Lacustrine mollusc radiations in the Lake Malawi Basin: experiments in a natural laboratory for evolution

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Abstract. In terminal Pliocene–early Pleistocene times, part of the Malawi Basin was occupied by paleo-lake Chiwondo. Molluscan biostratigraphy situates this freshwater lake either in the East African wet phase between 2.7–2.4 Ma or that of 2.0–1.8 Ma. In-lake divergent evolution remained restricted to a few molluscan taxa and was very modest. The lacustrine Chiwondo fauna went extinct at the beginning of the Pleistocene. The modern Lake Malawi malacofauna is depauperate and descends from ubiquitous southeast African taxa and some Malawi basin endemics that invaded the present lake after the Late Pleistocene mega-droughts. The Pleistocene aridity crises caused dramatic changes, affecting the malacofauna of all East African lakes. All lacustrine endemic faunas that had evolved in the Pliocene rift lakes, such as paleo-lake Chiwondo, became extinct. In Lake Tanganyika, the freshwater ecosystem did not crash as in other lakes, but the environmental changes were sufficiently important to trigger a vast radiation. All African endemic lacustrine molluscan clades that are the result of in-lake divergence are hence geologically young, including the vast Lavigeria clade in Lake Tanganyika (ca. 43 species).

1 Introduction

The hypothesis that the large African lakes are “natural laboratories of evolution” and that their diversified molluscan fauna are prime examples of intense and ancient processes of in-lake evolution (Michel et al., 1991) has become generally accepted. The recent and fossil malacofauna of Lake Malawi does not corroborate this theory. In fact, all actual molecular and paleontological investigations provide evidence pointing to the contrary, namely that in Lake Malawi, as well as in other past and present rift lakes, molluscan in-lake divergence is surprisingly modest and young geologically speaking. This view is partly applicable even to the unique malacofauna of Lake Tanganyika. The present study links the late origin of the malacofaunas of the present rift lakes to the increasing climate destabilisation in East Africa since the late Pliocene.

2 The fossil record

This study is based on the collections of terminal Pliocene–early Pleistocene molluscs collected in the Chiwondo region (NW margin of Lake Malawi), respectively, in the 1960s during the Desmond Clark Palaeo-Antropological Investigation and in 1980–1990s during the Hominid Corridor Research Project led by Timothy Bromage and Friedemann Schrenk. Albrecht Gorthner, the HCRP malacologist, also sampled the Holocene assemblages near the Shire River outlet and this preliminary investigation was recently continued (Van Boeckelaer, 2004; Van Boeckelaer et al., 2012). All relevant fossil material collected in the Malawi Basin is provisionally stored at the Paleontological Research Unit, Ghent University, awaiting formal taxonomic description. The taxonomy of the Desmond Clark collection was studied by Gautier (A. Gautier, unpublished data, 1968) and taxonomy and biostratigraphy of the HCRP-collection by A. Gorthner (unpublished data, 1995). Copies of both manuscripts are kept at the Paleontological Research Unit Ghent as part of the collection.

The present paper is essentially a critical review of the fossil material cited above and the published literature on the modern and fossil Lake Malawi malacofauna in the light of our vastly improved knowledge on phylogeny and
paleontology of the molluscs of the African Great Lakes and of the paleoimnological evolution of these lake basins. The published literature in which provisional species lists of the Chiwondo fossil molluscs or part of them are provided is quite extensive and widely scattered. It includes Pain (1966), Gautier (1966), Gautier (1970), Van Damme (1984, 1988), Van Damme and Pickford (1999, 2003), Goethner (1994), Goethner et al. (1992), Schrenk et al. (1995), Schultheiß et al. (2009) and Van Bocxlaer (2010).

3 The evolution of taxonomic concepts about the modern Malawi malacofauna: from speciose 19th Century Lake Nyasa to species-poor 21st Century Lake Malawi

The spectacular thalassoid or marine-like fauna of Lake Tanganyika (area: 32 900 km²) greatly puzzled 19th Century scientists and led to heated debate as to their origin. The comparatively unspectacular molluscs of Lake Nyasa, as L. Malawi formerly was known, saddled them with the subsidiary question why in this equally vast lake (area: 29 600 km²) no such a diversified thalassoid fauna was found, for according to Bourguignat (1889) only the Nyasan thiarids did possess thalassoid characters. This author, recognizing about 40 thiarid species, divided them in five genera: Melania (= Melanoides) represented only by the ubiquitous M. tuberculata and the rest belonging to the endemic genera Nyassa, Nyassa, Micronyassa and Nyassomelania (Fig. 1). He believed that the relatively low species richness was due to insufficient sampling in the at the time virtually unexplored lake.

