# Heavy, Bulky, or Both: What Does "Large Prey" Mean to Snakes?

HARRY W. GREENE<sup>1,2,4</sup> AND KEVIN D. WISEMAN<sup>3</sup>

<sup>1</sup>Biodiversity Center and Department of Integrative Biology, The University of Texas at Austin, Austin, Texas, 78712, USA
 <sup>2</sup>Museum of Vertebrates and Department of Ecology and Evolutionary Biology, Cornell University, Ithaca, New York, 14853, USA
 <sup>3</sup>Department of Herpetology, California Academy of Sciences, Golden Gate Park, San Francisco, California, 94118, USA

ABSTRACT.—We lay out mass-bulk theory (MBT)—relationships for gape-limited predators among relative prey mass (RPM), relative prey bulk (RPB), prey shape, prey taxon, and feeding frequency. Elongate reptiles with narrow mouths eat tiny items; chunks of prey; or, as with many snakes, "large" animals ingested intact. RPM and RPB define item size, with implications for costs and benefits of feeding. Prey are heavy, bulky, both, or neither, only relative to consumers. Type I items are not heavy or bulky; they require minimal handling and gape, but many must be eaten. High RPM, high RPB, or both characterize types II, elongate (e.g., eels); III, fusiform to ovoid (e.g., mice); and IV, nonuniform in cross-sectional dimensions (e.g., some fishes), density (e.g., birds), and/or deformability (e.g., crustaceans). High handling costs and payoffs characterize types II and III; III and IV require wider gape, but IV comes with lower RPM, costs, and payoffs. RPM and RPB have implications for biology and conservation, such that heavy, bulky, or heavy and bulky—but not large—usefully describe prey size. We explore MBT with 1) natural history vignettes and graphical integration of RPM, RPB, prey shapes, and feeding frequency; 2) nonvenomous colubrids that vary in gape and diet; (3) front-fanged colubroids that consume lizards, centipedes, or earthworms; and (4) bird-eating snakes. Further testing of MBT is hampered by logistical and cultural challenges. Our explorations are bookended by reflections of a herpetologist enjoying his eighth decade, emphasizing how an early-career publication stemmed from youthful experiences and led to this review.

RESUMEN.—HWG reflexiona sobre como una publicación en los inicios de una carrera académica se originó de experiencias juveniles y derivó en la vida de un herpetólogo disfrutando su octava década. Relacionado con ello, definimos la teoría masa-volumen (TMV)-la relación para predadores limitados por el tamaño de su boca entre la masa relativa de su presa (MRP), el volúmen relativo de la presa (VRP), la forma de la presa, el taxon de la presa, y la frecuencia de alimentación. Los reptiles de cuerpos alargados y bocas estrechas obtienen su alimento de porciones o presas pequeñas, de partes pequeñas de presas grandes, o de presas "grandes" ingeridas completas, como lo hacen muchas serpientes. MRP y VRP definen el tamaño de la porción, y tienen implicaciones en los costos y beneficios de la alimentación. Las presas pueden ser pesadas, voluminosas, ambas o ninguna, en relación con su depredador. Las presas de tipo I, no son pesadas ni voluminosas; son fáciles de manipular y no requieren bocas grandes, pero deben consumirse muchas. MRP alta, VRP alta, o ambas, definien a las presas tipo II, alargadas, como las anguilas; tipo III, esferoides a ovoides, como los ratones; y tipo IV, no uniformes en sección transversal, como algunos peces, o no uniformes en densidad, como las aves. Costos de manipulación y beneficios nutricionales altos caracterizan a los tipos II y III; los tipos III y IV requieren bocas más grandes, pero el tipo IV implica una MRP baja, y costos y beneficios nutricionales más bajos. MRP y VRP tienen implicaciones biológicas y de conservación, de manera que es útil describir la presa como pesada, voluminosa, o pesada y voluminosa-pero no grande. Aquí exploramos la TMV con 1) viñetas de historia natural e ingración gráfica de MRP, VRP, tipos de formas de presas, y frecuencia de alimentación; 2) colúbridos no venenosos, que varían en tamaño de boca y en dieta; 3) colubroideos con colmillos frontales que consumen lagartijas, ciempiés, o gusanos de tierra; y 4) serpientes que consumen aves. Pruebas adicionales de la Teoría Masa-Volúmen son obstaculizadas por retos logísticos y culturales.

"[The *Python sebae*'s stretched] skin when dry was 25 feet 2 inches [7.7 m] long ... stomach of the snake contained not less than one peck [approximately nine liters] of brass, copper, and iron rings, such as the natives wear on the arms and legs ... A snake of that size would swallow an antelope as large as a cow, horns and all." (Johnston, 1908:270)

"Natural history is replete with observations of feeding, yet only recently have investigators begun to treat feeding as a device whose performance—as measured in net energy yield/feeding time or some other units assumed commensurate with fitness—may be maximized by natural selection." (Schoener, 1971:369)

"Too many workers continue to publish lists of prey species eaten, without regard to the size and seasonal energetic requirements of the snake or to the availability and nutritional content of prey in the environment." (Godley, 1980:411)

Observations of serpents ingesting humans and other "large" animals in one piece must be far older than written history (e.g., Isbell, 2009; Headland and Greene, 2011), although what prey size means in this context often has been vague, even among herpetologists-a 15-kg venison medallion or salmon fillet, immense by our standards, would be small if scaled to the masses of many snakes and their meals (Figs. 1, 2). These limbless reptiles generally feed infrequently, and their diets have been revealed by field observations (e.g., Trail, 1987; Ribble and Rathbun, 2018; Groen et al., 2020), necropsies and regurgitations (e.g., Fitch, 1960; Luiselli and Akani, 2003; Boback et al., 2016), and museum specimen stomach contents (e.g., Werner, 1909; Schmidt, 1932; Klauber, 1956). Now, data also flow from stable isotopes (e.g., Willson et al., 2010; Durso and Mullin, 2017), fecal DNA (e.g., Brown et al., 2014; Durso et al., 2022), roadkill (Hoefer et al., 2021), remote cameras (e.g., Robinson et al., 2005; Putman and Clark, 2015; Glaudas et al., 2017a), and community science (e.g., Maritz and Maritz, 2020; Durso et al., 2021; Putman et al., 2021). Following that brief preface, this coauthored perspective begins and ends in first person

<sup>&</sup>lt;sup>4</sup>Corresponding author. E-mail: harry.greene@austin.utexas.edu, *Mailing* address: Rancho Cascabel, P. O. Box 178, Pontotoc, Texas, USA 76869 DOI: 10.1670/22-068



FIG. 1. Mammals can be heavy and bulky prey—"large" in handling costs, nutritional payoffs, and required gape—as illustrated by Boa Constrictors (*Boa constrictor* sensu lato) that ate White-tailed Deer (*Odocoileus virginianus*) in Sector Santa Rosa, Guanacaste Conservation Area, Guanacaste Province, Costa Rica (data and photos: D. H. Janzen and W. Hallwachs). (A) During and (B) shortly after ingestion of a 4-kg fawn by a 10-kg boa, 5 April 2013 (masses estimated from similar-sized conspecifics). (C) Forced regurgitation of a 3.5-kg fawn by a 3-kg boa, 1983 (weighed in the field; right, D. H. Janzen; left, E. Carrillo). They are fusiform prey with relative prey mass (RPM) of ~0.4 (A, B), which is not unusual for snake meals, and RPM of ~1.17 (C), which is heavier than typically consumed by snakes other than boids, pythonids, and front-fanged species.

singular, whereby HWG details how an early-career publication on the evolution of feeding in snakes (Greene, 1983a) grew out of youthful experiences and then reflects on life for a herpetologist enjoying his eighth decade. In between, we (HWG and KDW) review research that helps better elucidate relationships among relative prey mass (RPM), relative prey bulk (RPB), prey shape, prey taxonomic identity (ID), and feeding frequency what we call mass-bulk theory (MBT).

I (HWG) first thought about snakes eating large meals as a recent high school graduate interning with Henry Fitch and Charles "Jay" Cole at the University of Kansas Museum of Natural History. My assignment that summer of 1963 was to dissect preserved skinks and assay their breeding cycles (Fitch and Greene, 1965; Greene, 1969), but accounts of snake prey (e.g., Schmidt, 1932; Klauber, 1956; Fitch, 1960) and field encounters with Western Massasauga Rattlesnakes (*Sistrurus tergeminus*) led me to also assess that species' diet with museum specimens (Greene and Oliver, 1965). Then, while off-duty in the military, I recorded scars on amphisbaenians and snakes in European museums to test hypotheses about their defensive tail displays (Greene, 1973a). For an M.A. at the

University of Texas at Arlington (UTA), advised by William F. Pyburn, I studied feeding in venomous New World coralsnakes (Micruroides and Micrurus), again with museum specimens (Greene, 1973b, 1976, 1984). For my Ph.D. at the University of Tennessee, Knoxville, supervised by Gordon M. Burghardt, I used observations of defense and constriction to address homology, convergence, and the origins of novel behavior in snakes (H.W. Greene, 1977, 1979, 1994, 1999; Greene and Burghardt, 1978). Upon completion of graduate work, I had pondered hundreds of natural prey items and more than a thousand captive feeding events, based on phylogenetically basal (e.g., pipesnakes [Cylindrophis, Uropeltidae], Mexican Burrowing Pythons [Loxocemus bicolor, Loxocemidae], and dwarf boas [Tropidophis, Tropidophiidae]) to highly derived taxa (e.g., stilettosnakes [Atractaspis, Atractaspididae], king cobras [Ophiophagus, Elapidae], and mock vipers [Psammodynastes, Lamprophiidae]). Those experiences, along with foundational papers on diet (Fitch, 1941; Fitch and Twining, 1946), functional morphology (e.g., Gans, 1961; Boltt and Ewer, 1964), phylogenetics of character variation (Rabb and Marx, 1973), and optimal foraging (MacArthur and Pianka, 1966; Schoener, 1971), led me to wonder why snakes eat some prey but not others.



FIG. 2. Aquatic chordate prey of natricine colubrids range from elongate to fusiform or bulky and from light to heavy. (A) Northern Watersnake (*Nerodia sipedon*) grasping a Chestnut Lamprey (*Ichthyomyzon castaneus*; identified by B. H. Bauer), 30 April 2021, Wolf River, Fayette County, Tennessee (data and photo: D. P. Hailey). (B) Terrestrial Gartersnake (*Thamnophis elegans*; TL ~43 cm) grasping a Mottled Sculpin (*Cottus bairdii*), late afternoon, 25 July 2019, Warm Springs Creek, Sun Valley, Blaine County, Idaho (data and photo: M. and J. W. Fitzpatrick). (C) Diamond-Backed Watersnake (*Nerodia rhombifer*; TL ~75 cm) ingesting a Gizzard Shad (*Dorosoma cepedianum*; identified by D. S. Hendrickson; evidently carrion, TL ~22 cm), 29 July 2009, Village Creek Heritage Park, Arlington, Tarrant County, Texas (data and photo: T. D. Hibbitts). (D) Mississippi Green Watersnake (*Nerodia cyclopion*) ingesting a sunfish (*Lepomis* sp.; identified by A. A. Echelle); Cane Bayou, Lacombe, St. Tammany Parish, Louisiana (data and photo: J. Schauer). Completed predation was not witnessed in these incidents. For (A), (C), and (D), the first likely had high relative prey mass (RPM) and low relative prey bulk (RPB), the latter two likely had low RPM and high RPB; (B) likely was intermediate in RPM and RPB, although the pectoral fins might have enhanced RPB.

At the 1977 American Society of Ichthyologists and Herpetologists meeting, I nervously presented "Behavioral, ecological, and morphological aspects of adaptive radiation in snakes" (Collette, 1977:814). My so-called "preliminary working model" specified item size with prey/predator mass ("weight ratio" [WR]) and prey diameter/predator head diameter ("ingestion ratio" [IR]). I expected handling costs and payoffs would increase with higher WR and gape with higher IR; prey types were described as small and any shape (low WR, low IR), elongate (high WR, low IR), ovoid (high WR and IR), irregular (low WR, high IR), or fusiform (moderate WR and IR). Pilot comparisons supported the model's predictions about the evolution of methods for subduing prey, gape, and foraging trade-offs, of which later explorations were published (Greene, 1983a, 1984, 1986a, 1992, 1997, 2013; Losos and Greene, 1988; Rodríguez-Robles et al., 1999a; Cundall and Greene, 2000; Wiseman et al., 2019). Meanwhile, early on, Shine (1977) and Godley (1980) had used mass to assess prey for six snake species and foraging trade-offs between prey ID within a species, respectively, and

Voris and Voris (1983) examined prey shapes and gapes in Seasnakes.

Subsequent decades have entailed an explosion of interest in snake biology, within which we (HWG and KDW) conclude that MBT has had significant but patchy effects. Beyond the studies cited in the previous paragraph, the deconstruction of prey size (Figs. 3, 4) into RPM (previously WR) and RPB (previously IR) have influenced some discussions of snake biology (exemplified by references cited in Appendix 1). However, often research on snakes has not used them or has done so ambiguously (Appendix 2). Referring to snake prey, for example, Brecko et al. (2011) assumed fish are less bulky than frogs regardless of mass; Mociño-Deloya et al. (2015) treated all lizards as "small" and mammals as "large"; and Moon et al. (2019) in a comprehensive review frequently alluded to "large" prey, usually without reference to RPM or RPB. Likewise, some taxon-focused reports have provided data pertinent to MBT, typically prey ID and RPM (Appendices 3-5), but many other diet studies mention neither RPM nor RPB (Appendix 6).



FIG. 3. Idealized prey size and shape types and the components of gape. Small type I prey are not heavy or bulky, regardless of shape and taxonomy; type II prey are elongate (e.g., eels); type III prey are fusiform to ovoid (e.g., rodents); and type IV prey are noncircular in cross-section (e.g., some fishes), nonuniform in density (birds), or both. (A) Dorsal view of a generalized snake with simple external measurements that might reflect gape; HL = head length from snout to retroarticular process of the quadrate, HW = head width at widest point. (B) Simplified view of some bony elements in a snake that might influence gape (G), illustrating supratemporals (Su), quadrates (Qd), and mandibles (Md), as well as the elastic tissue connection (Etc) where most amniotes have a firm mandibular symphysis (modified from Arnold, 1983; animal silhouettes in this and Fig. 4 were adapted from phylopic [http://phylopic.org/]).

Among "Natural History Notes" we surveyed in the first 2021 issue of Herpetological Review, 39 diet records for 33 snake species (27 genera) include 6 (15%) with RPM data; for 33 other prey (85%), RPM could have been recorded for at least 3 and perhaps 5 more because specimens were deposited in museums, so the total could have been 14 (36%). None of the 39 records addressed RPB.

Several goals justify gathering diet data, from answering questions about morphology, physiology, ecology, ethology, evolution, and conservation to furthering nature appreciation with public outreach. Moreover, different applications might prioritize certain information-prey ID for ecological questions (e.g., Greene and Jaksic, 1983; Luiselli, 2006a; Pinto-Cuelho et al., 2021), RPM for foraging behavior (e.g., Arnold, 1993; Andreadis and Burghardt, 2005; Loughran et al., 2013; Glaudas et al., 2019), RPM and RPB for evolutionary and functional morphology (Cundall and Greene, 2000; Vincent et al., 2006a; Cundall et al., 2014; Moon et al., 2019; Gripshover and Jayne, 2021; Cundall and Irish, 2022; Jayne et al., 2022), and all of them for conservation and education (e.g., Greene, 1997, 2003, 2013; Clayton and Myers, 2015; Mehta et al., 2020). MBT is clearly germane to many aspects of snake biology, and yet its key parameters often have gone unmeasured, perhaps in part because Greene (1983a) ineffectively portrayed them. Although diet records and broader studies absent MBT can be useful,



FIG. 4. Graphical model illustrating the relationship between prey types I–IV in terms of relative prey mass (x-axis), relative prey bulk (y-axis), and time between feedings (z-axis). See text for additional explanation.

Godley's complaint (1980; quoted above) still rings true—many accounts of snake diets are simply prey ID lists or are based upon them.

We believe in core roles for natural history within biology and art in clarifying science (e.g., Greene, 2005a, 2005b, 2013; Wiseman and Bettaso, 2007; Wiseman, 2018). This paper, therefore, first explores verbally and visually "large prey" and its implications for MBT. In three following sections, we illustrate MBT with nonvenomous colubrids that vary in diet and gape; front-fanged colubroids that feed on lizards, centipedes, and earthworms; and snakes that eat birds. We next emphasize gathering data for RPM and RPB in taxon-focused and broader studies and then comment on logistical and cultural impediments to that task. Throughout this paper, we detail specific predator-prey interactions to promote acquiring useful information for future syntheses; we provide extensive literature citations to support our conclusions, rather than as an exhaustive review of snake feeding biology (but see, e.g., Moon et al., 2019; Cundall and Greene, 2000; Grundler, 2020; Cundall and Irish, 2022).

Abbreviations refer to California Academy of Sciences (CAS); Museum of Vertebrates, Cornell University (CUMV); Robert W. Hansen field catalog (RWH); Harry W. Greene field catalog (HWG); Museum of Comparative Zoology, Harvard University (MCZ); Robert L. Seib field catalog (RLS); Museum of Vertebrate Zoology, University of California, Berkeley (MVZ); Texas Natural History Collection, University of Texas at Austin (TNHC); snout–vent length (SVL); total length (TL); and carapace length (CL). Among the taxa discussed here (see Pough et al., 2016), Scolecophidia (including Typhlopidae) and Alethinophidia are treated as basal lineages of Serpentes ("snakes"; Head et al., 2020). Within Alethinophidia, Colubroidea is successively more distantly related to Acrochordidae, Boidae plus Pythonidae, Loxocemidae, Uropeltidae (including

parentheses): Glossy Snakes (Arizona elegans; Rodríguez-Robles et al., 1999a), California

TABLE 1. Dietary attributes for the following six species of North American colubrids (sources in parentheses): Glossy Snakes (Arizona elegans; Rodríguez-Robles et al., 1999a), California Kingsnakes (Lampropeltis getula californiae; Wiseman et al., 2019; taxonomy follows Hillis, 2020), Scarlet Kingsnakes (Lelapsoides; Greene et al., 2010), California Mountain Kingsnakes (L. zonata)



A dozen or more ant larvae regurgitated by a Bibron's FIG. 5. Blindsnake (Afrotyphlops bibronii), 16 December 2022, KwaSani, Underberg, KwaZulu-Natal, South Africa (photo: M. da Fonseca). Assuming these insects were encountered as a single meal, overall RPM would have been  $>10\times$  that of each individual type I prey item; approximately the same gape would have been required for an elongate type II prey item with the same diameter as each larva.

Cylindrophis), and Aniliidae plus Tropidophiidae. Colubroidea encompasses Atractaspididae, Colubridae (including Colubrinae, Dipsadinae, Natricinae), Elapidae, Homalopsidae, Lamprophiidae, and Viperidae. Front-fanged colubroids include atractaspidids Atractaspis and Homoroselaps, elapids, and viperids.

# LONG BODIES, SMALL MOUTHS, AND MASS-BULK THEORY

Body elongation repeatedly preceded limb loss in tetrapod evolution (Mann et al., 2022), and reduced diameter entails a narrower mouth (Gans, 1961). Other than by lowering metabolic rates, attenuate squamates compensate for a narrow mouth by eating many tiny organisms (e.g., >50 ants/stomach in some typhlopids, Webb and Shine, 1993a; "nibblers," Andreadis and Burghardt, 2005; Fig. 5), parts of bigger ones (Appendix 7), or "spectacularly large prey" (Gans, 1961:217; "gorgers," Andreadis and Burghardt, 2005), as do many snakes (e.g., Moon et al., 2019; Cundall and Greene, 2000; Cundall and Irish, 2022) and a near-limbless gekkotan (Burton's Flap-footed Lizard, Lialis burtonis; Patchell and Shine, 1986). Conversely, most limbed lizards (including many varanids; Shine and Thomas, 2005; see Losos and Greene, 1988) frequently consume small items-the mean number of prey per stomach was 6.0-75.8 for six North American species (Pianka, 1970; Pianka and Parker, 1972; Parker and Pianka, 1973; Parker and Pianka, 1974; Pianka and Parker, 1975); means were 1.07-2.16 for five colubrid species from the same region, as predicted by MBT (see below), and mean RPMs were 0.19-0.33 (Table 1).

Gans (1961), by posing the small mouth problem in terms of food item value, implicitly identified RPM as crucial to understanding large prey. However, he construed biomechanical solutions (e.g., mandibular liberation, kinetic palatomaxillary arches, and unilateral feeding) in terms of prey "cross-sectional area" (Gans, 1961:220), a component of RPB. The scene

	A. elegans	L. getula californiae	L. elapsoides	L. zonata	P. catenifer	R. lecontei
Predator SVL (cm)	21–95 100	23–115	20-50	24-71	26–160 417	21–93
Snakes with prey Prev items	107	390 447	34 34	51	$^{41/}_{1066}$	110
Proportion of diet: reptiles, mammals, birds	0.50, 0.44, 0.04	0.59, 0.29, 0.11	0.97, 0.03, 0.0	0.84, 0.12, 0.04	0.03, 0.75, 0.20	0.73, 0.26, 0.0
Prev species	16	75	6	10	66	17
Preý items/stomach	$1-3 \ (\overline{x} = 1.07)$	$1-3 \ (\overline{x} = 1.13) \\ 0 \ 01 \ 0 \ 72 \ (\overline{x} = 0.24)$	$\begin{array}{c} 1-2 \ (\overline{\mathrm{X}}=1.1) \\ 0 \ 11 \ 0 \ 28 \ (\overline{\mathrm{Y}}=-0 \ 10) \end{array}$	$1-3$ 0.11 0.62 ( $\overline{v}$ - 0.22)	$1-35 (\overline{x} = 2.6)$	$\begin{array}{c} 1-2 \ (\overline{\mathrm{X}}=1.16) \\ 0.02 \ 0.62 \ (\overline{\mathrm{X}}=0.02) \end{array}$
RPB discussed?	V.07 - V.7 (V V V V V V V V.	$V.01-0.7 (\Lambda - 0.24)$	$(\alpha - \alpha) \circ \alpha$	V = V = V = 0	$V_{12} = V_{12} = V_{12} = V_{12}$	$V_{\rm CD} = V_{\rm CD} = V_{\rm CD} = V_{\rm CD} = V_{\rm CD}$
Proportion of head-first ingestion	1.0	0.95	1.0	0.88	0.89	1.0
Geographic variation	Not studied	Yes	No	No	Yes	Yes
Seasonal variation	Not studied	Yes	Not studied	Yes	Not studied	Not studied
Ontogenetic diet shift	Partial	Partial	No	Partial	Yes	Partial



Ways to be a type IV prey, with low relative prey mass FIG. 6. (RPM) and high relative prey bulk. (A) Nonuniform cross-sectional dimensions because of nondeformable shell: adult Cottonmouth (Agkistrodon piscivorus) eating a juvenile Slider (Trachemys scripta), Maxwell Air Force Base, Montgomery County, Alabama, 1102 h, 26 May 2014; snake TL, ~75-90 cm; turtle CL, ~7.5 cm; ingestion required ~30 minutes and RPM likely <0.1 (data and photo: R. Dowling). (B) Nonuniform cross-sectional dimensions because of wings and nonuniform density because of feathers: young adult female Western Diamond-backed Rattlesnake (Crotalus atrox) grasping adult male Acorn Woodpecker (Melanerpes formicivorus; identification, age, sex, and estimated mass of 80 g by W. D. Koenig), ingestion required ~180 min, Portal, Cochise County, Arizona, 2 August 1999; TL, 79 cm; mass, 380 g including prey; RPM, ~0.27 (CUMV 13952; photo: H. W. Greene). (C) Facultative increase in cross-sectional dimensions and decrease in density because of lung inflation: adult Common Gartersnake (Thamnophis sirtalis) grasping a Southern Toad (Anaxyrus terrestris), St. Augustine, St. Johns County, Florida, 7 May 2022; completed ingestion was not observed (data and photo: K. Glaser).

was, thus, set for ongoing confusion of two distinct and yet interactive size parameters, despite efforts to clarify these relationships (e.g., Greene, 1983a; Arnold, 1993; Greene, 1997:71–73; Cundall and Greene, 2000; King, 2002; Vincent et al., 2006a; Greene 2013:151–155). As an example of conflating mass and bulk subsequent to Gans (1961), "the largest prey item recorded for any snake is a 59 kg impala consumed by a 4.72 m African python [*Python sebae*] ... The shoulders of an adult man when collapsed forward may measure only 35–40 cm wide, and could probably be engulfed by pythons in excess of 5 m" (Branch and Hacke, 1980:306).

