Mummified fruits of *Canarium* from the upper Pleistocene of South China

*Canarium maomingense* sp. nov. described from the upper Pleistocene of South China

Micro-CT shows ovule abortion in *Canarium* existed at least since the late Pleistocene

Fossil assemblage suggests subtropical evergreen broad-leaved and mixed forests

Damage traces indicate possible interaction among plants, animals, and fungi

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Article

Mummified fruits of Canarium from the upper Pleistocene of South China

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SUMMARY

Canarium L. contains approximately 78 species distributed in low to middle altitudes of the Paleotropics and northern Australia. Canarium fruit fossils are known mainly from Paleogene to Neogene of North America, Africa, and Eurasia. Here, we described a new species Canarium maomingense sp. nov. from the upper Pleistocene of the Maoming Basin, Guangdong, South China. Similarly to extant Canarium species, each of three locules of C. maomingense possesses two ovules, but only one or two of six ovules develop into a seed, indicating that the ovules undeveloped into seeds in Canarium species have existed at least since the late Pleistocene. The natural habitats of extant relatives and associated fossil plants suggest subtropical evergreen broad-leaved and mixed forests in the late Pleistocene of this region. Some special damage traces are observed on pyrene surfaces, indicating possible plant interactions with animals and fungi.

INTRODUCTION

The family Burseraceae comprises ca. 700 species in 18 genera and is divided into three tribes: Canarieae, Proteae, and Bursereae. Canarium is the largest genus of the tribe Canarieae. It contains ca. 78 species that are mainly medium- to large-sized canopy trees, distributed in low- to middle-altitude areas of the Paleotropics and northern Australia. Seven Canarium species are native to the subtropical and tropical regions of southern China. Formally, the fruit of Canarium is a syncarpous pyrenarium of the Butia type with a trilocular pyrene composed of an inner mesocarp zone and the endocarp. Each pyrene locule contains two hanging campilotropous ovules. When the fruit matures, its peripheral soft tissues are eaten by animals or rot, and thus the hard pyrene is released. During seed germination, the pyrence surface cracks along distinct grooves bordering the lid (the germination valve) along the angle-ribs and the lid is pushed out from the inside by the growing seedling.

Currently known fossil remains of Canarium are represented by leaves, fruits, pollen, and wood. The fossil fruits have mainly been recorded from the lower and middle Eocene of North America, the Oligocene of Asia, Europe, and Africa, and the Miocene of Asia. Leenhouts supposed that the East Malaysian–northern Australia region was the most probable center of origin of Canarium, whereas analysis of the fossil records of this genus and its extant distribution suggests that Canarium originated in North America in early Eocene and migrated eastward to Europe during the middle Eocene.

Canarium is an economically important plant having medicinal and food uses and hence has a long history of cultivation and consumption in Guangdong Province, other provinces of China and in some South-East Asian countries. However, the fossil records of this genus have not been reported from Guangdong Province until now. Thus, the well-preserved mummified Canarium fossil fruits, which have been recently found in the upper Pleistocene of Maoming Basin (Figure 1), are the first megafossil record of this genus in Guangdong Province. Here, we provide a morphological, anatomical, and taxonomic study of these fossils and apply the approach of the nearest living relative (NLR) method to analyze the paleoclimate in the region. The research is aimed at revealing the phytogeographical and phylogenetic significance of the new fossil records of Canarium, as well as focusing on clarifying their probable interactions with animals and fungi.

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Figure 1. Location and lithology of the fossil site (modified from Huang et al.21)
(A) Geographic map showing the location of the Zhenjiang opencast mine within the Maoming Basin in Guangdong Province, South China (base map from d-maps: https://d-maps.com).
(B) View of the plant-bearing section. Red dashed line indicates the unconformity between the upper Pleistocene deposits and the upper Eocene Huangniuling Formation.
(C) Lithological column of the studied succession showing the position of the Canarium fossil fruits.
RESULTS
Systematic paleontology
Order
Sapindales Juss. ex Bercht. et J. Presl (1820).

Family
Burseraceae Kunth (1824).

Genus
Canarium Linnaeus (1759).

Species
C. maomingense H.-L.-L. Xiang et J.-H. Jin sp. nov. (Figure 3A–3J, 4A–4G, 4J, 4K, 4L, 4O, 5, 8–10).

Etymology
The specific epithet “maomingense” refers to the Maoming Basin from where the specimens were collected.

Holotype
MMRH–192 (Figure 3A–3D, 4J, 4K, and 4O).

Specimens examined
MMRH–192; MMRH–193; MMRH–194; MMRH–195; MMRH–196; MMRH–197; MMRH–198; MMRH–207; MMRH–208; MMRH–209; MMRH–210; MMRH–211; MMRH–212; MMRH–213; MMRH–245; MMRH–246; MMRH–247 (total of 17).

