**Kupeantha** (Coffeeae, Rubiaceae), a new genus from Cameroon and Equatorial Guinea

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**Abstract**

Two new coffee relatives (tribe Coffeeae, Rubiaceae), discovered during botanical expeditions to Cameroon, are examined for generic placement, and the placement of three previously known species (*Argocoffeopsis fosimondi*, *A. spathulata* and *Calycosiphonia pentamera*) is reinvestigated using plastid sequence (*accD-psa1, rpl16, trnL-F*) and morphological data. Seed biochemistry of the new species and pollen micromorphology (only one of the two species) are also studied. Based on the plastid sequence data, the new taxa are nested in a well-supported monophyletic group that includes *Argocoffeopsis* and *Calycosiphonia*. Within this clade, three well-supported subclades are recovered that are morphologically easy to diagnose: (1) *Calycosiphonia* (excluding *C. pentamera*), (2) *Argocoffeopsis* (excluding *A. fosimondi* and *A. spathulata*), and (3) a clade including the above excluded species, in addition to the new species. Based on the results, *Kupeantha*, a new genus of five species, is described, including two new Critically Endangered taxa from the Highlands of Cameroon: *Kupeantha ebo* and *K. kupensis*. Phytochemical analysis of *Kupeantha* seeds reveals compounds assigned as hydroxycinnamic acid derivatives, amino acids and ent-kaurane diterpenoids; caffeine was not detected. *Kupeantha* is the first new genus described in tribe Coffeeae in 40 years.

**Introduction**

Distribution data, although important to ensure effective conservation management and sustainable utilisation of natural resources [1], is lacking for many organisms in tropical Africa, including vascular plants necessitating renewed efforts for botanical exploration of the region [2]. Based on the RAINBIO dataset containing 600,000+ georeferenced records representing 22,000+ plant species, priority target areas for future sampling efforts were proposed in Tanzania, Atlantic Central Africa and West Africa [2]. One of the areas, which remained underexplored until recently, is Cameroon in West Central Africa. During the 1990s, the Royal Botanic Gardens, Kew and partners in Cameroon began a series of botanical expeditions to...
could input the codes provided by the authors and find all the needed information to contact the relevant herbarium.

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improve fundamental knowledge of plant species diversity and conservation management in western Cameroon and adjoining regions [3]. At that time, SW Cameroon was recorded as having the highest plant species diversity in Tropical Africa per degree square [4], with many species thought to be at risk of extinction. This was later confirmed on publication of the Red Data Book of the Flowering Plants of Cameroon [5], which showed that the highest concentrations of threatened plant species occurred in SW Cameroon.

Locations for botanical expeditions were selected in areas of intact natural vegetation considered of importance by local conservation NGOs that hosted the surveys. With the support of volunteers from the Earthwatch Institute, botanists collected 19,500+ herbarium specimens [6], resulting in the discovery of 100+ species new to science, most of which are threatened with extinction, e.g., Ancistrocladus grandiflorus Cheek [7] and Impatiens etiendensis Cheek & Eb.Fisch. [8], and included two new genera, Korupodendron Litt & Cheek (Vochysiaceae) [9] and Kupea Cheek & S.A.Williams (Triuridaceae) [10]. Many novel Rubiaceae were also discovered [11–16].

In Mt Kupe, in the Bakossi tribal area, specimens of a cloud forest tree species resembling Calycosiphonia Pierre ex Robbr. (Rubiaceae) were collected in 1995, but these had several characters not seen in that genus. Based on morphology, it was suggested as a possible new genus by Bridson (pers. comm. to Cheek), but designated as Calycosiphonia sp. A [6]. Independently, in 2005 and 2006, a series of botanical expeditions focussed on Rubiaceae were led in South Cameroon by Sonké. These resulted in further discoveries of new taxa [17–20]. Still, the study of the family Rubiaceae in Cameroon remains far from complete. Among the collections made were numerous specimens of two species similar to Calycosiphonia sp. A, one with spathulate leaf tips (e.g., Sonké & Nguembou 3783), the other drying black (e.g., Sonké & Djuikouo 4320). Specimen Sonké & Nguembou 3783 was included in the seminal study of tribe Coffeae [21]. In the result of the combined analysis of the molecular and morphological data sets, a clade containing Argocoffeopsis Lebrun, Calycosiphonia and ‘Calycosiphonia cf.’ (Sonké & Nguembou 3783) received strong bootstrap support (BS = 99) and a high Bremer (decay) value (b = 9). Monophyly was retrieved for Calycosiphonia (BS = 85; b = 3) and Argocoffeopsis (BS = 81; b = 2), but Calycosiphonia cf. was placed in an unresolved position in a polytomy with the genera Argocoffeopsis and Calycosiphonia [21]. It was suggested that ‘Calycosiphonia cf.’ might represent an additional genus in the tribe, but that further material and morphological investigations were required [21]. Prior to allocating ‘Calycosiphonia cf.’ to a genus, [22] reviewed the morphological features of Argocoffeopsis and Calycosiphonia, and concluded that there was “no logic in describing Calycosiphonia cf. in a new genus, as apart from the size and shape of its fruit, it shows no morphological differences when compared with either Argocoffeopsis or Calycosiphonia”. They described the taxon represented by Sonké & Nguembou 3783 as Argocoffeopsis spathulata A.P. Davis & Sonké [22]. The second species, represented by Sonké & Djuikouo 4320, was published as Calycosiphonia pentamera Sonké & Robbr., which necessitated broadening the generic concept [23]. Meanwhile, continued botanical expeditions in West Cameroon, this time led by Tchiengué in the cloud forest of the Lebialem Highlands in 2005–2006, led to the discovery of a fourth species of clear affinity with Calycosiphonia sp. A and Argocoffeopsis spathulata. Following [22] on generic placement, this Critically Endangered, attractive, flowering shrub was published as Argocoffeopsis fosimondi Tchiengué & Cheek [24]. A fifth species of ‘Calycosiphonia cf.’ was discovered in the proposed Ebo National Park of Cameroon’s Littoral Region with records in 2007 (Fenton 134) and 2015 (Alvarez 11).

The goal of this study is to use morphological and plastid sequence data to (1) place the newly discovered species from Mt Kupe and Ebo; and (2) reassess the generic placement of the recently described Argocoffeopsis fosimondi, A. spathulata and Calycosiphonia pentamera. Therefore, we aim at providing evidence for the delimitation of the new taxa and by publishing
them formally in accordance with the *International Code of Nomenclature* (ICN) [25], we hope to draw attention to their conservation. Furthermore, high-resolution LC-UV-MS/MS seed analyses of the new species was carried out in order to contribute knowledge of the seed chemistry of the genera in the coffee tribe, the chemotaxonomic significance of the chemical constituents, and their distribution across the group. While the chemistry of the genus *Coffea* L. has been extensively investigated [26–28], there is a lack of knowledge about the chemistry of other genera, with the exception of species currently placed in *Diplospora* DC. (previously assigned to *Tricalysia* A.Rich ex DC) [27, 29–32].

