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Termites (Isoptera): Their Phylogeny, Classification, and Rise to Ecological Dominance

MICHAEL S. ENGEL, DAVID A. GRIMALDI, AND KUMAR KRISHNA

ABSTRACT

Like ants, termites are entirely eusocial and have profound ecological significance in the tropics. Following upon recent studies reporting more than a quarter of all known fossil termites, we present the first phylogeny of termite lineages using exemplar Cretaceous, Tertiary, and Recent taxa. Relationships among Recent families were largely unaffected by the addition of extinct taxa, but the analysis revealed extensive grades of stem-group taxa and the divergence of some modern families in the Cretaceous. Rhinotermitidae, Serritermitidae, and the "higher" termites (family Termitidae), which comprise 84% of the world termite species, diverged and radiated entirely in the Tertiary, corresponding to a significant increase in termite individuals in the fossil record. Radiation of the higher termites may have affected the formation of terrestrial carbon reserves like oil and coal. The higher classification of Isoptera is slightly revised based on the phylogenetic results. The following new taxa are proposed: Cratomastotermitidae, new family; Euisoptera, new clade; Archotermopsidae, new family; and Neoisoptera, new clade. In addition, the families Stolotermitidae, Stylotermitidae, and Archotenhinotermitidae are newly recognized or resurrected, and the families Termopsidae and Hodotermitidae are significantly restricted in composition.

INTRODUCTION

"Success" of a species or group of species is typically either ecological or evolutionary. Insects, in particular, are evolutionarily very successful because of their early origins in the Devonian and subsequent radiation into millions of species. It is only certain insects, however, that account for the remarkable overall ecological dominance of insects in terrestrial ecosystems, in terms of biomass and impact on biological communities, and

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chief among them are the social insects. This report is a phylogenetic synthesis of many recent discoveries of fossil termites, which is then used to consider the origins of termite ecological success.

Advanced sociality, or eusociality, involves overlapping generations of siblings that generally share a durable domicile in which groups of individuals specialize in tasks (i.e., castes), especially reproduction, foraging, and commonly defense and brood care (Wilson, 1971). A colony's efficiency in mobilizing its foragers and soldiers fosters the superior competitive ability of major eusocial groups like ants, vespid wasps, honeybees, and termites. For example, there are approximately 19,340 modern species of bees (Apoidea: Anthophila), but it is the large, complex colonies of honey bees (genus *Apis*) that easily outcompete the native social species, even leading to the latter's localized extirpation (e.g., Roubik et al., 1986; Sugden et al., 1996). Unlike bees, all of the 12,516 living species of ants and 2,958 living species of termites are eusocial (termite species numbers valid as of 11 March 2009: Krishna et al., in press). Basal lineages of both these groups generally live in small colonies of several dozen to several hundred individuals with less caste differentiation, and the most recently derived lineages (like army ants, leaf cutter ants, and moundbuilding termites) form massive colonies of over one million individuals with extreme caste differentiation. Termites in tropical and subtropical ecosystems are the major consumers of the most abundant biomolecule on land, cellulose, and its more inert form, lignocellulose. It is estimated, for example, that termites ingest 50%–100% of the dead plant biomass in tropical ecosystems (Bignell and Eggleton, 2000). Their abundance, like that of ants, frequently exceeds 1,000 individuals/m² or 2,000 mg/m², and it is estimated that gas excretion from termites and their nests contributes 2%-5% of the world's atmospheric methane (Sugimoto et al., 2000). The ecological impact of termites, even apart from the commercial damage they cause, is prodigious.

There has been intensive study of the relationships and fossil record of bees (Engel, 2001, 2004; Danforth et al., 2006; Michener, 2007; Ohl and Engel, 2007) and ants (Grimaldi

et al., 1997; Grimaldi and Agosti, 2001; Dlussky and Rasnitsyn, 2002; Engel and Grimaldi, 2005; Moreau et al., 2006; Brady et al., 2006). Both of these groups appear to have originated in the late Early Cretaceous, ca. 100–120 Ma, with some modern subfamilies diverging in the Late Cretaceous. Though their evolutionary history has been studied less, it is now acknowledged that termites are highly modified, eusocial roaches (Cleveland et al., 1934; McKittrick, 1964; Lo et al., 2000; Deitz et al., 2003; Grimaldi and Engel, 2005; Klass and Meier, 2006), whose earliest fossils predate those of ants and bees by approximately 35 million years (Thorne et al., 2000; Engel et al., 2007a). Recent phylogenetic work on termites involves molecular and some morphologically based analyses of living species only (e.g., Kambhampati et al., 1996; Donovan et al., 2000; Thompson et al., 2000; Bitsch and Noirot, 2002; Klass and Meier, 2006; Inward et al., 2007a, 2007b; Legendre et al., 2008). Unique for any insect order, all early fossil termites have been classified into living families, particularly the Hodotermitidae, despite the fact that such fossils may be stem groups since they lack many derived features of living families.

Here we present the first analysis of relationships among fossil and living termite lineages, along with estimates of divergence times and ecological patterns of the major lineages. Recent studies have reported 18 new termites from the Cretaceous (Krishna and Grimaldi, 2000, 2003; Engel et al., 2007a; Grimaldi et al., 2008) and 38 species from the Tertiary (Nel and Bourguet, 2006; Wappler and Engel, 2006; Engel and Krishna, 2007a, 2007b; Engel et al., 2007b; Engel, 2008; Krishna and Grimaldi, 2009), comprising about one-quarter of all described fossil species. Nearly 80% of these species are preserved in amber, from the Cretaceous of Lebanon, the mid-Cretaceous of Myanmar and New Jersey, the Eocene of the Baltic Region and France, and the Miocene of Mexico and the Dominican Republic (deposits are reviewed in Rasnitsyn and Quicke, 2002; Grimaldi and Engel, 2005). The microscopic fidelity of preservation in amber allows uniquely detailed comparisons with living species, greatly facilitating phylogenetic analysis of extinct taxa. Insects in 110 Ma limestone from the Crato Formation of Brazil also have exceptional preservation, as mineralized replicas with cuticular microstructure and even some internal organs. As a result, Crato species preserved as series of specimens can be largely reconstructed (Grimaldi et al., 2008). Thus, now is an opportune time for deciphering nearly 140 million years of termite evolution.

MATERIALS AND METHODS

Termite specimens belonging to 38 exemplar living species of 36 genera, representing all seven traditionally recognized families and four subfamilies of the "higher" termites (family Termitidae), were dissected and directly examined; 38 extinct species in 25 genera were studied and compared to living taxa. The fossils ranged in age from Early Cretaceous (Berriasian, ca. 135 Ma, of Baissa, Siberia) to the Miocene (in amber from the Dominican Republic, ca. 17 Ma) (table 2). Species preserved as compressions or mineralized replicas were also included where bodies were preserved, not just isolated wings (we have omitted from the analysis taxa known only or largely from wings, such as *Ulmeriella* Meunier).

There were 108 morphological and biological attributes scored from imago, soldier, and worker castes where available (table 1 and appendix); outgroup taxa were other Dictyoptera, specifically the most basal mantis (Chaeteessa sp. Burmeister), the roaches Periplaneta Burmeister and Panchlora Burmeister, and the relict wood roaches (Cryptocercus spp. Scudder), the latter being the living sister group to termites. Within the matrix of 8,748 cells, 21% of the cells were coded as unknown (feature not observed in the fossils), and a further 9% as inapplicable (e.g., soldier characters in genera that lack this caste). Phylogenetic analysis employed NONA (Goloboff, 1997), using 500 replicates of the data set with random taxon addition and branch swapping on all shortest topologies. This yielded 2,088 equally most-parsimonious trees of 302 steps (C.I. 44, R.I. 86), the strict consensus of which resulted in 323 steps (unambiguous character changes mapped in figs. 1, 2, 4, 5). The relative abundance of termites through time was plotted as proportions (%) of the number of termite specimens to all insect specimens per fossil deposit (table 2); only deposits that yielded at least one termite specimen were plotted, in order to ensure the appropriate taphonomic conditions for termite preservation.

RESULTS AND DISCUSSION PHYLOGENY

Analysis of the paleontological data resulted in a remarkably resolved topology for Isoptera (fig. 3). Structure of the consensus tree generally agrees with that from prior morphological and molecular studies (Kambhampati et al., 1996; Donovan et al., 2000; Thompson et al., 2000; Bitsch and Noirot, 2002; Inward et al., 2007a, 2007b), but depart in some respects from the recent molecular study of Legendre et al. (2008) (e.g., the relatively basal placement of Kalotermitidae). Legendre et al. (2008) had a dramatically reduced taxon sampling relative to other, more comprehensive treatments (e.g., Inward et al., 2007a).

