Convergent evolution of chemical defense in poison frogs and arthropod prey between Madagascar and the Neotropics

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With few exceptions, aposematically colored poison frogs seques-
ter defensive alkaloids, unchanged, from dietary arthropods. In the
Neotropics, myrmicine and formicine ants and the siphonothrin millipede Rhinotus purpureus are dietary sources for alkaloids in
dendrobatid poison frogs, yet the arthropod sources for Mantella
poison frogs in Madagascar remained unknown. We report GC-MS
analyses of extracts of arthropods and microsympatric Malagasy
poison frogs (Mantella) collected from Ranomafana, Madagascar.
Arthropod sources for 11 “poison frog” alkaloids were discovered,
7 of which were also detected in microsympatric Mantella. These
arthropod sources include three endemic Malagasy ants, Tetramo-
rhrium electrum, Anochetus grandidieri, and Paratrechina amblyops
(subfamilies Myrmicinae, Ponerinae, and Formicinae, respectively),
and the pantropical tramp millipede R. purpureus. Two of these ant
species, A. grandidieri and T. electrum, were also found in Mantella
stomachs, and ants represented the dominant prey type (67.3% of
609 identified stomach arthropods). To our knowledge, detection
of 5,8-disubstituted (ds) indolizidine iso-217B in T. electrum repre-

tsents the first izidine having a branch point in its carbon skeleton

to be identified from ants, and detection of 3,5-ds pyrrolizidine
251D in A. grandidieri represents the first ponerine ant proposed
as a dietary source of poison frog alkaloids. Endemic Malagasy ants
with defensive alkaloids (with the exception of Paratrechina) are
not closely related to any Neotropical species sharing similar
chemical defenses. Our results suggest convergent evolution for
the acquisition of defensive alkaloids in these dietary ants, which
may have been the critical requisite for subsequent conver-
gence in poison frogs between Madagascar and the Neotropics.

alkaloid occurrence | dietary sequestration | nicotine

Amphibians use a variety of skin chemicals for protection
against predators, and anurans that store lipopholic basic
alkaloids in granular glands are collectively termed “poison frogs”
(1–3). Poison frogs have been recognized in four families: South
American Melanophryniscus (Bufonidae); Australian Pseudophryn-

edes (Myobatrachidae); Central and South American Dendrobate-

tas, Epipedobates, and Phyllobates (Dendrobatidae); and Malagasy
(Madagascar) Mantella (Mantellidae) (1, 4–8). Additionally, trace
amounts of such alkaloids have been detected in a Thai ranid frog,
Limnonectes kuhli (9). More than 500 frog skin alkaloids belonging
to 25 structural classes have so far been categorized, with each
coded by a boldface number representing the nominal mass and a
letter to distinguish among alkaloids of the same molecular weight
(MW code) (1).

Although wild poison frogs retain skin alkaloids for several years
in captivity (10–12), they do not seem to produce alkaloids; rather,

it seems that they sequester and accumulate such toxins from
dietary arthropods by using an as-yet-uncharacterized alkaloid
takeup system (11–13). Alkaloids are absent in dendrobatid and
Mantella frogs raised in captivity on a diet of Drosophila, but these
individuals will readily accumulate alkaloids added to their diet
(10–15). Some anurans also have the ability to modify ingested
alkaloids; pumiliotoxin 307A was metabolized by one species of the


genus Pseudophryne (8), and pumiliotoxin (+) 251D was efficiently
and stereoselectively hydroxylated by Dendrobates spp. into allopu-

miliotoxin (+)-267A, which is five times more toxic (14). The only

known example of direct alkaloid production is in the Australian
Pseudophryne frogs, which seem to be capable of synthesizing
indolic pseudophyridines (8).