### Material and methods

#### Ethical statement

This study is based mainly on herbarium specimens and field observations made in Cameroon during a series of botanical surveys beginning in 1991. These surveys were mainly led by the first author. So far, they have resulted in 52,450 specimens being studied at K and YA, of which 37,850 were newly collected. Data are stored in the Kew-Cameroon specimen Access database (Gosline, p. 11 in Cheek et al. 2004). The top set of specimens was initially deposited at SCA, and later at YA, and duplicates sent to K. The fieldwork was approved by the Institutional Review Board of the Royal Botanic Gardens, Kew, named the Overseas Fieldwork Committee (OFC), under permit number 807. The most recent official invitation to carry out research on the flora and vegetation of Cameroon has reference number 050/IRAD/DG/CRRA-NK/SSRB-HN/09/2016 and it was issued under the terms of the 5-year Memorandum of Collaboration between the Institute for Research in Agricultural Development (IRAD)-Herbier National du Cameroun and the Royal Botanic Gardens, Kew, signed 5th Sept 2014.

#### Taxon sampling and study area

Field collections were undertaken in the Mt Kupe area (SW Region, Cameroon) between June 1995 and October 2009, and the Ebo area (Littoral Region, Cameroon) between 2007 and 2015. Herbarium material and/or digitised specimens of *Argocoffeopsis*, *Belonophora* Hook.f., *Calycosiphonia*, *Coffea*, *Diplospora*, *Discospermum* Dalzell, *Empogona* Hook.f., *Sericantha* Robbr., *Tricalysia* and *Xantonnea* Pierre ex Pit. was consulted at BR, BRLU, K, MO, P, SCA, US, WAG and YA [33]. All specimens cited have been seen, unless otherwise indicated. Total sampling for the molecular phylogenetic study comprises 38 accessions, including 2/3 of *Calycosiphonia* species (DNA could not be extracted from material of *C. pentamera* and, therefore, its placement was inferred from morphological characters only), 6/10 of *Argocoffeopsis* species, and two species new to science. The samples with species names, voucher information, origin, and GenBank accession numbers, are given in S1 Appendix.

#### Phylogenetic analyses

In this study, previously published sequence data were used [21, 34] and supplemented with new sequences from selected plastid regions (*accD-psa1* intergenic spacer (IGS), *rpl16* intron, and *trnL-F* intron & *trnL-F* IGS) (S1 Appendix). The DNA extraction protocol and material and methods for amplification and sequencing followed [21].

Sequences were assembled and edited in Geneious R8 (http://www.geneious.com) [35], aligned using MAFFT 7 [36, 37], and subsequently, alignments were adjusted manually in PhyDE 0.9971 [38]. The concatenated alignment used in the phylogenetic analyses is provided in S1 File.
The matrices of the three chloroplast regions were concatenated for the downstream analyses. PartitionFinder 2.1.1 [39] was used to determine an appropriate data-partitioning scheme from potential partitions that were defined a priori (each marker treated as a separate partition), as well as the best-fitting model of molecular evolution for each partition, using the Bayesian Information Criterion. The optimal data-partitioning scheme was $\text{accD-\text{psa1}} + \text{rpl16}$ and $\text{trnL-F}$, and the GTR+$\Gamma$ (invgamma) model of sequence evolution was determined to be the best-fitting model for the $\text{accD-\text{psa1}} + \text{rpl16}$ partition, while the GTR+$\Gamma$ (gamma) model of sequence evolution was determined to be the best-fitting model for the $\text{trnL-F}$ partition in the concatenated data set.

Maximum likelihood (ML) analyses of the optimally partitioned data were performed using RAxML 8.2.10 [40]. The search for an optimal ML tree was combined with a rapid bootstrap analysis of 1000 replicates. Partitioned analyses were conducted using Bayesian Inference (BI) in MrBayes 3.2.6 [41]. The parameters of each of the partitions were the same as in the ML analysis. Rate heterogeneity, base frequencies, and substitution rates across partitions were unlinked. The analysis was allowed to run for 100 million generations across four independent runs with four chains each, sampling every 10000 generations. Convergence, associated likelihood values, effective sample size (ESS) values and burn-in values of the different runs were verified with Tracer 1.5 [42]. The first 25% of the trees from all runs were excluded as burn-in before making a majority-rule consensus of the 30000 posterior distribution trees using the “sumt” function. All phylogenetic analyses were run using the CIPRES portal (http://www.phylo.org/) [43]. Trees were drawn using TreeGraph2 [44].

**Morphology, palynology, distribution and conservation**

Measurements, colours and other details given in the descriptions of the new species *Kupeantha ebo* and *K. kupensis* are based on living material, spirit, and herbarium specimens, and data and photographs derived from field notes. Pollen samples were collected only from *Kupeantha kupensis* (Cheek 7882 K). Whole, unacetolysed anthers were placed on a stub using double-sided tape and sputter-coated with platinum in a Quorom Q150T coater for 30 seconds and examined in a Hitatchi 54700 scanning electron microscope at an acceleration voltage of 4kV.

The conservation status was assessed using the IUCN Red List Category criteria [45]. The distribution of the species was mapped using SimpleMappr [46].

**Nomenclature**

The new taxon names generated as part of this study satisfy the requirements of the International Code of Nomenclature for algae, fungi, and plants, and are hereby effectively published. In addition, they have been submitted to The International Plant Names Index (IPNI), from where they will be made available to the Global Names Index (http://gni.globalnames.org/). The IPNI LSIDs will resolve and the associated information viewed through any standard web browser by appending the LSID contained in this publication to the prefix http://ipni.org/.

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**Phytochemical analysis**

The extracts were prepared by extracting ground seed material from the newly described species *Kupeantha kupensis* (one accession) and *Kupeantha ebo* (two accessions). There was insufficient or no material available for the other species. Extracts were made in 80% aqueous methanol (100 mg/ml) for 24 h, prior to centrifugation. The supernatants were then subjected...
to LC–UV–MS/MS analysis. Analyses were performed on a Thermo Scientific system consisting of an ‘Accela’ U-HPLC unit with a photodiode array detector and an ‘LTQ Orbitrap XL’ mass spectrometer fitted with an electrospray source (Thermo Scientific, Waltham, MA, USA). Chromatography was performed on 5 μl sample injections onto a 150 mm x 3 mm, 3 μm Luna C-18 column (Phenomenex, Torrance, CA, USA) using the following 400μl/min mobile phase gradient of H₂O/CH₃OH/CH₃CN +1% HCOOH: 90:0:10 (0 min), 90:0:10 (5 min), 0:90:10 (60 min), 0:90:10 (65 min), 90:0:10 (67 min), 90:0:10 (70 min) followed by return to start conditions and equilibration in start conditions for 5 min before the next injection. The ESI source was operated with polarity switching and the mass spectrometer was set to record high resolution (30 k resolution) MS1 spectra (m/z 125–2000) in positive mode using the orbitrap and low resolution MS1 spectra (m/z 125–2000) in negative mode and data dependent MS2 and MS3 spectra in both modes using the linear ion trap. Detected compounds were assigned by comparison of accurate mass data (based on ppm), and by available MS/MS data, with reference to the published compound assignment system [47] and with supportive UV spectra.