Locality and age
Opencast mine near Zhenjiang Town, Maoming City, Guangdong Province; late Pleistocene.

Repository
The Museum of Biology of Sun Yat-sen University, Guangzhou, China.

Diagnosis
Pyrene spindle- to ovoid-shaped, with acute to attenuate apex and acute base, rounded (to hexagonal) in cross-section, trilocular, with two ovules per locule. Each locule with an outwardly projecting germination valve (lid), triangular at the top, with undulated or smooth surface and notably convex or slightly convex median longitudinal keel; all lids bordered along the angle-ribs by distinct grooves. Each of three angle-ribs (the septas comprising the three-rib central column) with an opening near the pyrene apex—the entrance into the septal vascular bundle channel. Pyrene wall composed of a thick mesocarp inner zone and the endocarp. The mesocarp inner zone consists of numerous layers of sclerenchyma and single innermost layer of thin-walled cells; the endocarp composed by single layer of obliquely radially elongated cells with wavy walls. Seeds one or two, the locules lacking the seed slightly undeveloped.

Description
The mummified fossils represent the pyrenes of syncarpous trilocular drupaceous fruits (Figure 2). The pyrenes are 24.78–33.38 mm long (mean 27.74 mm) and 9.52–14.35 mm wide (mean 12.66 mm), spindle- (Figures 3A, 3E, and 3F) to ovoid-shaped (Figure 3G), rounded (Figures 3B, 3C, 3I, and 3J) to hexagonal (Figures 3D and 3H) in cross-section; the apex is acute to attenuate (Figures 3A, 3E, 3F, and 3G), the base is acute and marked by a small attachment scar (Figure 3H). The pyrenes have three locules separated by a three-ribbed central column (Figures 2, 3D, 3I, and 3J). Each locule of the ovary has two ovules and axial placentation (Figures 4E, 4F, 4G, and 4L). Only one of the two ovules (or none) in each locule develops into a seed, and often only one per fruit, other ovules remain undeveloped into seeds (Figures 3D, 3I, 3J, 4J, 4K, and 4O). The locules with undeveloped seeds are slightly or significantly reduced
The ovular vascular bundle scar is observed on the placenta (Figures 4G and 4L). Each locule has an outwardly projecting lid, triangular at the top (Figures 3A, 3E–3G, 4B, and 4D), which is 15.38–22.10 mm long (mean 17.95 mm), 6.58–10.73 mm wide (mean 9.34 mm), and 1.48–3.03 mm thick (mean 2.39 mm), it extends for 53.80%–73.37% of the pyrene length (mean 64.71%) (Figure 4B). The outer surface of the lids is undulated (Figures 3A, 3E, and 3F) or smooth (Figure 3G), with a median keel, which is notably convex (Figures 3A and 3E) or slightly convex (Figure 3G). The septal ribs between the locules are usually distinctly visible on the surface as the angle-ribs (Figures 3B–3D), which are 1.41–2.38 mm wide (mean 1.93 mm), and each of the three angle-ribs have a septal vascular bundle channel on the surface near the pyrene apex (Figures 3B, 3C, and 4J). The germination valves are bordered along the angle-ribs by distinct grooves. During germination, lids crack naturally along the grooves and become detached from the pyrene (Figure 3F). The seed coat (Figure 4A) and detached endocarp (Figures 4C–4F, 5C, and 5D) are observed in some specimens. The inner zone of the mesocarp is composed of numerous layers of mainly radially elongated sclerenchymatous cells with intercellular pits in the walls, which appear like dense hemispherical ornamentation (Figures 5A and 5B). These sclerenchymatous cells are rather polygonal in outline on the periphery of the pyrene and mainly isodiametric toward the inside (Figures 5A and 5B). The innermost zone of the mesocarp is represented by single layer of rather thin-walled cells (Figure 5E). Along this layer, which becomes relatively easily destroyed in fossil pyrenes, the endocarps can be separated from the hard mesocarp tissue (Figures 5C and 5D). The endocarp is composed by radially or obliquely radially elongated cells in relation to the center of the locule (Figures 5C–5F). The cells of the endocarp have thickened and considerably wavy walls with numerous pits and large pores, and are comparable with hydrocites in structure. The seed coats fuse with the perispem and 1- to 2-layered endosperm, which contains idioblast cells visible on inner surfaces contacting the cotyledons (Figures 5C–5E).

DISCUSSION

Comparison of C. maomingense with extant species of Canarium

Fruits are considered to be conservative plant organs with a stable anatomical structure that is often used as an important taxonomic character. Canarium is distinguished from other genera in Burseraceae by its drupaceous fruit (pyrenarium) with three locules, and two hanging campilotropous ovules per locule.1,6,7 Usually only one or two ovules (and in all cases only one per locule) develop into seeds.

