Feathered dinosaurs, flying dinosaurs, crown dinosaurs, and the name "Aves"

Jacques Gauthier

Department of Geology and Geophysics, P. O. Box 208109 Yale University, New Haven, CT 06520-8109 U.S.A.

Peabody Museum of Natural History, Yale University P. O. Box 208118, New Haven, CT 06520-8118 U.S.A. *E-mail:* jacques.gauthier@yale.edu

Kevin de Queiroz

Department of Vertebrate Zoology, National Museum of Natural History Smithsonian Institution, Washington, DC 20560 U.S.A. *E-mail*: dequeirk@nmnh.si.edu

Abstract

The taxon name "Aves" is currently used for several different clades, a situation that violates the fundamental nomenclatural principle that, to minimize ambiguity, each taxon name should refer to a single taxon. To clarify this situation, we explore some general issues concerning the properties of the three classes of phylogenetic definitions, including: how names can be tied to clades through composition or characters; the relationships between "total" and "crown" compared to "stem" and "node" clades; how taxa can be referred to clades whose names are defined using different classes of phylogenetic definitions; perceived problems with apomorphy-based definitions; whether and in what senses taxon names have traditional meanings; and the consequences of restricting Aves to each one of the four clades to which it is most commonly applied. We then propose a taxonomy that resolves current nomenclatural ambiguity by using a different name for each of the clades that everyone wants to talk about—namely, those composed of bird-line archosaurs, the feathered dinosaurs, the flying dinosaurs, and the crown dinosaurs—instead of referring to all of them as "Aves." Our proposed taxonomy also fixes the clade references for six additional names—"Carinatae," "Ornithurae," "Palaeognathae," "Tinamidae," "Ratitae" and "Neognathae"—that are no less than a century old, as well as two newer ones—"Galloanserae" and "Neoaves."

Keywords

Birds, taxonomy, nomenclature, feathers, flight, phylogeny, systematics, evolution, node-based, stem-based, apomorphy-based, phylogenetic definitions, *PhyloCode*, taxon names, Reptilia, Archosauria, Panaves, Dinosauria, Theropoda, Coelurosauria, Avifilopluma, Maniraptora, Avialae, Carinatae, Ornithurae, Aves, Palaeognathae, Tinamidae, Ratitae, Neognathae, Galloanserae, Neoaves

Introduction

What is a bird? This question was raised in the latter half of the 19th century by the discovery of *Archaeopteryx* fossils in late Jurassic sediments from southern Germany. The ensuing debate focused on where to "draw the line" between reptiles (in the paraphyletic sense) and birds. An analogous debate has resurfaced with the equally spectacular fossils of feathered dinosaurs recently discovered in early Cretaceous deposits of northeastern China. But in the modern context, in which higher taxa are conceptualized as clades, the question has been reformulated as: Which clade is to be designated by the taxon name "Aves"? In the recent literature, four different answers are commonly given to this question. That is to say, the name "Aves" (hereafter equiv-

alent to "birds") is used, either explicitly or implicitly, for at least four different, albeit internested, clades. Thus, "Aves" has been used for (1) the clade of *feathered dinosaurs* (e.g., Paul 1988; Lee and Spencer 1997), (2) the clade of *flying dinosaurs* (e.g., Ji and Ji 2001), (3) the clade stemming from the most recent common ancestor of *Archaeopteryx* and crown dinosaurs, hereafter referred to as the "*Archaeopteryx* node" (e.g., Padian and Chiappe 1998), and (4) the clade of *crown dinosaurs* (e.g., Perle et al. 1993). The name has occasionally been used in other ways: Patterson (1993), for example, applied "Aves" to the entire clade of bird-line Archosauria, while Thulborn (1975) applied it to only part of that clade, namely, Theropoda. Nevertheless, the four alternatives listed above continue to be the most popular uses of the name "Aves."

Even with all the remarkable fossils described in the past two decades, there are still sizable morphological and temporal gaps among these four internested clades. Nevertheless, even experts who are well aware of the differences among them have confused these clades. It has been argued, for example, that the smaller Eichstätt and larger London *Archaeopteryx* specimens represent different species (Martin 1991). The chain of inference on which this conclusion is based begins with the observations that both specimens have a full set of tail feathers (rectrices). Birds (i.e., Aves), however, as represented by extant species, do not have completely developed rectrices until they are fully grown. Because *Archaeopteryx* is considered a bird, it is inferred to have grown and developed in much the same way as other birds. It then follows that the presence of a full set of rectrices indicates that both a small and a large species have mistakenly been included in what is usually thought to be a single species, *Archaeopteryx lithographica* (e.g., Houck et al. 1990).

That inference would be justified if *Archaeopteryx* could be shown to be part of the crown clade, in which case it could reasonably be expected to have grown and developed like a living bird. But *Archaeopteryx* existed many tens of millions of years before the ancestor of the dinosaur crown (Gauthier 1986). And it may be that the temporal association between skeletal maturity and tail plumage development had itself evolved during the course of phylogeny, and along with it, the ontogeny of flight. In other words, it may have been the case that an individual *Archaeopteryx* with fully developed rectrices was able to fly long before it reached maximum adult size. That is quite unlike the case for species in the crown clade, in which flight capability and maximum size are tightly coupled in ontogeny, as Martin (1991) observed.

This class of errors—the "unjustified inferences" of de Queiroz and Gauthier (1992)—is common in the literature (for example, flight ability = adult size in Enantiornithes; Sereno and Rao 1992), particularly in cases where taxa in question are species. In Quaternary studies, for example, attributes observed in the crowns of extant species are regularly presumed to extend unchanged down into their stems (e.g., Campbell 1980; Van Devender and Bradley 1994). In the absence of additional information (such as correlated characters), parsimoniously justified, or "safe," inferences are those for which the taxon inferred to have possessed a particular trait is descended from an ancestral node whose primary subclades can be observed to possess that trait. For example, extinct pseudodontorns, which are thought to be pelecaniforms (Olson 1985) and thus part of the crown clade, can safely be inferred to have developed rectrices after they were fully grown. That inference follows from the observation that tail feather development and skeletal maturity are closely linked in all living members of the crown clade that develop rectrices, including other pelecaniforms. But without additional evidence such an inference cannot be extended to taxa—including *Ichthyornis, Confuciusornis* and *Archaeopteryx*—that diverged before the most recent common ancestor of taxa observed to possess that trait.

Such confusions are closely tied to taxonomy in that they involve the reference of the name "Aves." In this context, it seems relevant that part of scientific progress involves making distinctions not made previously, commonly accompanied by the introduction of terms reflecting those distinctions (e.g., Gould and Vrba 1982). Given that terminological precision is an ever-increasing phenomenon, some level of imprecision in scientific terminology is inescapable. It is nevertheless clear that, at this point in the history of taxonomy, there is little benefit in continuing to tolerate the current level of imprecision in the meaning of "Aves." So it seems that for the sake of effective communication—the ultimate goal of biological tax-

onomy—comparative biologists face an important (if arbitrary) decision: Which of the four currently popular references should be associated with the name "Aves"? Given that "Aves" has been applied to all four of these clades, we take it for granted that all four are worthy of being named. Nevertheless, even among those who adopt phylogenetic nomenclature, precisely which clade should be called "Aves" is currently a matter of debate (Gauthier 1986; Chiappe 1991, 1995, 1996; Perle et al. 1993, 1994; Chiappe and Calvo 1994; Holtz 1994, 1996; Lee 1996; Lee and Spencer 1997; Padian 1998a, 1998b; Padian and Chiappe 1998; Sereno 1998, 1999a, 1999b; Gauthier and Gall 2001).

In this paper we examine different classes of definitions as they bear on naming the four clades in question and the reference of the name "Aves." We first explore some general issues concerning the properties of three classes of phylogenetic definitions. These include how names can be tied to clades either through composition or characters, as well as the relationships between Hennig's (1965) "total" and "crown" clades compared to our "stem" and "node" clades. We then describe how taxa are to be referred to clades whose names are defined using different classes of phylogenetic definitions, and discuss perceived problems with apomorphy-based definitions. Then we address the issue of whether and in what sense taxon names have traditional meanings. We conclude with an examination of the consequences of restricting "Aves" to each of the four clades to which it is most commonly applied, and propose a novel taxonomy that resolves current nomenclatural ambiguity.

Properties of Different Classes of Phylogenetic Definitions

Classes of phylogenetic definitions based on composition and characters

To appreciate the consequences of attaching "Aves" to different clades, we must first consider the different ways in which names can be tied to clades. This can be accomplished in one of two general approaches: (1) by specifying an ancestor (and thus the clade stemming from it) through reference to a particular apomorphy, or (2) by specifying an ancestor through reference to particular descendant taxa. De Queiroz and Gauthier (1990) described three methods for defining taxon names conforming to these two general approaches. Apomorphy-based definitions specify clades in the first way; namely, "Aves" is defined as the clade stemming from the ancestor in which a particular character first evolved. Node- and stem-based definitions specify clades in the second way, either: (1) in the case of node-based definitions, by defining a name (for example, "Aves") as referring to the clade stemming from the most recent common ancestor of specified descendants (= internal "reference taxa" of Sereno 1999a, or "specifiers" of Cantino and de Queiroz 2000), which must necessarily share a single most recent common ancestor at some point in their genealogies; or (2) in the case of stem-based definitions, by defining a name (such as "Aves") as referring to the clade composed of a particular descendant or descendants (internal specifier), as well as all other taxa that share a more recent common ancestor with that descendant(s) than they do with some other taxon or taxa (external specifiers). Selecting one kind of definition rather than another amounts to applying that name to one clade rather than to another. This does not mean that clades described by node- or stem- or apomorphy-based definitions cannot be the same in theory; for example, it is possible that the origin of an apomorphy coincides with the initial divergence of one lineage from another. On the other hand, the origins of apomorphies and lineages need not, and usually will not, be coincident.

The distinctions made by these classes of definitions are obvious when they involve clades that differ markedly in inclusiveness (and therefore in secondary properties such as time and place of origin [de Queiroz and Gauthier 1990; Rowe and Gauthier 1992]). Consider, for example, the differences among the clade stemming from the most recent common ancestor of *Passer* and *Struthio* (= node based), the clade composed of *Passer* and *Struthio* plus all archosaurs more closely related to them than to *Crocodylus* (= stem-based), and the clade stemming from the first flying dinosaur (= apomorphy-based). But the distinctions obtain even when there are only minor differences in inclusiveness—for example, if *Ichthyornis* is used as

the external reference taxon in the stem-based definition, and an ossified supratendinal bridge on the tibiotarsus is regarded as the defining attribute in the apomorphy-based definition. In these cases, the three definitions might seem to refer to the same taxon in terms of *known* composition (that is, the crown clade). But the taxa specified by those definitions would nevertheless be different in *actual* composition, which would be revealed, for example, if a species were to be discovered that is outside of the crown but more closely related to the crown than is *Ichthyornis* (de Queiroz and Gauthier 1992). The distinction in question holds even when the external specifier of the stem-based definition is the true sister species (or one of its descendants) of the ancestral species terminating at the node specified by the node-based definition, provided that one is willing to allow the "last common ancestor" of the node-based definition to be a part of a species, rather than the entire species (see de Queiroz and Donoghue 1988).

Frost and Kluge (1994) and Sereno (1999a) argued for an alternative view in which nodeand stem-based taxa are regarded as the same, which is correct under the constraint that the last "common ancestor" refers to an entire species. We agree that all organisms in the final unitary lineage segment ancestral to Aves are parts of the same species (sensu de Queiroz 1998, 1999). Nevertheless, that lineage segment could have undergone considerable evolutionary change, in which case its later populations might show clear evidence of their closer relationship to the node and its descendants than to the sister species, or even to earlier parts of their own species (de Queiroz and Donoghue 1990). Moreover, in practice, one can never be sure to have found (for use as a reference point) the true sister species to any ancestral species. That is to say, a stem-based version of "Aves" using Ichthyornis, currently the closest known outgroup to the crown (Norell and Clarke 2001), as an external specifier may not (and probably does not) refer to the same entity as a node-based version of "Aves" attached to the crown. Some specimens currently referred to *Ichthyornis* may prove closer to the crown than to the type specimen of Ichthyornis (Clarke 2000). Or an as yet unknown species may be found that is closer to the crown than is *Ichthyornis*. Indeed, even under the constraint that ancestors are entire species, the only way to force a stem-based definition of "Aves" to refer to the crown clade (and thus be equivalent to a node-based definition) would be to formulate the definition using the following wording or some variant of it: "the clade including Passer and Struthio [for example] and all taxa more closely related to Passer and Struthio than to the sister species of their most recent common ancestor." Although technically stem-based, this definition incorporates elements of a node-based definition in referring to the most recent common ancestor of two descendant taxa. Moreover, it has the undesirable property of using a hypothetical rather than a known taxon as an external specifier, thus severely compromising its practical utility.

Crown and total clades are kinds of node and stem clades

Some authors (e.g., Patterson 1994) have confused stem- and node-based definitions of taxon names with Hennig's (1965, 1966, 1981) distinction between clades that Jefferies (1979) termed "total groups" and "crown groups." (The word "total" is an unfortunate choice if it implies that other clades are somehow incomplete.) "Total groups" are only a subset of the stems that can be designated by stem-based definitions, and "crown groups" are only a subset of the nodes that can be designated by node-based definitions (de Queiroz and Gauthier 1990). A node-based definition can as easily describe an entirely extinct clade, or one in which only one of the two primary subclades has survived to the present time, as one in which both primary subclades are represented by extant organisms. Likewise, a stem-based definition can describe either an entirely extinct clade, or a clade that contains only part of the stem of a crown, or a clade that contains the entire stem of a crown.

Moreover, just as stem- and node-based definitions are both valid, so are total-clade and crown-clade definitions in that both explicitly tie names to clades (de Queiroz and Gauthier 1990, 1992, 1994; Rowe and Gauthier 1992; Bryant 1994, Lee 1996, Sereno 1998, 1999a, 1999b). Thus, "total groups" are not the only clades, nor are they (contrary to the opinion of Smith [1994:97]) the only ones worthy of naming. Systematists may have good reasons—such as con-

flicting data or weak support—for naming some clades and not others, but such complications are hardly unique to crown-based names alone. Moreover, there are good reasons for naming both crown and total clades (de Queiroz and Gauthier 1992). Thus, although we are sympathetic with Patterson's (1993) view that applying the name "Aves" to the total stem clade containing *Struthio* and *Passer* but not *Crocodylus* might force ornithologists to confront the deep history of the lineage leading to the crown, that is not to say that the crown clade should go unnamed. Indeed, there may be equally compelling reasons for using "Aves" for the crown, which is the focus of so much biological research.

Given these differences, it should be clear that the three classes of phylogenetic definitions are not alternative ways of naming the same thing. On the other hand, they can be thought of as alternative ways of defining the same name; which is to say that "Aves" can be tied to different taxa using node-based, stem-based or apomorphy-based definitions. As it now stands, all three types of definition have, in effect, already been proposed: node-based for the "Archaeopteryx node" and the "crown clade," stem-based in the case of the "total clade," and apomorphy-based for "feathers" and "flight." A more complete understanding of the effects of fixing the name "Aves" to just one of these clades requires consideration of some consequences for taxon assignment attendant to the different classes of phylogenetic definitions.

Referring species to clades named using different classes of phylogenetic definitions. Each of the three classes of phylogenetic definitions has certain properties that dictate how its associated clades are to be diagnosed, thereby determining how species and organisms can be referred to them. These properties are necessary consequences of the particular kind of definitions chosen, so they must be kept in mind when using different definitions to avoid unjustified referrals of species and organisms to the clades specified by them. Homoplasy and incomplete data present general challenges for taxon reference; these issues will be discussed separately (see below under "Ambiguity from missing data" and "Ambiguity from homoplasy"). To simplify the rest of the discussion in this section, we will assume absence of homoplasy.

Node-based definitions. Although a node is diagnosed by the apomorphies arising on its subtending branch, possession of those apomorphies is insufficient for referral of specimens to the clade stemming from that node. Consider, for example, a node-based definition of "Aves" for the dinosaur crown. That clade may possess synapomorphies relative to various outgroups, including *Ichthyornis*. Nevertheless, a given fossil can only be referred to that "Aves" (node) if it also possesses at least one apomorphy of one of the crown's primary total subclades (i.e., there must also be evidence that the fossil is part of total clade versions of either "Palaeognathae" or "Neognathae"). Possession of crown-clade apomorphies alone is insufficient for referral to the crown because they could have arisen within the ancestral lineage after its divergence from *Ichthyornis* but before its divergence into palaeognath and neognath total subclades. Consequently, reference of a particular fossil to a node-based clade requires evidence of membership in a subclade. A corollary of this conclusion is that the common ancestor from which a node-based clade originates, which logically belongs to that clade but not to any of its subclades (Hennig 1966), can never be definitely referred to its own clade on the basis of character evidence alone.