Prey size should be defined by RPM and RPB because, as detailed below, they have different implications for costs and benefits of feeding. Prey can be "large" in one, both, or neither parameter (Figs. 1-5) but only relative to masses and gapes of individual snakes who subdue, consume, and process them or not. Prey taxa are not intrinsically heavy or bulky but can be described in terms of four types. Type I items with low RPM and RPB are not heavy or bulky, regardless of shape; their masses and cross-sectional areas are trivial to predators, so they require neither subduing nor big gapes to be swallowed; and they must be eaten often to provide adequate nutrition (Figs. 3-5). "Large" prey with high RPM, high RPB, or both define the following three additional idealized shape types: II, elongate (e.g., eels; Figs. 2-4, 9); III, fusiform to ovoid (e.g., mammals; Figs. 1, 3, 4); and IV, noncircular in cross-section, density, and/or deformability (e.g., many fishes and birds; Figs. 2–4, 6, 7). Among these shape types, with all else equal, high handling costs (e.g., Arnold, 1993; Andreadis and Burghardt, 2005; Mukerjee and Heithaus, 2013; Kornilev et al., 2022), high payoffs, and low feeding frequency characterize II and III. Types III and IV require wide gape to surmount high RPB; type IV items also come with lower meal payoff because of nonuniform cross-sectional dimensions or density and, thus, require increased feeding frequency or other compensation, e.g., low energy demands.

All else is rarely equal, and prey can vary in taxon-typical attributes such as retaliatory bite force (e.g., amphisbaenians, Barbo and Marques, 2003), nutritional content (e.g., Krause et al., 2003; Wiseman et al., 2019), and surface features (e.g., Godley, 1980; Savitzky, 1983; Voris and Voris, 1983; Arnold, 1993; Willson and Hopkins, 2011; Bringsøe, 2019; Wiseman et al., 2019; Hamanaka and Mori, 2020; Cleuren et al., 2021). They also might differ in ways evident only at high RPM, high RPB, or both, including toxicity (e.g., some amphibians, Feldman et al., 2012, 2020), social defense (e.g., carnivores, Janzen, 1970; primates, Gardner et al., 2015), and shape changes (e.g., lizard ring-forming, Fitch, 1935; Bowker, 1987; anuran body inflation, Ferreira et al., 2019). Even tiny RPM and RPB prey items can vary in ways that matter-Black Mambas (Dendroaspis polylepis, TL > 2 m, mass  $\sim$ 1.5 kg) eat lipid-rich termites (~2 mg each, RPM ~0.001) but not toxic ants (Dial and Vaughan, 1987; Branch, 1991; Branch et al., 1995; but see Evans and Alexander, 2021). Likewise, weasels (Mustela) might be more formidable prey than rodents for Old World ratsnakes (Elaphe; Prötzel et al., 2018), Bullsnakes (Pituophis catenifer sayi; Mulaik, 1938), and adders (Vipera; López Jurado and Caballero, 1981; Bringsøe, 2019), but data on RPM and handling times will be required to explore costs and benefits of eating those carnivores. We conclude that the term large prey is always ambiguous and should be replaced with the words heavy, bulky, or both, which in common parlance signify just what they mean here; Arnold's (1993:103-111) discussion of



FIG. 7. Frequency distribution of relative prey mass (RPM) for different prey shapes from 43 California Kingsnakes (*Lampropeltis getula californiae*; modified from Wiseman et al., 2019). Black bars = snake prey (type II), white bars = nonsnake prey (types III and IV); banded gray bar = digested captive snake prey and solid gray bars = regurgitated captive snake prey, from Jackson et al. (2004). Inset (A): Eastern Kingsnake (*L. g. getula*; mass, ~300 g) attempting to ingest a Eastern Mole (*Scalopus aquaticus*; adult mass, ~50 g), RPM of ~0.17, Beaufort, Carteret County, North Carolina, 1327 h, 21 July 2017 (data and photo: F. S. Boyce). Inset (B): *L. g. californiae* ingesting Mojave Rattlesnake (*Crotalus scutulatus*; RPM, ~0.80–1.0), Pima, Graham County, Arizona, 17 August 2006 (data and photo: R. White).

"varieties of useless prey" remains pertinent, as is Kornilev et al.'s (2022) review of snakes failing to survive ingesting harmful prey.

Besides foraging theory and other conceptual realms, MBT might be applicable to additional gape-limited predators. Possible examples include frogfishes (Antennariidae; Pietsch and Arnold, 2020:451), lizardfishes (Synodontidae; Soares et al., 2003), venomous deep-sea eels (Monognathidae; Bertelsen and Nielsen, 1987), morays (Muraenidae; Diluzio et al., 2017; Higgins et al., 2018; Mehta et al., 2020), some frogs (e.g., *Ceratophrys;* Duellman and Lizana, 1994), certain varanids and helodermatids (Greene, 1986a; Repp and Schuett, 2009), and some birds (e.g., Roadrunner [*Geococcyx californianus*]; Holte and Houck, 2000).

*Relative Prey Mass.*—Greene's (1983a) WR and IR suffered from the use of ratios (Atchley et al., 1976) and imprecisions of "weight" and "ingestion." RPM instead specifies equivalent aspects of predators and prey, which are measurable with simple tools (e.g., field-portable balances) in the same units (e.g., grams) and amenable to diverse comparisons (e.g., analyses of covariance on log-transformed data for hypothesis testing and percentages for outreach). Moreover, RPM has long been used for snakes (e.g., Fitch and Twining, 1946; Brown, 1958; Rodríguez-Robles and Greene, 1999; King, 2002; Andreadis and Burghardt, 2005; Vincent et al., 2006a) and is more directly related to costs and benefits than linear dimensions or volume (e.g., Henderson, 1993; Greene et al., 1994; Machio et al., 2010; Enge et al., 2022). Predator mass scales variably with length among species (e.g., Jayne et al., 2022) but also differs within species, even within an individual seasonally, depending on physiological condition (e.g., Fitch, 1949; Dobson, 1992; Cundall, 2000; Rivas, 2020:92). Finally, RPM measurements are subject to other errors and biases, particularly with preserved specimens and proxy estimates of live weights (for a careful example, Boback et al., 2016; for subsampling stomachs with hundreds of tiny prey items, Araújo et al., 2008).

Multiple similar items in a stomach could represent single meals in terms of search costs. Clumped prey eaten in rapid succession at one site might include schooling fishes (B. Greene et al., 1994), insect larvae (Webb et al., 2000; Fig. 5), reptile eggs (e.g., Rodríguez-Robles and Greene, 1999; Barends and Maritz, 2022a; Durso et al., 2022), roosting bats (Sorrell et al., 2011), nestling birds and mammals (e.g., Rodríguez-Robles et al., 1999b; Quick et al., 2005; Barends and Maritz, 2022b), and suckling mammals ingested with their mothers (e.g., Lanchi et al., 2012). As exemplars of payoffs from prey taken in one foraging bout, for a 50-g California Mountain Kingsnake (Lampropeltis zonata) that ate five 1-g nestling mice, RPM was 0.02/item and 0.1 combined; a 5-g rodent with the latter RPM would have entailed greater RPB and perhaps overall higher handling costs. For a 45-g L. zonata that ate 10 1-g squamate eggs, RPM was 0.022 per item and 0.222 combined; a single egg of equivalent value would require wider gape but not higher costs to subdue (data from Greene and Rodríguez-Robles, 2003). Note, however, that intact and well-digested young rabbits in the stomach of a Trans-Pecos Ratsnake (*Bogertophis subocularis*) could have been taken from separate nests (Moon and Rabatsky, 2004); likewise, a Reticulated Python (*Malayopython reticulatus*) simultaneously located and killed two children (Headland and Greene, 2011), but repeated predation by *P. sebae* on adult humans (as implied by Johnston, 1908) likely occurred over months or years. In each of these examples, the costs and benefits of ingestion might best be assessed for individual prey.

Above complexities notwithstanding, measuring RPM is straightforward compared to RPB, and perhaps its greatest challenge is to account for predator mass variation over time relative to length (e.g., Fitch, 1949; for length-mass relationships in snakes, Feldman and Meiri, 2013; Rivas, 2020:92; Jayne et al., 2022). A core importance of RPM is that higher values imply higher handling costs (heavier adversaries struggle harder) and higher payoffs (more grams of prey means more nutrition). Two predictions of MBT thus are that high RPM comes with the benefit of less frequent foraging—yielding fewer risks and time for other activities—but entails the cost of subduing heavier prey by brute force, constriction, and/or venom.

Relative Prey Bulk.-Beyond error and bias, RPB is conceptually and empirically more problematic than RPM; Fabre et al. (2016:635), for example, wrote of "large and bulky prey ..." as "heavy and/or ... relatively wide or tall for their length ...." Challenges arise because RPB might reflect a prey's cross-sectional dimensions (e.g., mouse versus shad; Fig. 2c), cross-sectional density (e.g., mouse versus bird; Fig. 6b), deformability (e.g., mouse versus turtle; Fig. 6a), or a combination of those variables, as well as structural components of predator gape (e.g., cranial bones, soft tissues; Fig. 3)-attributes difficult to measure for both prey and predators in ways that are functionally relevant, variable across taxa and methods, and controversial (e.g., King, 2002; Martins et al., 2002; Close and Cundall, 2012; Hampton and Moon, 2013; Hampton, 2018; Cundall, 2019; Moon et al. 2019; Gripshover and Javne, 2021; Cundall and Irish, 2022; Jayne et al., 2022). Nonetheless, qualitative comparisons and experimental studies indicate significant relationships between RPB and structural correlates of gape (e.g., Cundall and Greene, 2000:324; Close and Cundall, 2012; Gripshover and Jayne, 2021; Cundall and Irish, 2022; Jayne et al., 2022).

Having struggled with these intricacies when studying snake diets (e.g., Rodríguez-Robles et al. 1999a; Wiseman et al., 2019), we anticipate their clarification by other researchers (see below) and simply refer here to RPB because bulk is defined as "a lot of size or heft, though not necessarily heavy ... Pillows are bulky ... big in an inconvenient way" (www.vocabulary. com, accessed June 1, 2021). Bulk describes a key aspect of snake feeding and has been used in this sense (e.g., Marques et al., 2010; Passos et al., 2019:9; Barends and Maritz, 2022a; Solórzano and Sasa, 2022) and yet provides an umbrella for more precise terms and elaborations (e.g., Close and Cundall, 2012; Cundall, 2019; Moon et al., 2019; Gripshover and Jayne, 2021; Jayne et al., 2022); moreover, this overarching descriptor is useful in realms as different as functional morphology and public outreach (a child alerted us to the pillow example). High RPB implies high handling costs (more time and energy for ingestion and concomitant risks from other predators),

separate from but interacting with those imposed by high RPM (King, 2002; see especially Close and Cundall, 2012; Jayne et al., 2022; Kornilev et al., 2022). Another core prediction of MBT is that snakes feeding on high RPB prey are specialized for enhanced gape regardless of RPM, whereas feeding frequencies depend on eating type III (high RPM, less often) versus IV prey (low RPM, more often).

RPM, RPB, and Prey Shapes.-Prey shapes are defined by linear dimensions and geometry, which are often taxon specific; at high RPM and/or RPB, they have consequences for costs and benefits of feeding (e.g., Greene, 1983a; Voris and Voris, 1983; Gripshover and Jayne, 2021; Cundall and Irish, 2022; Javne et al., 2022). As discussed below, all else equal and at a given gape, type II prey (Figs. 2a, 7b, 9) will entail the highest handling costs and payoffs; heavy bulky type III prey (Fig. 1) will have high costs from RPM and RPB, as well as high payoffs from RPM. At constant RPM, however, items that are fusiform, ovoid, or asymmetric in cross-section (III and IV, Figs. 1, 6), rather than uniform and elongate (II), require increased gapes-so generalists should drop these from their diet at lower RPM than type II prey (potentially testable, e.g., with a South American Watersnake [Erythrolamprus miliaris] eating fishes, frogs, and caecilians, Eisfeld et al., 2021; see next section on colubrids).

In terms of prey ID and shape, earthworms, centipedes, and some chordates are elongate (type II); limbed squamates and mammals span fusiform to ovoid, and some amniote eggs are spherical (type III). Type IV prey vary in overall dimensions, density, and/or deformability relative to mass but are defined by a need for wide predator gape at lower RPM-note that fusiform or roundish prey can differ in density and/or deformability, such that a bird or a tortoise would yield lower RPM than a rodent with equivalent cross-sectional area (Close and Cundall, 2012; Jayne et al., 2022; see below). Thus, although high RPB is obvious for shad and many other fishes because of disparate major and minor cross-sectional axes (Fig. 2c; Voris and Voris, 1983), some other prey taxa have high RPB because of rigid or dangerous structures, including turtle shells (Fig. 6a), bird beaks and long feathered limbs (Fig. 6b), mole forefeet (Fig. 7a), porcupine quills (Duarte, 2003), deer antlers (e.g., Sunquist, 1982; Rivas, 2020:91-92), and inflatable lungs of anurans (Fig. 6c; Ferreira et al., 2019).

As another example of linking RPB with RPM, relevant to human-snake relationships and thus conservation (Pooley, 2022), some herpetologists have claimed our shoulders prevent ingestion by all but the longest snakes—but people coexisting with giant constrictors usually weigh less than adult Caucasians (at 90 kg, HWG has twice the mass of an adult male Indigenous Philippine Agta) and occasionally are attacked and eaten by these snakes (Branch and Hacke, 1980; Headland and Greene, 2011; Rivas, 2020:99–103; Natusch et al., 2021). Moreover, snakes can reduce RPB for at least some mammals by alternately deforming a prey item's shoulders during ingestion, such that they are swallowed sequentially rather than simultaneously (Close and Cundall, 2012).

*MBT and Fossils.*—Greene (1983a) estimated TL (~1.8 m and ~0.5 m, respectively) and RPM (~0.42) for an Eocene boid (*Eoconstrictor fischeri*, Georgalis et al., 2021) and its crocodilian prey. Subsequent researchers described the fossilized stomach contents of other arguably crown group snakes (e.g., *Pachyrhachis problematicus;* Scanlon et al., 1999; Greene and Cundall, 2000), and we anticipate further integration of paleontological evidence for RPM and RPB with data from extant taxa. A stem

serpent that plausibly ate clumped nestling dinosaurs (Zaher et al., 2022), an *E. fischeri* (SVL ~90 cm) containing a freshly ingested lizard (SVL ~8 cm, torso diameter ~17 mm; Smith and Scanferla, 2016), and a giant Pliocene adder (*Bitis* cf. *olduvaiensis*, TL ~1.45 m) that ate an immature hare (Rage and Bailon, 2011:473–476) exemplify possibilities for applying MBT to ancient prey-predator interactions in snakes and their closest extinct relatives.

# Mass-Bulk Theory and Nonvenomous Colubrid Snakes

Refining MBT could entail holding RPM, RPB, prey shape, or ID constant to test predictions of how other variables respond across a diverse range of snakes and prey. Ideally, this approach includes evaluating individual, ontogenetic, sexual, seasonal, and geographic variation before addressing specific questions (e.g., Greene, 1984; Bea et al., 1992; Luiselli, 2006b; Wiseman et al., 2019) in a phylogenetic framework (e.g., Greene, 1983a; Vincent et al., 2006a; Barends and Maritz, 2022b). For those reasons, and because it has a broad diet and is well represented in museum collections, California Kingsnakes (Lampropeltis getula californiae) provided special potential for testing MBT. We began a study of these serpents while KDW was in HWG's Berkeley herpetology course (Wiseman et al., 2019), of which the results are integrated here with research on certain other colubrids-collectively 1,840 prev items from 1,108 snakes (Table 1; for relationships, Zaher et al., 2019). We partly used data from museum specimen stomach contents (e.g., 55% of 447 L. g. californiae records) and attempted to address redundancy, bias, and sources of variation (e.g., Rodríguez-Robles and Greene, 1999:490). Our smallest samples were for Scarlet Kingsnakes (Lampropeltis elapsoides) and L. zonata, attractive snakes that collectors might not kill immediately, such that stomach contents were not preserved. Lampropeltis elapsoides, P. catenifer, and Long-nosed Snakes (Rhinocheilus lecontei) eat mostly skinks (Plestiodon), mammals, or whip-tailed lizards (Aspidoscelis), respectively; Glossy Snakes (Arizona elegans) and L. g. californiae have broader diets, encompassing squamates, birds, and rodents. For heuristic purposes, we subjectively characterize gapes as narrow or wide, with hopes that differences (or lack thereof) eventually will be quantified.

Several results from comparisons among these colubrids are consistent with predictions from MBT:

- Lampropeltis g. californiae, with a narrower gape than A. elegans and P. catenifer, drops high RPB prey types III and IV from the diet at much smaller RPM (~0.2) than for type II; L. g. californiae eats high RPM meals only in the form of snakes, and stout *Crotalus* provide the highest value (Fig. 7b). RPM thus helps explain how eating rattlesnakes, only 7% of prey by frequency, might select for immunity against viper venom.
- (2) Lampropeltis g. californiae, A. elegans, and P. catenifer have a maximum RPM ~0.7–1.0, but the broader gaped A. elegans and P. catenifer achieve higher values with bulky type III instead of elongate type II prey (Rodríguez-Robles, 2002:173; Wiseman et al., 2019:20). At distributional extremes, RPM was 0.01–0.73 for L. g. californiae and 0.02–0.86 for Eastern Kingsnakes (L. g. getula; Godley et al., 2017); captives regurgitated prey with RPM 1.17 and 1.35 but digested one with RPM 1.06 (Jackson et al., 2004).

These observations imply a maximum RPM ("upper breaking point" of Arnold, 1993) of ~1.0 (Fig. 7) for that species, which is rarely achieved in nature and only with type II prey. Likewise, two *P. catenifer* died during or shortly after ingesting type III rodents with RPMs of 0.82 and 1.36 (Rodríguez-Robles, 2002). Whether such success-failure bracketing can work for other species depends upon an adequate sample of field-based RPM data and the logistics of providing especially heavy prey to captive animals: Mole Snakes (*Pseudaspis cana*) likely would require a huge enclosure to seize an antelope (B. Maritz et al., 2020), for example, as might Gaboon Adders (*Bitis gabonica*) to ambush an ungulate or primate (Foerster, 2008; Warner and Alexander, 2011).

- (3) Arizona elegans and L. zonata that consume type IV birds are longer than other snakes that eat type III mammals, and the latter are longer, on average, than snakes taking less bulky types II and III lizards. Total length is correlated with gape within species (Jayne et al., 2022), and a similar relationship between snake TL and lizard, bird, or mammal prey also characterizes some other colubrids (e.g., Milksnakes [Lampropeltis triangulum sensu lato] Rodríguez and Drummond, 2000; Barten, 2010; Greene et al., 2010).
- (4) *Arizona elegans* has a wider gape than *R. lecontei* and consumes mammals at a smaller TL; among limbed squamate prey, the former mainly consumes stout-bodied type III phrynosomatids and the latter elongate type II whiptails.
- (5) Lampropeltis elapsoides and R. lecontei are slender, sharpsnouted diggers, with narrow gapes and diets that emphasize type II lizards in their diets. Longer L. elapsoides rarely add higher RPM and RPB items (among 34 individuals with prey, the longest individual's TL was 50 cm; a 44-cm TL snake ate the only rodent), whereas southerly R. lecontei with TLs of 38–97 cm occasionally eat type III mammals.
- (6) Head-first ingestion is typical for most snakes, perhaps because legs, scales, and other protuberances more easily fold that way (for taxa in which tail-first prevails see, e.g., Greene, 1976; Cobb, 2004). For a given gape, tail-first should be easier as RPB decreases at lower RPM (e.g., Greene, 1976; Pleguezuelos et al., 1994; Mehta, 2003). Among 187 L. g. californiae prey, 10 swallowed tail-first were "relatively small or attenuate" (e.g., nestling rodents; Wiseman et al. 2019:8). Of 25 L. zonata prey, "three neonate mammals, probably relatively small items, were eaten tail first" (Greene and Rodríguez-Robles, 2003:309). Thirtyseven of 321 P. catenifer prey were swallowed tail-first or bent double, with "a trend for smaller animals (i.e., nestlings) to be swallowed tail-first with a higher frequency than juvenile[s] or adult[s]" (Rodríguez-Robles, 2002:168). Lampropeltis elapsoides and R. lecontei have narrow gapes and always eat prey head-first.

