Redescription, molecular characterisation and taxonomic re-evaluation of a unique African monitor lizard haemogregarine *Karyolysus paradoxa* (Dias, 1954) n. comb. (Karyolysidae)

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**Abstract**

**Background:** Within the African monitor lizard family Varanidae, two haemogregarine genera have been reported. These comprise five species of *Hepatozoon* Miller, 1908 and a species of *Haemogregarina* Danilewsky, 1885. Even though other haemogregarine genera such as *Hemolivia* Pettit, Landau, Baccam & Lainson, 1990 and *Karyolysus* Labbé, 1894 have been reported parasitising other lizard families, these have not been found infecting the Varanidae. The genus *Karyolysus* has to date been formally described and named only from lizards of the family Lacertidae and to the authors’ knowledge, this includes only nine species. Molecular characterisation using fragments of the 18S gene has only recently been completed for but two of these species. To date, three *Hepatozoon* species are known from southern African varanids, one of these *Hepatozoon paradoxa* (Dias, 1954) shares morphological characteristics alike to species of the family Karyolysidae. Thus, this study aimed to morphologically redescribe and characterise *H. paradoxa* molecularly, so as to determine its taxonomic placement.

**Methods:** Specimens of *Varanus albigularis albigularis* Daudin, 1802 (Rock monitor) and *Varanus niloticus* (Linnaeus in Hasselquist, 1762) (Nile monitor) were collected from the Ndumo Game Reserve, South Africa. Upon capture animals were examined for haematophagous arthropods. Blood was collected, thin blood smears prepared, stained with Giemsa, screened and micrographs of parasites captured. Haemogregarine morphometric data were compared with the data for named haemogregarines of African varanids. Primer set HepF300 and HepR900 was employed to target a fragment of the 18S rRNA gene and resulting sequences compared with other known haemogregarine sequences selected from the GenBank database.

**Results:** *Hepatozoon paradoxa* was identified infecting two out of eight (25 %) *V. a. albigularis* and a single (100 %) *V. niloticus* examined. Phylogenetic analyses revealed that *H. paradoxa* clustered with the ‘Karyolysus’ clade, and not with those of reptilian *Hepatozoon* spp.

**Conclusions:** In addition to this being the first morphological and molecular characterisation of a haemogregarine within the African Varanidae, it is the first report of a species of *Karyolysus* infecting the monitor lizard family. Furthermore, this constitutes now only the third described and named *Karyolysus* species for which there is a nucleotide sequence available.

**Keywords:** Haemogregarine taxonomy, *Hepatozoon*, Monitor lizard, Haematozoa, *Hemolivia*
Background

Within the apicomplexan order Adeleiorina, representatives of two haemogregarine genera, *Hepatozoon* Miller, 1908 and *Karyolysus* Labbé, 1894, are commonly reported infecting saurians. The genus *Hemolivia* Petit, Landau, Baccam & Lainson, 1990 on the contrary, even though reported parasitising saurian hosts, has but a single described species *Hemolivia mariae* Smallridge & Paperna, 1997 [1, 2]. Representatives of *Hepatozoon* are the most common and are cosmopolitan parasites found parasitising a wide range of vertebrate hosts from amphibians and reptiles to birds and mammals [3, 4]. *Karyolysus*, conversely, is known mainly as a saurian haemogregarine genus that primarily parasitises lizards of the family Lacertidae, but has also been reported from lizards of the Scincidae [1, 5–7]. Besides this discrepancy in vertebrate host preference of the species in the above haemogregarine genera, species in these genera also demonstrate different developmental patterns. Even though species of all three of the haemogregarine genera may be transmitted to the saurian host through the ingestion of the infected invertebrate vector, *Hepatozoon* spp. may be transmitted through a wide range of arthropod vectors (mosquitoes to ticks), whilst transmission of *Hemolivia* spp. and *Karyolysus* spp. has been recorded only through a tick and mite vector, respectively [1].

Whilst more than 30 *Hepatozoon* spp. have been recorded from saurians throughout Africa [8], *Karyolysus* spp. have mainly been reported from lacertid lizards of Europe and Asia [7]. Until Smith’s [3] revision of the Hepatozoidae, the genus *Karyolysus* comprised 11 species, Smith [3] reassigning two of these to *Hepatozoon*, now *Hepatozoon berestnewi* (Finkelstein, 1907) and *Hepatozoon bicapsulata* (Franca, 1910). To the authors’ knowledge, to date, nine species of *Karyolysus* are considered valid: *Karyolysus lacertae* Danilewsky, 1886; *Karyolysus lacazei* Labbé, 1894; *Karyolysus biretortus* Nicolle, 1904; *Karyolysus zuluetai* Reichenow, 1920; *Karyolysus subtilis* Ricci, 1954; *Karyolysus octocromosomi* Alvarez-Calvo, 1975; *Karyolysus latus* Svahn, 1975; *Karyolysus minor* Svahn, 1975; and the only species reported from sub-Saharan Africa *Karyolysus poleensis* Mutunga & Diopelu, 1989 [3, 6, 7]. Descriptions of haemogregarine species were until recently based on morphological characteristics and life-cycle data [7]. This is particularly true of the haemogregarines described from saurians within southern Africa. Haemogregarines, specifically species of *Hepatozoon*, have been commonly recorded blood parasites of southern African saurians including those from lizard genera of the families Cordylidae and Varanidae [8, 9]. Within the African Varanidae, six species of *Hepatozoon* have been described, three of these from southern Africa (Table 1), the latter three comprising *Hepatozoon varani* (Laveran, 1905) from *Varanus niloticus* (Linnaeus, 1762) in South Africa [3, 6], *Hepatozoon camarai* (Dias, 1954) and *Hepatozoon paradoxa* (Dias, 1954) from *Varanus albicularis albicularis* Daudin, 1802 in Mozambique [3, 6].