In the middle of the 20th Century, during what Michel et al. (2003) call “the dawn of Mayrian optimism for the practicality of a single ‘biological’ species concept”, the number of molluscan species in all African lakes was taxonomically decimated. In Lake Malawi none of the endemic thiarid genera survived this taxonomic lumping event and only 16 gastropods are presently considered to live in the lake sensu stricto of which eight are endemic Melanoides (Brown, 1994; Darwall et al., 2005), but the number of the Melanoides endemics is still unresolved (see Table 1). Eldblom and Kristensen (2003) retain three endemics only in the last revision based on morphology. While molecular biologists speak of the “Melanoides polymorpha-complex”, considered to be a poly- or paraphyletic group of clones (Genner et al., 2004, 2007b; Von Gersdorff Sørensen et al., 2005). This led Michel et al. (2008) to raise a question about their equivalence to “standard” species, i.e., as defined according to the Biological Species Concept (BSC).

Questions on the morphological distinctiveness and the number of the endemic Bellamya (Viviparidae) and Lanistes (Ampullariidae) have not yet been fully resolved either. Recent molecular investigations of these two other prosobranch “species flocks”, considered to have diverged within the lake, revealed that both groups consist not exclusively of in-lake endemics, as was formerly assumed, but of in-lake endemics sensu stricto plus one or two paludal/fluvial species endemic to the whole of the Malawi Basin, i.e., basin endemics (Sengupta et al., 2009; Schultheiß et al., 2009, 2011). Both these Malawi species groups are monophyletic, young and the speciation processes still likely are going on in the viviparid flock (Schultheiß et al., 2011). In addition, the possibility that the clonal Melanoides lineages endemic to Lake Malawi are not the result of an in-lake radiation either, but that they are allopatric paleo-endemics that colonized Lake Malawi at different times, is considered a possible scenario in view of their para-/polyphyly (Genner et al., 2007b).

Dudley’s (2000) remark that “Malawi gastropod classification is in a continuing state of revision and that it will be some time before a system comes to be generally agreed for the groups of higher rank” is still painfully actual. Certain is that the recent genetic research on Lake Malawi molluscs does not lend support to the ingrained concept of “spectacular” species diversification/radiation in this supposedly long-lived “natural laboratory of evolution”. The largest Malawi group, that of Melanoides, can only partially result from an in-lake radiation and the two other “clades” are small and genetically little diversified.

Diversification is not spectacular and hence niche partitioning and occupation are neither. Most Malawi species are restricted to the shallow littoral zone with sandy substrate (above 20 m) and only a few are found in the deep sublittoral between 40 and 80 m. But at such depths their occurrence is sporadical and the only two species, Lanistes nasutus and Bellamya ecclesi (Fig. 2), possessing morphological adaptations for life at greater depths, are represented by a few rare and highly localized populations (Brown, 1994). The specific shells adaptations in both species consist of the persistence of essentially neotenous traits, namely a thin, rapidly
Growing shell, a strongly inflated body whorl and a mouth aperture that is very large compared to the total length and width of the animal, either being strongly elongated as in *L. nasutus* or wide as in *B. ecclesi*. Such morphological features increase buoyancy and permit the extension of a large foot, adaptations for life in or on fluid sediment. No special anatomical features are mentioned in literature except for strongly elongated tentacles in *L. nasutus* (Mandahl-Barth, 1972; Berthold, 1990).

### 4 The paleontological data reviewed: the end of Lake Malawi as an ancient lake

The paleolimnological, pre-Holocene evidence in the Malawi basin is fragmentary and localized. Pliocene–Pleistocene deposits, described as the Chiwondo and the Chitimwe Beds, are exposed at the NW fringe of Lake Malawi near its satellite lake Chiwondo (Fig. 3). For detailed stratigraphic information we refer to Sandrock et al. (2007) and Kullmer (2008). Age estimates are based on suid biochronostratigraphy (=chronostratigraphy based on teeth of fossil pigs (Suidae)). Since no mammal fossils are found at the mollusc bearing sites, their age was based, as well as possible, on stratigraphic correlations with nearby exposures yielding mammals.

According to the authors cited, five limnological stages can be discerned in the Chiwondo region ranging from terminal Miocene to Late Pleistocene times:

- **Chiwondo Beds, Unit 1** (time range: > 5 to ca. 4.0 Ma). Exclusively fluvial deposits, no molluscs.

- **Chiwondo beds, Unit 2** (time range: ≥ 4 to ca. 3.75 Ma). Lacustrine limestone and silt- to sandstone deposits formed in littoral and margin environments with rare gastropod fossils (*Bellamya*). At the top of the unit the facies changes to littoral marlstones with abundant gastropods and rare bivalves. The molluscan assemblages are found in consolidated shell beds cropping out as benches with a thickness up to several meters and extending over several hundred meters (Schrenk et al., 1995). Preservation in the coarse sandstone is poor with inner casts and partly dissolved outer casts. In the rare marl- and limestones fossilization is slightly better, consisting of recrystallized calcitic shells.

