

Dietary niche differentiation among three species of invasive rodents (*Rattus rattus*, *R. exulans*, *Mus musculus*)

Aaron B. Shiels · Caitlin A. Flores ·
Arthur Khamsing · Paul D. Krushelnycky ·
Stephen M. Mosher · Donald R. Drake

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Abstract The diets of sympatric rodents partially define their realized niches. Identifying items in stomachs of introduced rodents helps determine rodents' trophic positions and species most at risk of consumption. In the Hawaiian Islands, which lacked

rodents prior to human arrival, three rodents (*Rattus rattus* or black rat, *R. exulans* or Pacific rat, *Mus musculus* or house mouse) commonly coexist in native habitats where they consume a wide range of plants and animals. These three rodent species were trapped in montane forest for 2.5 years; their stomach contents were analyzed to determine short-term diets ($n = 12\text{--}95$ indiv. per species), and isotopic fractions of $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ in their bone collagen were analyzed to further estimate their trophic positions ($n = 11\text{--}20$ indiv. per species). For all three species, $>75\%$ of individuals had plants and $>90\%$ had arthropods in their stomachs, and significant differences in mean relative abundances were found for food items in stomachs among all three rodents. Rodents may be dispersing some native and non-native seeds, including the highly invasive *Clidemia hirta*. Most identifiable arthropods in rodent stomachs were non-native, and no stomachs contained birds, snails, or lizards. The $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ signatures were consistent with trophic feeding differences revealed from stomach contents. Dietary niche differentiation by coexisting rodent species is evident in this forest, with Pacific rats being intermediate between the mostly carnivorous house mouse and the mostly herbivorous black rat; such findings can help forecast rodent impacts and direct management efforts in ecosystems where these invasive animals coexist.

A. B. Shiels (✉)
USDA, APHIS, National Wildlife Research Center,
Hawaii Field Station, P.O. Box 10880, Hilo,
HI 96721, USA
e-mail: ashiels@hawaii.edu

C. A. Flores
Department of Microbiology, University of Hawaii at
Manoa, 2538 McCarthy Mall, Honolulu, HI 96822, USA

A. Khamsing
Department of Natural Resources and Environmental
Management, 1910 East-West Road, Honolulu,
HI 96822, USA

P. D. Krushelnycky
Department of Plant and Environmental Protection
Sciences, 3050 Maile Way, Honolulu, HI 96822, USA

S. M. Mosher
Naval Facilities Engineering Command Marianas, US
Navy, PSC 455, Box 195, FPO AP, Santa Rita,
GU 96540, USA

D. R. Drake
Department of Botany, University of Hawaii at Manoa,
3190 Maile Way, Honolulu, HI 96822, USA

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Introduction

How closely-related species coexist in a community has long intrigued ecologists. Resource use and resource competition are two of the many ecological factors that influence a species' niche (Elton 1927; Hutchinson 1957). Theory predicts that animals with similar life-history traits and close phylogenetic associations, such as different species of rodents in the same habitat, are able to coexist because they partition resources across time and space (Gause 1934). However, several factors complicate the ability to determine if resource partitioning is actively practiced by coexisting species, including past and present competition, arrival order, relative abundances, and resistance and resiliency to disturbance. Despite the difficulty in determining the mechanisms of differential resource use, dietary comparisons of similar, coexisting species can help define species niches (Kotler and Brown 1988; Biró et al. 2005).

Introduced animals can disrupt food webs by consuming native species and by altering the realized niches of native competitors (Fritts and Rodda 1998; Fukami et al. 2006). Identifying species consumed by introduced animals provides insight into native species' vulnerability and can inform strategies for managing native and non-native species (Stapp 2002; Caut et al. 2008a; Bonnaud et al. 2011; St Clair 2011). Rodents (*Rattus rattus*, black or ship rat; *R. norvegicus*, Norway rat; *R. exulans*, Pacific rat; *Mus musculus*, house mouse) have been introduced to many ecosystems and are among the most widespread and problematic invasive animals affecting islands (Towns et al. 2006; Angel et al. 2009; Drake and Hunt 2009). Introduced rodents may consume a wide variety of food items, including plants (e.g., fruits, seeds, vegetative material) and animals (e.g., arthropods, mollusks, birds; Sugihara 1997; Stapp 2002; Drake et al. 2011), and their diets can shift depending upon a number of factors, including food availability, the chemical and nutritional quality of food items, and the rodents' competitive ability relative to other animals that coexist in the environment (Clark 1981, 1982; Caut et al. 2008a; Ruffino et al. 2011).