The aim of the present study was thus to provide a morphological redescription of *H. paradoxa* and molecular data aiding in the correct taxonomic placement of this parasite.

Methods

Study area, *Varanus* spp. collection and blood preparation

Specimens of *Varanus albicularis albicularis* and *Varanus niloticus* were collected in daylight during the summer months of November 2013, February and November 2014, and February 2015 in the Ndumo Game Reserve (NGR) (26°52′00.0″S, 32°15′00.0″E), north-eastern KwaZulu-Natal (KZN), South Africa, bordering southern Mozambique [10]. Lizards were restrained by hand whilst blood and any haematophagous arthropods were collected in situ. A small volume of blood (approximately one drop) was collected from the ventral caudal vein using an appropriately gauged (depending on the size of the lizard) sterile needle and 1 ml syringe. A small portion of the collected blood was used to prepare 2–3 duplicate thin blood smears and the remainder dropped into an equal volume of 70 % ethanol for future molecular analysis. Thin blood smears once air-dried in a dustproof container were fixed in absolute methanol and stained thereafter using a modified solution of Giemsa stain (FLUKA, Sigma-Aldrich, Steinheim, Germany) according to the methods of [11, 12].

Screening of *Varanus* spp. blood smears

Smears were screened under a 100× oil immersion objective on a Nikon Eclipse E800 compound microscope (Nikon, Amsterdam, The Netherlands) and images were captured with an attached Nikon digital camera and accompanying software. Haemogregarines were identified to species level by comparing morphometric data to that of previous studies on African *Varanus* spp. haemogregarines [9, 13–15] (see Table 1). Parasitaemia was calculated per 100 erythrocytes, with c.10⁴ erythrocytes examined per blood smear [16–18].

DNA extraction, PCR amplification and 18S rDNA sequence analysis

Ethanol-preserved blood samples were used for molecular work. Genomic DNA of haemogregarine species was extracted from the samples using a rapid DNA extraction method as detailed in the KAPA Express Extract Kit (Kapa Biosystems, Cape Town, South Africa). Based on previous studies, amplifying fragments of the 18S rRNA gene of reptile haemogregarines of the genera *Karyolysus*
Hemolivia – Butaeva, 1996 (GenBank: Duméril & Bibron, 1836 (Wolbach, 1914) when first discovered was tentatively thought to be Dactylosoma Hepatozoon XR+ imaging system (Bio-Rad, Hemel Hempstead, UK) – 3

and (Kenya) V. niloticus 41

8.5 × 1.5

6

2.5 [Karyolysus Haemogregarina 3×1 Varanus 3 – [19], identification

V. niloticus 9

7

V. V. Adelina grylli 2[20 × 1.5 Adelina dimidiata μ et al. Parasites & Vectors Hepatozoon (syn. 3 – 6 b 14 30 V. griseus described from African varanids 15 1 – 2.5/6.0 and one V. niloticus V. niloticus 43 Hepatozoon toddi 30 Danilewsky, 1885 46 – Thermal Cycler PCR machine (Bio-Rad, UK). Reactions were undertaken in a Bio-Rad C1000 Touch™ Thermal Cycler PCR machine (Bio-Rad, Hemel Hempstead, UK). Resulting amplicons were visualized under ultraviolet light on a 1 % agarose gel stained with gel red using a Bio-Rad GelDoc™ XR+ imaging system (Bio-Rad, Hemel Hempstead, UK). Two PCR products from each sample were sent to a commercial sequencing company (Inqaba Biotechnical Industries (Pty) Ltd, Pretoria, South Africa) for purification and sequencing in both directions. Resultant sequences were assembled, and chromatogram-based contigs were generated and trimmed using Geneious Ver. 7.1 [21]. Sequences were identified using the Basic Local Alignment Search Tool (BLAST) [22], and deposited in the NCBI GenBank database under accession numbers KX011039 and KX011040.

Comparative sequences for species of Hemolivia, Hepatozoon, Karyolysus, Haemogregarina, Dactylosoma Labbé 1894 and Babesiosoma Jakowska & Nigrelli, 1956 parasitising reptiles, amphibians, mammals and ticks were downloaded from GenBank and aligned to the sequences generated in this study. Adelina dimidiata Schneider, 1875, Adelina grylli Butaeva, 1996 (GenBank: DQ096835–DQ096836) and Klossia helicina Schneider, 1875 (GenBank: HQ224955) were chosen as the outgroup to root the phylogeny.