- **Chiwondo Beds, Unit 3** (time range: ca. 3.75 to 2.0 Ma for Subunit 3A and ca. 2.0 to ca. 1.5 Ma for Subunit 3b on the basis of suid biochronostratigraphy). The overlying unit, separated from the earlier deposits by an unconformity, is characterized by the return of deltaic/fluvial conditions. Pronounced lake regressions took place between 2.3–2.0 Ma and 1.6–1.5 Ma. No molluscan assemblages.

- **Chiwondo Beds, Unit 4** (time range: ca. 1.5 Ma to > 20 ka). Deposits of eolian sands indicate periods of marked aridity but a brief lacustrine transgressive phase is recorded (no molluscs).

- **Chitimwe Beds** (time range: ≤ 20 ka). Alluvial fan deposits indicate a phase of lacustrine regression. Holocene lacustrine deposits are not present in the NW Lake Malawi region.

The known Chiwondo malacofauna comprises species belonging to the genera *Lanistes* (2–3 species; Ampullariidae), *Bellamya* (4 sp.; Viviparidae), *Gabbiella* (2–3 sp.; Bithyniidae), *Cleopatra* (1 sp.; Paludomidae), *Melanoideas* (2 sp.; Thiaridae), *Bulinus* (1 sp.; Planorbidae), *Coelatura* (2 sp.; Unionidae), *Pseudobovaria* (1 sp.; Unionidae), *Chambardia* (1 sp.; Iridinidae), *Etheria* (1 sp.; Etheriidae) and *Corbicula* (1 sp.; Cyrenidae) (see Table 1). Many assemblages are
species poor with mainly Bellamy, Gabbiiella, Melanoides or monospecific with only Bellamya. All appear to have been formed in the littoral storm wave zone and the higher part of the littoral.

Most Chiwondo species, i.e., those belonging to the genera Lanistes, Gabbiiella, Melanoides, Bulinus, Coelatura, Pseudobovaria, Chambardia, Etheria and Corbicula are morphologically similar to species that are or were (i.e., Pseudobovaria) widespread in East Africa or are endemic to the Malawi Basin (e.g., Chambardia nyassaensis (Graf and Cummings, 2007) and do not show lacustrine adaptations.

A marked diversification can be observed only in the genus Bellamya (Fig. 2.), which apart from an unornamented morph (“Bellamya capillata” auctores) is represented by (1) a form with sloping, flattened whorls and a basal carina, described by A. Gautier (unpublished data, 1968) and cited
in Van Damme and Pickford (1999) as Bellamya cf. pagodiformis for the likeness with the modern Lake Mweru species, (2) a carinated form, B. cf. trochlearis, resembling the modern Lake Victoria endemic and (3) B. cf. robertsoni, which is morphologically quite similar to the present-day Lake Malawi B. robertsoni but smaller. The unornamented B capillata is dominant, all others are relatively rare. The morphological differences suggest a para- or polyphyletic origin of the Chiwondo Bellamyla-group. That these Chiwondo lacustrine endemics are ancestral to the present ones is highly unlikely, considering the evidence for intermittent aridity and salinity peaks during the Pleistocene (see paleoenvironmental reconstruction of Pleistocene events). As to a continuity in Melanoides, the morphological divergence and dwarfism observed in the modern Malawi representatives is not found in the Chiwondo assemblages, where the taxon is represented by two large sized forms with no distinct correlations to the modern clade (see also further).

Generally the Chiwondo littoral malacofauna is surprisingly similar in diversity and composition to the early-middle Holocene fauna (Van Boxclaer, 2004), at least in comparison to the marked differences between Plioence and Holocene-modern faunas in paleo-lakes of the Turkana and the Albertine basins. It is safe to conclude that during late Pliocene times the Chiwondo lake fauna did already consist for an important part of the same ubiquitous and basin endemic taxa occurring in the present lake and that the composition of the late Pliocene faunal community was in many aspects already “modern”. The only “ancient” taxon present, be it rare, in the Chiwondo assemblages is the unionid Pseudobovaria. In the two other basins mentioned, Pliocene fauna elements remain dominant until the period ca. 1.8 Ma to ca. 1.2 Ma (see further).