Recently, the putative dietary sources of representative alkaloids
of several structural classes of “poison frog alkaloids” were re-
ported; these dietary arthropods include beetles, ants, and milli-
pedes. Dendrobatid and bufoinid frogs and coccinellid beetles all
share precoccinelline (193C), suggesting that these beetles repre-
sent a dietary source of this coccinelline-like tricyclic alkaloid and
others like it (1, 13, 16, 17–21). Poison-dart frogs, genus Phyllobates
of the Neotropics, contain highly toxic steroidal alkaloids, the
brachotoxins (22), as do passerine birds (23–24) and putative
dietary melyrid beetles of Papua New Guinea (25); all are brightly
colored.

Regarding ants as sources of dietary alkaloids, pumiliotoxins
were recently detected in formicine genera Brachymyrnex and
Paratrechina, where they occurred microsympatrically with the
poison frog Dendrobates pumilio in Panama (26). Poison frogs and
Neotropical ants of the subfamily Myrmicinae share several classes
of alkaloids, including 2,5-disubstituted (ds) pyrrolidines, 2,6-ds
piperidines, 3,5-ds pyrrolizidines, 3,5 ds indolizidines, 4,6-ds quino-

lizidines, and 2,5-ds decahydroquinolines (1, 13, 16, 27). Reports of
alkaloids from African ants are limited to the subfamily Myrmici-
dae: 2,5-dialkylpyrrolidines and 1-pyrrolines were detected in
Monomorium in South Africa and Kenya (28), and substituted pyrazine alkaloids detected in Eutetramorium mocquersyi, a genus
demic to Madagascar (29). Two other Malagasy myrmicine ants
of the genus Metapone use methyl pyrrole-2-carboxylate as a trail
pheromone and also contain pyrazines (30).

Seven alkaloids of the spiropyrrolizidine (SpiroP) class have been
detected in poison frogs; three of these alkaloids have also been
reported from two millipede species: (i) polyzonicine (151B) and
nitropolyzonicine (238) from Petesirpes cryptocoefalum (McNeill
1887) [Polyzoniiidae: Polyzoniiida, often misidentified as Polyzonium
rosalum (Cope 1879)] (31; see p. 32 of ref. 32) of Ithaca, New York
(33–34) and (ii) the 238 and SpiroP O-methylxizidine 236 from the
widespread Rhinotus purpureus (Pocock 1894) (Siphonotidae: Po-
yzoniiida) that occurs sympatrically in Panama with the poison frog
D. pumilio (35). Other alkaloids detected in millipedes include
glomerins (quinazolinones) from the Glomeridae (36–37) and the

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Abbreviations: ds, disubstituted; SpiroP, spiropyrrolizidine; Saha, Sahavondrona; Vato,
Vatomaranana; GCT, GC-TOF mass spectrometer; TAS, transcutaneous amphibian
stimulator.

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terpenoid alkaloid buzonamine from the Polyzoniidae, genus *Buzonium* (38).

More than 100 alkaloids of 12 classes have been detected in skin of *Mantella* (1, 4, 6, 39–40), and many also occur in Neotropical poison frogs. In addition to their ability to sequester and accumulate alkaloids from diet, *Mantella* and some dendrobatid frogs also share the following features: terrestrial eggs, small body size (<50-mm snout-vent length), toothless jaws, a specialist diet composed largely of ants, active diurnal foraging behaviors, and aposematic coloration; all features are considered to have been produced by convergent evolution (4, 41–45). In both groups, sequestered defensive chemicals appear to be closely associated with (i) the evolution of aposematism, as a visual warning of their toxicity to potential predators, and (ii) active diurnal foraging, a behavior that is generally rare in frogs (46). Although the foraging behavior and diet of *Mantella* is not yet well documented relative to dendrobatids (46–52), ants are known to dominate the diet of *Mantella* (42–43). A study of 774 prey items taken from the stomachs of 15 *Mantella* specimens of four species found that ants represented 74% of the total prey and that all prey items were <5 mm in length (43).

