</td>
<td>0.491**</td>
<td></td>
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<tr>
<td>Cypap28**</td>
<td>F: 6FAM-ACTACACACAGACGTAACACT &lt;br&gt; R: TACAGTGGCGCTGTCTGCT &lt;br&gt; (AG) 9</td>
<td>KT873451</td>
<td>57</td>
<td>100–120</td>
<td>6</td>
<td>4.6</td>
<td>0.431</td>
<td>0.782</td>
<td>0.478** null</td>
<td></td>
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<tr>
<td>Cypap34*</td>
<td>F: VIC-TCTATATCACTATATCATCCTACAGG &lt;br&gt; R: GACACAGGCAACACCGAGA &lt;br&gt; (AAAGG) 8</td>
<td>KT873452</td>
<td>57</td>
<td>90–118</td>
<td>10</td>
<td>6.3</td>
<td>0.508</td>
<td>0.834</td>
<td>0.446**</td>
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<tr>
<td>Cypap38*</td>
<td>F: 6FAM-GGAGTAAAATGGCCATGAGG &lt;br&gt; R: CCATCTCTTCTCCTCTGCTCAAA &lt;br&gt; (AG) 11</td>
<td>KT873453</td>
<td>57</td>
<td>90–112</td>
<td>9</td>
<td>6.6</td>
<td>0.474</td>
<td>0.447</td>
<td>-0.062**</td>
<td></td>
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<td>Cypap52*</td>
<td>F: 6FAM-CCAAACACAGAGAGAAACAAA &lt;br&gt; R: ACTGCAGGGGTTATGGTTG &lt;br&gt; (CCA) 9</td>
<td>KT873455</td>
<td>57</td>
<td>218–260</td>
<td>6</td>
<td>4.6</td>
<td>0.326</td>
<td>0.768</td>
<td>0.592**</td>
<td></td>
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<tr>
<td>Cypap56++</td>
<td>F: 6FAM-CTCTTGTCCTTGTTGTTAAAGG &lt;br&gt; R: TGGAGGACCCGCTTGTCCTCC &lt;br&gt; (GA) 15</td>
<td>KT873456</td>
<td>57</td>
<td>230–260</td>
<td>6</td>
<td>4.6</td>
<td>0.326</td>
<td>0.768</td>
<td>0.592**</td>
<td></td>
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<tr>
<td>Cypap62++</td>
<td>F: PET-GAGAGAGGCACCTGACTAGC &lt;br&gt; R: TGGATATGACTCCAAGCATT &lt;br&gt; (AG) 5</td>
<td>KT873454</td>
<td>57</td>
<td>130–140</td>
<td>3</td>
<td>2.4</td>
<td>0.247</td>
<td>0.552</td>
<td>0.573**</td>
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<tr>
<td>Cypap4*</td>
<td>F: 6FAM-AACAGGTCTTAATGACTGAGTG &lt;br&gt; R: TGGTAGTTCTGCTGCTGCTGA &lt;br&gt; (TG) 10</td>
<td>KC460659</td>
<td>57</td>
<td>138–169</td>
<td>10</td>
<td>7.0</td>
<td>0.494</td>
<td>0.856</td>
<td>0.371**</td>
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<tr>
<td>Cypap14*</td>
<td>F: 6FAM-CATGCACATGCTTTTGATGA &lt;br&gt; R: TGTTCATTGATGCTGCTTT &lt;br&gt; (GT)n</td>
<td>KC460660</td>
<td>57</td>
<td>186–200</td>
<td>5</td>
<td>4.5</td>
<td>0.384</td>
<td>0.776</td>
<td>0.559**</td>
<td></td>
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<tr>
<td>Cypap10*</td>
<td>F: VIC-GACAGCGGCTTGTTAAAT &lt;br&gt; R: TCTCTGCCTTTCACACACTC A &lt;br&gt; (GT)7</td>
