

Post-Weaning Cranial Growth in Shrew Opossums (Caenolestidae): A Comparison with Bandicoots (Peramelidae) and Carnivorous Marsupials

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Abstract The patterns of development and skull ontogeny in caenolestids have been poorly studied, resulting in a limited knowledge. In this work, we report and compare the allometric growth trends of 15 variables in the three living groups of the Family Caenolestidae, represented by *Caenolestes fuliginosus*, *Lestoros inca*, and *Rhyncholestes raphanurus*. We analyzed the bivariate and multivariate allometry in comparison with morphologically convergent Australasian peramelids, as well as with other marsupials and placentals previously studied. We also report the phylogenetic signal and

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optimization of the confidence intervals of the variables analyzed in two alternative hypotheses, where Ameridelphia is considered as monophyletic and paraphyletic. *Rhyncholestes raphanurus* and *C. fuliginosus* shared more allometric trends than any other between-taxa comparisons. Notwithstanding, several statistics were higher in *R. raphanurus*, except for those variables related to temporal muscles and bite. The close relationship between *R. raphanurus* and *L. inca* is also supported by the longitudinal growth of the rostrum, although with a clear growth extension in *R. raphanurus*. The allometric trends reported for *L. inca* reflect a more predaceous condition compared to other caenolestids. Bandicoots and caenolestids did not show a particularly shared growth pattern, with the latter being morphologically more conservative. Ameridelphia was paraphyletic in the shortest tree regarding the optimization of the confidence intervals. However, the growth of several variables supported monophyletic groups in both hypotheses. Skull ontogeny in marsupials is informative in several aspects of the mandible and neurocranium reflecting the high phylogenetic signal displayed by variables related to these cranial regions.

Keywords Caenolestidae · Ontogeny · Growth evolution · Marsupials

Introduction

Extant marsupials exhibit high morphological variation in skull morphology (e.g., Wroe and Milne 2007; Goswami et al. 2011, 2012; Bennett and Goswami 2013). Recent comparative studies in marsupials and placentals (e.g., Wilson and Sánchez-Villagra 2010; Wilson 2013) suggest that the former show less disparity in the general morphology of the skull, but higher integration, mostly in the oral region (e.g., Shirai and

Marroig 2010; Porto et al. 2013). In marsupials, the rostral region is initially constrained by its functional compromise from attachment for suckling during early stages of development (Bennett and Goswami 2013). Such differences are probably caused by intrinsic factors (especially the mode of marsupial reproduction and development, Goswami et al. 2012) and/or extrinsic ones (history of diversification and biogeography, Sánchez-Villagra 2013). However, in spite of the lower developmental disparity reported for marsupials in comparison with placentals, the former group exhibits high morphological diversity including both specialized (e.g., strictly herbivorous, nectarivorous, or carnivorous groups) as well as generalized forms showing morphological convergences on skull shape (see Wroe and Milne 2007 for carnivores).

The Australasian bandicoots (Peramelidae) and the South American shrew opossums (Caenolestidae), exhibit similar skull features such as a greatly elongated rostrum and large palatine vacuities, which distinguish both from other extant marsupials. This resemblance between bandicoots and shrew opossums, which is apparent in spite of well-marked differences in body size (e.g., Osgood 1921; Retief et al. 1995; Krajewski et al. 1997; Palma and Spotorno 1999), represented the basis for old hypotheses of a close relationships between both groups (see Osgood 1921; Gregory 1922). Several studies have placed caenolestids in different relationships with other marsupials (Beck 2008; Meredith et al. 2008). They have been recovered as the sister group to Didelphimorphia within Ameridelphia (Szalay 1982, 1994; Gemmell and Westerman 1994; Retief et al. 1995; Colgan 1999); sister group to the entire Australian marsupial radiation and separate from Ameridelphia (Springer et al. 1998; Amrine-Madsen et al. 2003; Horovitz and Sánchez-Villagra 2003; Asher et al. 2004; Cardillo et al. 2004; Nilsson et al. 2004; Munemasa et al. 2006; Beck 2008; Beck et al. 2008); sister to both Didelphimorphia and all Australian marsupials (Kirsch et al. 1991; Szalay and Sargis 2001); or as the sister group to Peramelemorphia and different from the rest of all marsupials (Palma and Spotorno 1999). Another recent phylogeny placed the caenolestids in a position closely related to bandicoots (Lavedèze and Muizon 2010). Despite this conflicting phylogenetic information, caenolestids exhibit a plesiomorphic tarsal morphology similar to that of other New World marsupials (Szalay 1982, 1994), paired spermatozoa as in most New World marsupials, albeit with unique characteristics (Biggers and DeLamater 1965; Temple-Smith 1987), enlarged and forwardly projected first lower incisors (Marshall 1980; Patterson 2008), large antorbital vacuities (Marshall 1980; Patterson and Gallardo 1987), and a shrew-like external appearance (Palma 2003). The pair of large, forwardly directed incisors in the lower jaw resemble those of Diprotodontia (Osgood 1924), but they are apparently homologous to i2 in other marsupials and not to

i1 (Hershkovitz 1995). In turn, bandicoots possess a tarsal morphology typical of Australasian marsupials (Szalay 1982), unpaired spermatozoa (Tyndale-Biscoe 2005), and a chorioallantoic placenta (Hughes et al. 1990; Tyndale-Biscoe 2005), which is functionally related to a fast rate of development in this group (Gordon and Hulbert 1989).

In the balance of all evidence, the two long-nosed clades of living marsupials appear to be convergent and not closely related phylogenetically. Caenolestids are the only extant representatives of the once diverse order Paucituberculata (Abello 2007; Patterson 2008), which includes several specialized forms that had their richness peak during the middle Tertiary (Abello 2007, 2013), convergent with other “pseudodiprotodont” fossils such as Polydolopimorphia (Goin et al. 2003).

Regarding bandicoots, until recently the scarce ontogenetic data published were limited to aging of specimens on the basis of tooth eruption (e.g., Kingsmill 1962; Lyne 1964; Lyne and Mort 1981). Recently, Travouillon et al. (2014) described some cranial ontogenetic changes in the fossil *Madju varia*, and Flores et al. (2013) quantified the pattern of skull growth in *Echymipera kalubu* and *Isoodon macrourus*. This study showed that positive allometries apply to longitudinal dimensions in *E. kalubu*, whereas positive allometries are restricted to vertical dimensions of the skull in *I. macrourus*. By contrast, information on ontogeny is lacking in caenolestids, beyond a reported slow rate of development for the group (Tyndale-Biscoe 2005), and just a few studies dealing with dental homologies and tooth replacement / eruption (e.g., Luckett and Hong 2000; Martin 2007, 2013). Breeding data are restricted to anecdotal records (see Patterson and Gallardo 1987).

Extant caenolestids represent excellent candidates for comparative studies of skull growth with convergent Australasian forms, the bandicoots, and with other marsupials, given the wealth of recent studies on the allometry of skull growth in marsupials from several groups from both the Australasian and American radiations (e.g., Abdala et al. 2001; Flores et al. 2003, 2006, 2010, 2013; Giannini et al. 2004). Flores et al. (2013) compared the allometric growth pattern of two bandicoots with previously studied marsupials and placentals (three didelphids, two australidelphians, two primates, and two carnivorans). The two bandicoots showed ontogenetic trajectories that differed in several measurements, suggesting a clear divergence in the growth patterns between them. However, and in agreement with those studies that suggest a low morphological disparity in the neurocranium of marsupials (e.g., Bennett and Goswami 2013), the allometric trends of sensorial capsules scaled mostly with negative allometry in all groups (Flores et al. 2013). The higher degree of integration of the viscerocranium in marsupials (Bennett and Goswami 2013) is in general agreement with the stability of the negative allometry of the palate width and the positive

allometry of the rostral length, across all previously studied species (see Flores et al. 2013).

In this work we report on the skull growth of one representative species per extant genus of caenolestids: *Caenolestes fuliginosus*, *Lestoros inca*, and *Rhyncholestes raphanurus*. We integrated our results with previously published data on cranial ontogeny of marsupials, particularly the convergent bandicoots, and established a phylogenetic framework of comparison to place the particulars of the caenolestid cranial ontogeny in the broader context of skull evolution in metatherians.

Materials and Methods

The Group Extant caenolestids comprise only three recent genera, *Caenolestes*, *Lestoros*, and *Rhyncholestes* (the first one polytypic) known only from South America (Cabrera 1957; Patterson 2008). Their broadly relictual and disjunct distribution (Patterson 2008) suggests a wider distribution and higher diversity of extinct taxa, which is supported by its substantial fossil record (see Abello 2007; Goin et al. 2007, 2009). In this study we included representatives of the three genera. *Caenolestes fuliginosus* is distributed in the Páramo (i.e., the Neotropical high mountain biome with a vegetation composed mainly of giant rosette plants, shrubs, and grasses; Luteyn 1999) of Colombia and Ecuador and is the species of *Caenolestes* best represented in systematic collections (Brown 2004; Patterson 2008; Solari et al. 2013). *Lestoros inca* inhabits wet, mossy elfin forests, as well as highly disturbed scrub and second-growth forests of southern Perú (Myers and Patton 2008; Medina et al. 2012). *Rhyncholestes raphanurus* is an extremely rare species that has been collected only in the temperate Valdivian rain forests of southern Chile and adjacent Argentina (Pearson 1995; Birney et al. 1996; Martin 2011), separated from the northern caenolestids by the arid diagonal of South America (Bruniard 1982). Although caenolestids exhibit an old evolutionary history in South America, and high species richness in the past (Abello 2007), their extant representatives show a very low diversity when compared to other marsupial groups including the sympatric didelphids (Gardner 2005; Patterson 2008). The extant species exhibit a relatively conservative skull morphology as well as little variation in body size (Patterson 2008). Old and recent evidence of development in shrew opossums only covered patterns of tooth replacement, function, and wear, which revealed a consistent pattern of tooth use and so important inferences on food habits and behavior were made in this poorly known group (Lockett and Hong 2000; Martin 2007, 2011, 2013). The principal aspects of reproduction in shrew opossums remain poorly understood, and the scarcity of young specimens in collections represents a serious limitation for ontogenetic studies. Extant caenolestids are very difficult

to capture in the field and almost impossible to study alive, given that no individual has survived for more than a few days in captivity (Patterson and Gallardo 1987; Tyndale-Biscoe 2005). In addition, capture of a female with attached young has never been reported. Knowledge about reproduction and breeding in the group is based upon scattered field data (e.g., Kirsch and Waller 1979; Patterson and Gallardo 1987; Tyndale-Biscoe and Renfree 1987), which suggested a pattern of seasonal breeding, a single reproductive event per year, small litter size, and a very slow rate of development (Tyndale-Biscoe 2005).