However, to date there had been no studies concerning arthropods as potential sources of frog skin alkaloids in Madagascar, and the potential convergence of alkaloid defenses in arthropod groups between the Neotropics and Madagascar had not yet been investigated. Here, we report results of our alkaloid survey conducted in and around Ranomafana National Park (in Fianarantsoa Province, southeast Madagascar) that targeted both *Mantella* poison frogs and potential dietary microsympatric leaf-litter arthropods.

### Materials and Methods

#### Field Collections

*Mantellid* frogs and arthropods were collected within and around Ranomafana National Park. Four collecting sites (all within 15 m of 2- to 4-m-wide streams) were surveyed: Vohiparara, 21°13.587’ S, 47°22.193’ E; Sahavondrona (Saha), 21°15.450’ S, 47°21.609’ E; Valoharanana (Vato), 21°17.444’ S, 47°25.569’ E; and Ampasimpoty, 21°28.796’ S, 47°33.424’ E, with sampling conducted during the latter part of the rainy season (March 13 to May 1, 2003). *Mantella* were photographed, killed with chloroform, and skinned into 100% methanol, and their bodies were fixed in 10% formalin within 1 h of capture to preserve stomach contents. In addition, a transcutaneous amphibian stimulator (TAS) (53) was used to obtain skin exudates from live frogs, which were then preserved. All frog voucher specimens have been deposited at the Department of Animal Biology at University of Antananarivo and at the American Museum of Natural History; frog stomach contents were removed for subsequent identification.

All *Mantella* capture sites were also surveyed for leaf-litter arthropods to produce samples for alkaloid analyses. Forceps-mediated collections were made by searching through leaf-litter on white cloth and removing arthropods with entomological forceps. Arthropod specimens were put in methanol in taxon-specific tubes, white cloth and removing arthropods with entomological forceps. Mediated collections were made by searching through leaf-litter on the same defensive alkaloids. (a) *M. madagascariensis* (shown actual size). (b) *M. bernhardi*. (c) *M. baroni*. (d) *R. purpureus*. (e) *T. electrunc*. (f) *A. grandidierii*. (Scale bars, 1 mm.)

Supelco Equity 5 column with 0.25-μm film thickness was used for all GCT injections, and the oven temperature of the GCT was increased at 10°C/min from 100°C to 280°C. GCT calibration samples supplied by John Daly (National Institutes of Health, Bethesda) for *Mantella baroni* (Saha, January 1993; Vato, December 1989) provided reference retention times for known alkaloids (4). Methanolic frog skin alkaloid fractions were prepared for individual frogs following the methodology of ref. 11. A volume of 1 μl, corresponding to 1 mg of wet weight frog skin, was injected for alkaloid fractions from *Mantella* skins, whereas 4 μl was injected for each TAS frog extract. Arthropod extracts were injected without work-up into alkaloid fractions. Alkaloids were identified by using the MS library of ref. 1.

### Results

Three species of *Mantella* poison frog were recorded during this survey of Ranomafana: *M. baroni*, *Mantella bernhardi*, and *Mantella madagascariensis* (Fig. 1). GC-MS analyses of the methanol extracts obtained from 22 individual *Mantella* representing these three species (18 skins and 4 TAS extracts) recovered 80 coded alkaloids (1). Additionally, nicotine (which we code as 162), previously undetected in any frog species, was detected in *M. baroni* of Saha. The occurrence of these 81 alkaloids in *Mantella* are given in Table 3, which is published as supporting information on the PNAS website, and the 33 alkaloids that were previously undetected in *Mantella* are highlighted; 9 of these 33 *Mantella* alkaloids are isomers not previously reported in frogs. Eleven of these 81 alkaloids are now known from a specific Malagasy arthropod source. TAS extracts yielded detectable alkaloids similar in diversity compared with skin extracts. The distributions of arthropod source alkaloids in individual *Mantella* frogs are presented by locality in Table 1.

