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Myrmecophagy in a Ranid Frog Rana rugosa: Specialization or Weak Avoidance to Ant Eating?

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ABSTRACT—We studied diets of an Asian ranid, Rana rugosa, inhabiting three different environments (reservoir, river shoreline, and paddy fields) in Kyoto, Japan. In all the three habitats, ants were the most frequently consumed prey, representing surprisingly similar proportions in both the frequency of occurrence (81.9–85.7%) and the number of total prey items (56.8–59.4%). These values are exceptionally large for Rana, and equivalent to those reported for ant specialists in other families of frogs such as dendrobatids or bufonids. However, R. rugosa consumed ants lower in proportions than those found in the environment, and could not be regarded as purely ant specialists. Instead, we conclude that this species tends to avoid ants more weakly than other species of Rana. Other than ants, larger prey were more and smaller ones less frequently taken in proportion to frog body size, indicating that the frog consumes ants because of its weak avoidance of these abundant potential prey.

INTRODUCTION

Frogs, in general, are generalist predators and consume a wide variety of prey in response to prey availability in the environment (Duellman and Trueb, 1986). Some frogs, however, are known to selectively feed on particular prey. Many dendrobatid and some bufonid species are myrmecophagous. These frogs are specialized for eating ants, and consume them in a higher proportion than found in surrounding environments (Toft, 1980; 1981; Lieberman, 1986). Within Dendrobatidae, suites of traits related to diet and foraging may have been a significant force driving radiation of the family at the generic level (Toft, 1995; Caldwell, 1996). Detailed studies of feeding habits in a number of frog families are needed to examine the evolution of myrmecophagy in anurans as a whole.

The family Ranidae contains more than 600 species and is distributed worldwide (Duellman and Trueb, 1986). Diets of ranid frogs have been studied by many workers from various regions of the world. Ranids are considered to be generalist predators (e.g., Houston, 1973; Premo and Atomowidjojo, 1987) and to change their diets in response to natural fluctuations of prey availability (Turner, 1959; Tyler and Hoestenbach, 1979; Hirai and Matsui, 1999). Ranid frogs are gape-limited predators that regulate size of their prey by gape width or body size (Kramek, 1972) and their diets change ontogenetically (Werner et al., 1995). Myrmecophagy, common in dendrobatids and bufonids, has never been reported for any species of ranid, although the family is speciose and contains morphologically and ecologically diverse species and genera. Is myrmecophagy really absent among ranid frogs?

Rana rugosa is distributed in east Asia from Japan to

northeastern China, and commonly found from city areas to montane regions. The frog occupies various habitats and breeds both in still water and in running water. Differing from the other ranids, R. rugosa has been reported to consume large numbers of ants (Maeda and Matsui, 1989), but this previous report is anecdotal, based on few quantitative data. In order to assess myrmecophagy in this species, we conducted an extensive diet analysis placing particular emphasis on anteating. Specifically, we examined (1) ontogenetic dietary change, (2) variation in diet among different habitats, and (3) relationships between diet composition and prey availability in the environment of the frog's habitat.

MATERIALS AND METHODS

Field work

For stomach content analyses, we collected frogs from three distinctly different environments in Kyoto, central Japan: (1) a reservoir at Iwakura (35°06'N, 135°52'E) between June and August in 1996; (2) a shoreline of the Kiyotaki River at Kiyotaki (35°03'N, 135°47'E) between June and August in 1997; (3) paddy fields at Shizuhara (35°07'N, 135°52'E) from May to September in 1995 and 1996. As a result, we examined a total of 77, 67, and 16 stomachs from each population. We made each collection at night between 1800 hr and 0200 hr because we observed frogs more frequently at night than during the daytime.

Within two hours of capture, we anesthetized frogs in 1% solution of MS-222 (methane tricaine sulfonate) and extracted their stomach contents with forceps. Contents were preserved in 10% buffered formalin for later identification and analyses. For each frog, we recorded snout-vent length (SVL) to nearest 0.1 mm and toe-clipped for individual identification. After these procedures, we released frogs where they were captured.