Stem-based definitions. At least one apomorphy is sufficient when referring specimens to stem-based clades. That apomorphy could be any one of those arising in the stem lineage subsequent to its divergence from its sister group. Thus, in Patterson's (1993) stem-based version of "Aves," any one of the apomorphies of bird-line archosaurs would do. On the other hand, as is generally the case with stem-based names, one might not be able to assign the earliest organisms of the stem lineage; that is, those existing before any apomorphies arose—or became fixed—to such a stem-based version of "Aves." A consequence of this observation, in conjunction with the discussion above, is that these organisms would not even be referable to the larger node-based clade of which this "Aves" is a primary total subclade (that is, a crown version of "Archosauria").

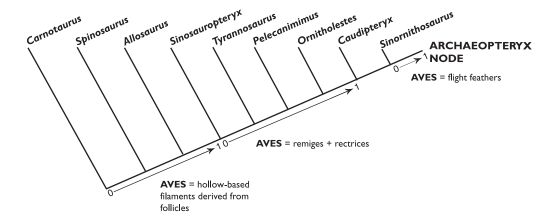


Figure 1. Depending on the definition of "feather," an apomorphy-based definition of "Aves" could be tied to one of three internested clades of theropod dinosaurs. Although the clade referred to by the name "Aves" in each instance is unambiguous (e.g., "Aves" = all theropods with remiges and rectrices), name application may not always be clear (for example, missing data render the avian status of tyrannosaurs uncertain by this definition). No system of taxonomy that depends on data for taxon assignment can escape the consequences of imperfect knowledge. Cladogram depicting phylogenetic relationships among select theropod dinosaurs modified from Sereno (1999b) and Norell et al. (2001).

Apomorphy-based definitions. Possession of the definitive apomorphy is all that is required for taxon reference in the case of apomorphy-based definitions. If "Aves" is defined with respect to "feathers," for example, then any organism possessing this definitive apomorphy would be part of Aves. That being said, the Chinese feathered dinosaurs remind us just how wrong we can be when equating a particular node (that is, the *Archaeopteryx* node) with the origin of a particular apomorphy. "[F]eathers, wings, and the capacity for flight" may have always been regarded as "key" avian characters (Sereno 1999a:332), but treating these characters as apomorphies of a single node was possible only because of the notorious incompleteness of the fossil record. In retrospect, it seems especially fortuitous that *Archaeopteryx*—only the second Mesozoic coelurosaur discovered—was fossilized in an exceptional depositional environment that preserved feather impressions. And for more than 100 years feathers have been regarded as the "key" apomorphy distinguishing the *Archaeopteryx* node from more distantly related dinosaurs (Lee and Spencer 1997). Nevertheless, new theropod fossils from the Liaoning lagerstätten (Chen et al. 1998; Ji et al. 1998, 2001; Xu, Tang et al. 1999; Xu, Wang et al. 1999) now reveal that feathers diagnose a far more inclusive clade than that stemming from the *Archaeopteryx* node.

Perceived problems with apomorphy-based definitions

Throughout the history of taxonomy, organismal traits have played central roles in taxon concepts, and by implication, the meanings of taxon names (Stevens 1994). Nevertheless, apomorphy-based definitions have been widely criticized, and it has been argued that this class of definitions should be avoided (Rowe and Gauthier 1992; Bryant 1994; Schander and Thollesson 1995; Padian et al. 1999; Sereno 1999a). Despite our sympathy with some of the problems raised, none is a compelling reason for avoiding apomorphy-based definitions.

There are three reasons why apomorphy-based definitions are thought to pose special obstacles (Sereno 1999a): (1) they are prone to ambiguities stemming from ambiguous character concepts (such as the precise definition of "feather"); (2) they are subject to ambiguities involving missing data (for example, integumentary structures are preserved in only a tiny fraction of fossil dinosaurs); and (3) they are vulnerable to problems caused by homoplasy (such as parallelism, convergence and reversal) or obliteration through subsequent evolutionary trans-

formation. One might be tempted to regard these as unique problems for apomorphy-based definitions to which node- and stem-based definitions are somehow immune. But as long as characters play a role in taxon assignment, systematists will face practical problems implementing any kind of definition. Indeed, the precise phylogenetic relationships of some specimens may simply not be determinable, regardless of how taxon names might be defined. Such problems are, however, general to phylogenetic inference, rather than being specific to phylogenetic taxonomy per se; regardless of taxonomic philosophy, there can be no guarantee that the available data will be sufficient to enable reference of every specimen to its exact phylogenetic position. Moreover, because phylogenetic taxonomy, in contrast with traditional practice, rejects the idea that certain categorical ranks are mandatory, specimens and taxa need not be referred to taxa beyond what the data will support (de Queiroz and Gauthier 1992).

Ambiguity in character concepts. Systematists might agree on all details of tree topology and loci of character transformation yet still disagree about which clade to name "Aves" because they cannot agree on which state in a transformation series should be called (for example) a "feather" (Figure 1). Systematists would be ill-advised to become too concerned with the "essence" or "true nature" of any apomorphy, but that still leaves the problem of arbitrarily deciding which state in a transformation series is to be designated by the name "feather."

In a situation analogous to the muddled state of affairs with the taxon name "Aves," the homologue name "feather" can be associated with several different morphologies (Figure 1). Evolutionists should have expected that such a suite of novel morphologies summarized by the term "feather" might not have arisen in a single macromutation. And from what is known of the ontogeny and phylogeny of feathers, that certainly seems to be the case (Prum 1999; Ji et al. 2001). Thus, short, hollow-based filaments occur in the basal coelurosaur Sinosauropteryx (Currie and Norell 1998) as well as in dromaeosaurs (Prum 1999; Ji et al. 2001). Branched appendages with long filaments (barbs) emanating from a short base (calamus) are also known in dromaeosaurs (Xu et al. 2001). Bipinnately branched filaments (barbs) emanating from a long, hollow, central rachis have been observed in oviraptors (Ji et al. 1998), dromaeosaurs (Ji et al. 2001; Xu et al. 2001), and Archaeopteryx (Owen 1863; Heilmann 1927). And those appendages may bear open vanes, as on the shank of Archaeopteryx, or closed vanes (and thus presumably barbules with hooked distal pennulae), as do those lying atop the second finger and trailing edge of the ulna and fanning from the tail in both Archaeopteryx and Caudipteryx (Ji et al. 1998; Wagner and Gauthier 1999). Finally, some closed-vaned pennaceous appendages may have stout, curved shafts—furrowed along the ventral side—supporting asymmetrically developed vanes, as do those borne on the hand of Archaeopteryx and many members of the dinosaur crown (Norberg 1985; Rietschel 1985).

Hollow-based, unbranched filaments—structures that Prum (1999) homologized with the thin epidermal sheaths that temporarily enclose developing feathers (pin feathers) in the crown clade—appear in basal coelurosaurs (Figure 1; Currie and Norell 1998). Unbranched filaments, as well as natal down and pennaceous feathers with open and closed vanes (= remiges and rectrices) appear in basal maniraptors (Figure 1; Ji et al. 1998, 2001; Xu et al. 2001). Fully functional flight feathers came later, originating after basal raptors such as *Sinornithosaurus*, but before the *Archaeopteryx*-node (Figure 1). In our estimation, any one of these morphologies could quite justifiably be called a "feather." Different workers may prefer one definition, such as the inferred presence of the feather follicle itself (Prum 1999), or another, such as the more readily observable herringbone pattern of pennaceous feathers (for example, *Caudipteryx*; Ji et al. 1998). The choice is arbitrary.

Of course, identifying feathers in fossils is sure to incite controversy. Ruben and Jones (2000), for example, expressed doubt that the structures identified as feathers in *Sinosauropteryx* (Currie and Norell 1998) were feathers, or indeed, that they were even of epidermal origin. Moreover, Jones, Ruben and co-workers (2000) claimed that the epidermal appendages in

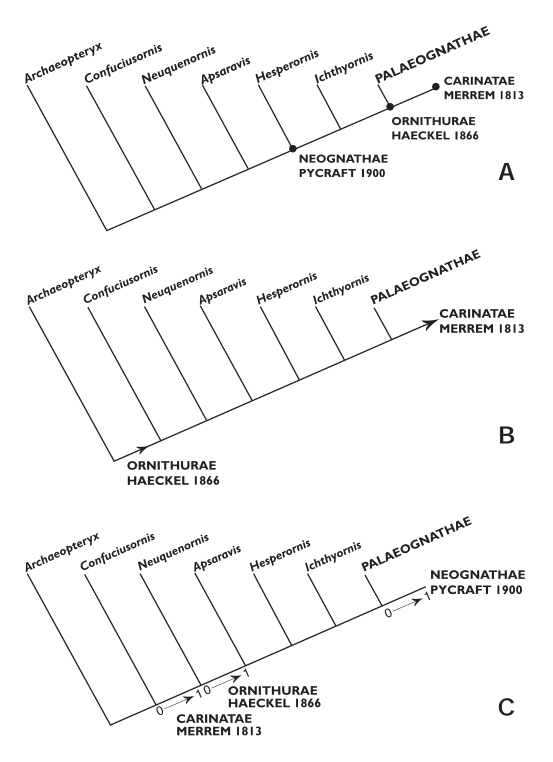


Figure 2. Three different versions of "Carinatae" (Merrem 1813), "Ornithurae" (Haeckel 1866), and "Neognathae" (Pycraft 1900) consistent with the original author's use. **A.** Node-composition-based versions using original set of included taxa as internal specifiers. **B.** Stem-composition-based versions using original set of included and excluded taxa as internal and external specifiers, respectively. **C.** Apomorphycharacter-based versions using eponymous synapomorphies as specifiers. Cladogram depicting phylogenetic relationships among select flying dinosaurs modified from Clarke (2001).

Longisquama are pennaceous feathers (or their "precursors"), while Reisz and Sues (2000) argued that although these have some superficially feather-like attributes—such as a thickened central axis with ridges diverging from it—they are nonetheless typical reptilian scales. Controversies of this sort stem from conflicting primary homology assessments, exacerbated by incomplete data, rather than the definition of the homologue name "feather" per se. The structures in question could be called feathers if they are found to satisfy criteria for primary and secondary homology assessment (Patterson 1982; de Pinna 1992), thus making these taxa part of a feather-based "Aves." But if those structures fail to satisfy homology criteria, then there would be no basis for calling them feathers, and no basis for referral of the taxa bearing them to "Aves." And finally, if the data are too poor or contradictory to enable a decision one way or the other, then homology of these structures, and referral of the taxa that bear them, would likewise be uncertain (barring other evidence of relationships).

Ambiguity from missing data. Another potential problem with apomorphy-based definitions is that they could be based on attributes that are unlikely to leave a trace in the fossil record, and thus be inapplicable to many extinct taxa (Lee 1998). If "Neoaves," for example, was defined in terms of the unique 5-codon deletion in the RAG-1 nuclear gene identified by Groth and Barrowclough (1999), one might never be able to refer certain fossil members to that clade. Nevertheless, if any "soft" (less preservable) character can contain phylogenetic information, the absence of these data in fossils seems an inadequate justification for rejecting their use in apomorphy-based definitions. Indeed, feathers have commonly been regarded as a defining avian character despite their low preservation potential. And even the "hard" (more preservable) apomorphies favored by paleontologists are not preserved in every fossil. For example, the only clearly diagnostic character of *Archaeopteryx*—a foramen passing through the ischium—can only be scored in one (the London specimen) among the seven skeletons currently referred to that taxon. Likewise, the distinctive palatal morphologies that characterize major avian clades (Huxley 1867) cannot be observed in most fossil specimens.

Related to the problem of missing data is that the precise point of origin of any character will always be beset with some uncertainty. For example, if feathers are defined as hollow-based, follicular epidermal appendages, we know only that they arose after the divergence of *Carnotaurus* and before that of *Sinosauropteryx* (Figure 1). Furthermore, although systematists often distinguish between "ambiguous" and "unambiguous" synapomorphies, even the latter are subject to this problem (see below). For example, although considered an unambiguous synapomorphy of the crown clade (Gauthier 1986), an ossified supratendinal bridge on the tibiotarsus can only be inferred to have arisen sometime after the divergence of *Ichthyornis* but before the origin of the crown (Figure 2).

Even with "complete" knowledge one could still be left with making arbitrary divisions along a continuum for characters exhibiting continuous variation or for those exhibiting discontinuous variation and arising as population-level polymorphisms. The latter case is nicely illustrated by Confuciusornis, in which a few specimens possess a slight ridge on the posteroventral midline of the sternum, while most others do not (Chiappe et al. 1999). This ridge hardly matches the prominent keel (carina)—the site of origin for hypertrophied pectoral muscles—that originated along the branch subtending the Neuquenornis node (Chiappe and Calvo 1994; Figure 2). Nevertheless, Chiappe and co-workers (1999) regarded the feature in some specimens of *Confuciusornis* as a potential homologue, even if only in an incipient form, of the carina seen in the clade stemming from the *Neuquenornis* node. But even if one is willing to accept the condition seen in *Confuciusornis* as a "carina," there is still ambiguity regarding criteria for considering a character as present or absent, which affects inclusion of that taxon in an apomorphy-based version of "Carinatae." Thus, if one elects to view a character as being absent until it becomes fixed in a lineage (Nixon and Wheeler 1992), then Confuciusornis would not be part of "Carinatae." Alternatively, one could treat the polymorphism as missing data (Kluge 1995), in which case taxon assignment will be uncertain. Finally, if one considers a character to

be present if any part of a population has it (Kron and Judd 1990), then *Confuciusornis* would be part of "Carinatae."

Despite these potential problems with apomorphy-based definitions, stem- and node-based definitions are no less subject to ambiguity in specimen referral, because that activity depends on character data, which may be fallible for all the reasons outlined above. For example, under a stem-based version of "Carinatae"—defined with respect to (keel-less) *Archaeopteryx* as an external reference taxon—at least some *Confuciusornis* specimens could mistakenly be excluded (for example, if the sternal keel was the only informative attribute or the sternum the only fossil remains). Similarly, under a node-based version of "Carinatae"—defined using a member of the crown and *Confuciusornis* as internal reference taxa—those same specimens might once again be mistakenly excluded. Even without the complications resulting from polymorphism and continuous variation, the earliest members of a stem lineage might have none of the apomorphies arising subsequently in their descendants, so they will often be mistakenly excluded from the stem-based clades of which they are parts and, thereby, from the node-based taxon composed of those stems.

Evolutionists are obliged to accept that common ancestry relations hold even when they might never be discovered in particular instances (Ghiselin 1984). Thus, all flying dinosaurs arising from an ancestor with a keeled sternum would, by definition, be part of a "Carinatae" based on that apomorphy. Whether one can correctly refer specimens in every case is beside the point, as no phylogenetic definition or, for that matter, any other nomenclatural system, can guarantee that.

Ambiguity from homoplasy. Some authors think that homoplasy places a special burden on apomorphy-based definitions (Bryant 1994; Schander and Thollesson 1995; Sereno 1998, 1999a). Suppose, for example, that "Aves" is defined using the flight apomorphy and that the capacity for aerial flight arose more than once in dinosaurs. Some have mistakenly interpreted this to mean, using this example, that "Aves" would be polyphyletic (Bryant 1994). Others have suggested that it would generate ambiguity (Sereno 1999a); namely, if two dinosaur clades evolved flight, which one should be called "Aves"? These problems are perceived rather than real, though they are related to a genuine problem. As originally proposed by de Queiroz and Gauthier (1990:310), an apomorphy-based definition is used to associate a name with "the clade stemming from the *first* ancestor to possess a particular synapomorphy" (emphasis added). The qualifier "first" is needed because there will often be many ancestors possessing the character. But it also ensures that the name will neither refer to a polyphyletic taxon nor be ambiguous. Thus, two origins of flight in dinosaurs would not result in "Aves" referring to a diphyletic taxon, because only one of two clades would be properly referred to by that name (that is, the one that first evolved flight). And for the same reason, the reference of "Aves" would not be ambiguous. Of course, it might not always be possible to infer the relative times of origin of states that arose homoplastically, but that is a problem with the data rather than the definition.

Nevertheless, there is a potential problem with apomorphy-based definitions as originally stated by de Queiroz and Gauthier (1990)—namely, that such definitions could result in the application of a name to an unintended clade. For example, the name "Aves" might end up referring to a small early clade of extinct flying dinosaurs rather than to the large clade that includes all living taxa currently considered birds. But this problem can be remedied by adding a clause to the definition specifying that the "flight" apomorphy must be homologous to flight in a particular descendant, such as *Passer*. If flight was then seen to have arisen twice in dinosaurs, then the clade of flying dinosaurs including *Passer* would be "Aves," and the other clade of flying dinosaurs would not, even if it evolved flight earlier.

It should be clear that if some "flying dinosaurs" were to lose the capacity for aerial flight, they would still be part of Aves (Gauthier 1986). That is to say, even in the case of apomorphy-based definitions, the possession of characters per se is secondary to common ancestry rela-

tionships (de Queiroz and Gauthier 1990). To paraphrase Darwin (1859), who was himself paraphrasing Linnaeus, a canary has feathers because it is a bird, rather than being a bird because it has feathers.

In conclusion, incomplete, variable and conflicting data pose genuine problems when referring specimens to taxa of any kind. Apomorphy-based definitions are no more or less prone to such problems than are node- or stem-based definitions. Moreover, in terms of communicating ideas about common ancestry relationships, none of the three classes of phylogenetic definitions is any less clear, stable or universal than is any other (Lee 1996). Finally, apomorphy-based definitions should not be avoided, if for no other reason that many of the clades people want to talk about and name are conceptualized in terms of apomorphies.