Analyses of the 154 extracts of arthropod morphospecies samples (65 ants and 89 others) detected 11 known poison frog alkaloids...
Table 1. Distribution of selected alkaloids in individual *Mantella* frogs at four survey sites in the Ranomafana region of Madagascar

<table>
<thead>
<tr>
<th>Alkaloid class and MW code (1)</th>
<th><em>M. baroni</em></th>
<th><em>M. mad.</em></th>
<th><em>M. bernhardi</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Saha</td>
<td>Vato</td>
<td>Vohi</td>
</tr>
<tr>
<td>Spiropyrrrolizidines</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>151B</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>222</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>236</td>
<td>2</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Iso-236</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>238</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>3,5 Indolizidines</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>cis-223H</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Iso-223H(^1)</td>
<td>—</td>
<td>—</td>
<td>3</td>
</tr>
<tr>
<td>251O</td>
<td>4</td>
<td>3*</td>
<td>8</td>
</tr>
<tr>
<td>3,5 Indolizidine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>195B</td>
<td>—</td>
<td>—</td>
<td>3</td>
</tr>
<tr>
<td>5,8 Indolizidines</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>217B</td>
<td>4</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>217B(^3) (third isomer)</td>
<td>1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Pumiliotoxins</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>307A</td>
<td>—</td>
<td>1</td>
<td>—</td>
</tr>
<tr>
<td>Picrococcinelline</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>193C</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Skin samples</td>
<td>3</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>TAS samples</td>
<td>1</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Total frogs</td>
<td>4</td>
<td>3</td>
<td>8</td>
</tr>
</tbody>
</table>

\(^{-}\) —, not detected; Vohi, Vohiparara; *M. mad.*, *M. madagascariensis*.

\(^{1}\) Data are combined for three additional isomers of 223H with earlier retention times in *Mantella* than the ant cis-223H, which has the same retention time as cis-223H in Daly's Saha *Mantella* standard (see Table 3).

(Tables 2 and Fig. 2); 7 of these arthropod alkaloids were also detected in microsymptactic *Mantella* frogs. However, alkaloids were generally rare among these arthropod samples, with at most five taxon-specific ant samples and one millipede sample yielding detectable products (one ant morphospecies sample later proved to contain two species). These alkaloids were confined to ant species representing three different subfamilies (Formicinae, Myrmicinae, and Ponerinae) and one millipede (*R. purpureus*; family Siphoniulidae). The negative alkaloid ant samples have not yet been identified, but the negative millipede samples (all Pachybolidae) included two pantropical species, *Trigonitius corollinus* (Gervais 1847) and *Lepiogramma sororius* (Butler 1876) and a Malagasy endemic *Aphistogoniulus* sp. (Silvestri) (32, 57, 58).

A single female specimen of the synanthropic millipede *R. purpureus*, collected from Vato, yielded five alkaloids of the SpiroP class (Table 2 and Fig. 2). Areas under the chromatograms reveal the following relative concentrations: 2% 151B, 17% 236, 77% iso-236, 3% 238, and 1% 254. The previously undetected isomer, iso-236, eluted 0.16 min after 236, but the fragmentation pattern was the same for both. The configuration of the oxime group of 236 was assigned as syn (Z) based on nuclear Overhauser effect NMR experiments (59).

The ant species *Tetramorium electum* Bolton (subfamily Myrmicinae, tribe Tetramorini; Fig. 1) collected from Ampasimpoty yielded a branched chain isozidine: 5,8-ds indolizidine iso-217B (Fig. 2). *Anochetus grandidieri* Forel (subfamily Ponerinae, tribe Ponerini; Fig. 1) collected from Vato contained 3,5-ds pyrrolizidine 251O (Fig. 2). *Paratrechina amyllops* (Forel) (Formicinae, Plagioplectridi) collected from Vato contained a variety of alkaloids, including 3,5-ds pyrrolizidines 195F and 223H, 2,5-ds pyrrolidine 197B (Fig. 2), and other uncharacterized alkaloids. Another 2,5-ds pyrrolidine, 225C, was detected in a mixed sample of larger (>7-mm length) black ants from Saha: Pachycondyla camboieu (Forel) (Ponerinae, Ponerini) and *Camponotus* sp. 1 (Formicinae, Camponotini).