Sample We studied aged skull series of species representative of the three extant genera in the Family Caenolestidae deposited in the mammal collections of the Universidad de Antioquia (CTUA), Field Museum of Natural History (FMNH), Instituto de Ciencias Naturales, Universidad Nacional de Colombia (ICN), and United States National Museum, Smithsonian Institution (NMNH): *Caenolestes fuliginosus* ($n=119$; 49 females, 70 males), *Lestoros inca* ($n=62$; 31 females, 31 males), and *Rhyncholestes raphanurus* ($n=26$; 13 females, 13 males; [Supplementary Information](#)). The lack of specimens with deciduous dentitions in systematic collections worldwide limited our sample to subadult and adult specimens, defined as individuals with emerging P3/p3, and individuals with fully erupted dentition, respectively (Lockett and Hong 2000). As discussed by Lockett and Hong (2000) and Martin (2013), the retention of a late erupting P3, associated with the derived condition of a greatly reduced dP3 that probably does not erupt, is a combination unique to caenolestids. However, it is still possible to establish a relationship between size and age. The larger specimens in the three species exhibit advanced degrees of molar cusp wear. For instance, in *L. inca* a small cusp anterolingual to stylar cusp B (apparently the paracone, see Goin et al., 2007) forms a small socket increasing in size from M1 to M3, which is highly susceptible to wear and can only be observed in young specimens (Martin 2013). Other dental characters for subadult specimens of *C. convelatus* and *C. fuliginosus* were described by Lockett and Hong (2000). The youngest specimen of *C. fuliginosus* (FMNH 70858, condylo-inscive length [CIL] 25.24 mm) exhibited the lower third premolar in the process of eruption (see Lockett and Hong 2000: fig. 9) whereas our largest specimen (ICN-PSP1247, unknown sex, CIL 33.85) presented heavily worn molar cusps. Additionally, seven specimens (FMNH 70832, 70888, 70859, ICN 16534-6, 11654) showed incompletely erupted third lower premolars. Our youngest specimen of *L. inca* (FMNH 172048, male, CIL 25.25) showed its fourth upper molar barely emerging above the alveolar line, and its third upper premolar not fully erupted with its main cusp still not reaching the height of the stylar cusp B of the next molar (lateral view; see Martin, 2013). Two additional specimens (FMNH 75113, USNM 194407) were of

small size, showing similar dental conditions. By contrast, our larger specimen (FMNH 75115, male, CIL 30.75) presented heavily worn molar cusps. Finally, the youngest specimen in our sample of *R. raphanurus* (FMNH 129828, female, CIL 28.8) exhibited an adult dentition with little cusp wear, whereas our largest specimen (FMNH 50071, male, CIL 34.78) showed heavily worn molar cusps.

Sexual dimorphism was detected in *L. inca* and *C. fuliginosus* involving size, whereas shape dimorphism was detected in *C. fuliginosus* and partially in *L. inca* (Astúa 2010; Martin 2013). The study of Astúa (2010) did not find evidence for sexual dimorphism in size or shape for *R. raphanurus*. Despite sexual variation detected in *L. inca* and *C. fuliginosus*, both males and females clearly shared the same ontogenetic trajectory in the three species (i.e., non-significant slope or intercept differences in bivariate analyses of selected variables; see below). Therefore, within each species we pooled male and female specimens of all ages in a single sample per species, as in our recent study on bandicoots (Flores et al. 2013).

Study of Growth Following our previous studies on marsupials (Abdala et al. 2001; Flores et al. 2003, 2006, 2010, 2013; Giannini et al. 2004), we used 15 cranial measurements (Fig. 1) to estimate allometric growth in caenolestids. We used bivariate and multivariate analyses to estimate growth trends. In bivariate analyses, the scaling of any measurement can be affected by the choice of the independent variable (Smith 1981; Wayne 1986). Because our multivariate analyses indicated that the total length of the skull, or the condylo-incisive length (CIL), was isometric, this measurement was taken as a reliable indicator of size variation and therefore it was used as an independent variable in bivariate allometry. We used linear (log₁₀) transformation of the power equation of growth for each variable (see Abdala et al. 2001 for details) and tested deviations from isometry by means of 2-tailed t-tests, after corroborating that the independent variable was normally distributed (Shapiro–Wilk test, *C. fuliginosus*: $w=0.981$, $P=0.466$; *L. inca*: $w=0.958$, $P=0.44$; *R. raphanurus*: $w=0.987$, $P=0.987$). Statistically significant deviations from isometry represent cases of “negative” allometry if the slope $b < 1.0$ and “positive” allometry if $b > 1.0$. We applied standardized major axis regression (SMA) and followed Warton et al. (2006) in using a likelihood ratio test for the common SMA slope using a chi-squared distribution (Warton and Weber 2002). If the species shared a common slope, we compared the significance of the common normalization constant (y-intercepts) using the Wald test (as described in Warton et al. 2006). All these regression coefficients, statistical parameters, and tests were performed using the SMATR package of R (Warton and Weber 2002). We statistically compared the parameters of the trajectories among the three species.

Multivariate coefficients of allometry were obtained from the first (unit-scaled) eigenvector of a principal component analysis (PCA) based on a variance–covariance matrix of log₁₀ transformed data (Jolicoeur 1963). We tested the significance of multivariate coefficients of allometry with a resampling strategy based on the jackknife, implemented in such a way that each specimen was removed from the sample at a time, generating n pseudovalues to calculate confidence intervals (CIs) for the original coefficients (see Giannini et al. 2004 and Flores et al. 2006 for further details on this method). If the interval excluded an expected value of isometry, the variable was considered positively or negatively allometric. For all multivariate coefficients of allometry, the expected value of isometry, which depends only on the number of variables (p), is calculated as $1/p^{0.5}$ (0.258 for our set of 15 variables). Trimming the largest and smallest m pseudovalues (with $m=1$) for each variable may significantly decrease the standard deviations calculated under jackknife and allow for more accurate allometric estimations (Manly 1997; Giannini et al. 2004). Here we report untrimmed and trimmed calculations, but opting for the results with either lower average standard deviation or lower bias (with the latter defined as the difference between the observed and jackknifed allometry coefficient; Giannini et al. 2004). For the multivariate statistical analyses (PCA + jackknife resampling), we used the R script of Giannini et al. (2010; available from the authors).

Comparative Analysis Here we followed the quantitative approach of character optimization and phylogenetic signal that we used in our recent report on bandicoots (Flores et al. 2013), and integrated the results obtained herein for extant caenolestids with a taxonomic sample of seven additional marsupials (see below) and four placentals included as outgroup (*Lycalopex culpaeus*, *Puma concolor*, *Cebus apella*, and *Alouatta caraya*). We used the same dataset and included a re-analysis under multivariate allometry of the ontogenetic raw data of *Didelphis albiventris*, *Lutreolina crassicaudata*, and *Dromiciops gliroides* (previously reported using bivariate allometry), as well as placental outgroups (see Flores et al. 2013), considering the exact same 15 cranial variables as in caenolestids and bandicoots (Fig. 1).

Optimization is a tool used to determine the cost of a phylogenetic topology and ancestral character states; this way of mapping evolutionary information onto phylogenetic trees represents one of the essentials of comparative biology (see discussion in Giannini and Goloboff 2010). Here we followed Giannini (2014) who demonstrated that allometry should be mapped using the estimated confidence interval as a continuous character (see Giannini 2014 and Segura 2014 for applications in rodents and carnivorans, respectively). Optimization of continuous characters using either point estimates or intervals for terminal values is implemented in the computer program TNT (Goloboff et al. 2008) as an extension

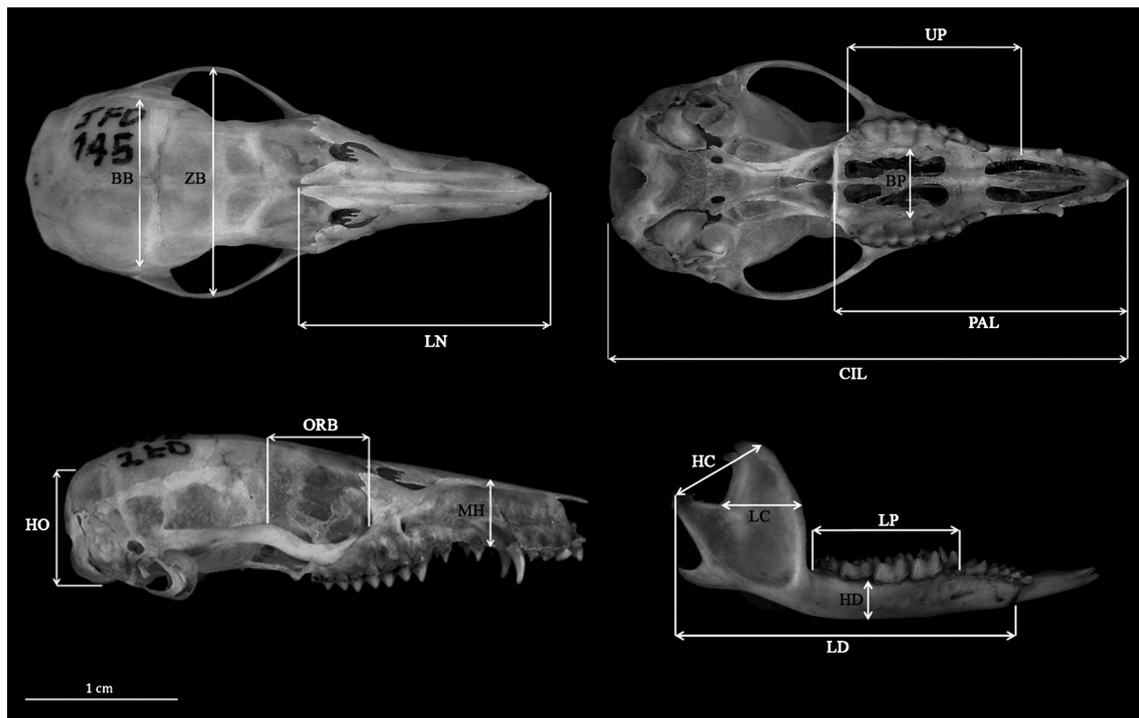


Fig. 1 Skull and mandible measurements considered in this study. *Abbreviations:* CIL Condylo-incipive length, BB Breadth of the braincase, BP Breadth of the palate, HC Height of the coronoid process, HD Height of the dentary, HO Height of the occipital plate, LC Length of the