Food consumption by introduced rodents is rarely observed directly, perhaps because they are shy, nocturnal, and often burrow belowground (Lindsey et al. 1999; Shiels 2010). Techniques commonly used to assess the diets of introduced rodents include field

observations of partially consumed prey (e.g., seeds, mollusks, arthropods; Norman 1970; McConkey et al. 2003; Meyer and Shiels 2009), captive-feeding trials (Bunn and Craig 1989; Williams et al. 2000; Pérez et al. 2008; Shiels 2011), and stomach content analysis (Clark 1981, 1982; Amarasekare 1994; Pisanu et al. 2011). Additionally, the analysis of naturally-occurring stable isotope ratios (i.e., $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$) of the rodents' tissues have been widely used to determine the diets and trophic levels at which animals have fed during tissue development; however, interpretations of diet using stable isotopes are complicated by variation in tissue turnover rates among organs (Peterson and Fry 1987; Lajtha and Michener 1994). For example, liver tissue has a higher turnover rate than blood cells or muscle, whereas bone collagen is deposited and modified throughout life, so its isotopic values represent a long-term average of an animal's diet (Lajtha and Michener 1994). The difference in isotopic composition between a consumer and its food (discrimination values) is presumed to average ca. 3 ‰ for $\delta^{15}\text{N}$ and 1 ‰ for $\delta^{13}\text{C}$ (Peterson and Fry 1987; Post 2002; Caut et al. 2009); however, discrimination values can differ widely among food sources (Caut et al. 2009). For example, Post (2002) found that the majority of the $\delta^{15}\text{N}$ discrimination values in lake organisms were 2–4.5 ‰, and Caut et al. (2008b) determined from lab trials that the $\delta^{15}\text{N}$ discrimination values for black rats ranged from –1.46 to 4.59 ‰. A disadvantage of stable isotope analysis is that it is often imprecise for identifying specific taxa that have been consumed (see Phillips 2012 for a review). Diet assessments that combine multiple techniques, such as stable isotope analysis with stomach content analysis, generally provide a more complete understanding of an animal's dietary niche than do assessments using only one technique (Drake et al. 2011).

The objectives of this study are to (1) determine the short-term diets, using stomach content analysis, of black rats, Pacific rats, and mice where they coexist in Hawaiian montane forest, and (2) investigate the trophic positions of the three rodent species via the analysis of stable isotopes occurring in low-turnover tissue (i.e., bone collagen) that has integrated the resources used by rodents over several months. These three rodents are widespread and coexist in many ecosystems in Hawaii from sea level to nearly 3,000 m a.s.l. (Amarasekare 1994; Shiels 2010). The first rodents (Pacific rats) arrived with the first humans ca.

800 years ago (Wilmshurst et al. 2011), and the others arrived with Europeans ca. 230 years ago. These rodents consume both plants and animals in Hawaii (Sugihara 1997; Cole et al. 2000; Shiels and Drake 2011; Pender et al. in press). However, the types of species most at risk of consumption by each of these three rodent species where they occur sympatrically have not been well established; such information can assist in native ecosystem management in areas where these rodents have invaded.

Methods

Study site

Rodents were obtained from Kahanahaiki Management Unit (21° 32' N, 158° 11' W), a 36 ha segment of mesic forest in highly dissected terrain (500–660 m a.s.l.) in the northern Waianae Mountains, on Oahu, Hawaii. Kahanahaiki is managed for native species conservation by the US Army, and the forest was fenced in 1996 to exclude feral goats and pigs. Annual precipitation at the site is approximately 1265 mm (Giambelluca et al. 2011), and the daily air temperature is 16–24 °C (Shiels 2010). The forest is a mixture of native and non-native vegetation. There are >30 tree species common to the forest, and the five dominant tree species include three natives (*Diospyros hillebrandii*, *Psyrdrax odorata*, and *Sapindus oahuensis*) and two non-natives (*Psidium cattleianum* and *Schinus terebinthifolius*; Shiels 2010). Fruit production occurs year-round, with the greatest fruit-fall observed in November–March (fruit numbers) and June–September (biomass) (Shiels 2010).

The black rat, Pacific rat, and house mouse occur at Kahanahaiki; Norway rats are absent from this forest and most others in Hawaii (Shiels 2010). Mean relative abundance (No. indiv. 100 trap nights⁻¹) estimated from bi-monthly mark-and-recapture sampling over 26 months during 2007–2009 were (mean ± SE) 13.5 ± 2.7 for black rats, 0.7 ± 0.4 for Pacific rats, and 7.9 ± 3.3 for house mice (Shiels 2010). Other vertebrate consumers in the forest include native and non-native birds, and non-native reptiles (e.g., *Lampropholis delicata*), mongooses (*Herpestes auropunctatus*), and house cats (*Felis catus*) (Shiels 2010). Invertebrate consumers include native and non-native arthropods and snails, and non-

native slugs (Joe and Daehler 2008; Meyer and Shiels 2009).