Sequences were aligned using the MUSCLE alignment tool [23] implemented in Geneious 7.1. The

| Table 1 Species of haemogregarines of the genus Hepatozoon described from African varanids |
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| **Species** | **Type-host** | **Type-locality** | **Other hosts (localities)** | **Peripheral gamont/nucleus dimensions** | **Reference** |
| Hepatozoon borelli (Nicolle & Comte, 1906) Smith, 1996 | Varanus griseus (Daudin, 1803) | Tunisia | 7–8 × 2/1–3 × 1–2 | [3], [6, 14] |
| Hepatozoon camarai (Dias, 1954) Smith, 1996 | Varanus albogularis albogularis (Daudin, 1802) | Mozambique | 2 forms observed: banana-shaped: 11.75 × 5.00; long curved: 14.25–18.25 × 1.25–4.25/nucleus irregular | [3], [9] |
| Hepatozoon paradoxus (Dias, 1954) Smith, 1996 | V. a. albogularis | Mozambique | V. niloticus (Kenya); V. a. albogularis and V. niloticus (South Africa) | 6.75–7.50 × 4.25–5.50/nucleus not visible1; 8.1 × 5.2/nucleus irregular or not visible2; 6.99 × 4.39/nucleus not visible3 | [9], [30], this study, [3] |
| Haemogregarina roshdyi Ramdan, Sauod, Mohammed & Fawzi, 1996 (probably Hepatozoon roshdyi) (Ramdan, Sauod, Mohammed & Fawzi, 1996)] | V. griseus | Egypt | 13–20 × 1.5–2.5/6.0/8 × 1.5–2.5 | [40] |
| Hepatozoon toddi (Wolbach, 1914) Smith, 1996 | Varanus niloticus (Linnaeus, 1762) | The Gambia | 10.3 × 2.5/not given | [3], [6, 15] |
| Hepatozoon varani (Laveran, 1905) Smith, 1996 | V. niloticus | South Africa | V. niloticus and V. griseus (Senegal); V. niloticus (Portuguese Guinea); V. niloticus (The Gambia); V. niloticus (Senegal); Varanus arenarius Duméril & Bibron, 1836 (French West Africa); V. griseus (French Sudan); V. niloticus (Kenya) | 14 × 3/not given1; 11–15 × 3/not given2; 12.0 × 13.5/4–5 × 3.0; 10.3 × 2.5/not given2; two forms: 12–14 × 2/not given, 10–12 × 4–5; 5–6 × 4–5; report only8; report only2; 12.7 × 4.6/not given | [13], [41], [15], [42], [43], [44], [45], [46], [30], [3] |

1 Corresponding description or report of the haemogregarine species and its bibliographic reference

[7], Hemolivia [12] and Hepatozoon [19], identification of the parasite of the two Varanus species, two V. a albogularis and one V. niloticus (n = 3) from the current study was completed using the primer set HepF300 (5′–GTT TCT GAC CTA TCA GCT TTC GAC G-3′) and HepR900 (5′–CAA ATC TAA GAA TTT CAC CTC TGA C-3′). The PCR reactions were run targeting a fragment (approximately 600 nt) of the 18S rRNA gene [20]. Conditions for PCR were as follows: initial denaturation at 95 °C for 3 min, followed by 35 cycles, entailing a 95 °C denaturation for 30 s, annealing at 60 °C for 30 s with an end extension at 72 °C for 1 min, and following the cycles a final extension of 72 °C for 10 min as detailed according to previous methods [12, 18]. PCR reactions were performed with volumes of 25 µl, using 12.5 µl Thermo Scientific DreamTaq PCR master mix (2×) (2× DreamTaq buffer, 0.4 mM of each dNTP, and 4 mM MgCl2), 1.25 µl of each primer, and at least 25 ng DNA. The final reaction volume was made up with PCR-grade nuclease free water (Thermo Scientific, Vilnius, Lithuania). Reactions were undertaken in a Bio-Rad C1000 Touch™ Thermal Cycler PCR machine (Bio-Rad, Hemel Hempstead, UK). Resulting amplicons were visualized under ultraviolet light on a 1 % agarose gel stained with gel red using a Bio-Rad GelDoc™ XR+ imaging system (Bio-Rad, Hemel Hempstead, UK). Two PCR products from each sample were sent to a commercial sequencing company (Inqaba Biotechnical Industries (Pty) Ltd, Pretoria, South Africa) for purification and sequencing in both directions. Resultant sequences were assembled, and chromatogram-based contigs were generated and trimmed using Geneious Ver. 7.1 [21]. Sequences were identified using the Basic Local Alignment Search Tool (BLAST) [22], and deposited in the NCBI GenBank database under accession numbers KX011039 and KX011040.
alignment consisted of 47 sequences, manually trimmed to a total length of 968 nt. Uncorrected pair-wise distances (p-distance), base pair differences as well as parsimony informative sites and the number thereof were identified or determined with the MEGA6 bioinformatics software program [24] for the aligned 18S rDNA sequences between all available species appearing in the phylogenetic analyses.

To infer phylogenetic relationships of the aligned dataset both Bayesian inference (BI) and Maximum likelihood (ML) methods were used. A comprehensive model test was performed to determine the most suitable nucleotide substitution model, according to the Akaile information criterion using jModelTest 2.1.7 [25, 26]. The best model identified was the Transversion Model plus with estimates of invariable sites and a discrete Gamma distribution (TVM+I+Γ). This model was substituted with the General Time Reversible model (GTR+I+Γ) for phylogenetic analysis, as this was the most appropriate model available with the best AICc score. The BI analysis was implemented from within Geneious 7.1 using MrBayes 3.2.2 [27]. The analysis was run twice over 10 million generations for the Markov Chains Monte Carlo (MCMC) algorithm. The Markov chain was sampled every 100 cycles, and the MCMC variant contained 4 chains with a temperature of 0.2. The log-likelihood values of the sample point were plotted against the generation time and the first 25 % of the trees were discarded as ‘burn-in’ with no ‘burn-in’ samples being retained. The ML analysis was performed using RAxML Ver. 8.1.22 [28] implemented in the raxmlGUI Ver. 1.3 [29]. The alpha-parameter selected was the GTR+I+Γ model, with support assessed using 1,000 rapid bootstrap inferences. Resulting trees were combined in a 50 % majority consensus tree.