In our analysis unequivocal relationships include Cryptocercus as the living sister group to the termites and *Mastotermes* Froggatt as the living sister group to all other termites the lineage we are designating here as Euisoptera. Mastotermes darwiniensis Froggatt, from northern Australia and southern New Guinea, is the sole survivor of the formerly global Mastotermitidae (Thorne et al., 2000; Wappler and Engel, 2006), and it retains striking plesiomorphic features with roaches, such as laying its eggs in a vestigial pod or ootheca. Interestingly, the Mastotermitidae as it has historically been classified was recovered as monophyletic, despite opinion to the contrary (e.g., Jarzembowski, 1981). Cratomastotermes Bechly, from the Early Cretaceous Crato Formation of Brazil, was formerly placed in the Mastotermitidae (Bechly, 2007); in our analysis it is the most basal species of termite since it retains even more plesiomorphic features than Mastotermitidae (Grimaldi et al., 2008). This genus is accordingly placed in a new, extinct family, Cratomastotermitidae, new family (see Classification). Cretaceous fossils traditionally classified as Hodo-

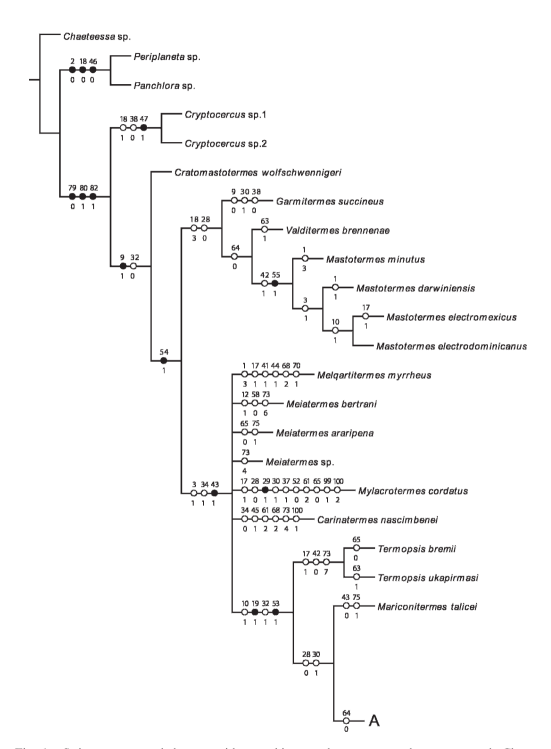


Fig. 1. Strict consensus cladogram with unambiguous character-state changes mapped. Character numbers appear above the branch, with the corresponding state beneath the branch. Branch "A" continued in figs. 2, 4, and 5. *Chaeteessa* (Mantodea), *Panchlora* (Blattaria), *Periplaneta* (Blattaria), and *Cryptocercus* (Blattaria) are the outgroup taxa.

TABLE 1 Character and Character-State Descriptions

The following list enumerates those characters and character states used in the analysis of relationships within Isoptera (refer to appendix 1 for codings). All characters were considered nonadditive and of equal weights

- 1. Number of alate/imago antennomeres: 50 or more antennomeres = 0; 30–40 antennomeres = 1; 23–28 antennomeres = 2; 11–22 antennomeres = 3.
- 2. Structure of imago flagellum: flagellomeres filiform = 0; flagellomeres moniliform = 1.
- 3. Apex of imago flagellum: distal 7-8 articles tapered = 0; distal 7-8 articles not tapered = 1.
- 4. Ocelli of alates: three present (homologous with other orders) = 0; absent = 1.
- 5. Ocelli of soldiers: present = 0; absent or vestigial = 1.
- 6. Pigmentation of soldier compound eyes: present = 0; absent = 1.
- 7. Soldier compound eyes: rudimentary = 0; absent = 1.
- 8. Frontal gland developed into distinct fontanelle: absent = 0; present = 1.
- 9. Position of alate head: hypo-/orthognathous = 0; prognathous = 1.
- 10. Ventral cervical sclerite of alate: present = 0; absent = 1.
- 11. Marginal teeth of left mandible of alate: greater than or equal to three = 0; two teeth = 1.
- 12. Subsidiary tooth of right mandible of alate: absent = 0; present = 1.
- 13. Lacinial teeth of maxilla: both teeth apical = 0; one tooth subapical = 1.
- 14. Postclypeal furrow (workers and alates): absent = 0; present as shallow longitudinal furrow = 1.
- 15. Clypeus (workers only) in profile: not keeled = 0; with keel = 1.
- 16. Shape of occipital foramen in imagos: rounded = 0; trapezoidal = 1.
- 17. Y-shaped coronal ecdysial cleavage line in imago: present = 0; absent or highly vestigial = 1.
- 18. Pair of ocelluslike structures (= ocelloids) near inner margin of compound eye: present as circular tympanalike areas = 0; vestigial as areas of weakened or pale cuticle = 1; absent = 2; present and nearly lenslike (superficially resembling ocelli and historically termed ocelli by isopterists) = 3.
- 19. Occipital sulcus: present = 0; absent or highly vestigial = 1.
- 20. Compound eye: lenticular = 0; circular = 1.
- 21. Notch between first and third marginal teeth: present = 0; absent = 1.
- 22. Mandibular excavation between apical and first marginal teeth: absent = 0; present = 1.
- 23. Soldier mandible: marginal teeth: distributed along length, with 1-4 teeth along middle of margin = 0; lost except small teeth/serrations at base = 1.
- 24. Soldier mandible: short to moderate in length, length ca 2 or less × basal width = 0; greatly elongate and narrow, length ca. 3 or more × basal width = 1; vestigial = 2.
- 25. Soldier mandibles: symmetrical, not clicking = 0; asymmetrical, clicking = 1.
- 26. Soldier head capsule: rectangular in dorsal aspect = 0; phragmotic, plug shaped = 1; nasutiform = 2.
- 27. Diagonal grooves between fontanelle and postclypeus: absent = 0; present = 1.
- 28. Pronotal posterolateral corners of imago: broadly arched = 0; acutely rounded, nearly orthogonal = 1.
- 29. Pronotal posterior margin of imago: straight or slightly indented = 0; rounded = 1.
- 30. Pronotal lateral margins of imago: subparallel = 0; converging = 1.
- 31. Pronotal size of imago: covering head dorsally = 0; not covering head but with width greater than or equal to head = 1; width significantly less than head width = 2.
- 32. Pronotal anterior margin of imago: concave = 0; relatively straight = 1; convex, with anterolateral corners developed = 2.
- 33. Tibial macrosetae and spurs: heavily serrate = 0; slightly pimplate = 1; smooth = 2.
- 34. Tibial macrosetae and spurs: asymmetrical, with one side flattened = 0; symmetrical = 1.
- 35. Tarsomeres: pentamerous, fully developed = 0; pentamerous, second tarsal article reduced = 1; tetramerous (second tarsal article lost) = 2; trimerous = 3.
- 36. Metabasitarsomere length: less than twice the width = 0; more than $3 \times$ the width = 1.
- 37. Pretarsal arolium: present = 0; absent = 1.
- 38. Plantular pads: present = 0; absent = 1.
- 39. Procoxal ventral keel of alate: present = 0; absent = 1.
- 40. Protibial spines along length: present and extensive = 0; absent, reduced to apical spines/spurs = 1.
- 41. First Rs fork: in basal half = 0; near midlength or beyond = 1.
- 42. Length of R1: short, extending in quarter of wing length = 0; median, extending in third of wing length = 1; long, extending to or past wing midlength = 2.
- 43. Forewing Rs: branched basally in scale with 2-3 branches = 0; simple in scale = 1.
- 44. Forewing CuA: shorter, to around wing midlength = 0; long, to point within apical third of wing = 1; elongate and extensively developed, extending to apex or subapex, posteriorly along one-half wing, with 6 or more branches = 2.

