Ontogenetic Relationships Between Cranium and Mandible in Coyotes and Hyenas

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ABSTRACT Developing animals must resolve the conflicting demands of survival and growth, ensuring that they can function as infants or juveniles while developing toward their adult form. In the case of the mammalian skull, the cranium and mandible must maintain functional integrity to meet the feeding needs of a juvenile even as the relationship between parts must change to meet the demands imposed on adults. We examine growth and development of the cranium and mandible, using a unique ontogenetic series of known-age coyotes (Canis latrans), analyzing ontogenetic changes in the shapes of each part, and the relationship between them, relative to key life-history events. Both cranial and mandibular development conform to general mammalian patterns, but each also exhibits temporally and spatially localized maturational transformations, yielding a complex relationship between growth and development of each part as well as complex patterns of synchronous growth and asynchronous development between parts. One major difference between cranium and mandible is that the cranium changes dramatically in both size and shape over ontogeny, whereas the mandible undergoes only modest shape change. Cranium and mandible are synchronous in growth, reaching adult size at the same life-history stage; growth and development are synchronous for the cranium but not for the mandible. This synchrony of growth between cranium and mandible, and asynchrony of mandibular development, is characteristic of a highly specialized carnivore, the spotted hyena (Crocuta crocuta), but coyotes have a much less protracted development, being handicapped relative to adults for a much shorter time. Morphological development does not predict life-history events in these two carnivores, which is contrary to what has been reported for two rodent species. The changes seen in skull shape in successive life-history stages suggest that adult functional demands cannot be satisfied by the morphology characterizing earlier life-history stages. J. Morphol. 272:662–674, 2011.

INTRODUCTION

The coordination between form and function is of particular interest during ontogeny because each life-history stage represents a temporal ecological niche, and animals must be adapted to that niche to survive to the next stage. Moreover, because developing organisms continually change their morphologies, they must reconcile the conflict between demands of survival within the current life-history stage with those of maturation as they negotiate the transitions through succeeding stages (Wainwright and Reilly, 1994; Monteiro et al., 1999; Herrel and Gibb, 2006). The ontogenetic relationship between cranium and mandible is of special interest because these two parts must function in concert while both change along steep growth trajectories. Not only does each part exhibit its own ontogenetic change but also their ontogenetic coordination is necessary to ensure continuous functionality. Understanding the ontogenetic changes in mandibular shape and their integration with the cranium is basic to understanding the function of the developing feeding apparatus especially the way in which young mammals negotiate the challenges posed by competition with adults.

Young mammals do not compete directly with adults for food early in postnatal development but they must often be able to do so after weaning. During their initial postnatal development, the skull undergoes tremendous changes in both size and shape (Fig. 1), and the cranium and mandible must function to satisfy the feeding needs of a
juvenile, which are not necessarily those of an adult. To investigate the ontogeny of the feeding apparatus, and the relationships among its components, a complete description is needed of both the developing cranium and mandible. Although there are numerous descriptions of the developing cranium, there are presently few detailed descriptions of the ontogeny of mandibular shape (Cardini and Tongiorgi, 2003; Boughner and Dean, 2008; Christiansen, 2008), much less of the two parts taken together (but see, Cardini and O’Higgins, 2005).

The cranial shape maturity predicts the timing of life-history, and developmental milestones have already been shown for two rodent species with disparate life-history strategies (Mus and Sigmodon spp., Zelditch et al., 2003). It is not clear whether cranial shape maturity is equally predictive in other mammalian orders because only one comparable study has been published (Order Carnivora, Tanner et al., 2010); patterns of growth and development may be as myriad as life-history strategies. Further, it is not known whether mandibular shape maturity can likewise predict life-history scheduling. Understanding the relationship between development of the feeding apparatus (the skull) and life-history schedules is important because it has obvious implications for fitness.

We capitalized on a unique opportunity to assemble a large ontogenetic series of known-age skulls of coyotes (Canis latrans) and provide detailed descriptions of the ontogenetic allometry of a nondomesticated canid utilizing shape-based geometric morphometrics. Our investigation considers the ontogeny of the cranium and mandible concurrently, which is vital to understanding their temporal relationship and integration across postnatal development. These descriptive data serve as a useful contrast with those obtained from more specialized carnivores (e.g., spotted hyena, Tanner et al., 2010), a baseline for studies of variation within the genus Canis (e.g., Wayne, 1986; Kieser and Groeneveld, 1992), and a groundwork for future studies of the ontogeny of coyote feeding performance (La Croix et al., unpublished data).