**Ethics statement**

This study received the relevant ethical approval (North-West University ethics approval no: NWU-00005-14-S3).

**Results**

**Prevalence, parasitaemia and general observations of H. paradoxa in peripheral blood smears**

Six adult and two juvenile *Varanus albigularis albigularis* (Fig. 1a, b) and one adult *Varanus niloticus* were captured, sampled for blood parasites and examined for possible haematophagous arthropods. The two juveniles were found to be negative for blood parasites, whilst 4/6 (67 %) adult *V. a. albigularis* and 1/1 (100 %) adult *V. niloticus* were found positive for haemogregarine infections. Two adult (33 %) (with a parasitaemia of c.5 and 20 %, respectively) *V. a. albigularis* and the *V. niloticus* (with a parasitaemia of c.0.2 %) were parasitised by a haemogregarine alike to *Hepatozoon paradoxa* described by Dias [9] (Figs. 1c–f and 2b–i). One of the two (17 %) *V. a. albigularis* (parasitaemia of 5 %) and the single *V. niloticus* were found to have a co-infection with another haemogregarine of a different *Hepatozoon* spp. (Cook, Netherlands & Smit, unpublished observations).
parasitaemia of c.0.4 and 3 %, respectively). The latter unidentified Hepatozoon species was also found infecting the remaining two parasitised V. a. albigularis (parasitaemia of c.0.1 and 0.5 %, respectively). All specimens of both species of Varanus were infested with adult and juvenile stages of the Leguan tick Amblyomma exornatum (Koch, 1844) (Fig. 1b), the two juvenile V. a. albigularis and V. niloticus demonstrating lower densities of these ectoparasites. These ectoparasites were found all over the animal, with the highest densities in the nostrils, often blocking them, also on the area surrounding the eyes, and along the edges of the mouth. It was not uncommon to find dead ticks and their remains deep within the nostril, their ability to exit blocked by later arrivals. Squashes of nymphal and adult female and male ticks provided no parasitic stages. No other haematophagous arthropods, including mites, were observed.

Stages of the H. paradoxa-like haemogregarine observed in peripheral blood smears from this study were compared morphologically with those observed in previous blood parasite studies of African varanids [9, 13–15] (Table 1). In size and morphology, the H. paradoxa-like stages observed during the current study conformed to those described by Dias [9] and Ball [30] (see Table 1, Figs. 1a–f, 2b–i and 3a–d, respectively). Two stages of the parasite were observed: a stage unreported by Dias [9], but possibly by Ball [30], a rare possible trophozoite stage (Figs. 1c and 3a), and what was identified as a mature intra-erythrocytic gamont stage (Figs. 1d–f, 2b–i and 3d, e).

Karyolysus paradoxa (Dias, 1954) Cook, Netherlands & Smit, 2016
Syns Haemogregarina paradoxa Dias, 1954; Hepatozoon paradoxa Smith, 1996.

Type-host: Varanus albigularis albigularis Daudin, 1802, Squamata: Varanidae [9].
Other hosts: Varanus niloticus (Linnaeus in Hasselquist, 1762), Squamata: Varanidae [30]; present study.
Vector: Unknown.
Type-locality: Ndumo Game Reserve (26°54′18.5″S, 32°19′24.7″E), KwaZulu-Natal, South Africa (present study).
Other localities: Maputo, Mozambique [9]. Ball [30] also described a haemogregarine morphologically alike to K. paradoxa in Marimonti, near Meru, Kenya.

Fig. 2 Illustration of Haemogregarina paradoxa Dias, 1954 in Varanus albigularis albigularis Daudin, 1802 [9]. a–i Redrawn and adapted from Dias (1954). Illustrations representing the original description of Karyolysus paradoxa (syns. Hepatozoon paradoxa and Haemogregarina paradoxa) ex Varanus albigularis albigularis from Mozambique. a Healthy non-parasitised erythrocyte. b–i Parasitised erythrocytes, note the shrinkage of the host cell, the heavy vacuolization of the host cell nucleus, and the thick capsule surrounding the gamont which results in the gamont nucleus being invisible. Scale-bar: 10 μm.
Type-material: Neohapantotype, 1× blood smear from the type-host Varanus niloticus (Linnaeus in Hasselquist, 1762), deposited in the protozoan collection of the National Museum, Bloemfontein, South Africa under accession number NMB P 410. Other voucher material deposited that includes stages of Haemoproteus paradoxa 1× blood smear from Varanus niloticus, deposited in the protozoan collection of the National Museum, Bloemfontein, South Africa under accession number NMB P 411.

Representative DNA sequences: Two sequences representing a 611 and 613 nt fragment of the 18S rRNA gene of K. paradoxa isolated from the type-host Varanus albignularis albignularis, deposited in the NCBI GenBank database under the accession numbers KX011039 and KX011040, respectively.

Redescription (Figs. 1, 2 and 3)

Trophozoite. Rare, ovoid, with vacuolated cytoplasm, measuring 6.5–6.9×4.3–4.7 (6.7×4.5) μm (n = 2); nucleus with loose chromatin, staining pink (Fig. 1c). Both trophozoites parasitising young erythrocytes, no host cell distortion visible.