TABLE 1 (Continued)

- 45. Multiple branches of R1: present = 0; absent = 1.
- 46. Forewing tegmenization: developed as tegmina = 0; not developed as complete tegmina = 1.
- 47. Wings: present = 0; absent = 1.
- 48. Forewing vein M: present = 0; absent = 1.
- 49. Forewing scale relative to hind wing scale: apical margin of forewing scale meeting or overlapping hind scale = 0; scales well separated = 1.
- 50. Wing membrane setae: absent = 0; present, microsetulose = 1.
- 51. Fore- and hind wings: not dehiscent = 0; dehiscent = 1.
- 52. Forewing basal cleavage suture: absent = 0; present = 1.
- 53. Hind wing basal cleavage suture: absent = 0; present but rudimentary = 1; completely developed = 2.
- 54. Cross veins connecting longitudinal veins: present = 0; absent = 1.
- 55. Sclerotization of veins: all uniformly sclerotized = 0; Sc, R, and M thick and sclerotized relative to CuA = 1; Sc and R sclerotized relative to M and CuA = 2.
- 56. Anal lobe of hind wing: present = 0; absent (wing homonomous) = 1.
- 57. Hind wing vein A1: present = 0; absent = 1.
- 58. Wing membrane reticulations: present, not pigmented = 0; present, pigmented = 1; absent = 2.
- 59. Wing membrane surface: smooth = 0; nodulose or pimplate = 1.
- 60. Radial fracture of forewing scale: present = 0; absent = 1.
- 61. Reticulations between CuA and CuP on forewing scale: veins = 0; reticulations = 1; reticulations only anterior to CuP = 2; absent = 3.
- 62. Distal margin of forewing scale: evenly convex = 0; straight to slightly convex = 1; straight and diagonal = 2.
- 63. CuP in forewing scale: convex = 0; straight or concave = 1.
- 64. Humeral margin of forewing scale: flat = 0; swollen beyond level of costal margin = 1.
- 65. Forewing CuP: terminates prior to posterior tip of basal suture = 0; terminates in basal suture = 1.
- 66. Position of forewing vein M: close to Rs = 0; midway between Rs and CuA or closer to latter = 1.
- 67. Costalization of forewing: not costalized = 0; C+Sc+R and Rs extremely close and simple and parallel = 1.
- 68. Branches of Rs: with dorsal and ventral branches, inferiors irregular (acute go tip) = 0; with dorsal and ventral branches, inferiors angled & parallel = 1; with short dorsal branches only or simple = 2.
- 69. Branches of vein M: with two or more branches = 0; with one apical branch = 1; simple (unbranched) = 2.
- 70. Radial field: encompassing apex or terminating at apex = 0; terminating anterior to apex = 1.
- 71. Development of CuP: extensive, developed as claval furrow = 0; reduced, confined to short, simple vein near wing base = 1.
- 72. Dichotomous branching of R and M: extensive = 0; reduced = 1.
- 73. Number of superior branches of Rs: none = 0; one = 1; two = 2; three = 3; four = 4; five = 5; six = 6; seven or more = 7.
- 74. Female styli: present = 0; absent = 1.
- 75. Imago cercus segmentation: four or higher = 0; three = 1; one or two = 2.
- 76. Castes: absent = 0; present, with workers = 1; present, without workers = 2.
- 77. Ootheca: present = 0; absent = 1.
- 78. Presence of *Blattabacterium* in fat body: present = 0; absent = 1.
- 79. Presence of flagellates and ciliates: present = 0; absent = 1.
- 80. Wood feeding: absent = 0; present = 1.
- 81. Social organization: absent = 0; extended parental care = 1; eusocial = 2.
- 82. Living in structures/nests: absent = 0; present = 1.
- 83. Soldiers: present = 0; lost = 1.
- 84. Soldier nasus: short = 0; very long, length several times width = 1.
- 85. Nasute fontanelle: rimmed or slitlike = 0; minute, not rimmed or slitlike = 1.
- 86. First proctodeal segment: expanded = 0; tubular, not dilated = 1.
- 87. Soldier labral apex: sclerotized = 0; hyaline = 1.
- 88. Soldier heads: normal = 0; flattened = 1.
- 89. Soldier labrum: well developed = 0; vestigial = 1.
- 90. Soldier pronotum: flat = 0; saddle shaped = 1.
- 91. Malpighian tubule number: eight or more = 0; four or fewer = 1.
- 92. Imago/worker fontanelle: small and round = 0; slit-, drop-, or Y-shaped = 1.
- 93. Anterior margin of worker/alate postclypeus: flat or concave = 0; convex = 1.
- 94. Imago compound eye: protruding well beyond lateral margin of head = 0; small, not protruding beyond lateral margin of head in frontal view = 1.

TABLE 1 (Continued)

- 95. Imago ocelloid: large, ca. 7–8x diameter of compound eye facet = 0; small, ca. 2–3x diameter of compound eye facet = 1.
- 96. Soldier fontanelle: normal, dorsal-facing = 0; enlarged and facing anteriad = 1.
- 97. Nasute head capsule: not constricted = 0; slightly constricted = 1.
- 98. Proventricular teeth: present = 0; highly reduced or lost = 1.
- 99. Protibial apical spur number: three or more = 0; two = 1.
- 100. Mesotibial apical spurs: four or five = 0; three = 1; two = 2.
- 101. Metatibial apical spurs: four = 0; three = 1; two = 2.
- 102. Sternal gland on third sternum: present = 0; absent = 1.
- 103. Sternal gland on fourth sternum: present = 0; absent = 1.
- 104. Sternal gland on fifth sternum: present = 0; absent = 1.
- 105. Soldier labral brush: absent = 0; present = 1.
- 106. Setulae surrounding fontanelle (directed toward fontanelle): absent = 0; present = 1.
- 107. Hind wing vein M: present = 0; absent = 1.
- 108. Soldier frontal groove: absent = 0; present (ridges anterior from fontanelle with width of fontanelle) = 1.

termitidae, such as the extinct "genus" Meiatermes Lacasa-Ruiz and Martínez-Delclòs, actually comprise a grade of stemgroup species that fall basal to the divergence of Termopsidae, true Hodotermitidae, and all other termites. Termopsidae stat. n. (see Classification) should be restricted to the Baltic amber species of *Termposis* Heer (Engel et al., 2007b). What we are designating as the true Hodotermitidae (see Classification)

TABLE 2
Termite Abundance (% of all insect specimens) in Major Insect
Deposits throughout the Cretaceous and Cenozoic

Deposit	Age	% termites	References
		CRETACEOUS	
Baissa, Siberia	137 Ma	0.1%	Rasnitsyn, 2008
Montsec, Llerida, Spain	129 Ma	0.2%	Lacasa-Ruiz and Martínez-Delclòs, 1986
Weald clay, UK	127 Ma	0.1%	Jarzembowski, 1977, 1984, 1991
Lebanese amber	120 Ma	0.2%	Azar, 2000
Crato Formation, Brazil	110 Ma	1.0%	Bechly, 2007; Grimaldi, unpubl. data
Burmese amber	100 Ma	0.2%	Grimaldi, unpubl. data
French amber (Archingeay)	100 Ma	0.03%	Perrichot et al., 2007
New Jersey amber	90 Ma	0.4%	Grimaldi et al., 2000
		CENOZOIC	
Baltic amber	45 Ma	0.2%	Sontag, 2003
Baltic amber	45 Ma	0.2%	Perkovsky et al., 2007
Baltic amber	45 Ma	0.3%	Krzemińska et al., 1992
Eckfeld Maar, Germany	45 Ma	0.5%	Wappler, 2003
Kŭclín, Czech Rep.	40 Ma	6%	Prokop, 2003
Bembridge Marls, UK	34 Ma	2.5%	Jarzembowski, 1980
Seifhennersdorf, Germany	30 Ma	6%	Prokop, 2003
Krottensee, Bohemia	18 Ma	5%	Prokop, 2003
Dominican amber	18 Ma	5.6%	Grimaldi, unpubl. data
Dominican amber	18 Ma	2.8%	Poinar and Poinar, 1999
Andance, France	15 Ma	10%	Prokop, 2003
Oeningen, Germany	15 Ma	2%	Prokop, 2003
Radoboj, Croatia	15 Ma	2%	Prokop, 2003
Zanzibar copal	1000 y.	8%	Grimaldi, unpubl. data
Colombian copal	300 y.	10%	Grimaldi, unpubl. data

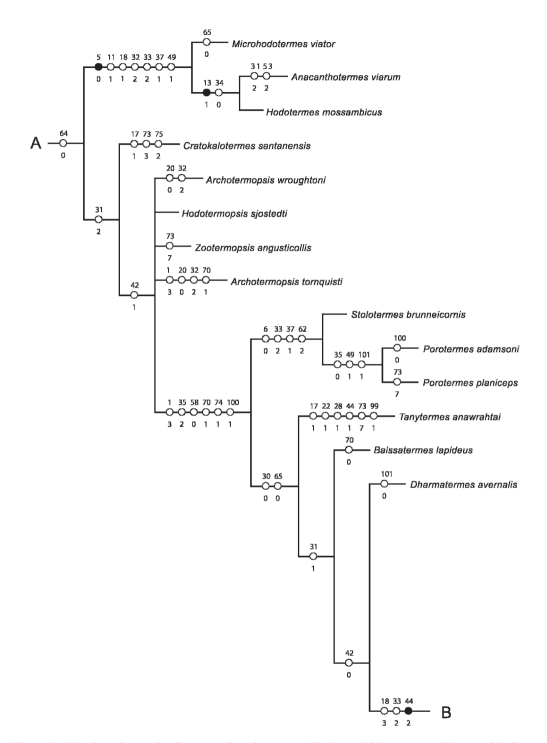


Fig. 2. Continuation of figure 1 focusing on Hodotermitidae s.s., "Termopsidae" s.s., Archotermopsidae, n. fam., Stolotermitidae, stat. n., and various stem-group lineages. Branch "B" (Kalotermitidae + Neoisoptera) is depicted in fig. 4.

is a group of three genera of the "harvesters"—Anacanthotermes Jacobson, Hodotermes Hagen, Microhodotermes Siöstedt—that feed on grasses in savanna and steppe biomes of Africa and Eurasia; monophyly of this group is confirmed by other analyses (Inward et al., 2007a, 2007b; Legendre et al., 2008). Three genera of highly disjunct wood feeders that are traditionally placed in the Termopsidae s.l.—Archotermopsis Desneux, Zootermopsis Emerson, and *Hodotermopsis* Holmgren—were not found to be monophyletic, contrary to other studies. Inclusion or exclusion of the fossils made no difference to the definition of "Termopsidae s.l." except that Archotermopsis was sometimes found to be monophyletic. As such, the traditional concept of Termopsidae s.l. could not be supported and these genera are accordingly classified in the new family Archotermopsidae, new family (see Classification).