In paddy fields, we sampled potential prey invertebrates on the Aze (slightly elevated narrow trail between adjoining rice fields) using two different techniques to estimate prey availability. To sample prey above ground, we made 1×80 m sweeps through the air and vegeta-

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tion, 0–0.5 m above the ground with an insect sweep net. For sampling terrestrial prey, we directly collected all visible animals on the ground or beneath stones in a 1×1 m plot using forceps. We collected samples before sunset (1800 hr–1900 hr, overlapping the time when frogs were active) on three days (spring: 27 May; summer: 26 July; autumn: 27 September 1995). These sampling dates largely covered the periods when frogs were actually collected. The sampled prey were killed with ethyl acetate and stored in ethylene glycol for later identification and analyses.

Diet analysis

We identified stomach contents and potential prey to the level of class or order except for Hymenoptera, which was classified into Formicidae and non-Formicidae. For holometabolous insects, larvae and adults were separated. The occurrence of plant materials or minerals were recorded for each stomach. Maximum length and width of prey, excluding antennae and cerci, were measured to the nearest 0.1 mm using a caliper or a calibrated ocular micrometer fitted to a dissecting microscope. For partially digested prey items, lengths (L) were estimated by measuring widths (W) and using predetermined length-width regressions from intact prey. Volumes of prey items were calculated by the formula for an ellipsoid (Dunham, 1983):

V = 4/3 π (L/2)(W/2)²

To detect ontogenetic diet change, we examined predator-prey size relationships by regressing volumes of the largest and smallest prey in a stomach to frog SVL and calculating correlation coefficients. Only frogs with at least three prey items in the stomach were included in this analysis. In addition, we examined relationships between frog SVL and the frequency of occurrence of the major (>25.0%) prey taxa using the reservoir population. The taxa included ants, beetles, dipterans, hemipterans, and spiders. For this analysis, we divided frogs by their SVL into size classes with intervals of 5.0 mm, and excluded the classes consisting of less than three individuals. For these examinations, we used the data from the reservoir because it had the largest number of stomach samples and contained frogs with a broad range of SVLs.

To examine variation in diets among populations inhabiting different environments, we compared the presence or absence of the five major prey taxa mentioned above with chi-squared 2×3 contingency table test. We also tested the difference in the numeric proportion of these taxa by using χ^2 test (Zar, 1999). The five major taxa found in the reservoir were commonly seen in stomachs of frogs from the river shoreline and the paddy fields (>18.0% in frequency of occurrence in all populations).

To determine the relationship between prey availability and diet composition, we calculated Kendall's rank correlation coefficients (τ), corrected for ties (Siegel, 1956) between the relative abundances of prey taxa in the habiat and their abundances in frog stomach contents. We approximated the prey availability within an area of 80 $m²$ by combining the total abundances of sweep and ground plot samples. In this analysis, we used only taxa that were found in both potential prey samples and the stomach contents.

Simon and Toft (1991) defined ant specialists as those frogs that consume ants in higher proportion than those found in the environment. We adopted this definition and calculated electivity indices (E of Ivlev, 1961 and E* of Vanderploeg and Scavia, 1979) to determine whether or not R . rugosa is an ant specialist. E is the most commomly used index, while E* is currently the most recommended one according to Lechowicz (1982). In this analysis, we calculated E and E* only for ants, because these indices are vulnerable to sampling errors for prey that are rare in the diet or in the environment (Lechowicz, 1982), and relative abundance of prey taxa other than ants greatly fluctuated seasonally, thus leading to large sampling errors. Both index values deviate symmetrically from zero between –1.0 to +1.0 as a prey taxon is respectively, avoided or preferred.

RESULTS

Diet composition

We identified 1577 prey items extracted from stomachs of 72 frogs collected at the reservoir; the remaining five frogs had empty stomachs. Arthropoda contained five classes (Arachnida, Crustacea, Chilopoda, Diplopoda, and Insecta) and constituted 97.3% in number and 96.7% in volume of the total stomach contents. Insecta contained 13 orders, and constituted 91.8% in number and 75.5% in volume (Table 1).

Ants (Formicidae) were not only the most frequently consumed prey taxon (81.9%), but also constituted the largest proportion in number (56.8%)(Table 1). In volume, however, ants constituted only a minute fraction (10.8%). Besides arthropods, gastropods, mostly lymnaeid pond snails, were present. Plant materials (plant pieces) and minerals (pebbles and dirt) were found in 19.4% and 25.0% of stomachs examined, respectively.