Analyses of the stomach contents of 21 *Mantella* specimens yielded 609 identified arthropod specimens (Table 4, which is published as supporting information on the PNAS web site), with the largest fraction (67.3%) representing ants (29 species of nine genera). The remaining stomach arthropods included 12.2% Acari (mites), 5.5% Collombola (springtails), 4.1% Amphipoda (lawn shrimps), 4.1% various larva, 2.5% Coleoptera (beetles), and 9.3% other arthropod groups, including one millipede individual identified as a pantropical species, *Prosopodesmus jacobseni* (Silvestri 1910) (Haplonedinidae: Polydesmida). The proportion of ants in the arthropod stomach samples ranged from 18% to 93% among the 21 *Mantella* frogs (mean = 63%). The 251O alkaloid-containing ant *A. grandidieri* was found in stomachs of *M. madagascariensis* of Vohiparara and *M. baroni* of both Saha and Vato. Nine species of *Tetramorium* ants were found (10% of the total ingested ants), including the iso-217B-containing *T. electum* in *Mantella* at Saha.

Discussion

Previously Undetected Alkaloids in *Mantella*. To our knowledge, our results represent the first report of four SpiroPs for *Mantella*: 151B, 222, iso-236, and 238 (Fig.2). Two SpiroPs, 236 and 252A, have been previously reported in *M. baroni* (4), and 252B has been reported in Australian *Pseudophryne* frogs (5). Ant alkaloid 3,5-ds indolizidine 195B and beetle alkaloid precocinelline 193C, both of which have also been reported in Neotropical dendrobatid and bufonid poison frogs, were detected here for what we believe to be the first time in poison frogs of Madagascar (Tables 1–3). A total of 23 known coded poison frog alkaloids (1), plus nine other isomers and nicotine 162, were not previously known from mantellids (1) but were detected in *Mantella* of our study (Table 3). To our knowledge, our finding of 2,6-dipiperidone 197E in *M. baroni* of Vato represents the first report of this class in *Mantella* (Table 3), thus demonstrating that *Mantella* can indeed sequester this alkaloid class.

The well known plant alkaloid nicotine (162) was detected in a single *M. baroni* specimen, representing what we believe to be the first report of nicotine from any frog source and the 14 th class of alkaloids (pyridyls) known from *Mantella*. However, other pyridyl alkaloids (e.g., the epibatidines and noranabasamine) have been reported from dendrobatid poison frogs (1). Nicotine is biosynthesized by a variety of plants of the family Solanaceae (60) and can also be sequestered by insects (61) and biosynthesized by lepidopteran larvae (62). However, we have not yet detected nicotine in any of our Malagasy arthropod extracts, and it is unknown to us whether suitable nicotine source plants occur at Saha. Still, our nicotine discovery suggests a sequestration food chain where nicotine is sequestered by an herbivorous arthropod subsequently eaten by *Mantella*. We also detected in these *Mantella* several other previously undetected alkaloids, not yet coded.

Individual Variation in *Mantella* Frogs. Our results show variation in alkaloid profiles between individual *Mantella* frogs of the Ranomafana region, within the same species, and at the same collecting site. However, at Vohiparara, the only site that included two sympatric *Mantella*, six of eight alkaloids found in each species were also present in the other sympatric species (Table 1). This finding tentatively suggests that alkaloid profiles of microsymptactic *Mantella* species may be quite similar and that *Mantella* alkaloid profiles, in general, may be more heavily influenced by the distribution of local arthropod alkaloid sources rather than ecological or evolutionary differences between *Mantella* species.