coronoid process, LD Length of the dentary, LN Length of the nasals, LP Lower postcanine tooththrow, MH Muzzle height, ORB Orbit length, PAL Palate length, UP Upper postcanine tooththrow, ZB Zygomatic breadth

of Farris' (1970) multistate character optimization (Goloboff et al. 2006). Thus, allometric similarities can be explained by common ancestry and ancestral allometry vectors can be optimally estimated (see Giannini 2014 and citations therein). We reconstructed the allometric vector at all internal nodes, calculated the optimal cost of reconstruction (steps on the tree), and examined the evolutionary changes (increases and decreases in individual allometric coefficients) as implied by the tree topology, to uncover the general phylogenetic pattern of cranial ontogeny evolution and its functional implications.

We included the didelphids *Didelphis albiventris*, *Lutreolina crassicaudata*, and *Caluromys philander*, the microbiotheriid *Dromiciops gliroides*, the dasyurid *Dasyurus albopunctatus*, the peramelid *Echimypera kalubu* and *Isoodon macrourus*, and the caenolestids studied herein, and four placental outgroups, arranged in two possible topologies. The first one (called H1 hereafter) considered Ameridelphia as monophyletic (i.e., caenolestids sister to Didelphidae; Szalay 1982; Beck 2008). The second topology (called H2 hereafter) considered Ameridelphia as paraphyletic (i.e., caenolestids were sister to Australidelphia; Horovitz and Sánchez-Villagra 2003; Asher et al. 2004; Nilsson et al. 2004; Beck et al. 2008; Meredith et al. 2008). In both topologies, *Dromiciops* was placed as sister to the remaining australidelphian marsupials (Meredith et al. 2008).

Phylogenetic signal was obtained using random swapping of characters (allometric coefficients) in terminal branches (Laurin 2004). The distribution of character states (i.e., resampled allometric coefficients) on the observed phylogenies was compared with the distribution of states on 20,000 randomly generated trees. The number of evolutionary steps of each character on the randomly generated trees was also compared with the number of steps of that character in the observed phylogenies. We considered a character to have significant phylogenetic signal if the observed number of steps was larger than the steps calculated for at least 95 % of the randomly generated trees. For phylogenetic signal calculation, we used the R script of Prevosti et al. (2010), computed with the program TNT v. 1.1 (Goloboff et al. 2008).

Results

Cranial Allometry in *Caenolestes fuliginosus*

Bivariate Analysis All variables were correlated with the total length of the skull; variables with $r^2 < 0.7$ included UP (upper postcanine length; $r^2 = 0.63$); ORB (orbital length; $r^2 = 0.41$); BB (breadth of the braincase; $r^2 = 0.31$); HO (height of the occipital plate; $r^2 = 0.38$); LC (length of the coronoid process; $r^2 = 0.52$);

HD (height of the dentary; $r^2=0.54$); and LP (lower postcanine length; $r^2=0.64$) (Table 1). UP, LD (length of the dentary), and LP were the only isometric variables (Table 1). Positive allometry was observed in ZB (zygomatic breadth); HM (height of the muzzle), PAL (length of the palate), ORB, LN (length of the nasals), HC (height of the coronoid process), LC, and HD. Three variables scaled negatively with condylo-incisive length (CIL), from both the neurocranium and the splanchnocranium: BB, HO, and BP (breadth of the palate). The latter exhibited the slowest growth rate ($b_{\text{ma}}=0.64$).

Multivariate Analysis The first principal component explained 64.46 % of the total variation. The multivariate approach showed a pattern similar to that of the bivariate analysis, although with weaker trends in five variables (Table 2). Indeed, the only differences among multivariate (untrimmed, see below) and bivariate approaches were limited to the negative allometry of LP and UP, and the isometry of ZB, PAL, and HC. Remarkably, CIL and LD (the two variables representing overall length of the skull and mandible, respectively) departed less from the point estimate of isometry (between -0.005 and -0.004), while BP, BB, HO, and LP departed the most (between -0.12 and -0.18).

Cranial Allometry in *Lestoros inca*

Bivariate Analysis Variables were not strongly correlated with the total length of the skull, with $r^2>0.7$ in just two variables (LN, and LD). The lowest correlations were detected in ORB ($r^2=0.14$), LP ($r^2=0.22$), HD ($r^2=0.32$), LC ($r^2=0.06$), HO ($r^2=0.05$) and BB ($r^2=0.31$; Table 1). Variables showing isometry with respect to the condylo-incisive length included measurements from both neurocranium and splanchnocranium (i.e., ZB, BP, BB, HO, LD, HC, and LC). Positive allometry was observed in seven variables: HM, PAL, UP, ORB, LN, HD, and LP. Finally, no variables scaled negatively in *L. inca*, although BP exhibited the slowest growth rate ($b_{\text{ma}}=0.89$).

Multivariate Analysis The first principal component explained 49.81 % of the total variation. The smallest departure from isometry was PAL (-0.028) and the largest departure was ORB (0.44). Multivariate analysis showed several different trends as compared with its bivariate counterpart, as only three variables (ORB, LD, and HC) shared allometric trends (Table 2). In a general sense, the allometric coefficients observed under multivariate analysis were lower than observed under bivariate analysis, that is, a scale shift was apparent: those variables with positive trends in bivariate analysis showed isometry in multivariate analysis (i.e., HM, PAL, UP, LN, HD, and LP), whereas those variables that were isometric under bivariate allometry, became negatively allometric under multivariate analysis (i.e. ZB, BP, BB, OH, and

LC). Finally, only ORB conserved its positive allometry also in this analysis.

Cranial Allometry in *Rhyncholestes raphanurus*

Bivariate Analysis All variables were well correlated with CIL, with $r^2>0.5$, with the lowest correlation obtained in HD ($r^2=0.51$) and LP ($r^2=0.57$), and the highest in LD ($r^2=0.92$). Eight variables were isometric with respect to CIL, most of which were related with trophic function (ZB, PAL, BP, UP, LN, LD, and HC) plus one neurocranial variable (BB). Positive allometry was detected in three splanchnocranial (HM, LC, and HD) and one neurocranial (ORB) variables. Only two variables were negatively allometric, also belonging to both cranial regions (LP and HO).

Multivariate Analysis The first component explained 75.29 % of total variation. The lowest departure from isometry was in MH (0.029). These results differed from the bivariate analysis in five variables (HM, HD, UP, ORB, and BB; Table 2). As in *L. inca*, the allometric coefficients observed under multivariate analysis were lower than those obtained with bivariate analysis. Most variables with positive trends in bivariate analysis were isometric in multivariate analysis (i.e., HM, ORB, and HD), whereas some isometric variables under bivariate allometry, became negatively allometric in multivariate analysis (i.e., UP and BB). Finally, several variables conserved their allometric trend in both analyses (i.e., positively allometric in LC; negatively allometric in HO and LP, and isometric in ZB, PAL, BP, LN, LD, and HC).

Comparative Allometry of Bivariate Results in Extant Caenolestids

In all multivariate analyses the condylo-incisive length (CIL) was recovered as isometric, for which our choice of it as independent variable in bivariate analyses was supported. *Caenolestes fuliginosus* and *L. inca* shared eight out of 14 allometric trends (i.e., ZB, HM, PAL, BB, LN, HO, LD, and HC; Table 3; Fig. 2a). Statistical differences in slopes between both species were found in the remaining six variables (i.e., BP, UP, ORB, LC, HD, and LP; Fig. 2b); in all but one case (LC), slopes were lower in *C. fuliginosus*. *Y*-intercepts were statistically different in those variables sharing the same slope (i.e., ZB, HM, PAL, BB, LN, HO, and HC) except in LD, and those of *L. inca* were lower except in ZB and HC.