(7) HWG and collaborators scarcely addressed RPB because of uncertainties regarding what to measure (e.g., Rodríguez-Robles et al., 1999a; Rodríguez-Robles, 2002; Wiseman et al., 2019). MBT nonetheless predicts that small individuals exclude high RPB items from their diets and longer snakes eat those same prey taxa when low RPB correlates with low RPM. The following two kinds of type IV prey demonstrate opportunities for future studies of these tradeoffs. (a) Among 447 items for *L. g. californiae*, ingestion of the only horned lizard (*Phrynosoma*) was fatal to predator and prey (Wiseman et al., 2019:14). Conversely, less than a fourth as many diet items for the wider-gaped *A. elegans*  included two fatal and two successful consumptions of Phrynosoma (Rodríguez-Robles et al., 1999a). Among other colubrids, Coachwhips (Masticophis flagellum) eats items as heavy and bulky as rabbits (Whiting et al., 1992), and yet RPM for *Phrynosoma* was low ( $\bar{\mathbf{x}} = 0.04$ ) and averaged half that of type II whiptails ( $\overline{x} = 0.08$ ; Appendix 8). Consistent with MBT, however, a Desert Night Snake (Hypsiglena chlorophaea) ate an essentially hornless Pigmy Short-horned Lizard (Phrynosoma douglasii) with an RPM ~0.5 (O'Connor et al., 2010). (b) Moles (Talpidae) have semirigid, outwardturned forelimbs with stout claws (Lin et al., 2019), to which snakes have responded as follows: a L. g. getula failed to ingest one with an RPM less than the predicted maximum for types III and IV prey (Fig. 7a); shorter Copperheads (Agkistrodon contortrix), with wider viperid gapes, ate adult moles (Uhler et al., 1939; Graves, 2002); and a Rubber Boa (Charina bottae) consumed three nestlings with an RPM of  $\sim$ 0.1, such that their combined RPM was  $\sim$ 0.3 (Rodríguez-Robles et al., 1999b).

# Mass-Bulk Theory and the Evolution of Venomous Snakes

Preliminary assessments are consistent with MBT's prediction that snakes with high RPM will be adapted for subduing high-cost prey. Scolecophidians are nonconstrictors, are nonvenomous, and generally take tiny type I items (e.g., Shine and Webb, 1990; Webb and Shine, 1993a,b; Webb et al., 2000; Fig. 5). Constricting basal alethinophidians-aniliids, uropeltids, boids, and pythonids-often eat types II or III prey with RPM > 0.5, and individuals of the latter two taxa occasionally eat type III prey with RPM > 1.0 (Fig. 1c; Appendix 4). An acrochordid contained an "enormous" fish with RPM of 0.3 (Shine, 1986:427). Nonconstricting, non-front-fanged colubroids typically take types I–IV with RPM < 0.5 (Fig. 2; for an exception, see Linares and Eterovick, 2012), and constricting colubrids rarely exceed RPM of 1.0 (Fig. 7, Table 1, Appendix 3). Only elapids and viperids with some frequency have an RPM of  $\sim$ 1.0–1.7 (Figs. 8–10; Appendix 5). These patterns exist despite biases that might obscure them and have not been evaluated for the energetic effect of rarely eaten but unusually heavy or otherwise nutritious prey (e.g., Greene, 1986a; Wiseman et al., 2019).

Cundall and Greene (2000) further suggested that frontfanged snakes with tranquilizing toxins (e.g., most elapids) often consume type II prey (Fig. 9), whereas those that tranquilize and tenderize (e.g., many viperids; Figs. 8b, 10) emphasize type III items (toxin terminology from Mackessy, 2010); they inferred this reflects lower surface area relative to mass for heavy bulky meals, such that tenderizers facilitate digestion, especially in cold climates (e.g., Greene, 1992; Lutterschmidt et al., 1996). Here, we show how MBT can elucidate the roles of diet in venomous snake evolution and emphasize that although prey ID matters (e.g., Daltry et al., 1996; Gibbs and Rossiter, 2008; Barlow et al., 2009; Modahl et al., 2018; Davies and Arbuckle, 2019; Zancolli et al., 2019; Lyons et al., 2020; Holding et al., 2021), RPM and RPB are central to this topic (see also, e.g., Hayes et al., 2002; Bringsøe, 2019; Hamanaka and Mori, 2020).

Assessing RPM While Controlling for Other Variables.—If venoms tranquilize and tenderize especially dangerous and heavy prey, venomous snakes should take higher RPM items than nonvenomous species. Broad comparisons, however, as summarized above, risk confounding venom effects with the availability of equivalent RPM prey (Tsai et al., 2016) and vulnerability of particular prey taxa (e.g., Arnold, 1993). With respect to availability, RPM for sympatric aquatic nonvenomous Banded Watersnakes (*Nerodia fasciata*; 0.01–0.39,  $\bar{x} = 0.11$ ) and venomous Cottonmouths (Agkistrodon piscivorus; 0.19–0.53,  $\bar{x} =$ 0.16) indeed differed as predicted by MBT (data from Camper, 2022). To control for vulnerability, we compared pairs of sympatric nonvenomous and venomous snakes and found that when colubrids (Masticophis) and rattlesnakes (Crotalus) eat the same lizard species, mean RPMs are four to five times higher for the latter (Fig. 8, Appendix 9). Future applications of this approach could encompass nonconstrictors, constrictors, nonfront-fanged, and front-fanged snakes (for categories, see Sullivan and Weinstein, 2017), comparing RPM and RPB for multiple prey types in different habitats and at local to global scales.

Elongate Nonvertebrates as Diversely Different Prey.—Centipedes, despite their conveniently attenuate shape, are never eaten by most snakes, presumably because of sharp-legged struggling abilities and venomous forcipules. Exceptions include several Old and New World viperids (e.g., Clark, 1967; Bea and Braña, 1988; Revault, 1996; Holycross et al., 2002; Hamanaka and Mori, 2020), black-headed and crowned snakes (Tantilla), and certain other rear-fanged New World colubrids (e.g., Solórzano et al. 2012; Rorabaugh et al., 2020; Enge et al., 2022) and one clade of African rear-fanged lamprophilds (centipede-eaters [Aparallactus]; Maritz et al., 2021a). If RPM and RPB were available for diverse centipede-eaters—e.g., generalists versus specialists, front-fanged versus not-one might test hypotheses about convergent evolution and adaptive significance of venom delivery systems (e.g., Hofmann et al., 2021). For example, among vipers, Terciopelos (Bothrops asper) have consumed centipedes with RPMs of 0.07 and 0.65 (Greene, 1992; Boada et al., 2005), whereas Aparallactus and Tantilla evidently cannot match that latter value (RPM for a Rim Rock Crowned Snake [Tantilla oolitica] that died eating a centipede was  $\sim 0.3$ , assuming equal densities of predator and prey; Enge et al., 2022); Plains Black-headed Snakes (Tantilla nigriceps), however, subdue centipedes faster than Rock Rattlesnakes (C. Lepidus; Rodríguez-Robles, 1994; Greene, 1997:81) and Mamushi Pitvipers (Gloydius blomhoffii; Hamanaka and Mori, 2020).

Defensive abilities of centipedes are obvious to humans who handle them, whereas earthworms are slimy but seemingly harmless. Among relatively basal snakes, although most scolecophidians feed only on small social insects, one species of Australasian Blindsnake (Acutotyphlops subocularis) eats annelids (Shine and Webb, 1990), as do uropeltids other than Cylindrophis (Rajendran, 1985). Species in several nonfanged colubrid lineages, usually with TL < 0.3 m, consume earthworms (e.g., Atractus [e.g., Dixon et al., 1976; Cunha and Nascimento, 1978; Martins and Oliveira, 1998; Camper and Zart, 2014; Passos et al., 2019]; wormsnakes [Carphophis; Barbour, 1960; Clark, 1970; Quinn and Carmody, 2021]; coffeesnakes [Ninia; Greene, 1975], and some other goo-eaters [Dipsas, Sibon; Ray et al., 2012]). Moreover, eating annelids is correlated with secondary fang loss in Aparallactus modestus (Portillo et al., 2019) and the homalopsid Brachyorrhos (Murphy et al., 2012); among front-fanged snakes, only one bizarre viper (Atheris barbouri; Rasmussen and Howell, 1998) and a few Australasian elapids (e.g., Ogmodon vitianus; Zug and Ineich, 1993) eat them. Cundall and Greene (2000:323-324) stated that worm-eaters



FIG. 8. Relative prey mass (RPM) differs between venomous crotalines and nonvenomous, nonconstricting colubrines feeding on the same prey types in western North America. (A) Sidewinder Rattlesnakes (*Crotalus cerastes*) and Coachwhips (*Masticophis flagellum*) that ate Western Whip-tailed Lizards (*Aspidoscelis tigris*). (B) Northern Pacific Rattlesnakes (*Crotalus oreganus*) and California Striped Whipsnakes (*Masticophis lateralis*) that ate sceloporines (*Sceloporus occidentalis* and *Uta stansburiana*). Animal images (top) by R. W. Hansen; log-transformed comparisons (middle) and frequency distributions (bottom) provide "moderate evidence" that RPM is "positively associated" with venom use by pitvipers (for methods, data, and clarifications, see Appendix 9).

are "nonconstrictors and nonvenomous, whereas those taking elongate vertebrates constrict (e.g., *Cylindrophis* and *Lampropeltis getula*) or are venomous (e.g., various fossorial elapids), suggesting... differences between annelids...and vertebrates... in mass-specific struggling abilities." Earthworms, however, might not always be easy to handle, as "Loss of the diastema [gap between fangs and other teeth] in *Toxicocalamus* could thus be interpreted as...for feeding on soft-bodied invertebrates that must be teased into the gullet because of the lack of any vertebral column or exoskeleton to resist longitudinal compression" (McDowell, 1969:507). Certain New Guinea elapids (*Toxicocalamus*) that eat annelids (Shine and Keogh, 1996) have long puzzled herpetologists, either because venom is presumed unnecessary to immobilize such prey (McDowell, 1969:465, 467; Calvete et al., 2012:4095; O'Shea et al., 2015:256, 2018:404), or because those snakes, despite powerful toxins, are inoffensive when handled (Strickland et al., 2016:665 doubted their "small gapes and fangs... [can envenom] humans"; Kraus, 2017:574). The venom of *Toxicocalamus* nonetheless might be used defensively, given that the bright coloration of *Toxicocalamus ernstmayri* could be aposematic and Indigenous people believe its bite is deadly



FIG. 9. (A) Venomous New Guinea worm-eating elapid, *Toxicocalamus loriae*, collected 23 December 1969, Kundiawa, Waghi Valley, Papua New Guinea (MCZ R-111785); total length (TL) of 178 mm, with recently ingested earthworm of roughly equal TL and relative prey mass of ~0.75—far heavier than quantified for any nonfront-fanged worm-eating snakes; (B) snake's head with protruding earthworm posterior (see text for details; photos: M. O'Shea).

(O'Shea et al., 2018, 2020); moreover, other small elapids do kill people (e.g., Asian coralsnakes [*Sinomicrurus*; Kramer, 1977] and kraits [*Bungarus*; Moffett, 2002]).

We obtained data consistent with MBT's prediction that individual Toxicocalamus ingest earthworms with higher RPM than nonvenomous annelid-eaters. The holotype of T. ernstmayri (O'Shea et al., 2015), with an SVL of 1,100 mm and mass of 280 g, contained an earthworm with a TL of 436 mm ( $\sim$ 40% snake SVL), mass of 85 g, and RPM of  $\sim$ 0.3. Assuming proportionality with those data, a Toxicocalamus loriae, with an SVL of 162 mm, contains an earthworm of a TL of  $\sim$ 160 mm and RPM of ~0.8 (Fig. 9; O'Shea et al., 2015); likewise, a Toxicocalamus goodenoughensis (Roberts and Austin, 2020), with an SVL of 271 mm, regurgitated an earthworm with a TL of  $\sim$ 200 mm and RPM of  $\sim$ 0.6. Those three snakes thus had an RPM of ~0.3–0.8 ( $\overline{x}$  ~ 0.5), compared to an RPM of 0.03-0.2 for worms eaten by three species of nonvenomous colubrids (Seib 1985a), a mean RPM of 0.07 for those eaten by Long-Tailed Alpine Gartersnakes (Thamnophis scalaris; Venegas-Barrera and Manjarrez, 2001), and a mean RPM of 0.3 for three eaten by Atractus snethlageae (Martins and Oliveira, 1998; Camper and Zart, 2014). Passos et al. (2019), however, illustrated two Atractus with perhaps high RPM annelid prey, suggesting that, like some other colubrids that eat softbodied invertebrates, they might have tranquilizing toxins (e.g., Carl, 1978; Salmão and Laporta-Ferreira, 1994; Zaher et al., 2014).

### MASS-BULK THEORY AND BIRDS AS SNAKE PREY

Birds epitomize high RPB at low RPM because of their beaks, long forelimbs, and feathers (e.g., Fitch and Twining, 1946; King, 1975; Mata-Silva et al., 2011; Camera et al., 2014; Javne et al., 2022; Fig. 6b)—perhaps this is why so few snakes specialize on them, compared to hundreds of species that eat mainly amphibians, other reptiles, or mammals (e.g., Greene, 1997; Barends and Maritz, 2022a,b). Moreover, as detailed above for colubrids whose diets include prey types II-IV, often only longer individuals with wider gapes take birds (see also Rodríguez and Drummond, 2000). Nonetheless, serpent taxa for which feathered reptiles are dietary mainstays include anacondas (Eunectes; Rivas, 2020; Thomas and Allain, 2021), Asian catsnakes (Boiga; Greene, 1989a), African treesnakes (Toxicodryas; Greenbaum et al., 2021), Neotropical birdsnakes (Phrynonax; Robinson et al., 2005; Visco and Sherry, 2015; Zuluaga-Isaza et al., 2015), certain island vipers (e.g., Golden Lancehead [Bothrops insularis; Marques et al., 2012]), and Round Island Boas (Casarea dussumieri; Roesch et al., 2022); some of these same species or close relatives eat bats, another type IV prey (e.g., Esbérard and Vrcibradic, 2007; Szczygiel and Page, 2020). Future research thus could address whether ambushing versus searching snakes consume adults or nestlings and if birds and bats are functionally equivalent prey in terms of MBT. Ratsnakes (Pantherophis) and related colubrids discussed above warrant attention on both counts (e.g., Fitch, 1963; Plummer, 1977; Brown, 1979; Fitch, 1999; Rodríguez and Drummond, 2000; Rodríguez-Robles, 2002; Stake et al., 2005; DeGregorio et al., 2016; Wiseman et al., 2019; Barends and Maritz, 2022a,b).

Southwestern Speckled Rattlesnakes (Crotalus pyrrhus) eat birds more frequently than most other Pitvipers, and Cochran et al. (2021) insightfully explored geographic dietary variation in that context. Cochran et al. (2021) did not consider MBT, but two C. pyrrhus from California exemplify lower payoff for a House Sparrow (Passer domesticus; RPM 0.17, eaten by 107-g MVZ 229959) than a Desert Cottontail (Sylvilagus audubonii; RPM > 0.5, regurgitated by 991-g MVZ 229801). Ingestion times at constant RPM also are likely higher for type IV than the type II and III centipedes, lizards, and mammals that Rattlesnakes typically consume (Figs. 6b, 8; Fitch and Twining, 1946; Mata-Silva et al., 2011). Accordingly, are C. pyrrhus that emphasize birds in their diets behaviorally and/or morphologically specialized for high RPB and thus still obtain high payoff per meal, or do they compensate for lighter prey by more frequent feeding, slower growth, or lower fecundity?

Terrestrial, arboreal, and aquatic boids might also prove enlightening because although individuals of several species consume diverse types II and III prey with high RPM (Fig. 1, Appendix 4), some of them also eat birds. As predicted by MBT, island Boa Constrictors (*Boa constrictor* sensu lato) that consume passerines have lower RPM ( $\bar{x} \sim$ 0.07) than mainland individuals feeding with equal frequency on lizards, birds, and mammals ( $\bar{x} \sim 0.44$ ; Boback, 2005). Among the longer mainland snakes—who also eat iguanas and mammals as diverse as carnivores and primates (Greene, 1983b)—the occasional Turkey Vulture (*Cathartes auratus*) is thus likely a low RPM-high RPB item (Boback, 2004; Platt et al., 2021).

# Is there a Future for Mass-Bulk Theory in Snake Biology?

To summarize, 1) MBT seeks to explain how RPM, RPB, prey shape, prey ID, and feeding frequency interact to influence the evolution, morphology, ecology, and behavior of snakes (see also Camper and Dixon, 2000; King, 2002; Vincent et al., 2006a,b; Close and Cundall, 2012; Loughran et al., 2013; Glaudas et al., 2019; Gripshover and Javne, 2021; Barends and Maritz, 2022a,b; Cundall and Irish, 2022; Jayne et al., 2022; Kornilev et al., 2022). 2) Snakes encompass individual and phylogenetic differences in RPM, reflecting extensive taxonomic and shape diversity in their prey. An Australian Scrub Python (Simalia kinghorni) that ate a pademelon (Thylogale) with an RPM of 1.67 (Glaudas et al., 2019; S. Fearn, pers. com.) and a Sidewinder Rattlesnake (Crotalus cerastes) that contained a Western Whip-Tailed Lizard (Aspidoscelis tigris) with an RPM of 1.72 (Mulcahy et al., 2003) hold records for that parameter. Dendroaspis polylepis might exhibit the greatest RPM range, from  $\sim 0.001$  for termites to  $\sim 1.0$  or higher for duikers, galagos, and other mammals (Jackson, 1956; Branch et al., 1995; Phelps, 2002; Bourquin, 2021; Evans and Alexander, 2021). 3) RPB also shows great variation within and among species (Figs. 2, 8, 9; e.g., Voris and Voris, 1983; King, 2002; Martins et al., 2002; Close and Cundall, 2012; Hampton and Moon, 2013; Gripshover and Jayne, 2021; Jayne et al., 2022), although discerning patterns therein is daunting because of problems discussed above. 4) As for conservation, combining natural history with fanciful human parallels can enhance empathy for snakes among lay people; a 10.5-g Northern Pacific Rattlesnake's (Crotalus oreganus; MVZ 229849) likely first meal, an 11.2-g Western Fence Lizard (Sceloporus occidentalis; RPM 1.07), was roughly equivalent to HWG ingesting a 95-kg hotdog without using hands or cutlery.

What is Needed?—Scientific, logistical, and cultural factors are hampering snake research in ways that could not have been predicted decades ago. Greene (1986b, 2005a) identified a lack of publishing and archiving outlets as among impediments to natural history, but these problems now are minimized by journals devoted to individual observations (Teodoro et al., 2022), high-profile venues promoting descriptive studies (e.g., Maritz et al., 2021b; Enge et al., 2022), and public platforms for aggregating huge data sets (e.g., Grundler, 2020; Maritz and Maritz 2020; Putman et al., 2021). Moreover, theoretical considerations of biodiversity "knowledge short-falls" (Hortal et al., 2015), "next-gen natural history" (Tosa et al., 2021), and globalizing studies of snake diets (Maritz et al., 2021b) all portend a welcome increase in knowledge. The challenge for expanding MBT will be to gather more rich content and widely applicable data-but what would doing that look like, whence could they come, and what obstacles await?

Complete accounts of snakes feeding would include where and when; direction of ingestion and other behavioral contexts; ID, sex, linear measurements, and mass for predators and prey; and validating information, e.g., observer's name and contact, voucher photographs, and/or museum catalog numbers (Maritz et al., 2021b). If those data were available for taxonomically diverse samples of many snakes and meals, collected with multiple methods, we could better assess biases and measurement errors (e.g., Rodríguez-Robles, 1998; Glaudas et al., 2017a; Maritz and Maritz, 2020; Durso et al., 2022); with those data, we could examine individual, ontogenetic, sexual, seasonal, and geographic variation prior to posing other questions (e.g., Pleguezuelos et al., 1994; Luiselli, 2006b; Glaudas et al., 2019; Wiseman et al., 2019; Grundler and Rabosky, 2021; Durso et al., 2022). Likewise, we need RPM standardized for predator TL to transcend variation in reproductive, nutritional, and hydration status (Cundall, 2000; Rivas, 2020:92). We also hope that functional morphologists will fine-tune measuring RPB beyond lab conditions (e.g., Jayne et al., 2022), making their insights applicable to field observations and preserved specimens (see especially Close and Cundall, 2012:1046-1048). These are all technical matters, so gaining additional RPM and RPB data are, in principle, possible, although special considerations might sometimes prohibit some procedures (e.g., forced regurgitations; Reinert et al., 2008). Future projects could thus use massive, detailed datasets to explore MBT in terms of global patterns of snake evolution and ecology (e.g., Luiseilli, 2006a; Glaudas et al., 2019; Grundler and Rabosky, 2021; Cundall and Irish, 2022; Kornilev et al., 2022).

Remaining Obstacles .-- Gathering MBT data from live and preserved snakes might prove ever more difficult, as regulatory overburdens for field biology threaten to prevent all methods except photography (e.g., Greene and Losos, 1988; Alexander et al., 2021). Exemplifying this trend, one herpetologist, after decades of permit and protocol approvals, quit teaching with live reptiles and collects occasional specimens with a hunting license; a young researcher concluded that beyond agency and institutional compliance, consequences of mistakenly breaking laws are so severe he no longer saves roadkill for museums. Now add in that those touting new methods often minimize their shortcomings (e.g., fecal DNA requires facilities and funds and yet yields limited data; Brown et al., 2014), focus on prey taxonomy (Hoefer et al., 2021; Durso et al., 2022), or emphasize problems with museum specimens ("the traditional method to gather snake diet data," Glaudas et al., 2019:758; but see, e.g., Fitch, 1960; Arnold, 1993; Luiselli and Akani, 2003). The negative effect of these trends is shown by a curator who denied our request to examine common species of Lampropeltis because "new imaging technologies can explore stomach contents without damaging valuable specimens, new generations of students rarely contribute museum specimens, and many recently common species are now rare or extinct and irreplaceable." Of course, we decry the last two realities, having prepped thousands of specimens and focused our careers on conservation. More importantly, museum specimens offer unique prospects for studying geographic variation in snake diets compared to other data sources (e.g., Sparks et al., 2015; Wiseman et al., 2019), so adopting that curator's attitude would lead to less learned about snakes and museums failing to meet their potential for studying biodiversity.

We also are not optimistic about community science contributing to MBT, despite its many positive aspects (Maritz and Maritz, 2020; Durso et al., 2021; Putman et al., 2021; see Cooper et al., 2021, for "community" versus "citizen" science). Recall that in our "Natural History Notes" survey described above, all 33 records lacking RPM (85% of 39 total) were from field observations. Obtaining additional data would have necessitated touching snakes, which is usually illegal without a permit as well as problematic because of animal welfare and, with venomous species, includes safety considerations (e.g., Ribble and Rathbun, 2018). Three records for which prey were available still would have required an instrument to provide RPM, so we wondered



FIG. 10. (A) Terciopelo (*Bothrops asper*) shortly after ingesting moderately heavy, bulky prey (perhaps a Tropical Cottontail [*Sylvilagus gabbi*] seen earlier at the site); 7 December 2021, Estación Biológica La Selva, Heredia Province, Costa Rica (photo: W. Lopez). (B) Rock was 18 cm wide, implying pitviper's total length was ~1.26 m (data, interpretation, and photo: O. Vargas Ramírez).

whether lay naturalists might carry portable scales—costing and weighing less than cheap binoculars—but community science innovators told us that asking untrained, unlicensed people to touch live or dead animals would be poorly advised. Perhaps instead the most that can be promoted for community scientists to bolster MBT is putting scale bars in photos, such that linear dimensions and mass can be estimated by comparison with organisms of similar size, visible animal structures (e.g., a hindfoot), or objects (e.g., a rock, Fig. 10; see also Barten, 2010; Marques et al., 2010; Close and Cundall, 2012; Feldman and Meiri, 2013; McMartin, 2013; Schalk and Cove, 2018:2; Quinn and Carmody, 2021). Then again, if wild hummingbirds can weigh themselves (Carpenter et al., 1983), perhaps someday snakes will too.