Do Taxon Names Have Traditional Meanings?

For reasons described below (see "Aves = crown dinosaurs"), de Queiroz and Gauthier (1992; see also Gauthier 1986) proposed that widely used names be tied to crown clades. This proposal has been criticized for seeming to violate traditional associations between names and particular sets of taxa (Chiappe 1991) or particular apomorphies (Lee and Spencer 1997). But as Sereno (1999a) observed, it has seldom been clear whether a given taxon name was to be associated with a particular organismal trait or a specific set of included species. For example, when Huxley (1867:424) explained that Merrem's (1813) "Carinatae" included "all existing birds, except the Ratitae," in which the "sternum possesses a keel," did he conceptualize the taxon in terms of its composition or its characters (or both)? And if he had composition in mind, is that composition to be interpreted as node-based (that is, only in terms of included taxa) or stembased (that is, also considering the excluded taxa)? Uncertainty on these points undermines arguments in favor of maintaining "traditional use" advanced by some authors (Gauthier 1986; Benton 1990, 1999; Chiappe 1991, 1995, 1996; Chiappe and Calvo 1994; Lee 1996; Lee and Spencer 1997; Padian 1998a; Padian and Chiappe 1998; Sereno 1998, 1999a, 1999b).

Figure 2 illustrates this point on a cladogram for selected basal flying dinosaurs. Three taxon names—"Carinatae" (Merrem 1813), "Ornithurae" (Haeckel 1866), and "Neognathae" (Pycraft 1900)—are of special relevance because the names themselves (which translate as "keeled," "bird tail," and "new jaw," respectively) indicate that the original authors had specific characters (now thought to be apomorphies) in mind when they coined them. And it is no less clear from their lists of included taxa, and sometimes excluded taxa, that these authors also applied each name to a particular set of taxa. In Figure 2 each name is applied to three different clades in the context of a current phylogenetic hypothesis: the first is a node-composition-based version using the original set of included taxa as internal specifiers (Figure 2A); the second is a stem-composition-based version using the original set of included and excluded taxa as internal and external specifiers, respectively (Fig. 2B); and the third is an apomorphy-character-based version using the eponymic synapomorphies as specifiers (Fig. 2C).

Note that node-composition, stem-composition and apomorphy-character versions of the same names often differ markedly in terms of the clades to which they refer (Figure 2). Although these different interpretations generally coincided when the names were first proposed, they tended to diverge in light of new discoveries. For example, when Merrem (1813) coined "Carinatae," extant tinamous and extinct crown outgroups with keeled sterna were unknown to him, while the only keel-less birds that Merrem knew of were members of the "anomalous Ostrich tribe," which he called "Ratitae" (= Struthio of Linnaeus 1758 = ostriches, cassowaries and rheas). Thus, "Carinatae" first referred to all then-known members of the crown except Ratitae (Figure 2A and B). Because tinamous were not included in "Carinatae" by Merrem, and because they now appear closer to ratites than to other members of the crown (Cracraft and Clarke 2001), stem- and node-composition-based versions of "Carinatae" now refer to all or part of the sister group to the ratite-tinamid stem clade. Subsequent finds—tinamous, Ichthyornis, Neuquenornis—and phylogenetic analyses (Chiappe and Calvo 1994) led eventually to the conclusion that a keeled sternum applied to a far more inclusive set of taxa than Merrem originally

envisioned (Figure 2C). Consequently, an apomorphy-character version of "Carinatae" now refers to a clade that arose sometime between the *Confuciusornis* and *Neuquenornis* nodes. Therefore, Merrem's "Carinatae" would now be associated with very different clades depending on whether one retained its association with his original set of included and excluded taxa (Figure 2A and B), or with the character of a keeled sternum (Figure 2C).

Likewise, the stem-composition, node-composition, and apomorphy-character versions of "Ornithurae" have also diverged over time. When Haeckel (1866) coined "Ornithurae," the only known flying dinosaurs were *Archaeopteryx* and members of the crown. Subsequent discoveries of noncrown fossils that were closer than *Archaeopteryx* to the crown (such as *Ichthyornis* [Marsh 1872a]) posed several possible associations for that name. Thus, should "Ornithurae" be used for the clade with a "bird tail," as the name suggests (Figure 2C; see *Ornithurae* under "Proposed Taxonomy," below, for a discussion of "bird tail")? Or should it apply to all flying dinosaurs other than *Archaeopteryx*, the only taxon that Haeckel explicitly excluded from "Ornithurae" (Figure 2B)? Or should it refer to the crown clade, the only flying dinosaurs that he explicitly included in "Ornithurae" (Figure 2A)? Any one of these alternatives would be consistent with Haeckel's (1866) original use of that name.

Finally, an apomorphy-character version (Figure 2C) of Pycraft's (1900) "Neognathae" refers to the clade stemming from the first ancestor to possess Huxley's (1867) "pterygo-palatine...arthrosis" (Pycraft 1900:171), "a point of great difference between Palaeo- and Neognathine skulls" (Pycraft 1900:208). But Pycraft included Ichthyornis and Hesperornis in his "Neognathae" (Figure 2A), even though "Huxley's arthrosis" could not be determined in these fossils. Pycraft, like most ornithologists of that era, seems to have been swayed by Fürbringer's (1888) conclusion that Hesperornis and Ichthyornis were respectively allied to foot-propelled divers and shore birds, implying that they were parts of what is now termed the crown. Although it was acknowledged from the outset that these "toothed birds" were very primitive (Marsh 1880), ornithologists did not begin to take seriously the idea that *Ichthyornis* and *Hes*perornis were outside of the crown until the latter half of the 20th century (Wetmore 1956; Harrison 1973; Martin 1984). In this light, regarding the issue of composition versus characters, should "Neognathae" refer to the clade that includes *Ichthyornis* and *Hesperornis*, as specified by Pycraft? Or should "Neognathae" apply to the clade diagnosed by Huxley's arthrosis, and thus exclude those taxa? Given Pycraft's aversion to "inaccurate" names (he substituted "Neognathae" for "Carinatae" because some members of the latter taxon lack a keel), we think it likely that he would have excluded from "Neognathae" taxa (such as *Hesperornis* [Elzanowski 1991]) that could be shown to lack Huxley's arthrosis. Unfortunately, inferences about the intentions of original authors are not always so straightforward.

These examples indicate that node-composition, stem-composition, and apomorphy-character interpretations of the same names may differ markedly in terms of their references. For example, the node- and stem-composition versions of "Carinatae" (Figure 2A and B) refer to internested parts of the crown clade (which currently happen to have the same known composition). In the apomorphy-character version (Figure 2C), however, "Carinatae" refers to a much larger clade. Indeed, under the apomorphy-character interpretation, "Carinatae" includes "Ornithurae," just the opposite hierarchical relation among the taxa designated by these names as in the node- and stem-composition versions, in which "Carinatae" is included within "Ornithurae." Likewise, the node-composition version of "Ornithurae" (Figure 2A) refers to the crown, while its stem-composition version refers to a much larger clade that includes all flying dinosaurs other than Archaeopteryx. Finally, the crown is only one part of a node-composition interpretation of "Neognathae" (Figure 2A), whereas an apomorphy-character version of "Neognathae" (Figure 2C) refers to only a part of the crown. And "Neognathae" refers to nothing in the stem-composition version (Figure 2B) because, according to current thought, there is no clade of flying dinosaurs that excludes Palaeognathae but includes the rest of the crown as well as Hesperornis and Ichthyornis (Figure 2B).

Thus, although taxon names do indeed have traditional meanings or references, it is not al-

ways clear how those references are to be interpreted in the face of newly discovered taxa. In particular, it is unclear whether those references are to be interpreted as ideas about composition as opposed to ideas about characters. Thus, it is unclear which "tradition" should be honored first, the implied association between a name and a set of taxa, or the implied association between a name and a character or set of characters, and the issue is further complicated by the finer distinctions (that is, nodes compared to stems) of contemporary phylogenetic theory. Many early ornithologists may have conceptualized taxa as philosophical classes (that is, as groups defined by the attributes of their members), in which case associations between names and characters might be argued to have primacy (Pycraft 1900). There are, however, examples involving post-Darwinian authors (if not pre-Darwinian as well) that cast doubt on that interpretation. Huxley (1867), for example, was well aware that certain flightless birds lacked keeled sterna, yet he still considered them members of "Carinatae." In any case, different interpretations of the traditional meanings of taxon names are clearly possible.

In light of the range of possible interpretations concerning traditional meanings, it is not surprising that ornithologists currently seem divided regarding their preferences for characters versus composition. Sometimes they conserve "traditional" character associations and ignore "traditional" ideas about composition. For example, Martin's (1984) use of "Ornithurae" conserves Haeckel's (1866) association of that name with a group possessing a more bird-like tail than that seen in "Sauriurae" (i.e., Archaeopteryx). But that use disregards Haeckel's (1866) association of "Ornithurae" with a specific composition—at least under the node interpretation—in that Martin included a series of successive outgroups to the crown that were not included by Haeckel. In other cases, "traditional" character associations have been ignored in favor of "traditional" associations regarding composition. In the case of "Palaeognathae," for example, Pycraft's (1900) composition (under the node interpretation) has been conserved (Cracraft 1974; Lee et al. 1997), even though many characters marking an "old jaw" are now known to apply to far more inclusive vertebrate clades. Interestingly, conserving traditional ideas about composition under the node interpretation now seems of paramount concern for many authors (Benton 1990, 1999; Lee 1996; Lee and Spencer 1997; Sereno 1998, 1999a). For example, Chiappe and co-workers (1999) excluded Confuciusornis and Neuquenornis from "Ornithurae" to preserve "traditional" ideas about the composition of that taxon despite the occurrence of several distinctly "bird-like" apomorphies in the tails of those taxa, which could have been used to justify their inclusion within "Ornithurae" (Sereno 1998). (It is also worth noting that "traditional" ideas about composition for Chiappe and his colleagues [1999] refer to the tradition of Fürbringer [1888], Martin [1984] and Chiappe [1996], rather than the original tradition of Haeckel [1866].) Thus, different authors have different ideas concerning how taxon names ought to be applied to best preserve their "traditional" meanings. By now, it should also now be abundantly clear that taxon names defined by apomorphies and those defined based on composition seldom refer to the same clades.

Alternative Definitions of the Taxon Name "Aves"

The importance of the taxon name "Aves" (and its equivalent English vernacular, "bird") is indicated by the fact that the *Zoological Record* lists no less than 206,238 records for that name from 1978 to 1999. During that same 21-year period, there are only nine references to "Ornithurae" (Ornithur*) and seven to its junior synonym "Neornithes" (Neornith*; see comments under Ornithurae in "Proposed Taxonomy," below). With a ratio of use of Aves: Ornithurae + Neornithes of 12,890: 1, it seems clear that as a practical matter the distinctions between the clade named "Aves" and those named either "Ornithurae" or "Neornithes" are of little relevance to most ornithologists. On the other hand, it seems doubtful that *Archaeopteryx* and other fossils from outside of the dinosaur crown are the intended subjects of most recent publications (e.g., *Avian Development, Avian Molecular Systematics*, etc.). Such imprecision may not be particularly problematic in many instances; one may, for example, intend no more than to describe "avian" development in all species in which it can be studied. If so, it hardly matters whether

Archaeopteryx, or any other fossil whose prehatching development can seldom be studied, is called a "bird."

On the other hand, precision is vital when one is interested in deriving inferences about ancestral states, especially inferences about "soft" characters that are well justified only for the crown clade. Nevertheless, although it may be widely agreed that *Archaeopteryx* is part of Aves, the name "Aves" is often used when describing biological properties that cannot be studied in *Archaeopteryx*, or worse yet, are known not to occur in that taxon. For example, although Zusi (1993:411) considered *Archaeopteryx* and other Mesozoic flying dinosaurs to be birds, Zusi and Livezey (2000) described criteria for determining the ontogenetic stage of "bird skeletons" that apply only to the crown; thus, "[c]hicks are birds within a few days of hatching with fully evident suturae cranii" (Zusi and Livezey 2000:162) Of course, no *Archaeopteryx* specimen is small enough to qualify as a neonate, and most of its cranial sutures remain unfused throughout ontogeny, as is generally the case in basal flying dinosaurs.

To promote effective communication, everyone should use the same names for the same clades (insofar as they agree on the clades). The problem is, although there is broad conformity on many details of Mesozoic theropod phylogeny (Gauthier and Gall 2001), there is little agreement on which clade is to be designated as "Aves." A precise and universally accepted definition of "Aves" is desirable, if for no other reasons than that it facilitates communication and reduces purely semantic disagreements. Granted that traditions naturally lend themselves to that goal, in light of our previous discussion, it is not clear which "tradition" should be universal: ideas about the composition of "Aves," or ideas about its "key" character(s)? If the former, which composition is to be conserved: Linnaeus' (1758) original set of species, all of which belongs to the crown clade, or Haeckel's (1866) subsequent expansion to include *Archaeopteryx*? If it is the latter "tradition," which apomorphy, feathers or flight, is to be conserved? We next consider some consequences of each of the four popular definitions of "Aves."

Aves = Archaeopteryx *node*

A close association between "Aves" and the Archaeopteryx node has been accepted by most biologists. But that association is perhaps the most unsatisfactory alternative, for it seems that apomorphy—feathers, flight or both—lies at the heart of this connection (Lee 1996; Lee and Spencer 1997; Sereno 1998, 1999a), rather than a specific set of taxa per se. This interpretation is supported by the fact that the century-old link between "Aves" and a set of species belonging to the crown clade was so soon abandoned (Haeckel 1866) after the description of the London Archaeopteryx (von Meyer 1861b; Owen 1863), along with the contrasting fates of Compsognathus and Archaeopteryx. The first skeletal remains of these taxa—the Munich Compsognathus (Wagner 1861) and the London Archaeopteryx (von Meyer 1861b)—were described from the lithographic beds of Solnhofen in the same year. And both fossils had a mosaic of ancestral "reptilian" and derived "avian" features (Gauthier and Gall 2001). Indeed, Gegenbaur's (1863) comparisons of the hind limbs of Compsognathus with those of embryonic birds are now regarded as the origin of the "bird-dinosaur" hypothesis (e.g., Rowe in Dingus and Rowe 1998). Nevertheless, Compsognathus has always been classified as a "reptile," while Archaeopteryx ultimately came to be accepted as a bird. This cannot be attributed to a striking morphological gap, for as noted by Romer (1966:166), "[t]he remains [of Archaeopteryx] so closely resemble those of some of the smaller bipedal dinosaurs that they might well have been taken for reptiles were it not for the impressions of feathers." In fact, the Eichstätt Archaeopteryx was initially identified as a juvenile Compsognathus (Ostrom 1985). The keys here are the short arms and (apparently) featherless form of Compsognathus in contrast to the long arms and feathers of Archaeopteryx, suggesting the capability of flight.

One might well want to preserve this "traditional" association between "Aves" and either feathers or flight in dinosaurs. But it must be understood that neither node- nor stem-based definitions can be counted on to tie that name to the clade stemming from the ancestor that first evolved one or the other of these apomorphies. If one chooses, say, *Sinornithosaurus* as an ex-

ternal reference taxon for a stem-based definition, then the avian stem might include some (not yet discovered) dinosaurs from before the origin of flight. And tying the name to the *Archaeopteryx* node—that is, to the last ancestor shared by *Archaeopteryx* and *Gallus*—hardly guarantees that it will refer to all flying dinosaurs, and it certainly does not apply to all feathered dinosaurs. Indeed, the only way to ensure that all feathered or flying dinosaurs, and only feathered or flying dinosaurs, are included in the taxon named "Aves" is to use an apomorphy-based definition for that name.

In our view, the *Archaeopteryx* node is significant primarily from a historical rather than a biological perspective. That is to say, the significance of this node derives from the fact that *Archaeopteryx* was the earliest noncrown dinosaur discovered that clearly possessed feathers and was probably capable of flight. And that historical significance was further reinforced by the fact that for well over 100 years, *Archaeopteryx* held the privileged position of being the most distant relative of the crown known to possess feathers and (probably) flight. Nevertheless, more distantly related feathered dinosaurs were eventually discovered, so that now feathers are no longer diagnostic of the clade stemming from the *Archaeopteryx* node. More distantly related flying dinosaurs may eventually be discovered as well, in which case neither of these "key avian" features will be associated with the *Archaeopteryx* node.

Aves = feathered dinosaurs

As Linnaeus (1758) and many others have observed, birds are bipedal, warm-blooded creatures with feathers and wing-powered flight. But birds are not the only bipedal animals, nor are they the only animals that are warm-blooded, and some birds are incapable of flight. So in terms of "key" or "essential" avian characters, feathers have been central to traditional notions of "Aves," because in the extant biota at least, all 10,000 species of birds, and only birds, possess feathers (Paul 1988; Brush 1996, 2001; Lee and Spencer 1997). There is a certain appeal to using that most remarkable evolutionary novelty—the feather—which appears unique and unreversed in the long history of life, as the basis for an apomorphy-based definition of "Aves." Fossilized feather impressions were, after all, the reason why von Meyer (1861a, 1861b) concluded that the London *Archaeopteryx* was a bird in the first place.