Results
Phylogenetic analyses
The concatenated ML and BI analyses did not generate significantly different topologies, therefore, we present the relationships shown in the 50% majority consensus multiple-locus BI tree, with the associated PP values and the bootstrap values of the multiple-locus ML tree (Fig 1). Only BS values above 70% and posterior probabilities (PP) above 0.75 are shown. The best scoring ML tree is available for comparison in S1 Fig.

Tribe Coffeeae is recovered as monophyletic, with the monophyletic genus Coffea strongly supported as sister to the rest of the tribe (PP = 1, BS = 100). The rest of the Coffeeae is represented by two main clades. The first of these, the Argocoffeopsis clade (PP = 1, BS = 99) indicated in grey (Fig 1), includes the genus Argocoffeopsis (PP = 1, BS = 86), which becomes monophyletic after exclusion of Argocoffeopsis fosimondi and A. spathulata, the monophyletic genus Calycosiphonia (PP = 1, BS = 87), and a monophyletic group (PP = 0.99, BS = 92) here named Kupeantha comprising Argocoffeopsis fosimondi and A. spathulata in addition to two new species described in this study. The placement of Calycosiphonia pentamera in Kupeantha was inferred based only on its morphology because DNA could not be extracted.

The other main clade includes the rest of the generic diversity of the tribe. Included in this study are members of Sericanthe, Diplospora, Empogona, Discospermum, Xantonnea, Belonophora, and Tricalysia.

Morphology and palynology
Characters separating Argocoffeopsis, Calycosiphonia and Kupeantha gen. nov. are provided in Table 1. Characters separating Argocoffeopsis fosimondi, A. spathulata, Calycosiphonia pentamera, and the new species are documented in Table 2. The descriptions in the taxonomic treatment largely follow [48].

The pollen grains in Cheek 7882 (Fig 2) are spheroidal, 19–20 μm in diameter, tricolporate, with an apocolpium of 5–6 μm diameter (Fig 2A). The mesocolpium sculpturing is verruculately-reticulate (Fig 2B), the projecting elements rounded, about 400 nm wide. Reticulum units are about 800 nm in diameter. The apocolpium exine sculpturing grades to porate (Fig 2C). The colpal membrane is densely verruculate (Fig 2D) and the pore about 3 μm in diameter.
Phytochemical analysis

LC–UV–MS/MS seed analysis (Table 3) revealed the detection of a range of hydroxycinnamic acid derivatives, including caffeoyl-, coumaroyl- and feruloyl-quinic acids, in all three extracts, one of Kupeantha kupensis (Cheek 7882), and two of Kupeantha ebo (Alvarez 11, Fenton 134), which were assigned from their observed [M+H]^+ ions, with supportive UV spectra and MS/MS interpretation (Table 3). The main hydroxycinnamic acid detected in the positive ionisation mode in the three extracts with m/z 339.1070 eluted at the retention time (Rt) of 11.0 min was assigned as 5-O-coumaroylquinic acid. Amino acids were also detected in all three extracts and were assigned from their observed [M+H]^+ ions and supportive MS/MS interpretation as asparagine, phenylalanine and tryptophan, or their isomers (Table 3).

In the present study, a compound eluting at Rt 15.5 min with m/z 497.2737 was detected in both new species and was assigned with the molecular formula C_{26}H_{40}O_{9}, determined from the [M+H]^+ ion, which is that of the diterpenoid glucoside, villanovane I. Also, detected in all of the extracts analysed were compounds eluting at Rt 24.4 min and 38.8 min, with m/z 495.2575 and 516.3161, respectively. These compounds were assigned with the molecular formulae from their observed [M+H]^+ ions as C_{26}H_{38}O_{9}, which is that of tricalysioside B or C, and as C_{26}H_{42}O_{9}, which is that of tricalysioside P. Other compounds assigned as ent-kaurane diterpenoids or their glycosides were only detected in the seed extract of Cheek 7882. These included a compound eluting at Rt 19.6 min with m/z 333.2059 that was assigned with the
molecular formula C\textsubscript{20}H\textsubscript{28}O\textsubscript{4}, determined from the observed [M+H]\textsuperscript{+} ion, which is the molecular formula of tricalysiolide A. Also detected in the seed extract of Cheek 7882 were compounds eluting at 21.6 min and 23.1 min, with m/z 657.3110 and 349.2008, respectively. These compounds were assigned with the molecular formulae C\textsubscript{32}H\textsubscript{48}O\textsubscript{14} and C\textsubscript{20}H\textsubscript{28}O\textsubscript{5} from their observed [M+H]\textsuperscript{+} ions, which are the molecular formulae of tricalysioside E, and of tricalysiolide B or G (or tricalysin D), respectively.

### Discussion

#### Relationships in tribe Coffeeae

The relationships recovered in tribe Coffeeae (Fig 1) largely match those obtained in previous studies [21, 34], with Coffea sister to the rest of the tribe, but with little support for the relationships among most genera. [34] removed the species of Empogona from Tricalysia since they are more closely related to the genus Diplospora. This relationship is confirmed here (Fig 1). As in [21], we find a possible sister relationship between Discospermum and Xantonnea, although only a single taxon of each was sampled and therefore, this relationship needs further investigation. According to our results (Fig 1), species previously included in Argocoffeopsis and Calycosiphonia, in addition to two new taxa, form a well-supported monophyletic group. This clade is here indicated as the Argocoffeopsis clade, which is the generic name with priority.
Generic delimitation of *Argocoffeopsis* and *Calycosiphonia*

Within the *Argocoffeopsis* clade, three well-supported subclades are recovered: (1) *Calycosiphonia* (excluding *C. pentamera* due to unavailability of DNA, but transferred to *Kupeantha* based on morphology), (2) *Argocoffeopsis* (excluding *A. fosimondi* and *A. spathulata*), and (3) *Kupeantha*, a clade including the above excluded species, in addition to the newly described *K. kupensis* and *K. ebo* (Fig 1). These three subclades are supported by numerous morphological characters (Table 1).