Comparisons of bivariate analyses between *L. inca* and *R. raphanurus* (Table 3) indicate that both species shared slopes in eight out of 14 variables (i.e., ZB, PAL, BP, UP, BB, LN, LD, and HC; Fig. 2b). Among these eight variables, the *y*-intercept was higher in *L. inca* just in ZB and BP. In LN, BB, and LD, the intercept was higher

Table 1 Summary of bivariate regressions for skull elements of *Caenolestes fuliginosus*, *Lestoros inca*, and *Rhyncholestes raphanurus*, using condylo-insinive length as proxy of size (independent variable, see text for details)

Variable	Species	Range (mm)	R ²	B _{sma}	T _{iso}	P _{iso}	Y-intercept	Allometric trend
ZB	<i>Caenolestes fuliginosus</i>	11.9–17.5	0.81	1.11	-7.45	0.01	-0.49	+
	<i>Rhyncholestes raphanurus</i>	12.2–15	0.71	1.08	-2.66	0.50	-0.51	=
	<i>Lestoros inca</i>	12.2–15.1	0.62	1.05	-3.21	0.54	-0.39	=
HM	<i>Caenolestes fuliginosus</i>	3.5–5.1	0.70	1.60	-14.73	0.00	-1.78	+
	<i>Rhyncholestes raphanurus</i>	2.4–3.1	0.78	1.37	-7.60	0.01	-1.61	+
	<i>Lestoros inca</i>	3–4.3	0.36	1.86	-7.57	0.00	-2.12	+
PAL	<i>Caenolestes fuliginosus</i>	13.1–18.7	0.93	1.10	-10.30	0.00	-0.42	+
	<i>Rhyncholestes raphanurus</i>	16.7–20.3	0.87	1.09	-2.92	0.29	-0.37	=
	<i>Lestoros inca</i>	13.7–17.8	0.55	1.27	-4.02	0.01	-0.65	+
BP	<i>Caenolestes fuliginosus</i>	6.7–8.2	0.22	0.64	-1.03	0.00	-0.08	-
	<i>Rhyncholestes raphanurus</i>	6.2–7.7	0.65	0.98	-3.35	0.89	-0.64	=
	<i>Lestoros inca</i>	6.7–8.1	0.33	0.89	-3.05	0.25	-0.42	=
UP	<i>Caenolestes fuliginosus</i>	8.2–12	0.63	0.90	-4.29	0.06	-0.32	=
	<i>Rhyncholestes raphanurus</i>	9.2–11	0.80	1.00	-3.40	0.97	-0.51	=
	<i>Lestoros inca</i>	8.1–10	0.44	1.23	-4.85	0.03	-0.85	+
ORB	<i>Caenolestes fuliginosus</i>	2.2–4.2	0.41	1.94	-11.83	0.00	-2.40	+
	<i>Rhyncholestes raphanurus</i>	3.3–4.5	0.64	1.85	-6.00	0.00	-2.19	+
	<i>Lestoros inca</i>	2.9–5.3	0.14	3.48	-7.34	0.00	-4.46	+
BB	<i>Caenolestes fuliginosus</i>	9.8–12.3	0.31	0.84	-2.11	0.02	-0.20	-
	<i>Rhyncholestes raphanurus</i>	9–10.7	0.64	0.77	-1.04	0.05	-0.16	=
	<i>Lestoros inca</i>	9.7–12	0.31	1.00	-2.69	0.98	-0.42	=
LN	<i>Caenolestes fuliginosus</i>	11.1–17.3	0.84	1.31	-10.85	0.00	-0.77	+
	<i>Rhyncholestes raphanurus</i>	12.7–18.5	0.65	1.15	-2.27	0.28	-0.51	=
	<i>Lestoros inca</i>	11.4–15.7	0.80	1.45	-7.95	0.00	-0.96	+
HO	<i>Caenolestes fuliginosus</i>	6.6–8.4	0.38	0.79	-3.37	0.00	-0.28	-
	<i>Rhyncholestes raphanurus</i>	7.2–8	0.62	0.60	-0.19	0.00	-0.02	-
	<i>Lestoros inca</i>	7.5–9	0.05	1.01	-3.07	0.95	-0.57	=
LD	<i>Caenolestes fuliginosus</i>	16.2–22.1	0.90	1.03	-5.47	0.32	-0.24	=
	<i>Rhyncholestes raphanurus</i>	17.9–21.4	0.92	1.06	-3.21	0.31	-0.31	=
	<i>Lestoros inca</i>	15.7–19.6	0.72	1.08	-2.84	0.29	-0.32	=
HC	<i>Caenolestes fuliginosus</i>	5.8–8.5	0.73	1.23	-11.17	0.00	-0.98	+
	<i>Rhyncholestes raphanurus</i>	6.2–8	0.72	1.14	-4.41	0.25	-0.88	=
	<i>Lestoros inca</i>	6–7.9	0.49	1.19	-5.59	0.06	-0.89	=
LC	<i>Caenolestes fuliginosus</i>	2.4–4.5	0.52	1.70	-12.24	0.00	-1.97	+
	<i>Rhyncholestes raphanurus</i>	2.5–4.5	0.67	2.81	-6.96	0.00	-3.70	+
	<i>Lestoros inca</i>	7.5–9	0.06	1.01	-3.07	0.95	-0.57	=
HD	<i>Caenolestes fuliginosus</i>	1.7–2.9	0.54	1.47	-13.37	0.00	-1.82	+
	<i>Rhyncholestes raphanurus</i>	1.9–2.7	0.51	1.52	-4.71	0.03	-1.94	+
	<i>Lestoros inca</i>	1.8–2.9	0.32	2.36	-8.42	0.00	-3.08	+
LP	<i>Caenolestes fuliginosus</i>	6.2–10	0.64	1.10	-4.97	0.29	-0.73	=
	<i>Rhyncholestes raphanurus</i>	7.5–8.6	0.57	0.61	-0.02	0.00	0.00	-
	<i>Lestoros inca</i>	6–8.3	0.22	2.11	-6.36	0.00	-2.23	+

Abbreviations (variable acronyms) as in Fig. 1. R² Adjusted coefficient of correlation, b_{sma} Allometric coefficient in reduced major axis analysis, T_{iso} 2-tailed Student's t-value under the assumption of isometry (expected allometric coefficient for isometry = 1. n-2 d.f.), P_{iso} P-value for T_{iso}. = isometry; +, positive allometry; -, negative allometry

in *R. raphanurus*. Both slope and intercept were shared in PAL (Fig. 2a) and UP, but with an extended trajectory in

R. raphanurus. The remaining six variables (i.e., HM, ORB, HO, LC, HD, and LP) showed statistical differences

Table 2 Results of the multivariate analysis of cranial allometry in *Caenolestes fuliginosus* (n=119), *Lestoros inca* (n=62), and *Rhyncholestes raphanurus* (n=26)

Variable	Species	Observed	Departure	Untrimmed			Trend	Trimmed			Trend
				Resampled	Bias	99%CI		Resampled	Bias	99%CI	
CIL	<i>Caenolestes fuliginosus</i>	0.251	-0.007	0.253	-0.0007	0.237-0.268	=	0.252	-0.0006	0.238-0.266	=
	<i>Rhyncholestes raphanurus</i>	0.215	-0.043	0.213	0.0010	0.161-0.263	=	0.229	-0.0071	0.201-0.256	-
	<i>Lestoros inca</i>	0.158	-0.1	0.204	0.0232	0.057-0.35	=	0.121	0.0182	0.054-0.188	-
ZB	<i>Caenolestes fuliginosus</i>	0.266	0.008	0.267	-0.0007	0.236-0.297	=	0.264	0.0006	0.237-0.291	=
	<i>Rhyncholestes raphanurus</i>	0.218	-0.04	0.220	-0.0009	0.103-0.336	=	0.226	-0.0039	0.165-0.286	=
	<i>Lestoros inca</i>	0.149	-0.109	0.190	0.0206	0.051-0.228	-	0.112	0.0181	0.041-0.183	-
HM	<i>Caenolestes fuliginosus</i>	0.351	0.093	0.352	-0.0006	0.289-0.414	+	0.353	-0.0010	0.295-0.41	+
	<i>Rhyncholestes raphanurus</i>	0.287	0.029	0.287	-0.0001	0.223-0.35	=	0.294	-0.0038	0.245-0.342	=
	<i>Lestoros inca</i>	0.301	0.043	0.359	0.0294	0.177-0.54	=	0.267	0.0165	0.188-0.345	=
PAL	<i>Caenolestes fuliginosus</i>	0.271	0.013	0.272	-0.0009	0.245-0.299	=	0.272	-0.0005	0.247-0.296	=
	<i>Rhyncholestes raphanurus</i>	0.229	-0.029	0.221	0.0042	0.142-0.298	=	0.225	0.0019	0.172-0.278	=
	<i>Lestoros inca</i>	0.231	-0.028	0.258	0.0137	0.181-0.333	=	0.237	0.0031	0.189-0.283	=
BP	<i>Caenolestes fuliginosus</i>	0.090	-0.168	0.091	-0.0007	0.053-0.128	-	0.091	-0.0008	0.056-0.126	-
	<i>Rhyncholestes raphanurus</i>	0.183	-0.075	0.202	-0.0097	0.111-0.293	=	0.190	-0.0033	0.114-0.265	=
	<i>Lestoros inca</i>	0.056	-0.202	0.079	0.0114	0.076-0.234	-	-0.011	0.0338	0.075-0.053	-
UP	<i>Caenolestes fuliginosus</i>	0.178	-0.08	0.179	-0.0005	0.13-0.228	-	0.177	0.0009	0.133-0.219	-
	<i>Rhyncholestes raphanurus</i>	0.197	-0.061	0.190	0.0034	0.128-0.251	-	0.205	-0.0042	0.156-0.253	-
	<i>Lestoros inca</i>	0.184	-0.074	0.208	0.0123	0.134-0.282	=	0.194	0.0052	0.133-0.254	-
ORB	<i>Caenolestes fuliginosus</i>	0.362	0.104	0.363	-0.0006	0.258-0.468	+	0.364	-0.0008	0.266-0.46	+
	<i>Rhyncholestes raphanurus</i>	0.346	0.088	0.333	0.0064	0.166-0.498	=	0.357	-0.0058	0.227-0.487	=
	<i>Lestoros inca</i>	0.701	0.443	0.682	0.0098	0.42-0.942	+	0.808	0.0533	0.673-0.942	+
BB	<i>Caenolestes fuliginosus</i>	0.109	-0.149	0.109	0.0000	0.056-0.161	-	0.118	-0.0045	0.069-0.166	-
	<i>Rhyncholestes raphanurus</i>	0.144	-0.114	0.158	-0.0067	0.09-0.224	-	0.141	0.0014	0.108-0.174	-
	<i>Lestoros inca</i>	0.154	-0.104	0.174	0.0099	0.098-0.249	-	0.157	0.0017	0.109-0.205	-
LN	<i>Caenolestes fuliginosus</i>	0.305	0.047	0.308	-0.0011	0.268-0.346	+	0.302	0.0015	0.268-0.336	+
	<i>Rhyncholestes raphanurus</i>	0.207	-0.051	0.225	-0.0090	0.119-0.329	=	0.186	0.0104	0.135-0.235	-
	<i>Lestoros inca</i>	0.215	-0.043	0.275	0.0301	0.062-0.488	=	0.145	0.0348	0.061-0.229	-
HO	<i>Caenolestes fuliginosus</i>	0.133	-0.125	0.134	-0.0007	0.097-0.171	-	0.138	-0.0024	0.104-0.171	-
	<i>Rhyncholestes raphanurus</i>	0.099	-0.159	0.098	0.0005	0.029-0.165	-	0.111	-0.0063	0.077-0.145	-
	<i>Lestoros inca</i>	0.146	-0.112	0.138	0.0043	0.035-0.239	-	0.162	0.0079	0.093-0.23	-
LD	<i>Caenolestes fuliginosus</i>	0.252	-0.006	0.254	-0.0007	0.23-0.276	=	0.257	-0.0023	0.238-0.275	=
	<i>Rhyncholestes raphanurus</i>	0.217	-0.041	0.209	0.0040	0.146-0.272	=	0.220	-0.0013	0.173-0.266	=
	<i>Lestoros inca</i>	0.133	-0.125	0.180	0.0234	0.008-0.351	=	0.074	0.0294	0.007-0.14	-
HC	<i>Caenolestes fuliginosus</i>	0.289	0.031	0.290	-0.0008	0.252-0.328	=	0.292	-0.0020	0.257-0.327	=
	<i>Rhyncholestes raphanurus</i>	0.221	-0.037	0.240	-0.0095	0.152-0.326	=	0.225	-0.0022	0.163-0.286	=
	<i>Lestoros inca</i>	0.181	-0.077	0.226	0.0225	0.08-0.371	=	0.145	0.0179	0.083-0.206	-
LC	<i>Caenolestes fuliginosus</i>	0.350	0.092	0.354	-0.0019	0.268-0.439	+	0.340	0.0051	0.264-0.414	+
	<i>Rhyncholestes raphanurus</i>	0.589	0.331	0.607	-0.0090	0.469-0.744	+	0.607	-0.0086	0.491-0.721	+
	<i>Lestoros inca</i>	0.146	-0.112	0.138	0.0043	0.035-0.239	-	0.162	0.0079	0.093-0.23	-
HD	<i>Caenolestes fuliginosus</i>	0.317	0.059	0.321	-0.0017	0.262-0.379	+	0.317	0.0004	0.264-0.368	+
	<i>Rhyncholestes raphanurus</i>	0.225	-0.033	0.249	-0.0115	0.043-0.453	=	0.178	0.0238	0.059-0.296	=
	<i>Lestoros inca</i>	0.343	0.085	0.427	0.0419	0.204-0.648	=	0.350	0.0037	0.215-0.485	=
LP	<i>Caenolestes fuliginosus</i>	0.079	-0.18	0.080	-0.0007	0.005-0.156	-	0.077	0.0012	0.005-0.147	-
	<i>Rhyncholestes raphanurus</i>	0.113	-0.145	0.119	-0.0033	0.061-0.177	-	0.103	0.0048	0.069-0.137	-
	<i>Lestoros inca</i>	0.159	-0.099	0.144	0.0798	0.233-0.345	=	0.168	0.0760	0.02-0.356	=