Stomach content analysis

Black rats ($n = 95$), Pacific rats ($n = 12$), and mice ($n = 47$) were collected from kill-traps (Victor[®] rat traps) placed on the ground from February 2007 through September 2009. In February 2007, traps were established at 10–25 m intervals along a single 300 m transect and within two 50 × 50 m plots at the ends of the transect (where native tree snails (*Achatinella mustellina*) were relatively abundant). Each month, 15–32 traps were baited with coconut chunks or peanut butter, set for 2–5 consecutive days, and checked daily. From May to September 2009, approximately 400 kill-traps were added to the 36 ha site and arranged along multiple transects that circled the core interior. Each transect was ca. 50 m from the next closest transect, and trap spacing was 12.5 m along the perimeter and 25 m on all interior transects (Pender et al. in press). We used the same bait as described above, and traps were checked each 1–7 days. Only rodents that were freshly (<24 h) killed, evidenced by lack of obvious decay, were used in this study. These rodent body masses were (mean ± SE) 124 ± 5 g for black rats, 52 ± 4 g for Pacific rats, and 12 ± 1 g for mice; the sex ratio of each species was roughly 50:50. Carcasses were stored in a freezer until analyzed.

Stomach contents were extracted, swirled for 5 min in water and mild detergent (Joy[®] brand) to separate contents and dissolve gastric juices and oils, sieved through a 0.4 mm sieve, and preserved in 95 % ethanol (Sugihara 1997). A transparent grid (5 × 5 mm for rats; 3 × 3 mm for mice) was positioned beneath a Petri dish containing each sample and then the sample was inspected using a dissecting microscope with 10–20× magnification. Relative abundance (percent) of each food type was determined for each sample by scoring the number of grid-boxes containing a given food type and dividing by the total number of grid-boxes (i.e., 40 grid-boxes). If more than one food type was in a grid-box, the item nearest the center was recorded (Cole et al. 2000). For each rodent species the frequency of occurrence (percent) for each food type was calculated by the presence of each of the food types in a given sample (individual) divided by the total number of samples. There were three major food types: plants, arthropods, and other.

Plant food types were further categorized as fruit, seed, and other plant material (including leaves, flowers, stems, wood). Arthropod food types included caterpillar (Lepidoptera larvae), ant (Hymenoptera), burrowing bug (Hemiptera), spider (Araneae), and other arthropod material. The ‘other’ category (major food type) included rodent hair and flesh, and unknown material that did not fit any of the previously listed food types. Food items were classified to the lowest taxonomic level possible using voucher specimens collected from the study site.

Stable isotope analysis

To augment the short-term diet assessment from stomach contents, the trophic positions of the three rodent species were estimated using stable isotope analysis. On a random subset of the trapped rodents ($n = 20$ black rats, 12 Pacific rats, and 11 mice), bone collagen was extracted and analyzed for $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values using methods described in Lajtha and Michener (1994) where the femur (plus tibia and fibula for mice) of each individual was excised, cleaned of flesh, and soaked in 0.5 M HCl for 48 h at 4 °C; the remaining sample (now collagen) was rinsed with deionized water, dried at 60 °C for 5 days, and ground to a fine dust. Common food items (fruit, seed, arthropod) were collected from the study site on 15 December 2009 and analyzed for $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$; food items were chosen based on similar species (or life forms) identified in the rodent stomachs and those found to be attractive to rodents at this site during field trials (Shiels and Drake 2011). For plant items ($n = 5$), one sample from each of the following species was collected for analysis: *Alyxia stellata* (seed), *Clidemia hirta* (fruit + seed), *Diospyros hillebrandii* (seed), *P. cattleianum* (seed), and *Planchonella sandwicensis* (fruit). Three samples of each of three herbivorous or detritivorous arthropods were analyzed: caterpillar, isopod, and amphipod. Three predatory arthropods (spiders) were analyzed, including *Steatoda capensis* and two unknown species. Samples were dried at 60 °C, ground to homogenize either multiple individuals of the same species (e.g., plants, herbivorous/detritivorous arthropods) or single individuals (e.g., spiders), and, like the rodent bone collagen, analyzed isotopically using a Carlo Erba elemental analyzer (model NC2500) with an attached mass spectrometer (Finnegan DeltaS with source

upgrade). Stable isotope ratios were expressed in δ notation as parts per thousand (‰) deviation from international standards (Lajtha and Michener 1994).

Statistical analysis

The relative abundances of food types were compared among the three rodents by parametric and non-parametric ANOVAs. Fruit and seed met parametric assumptions of ANOVA after arcsine square-root transformations; the remaining comparisons used Kruskal–Wallis tests to assess significant differences among rodents for each food type. Post-hoc Tukey’s tests (for fruit and seed) or Mann–Whitney U tests were applied to assess significance between rodent species; significance was based on $P < 0.05$ (R Development Core Team 2010).

To test whether the diet of black rats changed during the time period when few (February 2007–April 2009), and the majority (May–September 2009), of Pacific rats and mice were trapped, ANOVAs, after square-root transformations, were used to compare the two time periods for each of three food types: fruit, seed, and arthropod.