Here, we first describe the ontogenetic trajectories for age-related change in shape of the coyote cranium and mandible across the entire lifespan. Next, we investigate whether these ontogenetic trajectories are simple or composed of multiphase stages and describe any temporally restricted shape change. Nonlinear growth models are fitted to our data to describe cranial and mandibular growth (i.e., changes in size over ontogeny) and development (i.e., changes in shape over ontogeny), examine sexual dimorphism, and assess growth and development within the context of the coyote’s life history. We then describe the ontogenetic patterns and temporal coincidence of skull growth and development including their synchrony, or lack thereof, and implications for functionality of the feeding apparatus. Finally, we contrast these patterns of development in coyotes with those in a highly specialized carnivore, the spotted hyena (Crocuta crocuta), to uncover more generalized patterns of skull development and examine their relationship to life-history events.

MATERIAL AND METHODS

Coyote Life-History Stages

The annual cycle of coyote life-history events begins with mating during January, February, and March (Bekoff, 1977). Following a 63-day gestation, pups are born beginning in March (Bekoff, 1977). Weaning of pups begins by four weeks of age and is usually completed by six weeks of age (Knowlton, personal communication; Snow, 1967; 5–7 weeks in Bekoff, 1977). The onset of deciduous tooth replacement occurs by 12 weeks of age, and eruption of the adult dentition is complete by 26 weeks of age (Bekoff, 1977; SLC, personal observation). Pups are
initially provisioned by both parents, and during the summer, the juveniles develop greater independence and self-reliance (Harrison and Gilbert, 1985; Harrison et al., 1991). Subadults of either sex may emigrate in the fall of the year; first breeding in nonpersecuted populations typically occurs at 22 months of age although both sexes are physiologically capable of breeding at 10 months of age (Bekoff, 1977; Bekoff and Gese, 2003).

Here, we divide the coyote’s life history into five stages (Table 1) based on documented life-history milestones. The preweaning stage lasts from birth until weaning, at six weeks of age (Bekoff, 1977). The postweaning stage lasts from weaning until the onset of deciduous tooth replacement, at ~12 weeks of age (Bekoff, 2001). The juvenile stage lasts from the onset of deciduous tooth replacement until eruption of adult dentition is complete, which occurs by 26 weeks of age (SLC, personal observation). The subadult stage starts with the complete eruption of adult dentition at 26 weeks of age and ends with first breeding, at ~93 weeks of age (Bekoff and Gese, 2003). The adult stage extends from first breeding to death, which may occur at 13 or more years.

Among canids, coyotes are usually categorized as omnivorous generalists, with young coyotes capitalizing on that portion of the adult diet that is most easily obtained or subdued (Clark, 1972; Bowen, 1978; Johnson, 1978; Andelt et al., 1987; Gese et al., 1996; Hidalgo-Mihart et al., 2001; Arjo et al., 2002; Hernandez et al., 2002). Following weaning, juvenile coyotes are initially provisioned by their parents; subsequently, they ingest mostly small mammals, vegetation, invertebrates, and birds, whereas adults consume larger mammals, fewer invertebrates, mostly small mammals, vegetation, invertebrates, and birds (Hawthorne, 1970). Fruit and insect ingestion by coyotes occurs most frequently in the summer, which corresponds temporally with a coyote pup’s increasing independence (Young et al., 2006).

Specimens

An ontogenetic series of 187 coyote skulls (102 males and 85 females), all of known age, was used in this study; sample sizes are shown in Table 1. Individuals ranged in age from 1 day to 13.3 years, including 79 specimens under the age of 6 months. All specimens were prepared by and catalogued into either the Michigan State University Museum collection or the United States Department of Agriculture/APHIS/Wildlife Services National Wildlife Research Center’s Logan Field Station collection in Millville, UT (Supporting Information Appendix Table A1.). Specimens were collected incidentally from a captive colony of coyotes comprised of animals of wild-caught and colony parent-age and maintained at the Logan Field Station between 1979 and 2006. Within the ontogenetic series, 179 specimens were captive born, including all specimens under the age of 1 year, whereas eight were wild born. All animals were captive reared, and although this study did not investigate whether the observed ontogenetic patterns are equivalent to those found in the wild, rearing conditions at the Logan Field Station were designed to be as natural as possible. All animals were maintained on a commercially produced wet food diet designed for fur bearing animals and containing chicken (beaks, feet, and feathers) and grain. USDA/APHIS/WS/NWRC IACUC approved the study protocol QA-1179.

Exact dates of death were known for all individuals, and exact dates of birth were known for all captive-born animals. For animals obtained from the wild as pups, dates of birth were estimated by the original collectors based on deciduous tooth eruption and den observations; those estimated dates of birth were used here. Collector and animal records for all specimens are maintained at the Logan Field Station.