<td>KC460662</td>
<td>57</td>
<td>143–159</td>
<td>5</td>
<td>3.3</td>
<td>0.565</td>
<td>0.529</td>
<td>-0.046**</td>
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<tr>
<td>Cypap13F</td>
<td>F: NED-CTGGGCTATGCGATCAANAT &lt;br&gt; R: AAGCACAGGGTTATGGTTG &lt;br&gt; (GT)9(AAT)4</td>
<td>KC460664</td>
<td>57</td>
<td>151–160</td>
<td>4</td>
<td>2.4</td>
<td>0.290</td>
<td>0.575</td>
<td>0.393**</td>
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<td>Cypap13S</td>
<td>F: NED-CTGGGCTATGCGATCAANAT &lt;br&gt; R: AAGCACAGGGTTATGGTTG &lt;br&gt; (GT)9 (AAT)4</td>
<td>KC460664</td>
<td>57</td>
<td>163–175</td>
<td>4</td>
<td>3.4</td>
<td>0.409</td>
<td>0.699</td>
<td>0.258**</td>
<td></td>
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<tr>
<td>Cypap5*</td>
<td>F: NED-TGGAGTTATGGGCCCTCACACG &lt;br&gt; R: ATCTGACGGGACCCTTCTCC &lt;br&gt; (CT)13</td>
<td>KC460660</td>
<td>57</td>
<td>219–248</td>
<td>7</td>
<td>3.1</td>
<td>0.326</td>
<td>0.666</td>
<td>0.583**</td>
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<tr>
<td>Cypap3*</td>
<td>F: PET-AAAAGGTCTGACGCTGCCACG &lt;br&gt; R: AAGGAGGAAATGTTTGTTG &lt;br&gt; (CT)14(GTGAA)2</td>
<td>KC460658</td>
<td>57</td>
<td>154–217</td>
<td>5</td>
<td>3.3</td>
<td>0.408</td>
<td>0.684</td>
<td>0.561**</td>
<td></td>
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<tr>
<td>Cypap7**</td>
<td>F: 6FAM-GAAGCCAGAGGGAAAGGTGTG &lt;br&gt; R: CAAAAGGACACCTGCTGCT &lt;br&gt; (GT)6GC(GT)6(GA)9</td>
<td>KC460661</td>
<td>57</td>
<td>172–188</td>
<td>7</td>
<td>4.6</td>
<td>0.429</td>
<td>0.783</td>
<td>0.453**</td>
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<tr>
<td>Cypap12**</td>
<td>F: VIC-TGATTTCTCTGGAGCGGCTTGA &lt;br&gt; R: AGACCCACACACCGAAAAA &lt;br&gt; (TC)8</td>
<td>KC460663</td>
<td>57</td>
<td>176–182</td>
<td>5</td>
<td>4.1</td>
<td>0.418</td>
<td>0.750</td>
<td>0.452**</td>
<td></td>
</tr>
<tr>
<td>Cypap15**</td>
<td>F: NED-CGAGAAGACATGCTCTTTAATGC &lt;br&gt; R: GGAGAAGAGAGGGTTTCCC &lt;br&gt; (CT)7(GA)6</td>
<td>KC460666</td>
<td>57</td>
<td>164–182</td>
<td>6</td>
<td>3.7</td>
<td>0.409</td>
<td>0.718</td>
<td>0.611**</td>
<td></td>
</tr>
<tr>
<td>Cypap1**</td>
<td>F: PET-AACGCCAAATGAGCAACAAA &lt;br&gt; R: TGGTTTGGTTGTTGTTGA &lt;br&gt; (CAGA)10</td>
<td>KC460657</td>
<td>57</td>
<td>173–217</td>
<td>11</td>
<td>7.3</td>
<td>0.466</td>
<td>0.857</td>
<td>0.583**</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>_</td>
<td>_</td>
<td>_</td>
<td>_</td>
<td>6.8</td>
<td>4.7</td>
<td>0.438</td>
<td>0.694</td>
<td>0.335</td>
<td></td>
</tr>
</tbody>
</table>