Mature gamont. Rounded in shape, gamont seemingly folded within with a well-developed capsule (Fig. 1d–f), measuring 6.3–7.9×3.6–5.2 (7.0×4.4) μm (n = 20). Cytoplasm staining whitish-blue; nucleus not visible. Notable destruction of host cell cytoplasm and karyolysis of the host cell nucleus, causing an observable heavily vacuolated and foamy appearance (Fig. 1d–f).

Remarks

The haemogregarine described in this study from South African Varanus albignularis albignularis and Varanus niloticus (Fig. 1d–f) was found to be morphologically similar to Hepatozoon paradoxa described by Dias [9] from a specimen of V. a. albignularis in neighbouring Mozambique (Fig. 2b–i). It shared a number of unique characteristics including destruction of the infected host erythrocyte, consisting of dehaemoglobinisation resulting in shrinkage of the host cell and destruction of the host cell nucleus (characteristic of a number of species of Karyolysus [1]) resulting in a heavily vacuolated appearance (Figs. 1d–f and 2b–i). Additionally, the haemogregarine in this study agrees well with the size of H. paradoxa in the original description of Dias [9] (mean 7.0×4.4 vs 7.0×4.9 μm) (Table 1).

The same unique characteristics were reported of a haemogregarine found infecting a V. niloticus by Ball [30] from Kenya, measuring on average 8.1×5.2 μm (Fig. 3c, d). However, in Ball’s [30] study, additional, presumably younger, stages were observed (similar to the young trophozoite stage found in our study) (Figs. 1c and 3a). Ball [30] also noted a single possibly dividing stage of these trophozoites (Fig. 3b). In cells parasitised by all these possible trophozoite stages, the host erythrocyte showed no shrinkage as of yet, but was according to Ball’s report abnormal in shape and staining. At first, Ball [30] did assume this parasite to represent younger stages of another haemogregarine that has been reported parasitising African varanids Hepatozoon varani (Laveran, 1905) Smith, 1996 (Table 1). However, based on the effects of the parasite resembling H. paradoxa as described by Dias [9], its destruction of the host cell and the host cell’s nucleus, he concluded that this parasite was not H. varani. Overall, for the K. paradoxa described in this study and the parasites described in the other two studies [9, 30], the nucleus and cytoplast was not visible owing to what appeared to be a thick enclosing capsule as seen with the gamonts of species of Hemolivia, see [12, 31] (Figs. 1d–f, 2b–i and 3c, d). Only on rare occasion in the present study and that of Ball’s [30] was the parasite seen to be folding.
over on itself (Figs. 1d and 3c). Otherwise, the only evidence of this behaviour was a crescent shaped stain at the centre of the oval parasite (as seen in all three reports) (Figs. 1, 2 and 3). It is based on the above unique characteristics, particularly of the mature gamont stages, that we suggest all three reports are of the same parasite species *K. paradoxa*.

No hapantotype, according to the International Code of Zoological Nomenclature (ICZN) Article 73.3, was designated and deposited during the original description of *K. paradoxa* by Dias [9]. Ball [30] also did not identify the parasite to taxon level and did not deposit any voucher material. Furthermore, our efforts to locate any original specimens or voucher material were unsuccessful. In this study, *K. paradoxa* was collected from Ndumo Game Reserve, northern KwaZulu-Natal, South Africa, bordering the south of Mozambique. The original description by Dias [9] was collected in the vicinity of Maputo in the southern parts of Mozambique approximately 300 km from the NGR. Additionally, *K. paradoxa* in the present study was collected from the same host species *Varanus albigularis albigularis* as in the original description by Dias [9]. Based on the above, the mature gamont size comparisons, and the unique characteristics of the mature gamonts of *K. paradoxa* (destruction and shrinkage of the infected host erythrocyte, destruction of the host cell nucleus resulting in vacuolation, and the thick non-staining capsule) as described above for all three reports of this parasite, which includes the original description by Dias [9], and in accordance with ICZN Article 75.3, we designate a neohapantotype. The present study also includes both the description of an additional stage of the parasite (a trophozoite) and provides sequence data (fragment 18S rDNA), which was not provided by Dias [9] in his original description of this parasite species. This neohapantotype is deposited in the protozoan collection of the National Museum, Bloemfontein, South Africa under accession number NMB P 410.

**Sequence identification and phylogenetic analysis**

Amplicons of 611 and 613 nt for the 18S rRNA gene of *K. paradoxa* were obtained from the *V. a. albigularis* with the seemingly pure and highest parasitaemia infection using primer sets HepF300 and HepR900. No *K. paradoxa* isolate was obtained from the *V. niloticus*, likely due to its low parasitaemia in comparison to the concurrent infection of the other unidentified *Hepatozoon* species (unpublished data). The details of the species used in the phylogenetic analyses and presented in the consensus tree are provided in Table 2. The topology of both the BI and ML analyses were overall similar, with discrete monophyletic clades of known and likely belonging to *Karyolysus* species, *Hepatozoon* spp. of mammals ('intraerythrocytic' *Hepatozoon* spp.), the herpatofauna ('intraerythrocytic' *Hepatozoon* spp.), *Hemolivia* spp., *Haemogregarina* spp., and the Dactylosomatidae (Fig. 4). Our results showed that *K. paradoxa* clustered within a major monophyletic clade containing both known (morphologically and ecologically confirmed) and likely belonging to *Karyolysus* species, and ‘intraerythrocyclic’ *Hepatozoon* species of mammals that were sister to the *Karyolysus* clade. This major clade was distinct from the major monophyletic clade containing the herpatofaunal ‘intraerythrocytic’ *Hepatozoon* spp. and *Hemolivia* spp.