Ontogenetic Shape Change

Ontogenetic changes in cranial and mandibular growth and development were analyzed by geometric morphometrics (Rohlf and Slice, 1990; Zelditch et al., 2004). Each cranium was photographed in ventral and lateral views. In ventral view, the skull was oriented with the palate parallel to the photographic plane, and in lateral view, the skull was oriented with the mid-sagittal plane parallel to the photographic plane. Mandibles were photographed in lateral view, oriented with the longest axis of the mandible parallel to the photographic plane; only mandibles with intact symphyses were photographed. To assess measurement error, an initial sample of 121 crania was photographed in ventral and lateral views at three different times by the same observer. Measurement error was determined to be negligible, accounting for only 1.2% of the variation in the sample. The remaining 66 specimens were thus photographed and digitized only once; mandibles were photographed and digitized only once. Twenty-seven landmarks visible on photographs of the ventral cranium provide the data for the analysis of shape in that view (Fig. 2a). Landmarks alone cannot fully capture the dorsal curve of the lateral cranium or the mandibular ramus, so these were analyzed using a combination of landmarks and semilandmarks. Semilandmarks, unlike landmarks, are not discrete anatomical loci that can be recognized as homologous points, and they contain less information than landmarks because their spacing along the curve is arbitrary. However, semilandmarks make it possible to study complex curving morphologies where landmarks are sparse. Fourteen landmarks and 32 semilandmarks were selected for the lateral cranium (Fig. 2b), and 11 landmarks and 75 semi-landmarks were selected for the mandible (Fig. 2c). Descriptions of these landmarks are contained in Supporting Information Appendix Table A2. Landmarks and semilandmarks were digitized using tpsDig2.10 (Rohlf, 2005). For semilandmarks, the curve-tracing function was applied, and semilandmarks were evenly spaced along the curves using the “resample” function of the curve-tracing tool.

Landmarks were superimposed using Generalized Procrustes Analysis to remove variation in scale, position, and orientation (Rohlf and Slice, 1990; Zelditch et al., 2004). Semilandmarks require a specialized superimposition method because their spacing is biologically arbitrary. Semilandmarks were superimposed to minimize the Procrustes distance from the mean shape; we use this criterion for sliding semilandmarks because the Procrustes distance is the metric underlying the general theory of shape. According to this method, the tangent to the curve at each semilandmark is estimated and then each semilandmark is slid toward the normal of its respective tangent,
of shape. Results are depicted using the thin-plate spline function.

Using linear regression to analyze the whole of ontogeny assumes that the same shape changes occur over all ontogenetic phases. To test the hypothesis that ontogeny is more complex than that, we subdivided our sample into five life-history stages (Table 1), and compared the ontogenetic trajectories of successive stages (so long as statistically significant age-related shape change was present within both single stages). Comparisons were done by measuring the angle between the vectors of ontogenetic change; should both points in the same direction, the angle between them is 0°. The statistical analysis tested the null hypothesis that the angle between the vectors was no greater than expected by chance. That was done by comparing the angle between successive life-history stages to the range of angles that could be obtained by resampling with replacement (400 times), as this is a bootstrap procedure, within a single stage. Should the angle between vectors exceed the 95% confidence intervals for both stages, the difference was judged statistically significant; such a difference does not imply that ontogeny changed direction at the boundary between stages. Regression was done in Regress6 (Sheets, 2003); comparisons between vectors were done in VecCompare (Sheets, 2003).

**Rates and Timing of Growth and Maturity**

To determine the age at which skull growth and development attained adult values, we used a series of nonlinear growth models to find the one that best fit the data for size and shape. Eight models, Chapman-Richards, Logistic, Monomolecular, Gompertz, German Gompertz, Von Bertalanffy, Quadratic, and Linear were fitted to the skull size and shape data and assessed for their relative goodness-of-fit using Akaike Information Criterion (AIC; following Zelditch et al., 2003); the model with the lowest AIC value was judged to be the best so long as the residuals were not serially autocorrelated. Several models were excluded because they induced autocorrelations among residuals in one or more of the analyses (Supporting Information Appendix A.3. and A.4.). Of those that remained, several fit equally well. We chose the logistic model (following Gaillard et al., 1997) because this model not only fit the data well in all views, but it was the only model that did not induce autocorrelations among residuals in any of the analyses. This model was the basis for estimating rates and timing of growth and development:

\[
x(t) = A / (1 + e^{-(T_0 - t)})
\]

where \(x(t)\) is the measurement of interest at time \(t\), \(A\) is the asymptotic maturity, \(k\) is the rate of growth (rate of approach to asymptotic value), and \(T_0\) is the age at the curve inflection point (where growth has attained 50% of asymptotic value; Gaillard et al., 1997; Zelditch et al., 2003). We report age at maturity as the estimated age at which the measurement of interest reaches 95% of asymptotic (adult) value. Data for individuals above the 95% breakpoint were subsequently regressed on age to ensure that age had no further significant impact on the measurement of interest. Evaluation of growth models and estimation of parameters were performed using GrowChoice (Sheets, 2003).

