Collected Papers in Avian Paleontology Honoring the 90th Birthday of Alexander Wetmore

STORRS L. OLSON
EDITOR

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Smithsonian Institution
Collected Papers
in Avian Paleontology
Honoring the 90th Birthday of
Alexander Wetmore

Storrs L. Olson
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ABSTRACT

Olson, Storrs L., editor. Collected Papers in Avian Paleontology Honoring the 90th Birthday of Alexander Wetmore. Smithsonian Contributions to Paleobiology, number 27, 211 pages, 91 figures, 88 tables, 1976.—Eighteen papers covering diverse aspects of avian paleontology—from the earliest known bird to extinct species found in Indian middens—are collected here to honor the 90th birthday of Alexander Wetmore. These are preceded by an appraisal of the current state of avian paleontology and of Alexander Wetmore's influence on it, including a bibliography of his publications in this field. John H. Ostrom analyzes the hypothetical steps in the origin of flight between Archaeopteryx and modern birds. Philip D. Gingerich confirms that Ichthyornis and Hesperornis did indeed bear teeth, that the palate in Hesperornis is paleognathous, and that these Cretaceous toothed birds appear to occupy a position intermediate between dinosaurs and modern birds. Larry D. Martin and James Tate, Jr. describe the skeleton of the Cretaceous diving bird Baptornis advenus and conclude that the Baptornithidae belong in the Hesperornithiformes, but are less specialized than Hesperornis. Pierce Brodkorb describes the first known Cretaceous land bird as forming a new order possibly ancestral to the Coraciiformes and Piciformes. E. N. Kurochkin summarizes the distribution and paleoecology of the Paleogene birds of Asia, with particular emphasis on the evolution of the gruiform families Eogruiformes and Ergilornithidae. Pat Vickers Rich and David J. Bohaska describe the earliest known owl from Paleocene deposits in Colorado. Alan Feduccia transfers the Eocene genus Neanis from the Passeriformes to the Piciformes and he and Larry D. Martin go on to refer this and four other genera to a new family of Piciformes, concluding that these were the dominant perching land birds of the Eocene of North America. Storrs L. Olson describes a new species of Todidae from the Oligocene of Wyoming and refers the genus Protornis from the Oligocene of Switzerland to the Momotidae, concluding that the New World Coraciiformes originated in the Old World. Charles T. Collins describes two new species of the Eo-Oligocene genus Aegialornis and presents evidence that the Aegialornithidae should be referred to the Caprimulgiformes rather than to the Apodiformes, although they might be ancestral to the swifts. In the following paper he shows that the earliest known true swifts (Apodidae) are three nominal forms from the Lower Miocene of France which prove to be but a single species of Cypseloides, a modern genus belonging to a primitive subfamily now restricted to the New World. Stuart L. Wafer describes a new osprey from the Miocene of California and concludes that the species became extinct through human agency less than 3800 years ago.

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Preface

Had contributions for this volume been sought from the associates and friends of Alexander Wetmore in all fields of ornithology, their number would have been much too great to permit the timely appearance of this festschrift, for the endeavor was conceived barely in time for its proper execution. It was decided, therefore, to limit the scope of this work to avian paleontology—a study which has been particularly dear to Alex Wetmore for three score years. That this collection could be assembled and set before the press in less than a year is a tribute not only to the eagerness of the contributors to honor their esteemed colleague in his 90th year, but also to the fact that there is currently an extensive and active interest in the study of fossil birds—a fact that must be particularly gratifying to Dr. Wetmore, who for so many years strived to keep such an interest alive. The editor is particularly indebted to Dorsey Dunn and Joanne Williams, who typed and retyped manuscripts with great patience and care, and Anne Curtis, who assisted in preparing numerous illustrations. He also wishes to express his appreciation for the fine cooperation of the contributors; their combined efforts have here produced what is certain to be a landmark in paleornithology.
Alexander Wetmore is so familiar a figure to scientists as the dean of American ornithology that it is difficult to realize that he has been directly associated with the Smithsonian Institution as an administrator since 1924. His first responsibilities were in connection with the National Zoological Park, of which he became Superintendent in 1924. Subsequently, Dr. Wetmore became Assistant Secretary for Science of the Institution and Director of the Museum of Natural History in 1925, and continued as Assistant Secretary until 1945, when he was elected by the Regents to serve as the sixth Secretary, succeeding Dr. Charles G. Abbot, who retired in that year.

Throughout this period, and after his own retirement from administrative responsibilities in 1952, Dr. Wetmore has continued an extraordinarily active career in ornithology. In addition to his many duties with the Smithsonian, he also served as Home Secretary of the National Academy of Sciences from 1951 to 1955 and has been for many years a Trustee and Vice-Chairman of the Research Committee of the National Geographic Society.

Throughout this career his publications on birds have continued in depth and in great volume. Following his retirement he has continued his monographic studies on the birds of Panama, which have culminated in the publication of three volumes of “The Birds of the Republic of Panama” (Smithsonian Miscellaneous Collections, volume 150), with a fourth part in preparation. Even now, Dr. Wetmore’s work is not completed and he continues to be a productive scientist in the laboratory of the Division of Birds.