The austral disjuncts, Stolotermes Hagen and Porotermes Hagen, are sometimes placed within Termopsidae s.l. (e.g., Donovan et al., 2000; Thorne et al., 2000; Legendre et al., 2008) but should clearly be the separate family Stolotermitidae, stat. n. (= Stolotermitinae + Porotermitinae: see Classification), since they are the living sister group to Kalotermitidae plus higher termites in this study and in some molecular analyses (Inward et al., 2007a, 2007b). The divergence between Stolotermitidae and Kalotermitidae + higher termites was certainly in the Early Cretaceous, since there are three intermediate genera from the Early and mid-Cretaceous. The family Kalotermitidae, or dry-wood termites, is a cosmopolitan group of 457 living species, which has two known Cretaceous and three Tertiary stem-group taxa (only two of the latter were included in our study), with living species comprising a monophyletic group of probably Tertiary origin. Interestingly, a kalotermitid-like nest exists from the Late Cretaceous Javelina Formation of Texas (Rohr et al., 1986). Kalotermitidae is the sister group to an unequivocally monophyletic group we are calling the Neoisoptera, which is defined in part by the distinctive opening of the frontal gland called the fontanelle. The Neoisoptera is comprised of the Rhinotermitidae (13 living genera, 380 living species, all of which are wood feeders), the monotypic family Serritermitidae from Brazil, and the largest family, Termitidae. It also includes Archeorhinotermes rossi Krishna and Grimaldi in 100 Ma Burmese amber, the most derived termite from the Cretaceous, formerly placed Rhinotermitidae (Krishna Grimaldi, 2003), but actually an extinct stem group to the rest of the Neoisoptera and here considered as the sole member Archeorhinotermitidae, stat. Classification). Parastylotermes Snyder and Emerson and Stylotermes Holmgren and Holmgren (neither genus studied in prior analyses [Donovan et al., 2000; Inward et al., 2007a, 2007b; Legendre et al., 2008]) should be separated from Rhinotermitidae s. str., the latter doubtfully monophyletic (Donovan et al., 2000; Inward et al., 2007a, 2007b; Legendre et al., 2008) but clearly closely related to the Termitidae. These genera have been at times considered a separate family, as Stylotermitidae, stat. rev. (see Classification), and this status should be reinstated. The earliest rhinotermitids are Reticulitermes antiquus (Germar) and Heterotermes eocenicus Engel in Baltic amber (Engel et al., 2007b; Engel, 2008); the divergence of both families probably occurred in the Early Tertiary.

CLASSIFICATION

Several taxonomic changes are required in order to have the classification of termite families reflect our cladistic results. The revised, higher-level classification of Isoptera is outlined in table 3 (modified from Engel and Krishna, 2004a, 2004b, 2007c). The classification is that which is employed for (and will be further elaborated in) the forthcoming world catalog of Isoptera (Krishna et al., in press).

CRATOMASTOTERMITIDAE, new family

Type genus: Cratomastotermes Bechly.

DIAGNOSIS: Diagnosed by primitive retention of distinct cross veins, archedictyon between veins, arched humeral margin of forewing scale, large rectangular pronotum, pentamerous tarsi, and absence of ocelloids and fontanelle (refer also to Grimaldi et al., 2008).

TABLE 3

Synonymical Hierarchical Classification of Isoptera

(modified and updated from Engel and Krishna, 2004a, 2004b, 2007c)

Order ISOPTERA Brullé

Family †Cratomastotermitidae Engel, Grimaldi, and Krishna, new family

Family Mastotermitidae Desneux

- = †Pliotermitinae Pongrácz
- = †Miotermitinae Pongrácz

EUISOPTERA Engel, Grimaldi, and Krishna, new clade

Family Incertae Sedis

(a grade of primitive termites whose relationships are not yet resolved)

- = †Cretatermitinae Emerson
- = †Lutetiatermitinae Schlüter
- = †Carinatermitinae Krishna and Grimaldi
- = †Caatingatermitinae Martins-Neto et al.

Family †Termopsidae Holmgren, sensu stricto

Family Hodotermitidae Desneux, sensu stricto

Family Archotermopsidae Engel, Grimaldi, and Krishna, new family

Family Stolotermitidae Holmgren, status novus

Subfamily Stolotermitinae Holmgren

Subfamily Porotermitinae Emerson

Family Kalotermitidae Froggatt

- = Glyptotermitinae Froggatt
- = †Electrotermitinae Emerson

NEOISOPTERA Engel, Grimaldi, and Krishna, new clade

Family †Archeorhinotermitidae Krishna and Grimaldi, status novus

Family Stylotermitidae Holmgren and Holmgren, status revivisco

Family Rhinotermitidae Froggatt

Subfamily Coptotermitinae Holmgren

= Arrhinotermitinae Sjöstedt

Subfamily Heterotermitinae Froggatt

= Leucotermitinae Holmgren

Subfamily Prorhinotermitinae Quennedey and Deligne

Subfamily Psammotermitinae Holmgren

Subfamily Termitogetoninae Holmgren

Subfamily Rhinotermitinae Froggatt

Family Serritermitidae Holmgren

= Glossotermitinae Cancello and DeSouza, new synonymy

Family Termitidae Latreille

Subfamily Sphaerotermitinae Engel and Krishna

Subfamily Macrotermitinae Kemner

- = Acanthotermitinae Sjöstedt
- = Odontotermitini Weidner

Subfamily Foraminitermitinae Holmgren

= Pseudomicrotermitinae Holmgren, new synonymy

Subfamily Syntermitinae Engel and Krishna

Subfamily Nasutitermitinae Hare

Subfamily Apicotermitinae Grassé and Noirot

= Indotermitidae Roonwal and Sen Sarma

Subfamily Cubitermitinae Weidner, status revivisco et novus

Subfamily Termitinae Latreille

- = Microcerotermitinae Holmgren
- = Amitermitinae Kemner
- = Mirocapritermitinae Kemner
- = Mirotermitini Weidner
- = Capritermitini Weidner

Incertae Sedis †Eutermitinae Holmgren

INCLUDED GENERA: The family presently comprises a single genus, *Cratomastotermes*.

Family Termopsidae Holmgren, sensu novum

Termopsinae Holmgren, 1911: 35. Type genus: *Termopsis* Heer, 1849.

COMMENTS: Since the time of Hagen (1858), the fossil Termopsis bremii Heer (1849) has been intricately linked to a group of otherwise plesiomorphic modern species. Over the intervening 150 years a few additional genera and several living species have been added to the group that became known as the Termopsinae and eventually Termopsidae (e.g., Emerson, 1933). Unfortunately, no singular, specialized (i.e., apomorphic) feature has truly united these taxa and the monophyly of the group has been suspect. Our analysis reveals that those fossils of the Tertiary genus Termopsis are, in fact, not related to the modern members otherwise classified in the family (namely Archotermopsis, Zootermopsis, and *Hodotermopsis*, below classified into a new family). In addition, those other fossil and living genera also traditionally classified in the family (e.g., Stolotermes and Porotermes of the Stolotermitinae and Porotermitinae, respectively) were similarly found to be unrelated to Termopsis, as well as unrelated to the aforementioned genera of Termopsinae. Accordingly, Termopsidae is here significantly restricted and considered to comprise only the genus Termopsis.

Archotermopsidae, new family

Type genus: Archotermopsis Desneux.