That the Chiwondo malacofauna dates from the early Pliocene, that is from 4–3.8 Ma as proposed by Sandrock et al. (2007) and Kullmer (2008), is difficult to accept, not primarily because that would infer that the Malawi fauna had reached its modern aspect millions of years earlier than those of other East African basins, but because of the presence of the Asiatic bivalve Corbicula. The presence of fossils of this Asian clam provides a terminus ante quem date of 2.6–2.5 Ma for African assemblages. Well-calibrated evidence comes from the Turkana Basin where Corbicula appears for the first time in deposits of the short-lived paleo-lake Lokeride closely overlying the Burgi Tuff, dated to 2.68 ± 0.06 Ma (Feibel et al., 1989; McDougall and Brown, 2008). Fossil Corbicula from the Kada Me’e Tuff Complex in the Hadar Formation are dated at ca. 3.4 Ma, but this in an estimate obtained by interpolation of six $^{40}$Ar/$^{39}$Ar analytical data and considered as unreliable (Campisano, 2007; Campisano and Feibel, 2008).

The terminus ante quem date based on the molluscs is 1.5 to 1.2 Ma, i.e., the last records of the extinct unionid Pseudobovaria (Van Damme and Pickford, 2010). The age of all molluscan Chiwondo assemblages should, considering the absence of any marked morphologically differences among the assemblages, be considered as being broadly similar, hence deposited either around 2.5 Ma or around 1.9 Ma. In East Africa the 2.7–2.4 Ma interval appears to be a major lake period suggested by the moisture history of the Ethiopian, Kenyan and Tanzanian rift basins (Trauth et al., 2005, 2007, 2010; Tiercelin et al., 2010). Important incursions of invasive species (e.g., Corbicula) synchronously in the Turkana Basin (Van Boxclaer et al., 2008) and the Albertine Basin (Van Damme and Pickford, 2003, 2010) indicate a significant increase of hydrological connections among different basins during the time segment. Paleolake Chiwondo may hence have existed synchronously with paleolake Lokeride (ca. 2.5 Ma) of the Turkana basin and with the earlier stages of paleo-lake Kaiso and paleo-lake Lusso (ca. 2.5 Ma) of the Albertine basin. An alternative possibility is that it existed during the same period as paleo-lake Lorenyang (ca. 2.0–1.8 Ma) in the Turkana Basin and the terminal stages of both aforementioned lakes in the Albertine Basin (ca. 2.0–1.8 Ma) (Van Damme and Van Boxclaer, 2009; Van Damme and Pickford, 2010), present during an equally wide spread East African wet phase (Trauth et al., idem). In the latter period the Pliocene fauna elements in the two basins were disappearing also.

As in other parts of East Africa, periods of increasingly aridification and climatic instability are recorded in the Malawi Basin throughout the whole Pleistocene period with extremely low lake levels between 2.3–2.0 Ma in late Pliocene times and between 1.6–1.5 Ma during Lower Pleistocene times, in Unit 3b of the Chiwondo Beds. In the overlying Unit 4, eolian sands equally indicate phases of hyper-aridity during the rest of the early and middle Pleistocene (Sandrock et al., 2007). The intermittent occurrence of discrete but extreme periods of aridity during the Late Pleistocene and the dramatic degradation of the freshwater lacustrine ecosystem has been reconstructed in detail via the multidisciplinary study of Malawi lake cores (Cohen et al., 2007). Hyper-aridity is noted in the period between 135 to 90 ka, when the strongly constricted lake was shallow, holomictic and saline, surrounded by semi-desert.

From Pleistocene times only a single, brief high lake level stand of unknown age, younger than 1.5 Ma, and without fossils is recorded (Sandrock et al., 2007). This event may possibly fall within the late in the early Pleistocene period of major global climatic transition, situated at 1.1 to 0.9 Ma by Trauth et al. (2007), or it results from local climatic events. The absence of macrofossils may be due to unfavourable conditions for fossilisation, but more likely indicates the absence of macroscopic life (hyper-salinity/hyper-alkalinity). For the whole of the Pleistocene, information about molluscan life in the Malawi Basin is hence missing. The lacustrine ecosystem during that epoch must have fluctuated between a Lake Baringo-Lake Naivasha type of lake, i.e., with a conductivity in excess of 4000–4500 µS cm$^{-1}$ without freshwater prosobranchs or bivalves.
The chances that populations of oligohaline taxa, such as the gastropod Bellamyia (Viviparidae) and Cleopatra (Pulnodomidae) and equally sensitive bivalves such as Coelatura and Nyassunio (Unionidae), Mutela and Pleiodon (Iriniidae) and Etheria (Etheriidae), could survive these intermittent aridity/hyper-aridity peaks in the restricted and endorheic Malawi Basin, i.e., prior to the connection with the Zambezi Basin via the River Shire, were nil. These groups must have re-invaded the Malawi Basin, when environmental conditions improved in the Late Pleistocene. The morphological likeness between modern Bellamyia robertsoni and terminal Pliocene–early Pleistocene B. cf. robertsoni (Fig. 2) does therefore not imply direct parentage, in particular since the modern robertsoni morph is not yet clearly discernible in the late Holocene Malawi deposits (Van Bocxlaer, 2004; Van Bocxlaer and Van Damme, 2009).