Fossilized skin impressions indicate that a plesiomorphic integument composed of typical archosaur scales was present in the basal neotheropod *Carnotaurus* (Bonaparte et al. 1990), in sauropods (Bonaparte and Powell 1980), and in ornithischians (Osborn 1912). Nevertheless, integumentary impressions are almost unknown among theropods that separated from the avian line between the divergence of scaled *Carnotaurus* and the feathered (according to some definitions) basal coelurosaur *Sinosauropteryx* (see Figure 1). The integumentary structures of extinct theropods that diverged after *Sinosauropteryx* but before *Caudipteryx* are also poorly known. But given that their preserved anatomy is adequate to infer that they are closer to the crown than is *Sinosauropteryx*, they can be safely inferred to belong in a feather-based version of "Aves." Reasoning of this sort is the basis on which *Tyrannosaurus* can be safely inferred, even in the absence of skin impressions, to have had feathers (Figure 1) and thus also to be part of a feather-based "Aves." The same cannot be said for taxa more distantly related than *Sinosauropteryx*. *Allosaurus*, for example, stands as one of the best known Mesozoic theropods, but so long as its integument is unknown (and feathers are unknown in any more distant outgroup), by this definition one would be unable to determine whether it was part of Aves.

On the other hand, if feathers are truly the "key character" for Aves, it is striking that so few authors have been tempted to call *all* of the Chinese feathered dinosaurs birds. Most of those who accept that *Sinosauropteryx* had feathers, or at least protofeathers, do not consider it a bird. And although no one disputes that the basal oviraptor *Caudipteryx* had pennaceous feathers, only those who consider it secondarily flightless entertain the possibility that it was a bird (e.g., Jones, Farlow et al. 2000). Looking back across the last 250 years of avian taxonomy, the reason seems clear enough. As Padian (1998b:729)—who did not consider either *Sinosauropteryx* or *Caudipteryx* to be birds—observed, "[t]hese animals were clearly not...capable of flight." *Con-*

fuciusornis, on the other hand, possesses an unquestioned—if still in many ways plesiomorphic (Norell and Clarke 2001)—flight apparatus, and this taxon is generally regarded as "avian."

Aves = flying dinosaurs?

It should be clear by now that those who want to use the name "Aves" for the clade of flying dinosaurs are really pointing at the "flight" apomorphy (using hypothesized morphological correlates) and not at a node or stem. Thus, when a flighted taxon outside of the *Archaeopteryx* node is eventually found, as we suspect it will, we would be surprised if it failed to supplant *Archaeopteryx* as "the earliest bird" in many minds. Unlike feathers, flight has the advantage of being correlated with modifications in hard anatomy that are more likely to be fossilized. That, in turn, would provide a more broadly applicable basis for taxon assignment to an apomorphy-based version of "Aves" defined with respect to flight. For example, although several modifications to the pectoral appendages that enable *Archaeopteryx* to fly are now known to antedate the origin of flight (Ostrom 1974a, 1974b, 1976, 1997; Gauthier 1986; Gishlick 2001), one flight-related feature that distinguishes *Archaeopteryx* is the enormous size of its forelimbs relative to body size.

Nevertheless, the utility of this definition still depends largely on a gap in the fossil record; there remains a sizable morphological gap between the shorter-armed early dinosaurs and the longer-armed flying dinosaurs that is only partly bridged by the currently known forms. Intermediates will likely be found. Indeed, some participants at the Graves Museum dinosaur symposium held 7–9 April 2000 suggested that *Bambiraptor* (Burnham et al. 2000) may have had arms long enough to support flight. So was *Bambiraptor* a bird? Opinions are sure to vary. And although *Archaeopteryx* is now generally regarded to have been a flyer (Feduccia and Tordoff 1979; Norberg 1985), that point has been a matter of contention over the years (Speakman and Thomson 1994, 1995; Norberg 1995). So although flight has an advantage over feathers in that its morphological correlates have a better potential for preservation, correlations between those features and flight are themselves open to question. As was argued above, however, no definition (phylogenetic or traditional) can be immune to problems with taxon referral.

Aves = crown dinosaurs?

We have previously advocated tying the name "Aves" to the dinosaur crown (Gauthier 1986; de Queiroz and Gauthier 1992). The advantage of a crown clade definition is that it maximizes safe inferences regarding the biology of extinct members of the clade designated by that name. This statement has been misinterpreted as making the claim—which we agree is false—that those inferred "avian" characters are "unambiguous" synapomorphies of the crown relative to all possible outgroups, extinct and extant, known or unknown (Lee 1996; Sereno 1998, 1999a). As pointed out by Lee (1996), those synapomorphies, which are currently unobservable and could remain so in extinct taxa, might well apply to more inclusive clades than the crown alone. Indeed, chains of inference can sometimes be developed to suggest that soft attributes were in fact present in extinct outgroups (for example, amniote egg in diadectomorphs; Lee and Spencer 1997). On the other hand, some characters (such as molecular sequences) may not lend themselves to such chains of inference. And even when possible, they sometimes require exceptional circumstances of fossilization that rarely occur in nature (for example, behavioral inferences based on finding an oviraptor sitting on its nest [Clark et al. 1999]). Finally, these inferential chains may themselves depend on questionable assumptions about morphological correlates. And even if the assumptions seem reasonable, the inferences are less safe in resting on an extra layer of assumptions (that is, beyond those involved in character optimization based on observed occurrences). All of these points lead to the conclusion that synapomorphy is in practice a relative concept. That is to say, homeothermic endothermy is no less a synapomorphy of the dinosaur crown clade relative to Crocodylia just because it is unclear whether it is synapomorphic of the crown relative to *Ichthyornis*. One may never know exactly when that physiology arose within dinosaurs. But that does not alter the fact that it is safe to infer that the last

common ancestor shared by all living dinosaurs was an endotherm, while the last common ancestor of living dinosaurs and crocodilians was an ectotherm. Similar inferences concerning the physiology of the *Archaeopteryx* node, the first flying dinosaur, and the first feathered dinosaur, though not impossible, are not nearly as secure. For these reasons, attaching the name "Aves" to the dinosaur crown maximizes the number of safe inferences that can be made about the ancestor of the clade designated by that name; it therefore also maximizes the number of generalizations that can safely be made about birds as the group composed of that ancestor and all of its descendants.

Of course, the crown clade definition of "Aves" would result in significant departures from certain traditions regarding both characters (that is, flight would not be a synapomorphy of Aves but of a more inclusive clade) and composition (that is, fossils such as Archaeopteryx, Hesperornis and Ichthyornis would not be included in Aves). On the other hand, these departures would be deliberate; the change would be made because of its advantages regarding safe inferences and generalizations as well as common use of the name in association with characters that are unknown, or are known not to be present, outside of the crown. Moreover, although the reference of "Aves" under the crown clade definition would depart from some traditions, it would nonetheless be consistent with others. That is to say, it would more closely approximate both the composition and the full set of diagnostic characters of Linnaeus' (1758) "Aves," which did not include any taxa outside of the crown clade and included such characters as "absence of teeth" in its diagnosis. In addition, under the crown clade definition, nothing would be called a bird that had not previously been considered a bird (unlike the case with a feather-based definition), and because all excluded taxa previously considered birds are known only from fossils, this change probably would not greatly affect the research of most ornithologists, who for the most part confine their studies to extant birds. Finally, the name "Aves" does not itself describe a particular character but is simply the Latin name of a kind of organism, so tying it to the crown would not create a misnomer.

Proposed Taxonomy

As a potential solution to the problems addressed above, we propose a new taxonomy, with new names as well as new definitions for old names. Our primary objective is to eliminate ambiguity caused by using the same name, "Aves," for several different clades. A second, closely related goal is to suggest names for clades that everyone seems to want to talk about: the feathered dinosaurs, the flying dinosaurs, and the crown dinosaurs. To accomplish these goals, we will use a different name for each of the clades that other authors have referred to previously by the taxon name "Aves." And to prevent other existing names from suffering from the confusion that previously plagued "Aves," we will explicitly tie five additional names, each of which has been a fixture in the ornithological lexicon for at least a century, to specific clades: "Carinatae" and "Ratitae" of Merrem (1813), "Ornithurae" of Haeckel (1866), and "Palaeognathae" and "Neognathae" of Pycraft (1900). We also take this opportunity to rename the stem clade of bird-line archosaurs and to define names for the basalmost subclades within the crown. It is beyond the scope of this paper to consider some names, including "Ornithopectae," "Ornithothoraces," "Metornithes" and "Euornithes," which have been proposed recently for various basal clades of (putatively) flying dinosaurs.

The principles underlying our choice of names and definitions are currently under development and will be described here only briefly. For reasons discussed in this paper and elsewhere (e.g., Gauthier 1986; Gauthier, Estes et al. 1988; Gauthier, Kluge et al. 1988; de Queiroz and Gauthier 1992), we retain widely used names for crown clades. As a help to remembering associations between names and particular clades, we tried to use names that accurately describe, or at least are consistent with, their references (though violation of this principle should not be taken as a reason for rejecting a name or a definition). Thus, in the case of apomorphy-based definitions, names describe the apomorphies of the clades whose names are defined by those apomorphies (e.g., "Avialae" [= bird wing] for the clade of flying [winged] dinosaurs). For

Panaves
 Avifilopluma
 Avialae
 Carinatae
 Ornithurae
 Aves
 Palaeognathae
 Tinamidae
 Ratitae
 Neognathae
 Galloanserae
 Neoaves

Figure 3. Indented, unranked, summary taxonomy of select clades of bird-line archosaurs.

crown node-based definitions, we tried to use Greek or Latin vernacular names (e.g., "Aves" [= bird]) or those based on the names of subordinate taxa (e.g., "Tinamidae" [based on *Tinamus*]). When there is a conflict between our principle of tying widely used names to crowns and associating descriptive names with the apomorphies that they describe, we allow the former to take precedence (e.g., "Neognathae" [= new jaw] is tied to a crown node rather than to an apomorphy). We also use names for crown nodes that describe plesiomorphies if those names are already well known (e.g., "Palaeognathae"). For total stem-based definitions, we introduce the prefix "pan" (Greek, all). This convention was inspired by Lauterbach's (1989) term "Pan-Monophylum," which is equivalent to Jefferies' (1979) "total group" and Hennig's "* group" plus "stem group" (Jefferies renamed Hennig's "* group" the "crown group").

The wording of our definitions follows rules and recommendations described in the draft of the *PhyloCode* (Cantino and de Queiroz 2000), as does our italicization of all taxon names. In addition to the standard forms of node-, stem- and apomorphy-based definitions, we also make use of a less well known form called the stem-modified node-based definition (de Queiroz in Wyss and Meng 1996). For names used by Linnaeus, our specifiers are selected from among the species, generally the first listed, in the 10th edition of his *Systema Naturae* (1758) (thus *Vultur gryphus* is used as a primary specifier for "Aves"). For names coined by others, we tried to use as specifiers taxa included in their studies, or the types of the subordinate taxa on which their names are based (e.g., *Gallus gallus* for "Galloanserae"). Figure 3 illustrates an indented, unranked, summary taxonomy that is based on applying the names defined below (through their definitions) in the context of currently accepted views of archosaur relationships, although the utility of those names and definitions does not depend on accepting those views about relationships.

Panaves (new clade name)

<u>Definition:</u> "Panaves" refers to the most inclusive clade containing Aves (Vultur gryphus Linnaeus 1758) but not Crocodylia (Crocodylus niloticus Laurenti 1768).

Referred taxa: In addition to Aves (see below) and all other Dinosauria (e.g., Sereno 1999b), Panaves is currently thought to include a variety of Triassic species (e.g., Benton 1999): Pseudolagosuchus major (= Lewisuchus admixtus?), Lagosuchus (= Marasuchus) talampayensis, Lagerpeton chanarensis and Scleromochlus taylori, as well as the Mesozoic Pterosauria (e.g., Wellnhofer 1991). Gauthier (1984, 1986) included Euparkeria and Ornithosuchidae in this clade (which he called "Ornithosuchia"; see "Comments," below). Benton (1984) presented evidence

that *Euparkeria* was outside of the bird plus crocodylian crown (*Archosauria*), and Sereno and Arcucci (1990) proposed that *Ornithosuchidae* was closer to crocodiles than to birds, so neither of these taxa are now referred to *Panaves*.

Comments: Gauthier (1986:42) defined the name "Ornithosuchia" as referring to the clade of "extant birds and all extinct archosaurs that are closer to birds than they are to crocodiles." This definition violates Art. 11.8 of the *PhyloCode* (Cantino and de Queiroz 2000), which requires that a name (e.g., Ornithosuchia von Huene 1908) derived from the stem of a preexisting genus name (Ornithosuchus Newton 1894) must use the type species of that genus (Ornithosuchus longidens Huxley 1877 = Ornithosuchus woodwardi Newton 1894 = Dasygnathus longidens Huxley 1877) as an internal specifier. Moreover, although Ornithosuchia could be redefined in accordance with that rule, according to current phylogenetic hypotheses (e.g., Sereno and Arcucci 1990), it would not refer to the same clade (because Ornithosuchus is closer to crocodiles than to birds). Therefore, we have coined the name Panaves for the clade of bird-line archosaurs. Other names for this taxon, or at least for a group approximating its hypothesized composition, including Patterson's (1993) "Aves," Benton's (1999) "Avemetatarsalia," and many authors' use of Gauthier's (1986) "Ornithodira," will not be discussed here.

Avifilopluma (new clade name)

<u>Definition:</u> "Avifilopluma" refers to the clade stemming from the first panavian with feathers homologous (synapomorphic) with those of Aves (Vultur gryphus Linnaeus 1758). "Feathers" here refers to hollow-based, filamentous, epidermal appendages produced by follicles.

Referred taxa: In addition to Aves (see below), Avifilopluma is currently thought to include the following taxa, based on fossilized impressions of pennaceous feathers: the basal oviraptors Caudipteryx and Protarchaeopteryx (Ji et al. 1998); basal dromaeosaurs such as Sinornithosaurus (Ji et al. 2001; Xu et al. 2001), and the basal flying dinosaurs Archaeopteryx (Owen 1863), Confuciusornithidae (Chiappe et al. 1999), some Enantiornithes (e.g., Eoalulavis [Sanz et al. 1996], Concornis [Sanz et al. 1995]), and Parahesperornis (Williston 1898). With some reservations, the apparently unbranched filamentous epidermal appendages reported in the basal coelurosaur Sinosauropteryx and the basal therizinosaur Beipiaosaurus (Xu, Tang et al. 1999) are regarded to be feathers. Given the phylogenetic relations of these taxa (Gauthier and Gall 2001), Avifilopluma includes all of Maniraptora, and it may include all of Coelurosauria. Indeed, pending more complete knowledge of the epidermis in dinosaurs, Avifilopluma might even contain all but the basalmost theropods (that is, abelisaurids [including scaled Carnotaurus] and coelophysids and their close relatives, as well as taxa more distantly related to birds, such as herrerasaurs and Eoraptor; Sereno 1997, 1999b).

<u>Comments:</u> Many authors have used the name "Aves" for this apomorphy-based clade. As noted above, it appears that at least four kinds of feathers are present in the basalmost dinosaurs that clearly possess feathers (Ji et al. 2001; Xu et al. 2001). If the "feather" phylogenetic—as opposed to developmental—transformation series is from a simple hollow filament, to "natal" down, to pennaceous feathers, to remiges and rectrices, it may be useful to name the clade stemming from the first panavian with "natal" down (Aviplumosa), with pennaceous feathers (Avipinna), and with remiges and rectrices, that is, enlarged, stiff-shafted, closed-vaned (= barbules bearing hooked distal pennulae), pennaceous feathers arising from the distal forelimbs and tail (Aviremigia).

Avialae Gauthier 1986

<u>Definition:</u> "Avialae" refers to the clade stemming from the first panavian with feathered wings homologous (synapomorphic) with those of Aves (Vultur gryphus Linnaeus 1758) and used for powered flight.

Referred taxa: In addition to Aves (see below), Avialae is currently thought to include Ichthyornis (Marsh 1872a), Hesperornithes (Marsh 1872b; Fürbringer 1888), Apsaravis (Norell and

Clarke 2001), *Patagopteryx* (Alvarenga and Bonaparte 1992), *Enantiornithes* (Walker 1981), *Confuciusornithidae* (Hou et al. 1995), *Archaeopteryx* (von Meyer 1861a, 1861b), and *Rahonavis* (Forster et al. 1998). "Flight" in avian outgroups can be inferred from a variety of anatomical modifications in the shoulders, chest, arms and hands, as well as in the morphology of the remiges and rectrices. Given their phylogenetic relationships (Chiappe 2001; Clarke 2001), *Hesperornithes* and *Patagopteryx* are regarded as secondarily flightless avialans.