To analyze ontogenetic change in shape across the entire life-span and within single life-history stages, shape was regressed on age; age was log-transformed because most shape change occurs early in ontogeny. The relationship between shape and age was tested for statistical significance using a generalized Goodall’s F test (Sheets, 2003; Rohlf, 2007), which measures the ratio of explained to unexplained variation in units of Procrustes distance. Adult coyotes are known to be sexually dimorphic in size; adult females are up to 20% smaller than males (Bekoff, 1977; Bekoff and Gese, 2003). We assessed whether the ontogeny of shape is also sexually dimorphic using Mancova, with sex as the main factor and age as the covariate. Because sexual shape dimorphism was not statistically significant in any view, we pooled males and females in the analyses minimizing the overall difference from the reference (Sampon et al., 1996; Anderson et al., 2000; Bookstein et al., 2002).

Following superimposition, semilandmarks can be used in conventional shape analyses provided that statistical tests take into account that they have only one degree of freedom. For the ventral cranium, bilaterally homologous landmarks were reflected and averaged after the Procrustes superimposition because bilaterally homologous landmarks are not independent of each other; to ease interpretation of the visual results, they are shown as whole skulls. To quantify skull size, we used centroid size, the square root of the summed squared distances from each landmark to the geometric center of the object. Superimposition of landmarks was done using CoordGen6 (Sheets, 2009); semilandmarks were superimposed in SemiLand (Sheets, 2003). Reflection and averaging of bilateral landmarks was done in Sage (Marquez, 2007).

To analyze ontogenetic change in shape across the entire life-span and within single life-history stages, shape was regressed on age; age was log-transformed because most shape change occurs early in ontogeny. The relationship between shape and age was tested for statistical significance using a generalized Goodall’s F test (Sheets, 2003; Rohlf, 2007), which measures the ratio of explained to unexplained variation in units of Procrustes distance. Adult coyotes are known to be sexually dimorphic in size; adult females are up to 20% smaller than males (Bekoff, 1977; Bekoff and Gese, 2003). We assessed whether the ontogeny of shape is also sexually dimorphic using Mancova, with sex as the main factor and age as the covariate. Because sexual shape dimorphism was not statistically significant in any view, we pooled males and females in the analyses.
et al., 2003; Tanner et al., 2010). A large Procrustes distance for a particular individual could mean that the individual is oddly shaped rather than mature, but we found no major outliers in our sample.

RESULTS
Changes in Shape Across the Entire Life Span

Statistically significant and visually dramatic ontogenetic changes in skull shape occur between 1 day and 13 years of age in all three views (Fig. 3). For all views, the null hypothesis of isometric growth can be rejected ($P < 0.01$). In ventral view, the dominant feature of shape change is the relative narrowing of the skull along the anteroposterior axis and lengthening of the rostrum (Fig. 3a). In lateral view, the cranial profile becomes less rounded, and the nuchal and occipital crests expand posterodorsally (Fig. 3b). The dominant features of mandible ontogeny are the relative expansion of the angular process and the marked anterodorsal reorientation of the coronoid process, resulting in a taller and more vertically oriented process in adults than neonates (Fig. 3c).

Changes in Shape Within Life-History Stages

Shape changes substantially throughout preweaning, postweaning, and juvenile stages for both cranial views; the null hypothesis of isometry can be rejected for those stages ($P < 0.01$). Isometry can also be rejected for the adult lateral view ($P = 0.011$), but not for the adult ventral view ($P = 0.060$) or the subadult cranial views ($P > 0.05$). For the mandible, age-related changes in shape are statistically significant in all but the adult life-history stage ($P < 0.001$). Comparisons between successive stages that exhibit statistically significant changes in shape reveal the complexity of ontogeny (Table 2). In all cases, the differences between stages are statistically significant and in some the angles are exceptionally large, especially in the case of the mandible, where the angles for pre- vs. postweaning, and juvenile vs. subadult, exceed $90^\circ$. It is clear from this that the dominant features of a complete ontogeny (as described above) cannot capture the ontogenetic dynamics of shape; we,

![Fig. 3. Ontogenetic changes from a linear regression of shape on log(age) for the cranium (a: ventral, b: lateral) and mandible (c). Vectors on landmarks and semilandmarks show the direction and magnitude of change from the youngest to the oldest specimens after centroid size is scaled to the same size for each specimen.](image)

<table>
<thead>
<tr>
<th>Skull view</th>
<th>Life history stages compared</th>
<th>Age-related change in shape</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Between (°)</td>
<td>Younger (°)</td>
</tr>
<tr>
<td>Ventral</td>
<td>Pre- vs. postweaning</td>
<td>34.6</td>
</tr>
<tr>
<td></td>
<td>Postweaning vs. juvenile</td>
<td>40.4</td>
</tr>
<tr>
<td>Lateral</td>
<td>Pre- vs. postweaning</td>
<td>46.7</td>
</tr>
<tr>
<td></td>
<td>Postweaning vs. juvenile</td>
<td>26.3</td>
</tr>
<tr>
<td>Mandible</td>
<td>Pre- vs. postweaning</td>
<td>110.7</td>
</tr>
<tr>
<td></td>
<td>Postweaning vs. juvenile</td>
<td>73.9</td>
</tr>
<tr>
<td></td>
<td>Juvenile vs. subadult</td>
<td>97.1</td>
</tr>
</tbody>
</table>

Comparisons were made between successive life history stages only in cases where both single stages exhibited significant age-related change in shape. Ontogenetic trajectories of two successive stages were determined to differ significantly when the angle between the vectors of the two stages exceeded that obtained by resampling (400 times) within each stage (younger and older). If the observed angle between stages exceeded the 95% confidence interval of the two within-stage ranges, the difference was judged to be statistically significant, here indicated in bold font.