We hope to have convinced readers that for many serpents, eating prey that are heavy, bulky, or both is at the core of their existence. If obstacles to data acquisition are not solved, however, Godley's (1980) complaint about data quality will still apply 40 more years hence—biologists might well have 100,000 diet records, encompassing 75% of the world's snake species and accessible with a few keystrokes (Grundler and Rabosky, 2021; Maritz et al., 2021b), but they mostly will document when, where, and what taxa were eaten. Much of that dietary information will be relevant to only a subset of potential applications, and MBT, however central to snake biology, will remain based mostly on data available now.

# Coda

Watching and writing about animals has blessed me (HWG), over the course of roughly seven decades—including during preparation of this paper—with a resilient sense of purpose as well as boundless pleasure and satisfaction. My childhood love of reptiles began in Texas at age 7, thanks to "dry-land terrapins" (Eastern Box Turtles [*Terrapene carolina*]) and "horned frogs" (Texas Horned Lizards [*Phrynosoma cornutum*]) on grandpa's piney woods dirt farm. Within a few years, I met a Western Diamond-backed Rattlesnake (Crotalus atrox) at a camp for military brats in the Hill Country and was impressed that our soldier-counselors did not kill the rattlesnake. Since that first venomous serpent, there have been countless others in more than a dozen countries, along with many good times and some so bad they still haunt me. As a civilian first responder during college years, I helped many people survive violence, sudden illness, and emergency childbirth. By the age of 27, I had pulled a headless teenager out of a wreck, failed to save a toddler in anaphylactic shock while her mother sat screaming next to me, and lost a favorite professor and a lover to murders. Luckier breaks during my youth included as an army medic being sent to Germany instead of Vietnam, and, at a time when few academics thought snakes worthy of study, having William Pyburn and Gordon Burghardt as graduate advisors.

After earning a Ph.D., my good fortune has included for 20 years teaching herpetology and vertebrate natural history at the University of California, Berkeley, while serving as curator of herpetology in the MVZ. A 1999 move to Cornell University brought new challenges, as I lectured on evolution and ecology to thousands of mostly business majors and then fine-tuned "walking and talking the Tree of Life" for biology undergraduates (Ballen and Greene, 2017). Along the way, I penned two books that bridged science and art, with an emphasis on serpents of course (Greene, 1997, 2013). More than a decade ago, I veered into anthropology and shifted research emphasis to snake-primate interactions (e.g., Headland and Greene, 2011; Gardner et al., 2015; Greene, 2017, 2018, 2020; Kazandjian et al., 2021).

Some of my most rewarding activities as a field biologist have occurred since retirement. In 2019, I realized a longstanding dream of observing big elapids by helping former Cornellians Bryan and Robin Maritz, along with South African biologist Graham Alexander, during their research on Cape Cobras (Naja nivea) in the Kalahari Desert. Spying on those magnificent yellow snakes as they foraged on Puff Adders (Bitis arietans) and Sociable Weavers (Philetairus socius) did not disappoint (Fig. 11a-c; Maritz and Maritz, 2019). Meanwhile, Emily Taylor, an English major in my Berkeley classes, had become a distinguished professor at Cal Poly State University, San Luis Obispo, and elected president of the American Society of Ichthyologists and Herpetologists. Two decades after Emily first visited the Mojave Desert with my herpetology course, I joined her class's trip there, overflowing with pride for the phenomenal enthusiasm she inspires in students (Fig. 11d). Mentors, mentees, and professional colleagues are not obligated to be friends, so I feel blessed to count these people, along with coauthor Kevin Wiseman, as among my dearest.

As 2019 ended, I began restoring a chunk of Hill Country, named Rancho Cascabel for its resident *C. atrox*. Among the many joys of rural existence is enhanced familiarity with a place and its biota, across seasons and years, as well as surprises. In 2020, for example, I encountered a pair of Texas Patch-nosed Snakes (*Salvadora lineata*) mating near my Longhorns' water trough (Fig. 12)—and thereby confirmed in nature the male of this species' head-biting behavior, which was previously documented only for captives (Burchfield et al., 1982). Strolling on down life's road, I hope to observe many more serpents, including some consuming meals that are heavy, bulky, or both.



FIG. 11. Friends, field trips, and retirement. (A) HWG observing adult Cape Cobra (*Naja nivea*; left arrow) and adult female Puff Adder (*Bitis arietans*; right arrow) at Tswalu Kalahari Reserve, South Africa; 26 February 2019 (photo: R. A. and B. Maritz). (B) Same *N. nivea* inspects *B. arietans*, which it repeatedly envenomed and eventually ate (photo: H. W. Greene). (C) Another *N. nivea* looks down from Sociable Weaver (*Philetairus socius*) nest colony (photo: H. W. Greene). (D) Cal Poly State University, San Luis Obispo herpetology class at Pisgah Lava Flow, San Bernardino County, California; 12 May 2019; arrows indicate Professor Emily Taylor (left) and HWG (photo: E. N. Taylor).

Acknowledgments.—Special thanks to Erin Muths for inviting and so patiently editing this contribution, published half a century after the Society for the Study of Amphibians and Reptiles (SSAR) boosted an insecure Ph.D. candidate's confidence with its first Outstanding Student Paper Award (for Greene, 1973b). This is the final installment in the Journal of Herpetology's senior researchers' perspectives series, and we look forward to future pieces by younger, more diverse SSAR members. We appreciate the many museum curators who facilitated our studies, which have been supported primarily by the MVZ, the Lichen Fund, a Cornell University Stephen H. Weiss Presidential Fellowship, and the National Science Foundation (BSR 83-00346, OPUS 1354156). For curatorial assistance with this paper, we thank C. Austin and J. Roberts (Louisiana State University Museum of Zoology), C. Dardia and

C. Dillman (CUMV), T. LaDuc (TNHC), E. Smith (UTA), and C. Spencer (MVZ); M. O'Shea, along with A. Baldinger, J. Hanken, and J. Rosado of MCZ, were especially helpful with the *Toxicocalamus* example. We are especially grateful for feedback on our manuscript from S. Boback, G. Burghardt, D. Cundall, L. Alencar, C. Feldman, B. Maritz, G. Pauly, K. Schwenk, and J. Sigala-Rodríguez. B. Halstead advised on statistics and J. Sigala-Rodríguez prepared the Resumen. For other assistance we thank K. Adler, G. Alexander, A. Bauer, B. Bauer, C. Bell, K. Bemis, E. Braker, H. Bringsøe, G. Burghardt, R. Dowling, A. Durso, A. Echelle, A. Echternacht, S. Fearn, J. Fitzpatrick, M. Fitzpatrick, K. Glaser, X. Glaudas, E. Greenbaum, D. Hailey, W. Hallwachs, R. Hansen, D. Hendrickson, T. Hibbitts, E. Hillman, R. Huey, D. Janzen, B. Jayne, D. Johnson, J. Jones, D. Kizirian, W. Koenig, J. Losos, T. Lott, L. Luiselli, R.



FIG. 12. Mating behavior of Texas Patch-nosed Snakes (*Salvadora lineata*) at Rancho Cascabel, Mason County, Texas; observations were made at a distance of 1–2 m from the pair and began a few minutes before 0953 h CDT, 5 May 2021 (photos: H. W. Greene). (A) Snakes remained within a  $\sim$ 1-m<sup>2</sup> patch of sparse low vegetation between corral gate post and light-colored rock, lower center in image. (B) When discovered and thereafter, the longer, thicker male had grasped the female's head in his jaws and their bodies were loosely aligned, in hairpin or irregularly semicircular coils; their cloacae initially were not juxtaposed (1010 h). (C) Male grasping female's head during copulation (0954 h). (D) Intromission involved the right hemipenis (here, at 1056 h).

Maritz, R. Mehta, D. Moore, C. Moreau, S. Mullin, J. Murphy, D. Natusch, P. Passos, T. Pietsch, H. Reinert, R. Repp, J. Rivas, B. Rothermal, A. Savitzky, J. Schauer, C. Sheehy III, T. Sinclair, K. Smith, S. Spawls, W. Starnes, B. Stein, E. Taylor, O. Vargas Ramírez, R. Voss, K. Warkentin, M. Westneat, Paul Weldon, W. Wüster, and K. Zamudio. The Dwight W. and Blanche Faye Reeder Centennial Fellowship in Systematic and Evolutionary Biology paid page and open access charges for this paper.

#### LITERATURE CITED

- ABALOS, J. W., E. C. BAEZ, AND R. NADER. 1964. Serpientes de Santiago del Estero. Acta Zoologica Lilloana 20:211–283.
- ALEXANDER, G. J., K. A. TOLLEY, B. MARITZ, A. MCKECHNIE, P. MANGER, R. L. THOMPSON, C. SCHRADIN, A. FULLER, L. MEYER, R. S. HETEM, ET AL. 2021. Excessive red tape is strangling biodiversity research in South Africa. South African Journal of Science 117:10787.
- ANDREADIS, P., AND G. M. BURGHARDT. 2005. Unlearned appetite controls: watersnakes (*Nerodia*) take smaller meals when they have a choice. Journal of Comparative Psychology 119:304–310.

- ARAÚJO, M. S., A. PINHEIRO, AND S. F. REIS. 2008. Gluttonous predators: how to estimate prey size when there are too many prey. Brazilian Journal of Biology 68:315–320.
- ARNOLD, S. J. 1983. Morphology, performance, and fitness. American Zoologist 23:347–361.
- ARNOLD, S. J. 1993. Foraging theory and prey-size—predator-size relations in snakes. Pp. 87–115 in R. A. Seigel and J. T. Collins (eds.), Snakes: Ecology and Behavior. McGraw Hill, USA.
- ATCHLEY, W. R., C. T. GASKINS, AND D. ANDERSON. 1976. Statistical properties of ratios. I. Empirical results. Systematic Zoology 25:137–148.
- BALLEN, C. J., AND H. W. GREENE. 2017. Walking and talking the tree of life: why and how to teach about biodiversity. PLoS Biology 5: e2001630.
- BANCI, K. R. S., N. F. TORELLO-VIERA, A. C. FREITAS, AND O. A. V. MARQUES. 2017. Feeding on elongate prey: additional data for the coral snake *Micrurus corallinus* (Merrem, 1820) (Elapidae) and comments on aposematism. Herpetology Notes 10:335–338.
- BARBO, F. E., AND O. A. V. MARQUES. 2003. Do aglyphous colubrid snakes prey on live amphisbaenids able to bite? Phyllomedusa 2:113–114.
- BARBOUR, R. W. 1960. A study of the worm snake, *Carphophis amoenus* Say, in Kentucky. Transactions of the Kentucky Academy of Science 21:10–16.
- BARENDS, J. M., AND B. MARITZ. 2022a. Snake predators of bird eggs: a review and bibliography. Journal of Field Ornithology 93:1.
- BARENDS, J. M., AND B. MARITZ. 2022b. Dietary specialization and habitat shifts in a clade of Afro-Asian colubrid snakes (Colubridae: Colubrinae). Ichthyology and Herpetology 110:278–291.
- BARLOW, A., C. E. POOK, R. A. HARRISON, AND W. WÜSTER. 2009. Coevolution of diet and prey-specific venom activity supports the role of selection in snake venom evolution. Proceedings of the Royal Society B: Biological Sciences 276:2443–2449.
- BARROS, M. M., J. F. DRAQUE, P. A. MICUCCI, AND T. WALLER. 2011. Eunectes notaeus (yellow anaconda). Diet/cannibalism. Herpetological Review 42:290–291.
- BARTEN, S. L. 2010. Red milk snake taking large prey late in season. Reptiles and Amphibians: Natural History and Conservation 17:94.
- BARTOSZEK, I. A., P. T. ANDREADIS, C. PROKOPERVIN, M. PATEL, AND R. N. REED. 2018. *Python bivittatus* (Burmese python). Diet and prey. Herpetological Review 49:139–140.
- BEA, A., AND F. BRANA. 1988. Nota sobre la alimentación de Vipera latastei, Boscá, 1878 (Reptilia, Viperidae). Munibe Ciencias Naturales 40:121–124.
- BEA, A., F. BRAÑA, J. P. BARON, AND H. SAINT-GIRONS. 1992. Régimes et cycles alimentaires des vipères Européennes (Reptilia, Viperidae): étude comparée. Année Biologique 31:25–44.
- BERG, P., J. BERG, AND R. BERG. 2020. Predator-prey interaction between a boomslang, Dispholidus typus, and a flap-necked chameleon, Chamaeleo dilepis. African Journal of Ecology 58:855–859.
- BERTELSEN, E., AND J. G. NIELSEN. 1987. The deep sea eel family Monognathidae (Pesces, Anguilliformes). Steenstrupia 13:141–198.
- BHUPATHY, S., AND V. S. VIJAYAN. 1989. Status, distribution and general ecology of the Indian python, *Python molurus molurus* Linnaeus in Keoladeo National Park, Bharatpur, Rajasthan. Journal of the Bombay Natural History Society 86:381–387.
- BOADA, C., D. SALAZAR-V, A. F. LASCANO, AND U. KUCH. 2005. The diet of Bothrops asper (Garman) in the Pacific lowlands of Ecuador. Herpetozoa 18:77–79.
- BOBACK, S. M. 2004. Boa constrictor (Boa Constrictor). Diet. Herpetological Review 35:175.
- BOBACK, S. M. 2005. Natural history and conservation of island boas (*Boa constrictor*) in Belize. Copeia 2005:879–884.
- BOBACK, S. M., E. BURROUGHS, C. UGARTE, AND J. WATLING. 2000. *Boa constrictor* (Boa Constrictor). Diet. Herpetological Review 31:244.
- BOBACK, S. M., R. W. SNOW, T. HSU, S. C. PEURACH, C. J. DOVE, AND R. N. REED. 2016. Supersize me: remains of three white-tailed deer (*Odo-coileus virginianus*) in an invasive Burmese python (*Python molurus bivittaus*) in Florida. Bioinvasion Records 5:197–203.
- BOLTT, R. E., AND R. F. EWER. 1964. The functional anatomy of the head of the puff adder, *Bitis arietans* (Merr.). Journal of Morphology 114:83–106.
- BOURQUIN, O. 2021. *Dendroaspis polylepis* Günther, 1864. Black mamba. Diet. African Herpetology News 77:33–34.
- BOWKER, R. W. 1987. *Élgaria kingi* (Arizona Alligator Lizard). Antipredator behavior. Herpetological Review 18:73–74.
- BRANCH, W. R. 1991. Unusual herpetological observations in the Kruger National Park. African Herp News 16:39–40.

- BRANCH, W. R., AND M. BURGER. 1991. Lamprophis guttatus. Spotted house snake. Diet. Journal of the Herpetological Association of Africa 39:24.
- BRANCH, W. R., AND W. HACKE [SIC, HAACKE]. 1980. A fatal attack on a young boy by an African rock python *Python sebae*. Journal of Herpetology 14:305–307.
- BRANCH, W. R., G. V. HAAGNER, AND R. SHINE. 1995. Is there an ontogenetic shift in mamba diet? Taxonomic confusion and dietary records for black and green mambas (*Dendroaspis*: Elapidae). Herpetological Natural History 3:171–178.
- BRANCH, W. R., A. M. BAUER, AND T. LAMB. 2002. Bitis caudalis (Horned Adder). Prey size. Herpetological Review 33:137–138.
- BRECKO, J., B. VERVUST, A. HERREL, AND R. VAN DAMM. 2011. Head morphology and diet in the dice snake (*Natrix tessellata*). Mertensiella 18:20–29.
- BRINGSØE, H. 2019. Observations of adder, Vipera berus (Squamata: Viperidae) preying on least weasel, Mustela nivalis (Carnivora: Mustelidae): an overlooked feeding habit. Herpetology Notes 12:401– 403.
- BRINGSØE, H., M. SUTHANTHANGJAI, W. SUTHANTHANGJAI, AND K. NIMNAUM. 2020. Eviscerated alive: novel and macabre feeding strategy in Oligodon fasciolatus (Günther, 1864) eating organs of Duttaphrynus melanostictus (Schneider, 1799) in Thailand. Herpetozoa 33:167–173.
- BROWN, E. E. 1958. Feeding habits of the northern water snake, *Natrix* sipedon sipedon Linnaeus. Zoologica 43:55–71.
- BROWN, E. E. 1979. Some snake food records from the Carolinas. Brimleyana 1:113–124.
- BROWN, D. S., K. L. EBENEZER, AND W. O. C. SYMONDSON. 2014. Molecular analysis of the diets of snakes: changes in prey exploitation during the development of the rare smooth snake *Coronella austriaca*. Molecular Ecology 23:3734–3743.
- BURCHFIELD, P. M., T. F. BEIMLER, AND C. S. DOUCETTE. 1982. An unusual precoital head-biting behavior in the Texas patch-nosed snake, *Sal*vadora grahamiae lineata (Reptilia: Serpentes: Colubridae). Copeia 1982:192–193.
- CABRAL, H., L. PIATTI, M. MARTINS, AND V. FERREIRA. 2020. Natural history of *Xenodon matogrossensis* (Scrocchi and Cruz 1993) (Serpentes, Dipsadidae) in the Brazilian Pantanal. Cuadernos de Herpetología 34:211–218.
- CALVETE, J. J., P. GHEZELLOU, O. PAIVA, T. MATAINAHO, A. GHASSEMPOUR, H. GOUDARZI, F. KRAUS, L. SANZ, AND D. J. WILLIAMS. 2012. Snake venomics of two poorly known Hydrophiinae: comparative proteomics of the venoms of terrestrial *Toxicocalamus longissimus* and marine *Hydrophis cyanocinctus*. Journal of Proteomics 75:4091–4101.
- CAMERA, B. F., D. J. DA SILVA, M. DOS SANTOS FILHO, V. A. CAMPOS, AND G. R. CANALE. 2014. Bothrops moojeni (Brazilian Lancehead). Diet. Herpetological Review 45:705.
- CAMPBELL, E. F., AND J. B. HEWLETT. 2021. Agkistrodon piscivorus (Cottonmouth). Diet. Herpetological Review 52:416–417.
- CAMPER, J. D. 2022. Comparative ecology of two species of semiaquatic snakes in southeastern North America. Pp. 77–93 in M. M. Shah, U. Sharif, T. R. Buhari, and T. S. Imam (eds.), Snake Venom and Ecology. IntechOpen, UK.
- CAMPER, J. D., AND J. R. DIXON. 2000. Food habits of three species of striped whipsnakes, *Masticophis* (Serpentes: Colubridae). Texas Journal of Science 52:83–92.
- CAMPER, J. D., AND D. J. ZART. 2014. *Atractus snethlageae* (Ground Snake). Diet. Herpetological Review 45:705.
- CAPULA, M., L. LUISELLI, L. RUGIERO, F. EVANGELISTI, C. ANIBALDI, AND V. T. JESUS. 1997. Notes on the food habits of *Coluber hippocrepis nigrescens* from Pantellaria Island: a snake that feeds on both carrion and living prey. Herpetological Journal 7:67–70.
- CARBAJAL-MÁRQUEZ, R. A., G. ARNAUD, M. MARTINS, AND G. E. QUINTERO-DÍAZ. 2016. Diet of *Crotalus enyo* (Serpentes: Viperidae) from the Baja California Cape region. Acta Zoológica Mexicana 32:45–48.
- CARBAJAL-MÁRQUEZ, R. A., J. R. CEDEÑO-VÁZQUEZ, D. GONZÁLEZ-SOLÍS, AND M. MARTINS. 2020. Diet and feeding ecology of *Crotalus tzabcan* (Serpentes: Viperidae). South American Journal of Herpetology 15:9–19.
- CARBAJAL-MÁRQUEZ, R. A., J. J. SIGALA-RODRÍGUEZ, J. A. ESCOTO-MORENO, J. M. JONES, AND C. MONTAÑO-RULVALCABA. 2022. New prey items of *Crotalus campbelli* (Serpentes: Viperidae) from Mexico. Phyllomedusa 21:95–98.
- CARL, G. 1978. Notes on worm-eating in the prairie ringneck snake, *Dia-dophis punctatus arnyi*. Bulletin of the Maryland Herpetological Society 14:95–97.