**Table 1.**

| Habit and height (m) | Leaf-blade dimensions (cm) | Leaf acumen | Secondary nerves | Petiole length (mm) | Lower calyculus dimension (mm) | Upper calyculus dimension (mm) | Corolla tube length (mm) | Corolla lobe length (mm) | Anther apex & length (mm) | Fruit shape, size (mm) & apex |
|---------------------|---------------------------|-------------|------------------|-------------------|--------------------------------|-------------------------------|-------------------------|-------------------------|--------------------------|-----------------------------|
| **K. ebo**           | 5                         | 18–22 x (3–)5.5–7.2 | Acuminate to caudate (long cm) (0.9–)1.38–1.6 – 2.3 | 6–8, with intramarginal hooped nerves | 1.3–1.9 x 1.9–2.5 (l x w) | 1.85–2.3 x 4.2–4.5 | Ellipsoid to Globose 1.0–1.6 x 1.1–1.5 cm | ? | ? | Ellipsoid, 22–27 x 14–17, rostrate |
| **K. fosimondi**     | Shrub (1.5–)2–3           | 16–21.5 x 4.5–9.5 | Tapering to acute apex | 6–9, intramarginal hooped nerve not apparent | 1.6–2 x 2–3.4 | 2–3 x 2.5–2.8 | Globose, c. 30 x 25, round | 8–10 | 16–20 | Apiculate, c. 9 |
| **K. kupensis**      | Small tree (2–)3–8        | 13–18(–20.5) x (5.7–)6.2–7.3 | Acumen tapering, slender, acute | 8–10, uniting near the margin forming a looping connecting nerve | c. 2 x 2.8 | 2–3 x 2.5–2.8 | Ellipsoid, c. 30 x 25, round | 10–12 | 12–17 | Apiculate, 9.2–10.3 |
| **K. pentamera**     | Shrub 0.5–1.5             | (13–)14–20 x 4–7 | Acumen tapering, slender, acute | (9–)10–13, ascending, straight to curved | 0.5–1 x 0.5–1 | 1–1.5 x 1–2 | Ellipsoid, 22–27 x 14–17, rostrate | 7–9, joining to form hooped intramarginal nerve | 0.5–1 x 0.5–1 | Apiculate, c. 2 |
| **K. spathulata**    | Shrub 0.5–2.5             | (8.5–)11–17 x 2.5–5.7 | Spatulate, apex swollen, orbicular | 7–9, joining to form hooped intramarginal nerve | 0.5–1 x 0.5–1 | 0.7–1 x 1.5–2 | Ellipsoid, 22–27 x 14–17, rostrate | 4.5–6(–10) | (2.5–)7–9 | Subacute, 2.5–3 |

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**Generic delimitation of *Kupeantha* and its species**

The morphological affinities of the new species, *K. kupensis* and *K. ebo*, shown in Table 2, appear to lie with *Calycosiphonia pentamera*, *Argocoffeopsis fosimondi* and *A. spathulata*. In all five species, distinctive supra-axillary buds are developed, and the distal internodes of the stem dry dull black, becoming abruptly white and spongy in one of the more proximal internodes. The lobes of the calyculi are minute, and never foliose, except in deviant cases [19]. The calyx tube is reduced to a minute rim shorter than the disc, and lobes are absent. These characters set this group apart from *Argocoffeopsis* and *Calycosiphonia*. Although these five species, now included in the new genus *Kupeantha*, resemble *Calycosiphonia* in being evergreen small trees or shrubs, with glabrous leaves and axillary, calyculate, 1-flowered inflorescences, they lack the locellate anthers and 7–8-merous flowers found in that genus. Instead, flowers are (5–)6-merous, with non-locellate anthers. Superficially, they seem unrelated to *Argocoffeopsis* since they appear not to bear flowers on short leafy spur shoots.

*Kupeantha kupensis* differs from all other species of *Kupeantha* in having ellipsoid-rostrate fruits (the other species lack rostrate fruits). The long (12–17 mm) corolla lobes are only
exceeded by those of *Kupeantha fosimondi* (16–20 mm) and these two species share characters not seen in *Kupeantha pentamera* and *Kupeantha spathulata*, such as large anthers 9–10 mm long (instead of anthers c. 2 mm long), and a submontane habitat (instead of a lowland habitat). The affinities of *Kupeantha ebo* are more difficult to ascribe since flowers were lacking. In shape and size of leaves and fruits it resembles most closely *K. pentamera*, but differs by lacking the finely reticulate nervation of that species (the quaternary nerves are inconspicuous), and by having 6–9 secondary nerves (not 10–13) on each side of the midrib. Keys to the genera, species descriptions, and nomenclatural changes are provided in the taxonomic treatment below.

**Phytochemical analysis**

Hydroxycinnamic acid derivatives are known constituents of the seeds of *Coffea* species, which are used to prepare the beverage coffee [49, 50], and were also detected in all of the extracts of *Kupeantha kupensis* and *Kupeantha ebo* analysed in the present study. However, whilst the purine alkaloid caffeine is widely documented to occur in *Coffea* seeds, it was not detected in any of the seeds analysed in this study, nor in other previously studied genera in the Coffeeae [32].

A compound assigned as villanovane I (or isomer) was detected in both *Kupeantha ebo* and *K. kupensis*, and is a diterpenoid glucoside previously reported to occur in *Coffea* species [27].
Additionally, compounds assigned as tricalysioside B or C, and as tricalysioside P (or their isomers) were detected. These ent-kaurane glycosides have been reported to occur in species previously placed in the genus *Tricalysia* and now included in *Diplospora* [27]. Other compounds assigned as *ent*-kaurane diterpenoids or their glycosides were only detected in *Kupeantha kupensis*. These included compounds assigned as tricalysiolide A, tricalysiolide B or G (or tricalysin D), and tricalysioside E (or their isomers). These *ent*-kaurane compounds have also been reported to occur in species now placed in *Diplospora* species, as indicated in Table 3.

In summary, the detection of hydroxycinnamic acid derivatives, amino acids and *ent*-kaurane diterpenoids, or their isomers, which also occur in *Diplospora* species, are reported here for the first time in seeds of species in the *Argocoffeopsis* clade.