$T_{ar}$ annealing temperature, $A_T$ total number of alleles, $A_E$ effective number of alleles, $H_O$ observed heterozygosity, $H_E$ expected heterozygosity, $F_{IS}$ inbreeding coefficient with associated tests of deviation from Hardy-Weinberg equilibrium (HWE, ns = not significant, ** significant at 0.01 level), F hex-labelled forward primer, R reverse primer, * locus in multiplex-1, ** locus in multiplex-2, null alleles, shaded primers developed by Triest et al. [58] and non shaded, new primers.

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swamp. Clumped ramets of HSR populations produced shorter spacer lengths over spreading ramets than did those in the LSR (Table 2), reflecting HSR favors phalanx strategy than the LSR that promotes the guerilla strategy. Papyrus under an HSR demonstrated significant temporal difference in ramet density and spacer length (S1 Fig).

**Dynamics of ramets and genotypes composition**

Permutation tests for genotypic resolution revealed an exponential increase with the number of loci and then an asymptotic trend subsequent to the addition of locus (S2 Fig). We identified a total of 408 distinct MLGs across the 636 ramets (N = 301 for 2014 and N = 335 for 2016). The likelihood of encountering identical MLGs twice because of random sexual recombination events was very low (P<sex < 0.01, Table 3). Frequency distribution of allelic distances between

### Table 2. Clonal growth traits of *C. papyrus* across sedimentation regimes.

<table>
<thead>
<tr>
<th>Swamp</th>
<th>Plot</th>
<th>Latitude (N)</th>
<th>Longitude (E)</th>
<th>Altitude (m)</th>
<th>Sediment regime</th>
<th>Ramet density (m⁻²)</th>
<th>Culm height (cm)</th>
<th>Girth diameter (cm)</th>
<th>Spacer length (cm)</th>
<th>Biomass (gm⁻²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Robit</td>
<td>1</td>
<td>11°44'24.51&quot;</td>
<td>37°25'22.42&quot;</td>
<td>1788</td>
<td>High</td>
<td>57.6 (1.85)</td>
<td>307.9 (11.2)</td>
<td>7.4 (0.54)</td>
<td>48.4 (2.66)</td>
<td>5749.9 (93.1)</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>11°44'16.13&quot;</td>
<td>37°25'18.00&quot;</td>
<td>1788</td>
<td>Low</td>
<td>17.0 (1.19)</td>
<td>463.5 (10.9)</td>
<td>16.2 (1.04)</td>
<td>64.3 (1.75)</td>
<td>4609.1 (96.1)</td>
</tr>
<tr>
<td>Sebatamye</td>
<td>1</td>
<td>11°30'57.02&quot;</td>
<td>37°30'36.22&quot;</td>
<td>1691</td>
<td>High</td>
<td>41.0 (1.93)</td>
<td>240.7 (14.5)</td>
<td>6.8 (0.34)</td>
<td>55.5 (1.79)</td>
<td>5127.6 (108)</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>11°30'57.63&quot;</td>
<td>37°30'53.54&quot;</td>
<td>1690</td>
<td>Low</td>
<td>11.8 (1.02)</td>
<td>366.5 (5.03)</td>
<td>13.6 (1.01)</td>
<td>73.7 (1.31)</td>
<td>3491.0 (71.6)</td>
</tr>
<tr>
<td>Gilgel Abay</td>
<td>1</td>
<td>11°52'00.22&quot;</td>
<td>37°06'51.17&quot;</td>
<td>1788</td>
<td>High</td>
<td>45.0 (1.02)</td>
<td>381.3 (6.73)</td>
<td>15.3 (1.55)</td>
<td>66.6 (1.52)</td>
<td>4199.6 (75.6)</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>11°51'43.88&quot;</td>
<td>37°06'41.17&quot;</td>
<td>1788</td>
<td>Low</td>
<td>14.0 (1.02)</td>
<td>326.2 (15.5)</td>
<td>5.3 (0.39)</td>
<td>53.0 (1.98)</td>
<td>9073.5 (241)</td>
</tr>
</tbody>
</table>