The first two data columns show results using all specimens. The remaining columns show jackknife results calculated with untrimmed and trimmed ($m=1$) sets of pseudovalues (see Material and Methods for details). The allometric coefficient of a variable is the corresponding element of the first (unit) eigenvector per variable. The expected coefficient is the value under the assumption of isometry (0,258 for all variables). The observed coefficient is the value obtained with all specimens included. The resampled coefficient is the value generated by first-order jackknife resampling. Bias is the difference between the resampled and observed coefficients. The jackknifed 99 % confidence interval (CI) is provided; allometric variables are those whose CIs exclude the expected value under the assumption of isometry (0,258)

Abbreviations (variable acronyms) as in Fig. 1. =, isometry; +, positive allometry, -, negative allometry

in slopes, these being higher in most variables for *L. inca* with the exception of LC.

Finally, our comparison of bivariate analysis between *C. fuliginosus* and *R. raphanurus* (Table 3) showed that both species shared slopes in ten out of 14 variables (ZB, HM, PAL, UP, ORB, BB, LN, HO, LD, and HD). Among these variables, six showed higher values of y-intercepts favoring *R. raphanurus* (HM, PAL, ORB, BB, LN, and HO), whereas the remaining four variables (ZB, UP, LD, and HD) favored *C. fuliginosus*. The remaining three variables showed differences in slopes favoring *R. raphanurus* in two variables (BP and LC) and one (LP) favoring *C. fuliginosus*.

Comparison of Ontogenetic Patterns with Extant Bandicoots

No variables showed a particular or unique trend in shrew opossums, which was true also for bandicoots (Flores et al. 2013), given that each trend was also observed in some other marsupial group (Table 4). However, in the comparative context of shrew opossums and bandicoots, two variables showed a somewhat noteworthy distribution of ontogenetic trends. First, the condylo-incisive length (CIL), an important measurement because it is frequently used as proxy of overall size and therefore is a usual independent term in bivariate analyses, was recovered as isometric in both bandicoots and shrew opossums, and also in the carnivorous didelphid *Lutreolina crassicaudata*. Second, the dentary length was recovered as isometric in the three shrew opossum species, but positively allometric in bandicoots (Table 4). The remaining variables showed no common or contrasting pattern between both long-nosed groups, and the allometric trends shared between both groups were actually widely distributed among other marsupial and placental groups, such as trends in neurocranial variables (e.g., HO, BB) or palate breadth (BP; Table 4).

Ontogenetic Character Mapping and Phylogenetic Signal

Tree length for the topology considering Ameridelphia as paraphyletic (i.e., H2) was slightly shorter than the topology where Ameridelphia was monophyletic (5.51 vs. 5.57 steps, Table 5). Both cases exhibited phylogenetic signal, although statistically significant in just four variables. For both hypotheses, significant phylogenetic signal was present in two variables related with the neurocranium (BB and HO), and two related with mandibular growth (LD and HC; Table 5). A marginal significance was observed for ZB in H2 (i.e., where Ameridelphia is paraphyletic; Table 5).

Our character mapping of ontogenetic trends as continuous characters showed that most changes concentrated in terminals. However, we also recovered phylogenetic adjustment in the trend of several variables on internal nodes in both systematic hypotheses (H1 and H2; see Material and Methods section). For instance, caenolestids as a group were supported

by two ontogenetic changes in both hypotheses (Figs. 3 and 4): a decrease in the ontogenetic interval of the breadth of the palate (BP) and length of the dentary (LD). Two additional characters supported caenolestids depending on their position in the marsupial tree: when Ameridelphia was considered paraphyletic (Fig. 4) we observed an increase of the orbit interval (ORB), but when caenolestids were placed as sister group to Australidelphia (Fig. 3) we also detected a decrease in growth rate of the length of the palate (PAL). The proposed sister relationship between *R. raphanurus* and *L. inca* (Ojala-Barbour et al. 2013) was supported herein by a decrease in the intervals of height of the coronoid process (HC) and in length of the nasal (LN) for both topologies (Figs. 3 and 4). Didelphids were supported by an increase in growth rate of the height of the dentary (HD), height of the occipital plate (HO), and upper postcanine toothrow (UP) for both hypotheses, whereas an increase in growth rates of the condylo-incisive length (CIL) and zygomatic breadth (ZB) were recovered as a synapomorphy for the group in the topology where Ameridelphia is monophyletic (Fig. 3). Bandicoots (Peramelidae) were supported by an increase in HO and PAL growth rate in both hypotheses, and a rate increase in CIL in H2 (Fig. 4). The clade Ameridelphia as recognized in H1 (Fig. 3) was supported by rate changes in two variables: increase of growth rate in breadth of the braincase (BB) and decrease in length of the coronoid process (LC). The clade Australidelphia was supported by only one change in growth rate affecting the mandible in each hypothesis: an increasing rate in LC in H2, and a decreasing rate in HC in H1 (Figs. 2 and 3). Finally, the clade including Caenolestidae + Australidelphia recognized in H1 was supported by three changes, all related to the trophic apparatus, with a decreasing rate of growth in BP, ZB, and HC (Fig. 4).