### Table 3. Compounds assigned from LC-UV-MS/MS analysis of the *Kupeantha kupensis* seed extracts.

| Assigned compound (or isomer) | Retention time (min) | Molecular formula | (m/z) | Ion | K. ebo (Fenton 134) ppm | K. ebo (Alvarez 11) ppm | K. kupensis (Cheek 7882) ppm |
|-------------------------------|-----------------------|-------------------|-------|-----|------------------------|------------------------|---------------------------|
| Asparagine                    | 1.6                   | C₄H₆N₂O₃         | 133.0607 | [M + H]⁺ | 0.441                | 2.320                | 0.742                     |
| Phenylalanine                 | 3.0                   | C₆H₁₁NO₂         | 166.0859 | [M + H]⁺ | 2.259                | 2.439                | 2.921                     |
| Tryptophan                    | 5.0                   | C₁₁H₁₂N₂O₂       | 205.0967 | [M + H]⁺ | 1.044                | 1.434                | 2.019                     |
| 3-O-Coumaroylquinic acid      | 6.9                   | C₁₆H₁₈O₆         | 339.1068 | [M + H]⁺ | 0.248                | 1.427                | 1.781                     |
| Caffeoylquinic acid           | 7.4                   | C₁₀H₁₆O₆         | 355.1018 | [M + H]⁺ | 0.004                | 0.531                | 1.545                     |
| 5-O-Coumaroylquinic acid      | 11.0                  | C₁₆H₁₈O₆         | 339.1070 | [M + H]⁺ | 1.339                | 2.223                | 2.577                     |
| Feruloylquinic acid          | 12.5                  | C₁₇H₂₀O₆         | 369.1173 | [M + H]⁺ | 0.403                | 0.240                | 1.974                     |
| 4-O-Coumaroylquinic acid      | 14.0                  | C₁₆H₁₈O₆         | 339.1071 | [M + H]⁺ | 0.955                | 0.601                | 0.018                     |
| Villanova I°                  | 15.5                  | C₂₆H₄₆O₉         | 497.2737 | [M + H]⁺ | 0.682                | 0.059                | 1.728                     |
| Tricalysiolide A°             | 19.6                  | C₂₀H₂₄O₄         | 333.2059 | [M + H]⁺ | Nd                    | Nd                    | 0.408                     |
| Tricalysioside E°             | 21.6                  | C₁₂H₂₆O₁₄        | 657.3110 | [M + H]⁺ | Nd                    | Nd                    | 1.084                     |
| Tricalysiolide B° or G° or tricalysin D° | 23.1 | C₂₀H₂₄O₆        | 349.2008 | [M + H]⁺ | Nd                    | Nd                    | 0.488                     |
| Tricalysiolide B° or C°       | 24.4                  | C₂₀H₂₄O₆         | 495.2575 | [M + H]⁺ | 0.103                | 1.190                | 2.664                     |
| Tricalysioside P°             | 34.8                  | C₂₆H₄₂O₉         | 516.3161 | [M + H]⁺ | 0.946                | 0.830                | 0.345                     |
| Feruloyl hexoside             | 36.4                  | C₁₆H₃₂O₉         | 357.1178 | [M + H]⁺ | Nd                    | Nd                    | 0.500                     |

All compounds assigned by comparison of accurate mass data (based on ppm = parts per million), and by interpretation of available MS/MS and/or UV spectra. Nd: Not detected / below level of detection.

°Reported to occur in coffee (*Coffea* species) and numerous other plant species [27, 49, 50].
°Reported to occur in puer coffee beans (*Coffea arabica* 'Yunnan') [27].
°Reported to occur in *Tricalysia dubia* (Lindl.) Ohwi [26], accepted name *Diplospora dubia* (Lindl.) Masam.
°Reported to occur in *Tricalysia fruticosa* (Hemsl.) K.Schum. ex E.Pritz. [27], accepted name *Diplospora fruticosa* Hemsl.

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Taxonomic treatment

A key to the genera in the *Argocoffeopsis* clade is provided in below.

**Key to the genera of the *Argocoffeopsis* clade**

1. Climbing shrubs (usually), stems and leaves hairy (5 of the 8 species), flowers terminal, single, on short lateral stems (brachyblasts), with 2–3 pairs of more or less slightly reduced leaves; calyx lobes present ...................................................... *Argocoffeopsis*

2. Tertiary nerves ± scalariform in leaf-blades; white-pale brown epidermis (dried specimens) developing within mm of the stem apex; axillary buds concealed within stipule sheaths; anthers locellate, lacking apical connective appendage, filaments 2–3mm, calyx tube well-developed, at least 1mm long, accrescent and conspicuous in fruit ..................... *Calycosiphonia*

2. Tertiary nerves absent or inconspicuous, or (*K. pentamera*) present with quaternary nerves forming reticulate, not scalariform nervation; white-pale, corky epidermis (dried specimens) developing (1–)2 (–4) internodes below the stem apex; buds supra-axillary, inserted on stem 1 or more mm above the stipule sheath; anthers entire, not locellate, with a terminal connective appendage; filaments 0.2–0.3 mm long, calyx tube c. 0.1 mm long or less, barely detectable, inconspicuous in fruit ...................................................... *Kupeantha*

**Kupeantha** Cheek, gen. nov. [urn:lsid:ipni.org:names:60476503–2] Type: *Kupeantha kupensis* Cheek & Sonké

Differs from *Calycosiphonia* Robbr. and *Argocoffeopsis* Lebrun in the conspicuous, spike-like, supra-axillary buds, which are raised 1–4 mm above the stipule sheaths (not axillary, concealed within the stipule sheaths); epidermis of leafy stems drying black in the distal (1–)2(–4) internodes, becoming white and spongy in proximal internodes (not pale brown or grey, or, if black, only for a few mm from the most distal internode); calyx tube a minute indistinct rim above the hypanthium, shorter than the disc; calyx lobes absent (not with a distinct tube longer than disc with minute lobes or with a well-developed tube, half as long or more than ovary); the calyculi with foliar lobes minute, c. 0.2 mm long (not with foliar lobes 3.5 mm or more long, or with calyculi not formed).

Shrubs or small trees, glabrous, hermaphrodite, evergreen. Stems drying black in the distal-most (1–)2(–4) internodes, terete, changing abruptly proximally to a thicker, spongy, bright white epidermis, at length becoming, in older, non-leafy stems, pale brown and flaking-fibrous; lenticels absent. Stem apex with translucent gum from stipule colleters, as in *Coffea*; internodes even in length; axillary buds spike-like 2–4 mm long, supra-axillary, raised above the stipule sheath by several millimetres. Leaves opposite, equal in size and shape at nodes, and distichous. Leaf-blades elliptic, oblong-elliptic, rarely slightly obovate (*K. pentamera*), apex acuminate, rarely acute, base broadly acute, usually slightly asymmetric, margins entire, often slightly revolute, secondary nerves 7–13 on each side of the midrib, domatia absent, nervation usually brochidodromous, tertiary and quaternary nerves usually obscure (except in *K. pentamera*, where together they form a fine reticulum). Petiole plano-convex in transverse section, convex below, and flat above, not articulated with stem. Stipules sheathing with a triangular limb, often extending into an awn-like projection, inner surface with scattered standard collers and simple hairs. Inflorescences above the axils of leaves, opposite, sometimes in successive nodes, 1(–3) per axil, or (*K. kupensis*) 2–4 per axil, each calyculate, 1-flowered, terminal. Calyculi 2–3, or (*K. kupensis*) 3(–4), completely concealing the inflorescence axis and the base of the ovary-hypanthium, 4-lobed (2 foliar lobes and 2 stipular lobes) or (proximal calyculi)
lacking lobes, glabrous outside, but with colleters, and sometimes hairs, inside, sometimes on the margin; calyculi cup-like or shortly cylindrical, the most distal largest; foliar lobes usually ovate, non-foliose; stipular lobes, if developed, ± triangular, about as long as foliar lobes. Peduncles absent; rachis completely concealed by calyculi, <1(–2) mm long. Flowers 5–6-merous, homostylous. Calyx (hypanthium) entirely contained within the upper calyculus at anthesis, rarely with up to 1 mm of the apex exserted, shortly cylindric, glabrous; calyx limb truncate, glabrous, lacking colleters inside, 0.1–0.2 mm long, shorter than disc, barely detectable. Corolla glabrous, white; corolla lobes contorted to the left in bud; corolla tube cylindric, widening slightly at the apex, about as long as the lobes; corolla lobes oblong-elliptic, apices asymmetrically shortly acuminate. Anthers introrse, fixed at apex of the corolla tube at the base of the lobes, completely exserted, submedifixed; filaments short (except in *K. fosimondi* to 2.5 mm), inserted c. 1 mm from the base of the anther; anther sacs oblong, not locellate, base shortly bifid, apex with connective minutely apiculate. Pollen globose, tricolporate, surface vermiculate-reticulate (Fig 2). Disc subcylindrical, glabrous. Ovary bilocular, style filiform, glabrous, fleshy, stigma arms 2, separated or partly appressed together, exserted. Fruits ripening orange, yellow, globose, obovoid or ellipsoid, 12–30 mm long, sometimes shortly stipitate or rostrate, disc and calyx limb not markedly accrescent; outer surface matt, subrugulose, mesocarp leathery (becoming fleshy?); endocarp distinct, brown, glossy, vascularized, translucent, membranous to thinly papery. Seeds 1–2, plano-convex and elliptic-orbicular in outline 8–13 mm long (where 2 seeds per fruit) lacking grooves, but sometimes with the radicle emerging slightly at one end. Seed coat not detected. Endosperm waxy, translucent orange-brown, forming 98% of the volume of the seed. Embryo visible within the intact seed in transmitted light, straight, linear, running along the axis of the seed, radicle pointing downwards (inferior).