The significance level is based on T-test for two samples, d.f = 1 and P < 0.05). Different letters indicate significant difference between sedimentation regimes. Values in the bracket show standard error of the mean based on n = 50 (five plants per quadrat for total of 10 quadrats per swamp).

https://doi.org/10.1371/journal.pone.0190810.t002

### Table 3. Clonal diversity and structure measures of *C. papyrus* ramets sampled in three papyrus swamps in Lake Tana under two sedimentation regimes during 2014 and 2016.

<table>
<thead>
<tr>
<th>Swamp</th>
<th>Plot</th>
<th>Sediment regime</th>
<th>Year</th>
<th>N</th>
<th>G</th>
<th>Psex max</th>
<th>R</th>
<th>D</th>
<th>E</th>
<th>β</th>
<th>NR</th>
<th>DC</th>
<th>Ac</th>
<th>Ee</th>
<th>CR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Robit</td>
<td>1</td>
<td>High</td>
<td>2014</td>
<td>60</td>
<td>48</td>
<td>9.11 x 10⁻¹⁹</td>
<td>0.79</td>
<td>0.99</td>
<td>0.92</td>
<td>3.58</td>
<td>12</td>
<td>0.186</td>
<td>0.194*</td>
<td>0.031*</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>Low</td>
<td>2014</td>
<td>60</td>
<td>52</td>
<td>4.13 x 10⁻¹⁵</td>
<td>0.86</td>
<td>0.79</td>
<td>0.79</td>
<td>2.95</td>
<td>8</td>
<td>0.119</td>
<td>0.146*</td>
<td>0.023*</td>
<td>9.2</td>
</tr>
<tr>
<td>Sebatamye</td>
<td>1</td>
<td>High</td>
<td>2014</td>
<td>60</td>
<td>39</td>
<td>4.12 x 10⁻¹⁶</td>
<td>0.64</td>
<td>0.96</td>
<td>0.96</td>
<td>2.61</td>
<td>21</td>
<td>0.339</td>
<td>0.358*</td>
<td>0.103*</td>
<td>25</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>Low</td>
<td>2014</td>
<td>60</td>
<td>53</td>
<td>1.64 x 10⁻²⁰</td>
<td>0.88</td>
<td>0.79</td>
<td>0.79</td>
<td>3.12</td>
<td>7</td>
<td>0.102</td>
<td>0.129*</td>
<td>0.061*</td>
<td>9.1</td>
</tr>
<tr>
<td>Gilgel Abay</td>
<td>1</td>
<td>High</td>
<td>2014</td>
<td>60</td>
<td>35</td>
<td>9.8 x 10⁻¹⁷</td>
<td>0.58</td>
<td>0.98</td>
<td>0.96</td>
<td>2.87</td>
<td>25</td>
<td>0.407</td>
<td>0.390*</td>
<td>0.289*</td>
<td>14.6</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>Low</td>
<td>2014</td>
<td>60</td>
<td>47</td>
<td>3.03 x 10⁻²⁰</td>
<td>0.78</td>
<td>0.95</td>
<td>0.88</td>
<td>3.317</td>
<td>13</td>
<td>0.203</td>
<td>0.277*</td>
<td>0.039*</td>
<td>10.4</td>
</tr>
<tr>
<td>Mean HSR</td>
<td>2</td>
<td>Low</td>
<td>2014</td>
<td>45</td>
<td>20</td>
<td>5.13 x 10⁻⁹</td>
<td>0.43</td>
<td>0.99</td>
<td>0.91</td>
<td>3.33</td>
<td>25</td>
<td>0.545</td>
<td>0.101*</td>
<td>0.053*</td>
<td>22</td>
</tr>
<tr>
<td>Mean LSR</td>
<td>2</td>
<td>Low</td>
<td>2014</td>
<td>46</td>
<td>22.3</td>
<td>0.48</td>
<td>0.97a</td>
<td>0.84a</td>
<td>2.31b</td>
<td>25.3</td>
<td>0.52b</td>
<td>0.243</td>
<td>0.07a</td>
<td>24.3</td>
<td></td>
</tr>
</tbody>
</table>