Results

Stomach content analysis

All three rodent species consumed both plants and arthropods (Fig. 1). Plant relative abundance in stomachs differed significantly among rodents ($P < 0.001$; $\chi^2 = 56.7$, $df = 2$), with black rats > Pacific rats > mice ($P < 0.015$ for each post hoc comparison; Fig. 1). Arthropod mean relative abundance also differed significantly among rodents ($P < 0.001$; $\chi^2 = 56.7$, $df = 2$), with mice > Pacific rats > black rats ($P < 0.035$ for each post hoc comparison; Fig. 1). Rodent hair, which dominated the ‘other’ category in Fig. 1, was found in most stomachs of each species (69 % of black rat individuals, 67 % of Pacific rats, and 57 % of mice), and mean relative abundance for rodent hair was not significantly different among rodents ($P = 0.775$; $\chi^2 = 0.5$, $df = 2$; Table 1). Rodent hair in stomachs probably resulted from grooming; rodent flesh with hair attached was rare (i.e., in $n = 1$ black rat, and $n = 1$ Pacific rat

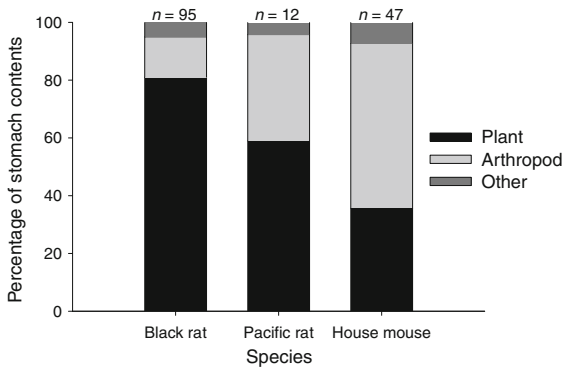


Fig. 1 Mean relative abundance (%) of major food types found in stomachs of black rats, Pacific rats, and house mice in Hawaiian mesic forest. There were significant ($P < 0.05$) differences between each species for the two major food types (plant and arthropod). The ‘other’ category is dominated by rodent hair, which was most likely a result of grooming rather than cannibalism

stomach). No evidence of birds, reptiles, slugs, snails, or fungi occurred in any stomachs.

All plant and arthropod food types analyzed were found in stomachs of at least some individuals of all three rodents (Table 1). There were significant differences among rodents for most food types found in stomachs, and burrowing bugs were the only prey whose relative abundance was not significantly different among rodents (Table 1). Fruit comprised the

majority of the plant material for both rats, but seed was the most abundant plant material in mice (Table 1). Caterpillars comprised the majority of the identifiable arthropods found in each of the rodents, and were ca. 94 % of the arthropod diet of mice (Table 1).

For each rodent species, >75 % of individuals had plants in their stomachs (Table 2). All black rats and Pacific rats had fruit in their stomachs, and >90 % also had seed. The frequency of mouse stomachs with fruit (40 %) tended to be less than those containing seed (64 %). The majority of seed in all three rodents appeared chewed and was probably destroyed, but intact seeds of some native and non-native species were found in black rats, and all three rodents had intact seeds of the invasive non-native *C. hirta*. The frequency of other plant material (mostly stems and leaves) was highest in Pacific rats, intermediate in mice, and lowest in black rats.

Arthropods, which mostly appeared as fragments rather than intact animals, were found in nearly all (>90 %) of the rodents examined (Table 2). Only four species of arthropod were found intact in rodent stomachs, and these were *Solenopsis papuana* (Papuan thief ant, Hymenoptera), *Xylosandrus compactus* (black twig-borer, Coleoptera), *Stelidota geminata* (strawberry sap beetle, Coleoptera), and Phthiraptera (rat lice) (Table 2). The two arthropod species that

Table 1 Mean \pm SE relative abundance (%) of plant, arthropod, and other food types identified in stomachs of invasive rodents in Hawaiian mesic forest

Food type	Black rat (n = 95)	Pacific rat (n = 12)	House mouse (n = 47)	P value
Plant				
Fruit	55.1 \pm 2.4 ^a	40.6 \pm 5.7 ^a	10.8 \pm 2.7 ^b	<0.001
Seed	24.9 \pm 2.2 ^a	15.9 \pm 4.3 ^{a,b}	19.0 \pm 3.3 ^b	0.037
Other plant material	1.1 \pm 0.4 ^a	2.5 \pm 0.8 ^b	5.8 \pm 1.5 ^b	0.002
Arthropod				
Caterpillar	3.2 \pm 0.7 ^a	27.8 \pm 3.8 ^b	53.8 \pm 4.9 ^c	<0.001
Ant	1.7 \pm 0.3 ^a	1.2 \pm 0.8 ^a	0.5 \pm 0.2 ^b	0.007
Burrowing bug	0.9 \pm 0.3	0.2 \pm 0.2	0.7 \pm 0.5	0.110
Spider	0.4 \pm 0.1 ^a	0.2 \pm 0.2 ^b	0.1 \pm 0.1 ^b	0.016
Other arthropod material	7.4 \pm 1.4 ^a	7.4 \pm 2.7 ^a	2.1 \pm 1.0 ^b	0.002
Other				
Rodent hair + flesh	4.9 \pm 0.7	4.2 \pm 1.4	7.2 \pm 1.5	0.775
Unknown	0.4 \pm 0.2	0.0 \pm 0.0	0.1 \pm 0.1	0.528