The parasite collected in the present study that morphologically conformed to the original description of *Hepatozoon paradoxa* formed a sister taxon to the larger ‘*Karyolysus*’ clade (containing the known and likely belonging to *Karyolysus* species) in the phylogenetic analysis (Fig. 4). According to evolutionary divergence estimates the present material was most closely related to known *Hepatozoon* spp. (at 97.7 %, p-distance = 0.02) than to species of the genera *Hepatozoon*, *Hemolivia* and *Haemogregarina* (Table 3).

**Discussion**

*Varanus albigularis albigularis* and *Varanus niloticus* are known to display somewhat different habitat preferences, preferring more terrestrial and aquatic environments respectively [32]. Both *V. a. albigularis* and *V. niloticus* can be found throughout South Africa from the more tropical Indian Ocean coastal belt in the East, west to the margins of the more arid Northern and Western Cape provinces [32]. Both species often occur sympatri- cally, particularly in Ndumo Game Reserve, rendering the finding of the same parasite in both these species not surprising.

As the morphological and developmental characteristics typical of *Haemogregarina* spp. had never been observed in haemogregarines of the herpatofauna, *K. paradoxa* was transferred along with many other species from the herpatofauna, birds and mammals, from the genus *Haemogregarina* (*Haemogregarinidae*) to the genus *Hepatozoon* (*Hepatozoa*) by Smith [3] during a systematic review of the Hepatozoidae. However, during the first detailed revision of *K. paradoxa* provided here in the present study, morphologically this species shares more characteristics with members of the family *Karyolysidae*. *Karyolysus paradoxa* peripheral blood gamonts appear to be encapsulated as in the *Hemolivia* and destroy the host cell nucleus as in a number of species of *Karyolysus* (as mentioned above in Remarks) [1, 7]. However, it is imperative to take into account that these morphological features, specifically for the latter genus, are not always present [7].

The phylogenetic position of *K. paradoxa* was shown to be at the base of a clade containing undescribed species of *Hepatozoon*, many of these the
| Clade                              | Organism                       | Host                          | Host Family | Common name                      | Accession number | Reference |
|-----------------------------------|--------------------------------|-------------------------------|-------------|---------------------------------|------------------|-----------|
| **Karyolysus**                    | Karyolysus latus               | Podarcis muralis             | Lacertidae  | Common wall lizard               | KJ461939         | [7]       |
|                                  | **Hepatozoon sp.**             | **Algyroides marchi**        | Lacertidae  | Spanish keeled lizard           | JX531933         | [34]      |
|                                  | **Hepatozoon sp.**             | **Podarcis vaucheri**        | Lacertidae  | Andalusian wall lizard          | HQ734803         | [33]      |
|                                  | **Karyolysus lacazei**         | **Lacerta trilineata**       | Lacertidae  | Balkan emerald lizard           | KJ461942         | [7]       |
|                                  | **Karyolysus sp.**             | **Zootoca vivipara**         | Lacertidae  | Viviparous lizard                | KJ461945         | [7]       |
|                                  | **Hepatozoon sp.**             | **Podarcis bocagei**         | Lacertidae  | Bocage's wall lizard            | JX531952         | [34]      |
|                                  | **Hepatozoon sp.**             | **Podarcis bocagei**         | Lacertidae  | Bocage's wall lizard            | JX531954         | [34]      |
|                                  | **Hepatozoon sp.**             | **Timon tangitanus**         | Lacertidae  | Moroccan eyed lizard            | HQ734799         | [33]      |
|                                  | **Hepatozoon sp.**             | **Atlantolacerta andreanskyi**| Lacertidae  | Andreansky's lizard             | HQ734798         | [33]      |
|                                  | **Hepatozoon sp.**             | **Ophionyssus sp. ex Lacerta viridis**| Lacertidae  | European green lizard          | KJ461944         | [7]       |
|                                  | **Hepatozoon sp.**             | **Psammophis schokari**      | Lamprophiidae | Schokari sand racer          | KC696565         | [47]      |
|                                  | **Hepatozoon sp.**             | **Hemorrhhois hippocrepis** | Colubridae  | Horseshoe snake                 | JX244269         | [35]      |
|                                  | **Hepatozoon sp.**             | **Eumeues algeniensis**      | Scincidae   | Algerian skink                  | HQ734796         | [33]      |
|                                  | **Hepatozoon sp.**             | **Sclerarcis perspicillata** | Lacertidae  | Menorca wall lizard             | HQ734791         | [33]      |
|                                  | **Hepatozoon sp.**             | **Podarcis vaucheri**        | Lacertidae  | Andalusian wall lizard          | HQ734804         | [33]      |
|                                  | **Hepatozoon sp.**             | **Podarcis hispanicus**      | Lacertidae  | Iberian wall lizard             | JX531917         | [34]      |
|                                  | **Hepatozoon sp.**             | **Podarcis lilfordi**        | Lacertidae  | Lilford’s wall lizard           | JX531920         | [34]      |
|                                  | **Karyolysus (syn. Hepatozoon)**| **Varanus altigularis**      | Varanidae   | Rock monitor lizard             | KX011039         | This study |
|                                  | **Karyolysus (syn. Hepatozoon)**| **Varanus altigularis**      | Varanidae   | Rock monitor lizard             | KX011040         | This study |
| **Intraleucocytic**              | **Hepatozoon americanum**      | Canis familiaris             | Canidae     | Domestic dog                    | AF176836         | [48]      |
| Hepatozoon                       | **Hepatozoon canis**           | Canis familiaris             | Canidae     | Domestic dog                    | AY461378         | [49]      |
|                                  | **Hepatozoon ursi**            | Ursus thibetanus japonicus  | Ursidae     | Japanese black bear             | EU041718         | [50]      |
|                                  | **Hepatozoon sp.**             | Martes martes                | Mustelidae  | Pine marten                     | EF222257         | [51]      |
|                                  | **Hepatozoon felis**           | Felis catus                  | Felidae     | Domestic cat                    | AY620232         | [50]      |
|                                  | **Hepatozoon catesbianae**     | Lithobates catesbianus       | Ranidae     | American bullfrog               | HQ224954         | [49]      |
|                                  | **Hepatozoon iaxa**            | Sclerocephrys (syn. Amietophrynus) maculatus | Bufonidae  | Flat-backed toad                | KP119772         | [52]      |
|                                  | **Hepatozoon theileri**        | Amietia queckettii           | Pyxicephalidae | Queckett's river frog       | KP119773         | [18]      |
|                                  | **Hepatozoon sipedon**         | Nerodia sipedon              | Colubridae  | Northern water snake            | JN181157         | [49]      |
|                                  | **Hepatozoon fitzsimonsi**     | Kinixys zambensis            | Testudinidae | Bell's hingeback tortoise       | KRO60084         | [12]      |
Table 2 List of organisms used in the phylogenetic analyses of this study, with associated host, host family and host common name, GenBank accession numbers and references (Continued)