Extant Canarium includes three taxonomic sections: Canarium, Canariellum, and Pimela.5 The pyrenes of C. maomingense sufficiently differ from those of the sect. Canarium (pyrenes are usually 5–7 cm long,
smooth, sometimes with angle- and median ribs); thus, we compared fossil pyrenes with those of the sections Canariellum and Pimela (Table S1). Pyrenes of the new fossil species are morphologically more similar with those of some extant Canarium species of sect. Pimela (Canarium album (Lour.) Raeuschel, C. bengalense Roxb., C. tonkinense Engl., C. pimela Leenh., C. parvum Leenh., C. subulatum Guillaumin, and C. strictum Roxb.). Among these extant species, C. album (Figures 3K–3P, 4H, 4I, 4M, and 4N) is...
Figure 4. Internal structure of pyrenes of fossil *Canarium maomingense* sp. nov. (A–G, J–L, O) and extant *Canarium album* (Lour.) Raeuschel (H, I, M, and N)

(A–D) MMRH–193. (A) Lateral view showing the endosperm inner surface (arrow). (B) The lid, the locule-faced surface. (C) Exfoliated endocarp. (D) Lateral view showing the endocarp (arrow).

(E) Lateral view showing the endocarp lining the locule (green arrow) and two placentas (blue arrows), MMRH–197.

(F) Enlarged view of two placentas in Figure 4E.

(G) Lateral view showing two placentas and ovular vascular bundle scars (arrows) (MMRH–213).

(H) Lateral view showing two placentas (arrows).

(I) Lateral view showing the endosperm inner surface.

(J) Longitudinal section showing the central vascular bundle channel (blue arrow), which merges at the top with the septal vascular bundle channel (yellow arrow), and the attachment of a normally developed ovule/seed (green arrow), holotype, MMRH–192, micro-CT.
most similar to the new fossil species in terms of the pyrene shape (spindle to ovoidal) and proportions as well as in anatomical features. The mean thickness of the lid in *C. maomingense* and *C. album* is basically the same. However, the diameter of the *C. maomingense* pyrene (9.52–14.35 mm) is smaller than in *C. album* (15–20 mm), and some of the studied fossil specimens have a smooth lid and median keel that is markedly convex, but the lid of extant *C. album* is undulated, and its median keel is slightly convex. The inner mesocarp sclereids that compose the outer part of the pyrenes are smaller and more isodiametric in extant species (50–90 × 20–30 μm in *C. album* vs. 80–140 × 20–30 μm in *C. maomingense*), and the endocarp thickness in *C. album* (up to 200–220 μm) noticeably exceeds that in the fossil species (70–90 μm) (Figures 6A, 6B, 6E, and 6F). The innermost layer of the mesocarp in *C. album* consists of thin-walled cells that are mostly flattened in a radial direction and contain dense substances of an unknown nature in the lumina with bagel-shaped outlines (Figure 6D). Along this thin-walled cell layer (Figures 6C and 6D), the separation of the sclerenchymatous inner mesocarp zone and the endocarp occurs in desiscent and partly decayed pyrenes, and the endocarps that are occasionally fused with seed coats, perisperm, and endosperm exfoliate out of the locale in *Canarium* species.

The new fossil species is also similar to *C. bengalense* in the shape of the pyrene cross-section and the thickness of the lid. However, both the length (24.78–33.38 mm) and width (9.52–14.35 mm) of the studied fossil specimens are smaller in comparison with those of *C. bengalense* (length 45–50 mm, width 17.5–20 mm). The large pyrene of the fruit of *C. bengalense* is spindle shaped with a triangular to circular cross-section; the smooth lid has unobvious median keel. The angle-ribs of *C. bengalense* often protrude over the pyrene surface up to 5 mm.

The pyrenes of *C. maomingense* and *C. tonkinense* are similar in having an undulated to smooth lid. However, the lid median keel in the fossil species generally protrudes markedly from the surface, whereas in *C. tonkinense* the keel is not seen. Nevertheless, in *C. tonkinense*, the pyrenes are elliptoidal in shape, with obtuse spines and bases, and are circular-triangular in cross-section, unlike in the Maoming fossils. The pyrene diameter (on average 12.7 mm) in *C. maomingense* is smaller than that in *C. tonkinense*.

Pyrenes of the new fossil species differ from those of *C. pimela*, which have a narrowly ovoidal shape, a nearly circular cross-section with a diameter of 17.5–20 mm, a smooth lid surface (3 mm thick), and an inconspicuous lid median keel.