N: number of ramets, G: number of MLGs, NR: clone size (m²), R: clonal richness. D and E: Simpson index and its equitability index. β: slope of Pareto distribution, DC: clonal dominance, Ac: aggregation index, Ee: edge effect, 

[1]multilocus genotypes reached maximum and CR: clonal subrange. All measures of clonal structure were obtained after 1000 permutations, ns, not significant and

[2]P < 0.05. Different letters indicate significant difference between sedimentation regimes collated for the three swamps (P < 0.05).

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MLGs confirmed that MLGs diverged neither as a result of scoring error nor from somatic mutation (S3 Fig). The number of individuals with distinct MLGs across sedimentation regimes ranged widely among the swamps studied (HSR: 39–53; LSR: 19–39; Table 3) and differed significantly between years (Mann–Whitney test: U = 55.5; P < 0.05). In general, clone composition differed between sedimentation regimes and demonstrated temporal dynamics within each papyrus swamp (Fig 1). Ramets belonging to each genet were distributed in a leptokurtic pattern (Fig 2). The majority of clones (particularly in LSR) were designated by more than two C. papyrus ramets, and eight clones were found in more than one sedimentation regime and swamp.

**Clonal diversity metrics**

Clonal richness ($R$), after being standardized for different sample sizes, ranged from 0.36 (Robit, 2014) to 0.67 (Gilgel Abay, 2014) and from 0.58 (Gilgel Abay, 2014) to 0.88 (Sebatamye, 2016) for LSR and HSR, respectively (Table 3). On the overall data set, mean $R$ of 2014 and 2016 were significantly different (Mann–Whitney test: $U = 45.3; P < 0.05$). We also found $R$ to be significantly different between sedimentation regimes (Mann–Whitney test: $U = 65; P < 0.05$). Simpson’s diversity ($D$) and evenness ($E$) indices were steady when compared across swamps and sampling periods (Table 3). The Pareto descriptor ($\beta$) of clonal diversity estimate showed significant spatial heterogeneity of genets distribution resulting different clone sizes in response to sedimentation (HSR: mean $\beta = 3.07$; LSR: mean $\beta = 2.31$). Similarly, Pareto distribution showed a temporal variability (mean 2014 $\beta = 2.19$ and mean 2016 $\beta = 2.75$). The lowest $\beta$ was observed in the Sebatamye swamp plot 2 ($\beta = 1.77$) in 2016.

**Clonal structure changes in space and time**

Clonal dominance ($Dc$) spanned from 0.10 (Sebatamye 2016, HSR) to 0.5 (Robit 2016, LSR) (Table 3), indicating that the majority of clones were intermingled. The difference in mean
Clone size was unevenly distributed among genotypes and the largest clone size \((NR = 38)\) was detected in Robit swamp under LSR in 2014 (Table 3). With the presumption that genotypes are randomly distributed in space, permutation analysis revealed significant aggregation of ramets within genets of each papyrus swamp plot (range \(Ac = 0.1–0.4\)). However, the mean \(Ac\) was neither significantly different between sedimentation regimes (Mann–Whitney test: \(U = 44.1; P > 0.05\); Table 3) nor between years (Mann–Whitney test: \(U = 47; P > 0.05\)). The test for the maximum spatial linear limits (clonal subrange, \(CR\)), where the chance of two identical MLGs of the same clone becomes null, varied from 9.1 m to 37 m. Plots with a high number of distinct genotypes and clonal richness were found to have a minimum clonal subrange (Table 3). There was no evidence of evidence of a significant edge effect detected within plots in either sedimentation regime or sampling year, except in Sebatamye during 2014. The edge effect did not have a sheer influence on the genotype distribution as observed.