| Clade | Organism          | Host                              | Family                      | Common name                          | Accession number | Reference |
|-------|-------------------|-----------------------------------|----------------------------|--------------------------------------|------------------|-----------|
|       | Hepatozoon ayorgbor | Python regius ("Lampropis (syn. Boaedon) fuliginous) | Pythonidae; Lamprophiidae | Ball python; brown house snake       | EF157822         | [53]      |
|       | Hepatozoon domerguei | Furcifer sp.                      | Chamaeleonidae              | Chameleon species                     | KM234649         | [54]      |
|       | Grandisonia alternans | Indotyphlidae                    | Caecilian                  |                                      | KF246566         | [55]      |
|       | Hepatozoon seychellensis | Python regius ("Lampropis (syn. Boaedon) fuliginous) | Pythonidae; Lamprophiidae | Ball python; brown house snake       | EF157822         | [53]      |
|       | Hemolivia domerguei | Furcifer sp.                      | Chamaeleonidae              | Chameleon species                     | KM234649         | [54]      |
|       | Grandisonia alternans | Indotyphlidae                    | Caecilian                  |                                      | KF246566         | [55]      |
|       | Hemolivia mauremys | Testudo graeca                    | Testudinidae                | Mediterranean spur-thighed tortoise   | KF992700         | [36]      |
|       | Hemolivia parvula  | Kinixys zambensis                 | Testudinidae                | Bell's hingeback tortoise             | KR069082         | [12]      |
|       | Hemolivia mariae   | Egeria stokesii                   | Scincidae                  | Gidgee spiny-tailed skink             | KF992711         | [36]      |
|       | Hemolivia sp.      | Rhinoclemmys pulcherrima mann    | Geoemydidae                 | Painted wood turtle                   | KF992714         | [36]      |
|       | Hemolivia mariae   | Grandisonia alternans            | Indotyphlidae              | Mediterranean spur-thighed tortoise   | KF992700         | [36]      |
|       | Hemolivia stellata | Amblyomma rotateum ex Rhinella marna | Amblyomma rotateum ex Rhinella marna | Cane toad                                      | KF992714         | [36]      |
|       | Haemogregarinidae | Haemogregarina sp_3140            | Pelomedusidae               | East African black mud turtle         | KF257925         | [56]      |
|       | Haemogregarina saccalea | Pelusios subniger                | Pelomedusidae               | Four-eyed turtle                      | KM887507         | [57]      |
|       | Haemogregarina pelligrini | Malaclemys subtrijuga            | Geoemydidae                 | Snail-eating turtle                   | KM887508         | [57]      |
|       | Haemogregarina balli | Chelydra serpentina serpentina   | Chelydridae                 | Common snapping turtle                | HQ224959         | [49]      |
|       | Haemogregarina stepanowi | Mauremys caspica                 | Geoemydidae                 | Caspian turtle                        | KF257926         | [56]      |
|       | Dactylosomatidae | Babesiosoma stableri             | Ranidae                     | Mink frog                             | HQ224961         | [49]      |
|       | Dactylosoma ranarum | Pelophylax lessonae (syn. esculentus) | Ranidae                     | Pool frog                             | HQ224957         | [49]      |
|       | Outgroup          | Klossia helicina                 | Cepaea nemoralis            | Helicidae                             | HQ224955         | [49]      |
|       | Adelina dimidiata | Scolopendra cingulata            | Scolopendridae              | Megarian banded centipede             | DQ096835         | [58]      |
|       | Adelina gryllith  | Gryllus bimaculatus              | Gryllidae                   | Field cricket                         | DQ096836         | [58]      |