Comments: Most authors (e.g., Haeckel 1866; Beddard 1898; Heilmann 1927; de Beer 1954; Ostrom 1974b, 1976; Sereno 1998, 1999a) conflate this apomorphy-based clade with the clade stemming from the *Archaeopteryx* node. The name "*Avialae*" was defined originally by Gauthier (1986:36) as referring to "Ornithurae plus all extinct maniraptorans that are closer to Ornithurae than they are to Deinonychosauria," which is a stem-based definition. However, unlike the case for definitions proposed for names of clades that emerged from his analysis, Gauthier (1986) did not always provide explicit definitions for his terminal taxa (e.g., Carnosauria) or use them consistently when he did. Thus, although Gauthier (1986) defined "*Avialae*" as the name of a stem-based clade, he also used it as a name for the *Archaeopteryx* node (for example, when listing the synapomorphies [1986:12]) and as an apomorphy-based name (when referring to avialans as "winged theropods" [1986:12]). Most subsequent authors (e.g., Perle et al. 1993, 1994; Norell et al. 2001) have used *Avialae* as a name for the *Archaeopteryx* node.

In spite of this precedent, "Avialae" is here redefined in keeping with its etymology (Avialae = bird wing) as an apomorphy-based name. As such, Avialae should not be equated with the clade stemming from the Archaeopteryx node. Although the two taxa currently seem identical in terms of known composition (assuming that Archaeopteryx was able to fly), future discoveries will likely reveal that Avialae is a more inclusive clade. The name "Aves" has also been associated with the Archaeopteryx node (e.g., Chiappe 1995, 2001), a practice that we would like to discourage. Although the Archaeopteryx node is of profound historical significance, its biological importance has been diminished by the discovery of taxa marking an extensive series of nodes along the avian stem lineage (Chiappe 2001). Moreover, although Archaeopteryx (probably) possessed at least some capacity for wing-powered flight, newly discovered fossils reveal several subsequent innovations in the step-wise assembly of the flight apparatus that characterizes the crown (Norell and Clarke 2001). For these reasons, and because we do not consider it particularly important to give every clade a formal name, we suggest that this clade be referred to simply, and informally, as "the Archaeopteryx node."

Carinatae Merrem 1813 (converted clade name)

Definition: "Carinatae" refers to the clade stemming from the first panavian with a keeled sternum homologous (synapomorphic) with that of Aves (Vultur gryphus Linnaeus 1758). The sternal keel is here defined as a bony crest extending from the ventral midline of the sternum (e.g., Concornis lacustris; see "Comments," below) for an hypertrophied pectoral (flight) musculature (= m. pectoralis and m. supracoracoideus in the crown; Baumel and Witmer 1993). Referred taxa: In addition to Aves (see below), Carinatae is currently thought to include Ichthyornis, Apsaravis and Enantiornithes. Based on other data, secondarily flightless (and therefore keel-less) Patagopteryx (Chiappe 1995, 1996) and Hesperornithes (Martin 1984), as well as various keel-less members of the crown (e.g., Ratitae), are likewise part of Carinatae. Confuciusornis is the closest known outgroup to Carinatae, although it is polymorphic for the most primitive stage in the evolutionary development of a sternal keel (Chiappe et al. 1999). Comments: Merrem's (1813) name "Carinatae" is based on the Latin carina, referring to the sternal keel. We conserve that association between name and character by proposing an apomorphy-based definition for "Carinatae." Note that although pterosaur sterna are also keeled, that keel is not homologous to the keel of Aves, and therefore pterosaurs are not part of Carinatae. The slight median ridge in some Confuciusornis, on the other hand, seems homologous

to that of *Vultur gryphus* (Linnaeus 1758). But according to Chiappe and co-workers (1999),

only as a sternal "ridge," rather than a sternal "keel." Doubtless other taxa will eventually be found with intermediate morphologies between keel-less (e.g., *Confuciusornis*) and keeled (e.g., *Concornis*) avialans as currently understood. We leave it to those describing such taxa to arbitrarily decide, as Sanz and Buscalioni (1992) and Chiappe and co-workers (1999) did with *Concornis* and *Confuciusornis*, respectively, what degree of development an "interpectoralis ridge" must have to qualify as a "keel."

The carina is known to undergo additional modification in the line leading toward the crown clade (Norell and Clarke 2001). In early Enantiornithes such as Concornis, the keel is confined to the caudal end of the sternum (Sanz and Buscalioni 1992; Sanz et al. 1995), rather than extending cranially to the level of the first sternal rib facets, as it does in *Ichthyornis* (Marsh 1880) and Neuquenornis (Chiappe and Calvo 1994). Moreover, in Concornis the ridges diverging anterolaterally from the keel are directed toward the forelimbs, thus appearing to mark the attachment of pectoralis muscles only; it seems that the supracoracoideus muscles remained confined largely to the coracoid and sternoclavicular membrane (e.g., Olson and Feduccia 1979). In later carinates, in which the keel is more strongly developed, there are craniocaudally oriented ridges on the carina medially and sternal plates laterally, sometimes delimiting a fossa, that mark the attachment line of the fibrous sheath surrounding the supracoracoideus muscle and separating it from the overlying pectoralis muscle. This sternal morphology indicates that the origin of the supracoracoideus has spread along the length and depth of the carina, deep to the pectoralis muscles, as can be seen in the crown (Baumel and Witmer 1993), as well as in a specimen that Marsh referred to Apatornis celer (YPM 1734; a specimen that Clarke [2000], considered closer to Aves than is Ichthyornis). And finally, Huxley (1867) observed that the carina ossifies from its own center in the crown clade, a condition that would be difficult to determine in a fossil. It may prove useful to name the clades associated with one of more of these synapomorphies, but we will leave that to other authors.

Ornithurae Haeckel 1866 (converted clade name)

<u>Definition:</u> "Ornithurae" refers to the clade stemming from the first panavian with a "bird tail," namely, a tail that is shorter than the femur (subequal to or shorter than the tibiotarsus) with a pygostyle of avian aspect—an upturned and ploughshare-shaped compressed compound element, fused in the adult, composed of less than six segments, and shorter than the up-curving free part of tail (the portion between pygostyle and pelvis composed of less than eight vertebrae) on which it is borne—that is homologous (synapomorphic) with that of *Aves* (*Vultur gryphus* Linnaeus 1758).

Referred taxa: Ornithurae is currently thought to include Aves (see below), Ichthyornis (i.e., YPM 1732 [Marsh 1880]), Hesperornithes Clarke (2001), and Apsaravis Norell and Clarke (2001). According to Clarke (2001), Hesperornithes is now thought to be closer to the Ichthyornis node than is Apsaravis (contra Norell and Clarke 2001). Hesperornis caudals are remarkably specialized (Marsh 1880), but several features of the "bird tail" are present in the basal species Baptornis advenus (Martin and Tate 1976). Given Clarke's (2001) hypothesis, Hesperornithes is considered to have a "bird tail," although modified to varying degrees within that clade. Comments: Haeckel (1866:140) originally applied the name "Ornithurae" to a group composed of "true birds" with the "characteristic tail morphology of all extant birds" (translation by Gauthier). Thus, "Ornithurae" ("bird tail") does not appear to describe a particular morphological novelty, but rather a suite of modifications separating the tail of the crown clade from that of Archaeopteryx (Haeckel's Sauriurae). Nevertheless, the name "Ornithurae" has long been associated with a subset of those modifications—namely, a shortened tail with a pygostyle of avian aspect (e.g., Fürbringer 1888; Gadow 1892, 1893; Sereno 1999a)—and our definition is consistent with that tradition. Gadow's (1892, 1893) "Neornithes" was originally proposed as a replacement name for, and thus a junior synonym of, Haeckel's (1866) "Ornithurae." Gauthier (1986:13) defined "Ornithurae" phylogenetically as a stem-based name, namely, "[e]xtant birds and all other taxa, such as *Ichthyornis* and *Hesperornithes*, that are closer to extant birds than is *Archaeopteryx*." However, because the name refers to a trait and the rules of priority for phylogenetically defined names are not yet in effect, we take the opportunity to redefine the name using an apomorphy-based definition.

Fossils found subsequent to Haeckel (1866) have done much to bridge the morphological gap between *Archaeopteryx* (Haeckel's "*Sauriurae*") and the crown clade (Chiappe 2001). Thus, it is now evident that a frond-like tail, as seen in *Archaeopteryx*, evolved into the fan-like tail of modern birds through a series of anatomical transformations, during the course of which the tail was divorced functionally from the hind limbs and entrained to the wings (Gatesy and Dial 1996; Carrano 1998; Gatesy 2001). Tail shortening, which began prior to flight in theropods (Gauthier 1986), as well as evolution of the pygostyle, continued in the line leading to the crown clade (e.g., Norell and Clarke 2001). Some conspicuous morphological gaps remain, especially between the relatively long tail of *Archaeopteryx* in the late Jurassic and the shorter tail—with a pygostyle composed of more than seven fused segments—of *Confuciusornis* in the early Cretaceous (Chiappe et al. 1999). Given that the time separating those fossils is some tens of millions of years, however, the prominence of that gap is almost certainly a consequence of missing data.

The pygostyle of *Confuciusornis* and certain noncrown carinates was by no means in its crown form (sensu Baumel and Witmer 1993). The pygostyle varies considerably within the crown (van Oort 1904), and its degree of differentiation is coupled to rectricial development (that is, it is poorly developed in birds with reduced or absent tail feathers, such as ratites and some tinamous; Gatesy and Dial 1996). Nevertheless, in contrast with its crown morphology, the pygostyle is quite long, and it extends posteriorly in line with the rest of the caudals, in Confuciusornis (Chiappe et al. 1999), Enantiornithes (e.g., Sanz and Bonaparte 1992; Sereno and Rao 1992), and Hesperornithes (e.g., Martin and Tate 1976). In contrast, a pygostyle and tail of avian aspect seems to be present in Apsaravis (Norell and Clarke 2001) and Ichthyornis (YPM 1732; Marsh 1880), so far as they are preserved. That is to say, the pygostyle is shortened—much shorter, rather than longer, than the free part of the tail—and directed upwards at the tail tip. Indeed, Marsh (1880: pl. 28) figures the angled posterior face of the penultimate free caudal in YPM 1732—a morphology that would tip the pygostyle up, as in living birds. Moreover, the upward flexion of the entire tail appears fully developed in the free caudal elements of *Ichthyornis* (YPM 1732; Marsh 1880, pl. 28). It seems that a fan-like tail like that seen in the crown, in which the tail is short and the tail feathers are aligned side-by-side in the rectricial bulb, instead being long with the tail feathers extending in pairs from each caudal vertebra, was already in place by the Apsaravis node. In Aves, encircling bands of muscles within the rectricial bulb can compress these aligned rectrices at their bases, thereby fanning the tail feathers independent of all other tail movements (Gatesy and Dial 1996; Gatesy 2001); this capability may also have been present in the ancestor at the Apsaravis node. The tails of Hesperornithes are difficult to interpret owing to the fact that aquatic birds tend to have longer tails and flightless birds tend to have poorly developed rectrices (Van Oort 1904), and foot-propelled divers have shorter femora and tarsometatarsi as well (e.g., Marsh 1880), thus making it difficult to determine whether hesperornithine tails are primitive or secondarily modified. Nevertheless, given Clarke's (2001) placement of Hesperornithes inside of the Apsaravis node, coupled with the fact that the basalmost Hesperornithes in which the tail is known (i.e., Baptornis; Martin and Tate 1976), had an avian arc to it, hesperornithine tails, especially the pygostyle, are here considered secondarily modified from the ancestral ornithurine condition.

It may eventually prove useful to coin an apomorphy-based name for avialans with fused distal caudals, and we recommend conversion of Chatterjee's (1997) "Pygostylia" to that end (rather than Chiappe's [2001] node-based definition). We recognize that there may be as yet unidentified differences among the tails of Apsaravis, Hesperornithes and Ichthyornis, and between one or more of these taxa and the crown, and that the differences between these and earlier-diverging taxa likely represent a series of transformations. Thus, our definitions of "bird

tail" and "pygostyle of avian aspect" are based on current knowledge and are necessarily arbitrary. We leave it to those describing new taxa that bridge current morphological gaps to refine the definitions of these characters as necessary.

Gadow (1892) proposed the name "Neornithes" as an explicit substitute for, and thus a junior synonym of, Haeckel's (1866) "Ornithurae": for example, "...Neornithes (Ornithurae vieler Autoren)..." (Gadow 1893:99). In contrast to Haeckel, however, Gadow (1892 1893, 1898) explicitly included taxa in his "Neornithes" that are now thought to lie outside of the crown (i.e., Hesperornis and Ichthyornis), as did Fürbringer (1888) for "Ornithurae" before it. Gadow's (1892) "Neornithes" largely supplanted Haeckel's "Ornithurae" in popular use (Gauthier 1986), although it eventually came to be used for the crown (e.g., Cracraft 1986; in Gadow's time Ichthyornis and Hesperornis were thought to be related to gaviiforms and charadriiforms, respectively). Because we use "Aves" for the crown (see below), "Neornithes" could be applied to one of several different clades while remaining consistent with Gadow's (1892) original use; an obvious candidate is the clade stemming from the Hesperornithes node.

Aves Linnaeus 1758 (converted clade name)

<u>Definition:</u> "Aves" refers to the crown clade stemming from the most recent common ancestor of Ratitae (Struthio camelus Linnaeus 1758), Tinamidae (Tetrao [Tinamus] major Gmelin 1789), and Neognathae (Vultur gryphus Linnaeus 1758).

<u>Referred taxa:</u> Aves is currently thought to include some 10,000 living species, including all those listed in Clements (2000). Partial lists of extinct taxa can be found in Unwin (1993).

Comments: Linnaeus (1758) used the name "Aves" for a group composed entirely of extant species and diagnosed by their shared characters. When the species of Linnaeus' "Aves," which included both neognaths and ratites, are placed on a modern phylogenetic tree, the last common ancestor of those species is the last common ancestor of the crown clade (Gauthier 1986). Thus, despite departing from more recent traditions concerning the reference of the name "Aves" (such as inclusion of some fossil outgroups), our definition is consistent with Linnaeus' use of that name. Various authors (e.g., Cracraft 1986; Sereno and Rao 1992; Chiappe 1995, 2001; Cracraft and Clarke 2001) have used the name "Neornithes" for the panavian (and dinosaurian) crown clade. As discussed above, however, Gadow (1892) proposed his "Neornithes" as an explicit replacement for, and in that sense as a junior synonym of, "Ornithurae" (Haeckel 1866). Although use of "Neornithes" for the crown is consistent with Gadow's assumption that Hesperornis and Ichthyornis were closely related to gaviiforms and charadriiforms, respectively (thus implying that they were nested within the crown), for reasons discussed in this and previous papers (e.g., Gauthier 1986; Gauthier, Estes et al. 1988; Gauthier, Kluge et al. 1988; de Queiroz and Gauthier 1992), we use the better known name "Aves" for this clade.

Palaeognathae Pycraft 1900 (converted clade name)

<u>Definition:</u> "Palaeognathae" refers to the crown clade stemming from the most recent common ancestor of *Tinamidae* (*Tetrao* [*Tinamus*] major Gmelin 1789) and *Ratitae* (*Struthio camelus* Linnaeus 1758).

Referred taxa: Tinamidae and Ratitae (see below).

<u>Comments:</u> As described above, not only did Pycraft (1900) base his "*Palaeognathae*" on a "primitive character" (1900:150), his "phylogenetic diagram" (1900:265) indicated palaeognath paraphyly (although he described *Palaeognathae* as a "monophyletic group" [1900:266], it is clear that he did not use that term as we do today). Nevertheless, Pycraft's "*Palaeognathae*" became a nomenclatural fixture. Currently available evidence, although sparse, supports monophyly (in Hennig's 1966 sense) of the set of taxa included in Pycraft's "*Palaeognathae*" (e.g., Cracraft and Clarke 2001), and data in support of alternative arrangements—such as a tinamou–landfowl grouping—are even more tenuous. Because we have restricted this node-based name to a crown

clade, a stem-based counterpart name for the total clade will be useful when members of the palaeognath stem lineage are discovered. Therefore, we propose the name "Panpalaeognathae" (new clade name), defined as the most inclusive clade containing both Tinamidae (Tetrao [Tinamus] major Gmelin 1789) and Ratitae (Struthio camelus Linnaeus 1758) but neither Galloanserae (Phasianus [Gallus] gallus Linnaeus 1758) nor Neoaves (Vultur gryphus Linnaeus 1758). With the possible exceptions of some "lithornids," the taxa currently referred to Panpalaeognathae are the same as for Palaeognathae (namely, Ratitae and Tinamidae).

Tinamidae Gray 1840 (converted clade name)

<u>Definition:</u> "*Tinamidae*" refers to the crown clade stemming from the most recent common ancestor of *Tetrao* (*Tinamus*) *major* Gmelin 1789 and all extant birds sharing a more recent common ancestor with that species than with *Struthio camelus* Linnaeus 1758 and *Vultur gryphus* Linnaeus 1758.