Etymology: *Kupeantha* means “flower of Kupe”.

Five species in lowland and submontane evergreen forest in Cameroon, one species extending to Rio Muni (Equatorial Guinea). A key to the species of *Kupeantha* is provided in below.

### Key to the species of *Kupeantha*

1. Fruit obovoid; acumen spatulate; petiole <1cm long.............................................*K. spathulata*
   1. Fruit globose or ellipsoid; acumen with apex acute; petiole >1cm long.................................2
2. 0.5–1m tall; quaternary nerves conspicuous, forming fine reticulation on lower surface of leaf-blade; secondary nerves 10–13 on each side of midrib; leaves drying black.....*K. pentamera*
   2. 2–5m tall; quaternary nerves absent or inconspicuous, fine reticulation absent; secondary nerves <10 on each side of midrib; leaves drying green .................................................................3
3. Fruits ellipsoid, ripening black, with a short rostrum SW Region, Mt Kupe.....*K. kupensis*
3. Fruit globose, ripening orange-red, stipe and rostrum absent or inconspicuous. SW Region, Lebialem or Littoral Region, Ebo ............................................................4
4. Fruit 25–30 mm diameter. Lebialem Highlands of S.W. Region; 1300–1400 m elev..........................*K. fosimondi*
4. Fruit 10–15 mm diameter. Ebo Highlands of Littoral Region; 770–830 m elev...............................*K. ebo*

1. **Kupeantha ebo** M. Alvarez & Cheek, [urn:lsid:ipni.org:names:60476504–2](https://doi.org/10.1371/journal.pone.0199324)

   ([Fig 3](https://doi.org/10.1371/journal.pone.0199324)).

   **Type:** CAMEROON, **Littoral Region**: Ebo Proposed National Park, near the Njuma River, near Njuma Camp, 770 m elev., 4°21’26” N, 10°15’01” E, fr., 3.x.2015, *M. Alvarez 11* (holotype: K; isotype: YA).

   **Diagnosis:** Differs from *K. pentamera* by being a small tree or shrub 2–5m high, not a shrublet 0.5–1m tall, and having fruits 10–16 mm long (not 17–25mm long); secondary nerves
**Fig 3. Kupeantha ebo.** A. fruiting branch, B detail of stipule and supra-axillary buds, C, D. detail of stipule and stipule scar at an older nodes, E. transverse section of petiole, F. nervation, abaxial surface of leaf, G. fruit, H. fruit apex showing disc, I. Calyculi, J. opened fruit showing the two endocarps with sagittal crest surrounding the seeds, K. half fruit showing the vascularised membrane (endocarp) separating the two seeds, L. diagram of median longitudinal section of a fruit, showing the exocarp (shaded area), mesocarp (white areas), endocarp, cavities (dotted areas)
6–8 (not 10–13), on each side of the midrib; tertiary nerves not conspicuous, versus conspicuous and forming a fine reticulum.

Description: Small tree or shrub, 2–5 m high. Stem dark, greyish brown to lighter reddish brown. Stem terete, glabrous, smooth to flaky and peeling off on older stems, 4–5 mm in diameter. Stipules connate, subcoriaceous, glabrous outside, puberulous inside, with gum exudate, rectangular to trapezoid, 1.5–2.8 mm long × 2.6–4.5(–6) mm wide, apex mucronate to narrowly triangular, crustaceous, 0.8–2 mm long. Leaves opposite, equal; petioles glabrous, drying darker and shinier than stems, (6–)9–13 mm long; leaf–blades drying dark green above and light green below, narrowly elliptic to oblong, glabrous above and below, cartaceous, (9–)18–22 × (3–)5.5–7.2 cm; apex tapering, acuminate to caudate (9–)13–16(–23) mm long, base attenuate; length/width ratio 3:1 to 4:1. Midrib sub-prominent above and prominent below; secondary nerves impressed above and prominent below, 6–8 on each side, ascending and curving to make a looping intramarginal nerve (brochidromous); midrib and secondary ribs drying dark brown below. Tertiary nerves conspicuous, reticulate above and weakly percurrent below. Quaternary nerves inconspicuous. Domatia absent, bacterial infections like ulcers of irregular form may appear on blades or on veins. Inflorescences on leafy branches. Flowers not seen. Inflorescences in pairs at each node, 1 per axil, supra-axillary, situated 1.5–1.8 mm above the base of petioles. Calyculi 2 to 3, sessile, cupuliform glabrous. The first calyculus (basal), may be corky, without apparent lobes, sometimes it forms part of the stem, 1.3–1.9 × 1.9–2.5 mm, the second one smooth, with 2 broadly triangular stipular lobes 2.1–2.3 × 1.8–3 mm; the third distal one cupuliform at the base, smooth, with a lenticular structure at the top, 1.8–2.3 × 4.2–4.5 mm, sinuate margin with 2 very fragile subulate lobes, c. 1.5 x 0.2 mm. Fruits berry like, ellipsoid when immature to globose when mature, 1.0–1.6 × 1.1–1.5 cm. Orange to red at maturity, glabrous, exocarp drying hard, leathery, dark brown; disc ± circular, 2.8–4 mm in diameter; calyx limbs absent. Disc exserted in young fruits, becomes ± flat in older fruits. Endocarp membrane very vascularised, separating the seeds. Seeds: 1–2, apparently without seed coat, surface ± smooth, endosperm entire cream. Where 2 seeds, the seed is plano–convex, broadly ovate in outline, c. 1 × 0.8 × 0.4 cm. Where 1 seed, ellipsoid with irregular surface, circular in cross section, c. 1.1 × 0.7 cm. Micropyle visible on seed embryo can be seen germinating prematurely in some seeds.