Ontogenetic change

As R. rugosa increases in body size, it consumes larger prey. There were significant positive correlations between frog SVL and volumes of the largest and smallest prey ingested (Fig. 1; rmin=0.561, P<0.01; rmax=0.287, P<0.05; n=71). Examination of the relationships between frog SVL and the frequency of occurrence of major prey taxa revealed that beetles with generally large body size exhibited a significantly positive correlation to SVL (Fig. 2b; r=0.767, P<0.05), but generally small-sized dipterans had a significantly negative correlation (Fig. 2c; $r=-0.776$, $P<0.05$). On the other hand, ants were consumed frequently regardless of frog body size, and their occurrence did not significantly correlate with SVL (Fig. 2a; r=–0.230, NS). Hemipterans and spiders also showed no significant correlations (Fig. 2d, e; r=0.753, r=0.675, respectively, both NS).

Variation among habitats

Ants were frequently consumed in all three habitats (Table 1). The frequency of occurrence of ants did not significantly differ among the habitats (χ^2 test, χ^2 =0.84, df=2, P>0.05). By contrast, beetles were consumed significantly more often in the paddy fields (χ^2 =9.00, df=2, P<0.05). Ants were numerically the most dominant in all the habitats, and values did not differ significantly among the habitats (χ^2 =0.93, df=2, P>0.05). The next largest proportion was occupied by beetles in the paddy fields, but by dipterans in the reservoir and the river shoreline; the values differed significantly among the habitats (beetles: χ^2 =94.37, df=2; dipterans: χ^2 =16.88, df=2, P<0.01 for both). The numeric proportions of hemipterans and spiders did not differ significantly among the habitats (hemipterans: χ^2 =94.37, df=2; spiders: χ^2 =16.88, df=2, P>0.05 for both). In volume, the largest proportion was occupied by beetles in both the river shoreline and the reservoir, but by beetle larvae in the paddy fields. Ants were represented volumetrically by 13.2% on the river shoreline, 4.1% in the paddy fields, and 10.8% in the reservoir.

		Frequency of occurrence (%)			Numeric proportion (%)			Volumetric proportion (%)		
Prey taxa	Reser- voir	Shore- line	Paddy fields	Reser- voir	Shore- line	Paddy fields	Reser- voir	Shore- line	Paddy fields	
Insecta										
Hymenoptera										
Formicidae	81.9	84.9	85.7	56.8	59.4	56.9	10.8	13.2	4.1	
non-Formicid	8.3	7.6	21.4	0.4	0.9	1.9	0.7	4.7	0.5	
larvae	5.6	1.9	7.1	0.3	0.2	0.4	1.2	0.6	0.2	
Coleoptera	45.8	41.5	85.7	3.8	7.7	19.5	17.3	20.9	9.9	
larvae	12.5	17.0	7.1	1.1	3.5	1.5	3.0	0.9	32.7	
Diptera	75.0	47.2	50.0	15.0	12.5	5.7	4.2	4.9	1.1	
larvae	22.2	7.6	14.3	5.6	1.2	0.8	1.9	1.3	4.9	
Lepidoptera	15.3	3.8	$\qquad \qquad -$	0.9	0.5	$\overline{}$	4.4	1.1	$\overline{}$	
larvae	18.1	11.3	14.3	1.2	1.6	1.5	22.7	23.1	12.5	
Trichoptera	1.4	$\qquad \qquad -$	$\qquad \qquad -$	0.1	-	$\overline{}$	< 0.1	$\overline{}$	-	
Neuroptera	6.9	$\overline{}$	-	0.3	-	-	0.4	$\overline{}$		
larvae	8.3	9.4	\equiv	0.4	2.3	$\overline{}$	0.8	1.7	$\overline{}$	
Hemiptera	38.9	18.9	28.6	4.1	2.8	2.3	4.4	7.3	2.3	
Dermaptera	$\qquad \qquad -$	1.9	$\overline{}$	$\overline{}$	0.2	-	$\overline{}$	1.1	$\overline{}$	
Orthoptera	6.9	1.9	7.1	0.4	0.2	0.4	3.1	0.7	0.1	
Plecoptera	1.4	$\overline{}$	$\overline{}$	0.1	$\overline{}$	$\overline{}$	< 0.1	$\overline{}$	$\overline{}$	
Odonata	1.4	1.9	$\overline{}$	0.1	0.2	\equiv	0.6	3.0	$\overline{}$	
larvae	-	1.9	-	$\qquad \qquad -$	0.2	-	-	3.9	-	
Thysanoptera	1.4	$\overline{}$	$\overline{}$	0.1	-	$\qquad \qquad -$	< 0.1	-	$\overline{}$	
Collembola	11.1	5.7	7.1	1.0	1.6	0.4	< 0.1	0.2	< 0.1	
Protura	2.8	$\qquad \qquad -$	$\qquad \qquad -$	0.2	$\qquad \qquad -$	$\overline{}$	< 0.1	$\overline{}$	-	
Arachnida										
Araneae	26.4	20.8	28.6	1.7	2.6	1.9	3.2	1.4	1.4	
Opiliones	$\overline{}$	$\overline{}$	$\qquad \qquad -$	\equiv	-	$\overline{}$	$\overline{}$	—	$\overline{}$	
Acarina	4.2	$\overline{}$	$\overline{}$	0.2	$\overline{}$	$\overline{}$	< 0.1	$\overline{}$	$\overline{}$	
Isopoda	5.6	5.7	7.1	1.5	0.7	0.4	8.8	4.8	0.2	
Decapoda	1.4	5.7	$\overline{}$	0.1	0.7	$\qquad \qquad -$	2.2	4.8	$\overline{}$	
Amphipoda	$\overline{}$	1.9	$\overline{}$	$\overline{}$	0.5	$\overline{}$	-	0.4	$\overline{}$	
Chilopoda	2.8	\equiv	$\overline{}$	0.1	-	$\overline{}$	0.3	-	$\overline{}$	
Diplopoda	20.8	1.9	42.9	1.9	0.2	3.8	6.7	0.1	6.3	
Gastropoda	25.0	1.9	21.4	2.7	0.2	1.2	3.3	0.2	0.8	
Oligochaeta		$\overline{}$	21.4	$\qquad \qquad -$	$\qquad \qquad -$	1.5	$\overline{}$	$\overline{}$	23.2	
Plant materials	19.4	30.2	71.4	$\overline{}$	-	$\overline{}$	$\overline{}$	÷	$\overline{}$	
	25.0	39.6	42.9		-					
Minerals				$\overline{}$		$\overline{}$	$\overline{}$	$\overline{}$	-	