Referred taxa: See Clements' (2000) list of extant species (under *Tinamiformes* and *Tinamidae*). Brodkorb (1963; and see Houde 1988, and references therein) refers several fossil species to Tinamidae, but it is by no means clear if their relations lie within or outside of the crown. Comments: As Newton (1896) observed, tinamous have held a particular fascination for avian taxonomists out of all proportion to their numbers. As these birds became better known to science (de Buffon 1778), they presented a confusing mix of gallinaceous and ratite characters and thus were regarded to stand near the "bond of union between those two great divisions," namely, Huxley's versions of "Ratitae" and "Carinatae" (Newton 1896:964). Disagreements about the precise relationships of tinamous stemmed from disagreements over which of their characters were considered most important. Huxley (1867), for example, chose to emphasize the (plesiomorphic) keeled sternum, placing tinamous in Merrem's "Carinatae." In contrast, Pycraft (1900) focused mainly on shared (plesiomorphic) features of the palate, allying tinamous with ratites in his "Palaeognathae." Wagler (1830), a pre-Darwinian, was the first to classify tinamous with ratites. Among evolutionists, Beddard (1898) acknowledged an implicit phylogenetic link between Tinamidae and Ratitae, and explicit statements to that effect are found in Bock (1963), Cracraft (1974) and Cooper et al. (2001).

Relationships among species of *Tinamidae* are poorly understood; therefore, we use a stem-modified node-based definition to define the name (de Queiroz in Wyss and Meng 1996). To accommodate the possibility that some of the extinct taxa referred to *Tinamidae* by Brodkorb (1963), or some of Houde's (1988) "lithornids," are stem relatives of *Tinamidae*, we propose the stem-based counterpart name "*Pantinamidae*" (new clade name) defined as the most inclusive clade containing *Tinamidae* (*Tetrao* [*Tinamus*] *major* Gmelin 1789), but not *Ratitae* (*Struthio camelus* Linnaeus 1758). Currently, the taxa referred to *Pantinamidae* are the same as for *Tinamidae* (but see preceding comments).

Ratitae Merrem 1813 (converted clade name)

<u>Definition:</u> "*Ratitae*" refers to the crown clade stemming from the last common ancestor shared by *Struthio camelus* Linnaeus 1758 and all extant birds sharing a more recent common ancestor with that species than with *Tetrao* (*Tinamus*) *major* Gmelin 1789 and *Vultur gryphus* Linnaeus 1758.

Referred taxa: Clements (2000) provides a list of extant species (under *Struthioniformes*). Although neither extinct moas and elephantbirds, nor extant kiwis, were known to Merrem (1813) when he coined the name *Ratitae*, they were placed in *Ratitae* soon after their discovery (Lesson 1831; Owen 1839). Nevertheless, the position of the extinct taxa inside or outside of the crown is controversial (e.g., Lee et al. 1997; Cooper et al. 2001). The Eocene taxon *Palaeotis weigelti* has been considered an ostrich relative (e.g., Houde and Haubold 1987), but it may well be a representative of the ratite stem (Gauthier, personal observation).

<u>Comments:</u> The name "*Ratitae*" begs for an apomorphy-based definition, referring as it does to a "raft-like" keel-less sternum. But like other widely used names, *Ratitae* is probably best reserved for a crown clade. Relationships within this clade are uncertain (Lee et al. 1997; Cooper et al. 2001), so we propose a stem-modified node-based definition to ensure that *Ratitae* will be tied to the crown regardless of (ingroup) tree topology (de Queiroz in Wyss and Meng 1996). Ratite monophyly ranks among the most enduring controversies in avian systematics, primarily due to the possibility that their flightless habits (and associated morphological correlates) evolved convergently (e.g., Feduccia 1996.). Nevertheless, there seems to be abundant data in support of this clade (e.g., Lee et al. 1997; Cooper et al. 2001; Cracraft and Clarke 2001), and no characters supporting alternative positions for these taxa within *Aves*. In the absence of character conflict, there is nothing for the hypothesis of convergence to explain. We accordingly accept the hypothesis that the flightless habit of extant ratites was inherited from a flightless common ancestor.

Given that a keel-less sternum is associated with flightlessness, and that both characters arose prior to the last common ancestor of *Ratitae*, it might be useful to name the more inclusive clade based on that apomorphy using a name that brings this attribute to mind. We refrain from naming that clade at the present time. On the other hand, we propose the name "*Panratitae*" (new clade name) for the total stem clade of the ratite crown, defined as the most inclusive clade containing *Ratitae* (*Struthio camelus* Linnaeus 1758) but not *Tinamidae* (*Tetrao [Tinamus] major* Gmelin 1789). Currently, taxa referred to *Panratitae* are the same as for *Ratitae*, although a consensus is lacking on whether the Holocene moas and elephantbirds and the Eocene *Palaeotis* are to be referred to both taxa or only to *Panratitae*.

Neognathae Pycraft 1900 (converted clade name)

<u>Definition:</u> "Neognathae" refers to the crown clade stemming from the last common ancestor of Charadrius pluvialis (Pluvialis apricaria) Linnaeus 1758 and all extant birds sharing a more recent common ancestor with that species than with Struthio camelus Linnaeus 1758 and Tetrao (Tinamus) major Gmelin 1789.

Referred taxa: Neognathae is currently thought to include two primary crown clades, Galloanserae and Neoaves (see below).

<u>Comments:</u> "Neognathae" has long been associated with an apomorphic joint between pterygoid and palatine bones; we nevertheless tie it to a crown clade because it is so widely used. Neognath phylogeny is poorly understood; therefore, we use a stem-modified node-based definition to define "Neognathae" (de Queiroz in Wyss and Meng 1996). Neognathae is one of the oldest recognized, and most strongly supported, deep clades within Aves. Given the number of synapomorphies diagnostic of this clade (Cracraft and Clarke 2001), naming one or more apomorphy-based clades may prove useful, pending finds of extinct neognath outgroups. A stembased counterpart name will clearly be useful; therefore, we propose the name "Panneognathae" (new clade name), defined as the most inclusive clade containing Neognathae (Charadrius pluvialis [Pluvialis apricaria] Linnaeus 1758) but neither Palaeognathae (Struthio camelus Linnaeus 1758) nor Tinamidae (Tetrao [Tinamus] major Gmelin 1789). Currently, the taxa referred to Panneognathae are the same as for Neognathae.

Galloanserae Sibley et al. 1988 (converted clade name)

<u>Definition:</u> "Galloanserae" refers to the crown clade stemming from the last common ancestor shared by Galliformes (Phasianus [Gallus] gallus Linnaeus 1758) and Anseriformes (Anas [Anser] anser Linnaeus 1758).

<u>Referred taxa:</u> Galloanserae is currently thought to include Galliformes and Anseriformes, whose included extant species are listed in Clements (2000). Unwin (1993) provides a partial list of referred fossils.

Comments: Monophyly of a clade composed of galliforms and anseriforms was first proposed

on the basis of DNA hybridization data (Sibley et al. 1988; Sibley and Ahlquist 1990). Subsequent research on diverse character systems continues to support this clade (Cracraft 1988; Livezey 1997; Mindell et al. 1997; Zusi and Livezey 2000; Ericson et al. 2001). A name for the total clade will be useful when members of the galloanseran stem are discovered, so we propose "Pangalloanserae" (new clade name), defined as the most inclusive clade containing Galloanserae (Phasianus [Gallus] gallus Linnaeus 1758 and Anas [Anser] anser Linnaeus 1758) but not Neoaves (Fringilla domestica [Passer domesticus] Linnaeus 1758).

Neoaves Sibley et al. 1988 (converted clade name)

<u>Definition:</u> "Neoaves" refers to the crown clade stemming from the last common ancestor of *Fringilla domestica* (*Passer domesticus*) Linnaeus 1758 and all extant neognaths sharing a more recent common ancestor with that species than with *Galloanserae* (*Phasianus* [*Gallus*] *gallus* Linnaeus 1758 and *Anas* [*Anser*] *anser* Linnaeus 1758).

<u>Referred taxa:</u> Neoaves is currently thought to include all extant bird species listed in Clements (2000) under Sphenisciformes, Gaviiformes, Podicipediformes, Procellariiformes, Pelecaniformes, Ciconiiformes, Falconiformes, Gruiformes, Charadriiformes, Pterocliformes, Columbiformes, Psittaciformes, Opisthocomiformes, Cuculiformes, Strigiformes, Caprimulgiformes, Apodiformes, Coliiformes, Trogoniformes, Coraciiformes, Upupiformes, Piciformes and Passeriformes. Unwin (1993) provides a partial list of referred fossils.

Comments: Sibley et al. (1988) first proposed "Neoaves" for Holocene neognaths other than Galloanserae. Later, Sibley and Ahlquist (1990) changed the reference of this name by including Galloanserae in it, thereby rendering "Neoaves" a junior synonym of "Neognathae." Our definition retains Sibley and co-workers' (1988) original use of "Neoaves" for one of the two primary crown subclades of Neognathae. As Cracraft and Clarke (2001) observe, support for Neoaves comes exclusively from molecular data sets, including a 5-codon deletion in the RAG-1 gene (Groth and Barrowclough 1999). Currently, there is little agreement among trees proposed for relationships among neoavians (Cracraft 1988; Sibley et al. 1988; Sibley and Ahlquist 1990; McKitrick 1991; Cooper and Penny 1997; Mindell et al. 1997; Groth and Barrowclough 1999); consequently, we have used a stem-modified node-based definition to define the name. For the neoavian total-stem clade, we propose "Panneoaves" (new clade name), defined as the most inclusive clade containing Fringilla domestica (Passer domesticus) Linnaeus 1758, but neither Phasianus (Gallus) gallus Linnaeus 1758 nor Anas (Anser) anser Linnaeus 1758.

Conclusions

The nomenclatural inconsistency and ambiguity that results from using the name "Aves" for several different nested clades needs to be corrected. If the primary concern is naming clades, and if names are merely arbitrary labels, then as long as "Aves" refers to some clade, the particular clade is not so important. On the other hand, insofar it is possible to maintain continuity—in other words, "tradition"—when restricting the reference of a name to one of its several prior uses, continuity ought to be maintained. Nevertheless, when it comes to calls for maintaining continuity there can be no simple answers, because traditions can take several different forms. Thus, are we to grant precedence to the use established by the original author, or to later-established uses? And in either case, which associations are we to consider more important, those involving composition or those based on diagnostic characters? Finally, if we choose to emphasize composition over characters, are we to interpret composition in a node-based or a stem-based fashion? All of these alternatives can be defended, which accounts for the current use of the name "Aves" to designate several different clades.

Feathers and flight are generally regarded as "key" evolutionary innovations and therefore particularly worthy of being the basis of the name "Aves." It is now clear, however, that only one of these key innovations can play this role, for evidence is mounting that "flight" and "feathers" arose at different times in the ancestral lineage of birds (contra Feduccia 1996). These evolu-

tionary novelties may have profound biological significance, thus qualifying as key innovations (Galis 2001), but their apparent distinctiveness has rested in no small part on missing data. Indeed, it seems that most characters that are sufficiently distinctive to have been associated traditionally with "higher" taxa are distinctive because they are complex. And most complex characters—including both feathers (e.g., Prum 1999) and the avian flight apparatus (e.g., Ostrom 1997; Norell and Clarke 2001)—turn out to be character complexes once a sufficiently dense series of intermediates comes to light. Many, if not most, conspicuous morphological gaps are eventually bridged by intermediates, regardless of whether that bridging is accomplished continuously or in discrete steps.

Preferentially associating higher taxon names such as "Aves" with marked discontinuities in phylogeny has a long and venerable tradition in biological taxonomy (Darwin 1859; Simpson 1945; Thulborn 1984; Lucas 1992; Lee and Spencer 1996; Sereno 1998, 1999a). Nevertheless, using such gaps to justify the traditional assignment of high taxonomic rank seems ironic for evolutionists, who must concede that many commonly recognized "higher" taxa testify as much to the influence of missing data as to important events in the history of life (Rowe and Gauthier 1992). Perceived disparity between the extant representatives of "higher" taxa often depends heavily on extinction of intermediate forms combined with an incomplete fossil record (Darwin 1859; Gauthier, Kluge et al. 1988). The traditional Class Aves is no exception. Granted that extant birds are separated from other extant reptiles by a considerable morphological gap, the same cannot be said for Caudipteryx, Sinornithosaurus and Archaeopteryx, long-extinct species in which traditional conceptions of "bird" and "reptile" converge and blur by anyone's standards. Indeed, traditional ideas about "key innovations" and "conspicuous morphological gaps" separating birds from other vertebrates lose much of their force in light of the Liaoning theropods. And attempting to accommodate such finds by shifting the boundaries of *Aves* to preserve the association of that taxon with a morphological gap effectively glorifies ignorance and fails to confront the continuity of the evolutionary process. That practice is best avoided.

But that still leaves the fundamental problem of which clades to name and what to name them. In our view, at least five clades formerly designated as "Aves" deserve to be named in one way or another: (1) the clade of bird-line archosaurs; (2) the clade of feathered dinosaurs; (3) the clade of flying dinosaurs; (4) the clade stemming from the *Archaeopteryx* node; and (5) the clade of crown dinosaurs. The reason is not that the speciation events underlying the origins of these clades were especially remarkable. Nor is it because the initial or the final evolutionary novelties diagnosing them are especially significant. It is just that everyone wants to talk about these clades, and every one of them has an advocate who wants to call it by the well known, and thus important, name "*Aves*." The question can therefore be narrowed to the following: what names shall we use for these five clades, and which of those clades is to be designated as "*Aves*"?

Granted that maintaining continuity of use ("traditional meanings") in one form or another is important, at least one of the five candidates can be eliminated. Tying the name "Aves" to the Archaeopteryx node will not do, as that practice cannot conserve the "traditional" reference of the name in the sense of either the original composition of the taxon or the characters most commonly associated with it—that is, flight and feathers. In the case of original composition, Linnaeus referred only members of the crown to Aves. Thus, if tradition is equated with Linnaean composition and interpreted as stem-based, Aves includes all bird-line archosaurs, not just those stemming from the Archaeopteryx node. Moreover, as a stem-based name, it cannot correspond with any node. And if that tradition is interpreted as node-based, it includes only a subset of the descendants of the Archaeopteryx node. Alternatively, if tradition is equated with the characters flight or feathers, defining "Aves" as referring to the Archaeopteryx node fails to ensure an association with either character; that definition merely points to the *known* flying dinosaurs, not to all flying dinosaurs, and it no longer comes close to designating even the known feathered dinosaurs. Finally, for those who wish to maintain the Haeckelian tradition of including Archaeopteryx within Aves yet interpret that tradition in terms of composition rather characters (that is, by tying "Aves" to the Archaeopteryx node rather than to feathers or flight), one must question the logic of appealing to a tradition that itself would have to be interpreted as violating an earlier (Linnaean) tradition.

So in our view that leaves four alternatives: "Aves" could be tied either to feathers, or to flight, through an apomorphy-based definition; to the total clade of bird line archosaurs through a stem-based definition; or to the dinosaurian crown through a node-based definition. Based on a perspective that emphasizes the naming of clades (de Queiroz 1994), and the premise that comparative biology can no longer function effectively by applying the same name to all four of these clades (see de Queiroz and Gauthier 1992), we develop a taxonomy that uses a different name for each. In that taxonomy, we define "Aves" as referring to the dinosaurian crown, the clade whose ancestor we can know the most about. We use "Panaves," a name that evokes the avian pan-monophylum for the total stem clade of bird-line archosaurs. And we use "Avifilopluma" and "Avialae," names that describe the apomorphies of interest, for the clades of feathered and flying (winged) dinosaurs, respectively. But regardless of whether biologists in general and ornithologists in particular adopt our proposed taxonomy, one conclusion is inescapable: the Chinese feathered dinosaurs have forced us to confront the fact that feathers and flight did not arise simultaneously, thus exposing the inadequacies of current taxonomies.

Acknowledgments

We would like to express our gratitude to Julia Clarke and Phil Cantino for the many and varied discussions of issues in phylogenetic taxonomy and nomenclature. We are particularly grateful to Julia for sharing her knowledge of basal avialan phylogeny; she will see many of her ideas in this work. We would also like to thank Lyndon Murray and Matthias Bernett for their help translating the German literature, Walter Joyce for his translation of Linnaeus' Latin diagnosis of Aves, and Lyndon Murray, Daniel Brinkman and Kristof Zyskowski for their help finding 18th and 19th century literature. We are grateful to Ben Creisler for his scholarly advice on issues of classical grammar related to the scientific names used in our taxonomy.