Distribution and habitat: Kupeantha ebo is endemic to the Ebo Proposed National Park in the Littoral Region in Cameroon. The species is found in submontane, evergreen forest, on well-drained sandy to loamy and rocky soil from 770 to 832 m in elevation. The diversity is represented by large trees of Aphanocalyx microphyllus (Harms) Wieringa, Berlina bracteosa Benth., Cynometra hankei Harms, Distemonanthus benthamianus Baill., Hymenostegia viridi-flora Mackinder & Wieringa, Plagiosiphon longitubus (Harms) J.Léonard, Pterocarpus soyauxii Taub. (all Fabaceae); Terminalia superba Engl. & Diels (Combretaceae) and Vitex lokundjensis W.Piep. (Lamiaceae).

Phenology: Flowering unknown; fruiting in October.

Conservation status: Kupeantha ebo is restricted to the Littoral Region in Cameroon and it is known from only two sites in Ebo Proposed National Park [51,52]. The forest, where this new species was found, was a logging and farming area and it currently seems to be slowly regenerating, since these activities stopped some years ago, although there are still gaps of secondary vegetation among the primary forest. Considering the Ebo Forest does not have a
protected status, the survival of this species will depend on the collaboration between locals and the scientific community. There are threats from oil palm plantations and iron ore mining, in addition to the resumption of slash and burn small-holder agriculture. Given that several botanists have spent cumulatively many months over several years collecting specimens at Ebo, and that only two specimens of this species have resulted, it is clear that *K. ebo* is genuinely infrequent and rare. Accordingly, *K. ebo* is here assessed as Critically Endangered CR B2ab(iii) [45].

Additional specimens examined: CAMEROON, Littoral Region: Ebo Proposed National Park, Yabassi, Bekongo trail. 832 m elev. 4°20'45 N, 10°24'40 E. fr. 08 Oct. 2007. *E. Fenton 134* (K).

2. *Kupeantha fosimondi* (Tchiengué & Cheek) Cheek, **comb. nov.** [urn:lsid:ipni.org:names:60476505–2] ≡ *Argocoffeopsis fosimondi* Tchiengué & Cheek, Pl. Lebialem Highlands: 54 (2010).

   Type: CAMEROON, W Bamboutos Mts, Ntoo forest at Fosimondi Forest, 1350 m elev., 24. ii.2006, *B. Tchiengué* 2597 (holotype: sheet 1/2 K000518627, sheet 2/2 K000265593; isotypes: K000265585, WAG0247275).

   Distribution: Cameroon, only known from the Lebialem Highlands (Fig 4).

3. *Kupeantha kupensis* Cheek & Sonké, **sp. nov.** [urn:lsid:ipni.org:names:60476506–2] (Figs 5 and 6).

   Type: CAMEROON, Cameroon Highlands, Mt Kupe, above Kupe Village, 1550 m elev., 4° 48’N, 9° 42’E, fl. fr., 17.xi.1995, *Cheek 7882* (holotype: K; isotypes: BR, MO, P, SCA, US, WAG, YA).

   Previously mentioned in literature as: *Calycosiphonia* sp. A [6: 369] and *Argocoffeopsis kupensis* Cheek & Sonké ined. [5: 260 & map 560E].

   Diagnosis: differs from *K. spathulata* and *K. fosimondi* by having ellipsoid and rostrate fruits (not globose, apex rounded), corolla tube 11–12 mm long (not 4–5 mm long), lobes 12–17 mm long (not 7–9 mm as in *K. spathulata*, nor 16–20 mm as in *K. fosimondi*).
Description: Shrub or small tree, (2–)3–8 m tall, glabrous; main axis brown. Stem terete, 2.5–4 mm diam., drying matt black in first and second internodes, in the third to fourth internodes the epidermis becoming white, spongy, subglossy and exfoliating as in a *Chazaliella* E. M.A. Petit & Verdc.; internodes (2.8–)5.2–6.8 cm long, apical bud with gum covering; buds supra-axillary, covered in gum, inserted 3–4.5 mm above the petiole bases. Petiole 0.7–1.7 cm long, drying black. Leaf blades opposite, equal, elliptic-oblong, rarely ovate-lanceolate, 13–18 (–20.5) × (5.7–)6.2–7.3 cm, acumen tapering, slender, acute, 1 cm long, 3–4 mm wide at base, base obtuse to acute, slightly to conspicuously asymmetric, midrib impressed above, lateral nerves 8–10 on each side of the midrib, arising at c. 50 degrees from the midrib, curving upwards and uniting near the margin to form a weak, looping, connecting nerve, tertiary and quaternary nerves sparse, reticulate, not conspicuous, domatia absent; margin entire; drying very dark green above, pale green below, with midrib nearly black; petiole flattened above, concave below, 0.7–1.7 cm long, drying black. Stipule with basal cylindrical sheath 2–4 × 4–6 mm; truncate, distal limb absent to narrowly triangular or awn-like 1–4 mm long, with central ridge absent. Inflorescences on leafy branches near nodes (2–)3–4 internodes from apex, subtended by leaves, inserted 2–4 mm above the leaf axil; fasciculate, inflorescences 1-flowered, 2–4 per axil at both axils of node; 2–4 axils fertile per stem. Calyculi 3(–4), the uppermost slightly larger, each calyculus cylindrical-campanulate, more or less 4-lobed (the uppermost: 2 foliar lobes and 2 stipular lobes) or 2-lobed (the lower calyculi, with two foliar lobes only), ± sessile, glabrous, colleters not seen; basal (1st) calyculus c. 2 × 2.8 mm, foliar lobes reduced to mucra, c. 0.7 mm long, stipular lobes absent or minute; middle (2nd) calyculus resembling the basal calyculus; upper (3rd) calyculus 2–3 × 2.5–2.8 mm, foliar lobes narrowly triangular or ligulate, 2.2 × 0.5 mm long, stipular lobes 0.75 mm long. Flowers hermaphrodite, homostylous (5– or) 6–merous, sessile. Calyx (hypanthium) entirely contained within the upper calyculus at anthesis, rarely with up to 1 mm of the apex exerted, c. 3.5 × 2.3 mm, glabrous; calyx limb truncate, glabrous, lacking colleters inside, 0.1–0.2 mm long, barely detectable. Corolla glabrous, white; corolla lobes contorted to the left in bud; corolla tube cylindrical, widening slightly in the apical 2 mm, (9–)10–12 mm long; corolla lobes oblong-elliptic, 12–17 × 3–4 mm, apices asymmetrically shortly acuminate. Anthers fixed at apex of the corolla tube at the
Fig 6. Kupeantha kupensis. A. habit, flowering branch, B. detail of stipule and node and supra-axillary buds, C. node with inflorescences, flowers in bud, D. calyculi, E. flower and calyculi, F. stamens, different views, G. gynoecium, left entire, showing disc and calyx, right, longitudinal section, H. fruits, I. fruit with a longitudinal section, showing empty locule (seed removed), J. seed, ventral surface. Scale bar: A = 4 cm; B, F, G = 7 mm; C, E, K = 1 cm; D = 4 mm; H = 2.5 mm; I = 2 mm. Drawn by Lucy T. Smith from Cheek 7882 (K).