Table 1. Dietary comparison of R. rugosa from the reservoir (1577 prey from 72 individuals, total volume 10902.4 mm³), the river shoreline (431 prey from 53 individuals, total volume 5691.5 mm³), and the paddy fields (262 prey from 14 individuals, total volume 4323.23 mm³).

Fig. 1. Relationships between frog SVL and minimum () and maximum () volumes of prey in stomach contents.

Prey selection

Dipterans and hemipterans were abundant prey taxa in the sweep samples, whereas ants were the most abundant prey taxon in the ground plot samples. Rana rugosa consumed prey animals occurring both above ground and on the ground. The relative abundances of prey taxa in the habitat were significantly correlated with their abundances in frog stomach contents ($τ=0.470, P<0.05$; Table 2).

The proportion of ants in the environment (81.3%) outweighed that found in the frog diet (56.9%), and both E (-0.18) and E^{*} (-0.50) indicated that R. rugosa avoided ants.

DISCUSSION

Regardless of differences in body size or habitat environments, R. rugosa proved to consume ants very frequently, and the prey represented surprisingly similar proportions in the frequency of occurrence (81.9–85.7%) and in number

Fig. 2. Relationships between frog SVL and the frequency of occurrence of major prey taxa.

(56.8–59.4%) among the three populations studied. From these large proportions of ants, R. rugosa may be viewed as an ant specialist. However, Simon and Toft (1991) suggested that ant specialists should consume ants higher in proportions than those found in the environment. Since there is no other definition for ant specialisits, we followed Simon and Toft (1991). Such a definition did not hold for R. rugosa, and this species is not regarded as a true ant specialist. Even so, the large proportion of ants taken by R. rugosa (56.8–59.4%) is within the range for those species that have been described as ant specialists (52–99%; Toft, 1980, 1981; Lieberman, 1986).

In fact, the proportion of ants was larger in the environment than in the frog diet, and both electivity indices (E and E^*) indicated that R. rugosa avoided ants. However, ant avoidance in R. rugosa was distinctively weaker than in another rice field dwelling ranid, R. nigromaculata ($E=-0.60$, $E^*=-0.86$; Hirai and Matsui, unpubl. data). This weak avoidance to ants by R. rugosa may result in its food habit to take prey in response to their relative abundance in the environment where ants were the most abundant.