- CARPENTER, F. L., D. C. PATON, AND M. A. HIXON. 1983. Weight gain and adjustment of feeding territory size in migrant hummingbirds. Proceedings of the National Academy of Sciences USA 80:7259–7263.
- CARREIRA VIDAL, S. 2002. Alimentación de los ofidios de Uruguay. Asociación Herpetológica Española, Monografías de Herpetologia 6:1– 126.
- CASPER, G. C., J. B. LECLERE, AND J. C. GILLINGHAM. 2015. Thamnophis sirtalis (Common Gartersnake). Diet/scavenging. Herpetological Review 46:653–654.
- CLARK, D. R. 1970. Ecological study of the worm snake Carphophis vermis (Kennicott). University of Kansas Publications Museum of Natural History 19:85–194.
- CLARK, R. J. 1967. Centipede in the stomach of young *Vipera ammodytes meridionalis*. Copeia 1967:224.
- CLARK, R. W. 2002. Diet of the timber rattlesnake, Crotalus horridus. Journal of Herpetology 36:494–499.
- CLAYTON, S., AND G. MYERS. 2015. Conservation Psychology: Understanding and Promoting Human Care for Nature. 2nd ed. Wiley, USA.
- CLEUREN, S. G. C., D. P. HOCKING, AND A. R. EVANS. 2021. Fang evolution in venomous snakes: adaptation of 3D tooth shape to the biomechanical properties of their prey. Evolution 75:1377–1394.
- CLOSE, M., AND D. CUNDALL. 2012. Mammals as prey: estimating ingestible size. Journal of Morphology 273:1042–1049.
- COBB, V. A. 2004. Diet and prey size of the flathead snake, *Tantilla gracilis*. Copeia 2004:397–402.
- COCHRAN, C., K. L. EDWARDS, Z. D. TRAVIS, L. R. POMPE, AND W. K. HAYES. 2021. Diet and feeding frequency in the southwestern speckled rattlesnake (*Crotalus pyrrhus*): ontogenetic, sexual, geographic, and seasonal variation. Journal of Herpetology 55:77–87.
- COLBERT, J. E., K. M. ANDREWS, AND T. M. NORTON. 2014. Agkistrodon piscivorus (cottonmouth). Diet and prey size. Herpetological Review 45:703–704.
- COLLETTE, B. B. 1977. Summary of the meetings. Copeia 1977:804-823.
- CONRADIE, W., AND P. V. PINTO. 2021. A snake with an appetite for the rare: *Amblyodipsas polylepis* (Bocage, 1873) feeding on the amphisbaenid *Monopeltis luandae* Gans, 1976. Herpetology Notes 14:205– 207.
- COOPER, C. B., C. L. HAWN, L. R. LARSON, J. K. PARRISH, G. BOWSER, D. CAV-ALIER, R. R. DUNN, M. HAKLAY, K. K. GUPTA, N. O. JELKS, ET AL. 2021. Inclusion in citizen science: the conundrum of rebranding. Science 372:1386–1388.
- CORREA-SANCHEZ, F., M. A. CASARIEGO-MADORELL, AND F. LUNA-CASTELLA-NOS. 2001. Porthidium dunni (Dunn's hognosed pitviper). Diet. Herpetological Review 32:264.
- CUNDALL, D. 2000. Drinking in snakes: kinematic cycling and water transport. Journal of Experimental Biology 203:2171–2185.
- CUNDALL, D. 2019. A few puzzles in the evolution of feeding mechanisms in snakes. Herpetologica 75:99–107.
- CUNDALL, D., AND H. W. GREENE. 2000. Feeding in snakes. Pp. 293–333 in K. Schwenk (ed.), Feeding: Form, Function, and Evolution in Tetrapod Vertebrates. Academic Press, USA.
- CUNDALL, D., AND F. IRISH. 2022. Macrostomy, macrophagy, and snake phylogeny. Pp. 438–454 in D. J. Gower and H. Zaher (eds.), The Origin and Early Evolutionary History of Snakes. Cambridge University Press, UK.
- CUNDALL, D., C. TUTTMAN, AND M. CLOSE. 2014. A model of the anterior esophagus in snakes, with functional and developmental implications. Anatomical Record 297:586–598.
- CUNHA, O. R., AND F. P. NASCIMENTO. 1978. Ofidios da Amazonia X. As cobras da região leste de Pará. Museu Paraense Emelio Goedi, Publicações Avulsas 31:1–218.
- DALTRY, J. C., W. WÜSTER, AND R. S. THORPE. 1996. Diet and snake venom evolution. Nature 379:537–540.
- DARTEZ, S. F., P. M. HAMPTON, N. E. HAERTLE, AND C. S. MONTEIRO. 2011. Lampropeltis getula holbrooki (speckled kingsnake). Diet. Herpetological Review 42:292.
- DA SILVA, A. S., E. F. DA SILVA, J. M. DA S. AMARAL, V. N. BARBOSA, AND F. G. R. FRANÇA. 2021. *Bothrops leucurus* (white-tailed lancehead). Diet. Herpetological Review 52:150.
- DA SILVA, C. F., E. P. ACANTARA, H. F. OLIVEIRA, M. A. S. OLIVEIRA, AND R. W. AVILA. 2015. Oxybelis aeneus (brown vinesnake). Diet. Herpetological Review 46:648.
- DA SILVA, F. O., A.-C. FABRE, Y. SAVRIAMA, J. OLLONEN, K. MAHLOW, A. HERREL, J. MÜLLER, AND N. DI-POI. 2018. The ecological origins of snakes as revealed by skull evolution. Nature Communications 9:376.

- DAVIES, E. L., AND K. ARBUCKLE. 2019. Coevolution of snake venom toxic activities and diet: evidence that ecological generalism favours toxicological diversity. Toxins 11:711.
- DEGREGORIO, B. A., P. J. WEATHERHEAD, AND J. H. SPERRY. 2016. Ecology and behavior of corn snakes (*Pantherophis guttatus*) on avian nests. Herpetological Conservation and Biology 11:150–159.
- DIAL, K. P., AND T. A. VAUGHAN. 1987. Opportunistic predation on alate termites in Kenya. Biotropica 19:185–187.
- DILUZIO, A. R., V. B. BALIGA, B. A. HIGGINS, AND R. S. MEHTA. 2017. Effects of prey characteristics on the feeding behaviors of an apex marine predator, the California moray (*Gymnothorax mordax*). Zoology 122:80–89.
- DIXON, J. R., R. A. THOMAS, AND H. W. GREENE. 1976. Status of the neotropical snake *Rhabdophis poeppigi* Jan, with notes on variation in *Atractus elaps* (Günther). Herpetologica 32:221–227.
- DOBSON, S. 1992. Body mass, structural size, and life history patterns of the Columbian ground squirrel. American Naturalist 140:109–125.
- DUARTE, M. R. 2003. Prickly food: snakes preying upon porcupines. Phyllomedusa 2:109–112.
- DUARTE, M. R. 2012. Elapomorphus quinquelineatus (Raddi's lizard-eating snake). Diet. Herpetological Review 43:146.
- DUELLMAN, W. E., AND M. LIZANA. 1994. Biology of a sit-and-wait predator: the leptodactylid frog *Ceratophrys cornuta*. Herpetologica 50:51– 64.
- DUGAN, E. A., AND W. K. HAYES. 2012. Diet and feeding ecology of the red diamond rattlesnake, *Crotalus ruber* (Serpentes: Viperidae). Herpetologica 68:203–217.
- DURSO, A. M., AND N. KIRIASZIS. 2011. Coluber constrictor (North American racer). Prey size. Herpetological Review 42:285.
- DURSO, A. M., AND S. J. MULLIN. 2017. Ontogenetic shifts in the diet of the plains hog-nosed snakes (*Heterodon nasicus*) revealed by stable isotope analysis. Zoology 120:83–91.
- DURSO, A. M., R. RUIZ DE CASTAÑEDA, C. MONTALCINI, M. R. MONDARDINI, J. L. FERNANDEZ-MARQUES, F. GREY, M. M. MÜLLER, P. UETZ, B. M. MAR-SHALL, R. J. GRAY, ET AL. 2021. Citizen science and online data: opportunities and challenges for snake ecology and action against snakebite. Toxicon X 9–10:100071.
- DURSO, A. M., T. J. KIERAN, T. C. GLENN, AND S. J. MULLIN. 2022. Comparison of three methods for measuring dietary composition of plains hog-nosed snakes. Herpetologica 78:119–132.
- DWYER, C. M., AND H. KAISER. 1997. Relationship between skull form and prey selection in the thamnophine snake genera *Nerodia* and *Regina*. Journal of Herpetology 31:463–475.
- EISFELD, A., L. PIZZATTO, AND D. VRCIBRADIC. 2021. Diet of the semiaquatic snake *Erythrolamprus miliaris* (Dipsadidae, Xenodontinae) in the Brazilian Atlantic Forest. Journal of Herpetology 55:330–337.
- ENGE, K. M., J. A. GRAY, C. M. SHEEHY III, T. FERRARO, D. M. MARTIN, AND J. D. MAYS. 2022. What killed the rarest snake in America? Ecology 104:e1857.
- ESBÉRARD, C. E. L., AND D. VRCIBRADIC. 2007. Snakes preying on bats: new records from Brazil and a review of recorded cases in the neotropical region. Revista Brasileira de Zoologia 24:848–853.
- ESCALANTE, R. N., AND D. G. ACUÑA. 2020. Predation of a plantation glass frog, Hyalinobatrachium colymbiphylum (Anura: Centrolenidae), ornate cat-eyed snakes, Leptodeira ornate (Squamata: Dipsadidae), in Costa Rica. Reptiles and Amphibians 27:489–490.
- EVANS, N., AND G. J. ALEXANDER. 2021. A natural test for the "endotherm diet hypothesis." African Herp News 78:49–51.
- FABRE, A.-C., D. E. BICKFORD, M. SEGALL, AND A. HERREL. 2016. The impact of diet, habitat use, and behavior on head shape evolution in homalopsid snakes. Biological Journal of the Linnean Society 118:634– 647.
- FARAONE, P. F., S. RUSSOTTO, G. GIACALONE, M. L. VALVO, I. BELARDI, AND E. MORI. 2021. Food habits of the javelin sand boa *Eryx jaculus* (Linnaeus 1758; Serpentes, Erycidae) in Sicily, Italy. Journal of Herpetology 55:452–458.
- FEARN, S. 2002. Morelia amethistina (Scrub Python). Diet. Herpetological Review 33:58–59.
- FELDMAN, A., AND S. MEIRI. 2013. Length-mass allometry in snakes. Biological Journal of the Linnean Society 108:161–172.
- FELDMAN, C. R., AND J. A. WILKINSON. 2000. Thamnophis sirtalis fitchi (Valley Garter Snake). Diet. Herpetological Review 31:248.
- FELDMAN, C. R., E. D. BRODIE JR., E. D. BRODIE III, AND M. W. PFRENDER. 2012. Constraint shapes convergence in tetrodotoxin-resistant sodium channels of snakes. Proceedings of the National Academy of Sciences USA 109:4556–4561.

- FELDMAN, C. R., R. W. HANSEN, AND R. SIKOLA. 2020. Thamnophis elegans terrestris (coast gartersnake). Tetrodotoxin poisoning. Herpetological Review 51:630-631.
- FERREIRA, R. E., R. LOURENÇO-DE-MORAES, C. ZOCCA, C. DUCA, K. H. BEARD, AND E. D. BRODIE JR. 2019. Antipredator mechanisms of post-metamorphic anurans: a global database and classification system. Behavioral Ecology and Sociobiology 73:69.
- FITCH, H. S. 1935. Natural history of the alligator lizards. Transactions of the Academy of Sciences of Saint Louis 29:1–38.
- FITCH, H. S. 1941. The feeding habits of California garter snakes. California Fish and Game 27:2–32.
- FITCH, H. S. 1949. Study of snake populations in central California. American Midland Naturalist 41:513–579.
- FITCH, H. S. 1960. Autecology of the copperhead. University of Kansas Publications, Museum of Natural History 13:85–288.
- FITCH, H. S. 1963. Natural history of the black rat snake (*Elaphe o. obsoleta*) in Kansas. Copeia 1963:649–658.
- FITCH, H. S. 1999. A Kansas Snake Community: Composition and Changes over 50 Years. Krieger Publishing Company, USA.
- FITCH, H. S., AND H. W. GREENE. 1965. Breeding cycle in the ground skink, *Lygosoma laterale*. University of Kansas Publications, Museum of Natural History 15:565–575.
- FITCH, H. S., AND H. TWINING. 1946. Feeding habits of the Pacific rattlesnake. Copeia 1946:64–71.
- FOERSTER, S. 2008. Two incidents of venomous snakebite on juvenile blue and Sykes monkeys (*Cercopithecus mitis stuhlmanni* and *C. m. albogularis*). Primates 49:300–303.
- FORSMAN, A., AND L. E. LINDELL. 1993. The advantages of a big head: swallowing performance in adders, *Vipera berus*. Functional Ecology 7:183–189.
- FREDRIKSSON, G. M. 2005. Predation on sun bears by reticulated python in East Kalimantan Indonesian Borneo. Raffles Bulletin of Zoology 53:165–168.
- GAIARSA, M. P., L. R. V. ALENCAR, AND M. MARTINS. 2013. Natural history of pseudoboine snakes. Papéis Avulsos de Zoologia 53:261–283.
- Gans, C. 1961. The feeding mechanism of snakes and its possible evolution. American Zoologist 1:217–227.
- GARDNER, C. J., P. RADOLALAINA, M. RAJERISON, AND H. W. GREENE. 2015. Cooperative rescue and predator fatality involving a group-living strepsirrhine, Coquerel's sifaka (*Propithecus coquereli*) and a Madagascan ground boa (*Acrantophis madagascariensis*). Primates 56:127– 129.
- GARDNER, S. A., AND J. R. MENDELSON. 2003. Diet of the leaf-nosed snakes, *Phyllorhynchus* (Squamata: Colubridae): squamate egg specialists. Southwestern Naturalist 48:550–556.
- GATICA-COLIMA, A., AND N. CÓRDOBA-REZA. 2012. Salvadora hexalepis deserticola (Big Bend patch-nosed Snake). Diet. Herpetological Review 43:350–351.
- GAVIRA, R. S. B., AND D. LOEBMANN. 2011. Bothrops sp. (GR. atrox) (Jararaca/Lancehead). Diet. Herpetological Review 42:436.
- GEORGALIS, G. L., M. RABI, AND K. SMITH. 2021. Taxonomic revision of the snakes of the genera *Palaeopython* and *Paleryx* (Serpentes, Constrictores) from the Paleogene of Europe. Swiss Journal of Palaeontology 140:18.
- GIBBS, H. L., AND W. ROSSITER. 2008. Rapid evolution by positive selection and gene gain and loss: PLA 2 venom genes in closely related *Sistrurus* rattlesnakes with divergent diets. Journal of Molecular Evolution 66:151–166.
- GLAUDAS, X., T. JEZKOVA, AND J. A. RODRÍGUEZ-ROBLES. 2008. Feeding ecology of the Great Basin rattlesnake (*Crotalus lutosus*, Viperidae). Canadian Journal of Zoology 86:723–734.
- GLAUDAS, X., T. C. KEARNEY, AND G. J. ALEXANDER. 2017a. Museum specimens bias measures of snake diet: a case study using the ambushforaging puff adder (*Bitis arietans*). Herpetologica 73:121–128.
- GLAUDAS, X., T. C. KEARNEY, AND G. J. ALEXANDER. 2017b. To hold or not to hold? The effects of prey size and type on the predatory strategy of a venomous snake. Journal of Zoology 302:211–218.
- GLAUDAS, X., K. L. GLENNON, M. MARTINS, L. LUISELLI, S. FEARN, D. F. TREMBATH, D. JELÍC, AND G. J. ALEXANDER. 2019. Foraging mode, relative prey size and diet breadth: a phylogenetically explicit analysis of snake feeding ecology. Journal of Animal Ecology 88:757–767.
- GODLEY, J. S. 1980. Foraging ecology of the striped swamp snake, *Regina alleni*, in southern Florida. Ecological Monographs 50:411–436.
- GODLEY, J. S., B. J. HALSTEAD, AND R. W. MCDIARMID. 2017. Ecology of the eastern kingsnake (*Lampropeltis getula*) at Rainey Slough, Florida: a vanished Eden. Herpetological Monographs 31:47–68.

- GOMEZ-MESTRE, I., AND K. M. WARKENTIN. 2007. To hatch and hatch not: similar selective trade-offs but different responses to egg predators in two closely related, syntopic treefrogs. Oecologia 153:197–206.
- GRAVES, G. R. 2002. Copperhead preys on star-nosed mole in the Great Dismal Swamp. Banisteria 20:70.
- GREENBAUM, E., K. E. ALLEN, E. R. VAUGHAN, O. S. G. PAUWELS, V. WALLACH, C. KUSAMBA, W. M. MUNINGA, M. M. ARISTOTE, F. M. M. MALL, G. BADJEDJEA, J. PENNER, M.-O. RÖDEL, J. RIVERA, V. STERKHOVA, G. JOHNSON, W. P. TAPONDJOU, AND R. M. BROWN. 2021. Night stalkers from above: a monograph of *Toxicodryas* tree snakes (Squamata: Colubridae) with descriptions of two new cryptic species from Central Africa. Zootaxa 4965:1–44.
- GREENE, B. D., J. R. DIXON, J. M. MUELLER, M. J. WHITING, AND O. W. THORNTON JR. 1994. Feeding ecology of the Concho water snake, *Nerodia harteri paucimaculata*. Journal of Herpetology 28:165–172.
- GREENE, H. W. 1969. Reproduction in a Middle American skink, *Leiolopisma cherriei* (Cope). Herpetologica 25:55–56.
- GREENE, H. W. 1973a. Defensive tail display by snakes and amphisbaenians. Journal of Herpetology 7:143–161.
- GREENE, H. W. 1973b. The Food Habits and Feeding Behavior of New World Coral Snakes. M.A. Thesis, University of Texas at Arlington, USA.
- GREENE, H. W. 1975. Ecological observations on the red coffee snake, *Ninia sebae*, in southern Veracruz, Mexico. American Midland Naturalist 93:478–484.
- GREENE, H. W. 1976. Scale overlap, a directional sign stimulus for prey ingestion by ophiophagous snakes. Zeitschrift f
  ür Tierpsychologie 41:113–120.
- GREENE, H. W. 1977. Phylogeny, Convergence, and Snake Behavior. Ph. D. diss., University of Tennessee, USA.
- GREENE, H. W. 1979. Behavioral convergence in the defensive displays of snakes. Experientia 35:747–748.
- GREENE, H. W. 1983a. Dietary correlates of the origin and radiation of snakes. American Zoologist 23:431–441.
- GREENE, H. W. 1983b. Boa constrictor (Boa, Bequer, Boa constrictor). Pp. 380–382 in D. H. Janzen (ed.), Costa Rican Natural History. University of Chicago Press, USA.
- GREENE, H. W. 1984. Feeding behavior and diet of the eastern coral snake, *Micrurus fulvius*. Special Publications of the Museum of Natural History, University of Kansas 10:147–162
- GREENE, H. W. 1986a. Diet and arboreality in the emerald monitor, Varanus prasinus, with comments on the study of adaptation. Fieldiana, Zoology (New Series) 31:1–12.
- GREENE, H. W. 1986b. Natural history and evolutionary biology. Pp. 99– 108 in M. E. Feder and G. V. Lauder (eds.), Predator-Prey Relationships: Perspectives and Approaches from the Study of Lower Vertebrates. University of Chicago Press, USA.
- GREENE, H. W. 1989a. Ecological, evolutionary, and conservation implications of feeding biology in Old World cat snakes, genus *Boiga* (Colubridae). Proceedings of the California Academy of Sciences 46:193–207.
- GREENE, H. W. 1989b. Defensive behavior and feeding biology of the Asian mock viper, *Psanmodynastes pulverulentus* (Colubridae), a specialized predator on scincid lizards. Chinese Herpetological Research 2:21–32.
- GREENE, H. W. 1992. The behavioral and ecological context for pitviper evolution. Pp. 107–117 in J. A. Campbell and E. D. Brodie, Jr. (eds.), Biology of the Pitvipers. Selva, USA.
- GREENE, H. W. 1994. Homology and behavioral repertoires. Pp. 369–391 in B. K. Hall (ed.), Homology: The Hierarchical Basis of Comparative Biology. Academic Press, USA.
- GREENE, H. W. 1997. Snakes: The Evolution of Mystery in Nature. University of California Press, USA.
- GREENE, H. W. 1999. Natural history and behavioural homology. Pp. 173–188 in G. R. Bock and G. Cardew (eds.), Homology (Novartis Foundation Symposium 222). John Wiley and Sons, UK.
- GREENE, H. W. 2003. Appreciating rattlesnakes. Wild Earth 13:28–32.
- GREENE, H. W. 2005a. Organisms in nature as a central focus for biology. Trends in Ecology and Evolution 20:23–27.
- GREENE, H. W. 2005b. Historical influences on community ecology. Proceedings of the National Academy of Sciences USA 102:8395–8396.
- GREENE, H. W. 2013. Tracks and Shadows: Field Biology as Art. University of California Press, USA.
- GREENE, H. W. 2017. Evolutionary scenarios and primate natural history. American Naturalist 190 (suppl.):S69–86.
- GREENE, H. W. 2018. Re-wilding the lifeboats. Pp. 360–369 in B. A. Minteer, J. Maeienschein, and J. P. Collins (eds.), The Ark and Beyond:

The Evolution of Zoo and Aquarium Conservation. University of Chicago Press, USA.

- GREENE, H. W. 2020. Pomegranates, peccaries, and love. Ecopsychology 12:166–172.
- GREENE, H. W., AND G. M. BURGHARDT. 1978. Behavior and phylogeny: constriction in ancient and modern snakes. Science 200:74–77.
- GREENE, H. W., AND D. CUNDALL. 2000. Limbless tetrapods and snakes with legs. Science 287:1939–1941.
- GREENE, H. W., AND F. M. JAKSIC. 1983. Food niche relationships among sympatric predators: effects of level of prey identification. Oikos 40:151–154.
- GREENE, H. W., AND F. M. JAKSIC. 1992. The feeding behavior and natural history of two Chilean snakes, *Philodryas chamissonis* and *Tachymenis chilensis* (Colubridae). Revista Chilena de Historia Natural 65:485–493.
- GREENE, H. W., AND J. B. LOSOS. 1988. Systematics, natural history, and conservation. BioScience 38:458–462.
- GREENE, H. W., AND G. V. OLIVER, JR. 1965. Notes on the natural history of the western massasauga. Herpetologica 21:225–228.
- GREENE, H. W., AND J. A. RODRÍGUEZ-ROBLES. 2003. Feeding ecology of the California mountain kingsnake, *Lampropeltis zonata* (Colubridae). Copeia 2003:308–314.
- GREENE, H. W., E. J. ZIMMERER, W. M. PALMER, AND M. F. BENARD. 2010. Diet specialization in the scarlet kingsnake, *Lampropeltis elapsoides* (Colubridae). Reptiles and Amphibians: Natural History and Conservation 17:18–22.
- GRIPSHOVER, N. D., AND B. C. JAYNE. 2021. Crayfish eating in snakes: testing how anatomy and behavior affect feeding performance. Integrative and Organismal Biology 3:obab001.
- GROEN, J., G. KAASTRA-BERGA, AND S. KAASTRA. 2020. First documented case of arboreal foraging by two male adders (*Vipera berus*) raiding the nest of a blue tit (*Cyanistes caeruleus*). Herpetology Notes 13:583–586.
- GRUNDLER, M. 2020. SquamataBase: a natural history database and R package for comparative biology of snake feeding habits. Biodiversity Data Journal 8:e49943.
- GRUNDLER, M. C., AND D. L. RABOSKY. 2021. Rapid increase in snake dietary diversity and complexity following the end-Cretaceous mass extinction. PLoS Biology 19:e3001414.
- HAAGNER, G. V. 1991. *Aspidelaps scutatus*. Shield-nosed snake. Diet and reproduction. Journal of the Herpetological Association of Africa 39:26.
- HAMANAKA, K., AND A. MORI. 2020. Toxicity of venom from mamushi, *Gloydius blomhoffii*, (Squamata: Crotalinae) to centipedes. Toxicon 188:11–15.
- HAMPTON, P. M. 2011. *Micrurus fulvius* (Harlequin Coralsnake). Diet. Herpetological Review 42:294.
- HAMPTON, P. M. 2018. Morphological indicators of gape size for redtailed pipe snakes (*Cylindrophis ruffus*). Journal of Herpetology 52:425–429.
- HAMPTON, P., AND B. R. MOON. 2013. Gape size, its morphological basis, and the validity of gape indices in western diamond-backed rattlesnakes (*Crotalus atrox*). Journal of Morphology 274:194–202.
- HARDY, D. L., SR., AND H. W. GREENE. 1999. Borderland blacktails: radiotelemetry, natural history, and living with venomous snakes. Pp. 117–121 in G. J. Gottfried, L. Eschew, C. G. Curtin, and C. B. Edminster (eds.), Toward Integrated Research, Land Management, and Ecosystem Protection in the Malpai Borderlands: Conference Summary; 6–8 January 1999; Douglas, Arizona. Proceedings RMRS-P-10, U.S. Department of Agriculture, Forest Service, Rocky Mountain Research Station, Fort Collins, Colorado, USA.
- HARTMANN, M. T., P. A. HARTMANN, S. Z. CECHIN, AND M. MARTINS. 2005. Feeding habits and habitat use in *Bothrops pubescens* (Viperidae, Crotalinae) from southern Brazil. Journal of Herpetology 39:664– 667.
- HAY, P. W., AND P. W. MARTIN. 1966. Python predation on Uganda kob. East African Wildlife Journal 4:151–152.
- HAYES, W. K., S. S. HERBERT, G. C. REHLING, AND J. F., GENNARO. 2002. Factors that influence venom expenditure in viperids and other snake species during predatory and defensive contexts. Pp. 207–234 in G. W. Schuett, M. Höggren, M. E. Douglas, and H. W. Greene(eds.), Biology of the Vipers. Eagle Mountain Publishing, USA.
- HEAD, J., K. DE QUEIROZ, AND H. GREENE. 2020. Serpentes C. Linnaeus 1758. Pp. 1131–1134 in K. de Queiroz, P. D. Cantino, and J. A. Gauthier (eds.), Phylonyms: A Companion to the PhyloCode. CRC Press, USA.