DIAGNOSIS: The new family can be characterized by the following combination of attributes: absence of ocelloids and fontanelle, antennae with 22–27 articles, pronotum distinctly narrower than head, tarsi pentamerous (sometimes cryptically), fourth sternite with sole sternal gland, forewing scale overlapping hind-wing scale, humeral margin of scale flat, imago-worker mandibles with three marginal teeth (left side) and subsidiary tooth between apical and first marginal teeth (right side).

INCLUDED GENERA: Archotermopsis, Zootermopsis, Hodotermopsis, and tentatively †Parotermes Scudder. The Late Miocene genus

† Gyatermes Engel and Gross (2009) may belong herein but must await more completely preserved material.

Family Hodotermitidae Desneux, sensu novum

Hodotermitini Desneux, 1904: 284. Type genus: *Hodotermes* Hagen, 1853.

Comments: Numerous genera of plesiomorphic fossil termites have been historically assigned to Hodotermitidae, leaving the impression that this group was once diverse in the past but has experienced significant extinction and that the modern taxa are relicts of this former diversity. In fact, our study demonstrates that none of the fossils assigned to Hodotermitidae can be considered actual hodotermitids. Instead, this assemblage represents a grade between several families and lineages of Isoptera. Hodotermitidae was recovered as a monophyletic group but strictly for the modern genera. We have accordingly restricted the sense of Hodotermitidae to those genera (listed below) and consider the remaining groups such as Cretotermitinae, Carinatermitinae, Lutetiatermitinae, and Caatingatermitinae (the latter two ill defined on teratologies and misinterpreted characters, respectively), among numerous other genera Meiatermes; *Melgartitermes* (e.g., Grimaldi, and Krishna; Mylacrotermes Engel, Grimaldi, and Krishna; Mariconitermes Fontes and Vulcano; Cratokalotermes Bechly) as incertae sedis among basal Euisoptera (fig. 3).

INCLUDED GENERA: *Hodotermes, Anacanthotermes,* and *Microhodotermes.*

Family Stolotermitidae Holmgren, status novus

Stolotermitinae Holmgren, 1910: 285. Type genus: Stolotermes Hagen, 1858.

COMMENTS: The Stolotermitidae is here recognized to encompass the former subfamilies Stolotermitinae and Porotermitinae of Termopsidae s.l. For the moment the two subfamilies are retained despite each being monogeneric.

INCLUDED GENERA: *Stolotermes* (in Stolotermitinae) and *Porotermes* (in Porotermitinae).

Family Archeorhinotermitidae Krishna and Grimaldi, status novus

Archeorhinotermitinae Krishna and Grimaldi, 2003: 2. Type genus: *Archeorhinotermes* Krishna and Grimaldi, 2003

COMMENTS: Although previously classified as a primitive lineage in the Rhinotermitidae, affinities with this group are entirely plesiomorphic. As our analysis reveals, *Archeorhinotermes* is actually a stem group, basal to all Euisoptera. Accordingly we have removed the genus from Rhinotermitidae and elevated Archeorhinotermitinae to familial rank.

INCLUDED GENERA: The family includes only *Archeorhinotermes* at present.

Family Stylotermitidae Holmgren and Holmgren, status revivisco

Stylotermitinae Holmgren and Holmgren, 1917: 141. Type genus: *Stylotermes* Holmgren and Holmgren, 1917.

Comments: Stylotermes and the Tertiary genus Parastylotermes have historically been classified in the Rhinotermitidae. These genera are particularly distinctive in their possession of trimerous tarsi, a feature otherwise known only in Indotermes of the Termitidae. The significance of this tarsal reduction was used by some authors in the past to accord Stylotermitinae familial rank, in the same fashion that *Indotermes* was placed in a monogeneric family of its own (e.g., Roonwal, 1958). Herein we resurrect the familial status of the former. Although the trimerous condition of the tarsi is truly a distinctive synapomorphy for the group, the classificatory alteration is based on the fact that Stylotermitinae comprises a grade, along with Archeorhinotermitinae leading to Rhinotermitidae + Serritermitidae + Termitidae (fig. 3). As such, its inclusion within Rhinotermitidae renders the assemblage demonstrably paraphyletic.

INCLUDED GENERA: Stylotermes and Parastylotermes.

ECOLOGY AND EVOLUTION

The circumtropical family Termitidae, or "higher termites," comprises approximately 70% of all termite species and appears to be

one of the most recent radiations of all insect groups that are ecologically significant. Monophyly of the Termitidae is well established; the family includes such familiar groups as the Macrotermitinae and Nasutitermitinae, some of which build huge mounds in grassland and scrub biomes; other nasute taxa build large arboreal nests of cartonlike, fecal material in tropical forests. The diets of Termitidae are extremely diverse, primitively being sound and rotting wood but also including humus, leaf litter, soil, grass, herbivore dung, and even the mycelia of a symbiotic fungus, Termitomyces R. Heim, that they cultivate in the nest like attine ants (e.g., Sands, 1969). The huge colonies and diverse diets of the Termitidae account for the overwhelming biomass of termites in tropical and subtropical environments. The earliest apparent termitid is an incomplete compression of an imago from the Oligocene of Brazil, ca. 30 Ma (Martins-Neto and Pesenti, 2006;, a putative termitid from the Bembridge Marls [Jarzembowski, 1980] is probably a rhinotermitid). The first diverse paleofaunas of Termitidae—more than 30 species—are in Miocene amber from the Dominican Republic (Krishna and Grimaldi, 2009) and Mexico, which are very similar to modern Neotropical faunas. This dramatic appearance is probably due to the fact that these ambers are the only major fossil insect deposits from the Neogene that were formed in the tropics. Were Termitidae abundant during the Eocene they should have been preserved in Baltic amber (Lutetian: ca. 45 Ma), since this deposit has yielded thus far most other living termite families (Engel et al., 2007b; Engel, 2008), as well as species belonging to a diversity of other tropical insect groups (Grimaldi and Engel, 2005). We estimate that Termitidae diverged from Rhinotermitidae sometime in the Early Paleogene (perhaps Late Paleocene or Early Eocene) and subsequently began its diversification in the latest Eocene (Priabonian-Bartonian: ca. 40 Ma) to Early Oligocene, continuing to radiate throughout the remainder of the Neogene and Quaternary.

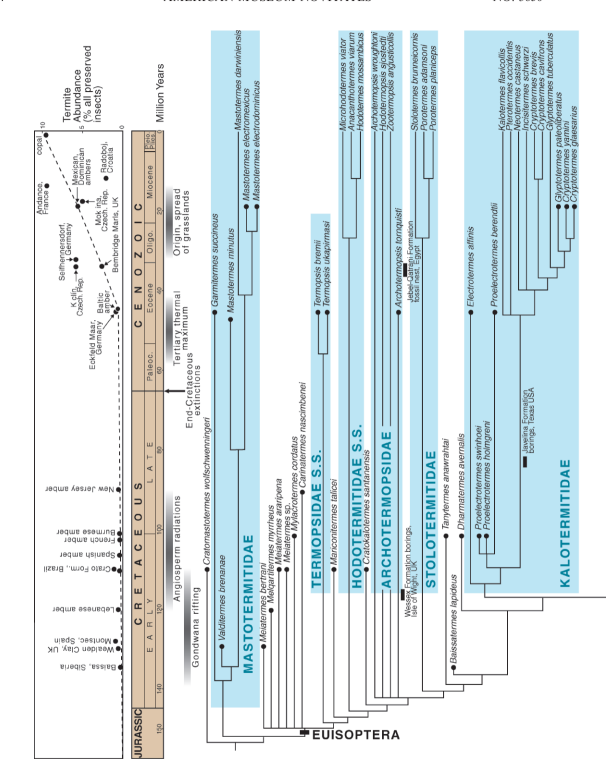
Though Jurassic remains of Isoptera have not been found, Isoptera appear to have diverged from cryptocercid roaches in the Late Jurassic. This would make termites the oldest group of eusocial animals, predating the origins of ants by some 35 million years. Major geological and biotic events in the Cretaceous probably had little effect on termites, since basal divergences appear to have preceded the drift of Gondwanan contiangiosperm radiations. nents and the Unfortunately, the stratigraphic sampling is too poor in the Late Cretaceous and Paleocene to determine any effects of the end-Cretaceous extinctions. The Tertiary thermal maximum of the late Paleocene and Eocene, however, probably had a profound effect on termites, specifically on the global spread of *Mastotermes* and the radiations of living Kalotermitidae and Neoisoptera. The rapid spread of C4 grasslands in the Miocene (Jacobs et al., 1999) doubtless promoted a minor diversification of the harvesters and the explosive diversification of many Termitidae, such as the Macrotermitinae. A macrotermitine nest, in fact, is known from the Miocene of Chad (Duringer et al., 2006).