Discussion

Ontogenetic Patterns of Cranial Growth in Shrew Opossums

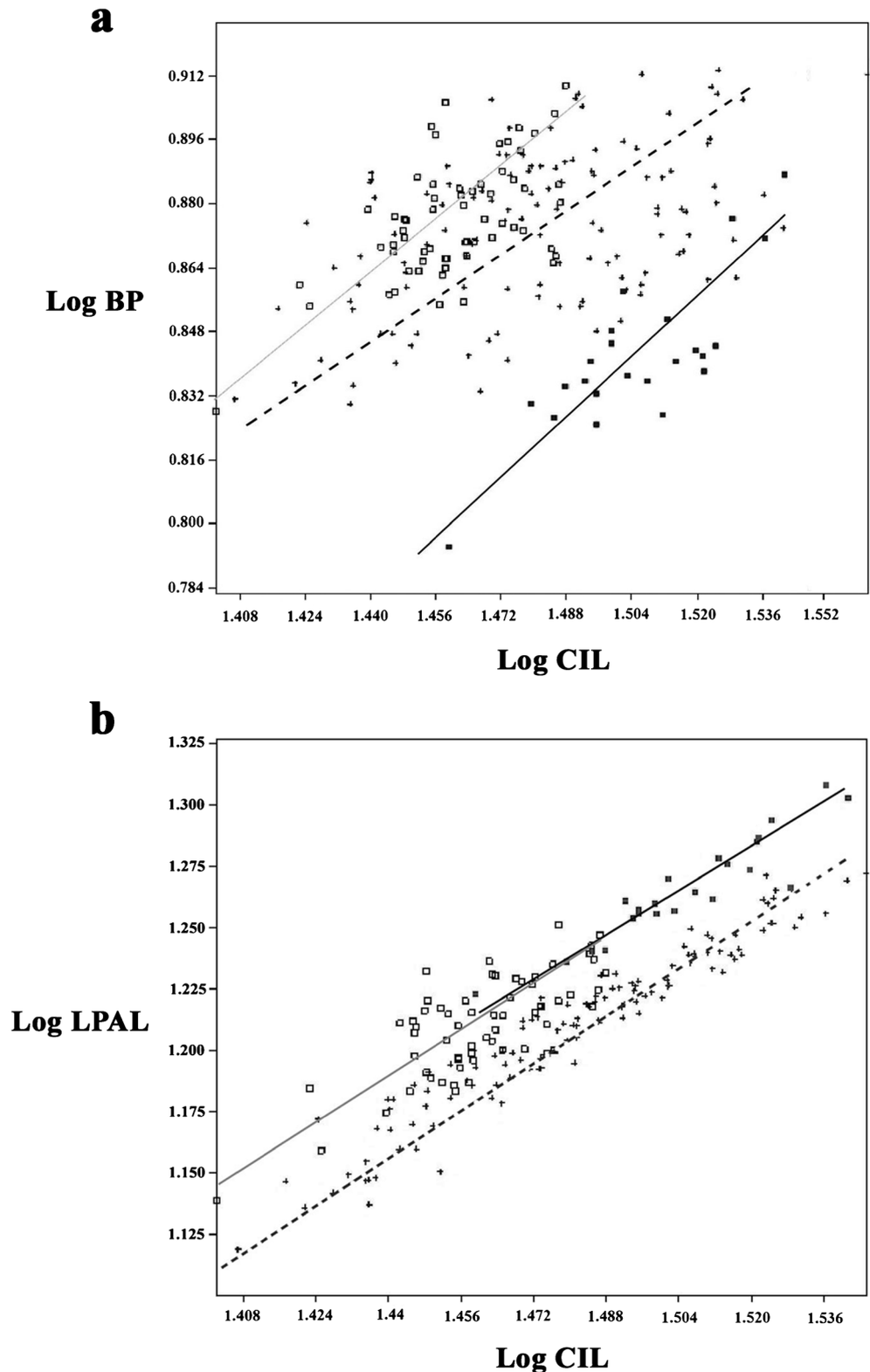
Our comparison of ontogenetic trends among the three extant caenolestid groups shed light onto the evolutionary and functional aspects of skull development in the group. Our ontogenetic evidence supports the clear distinction between *Lestoros* and *Caenolestes*, formerly suggested as synonyms (Bublitz 1987) but now considered different genera as firmly demonstrated in several studies (Myers and Patton 2008; Timm and Patterson 2008; Martin 2013; Ojala-Barbour et al. 2013). In fact, we observed more allometric trends shared between *R. raphanurus* and *C. fuliginosus* (ten out of 14 variables; Table 3) than between *R. raphanurus* and *L. inca*, which were recovered as sister taxa (Ojala-Barbour et al. 2013). However, *R. raphanurus* and *L. inca* shared the same slope and intercept

Table 3 Test for common slopes and common intercepts for the caenolestid taxa *Caenolestes fuliginosus* (C.f.), *Lestoros inca* (L.i.), and *Rhyncholestes raphanurus* (R.r.)

Variable	Comparison	Common slope			Common intercept			Shift
		Lr	P b ₁	b ₁ com	W log b ₀	P log b ₀	Log b ₀ comm	
ZB	<i>C.f.</i> - <i>L.i.</i>	0.4207	0.5165	1.09	26.76	<i>p</i> <0.01	<i>L.i.</i> > <i>C.f.</i>	
	<i>C.f.</i> - <i>R.r.</i>	0.0466	0.8289	1.10	413.40	<i>p</i> <0.01	<i>C.f.</i> > <i>R.r.</i>	
	<i>L.i.</i> - <i>R.r.</i>	0.0457	0.8306	1.06	285.84	<i>p</i> <0.01	<i>L.i.</i> > <i>R.r.</i>	
HM	<i>C.f.</i> - <i>L.i.</i>	1.5811	0.2085	1.65	44.26	<i>p</i> <0.01	<i>C.f.</i> > <i>L.i.</i>	
	<i>C.f.</i> - <i>R.r.</i>	1.8861	0.1696	1.55	1764.36	<i>p</i> <0.01	<i>R.r.</i> > <i>C.f.</i>	
	<i>L.i.</i> - <i>R.r.</i>	4.247	0.0392	<i>L.c.</i> > <i>R. r.</i>				
PAL	<i>C.f.</i> - <i>L.i.</i>	2.4399	0.1182	1.11	123.88	<i>p</i> <0.01	<i>L.i.</i> > <i>C.f.</i>	
	<i>C.f.</i> - <i>R.r.</i>	0.0281	0.8667	1.10	219.11	<i>p</i> <0.01	<i>R.r.</i> > <i>C.f.</i>	
	<i>L.i.</i> - <i>R.r.</i>	1.7548	0.1852	1.16	0.60	0.43	-0,55	<i>R.r.</i> > <i>L.i.</i>
BP	<i>C.f.</i> - <i>L.i.</i>	5.5413	0.01857	<i>L.i.</i> > <i>C.f.</i>				
	<i>C.f.</i> - <i>R.r.</i>	6.9999	<i>p</i> <0.01	<i>R.r.</i> > <i>C.f.</i>				
	<i>L.i.</i> - <i>R.r.</i>	0.3769	0.5392	0.92	278.46	<i>p</i> <0.01	<i>L.i.</i> > <i>R.r.</i>	
UP	<i>C.f.</i> - <i>L.i.</i>	7.8140	<i>p</i> <0.01	<i>L.i.</i> > <i>C.f.</i>				
	<i>C.f.</i> - <i>R.r.</i>	0.9407	0.3320	0.92	97.49	<i>p</i> <0.01	<i>C.f.</i> > <i>R.r.</i>	
	<i>L.i.</i> - <i>R.r.</i>	2.1662	0.1410	1.11	0.06	0.81	-0,65	<i>R.r.</i> > <i>L.i.</i>
ORB	<i>C.f.</i> - <i>L.i.</i>	16.9046	<i>p</i> <0.01	<i>L.i.</i> > <i>C.f.</i>				
	<i>C.f.</i> - <i>R.r.</i>	0.1187	0.7303	1.92	67.45	<i>p</i> <0.01	<i>R.r.</i> > <i>C.f.</i>	
	<i>L.i.</i> - <i>R.r.</i>	11.6099	<i>p</i> <0.01	<i>L.i.</i> > <i>R.r.</i>				
BB	<i>C.f.</i> - <i>L.i.</i>	1.6759	0.1954	0.89	18.98	<i>p</i> <0.01	<i>C.f.</i> > <i>L.i.</i>	
	<i>C.f.</i> - <i>R.r.</i>	0.3630	0.5467	0.81	396.16	<i>p</i> <0.01	<i>R.r.</i> > <i>C.f.</i>	
	<i>L.i.</i> - <i>R.r.</i>	2.3579	0.1246	0.89	345.76	<i>p</i> <0.01	<i>R.r.</i> > <i>L.i.</i>	
LN	<i>C.f.</i> - <i>L.i.</i>	2.1992	0.1380	1.34	59.96	<i>p</i> <0.01	<i>C.f.</i> > <i>L.i.</i>	
	<i>C.f.</i> - <i>R.r.</i>	0.8373	0.3601	1.29	88.10	<i>p</i> <0.01	<i>R.r.</i> > <i>C.f.</i>	
	<i>L.i.</i> - <i>R.r.</i>	2.4435	0.1180	1.39	8.29	<i>p</i> <0.01	<i>R.r.</i> > <i>L.i.</i>	
HO	<i>C.f.</i> - <i>L.i.</i>	2.9252	0.0872	0.83	169.17	<i>p</i> <0.01	<i>C.f.</i> > <i>L.i.</i>	
	<i>C.f.</i> - <i>R.r.</i>	2.8219	0.0929	0.74	14.24	<i>p</i> <0.01	<i>R.r.</i> > <i>C.f.</i>	
	<i>L.i.</i> - <i>R.r.</i>	7.3165	<i>p</i> <0.01	<i>L.i.</i> > <i>R.r.</i>				
LD	<i>C.f.</i> - <i>L.i.</i>	0.3564	0.5504	1.03	2.74	0.09	-0,27	<i>L.i.</i> > <i>C.f.</i>
	<i>C.f.</i> vs. <i>R.r.</i>	0.2494	0.6174	1.03	89.41	<i>p</i> <0.01	<i>C.f.</i> > <i>R.r.</i>	
	<i>L.i.</i> vs. <i>R.r.</i>	0.0161	0.8990	1.07	26.13	<i>p</i> <0.01	<i>R.r.</i> > <i>L.i.</i>	
HC	<i>C.f.</i> vs. <i>L.i.</i>	0.1046	0.7462	1.22	104.58	<i>p</i> <0.01	<i>C.f.</i> > <i>L.i.</i>	
	<i>C.f.</i> vs. <i>R.r.</i>	0.3559	0.5507	<i>R.r.</i> > <i>C.f.</i>				
	<i>L.i.</i> vs. <i>R.r.</i>	0.0788	0.7788	1.17	112.23	<i>p</i> <0.01	<i>R.r.</i> > <i>L.i.</i>	
LC	<i>C.f.</i> vs. <i>L.i.</i>	13.4765	<i>p</i> <0.01	<i>C.f.</i> > <i>L.i.</i>				
	<i>C.f.</i> vs. <i>R.r.</i>	10.8487	<i>p</i> <0.01	<i>R.r.</i> > <i>C.f.</i>				
	<i>L.i.</i> vs. <i>R.r.</i>	27.3084	<i>p</i> <0.01	<i>R.r.</i> > <i>L.i.</i>				
HD	<i>C.f.</i> vs. <i>L.i.</i>	13.9302	<i>p</i> <0.01	<i>L.i.</i> > <i>C.f.</i>				
	<i>C.f.</i> vs. <i>R.r.</i>	0.0216	0.8829	1.47	75.33	<i>p</i> <0.01	<i>C.f.</i> > <i>R.r.</i>	
	<i>L.i.</i> vs. <i>R.r.</i>	4.2220	0.0399	<i>L.i.</i> > <i>R.r.</i>				
LP	<i>C.f.</i> vs. <i>L.i.</i>	19.1661	<i>p</i> <0.01	<i>L.i.</i> > <i>C.f.</i>				
	<i>C.f.</i> vs. <i>R.r.</i>	11.1627	<i>p</i> <0.01	<i>C.f.</i> > <i>R.r.</i>				
	<i>L.i.</i> vs. <i>R.r.</i>	34.3593	<i>p</i> <0.01	<i>L.i.</i> > <i>R.r.</i>				

b₁com, common slope from standardized major axis analysis; log(b₀)com, common intercept from standardized major axis analysis; Lr, likelihood ratio (Warton et al., 2006); W, Wald statistic (Warton et al., 2006); P b₁, p-value of Lr parameter; P (logb₀), p-value of W parameter; Shift, growth extension