- HEADLAND, T. N., AND H. W. GREENE. 2011. Hunter-gatherers and other primates as prey, predators, and competitors of snakes. Proceedings of the National Academy of Sciences USA 108:E1470–1474.
- HENDERSON, R. W. 1993. On the diets of some arboreal boids. Herpetological Natural History 1:91–96.
- HERO, J.-M., AND W. MAGNUSSON. 1987. Leptophis ahaetulla. Food. Herpetological Review 18:16.
- HERZOG, H. A. JR., AND B. D. BAILEY. 1987. Development of antipredator responses in snakes: II. Effects of recent feeding on defensive behaviors of juvenile garter snakes (*Thamnophis sirtalis*). Journal of Comparative Psychology 101:387–389.
- HIGGINS, B. A., C. J. LAW, AND R. S. MEHTA. 2018. Eat whole and less often: ontogenetic shift reveals size specialization on kelp bass by the California moray eel, *Gymnothorax mordax*. Oecologia 188:875– 887.
- HILL, M. M. A., G. L. POWELL, AND A. P. RUSSELL. 2001. Diet of the prairie rattlesnake, *Crotalus viridis viridis*, in southeastern Alberta. Canadian Field Naturalist 115:241–246.
- HILLIS, D. M. 2020. The detection and naming of geographic variation within species. Herpetological Review 51:52–56.
- HOEFER, S., S. MILLS, T. PINOU, AND N. J. ROBINSON. 2021. What the dead tell us about the living: using roadkill to analyze the diet and endoparasite prevalence in two Bahamian snakes. Ichthyology and Herpetology 109:685–690.
- HOFMANN, E. P., R. M. RAUTSAW, A. J. MASON, J. L. STRICKLAND, AND C. L. PARKINSON. 2021. Duvernoy's gland transcriptomics of the plains black-headed snake, *Tantilla nigriceps* (Squamata, Colubridae): unearthing the venom of small rear-fanged snakes. Toxins 13:336.
- HOLDING, M. L., J. L. STRICKLAND, R. M. RAUTSAW, E. P. HOFMANN, A. J. MASON, M. P. HOGAN, G. S. NYSTROM, S. A. ELLSWORTH, T. J. COLSTON, M. BORJA, ET AL. 2021. Phylogenetically diverse diets favor more complex venoms in North American pitvipers. Proceedings of the National Academy of Sciences USA 118:e2015579118.
- HOLTE, A. E., AND M. A. HOUCK. 2000. Juvenile greater roadrunner (Cuculidae) killed by choking on a Texas horned lizard (Phyrnosomatidae). Southwestern Naturalist 45:74–76.
- HOLYCROSS, A. T., L. K. KAMEES, AND C. W. PAINTER. 2001. Observations of predation on *Crotalus willardi obscurus* in the Animas Mountain, New Mexico. Southwestern Naturalist 46:363–364.
- HOLYCROSS, A. T., C. W. PAINTER, D. B. PRIVAL, D. E. SWANN, M. J. SHROFF, T. EDWARDS, AND C. R. SCHWALBE. 2002. Diet of *Crotalus lepidus klauberi* (banded rock rattlesnake). Journal of Herpetology 36:589–597.
- HORAN, R. V., III, R. IBÁÑEZ-D, AND A. HERNANDEZ. 2011. Micrurus nigrocinctus nigrocinctus (Central American coral snake). Diet. Herpetological Review 42:294–295.
- HORTAL, J., F. DE BELLO, J. A. F. DINIZ-FILHO, T. M. LEWINSOHN, J. M. LOBO, AND R. L. LADLE. 2015. Seven shortfalls that beset large-scale knowledge of biodiversity. Annual Review of Ecology, Evolution, and Systematics 46:523–549.
- HOVEY, T. E., AND L. A. COMRACK. 2011. Crotalus oreganus helleri (Southern Pacific rattlesnake). Diet. Herpetological Review 42:288.
- HUANG, W.-S., H. W. GREENE, T.-J. CHANG, AND R. SHINE. 2011. Territorial behavior in Taiwanese kukrisnakes (*Oligodon formosanus*). Proceedings of the National Academy of Sciences USA 108:7455–7459.
- ISBELL, L. A. 2009. The Fruit, the Tree, and the Serpent: Why We See So Well. Harvard University Press, USA.
- JACKSON, K., N. J. KLEY, AND É. L. BRAINERD. 2004. How snakes eat snakes: the biomechanical challenges of ophiophagy for the California kingsnake, *Lampropeltis getula californiae* (Serpentes: Colubridae). Zoology 107:191–200.
- JACKSON, W. T. 1956. The elusive little Piti. African Wildlife 10:295–300.
- JANZEN, D. H. 1970. Altruism by coatis in the face of predation by *Boa constrictor*. Journal of Mammalogy 51:387–389.
- JAYNE, B. C., H. K. VORIS, AND P. K. L. NG. 2002. Snake circumvents constraints on prey size. Nature 418:143.
- JAYNE, B. C., H. K. VORIS, AND P. K. L. NG. 2018. How big is too big? Using crustaecean-eating snakes (Homalopsidae) to test how anatomy and behaviour affect prey size and feeding performance. Biological Journal of the Linnean Society 123:636–650.
- JAYNE, B. C., A. L. BAMBERGER, D. R. MADER, AND I. A. BARTOSZEK. 2022. Scaling relationships of maximal gape in two large species of invasive snakes, brown treesnakes and Burmese pythons, and implications for maximal prey size. Integrative and Organismal Biology 4:1–18.
- JOHNSTON, H. 1908. George Grenfell and the Congo. Hutchinson and Co., UK.

- JONES, K. B., AND W. G. WHITFORD. 1989. Feeding behavior of free-roaming *Masticophis flagellum*: an efficient ambush predator. Southwestern Naturalist 34:460–467.
- KAZANDJIAN, T., D. PETRAS, S. ROBINSON, J. VAN DER THIEL, H. W. GREENE, K. ARBUCKLE, A. BARLOW, D. CARTER, G. WHITELEY, S. C. WAGSTAFF, ET AL. 2021. Convergent evolution of pain-inducing defensive venom components in spitting cobras. Science 371:386–390.
- KING, K. A. 1975. Unusual food item of the western diamondback rattlesnake (*Crotalus atrox*). Southwestern Naturalist 20:416–417.
- KING, R. B. 2002. Predicted and observed maximum prey size—snake size allometry. Functional Ecology 16:766–772.
- KLACZKO, J., E. SHARRATT, AND E. Z. F. SETZ. 2016. Are diet preferences associated to skulls shape diversification in xenodontine snakes? PLoS ONE 11:e0148375.
- KLAUBER, L. M. 1956. Rattlesnakes: Their Habits, Life Histories, and Influence on Man. University of California Press, USA.
- KOJIMA, Y., I. FUKUYAMA, T. KURITA, M. Y. B. HOSSMAN, AND K. NISHIKAWA. 2020. Mandibular sawing in a snail-eating snake. Nature Science Reports 10:12670.
- KORNILEV, Y. V., N. D. NATCHEV, AND H. B. LILLYWHITE. 2022. Perils of ingesting harmful prey by advanced snakes. Biological Reviews 98:263–283.
- KRAMER, E. 1977. Zur Schlangenfauna Nepals. Revue Suisse Zoologie 84:721–761.
- KRAUS, F. 2017. Two new species of *Toxicocalamus* (Squamata: Elapidae) from Papua New Guinea. Journal of Herpetology 51:574–581.
- KRAUSE, M. A., G. M. BURGHARDT, AND J. C. GILLINGHAM. 2003. Body size plasticity and local variation of relative head and body size sexual dimorphism garter snakes (*Thamnophis sirtalis*). Journal of Zoology 261:399–407.
- LABONTE, J. P. 2001. *Phrynosoma coronatum* (Coast Horned Lizard). Predation and telemetry. Herpetological Review 32:257–258.
- LACEY, H., C. H. SHEWCHUK, P. T. GREGORY, M. J. SARRELL, AND L. A. GREG-ORY. 1996. The occurrence of the night snake, *Hypsiglena torquata*, in British Columbia, with comments on its body size and diet. Canadian Field Naturalist 110:620–625.
- LANCHI, F. A., R. F.LANDIM, AND M. M. MARTINELLI. 2012. *Bothrops jararacussu* (Jararacussu). Diet. Herpetological Review 43:341.
- LASPIUR, A., J. C. ACOSTA, AND G. A. FAVA. 2012. *Philodryas trilineata* (Argentine Mousehole Snake). Diet. Herpetological Review 43:151– 152.
- LAYLOO, I., C. SMITH, AND B. MARITZ. 2017. Diet and feeding in the Cape cobra, Naja nivea. African Journal of Herpetology 66:147–153.
- LIN, Y.-F., N. KONNOW, AND E. R. DUMONT. 2019. How moles destroy your lawn: the forelimb kinematics of eastern moles in loose and compact substrates. Journal of Experimental Biology 222:182436.
- LINARES, A. M., AND P. C. ETEROVICK. 2012. Erythrolamprus aesculapii (southern mock coralsnake). Diet and prey size. Herpetological Review 43:146.
- LÓPEZ JURADO, L. F., AND M. R. CABALLERO. 1981. Predación de Vipera latastei sobre Mustela nivalis. Acta Vertebrata Doñana 8:298–299.
- LOUGHRAN, C. L., E. M. NOWAK, J. SCHOFER, K. O. SULLIVAN, AND B. K. SULLIVAN. 2013. Lagomorphs as prey of western diamond-backed rattlesnakes (*Crotalus atrox*) in Arizona. Southwestern Naturalist 58:502– 505.
- LOSOS, J. B., AND H. W. GREENE. 1988. Ecological and evolutionary implications of diet in monitor lizards. Biological Journal of the Linnean Society 35:379–407.
- LUISELLI, L. 2006a. Resource partitioning and interspecific competition in snakes: the search for general geographical and guild patterns. Oikos 114:193–211.
- LUISELLI, L. 2006b. Broad geographic, taxonomic, and ecological patterns of interpopulation variation in the dietary habits of snakes. Web Ecology 6:2–16.
- LUISELLI, L., AND G. C. AKANI. 1998. Aspects of the ecology of *Calabaria reinhardtii* (Serpentes, Booidea) in southeastern Nigeria. Herpetological Natural History 6:65–71.
- LUISELLI, L., AND G. C. AKANI. 2003. Diet of sympatric gaboon vipers (*Bitis gabonica*) and nose-horned vipers (*Bitis nasicornis*) in southern Nigeria. African Journal of Herpetology 52:101–106.
- LUISELLI, L., AND F. M. ANGELICI. 1998. Sexual size dimorphism and natural history traits are correlated with intersexual dietary divergence in royal pythons (*Python regius*) from the rainforests of southeastern Nigeria. Italian Journal of Zoology 65:183–185.
- LUISELLI, L., J. M. PLEGUEZUELOS, M. CAPULA, AND C. VILLAFRANCA. 2001. Geographic variation in the diet composition of a secretive

Mediterranean colubrid snake: *Coronella girondica* from Spain and Italy. Italian Journal of Zoology 68:57–60.

- LUTTERSCHMIDT, W. I., R. L. NYDAM, AND H. W. GREENE. 1996. County record for the woodland vole, *Microtus pinetorum* (Rodentia: Cricetidae), LeFlore County, OK, with natural history notes on a predatory snake. Proceedings of the Oklahoma Academy of Sciences 76:93–94.
- LYONS, K., M. M. DUGON, AND K. HEALY. 2020. Diet breadth mediates the prey specificity of venom potency in snakes. Toxins 12:74.
- MACARTHUR, R. H., AND E. R. PIANKA. 1966. On optimal use of a patchy environment. American Naturalist 100:603–609.
- MACHIO, G. F., A. L. C. PRUDENTE, F. S. RODRIGUES, AND M. S. HOOGMOED. 2010. Food habits of *Anilius scytale* (Serpentes: Aniliidae) in the Brazilian Amazonia. Zoologia 27:184–190.
- MACKESSY, S. P. 2010. Evolutionary trends in venom composition in the western rattlesnakes (*Crotalus viridis* sensu lato): toxins vs. tenderizers. Toxicon 55:1463–1474.
- MANN, A. J., J. D. PARDO, AND H. C. MADDIN. 2022. Snake-like limb loss in a Carboniferous amniote. Nature Ecology and Evolution 6:614– 621.
- MARITZ, B. 2012. *Bitis schneideri* relative prey size. African Herp News 57:16.
- MARITZ, B., AND G. J. ALEXANDER. 2014. Namaqua dwarf adders are generalist predators. African Journal of Herpetology 63:79–86.
- MARITZ, B., G. J. ALEXANDER, AND R. A. MARITZ. 2019. The underappreciated extent of cannibalism and ophiophagy in African cobras. Ecology 100:e0522.
- MARITZ, B., M. VAN HEERDEN, AND T. SLADE. 2020. *Pseudaspis cana* (Linnaeus, 1758). Mole snake. Diet. African Herp News 74:72–74.
- MARITZ, B., A. RAWOOT, R. VAN HUYSSTEEN. 2021a. Testing assertions of dietary specialization: a case study of the diet of *Aparallactus capen*sis. African Journal of Herpetology 70:61–67.
- MARITZ, B., E. P. HOFMANN, R. A. MARITZ, H. W. GREENE, M. GRUNDLER, AND A. M. DURSO. 2021b. Points of view: challenges and opportunities in the study of snake diets. Herpetological Review 52:769–773.
- MARITZ, R., W. CONRADIE, C. I. SARDINHA, A. PETO, A. H. D. CHECHENE, AND B. MARITZ. 2020. Ophiophagy and cannibalism in African vine snakes (Colubridae: *Thelotornis*). African Journal of Ecology 58:543– 547.
- MARITZ, R. A., AND B. MARITZ. 2019. Head to head. Wild Magazine 47:16–19.
- MARITZ, R. A., AND B. MARITZ. 2020. Sharing for science: high-resolution trophic interactions revealed rapidly by social media. PeerJ 8:e9485.
- MARQUES, O. A. V., R. Z. COETI, P. A. BRAGA, AND I. SAZIMA. 2010. A rotten choice: feeding attempt by a coral snake (*Micrurus frontalis*) on a dead pitviper (*Bothrops jararaca*) that had swallowed a bulky rodent. Herpetology Notes 10:137–139.
- MARQUES, O. Ä. V., M. MARTINS, P. F. DEVELEY, A. MACARRÃO, AND I. SAZIMA. 2012. The golden lancehead *Bothrops insularis* (Serpentes: Viperidae) relies on two seasonally plentiful bird species visiting its island habitat. Journal of Natural History 46:885–895.
- MARQUES, O. A. V., AND I. SAZIMA. 2021. The natural history of New World coralsnakes. Pp. 275–289 in N. J. da Silva, Jr., L. W. Porras, S. D. Aird, and A. L. da C. Prudente (eds.), Advances in Coralsnake Biology: with an Emphasis on South America. Eagle Mountain Publishing, USA.
- MARTÍNEZ-VACA LEÓN, O. I., AND J. E. MORALES-MÁVIL. 2021. Bothriechis lateralis (Green Palm Pitviper). Diet. Herpetological Review 52:148.
- MARTINS, M., AND M. E. OLIVEIRA. 1998. Natural history of snakes in forests of the Manaus region, central Amazonia, Brazil. Herpetological Natural History 6:78–150.
- MARTINS, M., O. A. V. MARQUES, AND I. SAZIMA. 2002. Ecological and phylogenetic correlates of feeding habits in neotropical pitvipers of the genus *Bothrops*. Pp. 307–328 in G. W. Schuett, M. Höggren, M. E. Douglas, and H. W. Greene (eds.), Biology of the Vipers. Eagle Mountain Publishing, USA.
- MATA-SILVA, V., J. D. JOHNSON, R. COUVILLON, W. LUKESFAHR, AND A. ROCHA. 2011. *Crotalus atrox* (Western Diamond-backed Rattlesnake). Diet. Herpetological Review 42:438–439.
- McDowell, S. B. 1969. Toxicocalamus, a New Guinea genus of snakes of the family Elapidae. Journal of Zoology 159:443–511.
- MCMARTIN, Ć. 2013. An analysis of the "Snake Measurer" software tool. Southwest Center for Herpetological Research Bulletin 3:24–26.
- MEANS, D. B. 2017. Diamonds in the Rough: Natural History of the Eastern Diamondback Rattlesnake. Tall Timbers Press, USA.
- MEBARKI, M. T., O. GUEZOUL, K. SOUTOU, F. MARNICHE, A. BOUZID, AND S. E. SADINE. 2021. Report of camel spiders (Solfugae: Galeodidae)

predation by Saharan horned viper *Cerastes cerastes* (Linnaeus, 1758) Northern Algerian Sahara. Serket 18:22–26.

- MEHTA, R. S. 2003. Prey-handling behavior of hatchling *Elaphe helena* (Colubridae). Herpetologica 59:469–474.
- MEHTA, R. S., K. E. DALE, AND B. A. HIGGINS. 2020. Marine protection induces variation in the California moray, *Gymnothorax morax*. Integrative and Comparative Biology 60:522–534.
- MENDELSON, J. R. III, AND A. J. ADAMS. 2014. Diadophis punctatus (Ringnecked Snake). Diet. Herpetological Review 45:709–710.
- MIZUNO, T., AND Y. KOJIMA. 2015. A blindsnake that decapitates its termite prey. Journal of Zoology 297:220–224.
- MOCINO-DELOYA, E., K. SETZER, M. HEASKER, AND S. PEURACH. 2015. Diet of the New Mexico ridge-nosed rattlesnake (*Crotalus willardi*) the Sierra Pan Duro, Mexico. Journal of Herpetology 49:104–107.
- MODAHL, C. M., F. S. MRINALINI, AND S. P. MACKESSY. 2018. Adaptive evolution of distinct prey-specific toxin genes in rear-fanged snake venom. Proceedings of the Royal Society B: Biological Sciences 285:20181003.
- MOFFETT, M. 2002. Bit. Outside Magazine (April):102 S. P. 105, 130.
- MONTEIRO, C., C. E. MONTGOMERY, F. SPINA, R. J. SAWAYA, AND M. MARTINS. 2006. Feeding, reproduction, and morphology of *Bothrops mattogrossensis* (Serpentes, Viperidae, Crotalinae) in the Brazilian Pantanal. Journal of Herpetology 40:408–413.
- MOON, B. R., AND A. M. RABATSKY. 2004. *Bogertophis subocularis* (Trans-Pecos rat snake). Prey. Herpetological Review 35:175.
- MOON, B. R., P. M. CONN, AND A. M. RABATSKY. 2004. Agkistrodon contortrix (Copperhead). Maximum prey size. Herpetological Review 35:174.
- MOON, B. R., D. A. PENNING, M. SEGALL, AND A. HERREL. 2019. Feeding in snakes: form, function, and evolution of the feeding system. Pp. 527–574 in V. Bels and I. Q. Whishaw (eds.), Feeding in Vertebrates. Springer Nature, Switzerland.
- MUFF, S., E. B. NILSEN, R. B. O'HARA, AND C. R. NATER. 2022. Rewriting results sections in the language of evidence. Trends in Ecology and Evolution 37:203–210.
- MUKERJEE, S., AND M. R. HEITHAUS. 2013. Dangerous prey and daring predators: a review. Biological Reviews 88:550–563.
- MULAIK, S. 1938. Notes on Mustela frenata frenata. Journal of Mammalogy 19:104–105.
- MULCAHY, D. G., J. R. MENDELSON III, K. W. SETSER, AND E. HOLLENBECK. 2003. Crotalus cerastes (Sidewinder). Prey/predator weight-ratio. Herpetological Review 34:64.
- MURPHY, J. C., MUMPUNI, R. DE LANG, D. J. GOWER, AND K. L. SANDERS. 2012. The Moluccan short-tailed snakes of the genus *Brachyorrhos* Kuhl (Squamata: Serpentes: Homalopsidae) and the status of *Calamophis* Meyer. Raffles Bulletin of Zoology 60:501–514.
- MUSHINSKY, H. R. 1987. Foraging ecology. Pp. 302–334 in R. A. Seigel, J. T. Collins, and S. S. Novak (eds.), Snakes: Ecology and Evolutionary Biology. Macmillan, USA.
- NAIK, H., M. M. KGADITSE, AND G. J. ALEXANDER. 2021. Ancestral reconstruction of diet and fang condition in the Lamprophildae: implications for the evolution of venom systems in snakes. Journal of Herpetology 55:1–10.
- NATUSCH, D. J. D., AND J. A. LYONS. 2012. Relationships between ontogenetic changes in prey selection, head shape, sexual maturity, and colour in an Australasian python (*Morelia viridis*). Biological Journal of the Linnean Society 107:269–276.
- NATUSCH, D., J. LYONS, L.-A. MEARS, AND R. SHINE. 2021. Biting off more than you can chew: attempted predation on a human by a giant snake (*Simalia amethistina*). Austral Ecology 46:159–162.
- NOGUEIRA, C., R. J. SAWAYA, AND M. MARTINS. 2003. Ecology of the pitviper, *Bothrops moojeni*, in the Brazilian cerrado. Journal of Herpetology 37:653–659.
- NOWAK, E. M., T. C. THEIMER, AND G. W. SCHUETT. 2008. Functional and numerical responses of predators: where do vipers fit in the traditional paradigms? Biological Reviews 83:601–620.
- O'CONNOR, A. P., J. L. WALLACE, R. E. WEAVER, AND M. P. HAYES. 2010. Pygmy short-horned lizard (*Phrynosoma douglasii*): unrecorded prey for the Great Basin nightsnake (*Hypsiglena chlorophaea deserticola*). Northwestern Naturalist 91:79–81.
- OLIVEIRA, M. E., AND M. MARTINS. 2003. *Bothrops atrox* (Common Lancehead). Prey. Herpetological Review 34:61–62.
- OLIVEIRA, L. C., A. K. LEITE, G. S. PAGEL, H. A. ARAÚJO, AND M. S. TINCO. 2019. Erythrolamprus miliaris merremi (Watersnake). Diet. Herpetological Review 50:800.