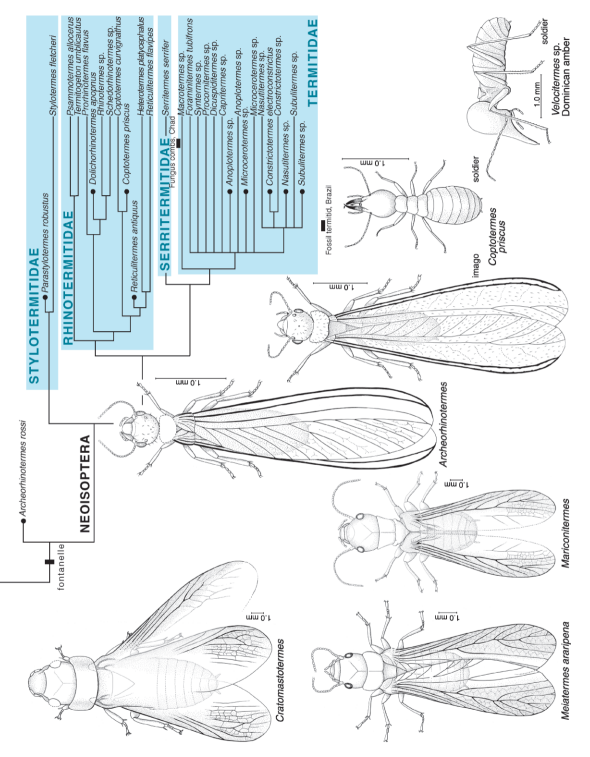
Throughout the Cretaceous and early Tertiary, including the mid-Eocene, termites represented less than 1% of all insect specimens in all fossil deposits (fig. 3). Their abundance rises in the late Eocene to approximately 2%, and then spikes from 5%–10% during the Miocene as both amber and compression fossils, to the present day (in copals, or subfossil resins) (table 2, fig. 3). This spike in abundance is due to the diversification of the Termitidae. The abundance of ants rises dramatically in the Eocene (Grimaldi and Agosti, 2001; Dlussky and Rasnitsyn, 2002), and ants are generally much more abundant in Tertiary insect deposits than are termites, probably because termites feed within the wood where they nest or they travel through tunnels from nest to food sources, so foragers are rarely exposed and imagoes are exposed only during brief nuptial flights. Interestingly, many termitids will forage in the open and these are concomitantly the most abundant termites in Dominican amber and copal.

Our analysis indicates the importance of including fossils in cladistic analyses rather than mapping their putative ages onto molecular-based trees for purposes of dating. The identity of stem groups is obscured in the latter method, resulting in overestimates for divergence times. In our analyses this overestimation is highlighted by the traditional taxonomic placement of Termopsis and most Cretaceous fossils as Termopsidae s.l. and Hodotermitidae, respectively (fig. 3). These taxa in fact represent either a grade to more nested termite lineages or stem groups to some modern families (fig. 3). Using such fossils to calibrate the basal nodes for Hodotermitidae or Termopsidae would result in significant overestimates of the ages of these groups. A failure to distinguish stem groups in an analysis probably explains prior overestimates of the age of other insect lineages (e.g., Moreau et al., 2006; Hunt et al., 2007).

Patterns in termite diversification are very similar to those of the ants (Grimaldi and Agosti, 2001): throughout the Cretaceous both groups were rare and consisted of basal lineages. The diversity and abundance of termites and ants spiked in the Tertiary when speciose groups that form large colonies with highly specialized castes (for subfamilies the Dolichoderinae, Formicinae, and Myrmecinae) eclipsed the smaller colonies of more basal taxa—the concept of "dynastic succession" (Wilson and Hölldobler, 2005). For ants, approximately 70 million years passed from origin to ecological dominance; in termites, this period was 100 million years. Thus, eusociality per se does not result in ecological success, but living in very large colonies with extreme division of labor does. Why, then, did it take so long for large colonies to evolve? We suggest that social evolution is like any other highly adaptive feature, such as the evolution of flight in feathered theropods, and thus may take tens of millions of years to refine.

Similarly, the symbiosis of termites with intestinal protozoa or bacteria which aid their break down of lignocellulose does not alone explain their ecological success as basal termite lineages exhibit the same mutualistic relationship. While their critical role as carbon recyclers is made possible by this symbiosis, this association existed for tens of millions of years before their rise in abundance and diversity (fig. 3).





Strict consensus cladogram of all termite species studied superimposed on geological time scale, with graph of termite abundance. 3 Fig.

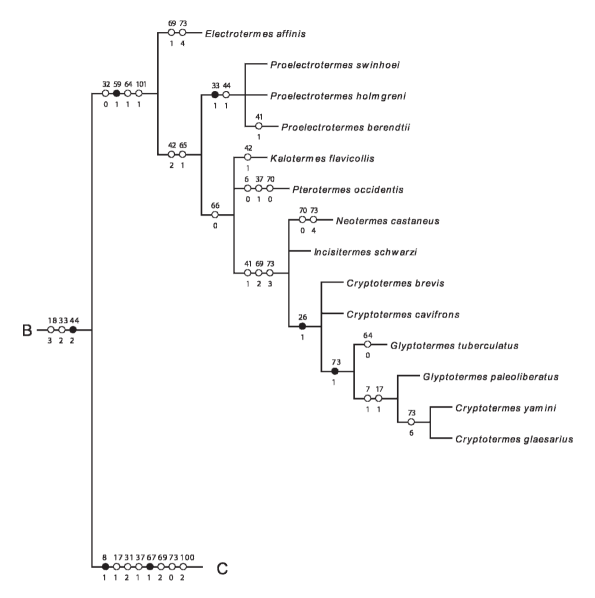


Fig. 4. Continuation of figs. 1-2 focusing on Kalotermitidae. Branch "C" (Neoisoptera) in fig. 5.

One question remains: how was wood decomposed in Mesozoic forests with few or no termites? Patterns of coal and oil deposition suggest that lignocellulose did not rapidly decompose prior to termites and the actions of fungi or other organisms at the time must have been either slow or negligible. Coal is formed from ombrotrophic, or waterlogged, peat (Scott, 1987). While there are some tropical peatlands, such as the coastal "moor" forests of western Borneo and southern Sumatra,

these are dwarfed in area compared to the boreal peatlands of sphagnum and heaths that comprise some 3% of earth's land surface. Tropical ecosystems produce more biomass, but much greater plant detritus accumulates in boreal forests and peatlands, which is traditionally explained by lower boreal evapotranspiration and because seasonality limits decomposition (Scott, 1987). Indeed, boreal peat lands lay hundreds of miles north of the most northern termites in the genus *Reticulitermes*

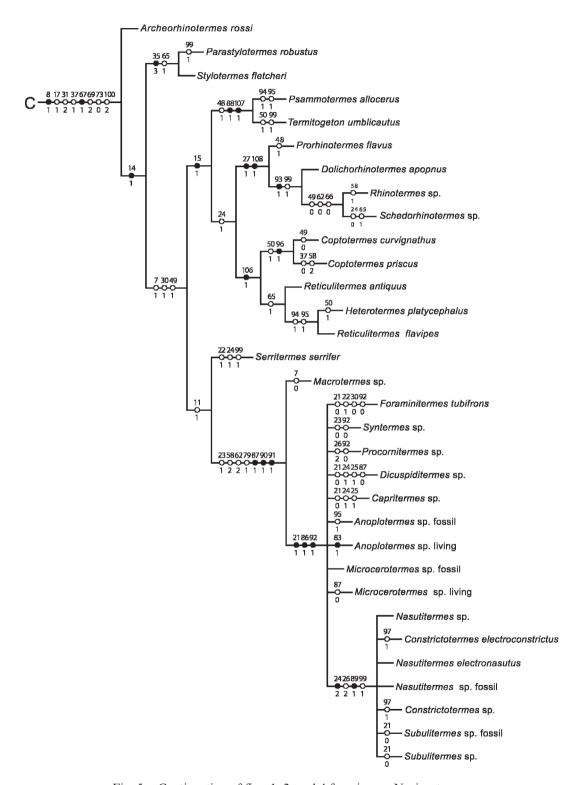


Fig. 5. Continuation of figs. 1, 2, and 4 focusing on Neoisoptera.

Holmgren (Rhinotermitidae). Maximum termite diversity is equatorial, and half of that diversity falls between 18° N and 30° S latitudes; by 48° N and S it is 1%-4% that of the diversity at the Equator (Eggleton, 2000). Tropical ecosystems where termites are most abundant and diverse have notoriously thin humus layers (Richards, 1996). This may explain why coals that were formed prior to the appearance of termites in the Paleozoic and Early Mesozoic, and in largely the same regions and habitats, decomposed less (i.e., contained significantly more vitrain) than Tertiary and modern peats (Shearer et al., 1995; Raymond et al., 2000), as well as the formation of some vast reservoirs of petroleum, like those in the Early Cretaceous Nubian sandstones of present-day Africa and the Middle East. While some Miocene coal formations are astonishingly thick (Shearer et al., 1995), these were formed in palaeoclimates that today would have very few or no termites.