In contrast to *C. maomingense*, pyrenes of *C. parvum* are spindle shaped with thin lids (1 mm vs. 1.48–3.03 mm in *C. maomingense*). In addition, the surfaces of *C. parvum* lids are smooth with obscure median keels, and the cross-sections are triangular, differing from the Maoming fossils.

The pyrene of *C. subulatum* is oval in outline, the lid surface is smooth with an obscure median keel, and the cross-section is circular-triangular, up to 15–20 mm in diameter, and so differing from the much smaller rounded-hexagonal cross-sections of the pyrenes of *C. maomingense*.

The pyrene lengths and diameters of *C. maomingense* are smaller than those of *C. strictum* (32.5–45 mm long and 17.5–22.5 mm wide), and the pyrene shape of the extant species is obovoid to ellipsoid, nearly circular to rounded triangular in cross-section, and blunt at the base and apex. The lid median keel of pyrenes of the Maoming fossil species and *C. strictum* are prominent or obscure, but the lid surface of the fossil specimens is smooth to undulated, whereas in *C. strictum* it is smooth.

Thus, the *Canarium* fossil fruit pyrenes studied in this paper are different in various ways from the extant Chinese and South-East Asian species mentioned above, and cannot be assigned to anyone of them.
Comparison of *C. maomingense* with fossil representatives of Burseraceae

Several extinct taxa are supposed to be closely related to *Canarium* (Table S2). One of them is *Tricarpellites communis* Bowerbank from the lower Eocene London Clay Formation, United Kingdom;\(^1\) *C. maomingense* is similar to *T. communis* in having syncarpous fruits with three locules along the main axis and axial placentation. However, the pyrenes of *T. communis* are sub-ovoid in shape, triangular in cross-section, and with a smooth lid surface. The lengths and widths of the pyrenes, and the widths of the lids (6.58–10.73 mm) of the Maoming specimens are...
larger than those of *T. communis* (7.5–12.5 mm × 4.25–7.5 mm in size, with 4.25–4.5 mm lid width). The lid length in *C. maomingense* is about 2/3 of the length of the pyrene, whereas in *T. communis* it is about 1/3 to 1/2 of the pyrene length. The specimens from the London Clay have tendency to split into separate pyrenes, possibly, the consequence of partial decay before fossilization.

Agarwal and Ambwani described carbonaceous angiosperm fruit fossils from the Miocene of Sindhudurg District, Maharashtra, India. The detailed morphological characteristics of these fruits, which they assigned to the species *Canariocarpus ratnagiriensis* Agarwal et Ambwani, indicate that they are related to the genus *Canarium*. The Maoming and Indian species are similar in the shape of their trilocular pyrenes; however, those of *C. ratnagiriensis* are larger (about 30–45 mm long and 13–20 mm wide), and the width of the lid is 7–20 mm. Moreover, some of our specimens have prominent median keels, whereas the median keels of *C. ratnagiriensis* are not prominent.
Pyrenes of Canarium parkii Tiffney from the lower Eocene of Virginia, USA\textsuperscript{13} were recently scanned with micro-CT.\textsuperscript{12} C. maomingense and C. parkii differ in their complex of characters. The pyrenes of C. parkii are much smaller (5.8–9.0 mm long and 2.5–5.6 mm wide), teardrop shaped, triangular in cross-section, and with an indistinct median keel on a smooth lid surface. Moreover, the point of vascular attachment in C. parkii is approximately in the middle of the seed length, whereas in C. maomingense it is located about 1/5 of the seed length from its apex.

Canarium sp. from the middle Eocene Wagon Bed Formation of Wyoming, USA\textsuperscript{14} is comparable to C. maomingense in size being 27 mm long. However, the fossil specimen from Wyoming is preserved as a silica cast, lacking anatomical details for further comparison.\textsuperscript{12}

Unlike in the species described here, pyrenes of Canarium sp. from the middle Eocene Messel Formation, Germany\textsuperscript{15} are elliptical to oval in lateral view, with obtuse apices and bases, 12 mm long and 8 mm wide, triangular in cross-section, and with quite narrow grooves between the angle-ribs and smooth lids.

Pyrenes of C. maomingense are close in size to those of Canarium chandleri Gregor et Goth (18–28 mm long, 13–15 mm in diameter) from the Oligocene of the Czech Republic.\textsuperscript{16} However, pyrenes of C. chandleri are nearly oval in lateral view, with a triangular cross-section and lids that extend for about 3/4 of the total pyrene length, in contrast to about 2/3 in C. maomingense.