- O'SHEA, M., A. DE SILVA, AND S. A. M. KULARATNE. 2004. *Daboia russelii russelii* (Sri Lankan Russell's Viper). Large prey. Herpetological Review 35:64.
- O'SHEA, M., F. PARKER, AND H. KAISER. 2015. A new species of New Guinea worm-eating snake, genus *Toxicocalamus* (Serpentes: Elapidae) from the Star Mountains of Western Province, Papua New Guinea, with a revised dichotomous key to the genus. Bulletin of the Museum of Comparative Zoology 161:241–264.
- O'SHEA, M., A. ALLISON, AND H. KAISER. 2018. The taxonomic history of the enigmatic Papuan snake genus *Toxicocalamus* (Elapidae: Hydrophiinae), with the description of a new species from the Managalas Plateau of Oro Province, Papua New Guinea, and a revised dichotomous key. Amphibia-Reptilia 39:403–433.
- O'SHEA, M., P. BLUM, AND H. KAISER. 2020. Discovery of the second specimen of *Toxicocalamus ernstmayri* O'Shea et al. 2015 (Squamata: Elapidae), the first from Papua Provice, Indonesia, with comments on the type locality of *T. grandis* (Boulenger, 1914). Bonn Zoological Bulletin 69:395–411.
- PARKER, W. S., AND E. R. PIANKA. 1973. Notes on the ecology of the iguanid lizard, *Sceloporus magister*. Herpetologica 29:143–152.
- PARKER, W. S., AND E. R. PIANKA. 1974. Further ecological observations on the western banded gecko, *Coleonyx variegatus*. Copeia 1974:528– 531.
- PASSOS, P., A. SCANFERLA, P. R. MELO-SAMPAIO, J. BRITO, AND A. ALMEN-DARIZ. 2019. A giant on the ground: qnother large-bodied *Atractus* (Serpentes: Dipsadidae) from Ecuadorian Andes, with comments on the dietary specializations of the goo-eaters snakes. Anais da Academia Brasileira de Ciências 91:e20170976.
- PATCHELL, R. C., AND R. SHINE. 1986. Feeding mechanisms in pygopodid lizards: how can *Lialis* swallow such large prey? Journal of Herpetology 20:59–64.
- PAULY, G. B., AND M. F. BENARD. 2002. Crotalus viridis oreganus (Northern Pacific Rattlesnake). Costs of feeding. Herpetological Review 33:56– 57.
- PHELPS, T. 2002. A study of the black mamba (*Dendroaspis polylepis*) in Kwa-Zulu Natal, South Africa, with particular reference to longterm refugia. Herpetological Bulletin 80:7–19.
- PIANKA, E. R. 1970. Comparative autecology of the lizard *Cnemidophorus tigris* in different parts of its geographic range. Ecology 51:703–720.
- PIANKA, E. R., AND W. S. PARKER. 1972. Ecology of the iguanid lizard Callisaurus draconoides. Copeia 1972:493–508.
- PIANKA, E. R., AND W. S. PARKER. 1975. Ecology of horned lizards: a review with special reference to *Phrynosoma platyrhinos*. Copeia 1975:141–162.
- PIETSCH, T. W., AND R. J. ARNOLD. 2020. Frogfishes: Biodiversity, Zoogeography, and Behavioral Ecology. Johns Hopkins University Press, USA.
- PINTO-COELHO, D., M. MARTINS, AND P. R. GUIMĀREIS, JR. 2021. Network analyses reveal the role of large snakes in connecting feeding guilds in a species-rich Amazonian snake community. Ecology and Evolution 11:6558–6568.
- PIZZATTO, L., O. A. V. MARQUES, AND K. FACURE. 2009. Food habits of Brazilian boid snakes: overview and new data, with special reference to *Corallus hortulanus*. Amphibia-Reptilia 30:533–544.
- PIZZATTO, L., J. L. DE OLIVEIRA, O. A. V. MARQUES, AND M. MARTINS. 2018. Body shape and food habits of South American goo-eater snakes of the genus *Sibynomorphus*. South American Journal of Herpetology 13:300–307.
- PLATT, J. S., T. R. RAINWATER, J. C. MEERMAN, AND S. M. MILLER. 2016. Notes on the diet, foraging behavior, and venom of some snakes in Belize. Mesoamerican Herpetology 3:162–170.
- PLATT, J. S., H. A. BARRETT, L. ASH, J. A. MARLIN, S. M. BOYLAN, AND T. R. RAINWATER. 2021. Predation on turkey vultures (*Cathartes aura*): a new observation and review. Journal of Raptor Research 55:455– 459.
- PLATT, S. G., A. W. HAWKES, AND T. R. RAINWATER. 2001. Diet of the canebrake rattlesnake (*Crotalus horridus atricaudatus*): an additional record and review. Texas Journal of Science 53:115–120.
- PLEGUEZUELOS, J. M., S. HONRUBIA, AND S. CASTILLO. 1994. Diet of the false smooth snake, *Macroprotodon cucullatus* (Serpentes, Colubridae) in the western Mediterranean area. Herpetological Journal 4:98–105.
- PLUMMER, M. V. 1977. Predation by black rat snakes in bank swallow colonies. Southwestern Naturalist 22:147–148.
- POMMER-BARBOSA, R. A., J. F. T. REIS, J. R. EVANGELISTA, W. P. FERREIRA, S. ALBUQUERQUE, M. A. OLIVEIRA, AND A. L. C. PRESTES. 2022. Predation on Amphisbaena fuliginosa Linnaeus, 1758 by Anilius scytale

(Linnaeus, 1758) in the southwestern Brazilian Amazon. Herpetology Notes 15:615–617.

- POOLEY, S. 2022. The challenge of compassion in predator conservation. Frontiers in Psychology 13:977703.
- PORTILLO, F., E. L. STANLEY, W. R. BRANCH, W. CONRADIE, M-O RÖDEL, J. PENNER, M. F. BAREJ, C. KUSAMBA, O. S. G. PAUWELS, W. M. MUNINGA, ET AL. 2019. Evolutionary history of burrowing asps (Lamprophiidae: Atractaspidinae) with emphasis on fang evolution and prey selection. PLoS ONE 14(4):e0214889.
- POUGH, F. H., AND J. D. GROVES. 1983. Specializations of the body form and food habits of snakes. American Zoologist 23:443–454.
- POUGH, F. H., R. M. ANDREWS, M. L. CRUMP, A. H. SAVITZKY, K. D. WELLS, AND M. C. BRANDLEY. 2016. Herpetology, 4th edition. Sinauer Associates, USA.
- PRÖTZEL, D., J. FORSTER, T. KRAUTZ, AND F. GLAW. 2018. Predator versus predator: four-lined snake (*Elaphe quatuorlineata*) feeding on a least weasel (*Mustela nivalis*) in Istria, Croatia. Spixiana 41:157–159.
- PRUDENTE, A. L. C., A. C. MENKS, AND G. F. MASCHI. 2014. Diet and reproduction of the western indigo snake *Drymarchon corais* (Serpentes: Colubridae) from the Brazilian Amazon. Herpetology Notes 7:99– 108.
- PUTMAN, B. J., AND R. W. CLARK. 2015. Habitat manipulation in hunting rattlesnakes (*Crotalus* species). Southwestern Naturalist 60:374–377.
- PUTMAN, B. J., R. WILLIAMS, E. LI, AND G. B. PAULY. 2021. The power of community science to quantify ecological interactions in cities. Nature Scientific Reports 11:3069.
- QUICK, J. S., H. K. REINERT, E. R. DE CUBA, AND R. A. ODUM. 2005. Recent occurrence and dietary habits of *Boa constrictor* on Aruba, Dutch West Indies. Journal of Herpetology 39:304–307.
- QUINN, A., AND S. CARMODY. 2021. Carphophis amoenus (Common Wormsnake). Diet and foraging behavior. Herpetological Review 52:864.
- RABB, G. B., AND H. MARZ. 1973. Major ecological and geographical patterns in the evolution of colubroid snakes. Evolution 27:69–83.
- RAGE, J.-C., AND S. BAILON. 2011. Amphibia and Squamata. Pp. 467–478 in T. Harrison (ed.), Paleontology and Geology of Laetoli: Human Evolution in Context. Volume 2. Fossil Homins and the Associated Fauna. Springer, Netherlands.
- RAJENDRAN, M. 1985. Studies in uropeltid snakes. Madurai Kamaraj University, India.
- RASMUSSEN, J. B., AND K. M. HOWELL. 1998. A review of Barbour's shortheaded viper, *Adenorhinos barbouri* (Serpentes: Viperidae). African Journal of Herpetology 47:69–75.
- RAY, J. M., C. E. MONTRGOMERY, H. K. MAHON, A. H. SAVITZKY, AND K. R. LIPPS. 2012. Goo-eaters: diets of the neotropical snakes *Dipsas* and *Sibon* in central Panama. Copeia 2012:197–202.
- REED, R. N. 1997. Trimorphodon biscutatus quadruplex (Lyre Snake). Diet. Herpetological Review 28:206.
- REED, R. N., AND G. H. RODDA. 2009. Giant constrictors: biological and management profiles and an establishment risk assessment for nine large species of pythons, anacondas, and the boa constrictor. U. S. Geological Survey Open-File Report 2009–1202.
- REID, J. R., AND T. E. LOTT. 1963. Feeding of *Leptotyphlops dulcis dulcis* (Baird and Girard). Herpetologica 19:141–142.
- REINERT, H. K., L. M. BUSHAR, C. L. ROCCO, AND R. A. ODUM. 2008. Ecology of the Aruba Island rattlesnake, *Crotalus durissus unicolor*. Pp. 335–352 in W. K. Hayes, K. R. Beaman, M. D. Cardwell, and S. P. Bush (eds.), The Biology of Rattlesnakes. Loma Linda University Press, USA.
- REINERT, H. K., A. E. LETO, J. A. TUMALIUAN, S. JACKREL, W. I. LUTTERSCH-MIDT, AND L. M. BUSHAR. 2021. A long-term dietary assessment of invasive *Boa constrictor* of Aruba. Herpetological Conservation and Biology 16:211–224.
- REPP, R. A., AND G. W. SCHUETT. 2009. *Heloderma suspectum* (Gila Monster). Diet and predatory behavior. Herpetological Review 40:343– 345.
- REVAULT, P. 1996. Scolopendra morsitans Linnaeus, 1758: a characteristic prey of the African carpet viper Echis ocellatus Stemmler, 1970. Memoirs Muséum National d'Histoire Naturelle 169:495–499.
- RIBBLE, D.O., AND G.B. RATHBUN. 2018. Preliminary observations on home ranges and natural history of *Scotinomys tenguina* in Costa Rica. Mammalia 82:490–493.
- RIBEIRO, M. A. JR., S. F. FERRARI, J. R. F. LIMA, C. R. DA SILVA, AND J. D. LIMA. 2016. Predation of a squirrel monkey (*Siamiri sciureus*) by an Amazon tree boa (*Corallus hortulanus*): even small boids may be a potential threat to small-bodied platyrrhines. Primates 57:317–322.

- RIVAS, J. A. 1998. Predatory attacks of green anacondas (*Eunectes muri-nus*) on adult human beings. Herpetological Natural History 6:157–159.
- RIVAS, J. A. 2020. Anaconda: The Secret Life of the World's Largest Snake. Oxford University Press, UK.
- ROBERTS, J. R., AND C. C. AUSTIN. 2020. A new species of New Guinea worm-eating snake (Elapidae: *Toxicocalamus* Boulenger, 1896), with comments on postfrontal bone variation based on micro-computed tomography. Journal of Herpetology 54:446–459.
- ROBINSON, W. G., G. ROMPRÉ, AND T. R. ROBINSON. 2005. Videography of Panama bird nests shows snakes are principal predators. Ornithología Neotropical 16:187–195.
- RODRÍGUEZ, M. C., AND H. DRUMMOND. 2000. Exploitation of avian nestlings and lizards by insular milksnakes, *Lampropeltis triangulum*. Journal of Herpetology 34:139–142.
- RODRIGUEZ-ROBLES, J. A. 1994. Are the Duvernoy's gland secretions of colubrid snakes venoms? Journal of Herpetology 28:388–390.
- RODRIGUEZ-ROBLES, J. A. 1998. Alternative perspectives on the diet of gopher snakes (*Pituophis catenifer*, Colubridae): literature records versus stomach contents of wild and museum specimens. Copeia 1998:463–466.
- RODRÍGUEZ-ROBLES, J. A. 2002. Feeding ecology of the North American gopher snakes (*Pituophis catenifer*, Colubridae). Biological Journal of the Linnean Society 77:165–183.
- RODRÍGUEZ-ROBLES, J. Á., AND H. W. GREENE. 1999. Food habits of the long-nosed snake, *Rhinocheilus lecontei*, a "specialist" predator? Journal of Zoology 248:489–499.
- RODRÍGUEZ-ROBLES, J. Á., C. J. BELL, AND H. W. GREENE. 1999a. Food habits of the glossy snake, *Arizona elegans*, with comparisons to the diet of sympatric long-nosed snakes, *Rhinocheilus lecontei*. Journal of Herpetology 33:87–92.
- RODRIGUEZ-ROBLES, J. A., C. J. BELL, AND H. W. GREENE. 1999b. Gape size and evolution of diet in snakes: feeding ecology of erycine boas. Journal of Zoology 248:49–58.
- RODRÍGUEZ-ROBLES, J. Á., D. G. MULCAHY, AND H. W. GREENE. 1999c. Feeding ecology of the desert nightsnake, *Hypsiglena torquata* (Colubridae). Copeia 1999:93–100.
- ROESCH, M. A., B. DYMOND, AND N. C. COLE. 2022. Feeding observations of the keel-scaled boa, *Casarea dussumieri* (Serpentes: Bolyeriidae), on Round Island, Mauritius, showing the use of its intramaxillary joint. Herpetology Notes 15:519–522.
- RORABAUGH, J. C., A. T. HOLYCROSS, AND T. C. BRENNAN. 2020. Tantilla nigriceps Plains Black-headed Snake. Pp. 372–378 in A. T. Holycross and J. C. Mitchell (eds.), Snakes of Arizona. Eco Publishing, USA.
- SALMÃO, M., G. DA, AND I. L. LAPORTA-FERREIRA. 1994. The role of secretions from the supralabial, infralabial, and Duvernoy's glands of the slug-eating snake *Sibynomorphus mikani* (Colubridae: Dipsadinae) in the immobilization of molluscan prey. Journal of Herpetology 28:369–371.
- SANTANA, S. 2011. Micrurus distans (Sinaloan coralsnake). Diet. Herpetological Review 42:294.
- SAVIOZZI, P., AND M. A. L. ZUFFI. 1997. An integrated approach to the study of the diet of *Vipera aspis*. Herpetological Review 28:23–24.
- SAVITZKY, A. H. 1983. Coadapted character complexes among snakes: fossoriality, piscivory, and durophagy. American Zoologist 23:397– 409.
- SAZIMA, I., AND M. MARTINS. 1990. Presas grandes e serpentes jovens: quando os olhos são maiores que a boca. Memórias do Instituto Butantan 52:73–79.
- SCANLON, J. D., M. S. Y. LEE, M. W. CALDWELL, AND R. SHINE. 1999. The palaeobiology of the primitive snake *Pachyrhachis*. Historical Biology 13:127–152.
- SCHALK, C. M., AND M. V. COVE. 2018. Squamates as prey: predator diversity patterns and predator-prey size relationships. Food Webs 16:e00103.
- SCHMIDT, K. P. 1932. Stomach contents of some American coral snakes, with the description of a new species of *Geophis*. Copeia 1932:6–9.
- SCHOENER, T. W. 1971. Theory of feeding strategies. Annual Review of Ecology and Systematics 2:369–404.
- SCHUETT, G. W., E. M. NOWAK, AND R. A. REPP. 2002. Crotalus cerberus (Arizona black rattlesnake). Diet and prey size. Herpetological Review 33:210–211.
- SEIB, R. L. 1984. Prey use in three syntopic neotropical racers. Journal of Herpetology 18:412–420.
- SEIB, R. L. 1985a. Feeding Ecology and Organization of Neotropical Snake Faunas. Ph.D. diss., University of California, USA.

- SEIB, R. L. 1985b. Euryphagy in a tropical snake, Coniophanes fissidens. Biotropica 17:57–64.
- SHEPARD, D. R., C. A. PHILLIPS, M. J. DRESLIK, AND B. C. JELLEN. 2004. Prey preference and diet of neonate eastern massasaugas (*Sistrurus c. catenatus*). American Midland Naturalist 152:360–368.
- SHEWCHUK, C. H., AND J. D. AUSTIN. 2001. Food habits of the racer (*Coluber constrictor mormon*) in the northern part of its range. Herpetological Journal 11:151–155.
- SHINE, R. 1977. Habitats, diets, and sympatry in snakes: a study from Australia. Canadian Journal of Zoology 55:1118–1128.
- SHINE, R. 1986. Ecology of a low-energy specialist: food habits and reproductive biology of the Arafura filesnake (Achrocordidae). Copeia 1986:424–437.
- SHINE, R. 1991. Why do larger snakes eat larger prey items? Functional Ecology 5:493–502.
- SHINE, R., AND J. S. KEOGH. 1996. Food habits and reproduction of the endemic Melanesian elapids: are tropical snakes really different? Journal of Herpetology 30:238–247.
- SHINE, R., AND J. THOMAS. 2005. Do lizards and snakes really differ in their ability to take large prey? A study of relative mass and feeding tactics in lizards. Oecologia 144:492–498.
- SHINE, R., AND J. K. WEBB. 1990. Natural history of Australian typhlopid snakes. Journal of Herpetology 24:357–363.
- SHINE, R., P. S. HARLOW, W. R. BRANCH, AND J. K. WEBB. 1996. Life on the lowest branch: sexual dimorphism, diet, and reproductive biology of an African twig snake, *Thelotornis capensis* (Serpentes, Colubridae). Copeia 1996:290–299.
- SHINE, R., W. R. BRANCH, P. S. HARLOW, AND J. K. WEBB. 1998. Reproductive biology and food habits of horned adders, *Bitis caudalis* (Viperidae), from southern Africa. Copeia 1998: 391–401.
- SHINE, R., W. R. BRANCH, J. K. WEBB, P. S. HARLOW, T. SHINE, AND J. S. KEOGH. 2007. Ecology of cobras from southern Africa. Journal of Zoology 272:183–193.
- SILER, C. D., R. L. J. WELTON, M. BROWN, C. INFANTE, AND A. C. DIESMOS. 2011. Ophiophagus hannah (king cobra). Diet. Herpetological Review 42:297.
- SLIP, D. J., AND R. SHINE. 1988. Feeding habits of the diamond python, *Morelia s. spilota*: ambush predation by a boid snake. Journal of Herpetology 22:322–330.
- SMITH, K. T., AND A. SCANFERLA. 2016. Fossil snake preserving three trophic levels and evidence for an ontogenetic dietary shift. Palaeobiology and Palaeoenvironments 96:589–599.
- SOARES, M. S. C., L. SOUSA, AND J. P. BARREIROS. 2003. Feeding habits of the lizardfish *Synodus saurus* (Linnaeus, 1758) (Actinopterygii). Aqua, Journal of Ichthyology and Aquatic Biology 7:29–38.
- SOLÓRZANO, A., AND H. W. GREENE. 2012. Predation in nature by a scorpion-hunter, *Stenorrhina freminvillei* (Serpentes, Colubridae). Cuadernos de Investigación UNED 4:31–32.
- SOLÓRZANO, A., AND M. SASA. 2022. Diet and feeding behavior of the mussurana (Clelia Clelia, Serpentes: Dipsadidae) in Costa Rica: report of five cases. Revista Latinoamericana de Herpetolgía 5:39– 42.
- SOLÓRZANO, A., L. D. WILSON, AND L. PORRAS. 2012. *Tantilla reticulata* (Lined Crowned Snake). Diet. Herpetological Review 43:153.
- SORRELL, G. G., S. M. BOBACK, R. N. REED, S. GREEN, C. E. MONTGOMERY, L. S. SOUZA, AND M. CHIARAVIGLIO. 2011. Boa constrictor (*Boa Constrictor*). Foraging behavior. Herpetological Review 42:281.
- SPARKS, A. M., C. LIND, AND E. N. TAYLOR. 2015. Diet of the northern Pacific rattlesnake (*Crotalus o. oreganus*) in California. Herpetological Review 46:161–165.
- STAKE, M. M., F. R. THOMPSON III, J. FAABORG, AND D. E. BURHANS. 2005. Patterns of snake predation at songbird nests in Missouri and Texas. Journal of Herpetology 39:215–222.
- STENDER-OLIVEIRA, F., M. MARTINS, AND O. A. V. MARQUES. 2016. Food habits and reproductive biology of tail-luring snakes of the genus *Tropidodryas* (Dipsadidae: Xenodontinae) from Brazil. Herpetologica 72:73–79.
- STRICKLAND, J. L., S. CARTER, F. KRAUS, AND C. L. PARKINSON. 2016. Snake evolution in Melanesia: origin of the Hydrophiinae (Serpentes, Elapidae), and the evolutionary history of the enigmatic New Guinean elapid *Toxicocalamus*. Zoological Journal of the Linnean Society 178:663–678.
- STRUSSMANN, C. 1997. Hábitos alimentares da sucuri-amarela, Eunectes notaeus Cope, 1862, no Pantanal Mato-Grossense. Biociéncias 5:35– 52.
- SULLIVAN, B. K., AND S. A. WEINSTEIN. 2017. Foraging and prey handling in the Western patch-nosed snake (*Salvadora hexalepis*), and

consideration of venom use in non-front-fanged colubroid snakes. Herpetological Review 48:19–23.