Literature Cited

- Alvarenga, H. M. F. and J. F. Bonaparte. 1992. A new flightless landbird from the Cretaceous of Patagonia. In: K. E. Campbell Jr., editor. Papers in avian paleontology honoring Pierce Brodkorb: proceedings of the 2nd international symposium of avian paleontology and evolution; 1988 Sept 28–30; Los Angeles, CA. Los Angeles: Nat. Hist. Mus. Los Angeles Cty. pp. 51–64. (Nat. Hist. Mus. Los Angeles Cty., Sci. Ser. 36.)
- Baumel, J. J. and L. M. Witmer. 1993. Osteologica.
 In: J. J. Baumel, A. M. King, J. E. Breazile and H. E. Evans, editors. Handbook of avian anatomy: nomina anatomica avium. 2nd ed. Cambridge, MA: Nuttall Ornithological Club. pp. 45–132. (Publ. Nuttall Ornithol. Club 23.)
- Beddard, F. E. 1898. Structure and classification of birds. Longmans, Green: London. 548 pp.
- Benton, M. J. 1984. The relationships and early evolution of the Diapsida. Zool. Soc. Lond. Symp. 52:575–596.
- ---1990. Origin and interrelationships of dinosaurs. In: D. B. Weishample, P. Dodson and H. Osmólska, editors. The Dinosauria. Berkeley: Univ. California Pr. pp. 11–30.
- ----1999. *Scleromochlus taylori* and the origin of dinosaurs and pterosaurs. Phil. Trans. Roy. Soc. Lond. B 354:1423–1446.
- Bock, W. J. 1963. The cranial evidence for ratite affinities. In: C. G. Sibley, editor. Proceedings of the 13th International Ornithological Congress; 1962; Ithaca, New York. Baton Rouge, LA: Am. Ornithol. Union, Mus. Zool., Louisiana State Univ. pp. 39–54.
- Bonaparte, J. F., F. Novas and R. A. Coria. 1990. *Carnotaurus sastrei* Bonaparte, the horned, lightly built carnosaur from the ?Albian of Patagonia. Nat. Hist. Mus. Los Angeles Cty. Contrib. Sci. 416:1–41.
- Bonaparte, J. F. and J. E. Powell. 1980. A continental assemblage of tetrapods from the Upper Cretaceous beds of El Brete, northwestern Argentina (Sauropoda–Coelurosauria–Carnosauria–Aves). Mem. Soc. Geol. France, Nov. Ser. 59 (139): 19–28.
- Brodkorb, P. 1963. Catalogue of fossil birds. Bull. Florida State Mus. 7(4):179–293.
- Brush, A. H. 1996. On the origin of feathers. J. Evol. Biol. 9:131–140.
- ——2000. Evolving a protofeather and feather diversity. Am. Zool. 40(4):631–639.
- ---2001. The beginnings of feathers. In: J. Gauthier and L. F. Gall, editors. New perspectives on the

- origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom; 1999 Feb 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. pp. 171–179.
- Bryant, H. N. 1994. Comments on the phylogenetic definition of taxon names and conventions regarding the naming of crown clades. Syst. Biol. 43(1):124–130.
- Burnham, D. A., K. L. Derstler, P. J. Currie, R. T. Baker, Z. Zhou and J. H. Ostrom. 2000. Remarkable new birdlike dinosaur (Theropoda: Maniraptora) from the Upper Cretaceous of Montana. Univ. Kansas Paleontol. Contrib. 13:1–14.
- Campbell, K. E. 1980. A review of the Rancholabrean avifauna of the Itchtucknee River, Florida. Nat. Hist. Mus. Los Angeles Cty. Contrib. Sci. 330: 119–129.
- Cantino, P. D. and K. de Queiroz. 2000. Phylo-Code: a phylogenetic code of biological nomenclature [online paper]. Athens, OH: Ohio Univ. [updated 3 May 2000]. Available at http:// www.ohiou.edu/phylocode/.
- Carrano, M. T. 1998. Locomotion in non-avian dinosaurs: integrating data from hind limb kinematics, in vivo strains, and bone morphology. Paleobiology 24:450–469.
- Chatterjee, S. 1997. The rise of birds. Baltimore: Johns Hopkins Univ. Pr. 312 pp.
- Chen, P., Z. Dong and S. Zhen. 1998. An exceptionally well preserved theropod dinosaur from the Yixian Formation of China. Nature 391:147–152.
- Chiappe, L. M. 1991. Cretaceous avian remains from Patagonia shed new light on the early radiation of birds. Alcheringa 15:333–338.
- ----1995. The phylogenetic position of the Cretaceous birds of Argentina: Enantiornithes and *Patagopteryx deferrariisi*. Cour. Forsch. Senkenb. 181:55–63.
- ——1996. Late Cretaceous birds of southern South America: anatomy and systematics of Enantiornithes and *Patagopteryx deferrariisi*. Münch. Geowiss. Abh. (A) 30:203–244.
- ——2001. Phylogenetic relationships among basal birds. In: J. Gauthier and L. F. Gall, editors. New perspectives on the origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom; 1999 Feb 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. pp. 125–139.
- Chiappe, L. M. and J. O. Calvo. 1994. *Neuquenornis* volans, a new Enantiornithes (Aves) from the

- Upper Cretaceous of Patagonia (Argentina). J. Vert. Paleontol. 14:230–246.
- Chiappe, L. M., S. Ji, Q. Ji and M. A. Norell. 1999. Anatomy and systematics of the Confuciusornithidae (Aves) from the late Mesozoic of northeastern China. Bull. Am. Mus. Nat. Hist. 242:1–89.
- Clark, J. M., M. A. Norell and L. M. Chiappe. 1999. An oviraptorid skeleton from the Late Cretaceous of Ukhaa Tolgod, Mongolia, preserved in an avian-like brooding position over an oviraptorid nest. Am. Mus. Novit. 3265:1–36.
- Clarke, J. 2000. Setting limits on phylogenetic inference from isolated elements. Vert. Palasiat. 38 (Suppl. 3):9–10.
- ---2001. The morphology and systematic position of *Ichthyornis* Marsh and the phylogenetic relationships of basal Ornithurae [dissertation]. New Haven: Yale University, Department of Geology and Geophysics. 457 pp.
- Clements, J. F. 2000. Birds of the world: a checklist. Vists, CA: Ibis. 867 pp.
- Cooper, A., C. Lalueza-Fox, S. Anderson, A. Rambaut, J. Austin and R. Ward. 2001. Complete mitochondrial genome sequences of two extinct moas clarify ratite evolution. Nature 409:704–707.
- Cooper, A. and D. Penny. 1997. Mass survival of birds across the Cretaceous–Tertiary boundary: molecular evidence. Science 275:1109–1113.
- Cracraft, J. 1974. Phylogeny and evolution of ratite birds. Ibis 116:494–521.
- ----1986. The origin and early diversification of birds. Paleobiology 12(4):383–399.
- ——1988. The major clades of birds. In: M. J. Benton, editor. The phylogeny and classification of the tetrapods. Volume 1, Amphibians, reptiles, birds. Oxford: Clarendon Pr. pp. 339–361.
- Cracraft, J. and J. Clarke. 2001. The basal clades of modern birds. In: J. Gauthier and L. F. Gall, editors. New perspectives on the origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom; 1999 Feb 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. pp. 143–156.
- Currie, P. J. and M. A. Norell. 1998. The anatomy of two feathered theropods from Liaoning, China. J. Vert. Paleontol. 18(Suppl. 3):36A.
- Darwin, C. 1859. On the origin of species by means of natural selection; or, the preservation of favoured races in the struggle for existence. London: John Murray. 502 pp.
- de Beer, G. 1954. *Archaeopteryx lithographica:* a study based on the British Museum specimen. London: Trustees of the British Museum (Nat-

- ural History) 224. 68 pp. [Reprinted 1967 Waterford, U.K.: Taylor Garnett Evans.]
- de Buffon, G. 1778. Histoire naturelle des oiseaux, Volume 4. Paris: De l'Imprimerie Royale.
- de Pinna, M. C. C. 1992. Concepts and tests of homology in the cladistic paradigm. Cladistics 7(4): 367–394.
- de Queiroz, K. 1994. Replacement of an essentialistic perspective on taxonomic definitions as exemplified by the definitions of "Mammalia." Syst. Biol. 43(4):497–510.
- ——1998. The general lineage concept of species, species criteria, and the process of speciation: a conceptual unification and terminological recommendations. In: D. J. Howard and S. H. Berlocher, editors. Endless forms: species and speciation. New York: Oxford Univ. Pr. pp. 57–75.
- 1999. The general lineage concept of species and the defining properties of the species category.
 In: R. A. Wilson, editor. Species: new interdisciplinary essays. Cambridge: MIT Pr. pp. 49–89.
- de Queiroz, K. and M. J. Donoghue. 1988. Phylogenetic systematics and the species problem. Cladistics 4(4):317–338.
- ——1990. Phylogenetic systematics or Nelson's version of cladistics. Cladistics 6(1):61–75.
- de Queiroz, K. and J. Gauthier. 1990. Phylogeny as a central principle in taxonomy: phylogenetic definitions of taxon names. Syst. Zool. 39(4):307–322.
- ---1992. Phylogenetic taxonomy. Annu. Rev. Ecol. Syst. 23:449–480.
- ---1994. Toward a phylogenetic system of biological nomenclature. Trends Ecol. Evol. 9:27–31.
- Dingus, L. and T. Rowe. 1998. The mistaken extinction: dinosaur evolution and the origin of birds. New York: W. H. Freeman. 332 pp.
- El zanowski, A. 1991. New observations on the skull of *Hesperornis* with reconstructions of the bony palate and otic region. Postilla 207:1–20.
- Ericson, P. G. P., T. J. Parsons and U. S. Johansson. 2001. Morphological and molecular support for nonmonophyly of the Galloanserae. In: J. Gauthier and L. F. Gall, editors. New perspectives on the origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom; 1999 Feb 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. pp. 157–168.
- Feduccia, A. 1996. The origin and evolution of birds. New Haven: Yale Univ. Pr. 420 pp.
- Feduccia, A. and H. B. Tordoff. 1979. Feathers of Archaeopteryx: asymmetric vanes indicate aerodynamic function. Science 203:1021–1022.

- Forster, C. A., S. D. Sampson, L. M. Chiappe and D. W. Krause. 1998. The theropod ancestry of birds: new evidence from the Late Cretaceous of Madagascar. Science 279:1915–1919.
- Frost, D. R. and A. G. Kluge. 1994. A consideration of epistemology in systematic biology, with special reference to species. Cladistics 10(3):259–294.
- Fürbringer, M. 1888. Untersuchungen zur Morphologie und Systematk der Vögel, Volumes 1–2. Amsterdam: Von Holkema. 1751 pp.
- Gadow, H. 1892. On the classification of birds. Proc. Zool. Soc. Lond. 17:229–256.
- ---1893. Vögel. Part 2, Systematischer Theil. Leipzig: C. F. Winter. 303 pp. (Dr. H. G. Bronn's Klassen und Ordnungen des Their-Reichs. Bd. 6. Abt. 4.)
- ——1898. A classification of Vertebrata recent and extinct. London: Adam & Charles Black. 82 pp.
- Gal is, F. 2001. Key innovations and radiations. In: G. Wagner, editor. The character concept in evolutionary biology. San Diego: Academic Pr. pp. 581–605.
- Gatesy, S. M. 2001. The evolutionary history of the theropod caudal locomotor module. In: J. Gauthier and L. F. Gall, editors. New perspectives on the origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom; 1999 Feb 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. 333–347 pp.
- Gatesy, S. M. and K. P. Dial. 1996. From frond to fan: *Archaeopteryx* and the evolution of short-tailed birds. Evolution 50:2037–2048.
- Gauthier, J. 1984. A cladistic analysis of the higher systematic categories of the Diapsida [dissertation]. Berkeley: University of California, Department of Paleontology. 564 pp. Available from University Microfilms, Ann Arbor, MI; 85-12825.
- ----1986. Saurischian monophyly and the origin of birds. In: K. Padian, editor. The origin of birds and the evolution of flight. San Francisco: California Acad. Sci. pp. 1–55. (Mem. Calif. Acad. Sci. 8.)
- Gauthier, J., R. Estes and K. de Queiroz. 1988. A phylogenetic analysis of Lepidosauromorpha. In: R. Estes and G. K. Pregill, editors. Phylogenetic relationships of the lizard families: essays commemorating Charles L. Camp. Palo Alto: Stanford Univ. Pr. pp. 15–98.
- Gauthier, J. and L. F. Gall, editors. 2001. New perspectives on the origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom: 1999 Feb

- 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. 613 pp.
- Gauthier, J., A. Kluge and T. Rowe. 1988. Amniote phylogeny and the importance of fossils. Cladistics 4(2):105–209.
- Gegenbaur, C. 1863. Vergleichendend-anatomische Bemerkungen über das Fusskelet der Vögel. Arch. Anat. Physiol. Wiss Med. 1863:450–472.
- Ghiselin, M. T. 1984. "Definition," "character," and other equivocal terms. Syst. Zool. 33:104–110.
- Gishlick, A. D. 2001. The function of the manus and forelimb of *Deinonychus antirrhopus* and its importance for the origin of avian flight. In: J. Gauthier and L. F. Gall, editors. New perspectives on the origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom; 1999 Feb 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. pp. 301–318.
- Gmel in, J. F. 1789. Caroli a Linné, equitis auranti de stella polari ... Systema Naturae per regna tria naturae, secundum classes, ordines, genera, species, cum characteribus, differentiis, synonymis, locis. Editio decimo tertia, aucta, reformata. Lipsiae: Impensis Georg. Emanuel. Beer (1788–1793). 3 vols. [Publication date sometimes seen as 1788; 2nd edition printed with alterations from Lugduni.]
- Gould, S. J. and E. S. Vrba. 1982. Exaptation—a missing term in the science of form. Paleobiology 8:4–15.
- Gray, G. R. 1840. A list of the genera of birds. London: R. & J. Taylor. 63 pp.
- Groth, J. G. and G. F. Barrowclough. 1999. Basal divergences in birds and the phylogenetic utility of the nuclear RAG-1 gene. Mol. Phylog. Evol. 12(2):115–123.
- Haeckel, E. 1866. Generelle Morphologie der Organismen. Berlin: Georg Reimer. 462 pp.
- Harrison, C. J. O. 1973. The humerus of *Ichthyornis* as a taxonomically isolating character. Bull. Br. Ornithol. Club. 93(3):123–126.
- Heilmann, G. 1927. The origin of birds. New York: Appleton. 209 pp. [Reprint of London: Witherby, 1926; reprinted New York: Dover, 1972.]
- Hennig, W. 1965. Phylogenetic systematics. Annu. Rev. Entomol. 10:97–116.
- ----1966. Phylogenetic systematics. Urbana: Univ. Illinois Pr. 263 pp.
- ---1981. Insect phylogeny. Chichester: J. Wiley. 514 pp.
- Holtz, T. R. Jr. 1994. The phylogenetic position of the Tyrannosauridae: implications for theropod systematics. J. Paleontol. 68:1100–1117.

- ——1996. Phylogenetic taxonomy of the Coelurosauria (Dinosauria: Theropoda). J. Paleontol. 70: 536–538.
- Hou, L., Z. Zhou, Y. Gu and Y. Sun. 1995. Introduction to the Mesozoic birds from Liaoning, China. Vert. Palasiat. 33(4):261–271.
- Houck, M. A., J. Gauthier and R. E. Strauss. 1990.
 Allometric scaling in the earliest fossil bird, Archaeopteryx lithographica. Science 247:195–198.
- Houde, P. W. 1988. Palaeognathous birds from the early Tertiary of the northern hemisphere. Publ. Nuttall Ornithol. Club 22:1–148.
- Houde, P. W. and H. Haubold. 1987. *Palaeotis weigelti* restudied: a small Eocene ostrich (Aves: Struthioniformes). Paleovertebrata 17:27–42.
- Huxley, T. H. 1867. On the classification of birds and on the taxonomic value of the modifications of certain of the cranial bones observable in that class. Proc. Zool. Soc. Lond. 1867:415–472.
- ——1877. The crocodilian remains found in the Elgin Sandstones, with remarks of on the ichnites of Cummingstone. Mem. Geol. Surv. U. K., Monogr. 3:1–58.
- Jefferies, R. P. S. 1979. The origin of chordates—a methodological essay. In: M. R. House, editor. The origin of major invertebrate groups. London: Academic Pr. pp. 443–477.
- Ji, Q., P. J. Currie, M. A. Norell and S. Ji. 1998. Two feathered theropods from the Upper Jurassic/ Lower Cretaceous strata of northeastern China. Nature 393:753–761.
- Ji, Q. and S. Ji. 2001. How can we define a feathered dinosaur as a bird? In: J. Gauthier and L. F. Gall, editors. New perspectives on the origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom; 1999 Feb 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. pp. 43–46.
- Ji., Q., M. A. Norell, K. Gao, S. Ji and D. Ren. 2001. The distribution of integumentary structures in a feathered dinosaur. Nature 410:1084–1088.
- Jones, T. D., J. O. Farlow, J. A. Ruben, D. M. Henderson and W. J. Hillenius. 2000. Cursoriality in bipedal archosaurs. Nature 406:716–718.
- Jones, T. D., J. A. Ruben, L. D. Martin, E. N. Kurochkin, A. Feduccia and P. F. A. Maderson. 2000. Nonavian feathers in Late Triassic archosaur. Science 288:2202–2205.
- Kluge, A. G. 1995. Cladistic relationships of sphaerodactyl lizards. Am. Mus. Novit. 3139:1–23.
- Kron, K. A. and W. S. Judd. 1990. Phylogenetic relationships within the Rhodoreae (Ericaceae) with

- specific comments on the placement of *Ledum*. Syst. Bot. 15:57–68.
- Laurenti, J. N. 1768. Specimen medicum, exhibens synopsin reptilium emendatam cum experimentis circa venena et antidota reptilium Austriacorum. Vienna: Joan. Thomæ Nob. de Trattnern. 214 pp.
- Lauterbach, K.-E. 1989. Das Pan-Monophylum— Ein Hilfsmittel für die Praxis der phylogenetischen Systematik. Zool. Anz. 223:139–156.
- Lee, K., J. Feinstein and J. Cracraft. 1997. The phylogeny of ratite birds: resolving conflicts between molecular and morphological data sets. In: D. P. Mindell, editor. Avian molecular evolution and systematics. San Diego: Academic Pr. pp. 173 –211.
- Lee, M. S. Y. 1996. Stability in meaning and content of taxon names: an evaluation of crown-clade definitions. Proc. Roy. Soc. Lond. B 263:1103–1109.
- ---1998. Phylogenetic uncertainty, molecular sequences, and the definition of taxon names. Syst. Biol. 47(4):719–726.
- Lee, M. S. Y. and P. S. Spencer. 1997. Crown-clades, key characters and taxonomic stability: when is an amniote not an amniote? In: S. S. Sumida and K. Martin, editors. Amniote origins: completing the transition to land. San Diego: Academic Pr. pp. 61–84.
- Lesson, R.-P. 1831. Traite d'ornithologie, Volume 1. Paris: Levrault. 659 pp.
- Linnaeus, C. 1758. Systema naturae per regna tria naturae. 10th ed. Volume 1, Regnum animale [photographic facsimile]. London: Trustees, British Museum (Natural History), 1939. 823 pp.
- Livezey, B. C. 1997. A phylogenetic analysis of basal Anseriformes, the fossil *Presbyornis*, and the interordinal relationships of waterfowl. Zool. J. Linn. Soc. 121: 361–428.
- Lucas, S. 1992. Extinction and the definition of the class Mammalia. Syst. Biol. 41:370–371.
- Marsh, O. C. 1872a. Notice of a new remarkable fossil bird. Am. J. Sci. 4:3–44.
- ----1872b. Discovery of a remarkable fossil bird. Am. J. Sci. 3:5657.
- ——1880. Odontornithes: a monograph on the extinct toothed birds of North America. Washington, DC: Dept. U. S. Army, Prof. Paper Engineers 18. 201 pp. [Also published as Volume 1, Memoirs of the Peabody Museum of Yale College, New Haven, CT.]
- Martin, L. D. 1984. A new hesperornithid and the relationships of the Mesozoic birds. Trans. Kans. Acad. Sci. 87(34):141–150.