Recently, two fossil species of Canarium were described from South and East China. Han et al.\textsuperscript{12} reported the Canarium guangxiense Han et Manchester from the Yongning Formation (upper Oligocene) of Nanjing City, the Erzitang Formation (Miocene) of Guiping City and the Foluo Formation (upper Miocene) of Beihai City, Guangxi Province. The original spelling of a specific epithet guangxiensis is modified here to correct the terminal ending of the specific epithet according to the grammatical gender of the generic name Canarium (lat. Neutrum) in conformity with Art. 23 of the International Code of Nomenclature for algae, fungi, and plants.\textsuperscript{24} The pyrenes of C. maomingense and C. guangxiense are similar in size and shape, although pyrenes of C. guangxiense often tend to be obovoid with an attenuate apex. In contrast to C. maomingense, the pyrenes of C. guangxiense are three- and four-locular, sometimes with triangular cross-sections. The pyrenes of C. maomingense are more elongated (length/width ratio 2.7 in average) as compared with those of C. guangxiense (length/width ratio 2.4 in average). Furthermore, in C. maomingense, the lid extends up to 2/3 of the pyrene length, whereas in C. guangxiense it reaches 4/5 of the pyrene length.

Canarium haominiae Yin, Wu, Wang et Shi from the middle Miocene Fotan Group in Fujian Province, China, distinctly differs from C. maomingense in having mainly ovoid (in lateral view) pyrenes, rounded triangular in cross-section, with acute to obtuse bases and slightly acuminate to acute apices and pyrene length/width ratios of about 1.6–2.2 (commonly 1.8).\textsuperscript{20}

Hence, the comparison of morphological characteristics displayed by the fruit fossils from the Maoming Basin studied in this paper shows that they differ from those of previously described fossil species and therefore we assign our fossils to a new species.

**Paleophytogeographical and paleoecological implications**

Based on the modern distribution of the Burseraceae in the American, African, and Indo-Asian tropics, and their highest generic diversity being in the Southern Hemisphere, Leenhouts\textsuperscript{5} and Raven and Axelrod\textsuperscript{25} considered that this family may have originated in tropical Gondwana and experienced vicariance as a result of Gondwanan fragmentation. McLaughlin\textsuperscript{26} suggested that the vicariance of Gondwanan angiosperms may be associated with more recent tectonic events, such as the collision of the Indian subcontinent with Asia followed by angiosperm dispersal to other regions of Northern and Southern Hemispheres. Lam\textsuperscript{27} also presumed that some genera of Burseraceae, in particular, Canarium may have originated in Africa and then reached Asia via the India connection. However, the estimates of divergence time for a number of tropical angiosperm lineages have indicated that Northern Hemisphere land connections had more influenced on the plant migration than previously hypothesized.\textsuperscript{1,28–30} Based on nuclear and chloroplast sequence data and fossil calibration of the molecular phylogeny, Weeks et al.\textsuperscript{1,30} suggested a North American origin of Burseraceae in the Late Cretaceous followed by migration of its lineages eastward over the North Atlantic land bridge to Europe and along
the northern margin of Tethys Sea to Southeast Asia, as well as long distance dispersals to Africa and South America.

Although fossil records of Canarium are scarce, their occurrences known to date are consistent, generally, with the Burseraceae diversification history proposed by Weeks et al. The oldest fruit fossils of Canarium come from the lower Eocene of western North America (Figure 7). By the middle Eocene, the genus extended throughout North America and had entered Europe. In the Oligocene, the reconstructed climate in most of Eurasia resembled that of a subtropical humid climate with distinctive seasonality, and frost-intolerant Canarium spreads across Europe, Asia, and northern Africa. During the Miocene, populations of Canarium from high- and mid-latitude regions of the Northern Hemisphere were presumably eliminated during global climate cooling at the Oligocene-Miocene transition and were forced to move southward into more favorable climates. The Miocene occurrences of Canarium are currently known only from southern Japan and southern China, while Mi-Pliocene Canarium fossils are restricted to India (Figure 7). For the Pleistocene and Holocene, Canarium is limited to southern Asia, Africa, northern Australia, and the Pacific Islands, which corresponds with the extant range of the genus.

Canarium first appears in South China in the late Oligocene and becomes more diverse in the Miocene. Pyrenes of both species from Guangxi and Guangdong provinces, previously described as the late Oligocene-Miocene C. guangxiense and the newly recognized late Pleistocene C. maomingense, morphologically closely resemble those of the extant C. album. Moreover, two pyrenes from the Miocene of Japan were assigned to the extant C. album. Hence, the fossil species related to C. album existed within a region close to the modern native range of this extant species from at least the late Oligocene.