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therefore, describe the changes occurring within each stage.

During the preweaning stage (Figs 4a and 5a), nearly half of the variance in cranial shape is related to age (ventral view: 46.6%, lateral view: 46.1%). In this first stage, the cranium changes shape dramatically as the initially bulbous neonate skull undergoes a tremendous elongation of rostrum and palate, and the basicranium narrows relative to its length and diminishes in relative height. Further, the frontal and parietal bones align dorsally, creating a continuous dorsal curve, visible in the lateral view of the cranium. During the subsequent postweaning stage (Figs. 4b and 5b), when feeding is accomplished with deciduous teeth, shape continues to change, but age accounts for a much lower proportion of the variance for ventral (30.1%) than for lateral view (51.9%). Rostral elongation continues, as does relative narrowing of the basicranium and flattening of the brain case, all of which further reduce the infant skull’s bulbosity. Specific to the postweaning stage, the basicranium lengthens relative to skull length as evidenced by the posterior displacement of the landmarks for the jugular foramen and the posterior point of the foramen magnum. In the next interval, the juvenile stage (Figs. 4c and 5c), the deciduous teeth are replaced by adult dentition. The proportion of ventral cranial shape accounted for by age here (29.1%) is similar to that seen in the previous stage, but a greater proportion of the variance is accounted for by age in lateral view (63%). During this stage, the general pattern of cranial elongation and dorso-ventral flattening continues, but now there is also expansion of the zygomatic arches (evident from the posterolateral displacement of the landmark at the jugal/squamosal suture), as well as enlargement of the sagittal crest, which lengthens and broadens, creating an increased area for origin of the temporal muscles. No statistically significant changes in cranial shape were detected during the subadult stage. However, in the adult stage (Figs. 4e and 5e), cranial shape in the lateral view changed significantly, with age accounting for 9.5% of the variance while cranial shape in the ventral view was nearly significant ($P = 0.060$). Over this interval, shape change is mostly subtle except for a distinct widening of the zygomatic arches.

Age consistently explains little of the variance in mandibular shape (9.4%, 10.8%, 16.3%, and 11.8% for preweaning, postweaning, juvenile, and subadult, respectively). During the preweaning stage (Fig. 5f), the anterior portion of the mandible body elongates and deepens and the vertical ramus dramatically broadens. In the subsequent, postweaning stage (Fig. 5g), the mandible body and ramus (anterior to the condyloid process) broaden and deepen, and the condyloid process expands posteriorly. During the next juvenile stage (Fig. 5h), the
coronoid process lengthens and broadens, whereas the vertical ramus elongates, becoming more perpendicular in orientation relative to the base of the ramus. During the subadult stage (Fig. 5i), there is an overall flattening and reduction in ventral curvature along the body of the ramus and ongoing growth at the extremes of the coronoid and condyloid processes. And as noted above, no

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statistically significant changes in shape are detected during the adult stage for the mandible.

**Rates and Timing of Growth and Maturity**

The best-fitting model for both growth and development of the cranium and mandible is logistic growth (Table 3; see also, Supporting Information Appendix Tables A.3. and A.4.). Asymptotic adult size \((A)\) differs significantly between male and female coyotes (Table 4), so each sex was analyzed separately when estimating the age at which size and shape reached adult maturity (95% of the asymptotic values). Among males, for both ventral and lateral views of the cranium, adult size is reached at 22.5 weeks, and for the mandible at 23 weeks. Females reach adult size within one week of males for all views of the skull (22 weeks). The age at which shape reaches adult maturity is not as consistent among views (Table 5). For the ventral skull, adult shape is reached at 17.2 weeks in males and 18 weeks in females, but the lateral skull takes three to four weeks longer, maturing by 21 weeks in both sexes. The mandible takes even longer, reaching adult shape at 28.2 weeks in both sexes.

Placed in the context of milestones regularly recorded in mammalian life-history studies, the cranium and mandible follow similar growth patterns (Table 6, Fig. 6a). Degree of maturation is similar at birth, and all views attain over half of their adult size by weaning (ventral view, 55% for males, 56% for females; lateral view, 60% for both sexes; mandible, 51% for males, 52% for females). By the onset of tooth replacement at 12 weeks, all have attained ~80% of the adult size (ventral view, 77% for males, 78% for females; lateral view, 79% for males, 80% for females; mandible, 74% for males, 76% for females). The crania and mandibles reach maturity up to three weeks before the adult dentition completes eruption at 26 weeks (Fig. 7).