- SUNQUET, M. E. 1982. An unusual death of an Indian python (*Python molurus*). Hornbill 1:9.
- SWANNACK, T. M., AND M. R. J. FORSTNER. 2003. Micrurus fulvius tener (Texas coral snake). Diet. Herpetological Review 34:376.
- SZCZYGIEL, H., AND R. A. PAGE. 2020. When the hunter become the hunted: foraging bat attacked by pit viper at frog chorus. Ecology 101:e03111.
- TAYLOR, E. N. 2001. Diet of the Baja California rattlesnake, *Crotalus enyo* (Viperidae). Copeia 2001:553–555.
- TEODORO, L. O., T. L. ANDREANI, M. A. G. SILVA, R. F. OLIVEIRA, J. F. R. TONINI, AND A. R. MORAIS. 2022. Patterns and trends in the publication of natural history notes in herpetology journals over the last decade. Journal of Herpetology 56:211–217.
- TETZLAFF, S. J., M. J. RAVESI, AND B. A. KINGSBURY. 2014. Sistrurus catenatus catenatus (Eastern massasauga). Diet. Herpetological Review 45:712–713.
- THOMAS, O., AND S. J. R. ALLAIN. 2021. A review of prey taken by anacondas (Squamata: Boidae: *Eunectes*). Reptiles and Amphibians 28:329– 334.
- TOSA, M. I., E. H. DZIEDZIC, C. L. APPEL, J. URBINA, A. MASSEY, J. RUPRECHT, C. E. ERIKSSON, J. E. DOLLIVER, D. M. LESMEISTER, M. G. BETTS, ET AL. 2021. The rapid rise of next-generation natural history. Frontiers in Ecology and Evolution 9:698131.
- TRAIL, P. W. 1987. Predation and antipredator behavior at Guianan Cock-of-the Rock leks. Auk 104:496-507.
- TSAI, C.-H., C.-H. HSIEH, AND T. NAKAZAWA. 2016. Predator-prey mass ratio revisited: does preference of relative prey body size depend on individual predator size? Functional Ecology 30:1979–1987.
- Tyler, J. D. 1977. Coachwhip preys on horned lizard. Southwestern Naturalist 22:146.
- UHLER, F. M., C. COTTAM, AND T. E. CLARKE. 1939. Food of snakes of the George Forest, Virginia. Transactions of the North American Wildlife Conference 1939:605–622.
- VALDUJO, P. H., C. NOGERIA, AND M. MARTINS. 2002. Ecology of *Bothrops neuwiedi paoloensis* (Serpentes: Viperidae: Crotalinae) in the Brazilian cerrado. Journal of Herpetology 36:169–176.
- VALENCIA-HERVERTH, J., R. VALENCIA-HERVERTH, AND L. FERNÁNDEZ-BAD-ILLO. 2021. *Coniophanes imperialis* (regal black-striped snake). Diet. Herpetological Review 52:151.
- VAN DEN BURG, M. P. 2020. How to source and collate natural history information: a case study of reported prey items of *Erythrolamprus miliaris* (Linnaeus, 1758). Herpetology Notes 13:739–746.
- VASQUEZ-CRUZ, V. 2020. New prey records for the Atlantic Central American milksnake *Lampropeltis polyzona* (Serpentes: Colubridae). Phyllomedusa 19:107–111.
- VELA, R. A., J. L. CASTILLO JUÁREZ, AND A. I. CONTRERAS CALVARIO. 2020. Predation on rainbow ameivas, *Holcosus undulatus* (sensu lato), and a second record of predation on *H. amphigrammus* (Smith and Laufe 1945) by a terciopelo (*Bothrops asper*) in Veracruz, Mexico. Reptiles and Amphibians 27:422–425.
- VENEGAS-BARRERA, C. S., AND J. MANJARREZ. 2001. Thamnophis scalaris (Mexican Alpine Blotched Garter Snake). Diet. Herpetological Review 32:266.
- VINCENT, S. E., AND A. MORI. 2008. Determinants of feeding performance in free-ranging pit-vipers (Viperidae: *Ovophis okinavensis*): key roles for head size and body temperature. Biological Journal of the Linnean Society 93:53–62.
- VINCENT, S. E., P. D. DANG, A. HERREL, AND N. J. KLEY. 2006a. Morphological integration and adaptation in the snake feeding system: a comparative phylogenetic study. Journal of Evolutionary Biology 19:1545–1554.
- VINCENT, S. E., B. R. MOON, R. SHINE, AND A. HERREL. 2006b. The functional meaning of "prey size" in water snakes (*Nerodia fasciata*, Colubridae). Oecologia 147:204–211.
- VINCENT, S. E., B. R. MOON, AND A. HERREL, AND N. J. KLEY. 2007. Are ontogenetic shifts in diet linked to shifts in feeding mechanics? Scaling of the feeding apparatus in the banded watersnake Nerodia fasciata. Journal of Experimental Biology 210:2057–2069.
- VISCO, D. M., AND T. W. SHERRY. 2015. Increased abundance, but reduced nest-predation in the chestnut-backed antbird in Costa Rican rainforest fragments: surprising impacts of a pervasive snake species. Biological Conservation 188:22–31.
- VORIS, H. K., AND M. W. MOFFETT. 1981. Size and proportion relationship between the beaked sea snake and its prey. Biotropica 13:15–19.

- VORIS, H. K., AND H. H. VORIS. 1983. Feeding strategies in marine snakes: an analysis of evolutionary, morphological, behavioral, and ecological relationships. American Zoologist 23:411–425.
- VRCIBRADIC, D., M. ALMEDA-GOMES, C. Č. SIGUEIRA, V. N. T. BORGES-JUNIOR, AND C. F. D. ROCHA. 2011. Oxyrhopus petola digitalis (false coral snake). Prev. Herpetological Review 42:299–300.
- WARKENTIN, K. M. 1995. Adaptive plasticity in hatching age: a response to predation risk trade-offs. Proceedings of the National Academy of Sciences USA 92:3507–3510.
- WARNER, J. K. 2011. Naja melanoleuca (Forest Cobra). Diet and foraging behavior. Herpetological Review 42:295.
- WARNER, J. K., AND G. ALEXANDER. 2011. Bitis gabonica (Gaboon Viper). Diet and prey size. Herpetological Review 42:280–281.
- WEBB, J. K., AND R. SHINE. 1993a. Dietary habits of Australian blindsnakes (Typhlopidae) Copeia 1993:762–770.
- WEBB, J. K., AND R. SHINE. 1993b. Prey-size selection, gape limitation and predator vulnerability in Australian blindsnakes (Typhlopidae). Animal Behaviour 45:1117–1126.
- WEBB, J. K., R. SHINE, W. R. BRANCH, AND P. S. HARLOW. 2000. Life-history strategies in basal snakes: reproduction and dietary habits of the African thread snake *Leptyphlops scutifrons* (Serpentes: Leptotyphlopidae). Journal of Zoology 250:321–327.
- WEBBER, M. M., T. JEZKOVA, AND J. A. RODRIGUEZ-ROBLES. 2016. Feeding ecology of sidewinder rattlesnakes, *Crotalus cerastes* (Viperidae). Herpetologica 72:324–330.
- WERNER, F. 1909. Über neue oder seltene Reptilien des Naturhistorischen Museums in Hamburg. Jahrbuch der Hamburgischen Wissenschaftlichen Anstalten 26:205–247.
- WHITING, M. J., B. D. GREENE, J. R. DIXON, A. L. MERCER, AND C. C. ECKER-MAN. 1992. Observations on the foraging ecology of the western coachwhip snake, *Masticophis flagellum testaceus*. The Snake 24:157– 160.
- WILLSON, J. D., C. T. WINNE, M. A. PILGRIM, C. S. ROMANEK, AND J. W. GIB-BONS. 2010. Seasonal variation in terrestrial resource subsidies influence trophic niche width and overlap in two aquatic snake species: a stable isotope approach. Oikos 119:1161–1171.
- WILLSON, J. D., AND W. A. HOPKINS. 2011. Prey morphology constrains the feeding ecology of an aquatic generalist predator. Ecology 92:744–754.
- WINCK, G. R., L. F. DANTAS, M. ALMEIDA-DANTOS, F. B. DA SILVA TELLES, L. MAGALHAÃES, M. R. PEREIRA, AND C. F. D. ROCHA. 2012. *Philodryas* olfersi. Diet. Herpetological Review 43:151.
- WISEMAN, K. D. 2018. Art in herpetology: Foothill Yellow-Legged Frog, Bungarus—Lycodon. Herpetological Review 49:782–783.
- WISEMAN, K. D., AND J. BETTASO. 2007. Rana boylii (Foothill Yellow-Legged Frog). Cannibalism and predation. Herpetological Review 38:193.
- WISEMAN, K. D., H. W. GREENE, M. S. KOO, AND D. J. LONG. 2019. Feeding ecology of a generalist predator, the California kingsnake (*Lampropeltis californiae*): why rare prey matter. Herpetological Conservation and Biology 14:1–30.
- WOSTL, L., T. J. HINKLE, B. LARDNER, AND R. N. REED. 2012. *Boiga irregularis* (brown treesnake). Diet. Herpetological Review 42:282.
- ZAHER, H., L. DE OLIVEIRA, F. L. GRAZZIOTIN, M. CAMPAGNER, C. JARED, M. M. ANTONIAZZI, AND A. L. PRUDENTE. 2014. Consuming viscous prey: a novel protein-secreting delivery system in neotropical snaileating snakes. BMC Evolutionary Biology 14:58.
- ZAHER, H., R. M. MURPHY, J. C. ARREDONDO, R. GRABOSKI, P. R. MACHADO-FILHO, K. MAHLOW, G. G. MONTINGELLI, A. B. QUADROS, N. L. ORLOV, M. WILKINSON, ET AL. 2019. Large-scale molecular phylogeny, morphology, divergence-time estimation, and the fossil record of advanced caenophidian snakes (Squamata: Serpentes). PLoS ONE 14(5):e0216148.
- ZAHER, H., D. M. MOHABEY, F. G. GRAZZIOTIN, AND J. A. WILSON MANTILLA. 2022. The skull of *Sanajeh indicus*, a Cretaceous snake with an upper temporal bar, and the origin of ophidian wide-gaped feeding. Zoological Journal of the Linnean Society 197:656–697.
- ZANCOLLI, G., J. J. CALVETE, M. D. CARDWELL, H. W. GREENE, W. K. HAYES, M. J. HEGARTY, H.-W. HERRMANN, A. T. HOLYCROSS, D. I. LANNUTTI, J. F. MULLEY, ET AL. 2019. When one phenotype is not enough: divergent evolutionary trajectories govern venom variation in a widespread rattlesnake species. Proceedings of the Royal Society B: Biological Sciences 286:20182735.
- ZUG, G. Ř., AND I. INEICH. 1993. Review of the biology and morphology of the Fijian bola *Ogmodon vitianus* (Elapidae). The Snake 25:9–20.

ZULUAGA-ISAZA, J. C., J. A. ROJAS-MORALES, R. F. DÍAZ-AYALA, AND V. A. RAMÍREZ-CASTAÑO. 2015. *Pseustes shropshirei* (Shropshire's Puffing Snake). Diet. Herpetological Review 46:649.

Accepted: 23 February 2023. Published online: 8 November 2023.

#### Appendix 1.

References to aspects of MBT for snakes include Voris and Moffett (1981), Pough and Groves (1983), Herzog and Bailey (1987), Mushinsky (1987), Sazima and Martins (1990), Shine (1991), Arnold (1993), Forsman and Lindell (1993), Martins and Oliveira (1998), Rodríguez-Robles et al. (1999a,b), King (2002), Martins et al. (2002), Rodríguez-Robles (2002), Mehta (2003), Andreadis and Burghardt (2005), Boback (2005), Vincent et al. (2006a,b; 2007), Glaudas et al. (2008), Nowak et al. (2008), Vincent and Mori (2008), Willson and Hopkins (2011), Close and Cundall (2012), Hampton and Moon (2013), Loughran et al. (2013), Cundall et al. (2014), Maritz and Alexander (2014), Ribeiro et al. (2016), Banci et al. (2017), Glaudas et al. (2017a,b), Means (2017), Bartoszek et al. (2018), Jayne et al. (2018), Glaudas et al. (2019), Maritz et al. (2019), Gripshover and Jayne (2021), Marques and Sazima (2021), Natusch et al. (2021), Reinert et al. (2021), Barends and Maritz (2022a), Cundall and Irish (2022), and Jayne et al. (2022).

# APPENDIX 2.

References to general aspects of snake feeding biology without data on RPM or RPB include Bea et al. (1992), Webb and Shine (1993a), B. Greene et al. (1994), Dwyer and Kaiser (1997), Saviozzi and Zuffi (1997), Rodríguez-Robles (1998), Fitch (1999), Scanlon et al. (1999), Luiselli et al. (2001), Carreira Vidal (2002), Luiselli and Akani (2003), Quick et al. (2005), Luiselli (2006a,b), Esbérard and Vrcibradic (2007), Shine et al. (2007), Dugan and Hayes (2012), Natusch and Lyons (2012), Brown et al. (2014), Fabre et al. (2016), Klaczko et al. (2016), Sullivan and Weinstein (2017), da Silva et al. (2018), Portillo et al. (2019), van den Burg (2020), Cleuren et al. (2021), Grundler and Rabosky (2021), Naik et al. (2021), and Durso et al. (2022).

### Appendix 3.

References with RPM data from nature for non-front-fanged colubroids include Brown (1958), Tyler (1977), Seib (1984, 1985b), Hero and Magnusson (1987), Greene (1989a,b), Branch and Burger (1991), Greene and Jaksic (1992), Whiting et al. (1992), Pleguezuelos et al. (1994), Lacey et al. (1996), Shine et al. (1996), Capula et al. (1997), Reed (1997), Martins and Oliveira (1998), Rodríguez-Robles and Greene (1999), Rodríguez-Robles et al. (1999a,b,c), Feldman and Wilkinson (2000), Camper and Dixon (2000), LaBonte (2001), Venegas-Barrera and Manjarrez (2001), Rodríguez-Robles (2002), Greene and Rodríguez-Robles (2003), Cobb (2004), Moon and Rabatsky (2004), O'Connor et al. (2010), Greene et al. (2010), Dartez et al. (2011), Durso and Kiriaszis (2011), Vrcibradic et al. (2011), Wostl et al. (2012), Duarte (2012), Gatica-Colima and Córdoba-Reza (2012), Laspiur et al. (2012), Linares and Eterovick (2012), Solórzano and Greene (2012), Winck et al. (2012), Camper and Zart (2014), Mendelson and Adams (2014), Casper et al. (2015), Da Silva et al. (2015), Zuluaga-Isaza et al. (2015), Stender-Oliveira et al. (2016), Pizzatto et al. (2018), Oliveira et al. (2019), Wiseman et al. (2019), Valencia-Herverth et al. (2021), and Camper (2022).

#### APPENDIX 4.

References with RPM data from nature for aniliids, uropeltids, boids, and pythonids include Hay and Martin (1966), Greene (1983a), Trail (1987), Slip and Shine (1988), Strussmann (1997), Martins and Oliveira (1998), Rivas (1998), Shine et al. (1998), Rodríguez-Robles et al. (1999b), Boback et al. (2000), Fearn (2002), Boback (2004, 2005), Fredriksson (2005), Pizzatto et al. (2009), Reed and Rodda (2009), Barros et al. (2011), Headland and Greene (2011), Boback et al. (2016), Ribeiro et al. (2016), Bartoszek et al. (2018), Glaudas et al. (2019), Rivas (2020), Natusch et al. (2021), Reinert et al. (2021), Cundall and Irish (2022), and Pommer-Barbosa et al. (2022).

### APPENDIX 5.

References with RPM data from nature for front-fanged colubroids include Fitch and Twining (1946), Fitch (1960), Abalos et al. (1964), Voris and Moffett (1981), Greene (1984, 1992), Haagner (1991), Branch et al. (1995), Lutterschmidt et al. (1996), Martins and Oliveira (1998), Fitch (1999), Hardy and Greene (1999), Cundall and Greene (2000), Correa-Sanchez et al. (2001), Taylor (2001), Branch et al. (2002), Martins et al. (2002), Pauly and Benard (2002), Rodríguez-Robles (2002), Schuett et al. (2002), Valdujo et al. (2002), Mulcahy et al. (2003), Nogueira et al. (2003), Oliveira and Martins (2003), Swannack and Forstner (2003), Moon et al. (2004), O'Shea et al. (2004), Shepard et al. (2004), Hartmann et al. (2005), Glaudas et al. (2008), Marques et al. (2010), Gavira and Loebmann (2011), Hampton (2011), Horan et al. (2011), Hovey and Comrack (2011), Mata-Silva et al. (2011), Santana (2011), Siler et al. (2011), Sorrell et al. (2011), Warner (2011), Warner and Alexander (2011), Maritz (2012), Loughran et al. (2013), Camera et al. (2014), Colbert et al. (2014), Maritz and Alexander (2014), Tetzlaff et al. (2014), Webber et al. (2016), Means (2017), Glaudas et al. (2019), Campbell and Hewlett (2021), Da Silva et al. (2021), Marques and Sazima (2021), Martínez-Vaca León and Morales-Mávil (2021), Camper (2022), and Carbajal-Márquez et al. (2022).

#### APPENDIX 6.

References on snake taxa that do not mention RPM or RPB include López Jurado and Caballero (1981), Bea and Braña (1988), Bhupathy and Vijayan (1989), Jones and Whitford (1989), Zug and Ineich (1993), Revault (1996), Shine and Keogh (1996), Luiselli and Akani (1998), Luiselli and Angelici (1998), Rodríguez and Drummond (2000), Hill et al. (2001), Holycross et al. (2001), Platt et al. (2001), Shewchuk and Austin (2001), Clark (2002), Holycross et al. (2002), Gardner and Mendelson (2003), Monteiro et al. (2006), Machio et al. (2010), Gaiarsa et al. (2013), Prudente et al. (2014), Sparks et al. (2015), Carbajal-Márquez et al. (2016, 2020), Platt et al. (2016), Layloo et al. (2017), Prötzel et al. (2018), Bringsøe (2019), Berg et al. (2020), Bringsøe et al. (2020), Cabral et al (2020), Escalante and Acuña (2020), Feldman et al. (2020), R. Maritz et al. (2020), Vela et al.

(2020), Vásquez-Cruz (2020), Cochran et al. (2021), Conradie and Pinto (2021), Eisfeld et al. (2021), Faraone et al. (2021), Hoefer et al. (2021), Maritz et al. (2021a), Mebarki et al. (2021), and Thomas and Allain (2021).

### APPENDIX 7.

That some snakes consume prey parts rather than intact animals scarcely could have been imagined at the time of Gans' (1961) landmark paper. Subsequent discoveries include Texas Blindsnakes (Rena dulcis) and Brahminy Blindsnakes (Indotyphlops braminus) removing termite heads prior to ingestion of their abdomens (Reid and Lott, 1963; Mizuno and Kojima, 2015), Northern Cat-eyed Snakes (Leptodeira septentrionalis) and Giant Parrot Snakes (Leptophis ahaetulla) tearing off portions of treefrog egg masses (Warkentin, 1995; Gomez-Mestre and Warkentin, 2007), Gerard's Watersnakes (Gerarda prevostiana) removing and ingesting crab legs (Jayne et al., 2002), Taiwan Kukri Snakes (Oligodon formosanus) slitting and swallowing contents of sea turtle eggs (Huang et al., 2011), Banded Kukri Snakes (Oligodon fasciolatus) slicing open and eating internal organs of Black-spined Toads (Duttaphrynus melanostictus; Bringsøe et al., 2020), Blunt-headed Slug-eaters (Aplopeltura boa) sawing off opercula before ingesting the bodies of snails (Kojima et al., 2020), and Queen Snakes (Regina septemvittata) eating appendages of crayfish (Gripshover and Jayne, 2021). In some of these examples, snakes perhaps consumed pieces to avoid toxic body parts (e.g., Mizuno and Kojima, 2015; Bringsøe et al., 2020), whereas a Cottonmouth (Agkistrodon piscivorus) pulling chunks off fish carrion (Campbell and Hewlett, 2021) is noteworthy because the intact RPM of  $\sim$ 3 was almost twice the recorded maximum for snakes.

## APPENDIX 8.

For five Coachwhips (*Masticophis flagellum*) that consumed five Texas Horned Lizards (*Phrynosoma cornutum*) and one Coast Horned Lizard (*Phrynosoma blainvilli*), RPMs were 0.028 (Tyler, 1977), 0.033 and 0.066 (Whiting et al., 1992), 0.041 (LaBonte, 2001), and 0.006 and 0.056 (TNHC 87848 contained two *P. cornutum*; T. LaDuc, pers. com.), thus a range of 0.006–0.056 ( $\bar{x} = 0.038$ ). For eight *M. flagellum* that ate type II Western Whip-tailed Lizards (*Aspidoscelis tigris*; Appendix 9), RPMs were 0.015–0.191 ( $\bar{x} = 0.078$ ). A conservatively two-tailed Mann-Whitney *U*-test implies this is "weak evidence" that the medians are different (P = 0.14; significance terminology of Muff et al., 2022:206).

# APPENDIX 9.

For six Northern Pacific Rattlesnakes (*Crotalus oreganus*) eating sceloporines (four Western Fence Lizards [*Sceloporus occidentalis*], two Side-blotched Lizards [*Uta stansburiana*]), masses (g) of prey and predator, respectively, followed by sources are as follows: 11.2, 10.05, MVZ 229849; 3.0, 11.2, MVZ 6842; 2.5, 22.5, MVZ 33913; 8.0, 20.5, MVZ 50213; 15.5, 45.0, HWG 1112; and 9.04, 11.91, Pauly and Benard (2002). For 10 California Striped Whipsnakes [*Masticophis lateralis*] eating sceloporines the values were as follows: 14.0, 118.1, MVZ 227; 14.0, 98.2, MVZ 5876; 14.0, 92.7, MVZ 60986; 7.0, 153.3, MVZ 43636; 16.0, 257.6, MVZ 21915; 14.0, 105.3, MVZ 26007; 5.0, 52.8, MVZ 36428; 1.0, 173.1, MVZ 35463; 5.0, 17.3, MVZ 93655; and 3.0, 11.7, MVZ 43638. For seven Sidewinder Rattlesnakes (*C. cerastes*) eating Western Whip-tailed Lizards (*Aspidoscelis tigris*), the results were as follows: 22.5, 55.6, MVZ 57606; 7.0, 75.8, MVZ 7458; 19.0, 34.5, HWG 851; 11.0, 135.5, HWG 967; 24.0, 46.2, MVZ 63664; 11.0, 25.5, RLS 6960; and 20.5, 33.0, RLS 8141. For eight Coachwhips (*M. flagellum*) eating *A. tigris* the values were as follows: 22.5, 118.0, RWH 942; 3.5, 224.2, MVZ 20394; 21.0, 247.0, CAS 10308; 5.0, 54.8, MVZ 6692; 5.0, 103.8, MVZ 13889; 24.0, 348.5, RWH 925; 1.5, 103.8, MVZ 13889; and 20.5, 215.1, MVZ 5546. We compared RPM medians with two-tailed Mann-Whitney *U*-tests and size effects with analyses of covariance on log-transformed masses; *P*-values were <0.05 (some <0.02) implying at least "moderate evidence" of differences (Muff et al., 2022). Note only small *C. oreganus* consumed lizards, whereas *C. cerastes* of all sizes ate them; both *Masticophis* species attain a TL of >1 m, such that lizard species with small adult masses cannot provide high RPM for adults of those snake species.