The P value reflects the comparison among species; within a row, means sharing the same letter are not significantly different ($P > 0.05$). When no amount of food item was present for a rodent (i.e., unknown), that rodent was not included in the statistical analysis

Table 2 Frequency (%) of rodent stomachs with identifiable seeds, fruit, and arthropod taxa in Hawaiian mesic forest

Species or group ^a	Life form	Native or non-native ^b	Black rat (n = 95)	Pacific rat (n = 12)	House mouse (n = 47)
<i>Plants</i>					
Intact seeds					
<i>Clidemia hirta</i>	Shrub	Non-native	30.5	25.0	6.4
<i>Rubus rosifolius</i>	Shrub	Non-native	7.4	0	0
<i>Paspalum conjugatum</i>	Grass	Non-native	2.1	0	0
<i>Cyrtandra dentata</i> ^c	Shrub	Native	1.1	0	0
<i>Delissea waianaensis</i> ^c	Shrub	Native	1.1	0	0
Unknown #1	–	–	1.1	0	0
Unknown #2	–	–	1.1	0	0
Unknown #3	–	–	2.1	0	0
Total intact seeds			42.1	25.0	6.4
Total seed			93.7	91.7	63.8
Fruit fragments					
<i>Clidemia hirta</i>	Shrub	Non-native	33.7	50.0	6.4
<i>Rubus rosifolius</i>	Shrub	Non-native	7.4	8.3	0
<i>Psidium cattleianum</i>	Tree	Non-native	25.3	2.1	0
Total fruit			100	100	40.4
Other plant material			16.8	58.3	34
Total			100	100	76.6
<i>Arthropods</i>					
<i>Solenopsis papuana</i>	Ant	Non-native	36.8	16.7	10.6
<i>Rhytidoporus indentatus</i>	Burrowing bug	Non-native	16.8	8.3	4.3
<i>Balta</i> spp.	Cockroach	Non-native	6.3	0	0
<i>Platyzoisteria soreri</i>	Cockroach	Non-native	1.1	8.3	0
<i>Steatoda capensis</i>	Spider	Non-native	3.2	8.3	0
<i>Mecaphesa</i> sp.	Spider	Native	0	0	1.1
<i>Blackburnia epicurus</i>	Beetle	Native	1.1	0	0
<i>Rhyncogonus</i> sp.	Beetle	Native	1.1	0	0
<i>Araecerus fasciculatus</i>	Beetle	Non-native	1.1	0	0
<i>Xylosandrus compactus</i>	Beetle	Non-native	1.1	0	0
<i>Stelidota geminata</i>	Beetle	Non-native	2.1	0	0
<i>Pantomorus cervinus</i>	Beetle	Non-native	0	0	1.1
<i>Banza</i> sp.	Katydid	Native	1.1	0	0
<i>Apis mellifera</i>	Honey bee	Non-native	2.1	0	0
Hemiptera	Aphid	Non-native	1.1	0	0
Phthiraptera	Louse/lice	Non-native	2.1	0	0
Orthoptera	Cricket/Katydid	–	1.1	0	0
Lepidoptera larvae	Caterpillar	–	33.7	100	83.0
Diptera larvae	Fly maggot	–	2.1	0	8.5
Araneae	Spider	–	11.6	0	0
Other arthropod material	–	–	9.5	75	23.4
Total			91.6	100	95.7

^a Many items were so damaged that they could not be classified more specifically than fruit, seed, plant, or arthropod

^b All taxa listed as native are endemic to Hawaii

^c Federally endangered species

were found in some individuals of all three rodents were *S. papuana* and *Rhytidoporus indentatus* (burrowing bug, Hemiptera). Most identifiable arthropods were non-native species; the only identifiable native arthropods found in stomachs were *Banza* sp. (bush cricket/katydid, Orthoptera), *Blackburnia epicurus* (ground beetle, Coleoptera), and *Rhyncogonus* sp. (weevil, Coleoptera) in black rats, and *Mecaphesa* sp. (crab spider, Araneae) in mice. Unknown species of caterpillars were found in all of Pacific rats, 83 % of mice, and 34 % of black rats (Table 2).

There was no evidence from stomach content analysis indicating a dietary shift in black rats between seasons when few (February 2007–April 2009), and the majority (May–September 2009), of Pacific rats and mice were trapped ($P = 0.709$ for fruit; $P = 0.860$ for seed; $P = 0.549$ for arthropod; Fig. 2). Clear evidence of seasonal patterns in consumed food items were generally absent in our study; fruit and/or seed of some common species at the site (e.g., *C. hirta*) were found each month in some rodent stomachs, whereas uncommon species (e.g., *Cyrtandra dentata* and *Delissea waianaensis*) were limited to stomachs of single individuals that were recovered during one collection period.