**Association between clonal growth traits and diversity**

We tested the relationship of clonal growth traits with clonal structure and with diversity (Table 4). Ramet density corresponded to a decrease in girth diameter \((r = -0.60; P < 0.05)\), spacer length \((r = -1.00; P < 0.01\); Fig 3), culm height \((r = -0.71; P < 0.05)\), clonal dominance \((r = -0.71; P < 0.05)\) and clonal subrange \((r = -0.72; P < 0.01)\) but to an increase in clonal richness \((r = 0.65; P < 0.05)\) and aboveground biomass \((r = 0.94; P < 0.01)\). Mean clone size was significantly negatively correlated with clonal richness \((r = -0.91; P < 0.01\); Table 4) and number of distinct genotypes \((r = -0.68; P < 0.05)\) and positively with \(Dc\) \((r = 0.92; P < 0.01)\), indicating that with an increase in of few, large dominant clones, there is a decrease in clonal richness.
Discussion

In this study, combining clonal traits and genetic data, we found a trade-off between phalanx strategy and guerrilla strategy under contrasting sedimentation regimes. Higher levels of sedimentation led to an increase in ramet density, enhanced phalanx strategy and favored higher clonal diversity which can be ascribed to seedling recruitment. Other wetland plant species showed the existence of a guerrilla strategy role to escape from several factors exists [7, 11, 70], but none of these studies addressed the point that this strategy also becomes reflected at levels of clonal diversity and structure at spatiotemporal scales. We discuss each of these findings with respect to the hypotheses set and conclude by highlighting the implications for restoration and conservation activities.

In line with our first hypothesis, clonal growth traits of *C. papyrus* varied significantly between sedimentation regimes where phalanx strategy was more prevalent in HSR and guerrilla strategy in LSR. *C. papyrus* overcomes the adverse effect of HSR through production of denser ramets at the cost of a short culm height and smaller girth diameter, reflecting a consolidation strategy (phalanx) to exploit available resources created by sediment dynamics. Production of short and dense ramets with heterogeneous genets points to compensatory growth [71] in HSR. The mechanisms of compensatory growth in response to sediment and sand burial differ in plants [72, 73]. In contrast, in accordance with long spacers, longer culm height and relatively larger girth diameter, ramets under LSR showed a higher intermingling pattern which might reflect the foraging strategy [74], and corresponds to a guerrilla-type architecture. A shift from phalanx to guerrilla growth strategy has been confirmed as a principal strategy to escape from low to high sediment burial in *C. brevicuspis* [11], competition due to neighbour density in *Elymus lanceolatus* [7], sand burial in dune shrub *Hedysarum laeve* [75], sand accretion and nutrient availability in costal *Sporobolus virginicus* [76] and under conditions of different nutrient supplies in *Leymus secalinus* [70]. Our results from papyrus under natural
conditions revealed a trade-off in clonal growth strategy from LSR to HSR that was the opposite of reports based on glasshouse and pot experiments [11, 70]. Clonal plants thus may have developed diverse clonal growth traits that underpin their adaptive strategies [7, 24, 25].