Taxa such as Melanoides (Thiaridae) and Gabbiella (Bithyniidae) tolerate relatively high salinity concentrations and taxa such as Lanistes (Ampullariidae) and Chambardia (Iriniidae) are able to aestivate during extended dry periods (Van Damme, 1984). Populations of these taxa may have survived the Pleistocene salinity crises and modern representatives in the basin or in the lake such as Lanistes ellipticus, Gabbiella stanleyi and Chambardia nyassaensis may derive from basin endemics already present during Chiwondo times. However, the molecular evidence concerning the Malawi Lanistes, indicates that the formation of this group is young (Middle Pleistocene?) (Schultheiß et al., 2009).

For the modern representatives of the genus Melanoides, the case is more complex. They are not yet present in Chiwondo times and may be dwarfed morphs that originated during Pleistocene times in the saline lakes, but that these morphs are descended from populations already present in the basin during Chiwondo times (Genner et al., 2007b) cannot be ascertained. The dominant fossil Chiwondo Melanoides was identified as M. cf. nodicincta, a modern Lake Malawi endemic, by A. Gautier (unpublished data, 1968), cited in Van Damme (1984) and Van Damme and Pickford (2003), but this identification is incorrect. The Chiwondo shells are too poorly preserved to provide a specific attribution. Presently it seems best to consider them to belong to the M. polymorpha–M. mweruensis–M. anomala species group, occurring in the south-eastern African region (Van Bocxlaer, 2010). Using these Chiwondo fossils to calibrate the molecular clock in calculating the age of the modern M. polymorpha-group should be avoided (Genner et al., 2007b).

5 Is the Pleistocene climatic destabilization responsible for major extinctions and major evolutionary radiations in the molluscs of the African rift lakes?

The reconstruction of the malacological history of the Malawi basin, though fragmentary, shows that as in the two other East African basins with a fossil record, Lake Turkana and Lake Albert basin, a diversified lacustrine fauna existed in the basin during late Pliocene times. It contained lacustrine endemics and definitely possessed already a distinct south-eastern African character, clearly different from that of the other basins mentioned. The age of this paleo-lake Chiwondo fauna is uncertain. It certainly is not older than 2.7–2.6 Ma but it could be younger, i.e., dating from the East African wet phase at 2.0–1.9 Ma.

After that period, possibly from ca. 1.8 Ma and definitely after 1.5 Ma, the old Pliocene malacoфаuna in the Turkana and Albertine basins shows signs of a severe deterioration. In the Albertine Basin the Miocene–Pliocene Congolian faunal elements are replaced by an impoverished eurytopic fauna of the “Nilotic” type around 1.8 Ma (Van Damme and Pickford, 1999, 2003, 2010). In the Turkana basin virtually all old Pliocene basin elements are definitely gone after ca. 1.3 to 1.2 Ma, i.e., after the brief lake interval around ca. 1.4 Ma at the base of Member L, Shungura Formation (Van Bocxlaer et al., 2008).

The paleo-environmental data for the Malawi Basin we possess indicate the instauration of hyper-arid phases possibly already from about 2.3 Ma, if the mammalian chronostratigraphy of Sandrock et al. (2007) is followed. It can be assumed that in the Malawi Basin all molluscan lake endemics must have become extinct prior to 1.8 Ma and that most of the basin endemics, possibly except those with adaptations to survive extended periods of aridity, e.g., Chambardia nyassaensis, or significant increases in salinity, e.g., Gabbiella stanleyi, also were gone after ca. 1.3 to 1.2 Ma.

In all larger East African basins, where according to the fossil evidence an endemic lake fauna developed during late Miocene–Pliocene times, all Pliocene lacustrine endemics and many basin endemics appear to have become extinct during the hyper-arid early Pleistocene times. In all smaller basins such as those in the Gregory and Ethiopian rift, fossil mollusc assemblages from Pliocene freshwater lakes are also known, but intense level fluctuations, endorheism and volcanic activity only permitted short lived incursions of eurytopic species and no evolution (Van Damme, 1984).