Stable isotope analysis

Examining trophic positions via isotopic signatures reveals that black rats have lower $\delta^{15}\text{N}$ values than

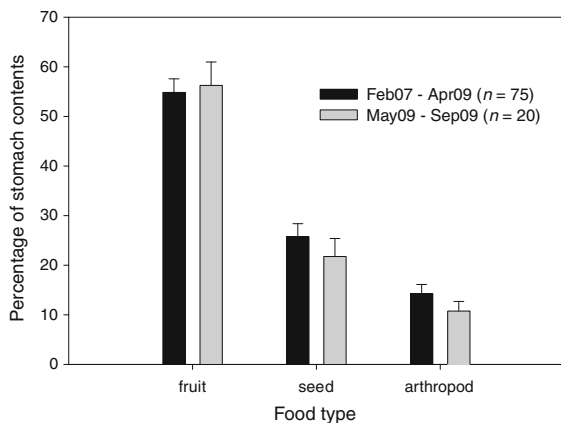


Fig. 2 Mean ± SE relative abundance (%) of fruit, seed, and arthropod identified in stomachs of black rats in Hawaiian mesic forest for the time periods when few (February 2007–April 2009), and the majority (May–September 2009), of Pacific rats and mice were trapped. There were no significant differences ($P > 0.05$) between time periods for any of the food items

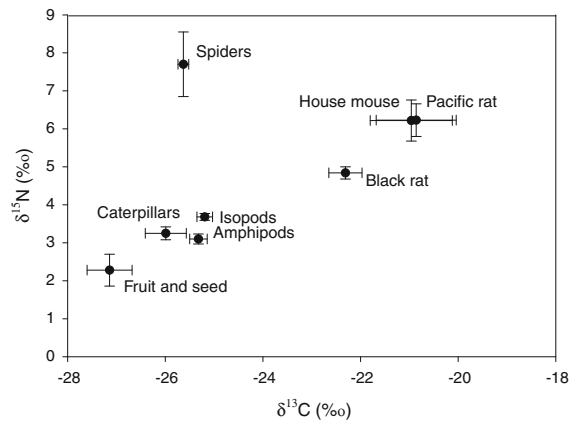


Fig. 3 Mean ± SE $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values for black rats ($n = 20$), Pacific rats ($n = 12$), and the house mouse ($n = 11$), and their potential prey items (spiders $n = 3$; isopods $n = 3$; amphipods $n = 3$; caterpillars $n = 3$; fruits and seeds $n = 5$), from Hawaiian mesic forest

Pacific rats and mice (Fig. 3), and therefore appear to generally be feeding at lower trophic levels than are Pacific rats and mice. The Pacific rat and mouse have similar $\delta^{15}\text{N}$ signatures. Spiders, which prey on arthropods but not plants, are generally feeding at higher trophic levels than all three rodents. Although all three rodents consume plants and animals, the $\delta^{15}\text{N}$ findings are consistent with the results from stomach contents, which depict black rats as mainly vegetarian and Pacific rats and mice as slightly more carnivorous.

The three rodents appear to form a distinct grouping from their potential prey when $\delta^{13}\text{C}$ is examined (Fig. 3). Although there is a relatively high amount of variability among Pacific rat and mouse samples for $\delta^{13}\text{C}$, these two rodents are nearly equal in $\delta^{13}\text{C}$ and tend to be slightly higher than the $\delta^{13}\text{C}$ signature of the black rat (Fig. 3). The herbivorous/detritivorous arthropods are about 1.5 ‰ from plants, but surprisingly the spiders are also aligned with the herbivorous/detritivorous arthropods for $\delta^{13}\text{C}$ (Fig. 3).

Discussion

Black rats, Pacific rats, and house mice each consume a variety of plants and animals where they coexist in this insular tropical forest. The dietary niches of these three rodents differ such that the house mouse is primarily carnivorous and feeds mainly on arthropods (especially caterpillars), the black rat is primarily

vegetarian and feeds mainly on fruit and seed, and the Pacific rat has an intermediate diet that, over its lifetime, is more closely related to the house mouse than to the black rat. An understanding of the trophic level overlap among these introduced rodents, as evidenced by stomach contents and stable isotope analysis, should help identify the types of species that may be vulnerable to rodent consumption and deserving of conservation attention.