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APPENDIX Data Matrix for Characters Described in Table 1

Below is provided the suite of character-state codings for characters used in the analysis of Isoptera relationships. Within the matrix "-" indicates inapplicable codings (e.g., soldier characters for species which lack this caste); "?" indicates unknown states; "*" indicates a subset polymorphism for character 73 with states 3, 4, 5, 6, and 7; while "\$" indicates a subset polymorphism for character 92 with states 0 and 1

		-	Character	2	4
T	100456700	1	2	3	4
Taxon	123456789	0123456789	0123456789	0123456789	01234
Chaeteessa sp.	010000	001020	1-010	0112100010	00200
Periplaneta sp.	000100	00001000	0-010	0010000010	00200
Panchlora sp.	100100	003-00	0-010	0010000010	00200
Cryptocercus sp. 1	110100	0000?010	0-010	0010000000	0
Cryptocercus sp. 2	110100	0000?010	0-010	0010000000	0
Mastotermes darwiniensis	11111-101	0100000030	1000000-00	0100000010	10100
Anacanthotermes viarum	211100001	1101001011	1000000-00	1222020111	10210
Microhodotermes viator	211100001	1100001011	1000000-00	1122121111	10210
Hodotermes mossambicus	211100001	1101001011	1000000-00	1122021111	10210
Archotermopsis wroughtoni	211111001	1010001021	000000-00	12210011	10110
Hodotermopsis sjostedti	211101	1010001021	1000000	10-11	10110
Zootermopsis angusticollis			1000000-00		
Porotermes adamsoni	311110001	1010001021	1000000-00	1212100111	10110
Porotermes planiceps	311110001	1010001021	1000000-00	1212100111	10110
Stolotermes brunneicornis			1000000-00		
Kalotermes flavicollis	311111001	1100001031	1000000-00	0102120011	10112
Cryptotermes brevis	311111001	1100001031	1000001-00	0102120011	11212
Cryptotermes cavifrons			1000001-00		
Neotermes castaneus			1000000-00		
Incisitermes schwarzi	311111001	1100001031	1000000-00	0102120011	11212
Pterotermes occidentis			1000000-00		
Glyptotermes tuberculatus	311111001	1100001031	1000001-00	0102120011	11012
Serritermes serrifer			1010100000		
Coptotermes curvignathus			1001100000		
Heterotermes platycephalus	31111-111	1010111131	1001100000	1212120111	112
Prorhinotermes flavus			1001100100		
Rhinotermes sp.			1000100100		
Psammotermes allocerus			1000000000		
Schedorhinotermes sp.			1000000100		
Reticulitermes flavipes			1001100000		
Termitogeton umblicautus			10000000		
Foraminitermes tubifrons			1011000000		
Macrotermes sp.			1001000000		
Syntermes sp.			1100000000		
Procornitermes sp.			110-002000		
Nasutitermes sp.			110-2-2000		
Dicuspiditermes sp.			10?1110000		
Capritermes sp.			1001110000		
Baissatermes lapideus			1??????-??		
Valditermes brennenae	?????????	?????????	?????????	??????????	?02?0
Melqartitermes myrrheus			1??????-10		
Meiatermes bertrani			100?-10		
Meiatermes araripena			1???????10		
Mariconitermes talicei	?1?1?????1	???????02?	1??????-00	TTT550005-	10200
Cratomastotermes				04000000	
wolfschwennigeri			1??????-10		
Cratokalotermes santanensis	?1?1????01	???????12?	1??????-00	121????????	?02??

Taxon	123456789	1 0123456789	Character 2 0123456789	3 0123456789	4 01234
Meiatermes sp.	????????	?????????	???????-??	??????????	?0??0
Proelectrotermes swinhoei	3111???01	???????3?	?????????	?????????	10?11
Proelectrotermes holmgreni	????????	?????????	?????????	???112001?	10211
Archeorhinotermes rossi	?1?1???11	?01?0??13?	100????000	?21212011-	112
Mylacrotermes cordatus	?111???01	0??????120	1??????-01	110010011?	1??1?
Dharmatermes avernalis	?1?1???01	?0?0????2?	100????-00	01101?0?1?	10010
Tanytermes anawrahtai	3111???01	??0????12?	101????-10	021012001?	10111
Carinatermes nascimbenei	?111???01	???????20	1??????-10	0100000010	10210
Garmitermes succineus	2101???00	??????030	1??????-00	1100000000	1??0?
Termopsis bremii			1??????-10		
Termopsis ukapirmasi	2111???01	1??????121	1??????-10	0110100011	10010
Archotermopsis tornquisti	3111???01	??????021	0??????-00	12210011	10110
Proelectrotermes berendtii	3111???01	??????03?	1??????-00	0101120011	11211
Electrotermes affinis	3111???01	1??????031	1??????-00	01020011	10012
Reticulitermes antiquus	3111???11	1???1??131	1??????000	12120111	112
Parastylotermes robustus	3111???11	1???1??131	1??????000	02130111	112
Mastotermes electromexicus	2111???01	1??????130	1??????-00	01000010	10100
Mastotermes electrodominicanus					
Mastotermes minutus			1??????-00		
Stylotermes fletcheri	311111011	10101-1131	10000000	0212130111	112
Constrictotermes					
electroconstrictus	311??????	????????1	1????02???	?????????	1????
Nasutitermes electronasutus	311??????	????????1	1????02???	?????????	1????
Dolichorhinotermes apopnus	3111???11	1??????1?1	1??????100	1212120111	112
Cryptotermes yamini	3111??101	1??????131	1??????-00	0102120011	11212
Cryptotermes glaesarius	3111??101	1??????131	1??????-00	0102120011	11212
Glyptotermes paleoliberatus	3111??101	1??????131	1??????-00	0102120011	11012
Coptotermes priscus	31111-111	1???11?131	1001???-00	1212120011	112
Anoplotermes sp. fossil	3111???11	11?????131	1??????000	1212120111	112
Anoplotermes sp. living	3111111	1100101131	11?000	1212120111	112
Microcerotermes sp. fossil			11?????000		
Microcerotermes sp. living	31111-111	1100101131	110-000000	1212120111	112
Nasutitermes sp. fossil	3111??111	110????131	1?0????000	1212120111	112
Constrictotermes sp.	31111-111	1100101131	110-202000	1212120111	112
Subulitermes sp. fossil	?1111-111	1??????31	10?-202000	?2?2120111	1?-??
Subulitermes sp. living	31111-111	1100101131	100-202000	1212120111	112

		Charae	cter	
	4 5	6	7	8
Taxon	56789 0123	456789 01234567	89 012345678	39 012345678
Chaeteessa sp.	0100-00000	000020 1010	-0 000000001	1100000-
Periplaneta sp.	0000- 00	000010 0010	-0 000700001	1100000-
Panchlora sp.	0000-00000	000010 0010	-0 000700001	1100000-
Cryptocercus sp.1				
Cryptocercus sp.2	1		00000	00 11100-
Mastotermes darwiniensis	01000 0110	110010 01000110	00 011610100	00 1210000
Anacanthotermes viarum	01001 0112	121010 13000110	10 011500111	10 1210000
Microhodotermes viator	01001 0111	121010 13000010	10 011500111	10 1210000
Hodotermes mossambicus	01001 0111	121010 13000110	10 011500111	10 1210000
Archotermopsis wroughtoni	-100-0111	121010 130-0-10	-0 011-00111	10 1210000