In striking contrast to their synchronous patterns of growth, cranial and mandibular shape show very disparate ontogenetic patterns. At birth, cranial shape is conspicuously immature, whereas mandibular shape is remarkably mature (Table 6, Fig. 6b). Over the first two life-history stages (preweaning and postweaning), cranial shape changes rapidly, such that, by the onset of tooth replacement, the cranium has progressed roughly 80% of the way to its mature shape for features visible in ventral view (83% for males; 80% for females) and 65% for features visible in lateral view (67% for males; 65% for females). Mandibular shape changes much more gradually, and although maturity at the onset of tooth replacement falls between the two cranial views (75% for both sexes), the mandible does not reach adult shape until several weeks after the cranium; indeed, maturity of the mandible is not attained until two weeks after the adult dentition is in place (Fig. 7).

**DISCUSSION**

At birth, the coyote cranium is small and immature; both size and shape change dramatically over ontogeny. The coyote mandible, like the cranium, is also small at birth. However, unlike the cranium, the shape of the mandible is not immature at birth, and it changes only modestly in shape over ontogeny. In its cranial development, the coyote skull follows the general mammalian pattern (Moore, 1981): bulbous at birth with subsequent elongation of the cranium relative to its width, flattening of the lateral profile, and broadening of the zygomatic arches. The ontogeny of the mandible also follows a general mammalian pattern: an elongation of the horizontal body relative to its height, vertical reorientation and broadening of the zygomatic arches. The ontogeny of the mandible is more complex than suggested by these general patterns; for each part there are features that change during only a specific phase of ontogeny. For example, the basicranium and nuchal crest are displaced posteriorly between weaning and the onset of deciduous tooth replacement, whereas the zygomatic arches...
TABLE 4. Estimates, based on best fitting model, for asymptotic size, $A$; growth rate constant, $k$; age (in weeks) at the curve inflexion point, $T_o$; and age (in weeks) at adult maturity, $M$; 95% confidence intervals given in parentheses

<table>
<thead>
<tr>
<th>Sex</th>
<th>Ventral cranium</th>
<th>Lateral cranium</th>
<th>Lateral mandible</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$A$</td>
<td>$k$</td>
<td>$T_o$</td>
</tr>
<tr>
<td>Female</td>
<td>273.095$^a$</td>
<td>0.168</td>
<td>4.501</td>
</tr>
<tr>
<td>Male</td>
<td>286.303$^a$</td>
<td>0.167</td>
<td>4.831</td>
</tr>
</tbody>
</table>

$^a$Asymptotic size is significantly different between male and female ventral crania, $t = 6.3767$, $p = 1.81e-010$.

$^b$Asymptotic size is significantly different between male and female lateral crania, $t = 7.5942$, $p = 3.0864e-014$.

$^c$Asymptotic size is significantly different between male and female lateral mandibles, $t = 7.5377$, $p = 4.774e-014$.

$^d$ $T_o$ for growth is highly suggestive of a trend for a difference between male and female lateral mandibles, $t = 1.9323$, $p = 0.05332$.

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TABLE 5. Estimates, based on best fitting model, for asymptotic shape, $A$; growth rate constant, $k$; age (in weeks) at the curve inflexion point, $T_o$; and age (in weeks) at adult maturity, $M$; 95% confidence intervals given in parentheses

<table>
<thead>
<tr>
<th>Sex</th>
<th>Ventral cranium</th>
<th>Lateral cranium</th>
<th>Lateral mandible</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>$A$</td>
<td>$k$</td>
<td>$T_o$</td>
</tr>
<tr>
<td>Female</td>
<td>0.124</td>
<td>0.270</td>
<td>6.991</td>
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<tr>
<td>(0.122–0.127)</td>
<td>(0.237–0.308)</td>
<td>(6.634–7.378)</td>
<td>(17.2)</td>
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<tr>
<td>Male</td>
<td>0.122</td>
<td>0.284</td>
<td>6.608</td>
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<td>(0.120–0.124)</td>
<td>(0.256–0.318)</td>
<td>(6.343–6.889)</td>
<td>(17.2)</td>
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Note: No significant difference between males and females for $A$, $k$ or $T_o$ for ventral cranium shape, lateral cranium shape or lateral mandible shape.
lengthen, relative to the basicranium, and the sagittal crest takes shape, between the onset of deciduous tooth replacement and the completion of adult dentition eruption. For the mandible, there is an elongation and deepening of the anterior-most part of the horizontal mandible body between birth and weaning, whereas there is an elongation and deepening of the posterior parts of the body of the mandible and ramus (anterior to the condyloid process) between weaning and the onset of deciduous tooth replacement. These age-specific patterns are distinctive enough to make ontogenetic trajectories during successive life-history stages significantly different from one another.