Plant and animal tissue fragments, and intact seeds, in the rodent stomachs provided evidence of the taxa that were consumed during the 2.5 year study. Invasive rodents are generally viewed as seed predators (Clark 1981; Towns et al. 2006; Angel et al. 2009), and most seeds consumed by rodents in our study appeared highly vulnerable to predation, having been chewed and fragmented to the point that species identification was impossible. However, some seeds in rodent stomachs were intact and identifiable, such as the highly invasive *C. hirta*, which was found in some individuals of all three rodents, and could potentially be dispersed by them. A greater range of seed sizes can pass intact through black rats than through the two smaller rodents (Williams et al. 2000). Intact seeds of eight plant species, including at least two endangered natives and two invasive non-natives, occurred in black rats (Table 2) and their small seeds (≤ 1.5 mm length) would likely be passed intact and germinate (Shiels 2011; Shiels and Drake 2011). Similarly, arthropods were more identifiable in black rats than in the other rodents, perhaps because many of the fragments were slightly larger. True bugs, spiders, ants, crickets and other orthopterans, beetles, and caterpillars are common prey of these three rodents elsewhere (Cole et al. 2000; Innes 2005; Ruscoe and Murphy 2005; St Clair 2011), and they occurred in rodents in our study. Approximately one-third of the identifiable arthropod taxa in black rats were native in our study, but only one native species (*Mecaphesa* sp.) occurred in the house mouse, and none were identified in Pacific rats. The frequency of native species among total arthropod prey items was low; however, this may be an artifact of the high percentage of prey items, such as caterpillars, whose species-level identity and therefore provenance could not be determined. Stomach content analyses cannot directly yield estimates of population-level impacts on prey species. Nevertheless, the fact that individuals of rare native arthropod species, such as *B. epicurus*,

were found in rodent stomachs suggests that rodents may represent important threats to the long-term viability of such species.

As expected, all three rodent species were highly omnivorous; however, the relative abundances of food items (e.g., fruit, seed, arthropod) and identifiable species consumed differed among rodents. Such niche differentiation is consistent with theory used to explain coexistence between closely related organisms (Gause 1934; Kotler and Brown 1988). Plant material often comprises 75–80 % of black rat diets within and outside of Hawaii, regardless of the types of coexisting rodent species (Kami 1966; Norman 1970; Clark 1981; Cole et al. 2000; Sweetapple and Nugent 2007; this study). In Hawaii, fruit can constitute the bulk of the black rats' plant diet in mesic forest (55 % relative abundance of stomach contents; this study), arid shrubland (44 %; Cole et al. 2000), and wet forest (23–53 %; Sugihara 1997). When Pacific rats are the only rodents present at insular sites, plants can be 65–90 % of their diet (Wirtz 1972; Mosby et al. 1973; Bunn and Craig 1989), yet fruit consumption by Pacific rats in Hawaii appears more variable (41 % in our study vs. 3–16 % in Sugihara 1997) than it does for black rats. Both rat species consume fruit of problematic invasive species in Hawaii, such as *C. hirta* and *Rubus rosifolius* (Beard and Pitt 2006; this study) and *P. cattleianum* (this study). Interestingly, fruit of *R. rosifolius* and *P. cattleianum* were not found in any mice at Kahanahaiki despite being abundant during the time when most mice were trapped. The amount of fruit (11 %) in mouse stomachs was much less than in both rat species in our study, but similar to the 10 % determined by Cole et al. (2000). Fruit was absent from all 25 mouse stomachs analyzed from gulches adjacent to sugar cane fields on Hawaii Island (Kami 1966). Studies within and outside Hawaii suggest that house mice consume relatively small portions of fruit (especially fleshy fruit) compared to seed, vegetative material, and arthropods (Kami 1966; Cole et al. 2000; Angel et al. 2009; this study). It is unclear why fruits and seeds from other common species from Kahanahaiki were not observed in rodent stomachs; it may be a result of food preference or simply a reflection of the difficulty in identifying microscopic material in rodent stomachs.

Arthropods are common in diets of introduced rodents, being found in at least 80 % of stomachs examined from Hawaii and elsewhere (Gales 1982;

Amarasekare 1994; Sugihara 1997; Miller and Webb 2001; this study). In a recent review, Angel et al. (2009) found that arthropods were the food of choice for house mice on islands in the Southern Ocean, a pattern consistent with that at Kahanahaiki, where arthropods accounted for an average of 57 % of their stomach contents. In high elevation (1600–3000 m) sites in Hawaii, arthropods comprised 33–54 % of mouse diets (Amarasekare 1994; Cole et al. 2000). Relative to mice, arthropods were a much smaller component of black rat stomach contents in our study (14 %) and that of Cole et al. (2000; 16 %). In Hawaiian lowland wet forest, there were no arthropods in black rat stomachs and only trace amounts of caterpillars in Pacific rat stomachs (Beard and Pitt 2006). Few data are available for arthropods in Pacific rat stomachs in Hawaii because these rats were not captured (Amarasekare 1994; Cole et al. 2000), or because arthropods were not segregated from other invertebrates when stomach contents were analyzed (Sugihara 1997). Despite the presence at Kahanahaiki of native and non-native birds and snails (Meyer and Shiels 2009; Shiels 2010), and non-native slugs, earthworms, and reptiles (Joe and Daehler 2008; Shiels 2010), there was no evidence of any of these organisms in rodent stomachs. The species composition and relative abundances of plants and animals available in rodent-occupied environments represent additional factors that can directly affect rodent diets (Kotler and Brown 1988; Ruffino et al. 2011); native birds, for example, are uncommon relative to non-native birds at our study site.