The clonal structure and diversity of *C. papyrus* revealed temporal and spatial fluctuation of genets. Those sites that experienced high sedimentation dynamics were more genotypically diverse in terms of clonal richness, Pareto distribution index and number of distinct genotypes, than were sites in LSR. The significant clonal subrange, $Dc$ and the occurrence of few genets spatially restricted to either HSR or LSR reflect clonal sorting. The size and spatial distribution of genets determine demographic and sorting patterns, and have been found to be a function of trade-offs between clonal growth and sexual reproduction [77]. Like various measures of clonal growth (high ramet density, long spacer length and short culm) on *C. papyrus* genets under HSR, low $Dc$, $CR$ and clone size related to a phalanx-type architecture. In addition, a skewed clone size distribution of *C. papyrus* implies dynamics in demographic processes at different stages as a response to sedimentation, which has also been reported for populations of the clonal seagrass, *Cymodocea nodosa* [69]. The latter only accounted for a difference at ramet level and genet level analysis associated with the reproductive modes, and did not include factors that could drive the demographic patterns in each population: yet historical contingencies during colonization are critical to understand factors driving genet distribution. A high $Dc$ and low contemporary ramet density in LSR might also be attributed to the influence of asymmetrical competition between clones of different sizes, which amplifies loss of genets and decrease in genotypic diversity [78]. Under a relatively stable sediment regime (LSR), the founder effect of a few dominant clones may have shaped clonal structure through selection of superior clones and clonal integration [28, 79, 80]. According to the "general-purpose hypothesis", persistence and resilience are common features of large clones [79, 81]. However, overall, our results on papyrus do not support this hypothesis, because of the significant genet turnover and high clonal richness observed in sites with HSR.

Despite the fact that our data are limited to two sampling periods, the change in clonal composition, richness and growth strategies suggests the potential of an “alle effect”. Such a phenomenon has also been proposed for stand establishment of the clonal macrophyte, *Phragmites australis* [82]. The capacity of a species to colonize, establish and maintain its population in a dynamic environment is a function of dispersal potential. Individuals with identical MLGs distributed across the swamps studied have supported long-distance clonal dispersal within the same lake, as reported for spatially isolated *C. papyrus* populations [83], and selective advantages that could facilitate the clonal spread [84]. While the latter is more prominent within the same swamp area it may be less likely to occur in Lake Tana across such geographically distant swamps (68 km apart).

Papyrus populations revealed a wide range of clonal richness, from low (with very few genets) to high (with many genets) levels across LSR to HSR, respectively. This discrepancy in the number of genets explicitly highlights how the relative contribution of sexual and asexual reproductive modes dictates distribution and recruitment patterns of genets [17] in response to sedimentation. Initial seedling recruitment (ISR) and repeated seedling recruitment (RSR) are the two contrasting dynamic life history traits governing the growth of clonal plants [18]. Our results are in full agreement with a previous study that has provided evidence for seedling recruitment differences in *C. papyrus* populations along drawdown and inundated regimes at Lake Naivasha, Kenya [85]. High genet turnover ensuring an increased number of new genets in plots under HSR did reflect the central role of sexual reproduction and a RSR strategy. Despite high ramet density in these plots, sediment dynamics open windows for seedling recruitment and induce chaotic distribution of genets. Microsites created during these sediment dynamics influence the earliest stage of seedling emergence [86]. In contrast, the
establishment of a few but dominant clones, coupled with maximum CR and longer spacer length in LSR revealed that an ISR model of *C. papyrus* was in place. In such a relatively stable regime, the presence of a cohort of a few large genets would show differential spatial spread (longer spacer length) and intragenet competition at the early stage of colonization [19]. Generally, our results showed that high sedimentation shifts the balance between seedling recruitment strategies (from ISR to RSR) and between the reproductive modes (from asexual to sexual). This, allows us a glimpse of the ultimate effect of environmental factors in driving microevolutionary forces [16] by influencing seedling establishment, and illuminates the comparative significance of reproduction modes [87, 88].