Fig. 2 Bivariate SMA regressions for *Caenolestes fuliginosus* (crosses), *Lestoros inca* (squares), and *Rhyncholestes raphanurus* (filled squares). **a** Same slope and intercepts for *L. inca* and *R. raphanurus*, but lower intercept for *C. fuliginosus* (PAL); **b** different slope in the growth trajectory of *C. fuliginosus*, and different intercepts between *R. raphanurus* and *L. inca* (BP). Lines SMA regression: dashed line *C. fuliginosus*; black line *R. raphanurus*; grey line *L. inca*. Abbreviations as in Fig. 1



for length of the palate (PAL) and the upper postcanine tooththrow (UP), although with a clear growth extension in the extremely long-nosed *R. raphanurus* (Table 3). *Y*-intercepts were larger in *R. raphanurus* than in *C. fuliginosus*, except for

those variables related to temporal muscles and bite (Osgood 1921), as zygomatic breadth (ZB), upper postcanine tooththrow (UP), and length (LD) and height (HD) of the dentary, which favored *C. fuliginosus* (Table 3). Such differences in growth

Table 4 Ontogenetic trends in the skulls of marsupials and placentals studied with multivariate analysis

Species	Source	CIL	ZB	LN	PAL	UP	LP	LD	LC	HC	HO	BB	HD	ORB	MH	BP
Marsupials																
<i>Didelphis albiventris</i>	Abdala et al. (2001)*	(+)	(+)	+	(=)	-	-	+	+	+	-	-	+	-	-	-
<i>Lutreolina crassicaudata</i>	Flores et al. (2003)*	=	+	-	-	-	-	+	+	+	-	-	+	-	=	-
<i>Caluromys philander</i>	Flores et al. (2010)	+	(=)	(=)	=	=	=	=	+	+	-	-	+	=	=	(-)
<i>Dromiciops gliroides</i>	Giannini et al. (2004)	-	-	-	-	=	-	=	+	=	-	-	+	(=)	-	-
<i>Dasyurus albopunctatus</i>	Flores et al. (2006)	-	+	=	-	-	(-)	=	+	+	-	-	+	-	=	(-)
<i>Isoodon macrourus</i>	Flores et al. (2013)	=	(=)	(=)	=	=	=	+	+	+	-	-	+	-	=	-
<i>Echymipera kalubu</i>	Flores et al. (2013)	=	-	+	+	-	-	+	+	=	-	-	+	=	=	-
<i>Caenolestes fuliginisus</i>	This report	=	=	+	=	-	-	=	+	=	-	-	+	(+)	+	-
<i>Rhyncholestes raphanurus</i>	This report	=	=	=	=	-	-	=	+	=	-	-	=	=	=	(=)
<i>Lestoros inca</i>	This report	=	-	=	=	=	=	=	(-)	=	-	-	=	(+)	=	-
Placentals																
<i>Cebus apella</i>	Flores and Casinos (2011)	(-)	+	+	-	-	-	+	+	+	-	-	+	-	-	-
<i>Alouatta caraya</i>	Flores and Casinos (2011)	=	-	+	=	-	-	+	+	+	-	-	+	-	=	-
<i>Lycalopex culpaeus</i>	Segura and Prevosti (2012)	=	+	+	-	(-)	=	+	+	+	-	-	+	-	(-)	-
<i>Puma concolor</i>	Giannini et al. (2010)	(+)	+	=	=	-	-	+	+	+	-	-	(-)	-	(=)	-

Marginally significant trends are listed in parentheses. Abbreviations of variables as in Fig. 1
 =, isometry, -, negative allometry, +, positive allometry. Asterisks indicate re-analyzed data sets under multivariate allometry

rate suggest possible differences in food habits of *Rhyncholestes* with respect to the more predaceous *Caenolestes*, as indicated in behavioral reports (Patterson and Gallardo 1987; Kirsch and Waller 1979; Patterson 2008). Despite the typical slender skull morphology exhibited

by *R. raphanurus* (Osgood 1924, 1943), the allometric trends of variables related with the length of the skull (e.g., LCI, PAL, LN, UP, LD, and LP) were mostly isometric or negatively allometric (Tables 1 and 2), suggesting that its elongated appearance is already

Table 5 Phylogenetic signal of allometric trends based on 20,000 permutations of two hypotheses of relationships

Variable	H1			H2		
	Steps	Permutation steps	P H1	Steps	Permutation steps	P H1
CIL	0,271	0,271	0,489	0,266	0,262	0,239
ZB	0,373	0,424	0,35	0,341	0,424	0,107
BB	0,228	0,298	0,012	0,242	0,298	0,031
HO	0,421	0,489	0,048	0,421	0,489	0,048
ORB	0,542	0,461	0,226	0,542	0,461	0,227
LN	0,383	0,283	0,297	0,383	0,267	0,298
PAL	0,197	0,172	0,694	0,197	0,19	0,692
BP	0,442	0,442	0,733	0,419	0,442	0,462
UP	0,29	0,309	0,121	0,29	0,306	0,121
LD	0,269	0,303	0,033	0,269	0,303	0,032
HD	0,454	0,498	0,145	0,454	0,498	0,146
HC	0,458	0,569	0,05	0,445	0,569	0,044
LC	0,589	0,608	0,611	0,593	0,608	0,738
LP	0,329	0,385	0,146	0,329	0,385	0,148
HM	0,325	0,272	0,775	0,325	0,272	0,777

In H1 Ameridelphia is monophyletic (i.e., Caenolestidae and Didelphidae are sister groups), whereas in H2 Ameridelphia is paraphyletic (i.e., Caenolestidae is sister group of Australidelphia). P indicates the probability of the found random trees being shorter than those of H1 or H2 (i.e. $p > 0.05$ means absence of pylogenetic signal)

latter being a morphologically conservative group, despite the longer rostrum of *R. raphanurus*. Extant bandicoots varied not only in size, but also in complex structures such as the tympanic region as a whole (e.g., Freedman and Joffe 1967; Freedman and Rightmire 1971; Archer 1976; Groves and Flannery 1990). However, this fact could be a consequence of the taxonomic level of our comparison, as bandicoots includes two extant families (Groves 2005, considering Chaeropodidae as extinct, Burbidge et al. 2008), whereas living caenolestids include only three closely related genera (Patterson 2008). The absence of a particular mode of growth in both long nosed groups (Table 4) can also be related to the basal position of both clades in the marsupial tree, either accepting or rejecting the monophyletic condition of Ameridelphia (e.g., Horovitz and Sánchez-Villagra 2003; Asher et al. 2004; Cardillo et al. 2004; Beck 2008; Beck et al. 2008; Meredith et al. 2008). The long nosed pattern of skull morphology exhibited in both basal groups was already a common aspect in ancient mammal taxa, such as the Cretaceous *Cronopio* (Rougier et al. 2011) or the Miocene dryolestid *Necrolestes* (Rougier et al. 2012; Chimento et al. 2012), for which such an ancient condition could easily result in convergent morphology.

Ontogenetic Evolution and Phylogenetic Significance

Our approach to the evolution of cranial ontogeny in marsupials took advantage of continuous character optimization with ontogenetic trends coded as confidence intervals extracted from intraspecific multivariate analyses (see Giannini 2014; Segura 2014). This avoids mistakes associated with discretization of ontogenetic trends, which is useful for some comparative purposes but not as much for reconstructing evolution of ontogeny in these characters. In addition, it makes use of the intraspecific variation around point estimates of allometry, as discovered by data simulation (e.g., our application of jackknife resampling; see Giannini 2014). We detected evolutionary changes both in the “sign” and “amount” of coefficients (i.e., changes along the continuous scale of allometry) at internal nodes. The shortest length recovered was that of the tree in which Ameridelphia was paraphyletic (i.e., H2; Table 5), thus providing additional support to the latest phylogenetic evidence from varied sources (e.g., Horovitz and Sánchez-Villagra 2003; Asher et al. 2004; Nilsson et al. 2004; Meredith et al. 2008). Our analysis contributed three unreversed synapomorphies from ontogenetic characters to the caenolestid + australidelphian clade (decreases in growth rate in HC, ZB, and BP; Fig. 4). H2 would indicate that the increase in braincase breadth (BB) growth rate and the decrease in length of the coronoid process (LC) growth rate in H1 (Fig. 3) are not actual synapomorphies of New World marsupials, but most likely common symplesiomorphies, as in other characters such as paired

sperm morphology (Biggers and DeLamater 1965). Indeed, BB was one of the variables with statistically significant phylogenetic signal (Table 5). As suggested by Flores et al. (2013), skull ontogeny in marsupials may be phylogenetically informative in several aspects, which is confirmed here given the significant signal observed both in splanchnocranial (LD, HC) and neurocranial (BB, HO) variables (Table 5). Indeed, adding living caenolestids to the previously available sample of marsupials generated phylogenetic support from almost all ontogenetic cranial variables to some clade. The marsupial root did not show changes in H1 or H2 with respect to the placental outgroups (as reported by Flores et al. 2013 using discretized ontogenetic characters of the skull), meaning that the divergent life-history strategies between these groups (in organogenesis, integration, development and reproduction; e.g., Clark and Smith 1993; Smith 1997; Shirai and Marroig 2010; Goswami et al. 2012) were not directly reflected in radical ontogenetic changes in skull development at the marsupial ancestor.