Caterpillars appear to be a highly attractive food item to all three rodents studied in Hawaii; some individuals of all species studied had caterpillars in their stomachs in high elevation environments (Amarasekare 1994; Sugihara 1997; Cole et al. 2000), and 100, 83 and 34 % of Pacific rats, mice, and black rats, respectively, had caterpillars in their stomachs in Kahanahaiki. The proportion of stomach contents comprised of caterpillars was greatest in mice (54 % in our study; 22 % in Cole et al. 2000) and least in black rats (3 % in our study; 4 % in Cole et al. 2000). On islands outside of Hawaii, caterpillars can also comprise the most common arthropod eaten by mice (Rowe-Rowe et al. 1989; Miller and Webb 2001; Ruscoe and Murphy 2005), and one of the most common groups eaten by Pacific rats (Bunn and Craig 1989; Atkinson and Towns 2005).

Although the three rodents in our study are omnivores, and thus appear to occupy the same general trophic level, they occupy different dietary niches. Stomach content analyses revealed that Pacific rats had an intermediate (short-term) diet between those of black rats and mice, yet the greater amount of caterpillars and unknown arthropods consumed by Pacific rats and mice relative to black rats may partially account for the slightly higher $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ for the two smaller rodents compared to black rats. Additionally, Pacific rats may be more similar to mice than to black rats in lifetime average diet (as indicated by $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$) as a result of isotopic incorporation rates of prey differing by rodent species (Gannes et al. 1997), or because Pacific rats shift their foraging microsites when the black rat is present (Lindsey et al. 1999; Atkinson and Towns 2005; Shiels 2010). The difference in $\delta^{15}\text{N}$ among rodent species averaged <1.5 ‰, which does not typically justify assigning distinct trophic levels to different species (Peterson and Fry 1987; Lajtha and Michener 1994; Post 2002). Many isotope studies have examined the degree to which introduced rodents ate seabirds and the proportion of diet attributable to marine and terrestrial sources (Stapp 2002; Caut et al. 2008a; Quillfeldt et al. 2008; Ruffino et al. 2011). Marine inputs to rodent diets are unlikely at Kahanahaiki because the site is >3 km from the ocean, there are no seabirds, and home-ranges are typically <4 ha for each rodent (Shiels 2010). Therefore, $\delta^{13}\text{C}$ differences in our study are more likely to involve unequal consumption of C_3 and C_4 plants (Gannes et al. 1997), which may help explain the slightly higher $\delta^{13}\text{C}$ values for Pacific rats and mice relative to black rats. The majority of the C_4 plants at Kahanahaiki are grasses (e.g., *Paspalum conjugatum*, *Megathyrus maximus*), and many mice and some Pacific rats in our study were captured near grassy patches (A. Shiels, personal observation). In an inland forest on Stewart Island, New Zealand, Harper (2006) used $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values to determine that diets of Pacific rats and black rats were similar. The only isotopic study available with wild house mice was by Quillfeldt et al. (2008) in the Falkland Islands where $\delta^{13}\text{C}$ were similar to those in our study (−21 and −24 ‰); yet the $\delta^{15}\text{N}$ for mice ranged from 12 to 31 ‰ and were indistinguishable from potential food items, including terrestrial plants (8–35 ‰), terrestrial invertebrates (14 ‰), and upland birds (16–19 ‰). Therefore, using isotopes to estimate trophic levels

and determine the types of species which consumers feed upon may be challenging without supplemental dietary analysis such as stomach contents (Stapp 2002; Caut et al. 2008a; Quillfeldt et al. 2008; this study).

Many factors can affect the dietary niches of coexisting rodents. Much evidence points to the largest of the three rodents, the black rat, as the dominant competitor of the three species studied (Yom-tov et al. 1999; Russell and Clout 2004; Shiels 2010). Stokes et al. (2009) in Australia, and Harris and Macdonald (2007) in the Galápagos Islands, demonstrated that native rats (*R. fuscipes* and *Nesoryzomys swarthi*, respectively) suffered from interference competition rather than resource competition with the larger, non-native black rats. Furthermore, removal of black rats can result in population increases in coexisting rodents such as house mice (Harper and Cabrera 2010; Ruscoe et al. 2011). The average body masses of the three coexisting rodents in our study differed 4–10 fold. Size differences >2 fold allow sympatric congeners to occupy different niches but co-occur in the same trophic level (Hutchinson 1959; Eadie et al. 1987). Diet variation is perhaps just one of the niche differences among these three rodent species that enable their coexistence at Kahanahaiki and in many other ecosystems.

Differential resource uses, or dietary niches, of sympatric black rats, Pacific rats, and house mice at Kahanahaiki reflect unequal consumption of species that occupy different trophic levels. The ecological and conservation implications for island habitats containing these three introduced rodents are that (1) fruit appears to be a main component of the diet for black rats and Pacific rats, (2) mice and Pacific rats likely exhibit greater predation pressure per capita than black rats on arthropod communities, and (3) all three rodents typically chew, and probably destroy, most consumed seeds >2 mm in length. The degree to which species and trophic levels are exploited by each introduced rodent in Hawaii and elsewhere may largely depend upon the assortment of rodent species, and the available food items, that are present at a given site.

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