The resilience of wetlands and their keystone clonal plants have been reported as challenging, due to the spatial dynamics of genets that can be decoupled to allow contraction and expansion in response to biotic and abiotic factors [89]. However, our findings have twofold implications for resilience through habitat protection and restoration of *C. papyrus* populations. Severe land degradation in the upper catchments of the Lake Tana basin, for instance, is a far-reaching problem that led to lake over-sedimentation [27, 44]. However, with the premise that papyrus also controls sediment loads and provides socio-ecological services [32, 34, 38, 42], revegetation of sedimentation-prone sites using this species would lessen over-sedimentation and promote restoration in a broader context. To do this, selection of individuals of genets (genotypes) that withstand sediment dynamics and temporally persist (such as those noted in HSR), and sites potentially suitable for creating stepping stones between threatened populations [90], have to be the priority for mitigation of over-sedimentation along a lake margin. For sustenance of the studied papyrus swamps, preventive measures have to counteract anthropogenic pressures encompassing sand-mining and drainage of swamps for agricultural purposes [27]: these have negative impacts on the substrate, affecting for clonal growth and the recruitment of new genets. Our findings suggest that wetland conservation and restoration of keystone clonal plants should consider the spatiotemporal dynamics of ramets and genets that shape the level of genotypic diversity and determine the shift in clonal growth strategies for both short-term responses and long-term evolutionary potential.

In summary, we observed that the paper reed (*C. papyrus*) was able to adapt to sediment regimes by shifting clonal growth strategy from guerrilla (in LSR) to phalanx (in HSR). The threshold level of sedimentation that limits seedling recruitment and clonal dynamics has not been established for *C. papyrus*, but our spatial and temporal observations highlight that HSR enhances clonal richness owing to RSR and genets turnover. HSR creates a heterogeneous microenvironment that can be occupied by coexisting multiple genotypes, and this could have implications for population dynamics and resilience in the long term. Finally, restoration and conservation of papyrus wetlands from over-sedimentation, using clonal plants, should also incorporate information about spatiotemporal dynamics in clonal growth strategy, and in clonal diversity and structure.

**Supporting information**

**S1 Fig. Clonal growth characteristics; ramet density (i) and spacer length (ii) of *Cyperus papyrus* under HSR and LSR observed for 2014 and 2016.** The values (mean ± SE) recorded for 18 quadrates within each plot collated over the three swamps. Different letters indicate significant difference between years within a sedimentation regime and ** significant difference between sedimentation regime regardless of years (P < 0.05).**

(TIF)
S2 Fig. Maximum number of distinct genotypes that adequately reached the number of markers used, thereby allowing the accurate estimation of clonal diversity of *C. papyrus*.

(TIF)

S3 Fig. Frequency distribution of the pairwise number of alleles differences between MLGs for sample of *C. papyrus*.

(TIF)

S1 Table. Genotypic data of *C. papyrus* from three papyrus swamps in Lake Tana under two sedimentation regimes over two years.

(XLS)

S1 Protocol. Table: Multiplex PCR protocol for *C. papyrus*.

(XLS)

S1 Text. Copyright permission granted to use the figure as shapefile.

(PDF)

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**Author Contributions**

**Conceptualization:** Addisie Geremew, Ludwig Triest.

**Data curation:** Addisie Geremew, Iris Stiers, Alemayehu Kefalew.

**Formal analysis:** Addisie Geremew, Iris Stiers, Ludwig Triest.

**Funding acquisition:** Addisie Geremew, Ludwig Triest.

**Investigation:** Addisie Geremew, Tim Sierens, Alemayehu Kefalew.

**Methodology:** Addisie Geremew, Iris Stiers, Tim Sierens, Alemayehu Kefalew, Ludwig Triest.

**Project administration:** Addisie Geremew, Ludwig Triest.

**Resources:** Tim Sierens, Alemayehu Kefalew, Ludwig Triest.

**Software:** Addisie Geremew, Iris Stiers, Tim Sierens.

**Supervision:** Ludwig Triest.

**Validation:** Addisie Geremew, Iris Stiers, Ludwig Triest.

**Visualization:** Addisie Geremew, Iris Stiers.

**Writing – original draft:** Addisie Geremew.

**Writing – review & editing:** Addisie Geremew, Iris Stiers, Tim Sierens, Alemayehu Kefalew, Ludwig Triest.

**References**