- ——1991. Mesozoic birds and the origin of birds. In: H.-P. Schultze and L. Trueb, editors. Origins of the higher groups of tetrapods: controversy and consensus. Ithaca: Comstock Pub. pp. 485–540.
- Martin, L. D. and J. Tate. 1976. The skeleton of *Baptornis advenus* (Aves: Hesperornithiformes). In: S. L. Olson, editor. Collected papers in avian paleontology honoring the 90th birthday of Alexander Wetmore. Smithson. Contrib. Paleobiol. 27:35–66.
- McKitrick, M. C. 1991. Phylogenetic analysis of avian hindlimb musculature. Univ. Mich. Mus. Zool. Misc. Pub. 179:1–85.
- Merrem, B. 1813. Tentamen systematis naturalis avium, Abh. Konigel. (Preussische) Akad. Wiss. Berlin 1812–1813(1816) (Physikal.):237–259.
- Mindell, D. P., M. D. Sorenson, C. J. Huddleston, H. C. Miranda Jr., A. Knight, S. J. Sawchuk and T. Yuri. 1997. Phylogenetic relationships among and within select avian orders based on mitochondrial DNA. In: D. P. Mindell, editor. Avian molecular evolution and systematics. San Diego: Academic Pr. pp. 213–247.
- Newton, A. 1896. A dictionary of birds, Volume 4. London: Adam & Charles Black. pp. 833–1088.
- Newton, E. T. 1894. Reptiles from the Elgin Sandstone: descriptions of two new genera. Phil. Trans. Roy. Soc. Lond. (B) 185(13):573–607.
- Nixon, K. C. and Q. D. Wheeler. 1992. Extinction and the origin of species. In: M. J. Novacek and Q. D. Wheeler, editors. Extinction and phylogeny. New York: Columbia Univ. Pr. pp. 119–143.
- Norberg, R. A. 1985. Function of vane asymmetry and shaft curvature in bird flight feathers: inferences on flight ability of *Archaeopteryx*. In: M. K. Hecht, J. H. Ostrom, G. Viohl and P. Wellnhofer, editors. The beginnings of birds: proceedings of the international *Archaeopteryx* conference; 1984; Eichstätt, Germany. Eichstätt: Freunde des Jura-Museums Eichstätt. pp. 303–318.
- ——1995. Feather asymmetry in Archaeopteryx. Nature 374:2–21.
- Norell, M. A., J. M. Clark and P. J. Makovicky. 2001. Phylogenetic relationships among coelurosaurian theropods. In: J. Gauthier and L. F. Gall, editors. New perspectives on the origin and early evolution of birds: proceedings of the international symposium in honor of John H. Ostrom; 1999 Feb 12–14; New Haven, CT. New Haven: Peabody Mus. Nat. Hist., Yale Univ. pp. 49–67.
- Norell, M. A. and J. Clarke. 2001. Fossil that fills a critical gap in avian evolution. Nature 409:181 –184.

- Olson, S. L. 1985. The fossil record of birds. In: D. S. Farner, J. R. King and K. C. Parkes, editors. Avian biology. New York: Academic Pr. pp.79–238
- Olson, S. L. and A. Feduccia. 1979. Flight capability and the pectoral girdle of *Archaeopteryx*. Nature 278:247–248.
- Osborn, H. F. 1912. Integument of the iguanodont dinosaur *Trachodon*. Mem. Am. Mus. Nat. Hist. 1:33–54.
- Ostrom, J. H. 1974a. The pectoral girdle and forelimb function of *Deinonychus* (Reptilia: Saurischia): a correction. Postilla 165:1–11.
- ---1974b. *Archaeopteryx* and the origin of flight. Q. Rev. Biol. 49:27–47.
- ---1976. *Archaeopteryx* and the origin of birds. Biol. J. Linn. Soc. 8:91–182.
- ----1985. The meaning of Archaeopteryx. In: M. K. Hecht, J. H. Ostrom, G. Viohl and P. Wellnhofer, editors. The beginnings of birds: proceedings of the international Archaeopteryx conference; 1984; Eichstätt, Germany. Eichstätt: Freunde des Jura-Museums Eichstätt. pp. 161–175.
- ——1997. How bird flight might have come about. In: D. L. Wolberg, E. Stump and G. D. Rosenberg, editors. Dinofest International: proceedings of a symposium held at Arizona State University; 1996 Apr; Phoenix. Philadelphia: The Academy of Natural Sciences. pp. 301–310.
- Owen, R. 1839. On the bone of an unknown Struthious bird from New Zealand. Proc. Zool. Soc. Lond. 1839:169–170.
- ---1863. On the *Archaeopteryx* of von Meyer, with a description of the fossil remains of a long-tailed species from the lithographic slate of Solnhofen. Phil. Trans. Roy. Soc. Lond. 153:33–47.
- Padian, K. 1998a. The origin and early evolution of birds. Biol. Rev. 73:1–42.
- ----1998b. When is a bird not a bird? Nature 393: 729–730.
- Padian, K. and L. M. Chiappe. 1998. The early evolution of birds. Biol. Rev. 73:1–42.
- Padian, K., J. R. Hutchinson, T. R. Holtz Jr. 1999. Phylogenetic definitions and nomenclature of the major taxonomic categories of the carnivorous Dinosauria (Theropoda). J. Vert. Paleontol. 19(1):69–80.
- Patterson, C. 1982. Morphological characters and homology. In: K. A. Joysey and A. E. Friday, editors. Problems of phylogenetic reconstruction. London: Academic Pr. pp. 21–74. (Spec. Vol. 21.)
- ---1993. Naming names. Nature 366:518.
- ----1994. Bony fishes. In: D. R. Prothero and R. M. Schoch, editors. Major features of vertebrate

- evolution. Knoxville, TN: Paleontol. Soc., Univ. Tennessee. pp. 57–84. (GSA Short Courses Paleontol. 7.)
- Paul, G. S. 1988. Predatory dinosaurs of the world. New York: Simon & Schuster. 464 pp.
- Perle, A., L. M. Chiappe, R. Barsbold, J. M. Clark and M. A. Norell. 1994. Skeletal morphology of *Mononykus olecranus* (Theropoda: Avialae) from the Late Cretaceous of Mongolia. Am. Mus. Novit. 3105:1–29.
- Perle, A., M. A. Norell, L. M. Chiappe and J. M. Clark. 1993. Flightless bird from the Cretaceous of Mongolia. Nature 362:623–626.
- Prum, R. O. 1999. The development and evolutionary origin of feathers. J. Exp. Zool. 285:291–306.
- Pycraft, W. P. 1900. On the morphology and phylogeny of the Palaeognathae (Ratitae and Crypturi) and Neognathae (Carinatae), Pt. 2. Trans. Zool. Soc. Lond. 15:149–290.
- Reisz, R. R. and H.-D. Sues. 2000. The "feathers" of *Longisquama*. Nature 408:428.
- Rietschel, S. 1985. Feathers and wings of Archaeopteryx. In: M. K. Hecht, J. H. Ostrom, G. Viohl and P. Wellnhofer, editors. The beginnings of birds: proceedings of the international Archaeopteryx conference; 1984; Eichstätt, Germany. Eichstätt: Freunde des Jura-Museums Eichstätt. pp. 251–260.
- Romer, A. S. 1966. Vertebrate paleontology, 3rd ed. Chicago: Univ. Chicago Pr. 468 pp.
- Rowe, T. and J. Gauthier. 1992. Ancestry, paleontology, and the definition of the name Mammalia. Syst. Biol. 41(3):372–378.
- Ruben, J. A. and T. D. Jones. 2000. Selective factors associated with the origin of fur and feathers. Am. Zool. 40:585–596.
- Sanz, J. L. and J. F. Bonaparte. 1992. A new order of birds (Class Aves) from the Lower Cretaceous of Spain. In: K. E. Campbell Jr., editor. Papers in avian paleontology honoring Pierce Brodkorb: proceedings of the 2nd international symposium of avian paleontology and evolution; 1988 Sept 28–30; Los Angeles, CA. Los Angeles: Nat. Hist. Mus. Los Angeles Cty. pp. 39–49. (Nat. Hist. Mus. Los Angeles Cty., Sci. Ser. 36.)
- Sanz, J. L., J. F. Bonaparte, L. M. Chiappe, B. P. Pérez-Moreno, A. D. Buscalioni, J. J. Moratella, F. Ortega and F. J. Poyato-Ariza. 1996. An Early Cretaceous bird from Spain and its implications for the evolution of avian flight. Nature 382:442–445.
- Sanz, J. L. and A. D. Buscalioni. 1992. A new bird from the Early Cretaceous of Las Hoyas, Spain

- and the early radiation of birds. Paleontology 35(4):829–845.
- Sanz, J. L., L. M. Chiappe and A. D. Buscalioni. 1995. The osteology of *Concornis lacustris* (Aves: Enantiornithes) from the Lower Cretaceous of Spain and a re-examination of its phylogenetic relationships. Am. Mus. Novit. 3133: 1–23.
- Schander, C. and M. Thollesson. 1995. Phylogenetic taxonomy—some comments. Zool. Scipta 24:263–268.
- Sereno, P. C. 1997. The origin and evolution of dinosaurs. Ann. Rev. Earth Planet. Sci. 25:435–489.
- ——1998. A rationale for phylogenetic definitions, with application to the higher-level taxonomy of Dinosauria. N. Jhrb. Geol. Palaontol. Abh. 210(1):41–83.
- ---1999a. Definitions in phylogenetic taxonomy: critique and rationale. Syst. Biol. 48(2):329–351.
- ---1999b. The evolution of dinosaurs. Science 284:2137–2147.
- Sereno, P. C. and A. B. Arcucci. 1990. The monophyly of crurotarsal archosaurs and the origin of bird and crocodile ankle joints. N. Jhrb. Geo. Palaeontol. Abh. 180:21–52.
- Sereno, P. C. and C. Rao. 1992. Early evolution of avian flight and perching: new evidence from Lower Cretaceous of China. Science 255:845–848.
- Sibley, C. G. and J. E. Ahlquist. 1990. Phylogeny and classification of birds: a study in molecular evolution. New Haven: Yale Univ. Pr. 976 pp.
- Sibley, C. G., J. E. Ahl quist and B. L. Monroe Jr. 1988. A classification of the living birds of the world based on DNA–DNA hybridization studies. Auk 105:409–423.
- Simpson, G. G. 1945. The principles of classification and a classification of mammals. Bull. Am. Mus. Nat. Hist. 85:1–350.
- Smith, A. B. 1994. Systematics and the fossil record: documenting evolutionary patterns. Oxford: Blackwell Sci. Pub. 223 pp.
- Speakman, J. R. and S. C. Thomson. 1994. Flight capabilities of *Archaeopteryx*. Nature 370:514.
- Speakman, J. R. and S. C. Thomson. 1995. Feather asymmetry in *Archaeopteryx*. Nature 374: 221–222.
- Stevens, P. F. 1994. The development of biological systematics: Antoine-Laurent de Jussieu, nature, and the natural system. New York: Columbia Univ. Pr. 616 pp.
- Thulborn, R. A. 1975. Dinosaur polyphyly and the classification of archosaurs and birds. Aust. J. Zool. 23:249–270.

- ——1984. The avian relationships of *Archaeopteryx*, and the origin of birds. Zool. J. Linn. Soc. 82: 119–158.
- Unwin, D. M. 1993. Aves. In: M. J. Benton, editor. The fossil record, Volume 2. London: Chapman & Hall. pp. 717–737.
- Van Devender, T. R. and G. L. Bradley. 1994. Late Quaternary amphibians and reptiles from Maravillas Canyon Cave, Texas, with discussion of the biogeography and evolution of the Chihuahuan desert herpetofauna. In: P. R. Brown and J. W. Wright, editors. Herpetology of the North American deserts. pp. 23–53. (Southwest. Herpetol. Soc. Spec. Pub. 5.)
- van Oort, E. D. 1904. Beitrag zur osteologie des vögelschwanzes. Tijdschr. Ned. Dierk. Vereen. 9:1–144.
- von Huene, F. 1908. Die Dinosaurier der europäischen Triasformation mit Berücksichtigung der aussereuropäischen Vorkommnisse. Jena, Germany: G. Fischer. 419 pp. (Geol. Paläontol. Abh. Suppl.-Bd. 1.)
- von Meyer, H. 1861a. Vögel-Federn und *Palpipes priscus* von Solnhofen. Neues Jhrb. Min. Geol. Paleontol. 1861:5–61.
- ----1861b. Archaeopteryx lithographica (Vögel-Feder) und Pterodactylus von Solnhofen. Neues Jhrb. Min. Geol. Paleontol. 1861:678–679.
- Wagler, J. G. 1830. Natürliches System der Amphibien, mit vorangehender Classification der Säugethiere und Vögel. München, Stuttgart und Tübingen: J. G. Cotta. 354 pp.
- Wagner, A. 1861. Neue Beiträge zur Kenntnis der urweltlichen Fauna des lithographischen Schiefers: Schildkröten und Saurier. Abh. Bayer. Akad. Wiss. 9:67–124.
- Wagner, G. and J. Gauthier. 1999. 1,2,3 =2,3,4: a solution to the problem of the homology of digits in the avian hand. Proc. Natl. Acad. Sci. 96: 5111–5116.
- Walker, C. 1981. New subclass of birds from the Cretaceous of South America. Nature 292:51–53.
- Wellnhofer, P. 1991. The illustrated encyclopedia of pterosaurs. New York: Crescent Books. 192 pp.
- Wetmore, A. 1956. A check-list of the fossil and prehistoric birds of North America and the West Indies. Smithson. Misc. Coll. 131(5):105.
- Williston, S. W. 1898. Birds. Univ. Geol. Surv. KS. 4(2):43–64.
- Wyss, A. and J. Meng. 1996. Application of phylogenetic taxonomy to poorly resolved crown clades: a stem-modified node-based definition of Rodentia. Syst. Biol. 45(4):559–568.

- Xu, X., Z. Tang and X. Wang. 1999. A therizinosaurid dinosaur with integumentary structures from China. Nature 399:350–354.
- Xu, X., X. Wang and X. Wu. 1999. A dromaeosaurid dinosaur with filamentous integument from the Yixian Formation of China. Nature 401:262 –266.
- Xu, X., Z. Zhou and R. Prum. 2001. Branched integumental structures in *Sinornithosaurus* and the origin of feathers. Nature 410:200–204.
- Zusi, R. L. 1993. Patterns of diversity in the avian skull. In: J. Hanken and B. K. Hall, editors. Volume 2, The skull. Chicago: Univ. Chicago Pr. pp. 391–437.
- Zusi, R. L. and B. C. Livezey. 2000. Homology and phylogenetic implications of some enigmatic cranial features in galliform and anseriform birds. Ann. Carnegie Mus. 69(3):157–193.