C. album is native to the subtropical and tropical regions of southern China (Yunnan, Guangxi, Guangdong, Fujian, and Taiwan) and Vietnam approximately between 22°N and 26°N. Evergreen trees of this species
Canarium fossil records reported from the upper Oligocene of Nanning, the Miocene of Guiping and the upper Miocene of Beihai, Guangxi, as well as the upper Pleistocene of Maoming, Guangdong, all fall within the modern distribution range of this genus. It can be speculated that, in the gradually intensifying Cenozoic global cooling, Canarium may have continued to exist in this region since the late Oligocene. In the past 66 Ma, four Earth’s climate states are identified based of their response to astronomical forcing depending on greenhouse gas concentrations and polar ice sheet volume: Warmhouse (66–56 Ma and the past 66 Ma), Hothouse (56–47 Ma), Coolhouse (34–3.3 Ma), and Icehouse (3.3–0 Ma).34 The late Pleistocene Icehouse climate in southern China was limited. It can be inferred from the genus distribution that the influence of Pleistocene Icehouse climate in southern China was limited.

Specific fruit characters of fossil and extant Canarium species

Extant Canarium fruits are trilocular with two hanging campilotropous ovules per locule, of which only one or rarely two locules contain a developed seed (Figure 3K and 3L), usually a single large seed with twisted cotyledons.67 Placenta (Figures 4H and 4M), the position of the ovule attachment, can be seen on the endocarp, and normally the seeds or/and the ovule(s) which do not develop into the seed can be observed in locule (Figures 4I and 4N) in extant Canarium. The micro-CT and macro-observations of C. maomingense pyrenes have revealed two ovules per locule, but only one ovule per locule (or none) develops into a seed and others remain undeveloped (Figure 4O). The ovule is attached to the placenta (Figures 4E, 4F, 4G, and 4L) and connected with the ovariolar vascular bundle which branches from the central vascular bundle (Figures 4J, 4K, and 4O). These characteristics are identical with the extant Canarium species and indicate that this phenomenon of Canarium fruit development had existed in the late Pleistocene.

Relationships with other organisms

Weeks et al.1 pointed out that the fruits of the Burseraceae do not have morphologies related to external and passive transmission by animals such as viscous mucus or hooks, but some genera have fruits more likely to be dispersed by animal feeding behavior. Animals either discard the inedible part of fruit immediately after eating the fleshy part of the pericarp or aril, or excrete some complete hard and stony parts after digestion. The fleshy and juicy mesocarp of Canarium fruits may attract animals, while some animals may destroy the pyrene in order to obtain nutritious seeds of Canarium.

Some damage can be seen on the surfaces of fossil pyrenes, such as dents (Figures 8A and 8B), scratches (Figures 8C, 9C, 9D, and 9E), relatively small rounded or oval holes (Figures 9A, 9B, 10D, 10F, and 10H), holes of various shapes (Figures 9C and 9D), traces of surface biodegradation of uncertain nature (Figures 8B, 9E, 9F, 10A, and 10B), and evidence of fungal attack (Figures 9G, 10C, 10E, and 10G). There is a certain regularity in the scratch damage on some pyrene surfaces, and many of them are a group of three marks (Figures 8B, 8C, 8E, 8F, and 9C), which is obviously different from the random and irregular scratch that may be caused by water transportation and other

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taphonomic processes. These damage types are probably caused by the teeth or claws of animals scraping the fleshy mesocarp. According to a range of studies, birds such as pigeons and hornbills as well as mammals of the order Chiroptera (e.g., bats and flying foxes), lemurs, squirrels, and terrestrial mammals such as rodents and deer consume the fleshy mesocarp and seeds of *Canarium* fruits.47–51 Flying foxes (bats) are considered to be active consumers of *Canarium* fruits,50 and originated in the early Pliocene and diversified during the Pleistocene inhabiting tropical regions of Asia, Australia, Africa, and some Pacific Islands.51 A variety of dents and holes can be created by animals as a result of attempts to gnaw the hard pyrene in search of a nutritious seed inside. Larger holes of various shapes, accompanied by cracks of the pyrene wall, may have resulted from pyrene splitting by mammalian teeth (Figures 9C and 9D). Single small holes that appear in some pyrenes (Figures 9A, 9B, 10D, 10F, and 10H) are similar to the damage caused by insects like weevil beetles (family Curculionidae). Some weevil beetles, for example *Ectatorhinus magicus* Gerstaecker,52 can lay their eggs in fruits by piercing fleshy tissue as well as the woody endocarps/shell of the fruit, and also feed on dying wood.52