In their study on the evolution of the modern endemic Lanistes clade of Lake Malawi, Schultheiß et al. (2009) extend the debate concerning possible evolutionary patterns in molluscs to all large African lakes. They discern three types of lacustrine ecosystems based on the criterion that tempo and mode of molluscan evolution are primarily determined by the degree of ecosystem stability on a geological timescale. At one end they place “seemingly” stable systems but which are, viewed on a geological timescale, instable and
### Table 1. Freshwater molluscs from terminal Pliocene–early Pleistocene Paleo-Lake Chiwondo and modern Lake Malawi. Taxonomy and range of modern species according to Brown (1994) and Graf and Cummings (2007) except when mentioned otherwise.

| Taxon | Paleo-Lake Chiwondo | Modern Lake Malawi | Remarks |
|-------|---------------------|--------------------|---------|
| **GASTROPODA** | | | |
| **Viviparidae** | | | |
| *Bellamya capillata* | x | | Wide spread SE-African species, but modern Malawi Basin populations differ genetically and the name *Bellamya simonsi* Bgt, a basin endemic, has preference |
| *Bellamya jeffreysi* | – | x | Modern lake endemic, absent in early Holocene L. Malawi deposits |
| *Bellamya robertsoni* | x | – | Modern lake endemic, absent in early Holocene L. Malawi deposits |
| *Bellamya cf. robertsoni* | x | – | Lake endemic. Morphological convergence with modern species from L. Malawi |
| *Bellamya ecclesi* | – | x | Modern lake endemic known only from deeper water in southern L. Malawi |
| *Bellamya cf. pagodiformis* | x | – | Lake endemic. Morphological convergence with modern species from L. Mweru |
| *Bellamya cf. trochlearis* | x | – | Lake endemic. Morphological convergence with modern species from L. Victoria |
| **Ampullariidae** | | | |
| *Lanistes ovum* Bgt Brown | x | x | Widespread in Africa but modern Malawi Basin populations differ genetically and the name *Lanistes ingens* Ancey, a basin endemic, hence has preference |
| *Lanistes ellipticus* | x | x | Basin endemic; not found in the lake proper |
| *Lanistes solidus* | – | x | Lake Malawi endemic (?) nearly indistinct from *L. ellipticus* |
| *Lanistes cf. solidus* | ? | – | Only a few badly preserved specimens possibly belong to this species |
| *Lanistes nyassanus* | – | x | Modern Lake Malawi endemic, not found in early Holocene deposits |
| *Lanistes nasutus* | – | x | Modern Lake Malawi deep water species, only known from a dozen specimens |
| **Bithyniidae** | | | |
| *Gabbiella stanley* | – | x | Mainly in vegetation in Lake Malawi shallows, probably basin endemic |
| *Gabbiella cf. stanley* | x | – | Morphological identical to modern species, probably basin endemic |
| *Gabbiella sp.* | x | – | Exceptionally large *Gabbiella* species, related to the above. Lake endemic? |
| **?Valvatidae** | | | |
| *?Valvata sp.* | x | – | Costulate ovate species, identified by Van Damme (1976) as *Valvata*, but more likely belonging to the Assiminaeidae |
| **Paludomidae** | | | |
| *Cleopatra sp.* | x | – | Basin endemic with modest carinae, probably belonging to the *Cl. smithi*/ *Cl. mweruensis* group from Zambia, Genus presently absent in the Malawi basin |
| **Thiaridae** | | | |
| *Melanoides tuberculata* | – | x | Widespread Oriental species |
| *Melanoides nodicincta* | – | x | Clone, endemic to modern L. Malawi |
| *Melanoides pergracilis* | – | x | Clone, endemic to modern L. Malawi |
| *Melanoides cf. pergracilis* | x | – | Very slender form, ca. twice the size of modern *M. pergracilis* |
| *Melanoides pupiformis* | – | x | Clonal dwarfed form, endemic to L. Malawi |
| *Melanoides turritispira* | – | x | Clonal dwarfed form, endemic to L. Malawi |
| *Melanoides polymorpha* | – | x | Modern Lake Malawi endemic. Extremely polymorphic species |
| *Melanoides cf. polymorpha* | x | – | Erroneously mentioned as *M. cf. nodicincta* by Van Damme and Pickford (2003) |
| *Melanoides nyassana* | – | x | Clonal dwarfed form, endemic to L. Malawi |
| *Melanoides truncatelliformis* | – | x | Clonal dwarfed form, endemic to L. Malawi |
| *Melanoides magnifica* | – | x | Clonal dwarfed form, endemic to L. Malawi |
| *Melanoides simonsi* | – | x | Considered distinctive from *M. nodicincta* by Eldblom and Kristensen (2003) |
| *Melanoides virgulata* | – | x | Invasive Oriental species in modern L. Malawi (Eldblom and Kristensen, 2003) |
| **Lymnaeidae** | | | |
| *Lymnaea natalensis* | – | x | Widespread pulmonate |
Table 1. Continued.