Neurocranium and splanchnocranium maintain a close developmental interaction both in growth pattern and function. Pattern of growth in neurocranial variables were negatively allometric in most mammals (Table 4). As evidenced in some works (e.g., Emerson and Bramble 1993; Maunz and German 1996; Wilson 2011; Segura et al. 2013; Koyabu et al. 2014; Tarnawski et al. 2014a, b), mammals exhibit an accelerated differentiation of the central nervous system and sensory capsules, producing newborns with large braincases, eyes, and auditory regions relative to the trophic components of the skull. Specifically in marsupials, much of neurogenesis occurs after birth and during lactation (Smith 1997), so during the post-weaning period most neurocranial variables scale negatively in all marsupials studied to date, except the orbit. The isometry of the orbit (which means that it grows faster than expected for a neurocranial component) in *D. gliroides*, *C. philander*, *R. raphanurus*, and *E. kalubu*, and the marginally positive allometry in *L. inca* and *C. fuliginosus*, suggests that this condition may be relatively common among marsupials. However, the marginally positive allometry of the orbit detected in *L. inca* and *C. fuliginosus* is likely an artifact of the low correlation index of this variable (Table 1), as the particular anatomy of the weak and thin zygomatic arch in shrew opossums makes it difficult to determine a clear landmark at the posterior margin of the orbit. Also, and in agreement with the consistent negative allometry in almost all neurocranial variables found in marsupials and placentals, the neurocranial region did not show any significant differences in distribution between groups in the morphospace reported by Bennett and Goswami (2013).

In spite of its alleged growth conservatism, character optimization of neurocranial variables was informative in several aspects (see Figs. 3 and 4), also demonstrated by significant phylogenetic signal (Table 5). The increase in braincase

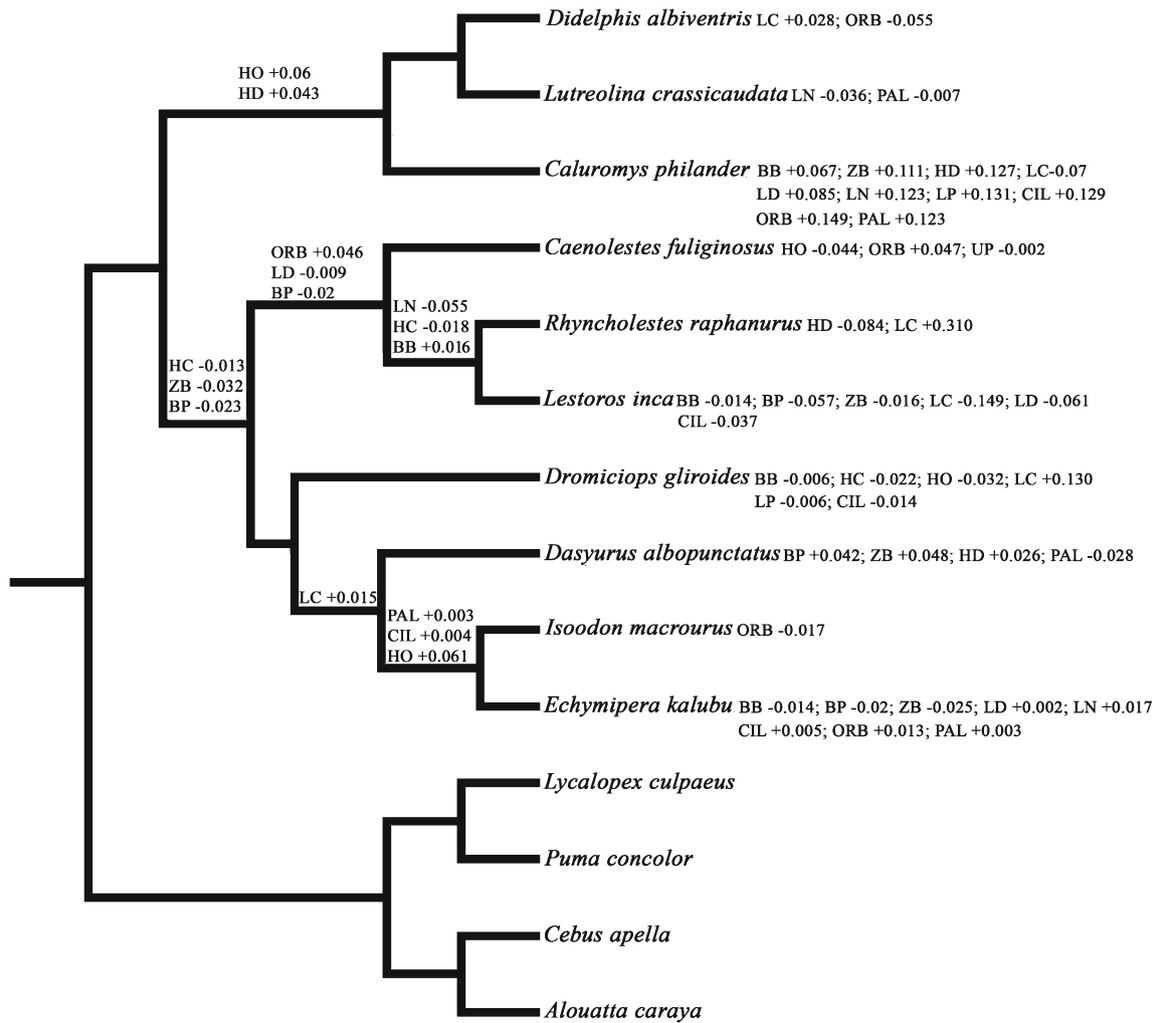


Fig. 4 Optimization of confidence intervals obtained from multivariate analysis of allometry for H2 hypothesis, where Ameridelphia is paraphyletic (see [Materials and Methods](#)). Abbreviations on internal nodes and terminals indicate variables as in Fig. 1. Numbers indicate

the difference between the confidence interval of allometry between ancestor and descendent node (or terminal). Sign + or – indicate increase or decrease of the confidence interval

growth rate in New World marsupials and the *R. raphanurus* + *L. inca* clade (in H1; Fig. 3), suggests some adjustment of neurocranial growth along the evolution of the group. However, the phylogenetic adjustment shown by the height of the occipital plate (i.e., growth rate increasing in Didelphidae and Peramelidae, Figs. 3 and 4) is likely associated with the development of the nuchal and sagittal crests on the supraoccipital bone in such groups (i.e., *D. albiventris*, *L. crassicaudata*, *C. philander*, *E. kalubu*, and *I. macrourus*). A recent report dealing with heterochronic patterns of prenatal skull ossification in mammals (Koyabu et al. 2014) suggested an accelerated timing in the ossification of the supraoccipital bone especially in those highly encephalized groups. However, the increase of the growth rate observed in our postnatal period is a consequence of the sagittal and nuchal crests in adults of larger species. Probably, the inclusion of mouse opossums and small sized dasyurids in our analysis, whose adults lack sagittal and nuchal crests, would reduce the

information content of this variable in a larger and more varied sample.

Growth of the trophic apparatus, including the rostrum and mandible, exhibited phylogenetic adjustment in several clades, some related to changes in food habits. For instance, changes in the rostral growth rate detected in both long-nosed groups (shrew opossums and bandicoots) demonstrated that both groups reach their typical rostral morphology in completely different ways. Bandicoots show an increased rate of palate growth, whereas shrew opossums exhibit a decreasing rate for dentary length and nasals, suggesting that, as already mentioned for *R. raphanurus*, its elongated morphology is reached early in its postnatal ontogeny. Indeed, the decreasing rate of palate breadth in shrew opossums (both topologies; Figs. 3 and 4) pointed in the same direction. In the larger carnivorous forms included in our sample (i.e., Didelphidae and the dasyurid *D. albopunctatus*), the height of the dentary (HD) increased its growth rate (Figs. 3 and 4),

thereby strengthening the jaw against mechanical stress imposed by chewing or biting in adults, as they shift to larger prey items as they grow. A similar condition was observed for the zygomatic breadth (ZB) and its increased rate of growth for didelphids and *D. albopunctatus* (on H1; Fig. 3). As already mentioned, the combination of the negative allometry in braincase growth and the positive allometry of ZB increases the space for the temporal musculature (Abdala et al. 2001).

Although extant shrew opossums and bandicoots exhibit a superficial convergence in general skull morphology, their postnatal allometries were different. The inclusion of Caenolestidae in the comparative analyses of cranial ontogeny in marsupials has demonstrated the importance of postnatal growth development as a source of information within a phylogenetic context. Even considering alternative topological relationships in the marsupial tree, ontogenetic characters (expressed as changes in growth rates) supported several monophyletic clades, such as Peramelidae, Didelphidae, Caenolestidae, Australidelphia, and Ameridelphia. By contrast, marsupials and placentals are not strictly supported by any specific change in growth rate (see also Flores et al. 2013). The integration of a broader taxonomic sample of small didelphids and dasyurids is crucial in order to detect if some functionally related characters (such as zygomatic breadth or height of dentary growth) are phylogenetically informative or depend on their association with alternative topologies. Finally, the further consideration of larger and smaller diprotodontians in a broad scale analysis would probably add new phylogenetic signals to this morphologically diverse group with strong anatomical specializations.

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