*Experimental laboratory animal*
results of molecular Hepatozoon spp. surveys [33, 34], and known species of Karyolysus [7]. This may suggest that these undescribed Hepatozoon spp. might rather be species of Karyolysus. The Karyolysus spp. clade is part of a larger clade including a sister clade of Hepatozoon spp. from mammals. This topology, as seen in this study, has been observed in a number of other studies [7, 33–37]. Karadjian et al. [37], in their attempt to understand the relationships of the different haemogregarine genera, particularly in respect to the conundrum of the polyphyletic Hepatozoon clade, proposed, based on their phylogenetic findings, that a number of Hepatozoon species (most of which have not been morphologically described) might rather represent species of Karyolysus. Given that it was only since the recent identification, description and molecular characterisation of Karyolysus spp. by [7], it is only now that we are beginning to realize this possibility. Moreover, it appears that the diversity of squamates parasitised by likely species of Karyolysus is fast increasing, a scenario

Fig. 4 Phylogenetic analysis of Karyolysus paradoxa (Dias, 1954) based on 18S rDNA sequences. Bayesian inference (BI) and Maximum Likelihood (ML) analysis showing the phylogenetic relationships for two Karyolysus paradoxa isolates from the current study represented in bold, 17 Hepatozoon and Karyolysus species (representing the Karyolysus clade in purple), eight herbatofaunal-infecting Hepatozoon species (representing the ‘intraerythrocytic’ Hepatozoon clade in green), five Hemolivia species (representing the Hemolivia clade in orange), five Haemogregarina species (representing the Haemogregarinidae clade in yellow), a Babesiosoma and Dactylosoma species (representing the Dactylosomatidae clade in pink) and two Adelina and one Klossia species (used as the outgroup in grey). All comparative sequences were downloaded from the GenBank database. Tree topologies for both the BI and ML trees were identical; the nodal support values (BI/ML) are represented on the BI tree
which is likely only to intensify in future. A recent molecular Hepatozoon spp. survey by [38] shows haemogregarine isolates from geckos of the genus Tarentola Gray, 1825 (Phyllodactylidae) are also falling within what may be seen as the 'Karyolysus' clade; a clade that at present also includes haemogregarines isolated from species of the families Colubridae and Lamprophiidae (both snakes), Lacertidae and Varanidae (both lizards), and Scincidae (skinks).

Furthermore, the present study shows that K. paradoxa is most closely related to known Karyolysus species, followed by species of the ‘intraleucocytic’ Hepatozoon clade and then species of the ‘intraerythrocytic’ Hepatozoon and Hemolivia clades. Karyolysus and Hemolivia morphologically still belong within the same family (Karyolysisidae), however, with the use of the 18S rRNA gene these two genera can be separated from one another. Clades generally fall in different major clades. Karyolysus clusters in a major clade with the ‘intraleucocytic’ mammal Hepatozoon, whilst Hemolivia clusters in a major clade with the ‘intraerythrocytic’ herpatofoanal Hepatozoon; this finding is apparent in the present study, and also in [36, 37]. It is clear that the relationship between these two genera may possibly only be resolved by using a multi-gene approach as in [39].

**Conclusions**

Based on the morphology and the molecular findings presented in this study, we recommend the following nomenclatural correction: Karyolysus paradoxa (Dias, 1954) (syn. Hepatozoon paradoxa (Dias, 1954) Smith, 1996, Haemogregarina paradoxa Dias, 1954) in the varanid lizards Varanus albigularis albigularis (type-host), and Varanus niloticus. Our results showed that Karyolysus paradoxa is as closely related to species within its current generic assignment in the ‘intraerythrocytic’ herpatofoanal Hepatozoon as it is with the more distantly related species of the Haemogregarina.

Besides this study representing the first morphological and molecular report of a haemogregarine within an African varanid, it is the first report of a species of Karyolysus infecting a host of the Varanidae. Furthermore, it represents the third described and named Karyolysus spp. for which there is a nucleotide sequence available. It is hoped that this study will encourage further molecular work on the Karyolysidae, particularly the genus Karyolysus.

This study also extends the host and distribution range of K. paradoxa from only a single specimen of V. a. albigularis in Mozambique to an additional two specimens in South Africa, as well as including V. niloticus as an additional host both in South Africa and Kenya. The distribution range of K. paradoxa falls within subtropical areas in South Africa, Mozambique and Kenya, and as such it would be interesting to see if this particular parasite is restricted to subtropical areas such as is the case with Hemolivia parvula (Dias, 1953) found parasitising Kinixys zombensis Hewitt, 1931 tortoises of South Africa and Mozambique; see [12], or if it is more widely distributed throughout different biomes as is the case with Hepatozoon fitsimonsi (Dias, 1953) found parasitising several tortoise species in South Africa and Mozambique; see [19].

Even though tick squashes did not result in any observable parasitic stages future studies will focus on identifying possible vectors. Parasitic stages found in these possible vectors will be identified to species level based on both morphological and molecular findings.

With the conundrum of the larger Hepatozoon clade being polyphyletic and absorbing the Hemolivia and Karyolysus, it is important to increase the number of taxa from which we can work and ask deeper phylogenetic questions. However, besides the molecular characterisation of these species, it is still important to focus on their morphology and where possible attempt to elucidate their life-cycles in order to resolve the complex taxonomy of these organisms. More importantly, it is necessary to include another faster evolving gene or even mitochondrial genomes of these groups following [39] before we can make any well-informed decisions.

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Authors’ contributions
All authors conceived and designed the project, participated in general data analysis and in drafting the manuscript. CAC and ECN carried out the fieldwork, prepared and examined blood smears, prepared light micrographs and compiled all measurement data. ECM participated in the molecular studies, and in the sequence alignment and provided support to the preparation of the manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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