The cranium and the mandible are synchronous in their growth, keeping pace with one another as they enlarge, and reaching adult size at the same life-history stage. The cranium itself is also synchronous with respect to growth and development; i.e., cranial size and shape show similar temporal patterns and reach adult levels at the same life-history stage. In contrast, whereas mandibular growth keeps pace with cranial growth, the developmental patterns evident in mandibular and cranial shape exhibit striking differences. The mandible is relatively mature in shape at birth, but despite this early maturity, it develops more slowly and attains adult shape later than the cranium (Fig. 7). Any mismatch between the growth and maturation of different parts belonging to a single functional complex can be disruptive unless function itself is also changing. When function is changing, however, constancy of the relationships between parts would not be expected. The functional relationships between cranium and mandible (as well as among regions within each part) do change over postnatal mammalian ontogeny. An

### Table 6. Relative maturity of size (CS/A) and relative maturity of shape (PD/A) at post-natal ages, based on parameters of the logistic model

<table>
<thead>
<tr>
<th>Age (Weeks)</th>
<th>Ventral cranium</th>
<th></th>
<th>Lateral cranium</th>
<th></th>
<th>Lateral mandible</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Female CS/A</td>
<td>Female PD/A</td>
<td>Male CS/A</td>
<td>Male PD/A</td>
<td>Female CS/A</td>
<td>Female PD/A</td>
</tr>
<tr>
<td>Birth</td>
<td>0.319</td>
<td>0.132</td>
<td>0.309</td>
<td>0.133</td>
<td>0.372</td>
<td>0.070</td>
</tr>
<tr>
<td>4</td>
<td>0.478</td>
<td>0.309</td>
<td>0.465</td>
<td>0.324</td>
<td>0.529</td>
<td>0.180</td>
</tr>
<tr>
<td>8</td>
<td>0.642</td>
<td>0.568</td>
<td>0.629</td>
<td>0.600</td>
<td>0.681</td>
<td>0.390</td>
</tr>
<tr>
<td>12</td>
<td>0.779</td>
<td>0.795</td>
<td>0.767</td>
<td>0.825</td>
<td>0.802</td>
<td>0.651</td>
</tr>
<tr>
<td>16</td>
<td>0.873</td>
<td>0.920</td>
<td>0.885</td>
<td>0.938</td>
<td>0.885</td>
<td>0.845</td>
</tr>
<tr>
<td>20</td>
<td>0.931</td>
<td>0.972</td>
<td>0.926</td>
<td>0.981</td>
<td>0.936</td>
<td>0.942</td>
</tr>
<tr>
<td>24</td>
<td>0.964</td>
<td>0.991</td>
<td>0.961</td>
<td>0.996</td>
<td>0.965</td>
<td>0.980</td>
</tr>
<tr>
<td>28</td>
<td>0.981</td>
<td>0.998</td>
<td>0.979</td>
<td>0.999</td>
<td>0.981</td>
<td>0.994</td>
</tr>
<tr>
<td>32</td>
<td>0.990</td>
<td>0.999</td>
<td>0.989</td>
<td>0.999</td>
<td>0.990</td>
<td>0.999</td>
</tr>
</tbody>
</table>

CS is centroid size, PD is Procrustes distance, and A is the corresponding asymptotic value for shape or size.

Fig. 6. Timeline illustrating the relative maturity in size (a) and shape (b) for the ventral and lateral views of the cranium and the mandible for coyotes in relation to major life-history events. There is no significant difference between males and females in rate of maturation of skull size or skull shape.

Fig. 7. Timeline illustrating the age at maturation for coyote skull morphology in relation to major life history events. Male and female skull maturation occurs within the same week for each morphological measure; the latest maturation age is indicated for each measure by a downward-pointing arrow.
obvious example is that suckling and mastication make different demands with respect to occlusion between upper and lower teeth. We might generally expect changes in the relationships among parts unless the demands of adult function are satisfied by the infant morphology. It is, therefore, not surprising that the coyote cranium and mandible each display the complex ontogeny described above, or that the mandible and cranium are asynchronous in development if not in growth.