| Taxon                          | Paleo-Lake Chiwondo | Modern Lake Malawi | Remarks                                                                 |
|--------------------------------|---------------------|--------------------|-------------------------------------------------------------------------|
| Planorbidae                    |                     |                    |                                                                         |
| *Biomphalaria pfeiffer*        | –                   | x                  | Ubiquistic. S. America genus that invaded Africa probably in Pleistocene times |
| *Ceratophallus natalensis*     | –                   | x                  | Widespread                                                              |
| *Gyraulus costulatus*          | –                   | x                  | Widespread                                                              |
| *Bulinus globosus*             | –                   | x                  | Widespread                                                              |
| *Bulinus cf. globosus*         | x                   | –                  | Species-level identification requires anatomical characters            |
| *Bulinus nyassanus*            | –                   | x                  | Modern Lake Malawi endemic, also present in early Holocene deposits      |
| *Bulinus succinoides*          | –                   | x                  | Modern Lake Malawi endemic                                              |
| *Bulinus forskalii*            | –                   | x                  | Widespread                                                              |

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| Unionidae                      |                     |                    |                                                                         |
| *Nyassuno nyassaensis*         | –                   | x                  | Lake endemic very close to *C. hypsiprymna*. Present in Holocene deposits |
| *Coelatura hypsiprymna*        | –                   | x                  | Basin endemic, also in modern L. Malombe and in Holocene L. Malawi deposits |
| *Coelatura mossambicensis*     | –                   | x                  | Widespread south-eastern African species                               |
| *Coelatura cf. mossambicensis* | x                   | –                  | Probably identical to the modern species                               |
| *Coelatura sp.*               | x                   | –                  | Very large *Coelatura*, resembling modern *C. briarti*, widespread in the Congo Basin |
| *Pseudohovaria mwayana*        | x                   | –                  | Extinct genus en species represented by the subsp. *tuberculata*, equally found in the L. Turkana and Albert Basin, where it is restricted to early Pleistocene strata. |

| Iridinidae                     |                     |                    |                                                                         |
| *Aspatharia subreniformis*     | –                   | x                  | Widespread                                                              |
| *Chambardia nyassaensis*       | x                   | x                  | Ranging from L. Rukwa (Tanganyikan basin) to L. Malombe (Malawi Basin) |
| *Chambardia wahlbergi*         | –                   | x                  | Widespread                                                              |
| *Mutela alata*                | –                   | x                  | In L. Malawi, Shire R., L. Malombe and in Malagarasi R. (Tanganyikan Basin) |

| Etheridae                      |                     |                    |                                                                         |
| *Etheria elliptica*            | x                   | –                  | The river oyster *Etheria* does presently not occur anymore in the Malawi Basin |

| Cyrenidae                      |                     |                    |                                                                         |
| *Corbicula fluminalis*         | x                   | x                  | Widespread Oriental species                                           |

| Sphaeriidae                    |                     |                    |                                                                         |
| *Pisidium pirothi*             | –                   | x                  | Widespread                                                              |
| *Pisidium reticulatum*         | –                   | x                  | Widespread                                                              |

intermittently cease to exist as large standing freshwater bodies. As an example, they cite Lake Victoria that dried out in Late Pleistocene times. In such type of lake radiation events may be triggered during renewed freshwater phases but there is no continuity in the molluscan lineages involved. On the other end they place the Lake Tanganyika ecosystem characterized by long-term stability of the abiotic setting. The authors claim that the endemic radiations within lakes like Lake Tanganyika or Lake Baikal “have regularly proved to be remarkably old”. The lacustrine ecosystem in the Malawi Basin is considered by them as an intermediate.

We fail to see why the lacustrine malacoфаunas in the Lake Malawi basin should be given this special status. The data advanced in this paper indicate that regardless of these depth and size, all African freshwater systems, that in the Lake Malawi Basin included, crashed during the Pleistocene and that their endemic intra-lacustrine faunas went extinct. The only exception is Lake Tanganyika (Fig. 3). However, while part of the Pliocene malacoфаuna survived in that lake, there are no indications that the intra-lacustrine molluscan radiations in it are remarkably old. Most Tanganyikan mollusc taxa are paleo-endemics, not neo-endemics, as shown by paleontological (Van Damme and Pickford, 2003, 2010), molecular (Wilson et al., 2004) and morphological (Glaubrecht, 2008) evidence. Though significant lake level drops did occur during the late Pliocene and Pleistocene
(ca. 435 m level drop during the early Late Pleistocene megadrought, McGlue et al., 2008), the Tanganyikan freshwater system did not crash, and the lake acted as a refuge for a number of taxa, e.g., the viviparid genus *Neothauma*, that became extinct in the Albertine basin around 1.8 Ma (Van Damme and Pickford, 1999). Consequently, the molluscan in-lake diversification processes observed in some of the Tanganyikan paludomid tribes such as *Lavigeria* may be quite young.