According to available dietary reports from various areas of the world, ranid frogs are grouped into generalist predators that consume a wide variety of prey organisms, responding to prey availability in the environment (e.g., Turner, 1959; Houston, 1973; Premo and Atomowidjojo, 1987). Rana rugosa also consumed a various kind of prey items, but ants made up a prominently large proportion in number (56.8–59.4%) that has never been reported for Rana (see Appendix 1). For example, in the diet of R. limnocharis and R. nigromaculata, that are often found syntopic with R. rugosa, ants represent at most 13.5–27.9% and 15.1–20.3%, respectively (Liu and Chen, 1933; Berry, 1965; Mohanty-Hejmadi and Acharya, 1982; Hirai and Matsui, 1999, see Appendix 1). Even the largest proportional value of ants in Rana that has ever been reported is only 35.0% (R. blythi: Inger and Greenberg, 1966), and therefore, it is clear that values of 56.8–59.4% found in R. rugosa are exceptionally large for Rana. This result suggests that R. rugosa has very peculiar feeding habits that are different from the other ranid frogs so far studied. This is probably due to weak ant avoidance by this species as mentioned above. Thus, we could confirm the validity of the anecdotal remarks that R. rugosa eats markedly numerous ants (Maeda and Matsui, 1989).

An animal must maintain a positive energy budget to grow, reproduce, and survive periods of inactivity. Optimal foraging theory suggests that larger animals should select large prey and overlook small prey which is too low to warrant energy involved in their pursuit and consumption (Schoener, 1979). In fact, a hylid frog, Pseudacris triseriata, excludes smaller prey from its diets as it ingests increasingly larger prey with its growth (Christian, 1982). Similar results have been obtained in R. nigromaculata; ants were important prey for metamorphosed juveniles of this species (SVL<40 mm), and represent 45% in their diets. However, subadults or adults (SVL≥40 mm) became less incorporated ants into their diets, and the proportion of ants decreased to less than 20% (Hirai and Matsui, 1999).

Rana rugosa also consumed larger prey such as beetles

		Sweep		Ground plot		Combined	
Prey taxa	N	%	N	%	N	%	℅
Insecta							
Hymenoptera							
Formicidae	179	14.2	499	83.0	40099	81.3	56.9
non-Formicid	37	2.9			37	< 0.1	1.9
larvae	5	0.4	$\overline{}$		5	< 0.1	0.4
Coleoptera	149	11.8	21	3.5	1829	3.7	19.5
larvae	8	0.6	2	0.3	168	0.3	1.5
Diptera	351	27.8			351	0.7	5.7
larvae							0.8
Lepidoptera	14	1.1			14	< 0.1	
larvae	10	0.8			10	< 0.1	1.5
Neuroptera	$\overline{2}$	0.2		-	$\overline{2}$	< 0.1	-
Hemiptera	327	25.9	4	0.7	647	1.3	2.3
Dermaptera	-	-	$\overline{7}$	1.2	560	1.1	-
Orthoptera	91	7.2	1	0.2	171	0.4	0.4
Collembola							0.4
Arachnida							
Araneae	89	7.1	32	5.3	2649	5.4	1.9
Acarina	$\overline{2}$	0.2	-	-	$\overline{2}$	< 0.1	-
Isopoda			21	3.5	1680	3.4	0.4
Diplopoda			10	1.7	800	1.6	3.8
Gastropoda							1.2
Oligochaeta			4	0.7	320	0.7	1.5

Table 2. Comparison of the relative abundances of prey taxa in the habitat with their abundances in the frog diet. See text for total abundances of prey taxa in combined samples.

more frequently and smaller prey such as dipterans less frequently with the increase of body size. However, ants were constantly consumed in a high frequency regardless of body size, and we could not detect ontogenetic change in ant consumption. Therefore, we presume that R. rugosa feeds on ants because it does not strongly avoid these most abundant potential prey in the habitat.

Prey availability is proved as one of the most important factors to determine what frogs eat (Labanick, 1976; Galatti, 1992; Hirai and Matsui, 1999). In generalist predators such as ranids, their diet compositions are known to vary among habitats. For example, R. catesbeiana inhabiting the pond consumes mainly insects, but the frog eats mostly crustaceans such as crayfish in the stream (Tyler and Hoestenbach, 1979). Similarly, R. cancrivora eats mostly crustaceans in brackish water habitats, but insects in fresh water habitats (Elliott and Karunakaran, 1974). We examined the diets of R. rugosa from three environmentally different habitats, and also detected differences in frequency of occurrence of beetles and in numeric proportions of beetles and dipterans among different habitats. Therefore, the diets of R . rugosa were judged to be influenced by prey availability in the habitats. However, differences were not detected for ants in either the frequency of occurrence or the numeric proportion; ants remained dominant in the diet in surprisingly similar proportions in all the three habitats (reservoir=56.8%, shoreline=59.4%: paddy fields=56.9%) despite their small volumetric contributions (4.1– 13.2%). This result appears to conform to the presumption that R. rugosa shows only weak avoidance to ants.