The individual variation seen for frog alkaloid profiles, coupled with the observation that *Mantella* retain alkaloids in their skin for years in captivity (12), suggests that some arthropod sources for...
alkaloids are rare prey items for *Mantella*. Rare prey types are also evident in our *Mantella* stomach content data (Table 4); for example, just one millipede is represented from this sample of 609 arthropods. Presumably, individual frogs missing alkaloids (otherwise represented in local populations) have never, or perhaps rarely, ingested the required source arthropod prey over the duration of their lifespan. Our own efforts at identifying arthropod sources available to these frogs (Table 2). Three of the four alkaloids missing in the frogs (254, 197B, and 195F) were found in two arthropod species (Table 2). *Mantella* can sequester other alkaloids of these classes, so we would expect that these alkaloids would be sequestered if ingested. An examination of trace alkaloids in Daly’s Vato standard revealed trace amounts of 197B (not in ref. 4); 197B was also present in our P. amblyops ant of Vato. The absence of 225C in Ranomafana *Mantella*, which was detected in a sample of large black ants (*Pachycondyla cambouei*, length = 10.7 mm and *Camponotus* sp., length = 7.9 mm) may reflect these ants being too large to serve as potential prey. Vences and Kniel (43) reported all *Mantella* prey as <5 mm in length, and the maximum sized prey specimen we recorded was a lepidopteran larva that was 4.7 mm long.

The discovery of seven alkaloids shared among microsympatric *Mantella* poison frogs and four leaf-litter arthropod species (Tables 1 and 2) provides data to identify potential dietary sources for alkaloids in Malagasy poison frogs. These findings are also partly corroborated by the stomach content data: two of the potential ant source species, *A. grandieri* and *T. electrum*, were found in the stomachs of the microsympatric *Mantella* (Table 4).

Surprisingly, the probable dietary source of the *Mantella* SpiroPs at Ranomafana appears to be the same millipede species, *R. purpureus*, as reported for dendrobatid poison frogs in Panama (35). This invasive tramp millipede has a pantropical distribution and, in

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**Table 2. The occurrence of alkaloids in arthropods and *Mantella* frogs recorded in this study with a comparison to published data for other *Mantella* in Madagascar and poison frogs and arthropods in the Neotropics**

<table>
<thead>
<tr>
<th>Alkaloid class and family</th>
<th>MW code</th>
<th>Arthropod species and family</th>
<th><em>Mantella</em> species</th>
<th>Other <em>Mantella</em></th>
<th>Arthropod species and family</th>
<th>Frog families</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spiropyrolizidines</td>
<td>151B</td>
<td><em>R. purpureus</em> S</td>
<td>1, 2, 3</td>
<td>—</td>
<td>—</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>222</td>
<td>—</td>
<td>3</td>
<td>—</td>
<td>—</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>236</td>
<td><em>R. purpureus</em> S</td>
<td>1, 3</td>
<td>4</td>
<td><em>R. purpureus</em> S</td>
<td>D, B</td>
</tr>
<tr>
<td></td>
<td>iso-236</td>
<td><em>R. purpureus</em> S</td>
<td>1, 3</td>
<td>—</td>
<td><em>R. purpureus</em> S</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>238</td>
<td><em>R. purpureus</em> S</td>
<td>3</td>
<td>—</td>
<td>—</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>254</td>
<td><em>R. purpureus</em> S</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>D</td>
</tr>
<tr>
<td>2,5 Pyrrolizidines</td>
<td>197B</td>
<td><em>Paratrechina amblyops</em> F</td>
<td>—</td>
<td>4</td>
<td><em>Monomorium pharaonis</em>, <em>Megalomyrmex gaeldii</em>, <em>Solenopsis punctaticeps</em> M</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>225C</td>
<td><em>Pachycondyla cambouei</em> P and/or <em>Camponotus</em> sp. F</td>
<td>—</td>
<td>4</td>
<td><em>Monomorium indicum</em>, <em>Megalomyrmex foreli</em>, <em>Solenopsis fugax</em>, <em>S. punctaticeps</em> M</td>
<td>D</td>
</tr>
<tr>
<td>3,5 Pyrrolizidines</td>
<td>195F</td>
<td><em>Paratrechina amblyops</em> F</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>223H</td>
<td><em>Paratrechina amblyops</em> F</td>
<td>1</td>
<td>4, +</td>
<td><em>Solenopsis</em> sp. M</td>
<td>D, B</td>
</tr>
<tr>
<td></td>
<td>231O</td>
<td><em>A. grandieri</em> P</td>
<td>1, 2, 3</td>
<td>4, +</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>3,5 Indolizidines</td>
<td>195B</td>
<td>—</td>
<td>1</td>
<td>—</td>
<td><em>Monomorium pharaonis</em> M</td>
<td>D, B</td>
</tr>
<tr>
<td>5,8 Indolizidines</td>
<td>217B</td>
<td>—</td>
<td>1, 2, 3</td>
<td>4, +</td>
<td>—</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>iso-217B</td>
<td><em>T. electrum</em> M</td>
<td>2</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Pumilioxotins</td>
<td>307A</td>
<td>—</td>
<td>1, 2, 3</td>
<td>4, +</td>
<td><em>Paratrechina steinhelli</em>, <em>Brachymyrmex spp.</em> F</td>
<td>D</td>
</tr>
<tr>
<td>Coccineiline-like tricyclics</td>
<td>193C</td>
<td>—</td>
<td>2</td>
<td>—</td>
<td><em>Coccinella septempunctata</em>, <em>Coleomegilla maculata</em>, <em>Chauliognathus pulcherellus</em> A*</td>
<td>D, B</td>
</tr>
</tbody>
</table>