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base of the lobes, completely exserted, submedifixed; filaments c.1 mm long, inserted c.1 mm from the base of the anther; anther sacs 9–10 mm long, base shortly bifid, apex with connective apiculate, 0.2–0.3 mm long. Disc subcylindrical, c. 0.4 × 1.8–2 mm, glabrous. Ovary bilocular, not dissected; style filiform, 9.8–1.15 cm × 0.5 mm, glabrous, fleshy, stigma arms 2, appressed together, 4–8 mm, only the apical 1–2 mm separated, exserted. Fruits black, at length ripening orange, ellipsoid, 22–27 × 14–17 mm including a 2–3 mm broad, 3–5 mm long rostrum with apex truncate, disc and calyx limb present but not markedly accrescent; outer surface matt, subrugulose, mesocarp leathery (becoming fleshy?), 0.7–1 mm thick when dried; endocarp distinct, brown, glossy, vascularized, translucent, membranous to thinly papery. Seeds 1–2, plano-convex and elliptic-orbicular in outline 10–13 × 8 × 2–3 mm (where 2 seeds per fruit) or c. 9 × 5 × 6 mm (where 1 seed per fruit), lacking grooves, but sometimes with the radicle emerging slightly at one end. Seed coat not detected. Endosperm waxy, translucent orange-brown, forming 98% of the volume of the seed. Embryo visible within the intact seed in transmitted light, straight, linear, running along the axis of the seed, c. 4 mm long.

Distribution and habitat: *Kupeantha kupensis* is restricted to the SW Region in Cameroon (Fig 4). This new species is known only from Mt Kupe. The area from which *K. kupensis* is known supports a submontane (cloud) forest with *Santiria trimera* (Oliv.) Aubrèv. (Burseraceae), *Cola verticillata* (Thonn.) Stapf ex A.Chev. (Sterculiaceae), *Garcinia smeathmannii* (Planch. & Triana) N.Robson ex Spirl. (Clusiaceae), *Zenkerella citrina* Taub. (Fabaceae-Detarioideae), *Chassalia laikomensis* Cheek, and *Pauridiantha paucinervis* (Hiern) Bremek. (both Rubiaceae), *Carapa Aubl.* spp. (Meliaceae). This evergreen forest has a closed-canopy with many epiphytes and a rich herb layer, classified as Biafran evergreen forests [53], rich in Fabaceae-Caesalpinoideae. The Kupe area is now well known as an area of high endemism in Cameroon [6]. Elevation 1100–1600(–2000?) m.

Phenology: Flowering in February, November; fruiting in January, June, July, October and November.

Etymology: Named for Mt Kupe, the only known location for the species. The survival of this species is in the hands of the Bakossi people and others who live around the mountain.

Conservation status: This assessment maintains that of Cheek [5: 260]. *Kupeantha kupensis* is only known from two sites at a single location. Many thousands of specimens have been collected at Mt Kupe [6] so the area cannot be considered undercollected. Although the Mt Kupe Integrated Ecological reserve has been created in the last 10 years, both sites for *Kupeantha kupensis* fall outside its boundary. One site is under threat of forest clearance (the type collection was made where a coffee plantation had been created inside the forest). *Kupeantha kupensis* is here assessed as Critically Endangered CR B2ab(iii) [45]. B2 indicates that the total area of occupancy is less than 10 km$^2$ (in fact the area of occupancy is 8 km$^2$ using the IUCN preferred 4 km$^2$ cells for each of the two sites known at this location), while b(iii) indicates “continuing decline inferred in area, extent and quality of habitat”. Further exploration is needed to find more locations for this tree, if they exist, however, the western slopes of Mt Kupe have been relatively intensively surveyed for plants. We hope that the people of Mt Kupe will seek to protect this beautiful species from extinction.

Note: *Kupeantha kupensis* is the latest of many narrowly endemic species recently described from Mt Kupe. Other examples are *Diospyros kupensis* Gosline [54], *Memecylon kupeanum* R. D.Stone, Ghogue & Cheek [55], *Dracaena kupensis* Mwachala, Cheek, Eb.Fisch. & Muasya [56] and *Psychotria ngollengollei* Cheek [57].

Additional specimens examined: CAMEROON, Cameroon Highlands, above Nyasoso, Max’s trail, 5 Feb 1984 (fl.), *Thomas* 3087 (K, MO n.v.); ibid., 9 Jul 1992 (young fr.), *Sunderland* 1539 (K, SCA n.v., YA n.v.); ibid., 29 Jan 1996 (fr.), *Etuge* 1664 (K, SCA n.v., WAG n.v., YA n.v.); ibid., 24 Jun 1996 (fr.), *Etuge* 2381 (YA); ibid., 24 Jun 1996 (fr.), *Etuge* 2401 (K, SCA n.v.,
4. *Kupeantha pentamera* (Sonké & Robbr.) Cheek, **comb. nov.** [urn:lsid:ipni.org:names:60476507–2] ≡ *Calycosiphonia pentamera* Sonké & Robbr., Nordic J. Bot. 25: 276 (2007 publ. 2008).

Type: CAMEROON, 3 km northwest of Mbikilikiki, (03˚11'24.9"N, 10˚31'35.3"E), 21 Jan 2006 (fl.), *B. Sonké* & *K. M.-N. Djuikouo* 4320 (holotype: BR; isotypes: BR including flowers in spirit, BRLU, K, MO, WAG, YA).

Distribution: Western Cameroon, Equatorial Guinea, widespread in the evergreen lowland rainforests of eastern Cameroon, extending into Rio Muni. (Fig 4).

5. *Kupeantha spathulata* (A.P.Davis & Sonké) Cheek, **comb. nov.** [urn:lsid:ipni.org:names:60476508–2] ≡ *Argocoffeopsis spathulata* A.P.Davis & Sonké, Blumea 53: 528 (2008).

Type: CAMEROON, Mvillé, 3 km NNW Ngovayang (03˚13'41"N, 10˚34'52"E), 30 Nov 2005, *B. Sonké* & *K. M.-N. Djuikouo* 4188 (holotype: K; isotype: BR, BRLU, MO, WAG, YA).

Distribution: Cameroon; relatively widespread in the lowland evergreen forests of eastern Cameroon. (Fig 4).

**Supporting information**

S1 Appendix. Voucher information and GenBank accession numbers for taxa used in this study. A dash indicates that the region was not sampled. Voucher specimens are deposited in the following herbaria: BR = National Botanic Garden of Belgium, Meise; DSM = University of Dar es Salaam; K = Royal Botanic Gardens, Kew; TAN = Parc de Tsimbazaza, Antananarivo; YA = National Herbarium of Cameroon, Yaoundé.

(DOC)

S1 File. Concatenated sequence alignment, with sequences from the plastid markers *rpl16*, *trnL-F* and *accD- psa1*.

(FAS)

S1 Fig. Best scoring ML tree for the concatenated dataset, with bootstrap values equal or higher than 70% displayed above the branches.

(TIF)

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