Ant specialists such as dendrobatids are known to have

toxic skin secretions that serve a defensive role. Daly (1995) and Daly et al. (1994) suggested that the toxic skin secretions were sequestered from their dietary sources. Since their diets principally consisted of ants, Toft (1995) and Caldwell (1996) suggested that anti-predator tactics were highly correlated with ant specialized diets. Rana rugosa is also presumed to have distasteful skin secretions by which the species is rarely predated by snakes in nature (Mori and Moriguchi, 1988). Actually, avoidance of R. rugosa by snakes has been demonstrated experimentally, by offering this species and Hyla japonica to Elaphe quandrivirgata, a generalist predator (Mori, 1989). From these knowledge, myrmecophagy by R. rugosa may also be correlated with predator defence in this species.

Feeding patterns of myrmecophagous dendrobatids were shown to be correlated with not only anti-predator tactics but also foraging strategy, morphology, and physiology (Toft, 1980, 1981). More detailed investigations of feeding habits in R. rugosa may help us to understand the ecological significance of weak ant avoidance by this species.

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Appendix 1

Published accounts of ant–eating in the genus Rana. Abbreviations: F=frequency of occurrence; N=numeric proportion; V= volumetric proportion.

Species	Size (mm) or sex	Proportion (%)				Sample size	Locality	Authority
		F	N	\vee	Frog	Prey		
	$27 - 58$	28.4	-	4	95	$\qquad \qquad -$	New York	Whitaker, 1962
		45*	4.3		20	184	Canada	Moore and Strickland, 1954
	$27 - 58$	28.4		4	95	$\overline{}$	New York	Whitaker, 1962
	$20 - 35$	14.2	-	2.9		-	New York	Linzey, 1967
	$50 - 80$	15.3		0.9	-	-	New York	Linzey, 1967
		60.9	14.8		23	162	Minnesota	Hedeen, 1972
		33.3			6	$\overline{}$	Nebraska	Frederick and Collette, 1959
			6.2	-	50	373	Montana	Miller, 1977
R. pirica			1.1		50	281	Japan	Inukai, 1925
R. pretiosa		< 27.5		-	142	$\overline{}$	Wyoming	Turner, 1959
		24.3	-	4.4	206	$\qquad \qquad -$	Oregon	Whitaker et al., 1983
	$28 - 33$	16.7	6		18	116	Canada	Licht, 1986
	>40	26.1	3.7		23	191	Canada	Licht, 1986
		22.0	4.6		41	307	Canada	Licht, 1986
			19.3		50	517	Montana	Miller, 1977
R. rugosa	$24.6 - 58.1$	$81.9 - 85.7$	$56.8 - 59.4$	$4.1 - 13.2$	$14 - 72$	262-1577	Japan	This study
R. septentrionalis		$27.0*$		$2.4*$	159	2503	New York	Kramek, 1972
		28.0	-	3.8			New York	Stewart and Sandison, 1972
		$10.5 - 52.2$	$1.2 - 6.8$		$18 - 23$	234-1421	Minnesota	Hedeen, 1972
R. temporaria		2	0.1	< 0.1	359	6681	England	Houston, 1973
		$29*$	$6*$		70	-	Ireland	Blackith and Speight, 1974
	30		$<15*$		$58 - 194$	$\overline{}$	Sweden	Loman, 1979
	$30 - 47$		$< 5*$		$46 - 89$	$\overline{}$	Sweden	Loman, 1979
			0.4		17	228	England	Savage, 1961
			10.1	$\overline{}$	30	503	France	Pilorge, 1982
R. tigerina	$100 - 143$	$<$ 10			819	-	Pakistan	Khan, 1973
R. vaillanti		0	$\mathbf 0$	-	819	-	Mexico	Ramirez et al., 1998

Appendix 1. (cont.)

+=Unspecified small amount

* =Proportion of Hymenoptera