*These beetles were not collected from the Neotropics; see refs. 18–21.
contrast to the endemic Malagasy millipedes, likely represents a relatively recent arrival in Madagascar mediated by accidental human introduction. The two reported records (58) for Madagascar, Toamasina (Tamatave) and Nosy Be, are both historical and current major trading ports. Although the genus Rhipinus is in need of revision, it is assumed that Neotropical and Afrotopical specimens of the genus are conspecific (63).

Although five SpiroPs were recorded in R. purpureus, the majority of Mantella had no more than two of these alkaloids, and in Panama, only two SpiroPs were found in R. purpureus and sympatric poison frogs (35). A possible explanation for this variability concerns the age of the millipedes and the environmental availability of compounds sequestered from food plants; Meinwald et al. (33) have suggested (for another Polyzoniida millipede) that a plant-origin pyrrolizidine may be sequestered, transformed into the spirocycle, and subsequently metabolized into 151B. The elongate mandibles of Rhipinus are suited for scraping (see figure 3 in ref. 63), and it is assumed that this species scrapes roots and shoots for plant juices (p. 819 of ref. 64). By contrast, spirobolid and spirostrepid millipedes that in our study did not yield detectable plant juices (p. 819 of ref. 64). This finding suggests that Neotropical and Afrotopical species of the genus are conspecific (63).

To our knowledge, our study reports the first known occurrence of six alkaloids for endemic Malagasy ants: 3,5-ds pyrrolizidines (195F, 223H, and 251O), 2,5-ds pyrrolizidines (197B and 225C), and the 5,8-ds indolizidine iso-217B. This latter finding also represents what we believe to be the first record of this alkaloid class for any ant species; previously, other 5,8-ds indolizidines were detected in mixed arthropod samples in Panama (68). However, only the iso-217B was detected in Malagasy ants, whereas both 217B and iso-217B were detected in Mantella. The occurrence of a 3,5-ds pyrrolizidine (251O) in A. grandidieri also represents, to our knowledge, the first ponerine ant to be proposed as a dietary source of poison frog alkaloids; however, Anochetus kempi of Puerto Rico contains a phenylpyrrole (69) not yet known from frogs.

The origin of defensive ant alkaloids is not yet clear; alkaloid biosynthesis has not yet been studied for any ant species endemic to Madagascar, and, globally, only tetraponerine and solenopsin ant alkaloids have so far been studied (ref. 70 and references therein). The ants produced these compounds themselves; however, it is also possible that ants sequester alkaloids from plants, as is known for complex pyrrolizidines in lepidopterans and coleopterans (71–72), or even that microsymbionts might serve a role (26). The alkaloids sequestered by poison frogs serve primarily as passive chemical defenses, and, presumably, these compounds also serve similar defensive functions for the source ant species.