Adult shape of the coyote feeding apparatus is attained long after the transition to solid food at six weeks of age: cranial shape does not mature for another 11–14 weeks, and mandibular shape, for another 22 weeks. At weaning, cranial shape is as little as 30% mature, whereas the mandible is already 61% mature, having been more mature at birth. Thus, juvenile coyotes are far from mature at weaning, so we would expect that their feeding performance at weaning would be poor relative to that of adults. Further, the ongoing maturation of the skull parts suggests that adult function is not achieved by the morphology characteristic of the postweaning stage. Synchrony of cranial and mandibular development may be unnecessary except to the extent that it affects growth and orientation of the masticatory muscles. Bone growth plays a pivotal role in orienting muscles and their action lines, in that regional bone growth differentially stretches the periosteum, with muscles being carried along with it via their periosteal attachments (Grimm and Katele, 1979; Carlson, 1983; Frankenhuys-van den Heuvel et al., 1992). Additionally, bones must grow to accommodate enlarging muscles, so we might anticipate synchrony in regions where masticatory muscles originate and insert, such as the sagittal crest (visible only in lateral view) and mandible. Indeed, our results show that association between mandibular development and sagittal crest; mandibular development in the subadult stage is marked by localized changes in condylar and coronoid shape and these changes are concurrent with localized cranial changes, particularly in the sagittal crest, that continue through adulthood. The influence of developing secondary and primary dentition on the synchrony of cranial and mandibular shape development remains unexplored, but tooth development and occlusion might well affect the synchrony of cranial and mandibular development.

It is not presently known whether the asynchrony shown here in coyote cranial and mandibular shape development reflects a more generalized mammalian pattern because only one other study, on the spotted hyena, Crocuta crocuta (Tanner et al., 2010), has concurrently examined ontogenetic patterns of cranial and mandibular shape change in relation to life history. Hyenas, unlike coyotes, are capable as adults of cracking open large bones using craniodental adaptations for durophy that include a vaulted forehead, a large sagittal crest, massive zygomatic arches and robust premolars with crack-resistant enamel. Another distinction between hyenas and coyotes may be related to those morphological demands of durophy: the hyena’s life history is notable for its protracted period of maternal dependence even in relation to other large carnivores (Watts et al., 2009). Skull ontogeny in the spotted hyena, like that in coyotes, demonstrates relative synchrony in cranial and mandibular growth but asynchrony in cranial and mandibular shape development, cranial size and shape maturation, and mandibular size and shape maturation (Tanner et al., 2010). What differs strikingly between coyotes and hyenas is both the sequence of adult size and shape maturation events and their timing within the life history (Fig. 8.). In the coyote, mandibular shape matures after cranial shape, and mandibular shape is the last skull measure to mature. By contrast, in the spotted hyena, cranial shape maturation is dramatically delayed, occurring long after mandibular shape maturation. In addition, all coyote skull maturation is completed before or shortly after the eruption of the adult dentition and long before reproductive maturity, but in the spotted hyena, skull maturation (except for cranial shape in the lateral view) occurs at or after reproductive maturity. Like the coyote skull, the spotted hyena skull matures long after weaning, and despite a late age-at-weaning in spotted hyenas (compared to other carnivores), the skull of a just-weaned spotted hyena is still notably immature (ventral, 73.5% mature; lateral, 90.3%; mandible, 83.5%, Tanner, unpublished data). Delayed skull maturation in the spotted hyena, especially late cranial maturation and delayed feeding performance maxima, may be explained as costs of the adult ability to crack open large bones. Indeed, the dominant feature of postweaning shape change in a spotted hyena skull is the development of the bony areas

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of muscle insertion: zygomatic arches expand and the sagittal and nuchal crests develop (Tanner et al., 2010). For coyotes and other mammalian carnivores lacking such specialized adult function, we might thus expect a more modest delay in skull maturation and achievement of feeding performance maxima than is seen in spotted hyenas. We have demonstrated the former here; future work is needed to determine whether a comparable delay in maturation also exists regarding achievement of feeding performance maxima and whether maturation schedules influence dietary choices.

Overall, comparison of ontogenetic patterns in coyotes and spotted hyenas reveals that both species share a synchronous pattern of maturation in growth between cranium and mandible, but temporally mismatched patterns of maturation in shape between cranium and mandible. That morphological development does not predict life-history events in these two carnivores distinguishes them from the two rodents studied to date; in those two rodents, one precocial, the cotton rat (Sigmodon fulviventris), and the other altricial, the house mouse (Mus musculus domesticus), cranial shape maturity predicted life-history scheduling (Zelditch et al., 2003). In contrast, coyote skull maturity occurs well before sexual maturity, and spotted hyena skull maturity occurs well after.

By looking at growth and development in the cranium and mandible concurrently, examining the relationship between their maturation patterns, and placing both within the context of life history, we can identify a synchronous pattern of maturation in growth across parts as well as a temporally mismatched pattern of shape maturation. It is likely that these patterns are complicated by the interaction between localized features in one part of the skull with those of another (such as the cranial origins and mandibular insertions of masticatory muscles). Clearly, additional studies on carnivores are needed to provide a richer context for interpreting the causes, consequences and functional bases for differences between cranial and mandibular ontogenetic patterns. Of particular importance for understanding the links between morphology, feeding performance, and life history is the protracted period of cranial development in carnivores, which makes juveniles handicapped relative to adults for a significant portion of their life history. Our findings demonstrate that an adult’s functional demands cannot be satisfied by the morphology characterizing earlier life-history stages, and this has significant consequences for life-history evolution whenever juveniles must compete with adults.

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