Convincing arguments that abiotic changes did have a severe impact on Lake Tanganyika’s fauna have recently been found by molecular phylogenetic research, e.g., on molluscivorous plathytelphusid crabs, limnochrome cichlids, *Synodontis* catfish and mastacembelid eels. The studies indicate that the radiations in these groups are geologically recent events (late Pliocene–early Pleistocene; Cumberlidge et al., 1999; Marijnissen et al., 2006, 2008; Duftner et al., 2005; Day and Wilkinson, 2006; Brown et al., 2010). Since speaking in geological terms, the specialized molluscivorous crabs evolved only recently, the onset of the radiation in the thalassoid *Lavigeria* and others, considered to result from an arms-race between crabs and their prey (West et al., 1992), must also date from late Pliocene or early Pleistocene times at the earliest. The quite impressive Holocene radiation of *Lavigeria* in Lake Rukwa (Fig. 3), when this lake was joined with Lake Tanganyika via the Karama Gap (Cox, 1939; Cohen et al., 2010) may indicate that the evolutionary divergence process in *Lavigeria* is still an ongoing process. On-going divergence is also suggested for the plathytelphusid crabs (Marijnissen et al., 2008).

In all East African lakes the current molluscan radiations are recent events, postdating the Pliocene–Pleistocene aridity crises. Lake Tanganyika excepted, the populations involved in the modest modern radiation events belong to Pleistocene invaders replacing older Pliocene in-lake lineages that went extinct. In Lake Tanganyika the increased environmental stress triggered a spectacular radiation but only in a single taxon, *Lavigeria*, the latest estimate of this clade being ca. 43 species (Ngereza, 2010). This marked radiation in Lake Tanganyika may also indicate that considerable ecospace became available due to the extinction of other groups. The *Lavigeria* radiation certainly is the result of an in-lake process, and the onset of this divergence-event probably falls in the same time segment as the extinction-events in the other lakes. That the genus is an ancient occupant of the lake is not even certain since a closely related taxon (*Potadomoides*) occurs in the Malagarasi and Luapula (Brown, 1994) and an invasion or multiple invasions could have occurred during transgressive periods throughout Pliocene–Pleistocene times.

6 Conclusions

The available data indicate that probably already during late Pliocene times a marked basin endemism had developed in the Malawi malacofauna, in *Lanistes*, *Gabbiiella* and *Chambardia*. In the paleo-lake that formed subsequently in this basin during an East African late Pliocene wet phase, intra-lacustrine divergence processes were initiated but ended abruptly due to increasing aridity crises. The modern lake fauna has no direct relation with the late Pliocene one, possibly except for some salinity or drought resistant species, such as respectively *Gabbiiella stanleyi* and *Chambardia nyassaensis*. The fact that in modern Lake Malawi the colonization of the lacustrine environment remains mainly restricted to sandy bottoms of the upper epilimnion suggests that even now, during the present freshwater optimum, exploitation of the available space and niches is suboptimal. The cycle of extinction and incipient radiation in the Malawi molluscs seems to have been the norm in the African rift lakes since late Pliocene times regardless of their size and depth. The only exception appears to be Lake Tanganyika, that seems to have acted primarily as a refugium for part of the older Pliocene malacofauna during Pleistocene megadroughts and not as a center of in-lake evolution. Apparently in Lake Tanganyika the Pleistocene climatological destabilization was insufficient to cause the freshwater ecosystem to crash, but it nevertheless altered the limnological environment so drastically that spectacular divergence processes started, e.g., in plathytelphusid crabs and in one thalassoid molluscan taxon, *Lavigeria*.

The terminal Pliocene–early Pleistocene aridity crises had continent-wide impact on the African malacofauna and this geological abrupt event did initiate major extinctions as well as radiations. Therefore, we propose that the beginning of the Pleistocene is used as a reference point in calibrating molecular clocks for African freshwater mollusc phylogeny instead of estimates of the age suggested for the earliest formation of a lake in any given rift basin. This proposal is consistent with recent molecular clock divergence estimates for several Tanganyikan non-molluscan groups (see higher), though some of these estimates place this event around the very onset of the Pleistocene or Gelasian Stage, i.e., between ca. 3–2 Ma. Here a slightly younger age of ca. 1.8–1.6 Ma, i.e., late in the early Pleistocene at the beginning of the Calabrian Stage, is proposed, since this coincides with a time of major molluscan extinctions and extreme aridity in East Africa (Trauth et al., 2005, 2007, 2010). A late in the early Pleistocene or even a Middle to Late Pleistocene date is also consistent with molecular clock divergence time estimates of modern Lake Malawi molluscs (Genner et al., 2007b; Schultheiß et al, 2009, 2011), modern Lake Malawi *Bathyclarias*, *Clariidae* (Agnèse and Teugels, 2001), Mbuna and non-Mbuna cichlids (Won et al., 2005, 2006) and of cichlids in paleo-lake Makgadikgadi and in modern L. Victoria (Genner et al, 2007a).
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