Convergent Evolution Between Madagascar and the Neotropics. A remarkable feature of the 16 coded alkaloids we report here in the arthropods and Mantella frog species of Ranomafana (Madagascar) is that 13 of them are also known in other ants, beetles, and frogs endemic to the Neotropics (Table 2; see also Table 3). Excluding R. purpureus (a likely recent invasive tramp) and possibly Paratrechina (phylogenetic relationships within this globally distributed genus remain uncertain), none of the other Malagasy and Neotropical endemic species that share these types of alkaloids are closely related to each other.

Three alkaloids (197B, 225C, and 223H) are each shared between at least one species of formicine or ponerine ant endemic to Madagascar and one species of myrmicine ant endemic to the Neotropics (Table 2 and refs. 27 and 73–75). These Malagasy and Neotropical ant species are classified in different subfamilies, and even at the subfamily level, these groups are not closely related based on molecular and morphological phylogenetic studies (76–77). Considering all Neotropical and Malagasy ant species known to contain defensive alkaloids shown in Table 2, just one genus (Paratrechina) is represented in both regions. Another ant species, Monomorium pharaonis, is a widespread tramp that could be of African origin (75) and could potentially serve as an alkaloid source in both Madagascar and the Neotropics; however, we did not recover any alkaloid-containing Monomorium in our survey (Table 2). The phylogenetic distribution of these defensive alkaloid-bearing species thus is suggestive of independent evolution of sequestration and/or biosynthesis within the endemic ant radiations of Madagascar and the Neotropics. Within each of these three ant subfamilies, the rarity of species containing these defensive alkaloids is striking. Very few ant species actually yield defensive alkaloids: only 4 of our 65 ant samples were positive, and similar low hit successes have also been reported for Neotropical ants and arthropods [e.g., 4 positive of 61 ant samples (16) and 3 formicine ant species containing pumiliotoxins from 512 arthropod samples (26)].

Of the 12 alkaloids we report here in Mantella from Ranomafana, 9 also occur in dendrobatid poison frogs (Dendrobates, Epipedobates, and Phylllobates), and 4 occur in bufonid toads (Melanophryniscus) of the Neotropics (see Table 2). The Malagasy Mantella species (Mantellidae) are not closely related to these Neotropical groups (Dendrobataeae and Bufonidae) but fall within a paraphyletic Mantidactylus group that itself is sister to other Malagasy endemic mantellid genera, and then rhacophorids (78–79). These shared alkaloids are unknown for all other genera within Mantellidae, Dendrobatidae, and Bufonidae. There is no doubt that alkaloid sequestration represents an apomorphic trait within each family and that the evolution of alkaloid sequestration for poison frogs in Madagascar and the Neotropics has independently occurred.
multiple times (4, 45). Evolutionary convergence of defensive alkaloid sequestration, similarly, has also driven evolutionary convergence for aposematic colorations, presumably in response to similar selective pressures from frog predators to this form of chemical defense. For Neotropical and Malagasy frogs, the convergent evolution of alkaloid uptake systems for chemical defense requires first the availability of suitable alkaloids in prey species to form toxic alkaloids and second the independent origin of mechanisms for achieving alkaloid sequestration. Because ants represent a common or even dominant part of poison frog diets (Table 4 and refs. 43, 49–52, and 80), the presence of suitable alkaloids in ants may have been the critical prerequisite for the evolution of alkaloid chemical defense in Malagasy and Neotropical poison frogs. Our demonstration that endemically toxic ants do indeed contain these toxic alkaloids supports this view and further suggests that the convergence seen in the poison frogs might itself have been first driven by convergent evolution in the endemic ant radiations of Madagascar and the Neotropics.

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