			Cl		
	4	5	Character 6	7	8
Taxon			0123456789	,	
Hodotermopsis sjostedti			130-0-10-0 13000110-0		
Zootermopsis angusticollis Porotermes adamsoni			13200110-0		
			1320011020		
Porotermes planiceps Stolotermes brunneicornis			1320011020		
Kalotermes flavicollis			1301110020		
Cryptotermes brevis			1301110020		
Cryptotermes cavifrons			1301110022		
Neotermes castaneus			1301110022		
Incisitermes schwarzi			1301110022		
Pterotermes occidentis			1301110020		
Glyptotermes tuberculatus	11000	0111121001	1301010022	1111122110	1210000
Serritermes serrifer	110	0112121100	1311-0112-	1110121110	1210?00
Coptotermes curvignathus			1311001122		
Heterotermes platycephalus			1311011121		
Prorhinotermes flavus			131100-12-		
Rhinotermes sp.			1301000121		
Psammotermes allocerus			13110?-12-		
Schedorhinotermes sp.			1301010120		
Reticulitermes flavipes			1311011120		
Termitogeton umblicautus			131100-12-		
Foraminitermes tubifrons			132100112 - 1321001122		
Macrotermes sp. Syntermes sp.			1321001122		
Procornitermes sp.			1321001122		
Nasutitermes sp.			1321001122		
Dicuspiditermes sp.			1321001122		
Capritermes sp.	11001	0112121120	1321001122	1110121111	1210110
Baissatermes lapideus	1100?	011?12-??0	??0?0?1020	0115??????	???????
Valditermes brennenae			??010?1000		
Melqartitermes myrrheus			??001?1020		
Meiatermes bertrani			?10???1000		
Meiatermes araripena			?10010-000		
Mariconitermes talicei	01000	01-?12-?10	?200111000	0115?1????	???????
Cratomastotermes	01000	01 000 010		011000000	0000 000
wolfschwennigeri			??????100?		
Cratokalotermes santanensis			???????102? ?10???1000		
Meiatermes sp.			130???1020		
Proelectrotermes swinhoei Proelectrotermes holmgreni			130???1020		
Archeorhinotermes rossi			131?0?1122		
Mylacrotermes cordatus			12?010????		
Dharmatermes avernalis			1311001020		
Tanytermes anawrahtai			1300001020		
Carinatermes nascimbenei			1200111020		
Garmitermes succineus			010011????		
Termopsis bremii			1100101000		
Termopsis ukapirmasi			11?1111000		
Archotermopsis tornquisti			130-0-1020		
Proelectrotermes berendtii			130111102?		
Electrotermes affinis	11000	0111121001	1301101021	1114?2????	???????

					Character			
	4	5		6		7		8
Taxon	56789	01234	56789	012	3456789	01234	156789	012345678
Reticulitermes antiquus								???????
Parastylotermes robustus								???????
Mastotermes electromexicus								???????
Mastotermes electrodominicanus								3333300
Mastotermes minutus								<pre>\$3.55555</pre>
Stylotermes fletcheri	11000	01121	21100	131	101112-	11101	.21110	1210?00
Constrictotermes								
electroconstrictus								???011?-0
Nasutitermes electronasutus	?????	?????	?????	355.	???????	?????	??????	???011?-0
Dolichorhinotermes apopnus	11001	01121	21100	131	100112-	1110?	2????	???? ???
Cryptotermes yamini	11000	01111	21001	130	111002-	1116?	2????	???????
Cryptotermes glaesarius	11000	01111	21001	130	111002-	1116?	2????	???????
Glyptotermes paleoliberatus	11000	01111	21001	130	111002-	1111?	2????	???????
Coptotermes priscus	11001	11121	21120	131	100112-	11101	21???	33.50303
Anoplotermes sp. fossil	11001	01121	21120	132	1001122	11101	2????	????????
Anoplotermes sp. living	11001	01121	21120	132	1001122	11101	21111	12111
Microcerotermes sp. fossil	11001	01121	21120	132	1001122	11101	2????	????????
Microcerotermes sp. living	11001	01121	21120	132	1001122	11101	21111	1210100
Nasutitermes sp. fossil	11001	01121	21120	132	1001122	11101	2????	????????
Constrictotermes sp.	11001	01121	21120	132	1001122	11101	21111	1210111-0
Subulitermes sp. fossil								????11??0
Subulitermes sp.	11001	01121	21120	132	1001122	11101	21111	1210111-0

	Character
	1
	8 9 0
Taxon	9 0123456789 012345678
Chaeteessa sp.	0-00000-
Periplaneta sp.	0-00000 000-
Panchlora sp.	0-00000 000-
Cryptocercus sp.1	0-00000 00
Cryptocercus sp.2	0-00000 00
Mastotermes darwiniensis	0 00-0000-00 000000-0-
Anacanthotermes viarum	0 00-00-0-00 001010-0-
Microhodotermes viator	0 00-00-0-00 001010-0-
Hodotermes mossambicus	0 00-00-0-00 001010-0-
Archotermopsis wroughtoni	0 00-00-0-00 021010-0-
Hodotermopsis sjostedti	0 00-00-0-00 001010-0-
Zootermopsis angusticollis	0 00-00-0-00 001010-0-
Porotermes adamsoni	0 00-00-0-00 011010-0-
Porotermes planiceps	0 00-00-0-00 111010-0-
Stolotermes brunneicornis	0 00-00-0-00 121010-0-
Kalotermes flavicollis	0 00-0000-00 111100-0-
Cryptotermes brevis	0 00-0000-00 111100-0-
Cryptotermes cavifrons	0 00-0000-00 111100-0-
Neotermes castaneus	0 00-0000-00 111100-0-
Incisitermes schwarzi	0 00-0000-00 111100-0-
Pterotermes occidentis	0 00-0000-00 111100-0-
Glyptotermes tuberculatus	0 00-0000-00 111100-0-
Serritermes serrifer	0 00-0000-01 221100000

	Character 1			
Taxon	8 9 0 9 0123456789 012345678			
Coptotermes curvignathus	0 0000001-00 221100100			
Heterotermes platycephalus	0 0000110-00 221100100			
Prorhinotermes flavus	0 0000000-00 221100001			
Rhinotermes sp.	0 0001000-01 221101001			
Psammotermes allocerus	0 0000110-00 221100010			
Schedorhinotermes sp.	0 0001000-01 221101001			
Reticulitermes flavipes	0 0000110-00 221100100			
Termitogeton umblicautus	0 0000000-01 221100010			
Foraminitermes tubifrons	0 1100000-10 221100000			
Macrotermes sp.	0 1100000-00 221100000			
Syntermes sp.	0 1100000-10 221100000			
Procornitermes sp.	0 1100000-?0 221100000			
Nasutitermes sp.	1 1110000001 221100000			
Dicuspiditermes sp.	0 11\$0000-00 221100000			
Capritermes sp.	0 1110000-?0 221100000			
Baissatermes lapideus	? ????????? ??????-0-			
Valditermes brennenae	? ????????? ??????-?-			
Melqartitermes myrrheus	? ???0-????0 00????-0-			
Meiatermes bertrani	? ????0???? ??????-0-			
Meiatermes araripena	? ???????? ??????-0-			
Mariconitermes talicei	? ???0???? ??????-?-			
Cratomastotermes wolfschwennigeri	? ???00????? ??????-?-			
Cratokalotermes santanensis	? ????0???? ??????-?-			
Meiatermes sp.	? ???0-????? ??????-?-			
Proelectrotermes swinhoei	? ????-0???? ??????-0-			
Proelectrotermes holmgreni	? ????-????? ?1????-?-			
Archeorhinotermes rossi	? ??00-0???0 22??????0			
Mylacrotermes cordatus	? ??-0???1 21????-?-			
Dharmatermes avernalis	? ??-0???? ?0????-?-			
Tanytermes anawrahtai	? ??-0???1 12????-?-			
Carinatermes nascimbenei	? ??-000???0 11????-?-			
Garmitermes succineus	? ??-000???0 00????-?-			
Termopsis bremii	? ??-00-???0 01????-0-			
Termopsis ukapirmasi	? ??-00-???0 01????-0-			
Archotermopsis tornquisti	? ??-00-???0 02????-0-			
Proelectrotermes berendtii	? ??-000???0 11????-?-			
Electrotermes affinis	? ??-000???0 11????-0-			
Reticulitermes antiquus	? ??0000???0 22????100			
Parastylotermes robustus	? ??0000???1 22????000			
Mastotermes electromexicus	? ??-000???0 00????-0-			
Mastotermes electrodominicanus	0 0?-0000-?0 00???0-0-			
Mastotermes minutus	? ??-000???0 00????-0-			
Stylotermes fletcheri	0 0000000?00 221100000			
Constrictotermes electroconstrictus	1 1?????01?1 22???00-0			
Nasutitermes electronasutus	1 1?????00?1 22???00-0			
Dolichorhinotermes apopnus	? ??0100???1 22?????0?			
Cryptotermes yamini	? ??00???0 11????-0-			
Cryptotermes glaesarius	? ??00???0 11????-0-			
Glyptotermes paleoliberatus	? ??00???0 11????-0-			
Coptotermes priscus	0 0?00001??0 22???0100			

	Character 1				
Taxon	8 9 0 9 0123456789 012345678				
Anoplotermes sp. fossil	? ??1001???0 22?????0?				
Anoplotermes sp. living	1100010 221100000				
Microcerotermes sp. fossil	? ??1000???0 22?????0?				
Microcerotermes sp. living	0 11\$0000-?0 221100000				
Nasutitermes sp. fossil	? ??1000???1 22?????0?				
Constrictotermes sp.	1 11100001?1 221100000				
Subulitermes sp. fossil	1 1?1??????1 22???00??				
Subulitermes sp.	1 1110000001 221100000				