Numerous different in size, shape, and depth superficial damage on the pyrenes are interpreted as a traces of surface biodegradation caused by unknown microorganisms (Figures 8B, 9E, 9F, 10A, and 10B). Such damage can be inflicted by bacteria or microscopic fungi whose metabolites initiate a set of plant tissue-destroying chemical and physical processes. A single fungal fruiting body was noted on the surface of *Canarium* pyrene (Figures 9B, 9G, 10C, and 10E). The fruiting body is immersed in the tissue of the plant host, rounded in shape, with a diameter of 1.2 mm. The ostiole is inconspicuous. Free hyphae occur on the surfaces of the endocarp and fruiting body (Figures 10E and 10G). This fruiting body can be compared to those of the genus Coleophoma Höhn. (Ascomycota) characterized by similar hemispherical pycnidia,
immersed in the host plant tissue, with a non-papillate inconspicuous ostiole. *Coleophoma* occurs on a wide range of plant hosts and includes species that are saprobic or plant pathogenic.\(^{53}\) This fungal genus also has been identified on the fruits of extant *Canarium indicum* L.\(^{50}\) Consequently, the damage types described here indicate possible co-evolutionary relationships between plants and some mammals and/or birds. These frugivores could use *Canarium* fruits as a food source, as well as serve as seed dispersal agents for this plant. The type of relationships between *Canarium* and microscopic organisms (micromycetes and/or bacteria) is not yet well understood.

**Figure 9.** Damage on the pyrenes of *Canarium maomingense* sp. nov
(A) Round hole (arrow) accompanied by cracks, MMRH–210.
(B) Fungal fruiting body (arrow), MMRH–213.
(C) Oval hole and three scratches (arrows), MMRH–212.
(D) Hole and scratches, MMRH–246.
(E) Scratches with the traces of surface biodegradation (arrows), MMRH–246.
(F) Shallow superficial damage (arrows) of unknown nature, MMRH–196.
(G) Enlarged view of Figure 9B showing fungal fruiting body (arrow). Scale bars: (A and E) = 5 mm; (B–D and F) = 3 mm; (G) = 0.3 mm.
Limitations of the study

Most research studies of paleobotany are limited by materials of study. First, the number of plant macrofossil is limited. Secondly, the limitation of preservation of plant macrofossils. These two factors may affect the data representativeness, more accurate identification and classification based on morphology, and, eventually, reconstruction of paleogeography and paleoclimate.

STAR methods

Detailed methods are provided in the online version of this paper and include the following:

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Figure 10. Damage on the pyrenes of *Canarium maomingense* sp. nov., SEM
(A) Shallow superficial damage of unknown nature, MMRH–196.
(B) Magnification of Figure 10A showing superficial damage.
(C) Fungal fruiting body, MMRH–213.
(D) Round hole, MMRH–213.
(E) Enlarged view of Figure 10C showing free hyphae and the body forming hyphae.
(F) Round hole, MMRH–213.
(G) Free hyphae on the endocarp surface (arrow), MMRH–213.
(H) Round hole, MMRH–213. Scale bars: (A) = 1 mm; (B and C) = 300 μm; (D–H) = 100 μm.
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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.105385.

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AUTHOR CONTRIBUTIONS

H.X., J.J., and X.W. conceived and designed the project. J.J., L.H., and X.W. organized field work and the collection of fossils. H.X. prepared, and imaged fossil and modern specimens, processed data. T.K., M.R., and N.M. interpreted the data on fossil fungi and participated in plate making. H.X., T.K., M.R., N.M., M.H., X.W., L.H., and J.J. contributed to initial manuscript preparation. All authors discussed results, read, revised, and approved the final manuscript.

DECLARATION OF INTERESTS

The authors declare there is no conflict of interest.

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STAR★METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Biological samples** |        |            |
| Canarium maomingense fossil specimens | Museum of Biology, Sun Yat-sen University, Guangzhou | MMRH–192; MMRH–193; MMRH–194; MMRH–195; MMRH–196; MMRH–197; MMRH–198; MMRH–207; MMRH–208; MMRH–209; MMRH–210; MMRH–211; MMRH–212; MMRH–213; MMRH–245; MMRH–246; MMRH–247 |
| C. album | Museum of Biology, Sun Yat-sen University, Guangzhou | EC192247001 |

| **Software and algorithms** |        |            |
| Post-processing of images and color markings were performed with Adobe Photoshop 2020 | Adobe Inc. | RRID: SCR_014199, URL: https://www.adobe.com/products/photoshop.html |
| DigiCamControl-Free Windows DSLR camera controlling solution | Duka, 2015 | http://digicamcontrol.com/ |
| Helicon Focus | Helicon Inc. | RRID:SCR_014462, URL: http://www.heliconsoft.com/heliconsoft-products/helicon-focus/ |
| Dragonfly v.4.1 | Comet Group | https://www.theobjects.com/dragonfly |
| GBIF | the Global Biodiversity Information Facility | RRID: SCR_005904, URL: https://doi.org/10.15468/dl.6nhprp |
| CUBG | ICCBG2022 | https://image.cubg.cn/ |

RESOURCE AVAILABILITY

Lead contact
Further questions should be directed to the lead contact, Xinkai Wu (kakukaikai@163.com).

Materials availability
Fossil specimens MMRH–192, MMRH–193, MMRH–194, MMRH–195, MMRH–196 MMRH–197, MMRH–198, MMRH–207, MMRH–208, MMRH–209, MMRH–210, MMRH–211, MMRH–212, MMRH–213, MMRH–245, MMRH–246, MMRH–247, and extant C. album EC192247001 are deposited in the Museum of Biology, Sun Yat-sen University, Guangzhou, China. The comparison of C. maomingense with Asian extant Canarium species is available from Table S1. The comparison of C. maomingense with fossil species of Canarium and other related taxa is available from Table S2. Fossil records used in phytogeographic analyses is available from Table S3.

Data and code availability
● All data reported in this paper will be shared by the lead contact upon request.
● No novel code was used in this study.
● Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

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All specimens used here were obtained from the source organizations listed in the key resources table.
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**Geological setting and terminology**

The *Canarium* fossil fruits were collected from an opencast mine (21°52′47.5″N; 110°40′06.3″E) within the Maoming Basin, located near Zhenjiang Town, northwest of Maoming City, southwestern Guangdong Province, South China (Figure 1A). The fossiliferous deposits overlying with an angular unconformity the upper Eocene Huangniuling Formation have been dated to the late Pleistocene using accelerator mass spectrometry $^{14}$C dating of fossil wood and fruit samples (28660 ± 140 and 26860 ± 130 BP, respectively), carried out at the Beta Analytic testing laboratory, Miami, Florida, USA. $^{21}$ The measured sedimentary succession with a total thickness about 2.5 m is composed of yellow sands with gravel and black, gray and grayish yellow mudstones (Figures 1B and 1C). Three-dimensionally preserved mummified fruits were obtained mainly from a layer of gray mudstone in the middle part of the studied succession. This sedimentary succession also includes confers of the genera *Pinus* and *Keteleeria* (Pinaceae) and angiosperms belonging to *Elaeocarpus* (Elaeocarpaceae), *L. formosana* Hance (Altingiaceae), and Fagaceae. $^{21,42,43}$

To describe the fruits of fossil and extant species of *Canarium*, carpological terminology was used based on those of previous studies. $^{5,9,11,27}$ The details of the fruit and pyrene anatomy of *Canarium* species are shown in Figure 2. The pyrene is treated as being composed of the inner zone of the mesocarp and the endocarp. The inner mesocarp zone is represented by thick-walled sclereids, while the innermost layer of the mesocarp is composed of relatively thin-walled cells, which separate the hard mesocarp tissue from the endocarp. Due to the decay of this layer of thin-walled cells, the endocarp may detach from the hard mesocarp tissue in fossil specimens. The endocarp is the innermost pericarp zone that develops from the locular (inner) epidermis and is represented in both studied extant and fossil species by one layer of stout palisade sclereids (*C. chandleri*, *C. indicum*, *C. luzonicum*) or by brachysclereids (*C. oleosum*). $^{8-11}$ The term “pyrene” is used in this study for identifying the inner hard fruit part instead of the term “endocarp”, which is often applied to the hard tissues of the fossil fruits.

**Specimens imaging and the nearest living relative method**

Fossil fruits were cleaned with an ultrasonic cleaner (JieMeng JP–880) at a frequency of 42 kHz and photographed using a Nikon SMZ25 stereo microscope at the Museum of Biology, Sun Yat-sen University (Guangzhou, China). Extant fruits were photographed by a Canon EOS 760D digital camera. DigiCamControl-Free Windows DSLR camera controlling solution, $^{54}$ Helicon Focus 6.6.1 (Helicon Soft Ltd., Kharkov, Ukraine) and Photoshop 2020 (Adobe, San Jose, California, USA) were used for image processing. Microstructure was examined using a thermal field emission environmental scanning electron microscope (Quanta 400) at the test center, Sun Yat-sen University (Guangzhou, China). The internal three-dimensional structure of fruit was observed by Zeiss Xradia 520 Versa X-ray microscope (CT) housed at the University of Science and Technology of China (Hefei, China) and image data were processed using Dragonfly software.

The assumption that Cenozoic extinct plants and their nearest living relatives (NLRs) have similar environmental requirements, is the base of the NLR method. This method was used to estimate the paleoclimatic and paleoecologic conditions under which the studied fossil taxon lived by identifying the NLRs of fossil species and the environmental characteristics predominating in the regions of their modern natural distribution. $^{55}$