In addition to the many research publications on fossil material specializing in birds, Dr. Wetmore is known today as one of the most outstanding systematic specialists. His renowned arrangement of the sequence of higher taxa of birds, “A Classification for the Birds of the World” (Smithsonian Miscellaneous Collections, 139 (11):1–37, 1960), still stands virtually unchallenged. He is a winner of the Brewster Medal of the American Ornithologists’ Union, and recently, in May 1975, of the Hubbard Medal of the National Geographic Society.

The amount of materials contributed by him to the collections of the National Museum is monumental. Indeed, present-day ornithologists would be staggered to think of the production of research and study material deposited by Dr. Wetmore in the National Collection: some 26,058 skins from North America, Puerto Rico, Hispaniola, the Hawaiian Islands, Uruguay, Paraguay, Argentina, Chile, Venezuela, and Central America, with more than half, some 14,291, from Panama alone. Of skeletal and anatomical specimens, Dr. Wetmore has prepared and contributed 4363, an enormously important increment to the anatomy collections in Washington. The majority of these are from North America and Puerto Rico, but nearly 1000 are from Central and South America and 540 from Panama.
alone. Of eggs, Dr. Wetmore has collected 201 clutches from North, Central, and South America. In this day and age when the collecting of birds has become markedly diminished due to the general knowledge of specimens in existing museums, as well as the varying directions taken in present-day studies in environment and ecology which tend to preclude such collecting, Dr. Wetmore's collections seem large in retrospect; but they form part of the fundamental resource on which present and future work will depend. The very magnitude of these collections would tend to make further collecting in most areas where he has worked superfluous. So today the specialist in taxonomic studies can be grateful for the efforts of meticulous collectors such as Dr. Wetmore, whose work has laid out in depth representative material. Thus, only highly specific additional collecting need be done in the future in areas where Wetmore's work has given us the foundation of our knowledge.

The number of species and subspecies described by Dr. Wetmore is equally impressive. Over the years since 1914 he has described as new to science some 189 species and subspecies of recent birds. Many of these, in fact most, are from Central and northern South America, but much of Dr. Wetmore's most significant early field work was done in the Caribbean, particularly in Puerto Rico, Hispaniola, and adjacent islands in the Greater Antilles. In addition, through the initiative of the late Dr. Casey Wood, Wetmore worked on and described a number of species from the Fiji Islands, as well as forms from other islands of the Pacific. His monographic revisions of a number of species of northern Central and North American birds, as well as Argentinian and southern South American birds, have produced many novelties for science. A great deal of his work was done in revising the avifauna of Venezuela with the late W. H. Phelps, Sr., with whom he co-authored a number of new species and subspecies.

At least one of Dr. Wetmore's discoveries, the population of Chilean Pintail found in the vicinity of Bogotá, Colombia, has subsequently gone extinct, due presumably to hunting pressure. Many of the environments in which he worked in Colombia and adjacent parts of northern South America are already so radically changed that one wonders whether additional forms may not have gone extinct as well. It is a sadness of our time that the development of tropical regions of the world, with the consequent destruction of forests and unique habitats, particularly in South and Central America, has been so rapid that many forms of the accompanying avifauna may never be seen again in life. In a spirit of pre-science, Alexander Wetmore was an early supporter of the Pan-American Section of the International Council for Bird Preservation, having joined T. Gilbert Pearson, Robert Cushman Murphy, Marshall McLean, William Vogt, and Hoyes Lloyd in helping to set up the original organization with Latin American colleagues.

Many of his admirers have named numbers of new birds after our beloved former Secretary, among them a long-billed rail of the Venezuelan coast, *Rallus wetmorei*, which I have recently considered in my own ornithological work. Including *Rallus wetmorei*, some 16 modern species and subspecies of birds have been named in honor of Alexander Wetmore, as well as 4 mammals, 7 reptiles and amphibians, 2 fishes, 9 insects, 5 molluscs, a sponge, a cactus, a glacier, and a canopy bridge in the Bayano River forest in Panama. Truly the incessant and intensive zeal which he has single-mindedly given to the study of birds over the years, often at very considerable personal expenditure in time and energy, will mark the career of Alexander Wetmore as one of the most memorable in the entire history of American ornithology.
I had been corresponding with Alexander Wetmore for several years before I had a chance to meet him. This I did in Washington, D.C., in the spring of 1926. Referring to a visit I made to the National Zoological Park at that time, I wrote as follows:

... the National Zoological Park is managed by the Assistant Secretary of the Smithsonian Institution, Dr. Alexander Wetmore, one of the youngest and most accomplished naturalists in the United States. Notwithstanding his heavy administrative obligations, Dr. Wetmore finds enough time for study in descriptive ornithology and technical work, and observations of birds in freedom and in captivity, all with remarkable results. I visited the Zoo under his kind guidance... (L'Oiseau, 7(1926):205).

Dr. Wetmore himself published in the same issue of that periodical (pages 324–325), a report of the first breeding in captivity at the National Zoo of the Blue Snow Goose, with several photographic plates. He was, therefore, awarded a special medal by the Société Nationale d'Acclimatation de France. Dr. Wetmore was Director of the National Zoo for two years, and before he exchanged that function for the Assistant Secretaryship he was responsible for choosing as his successor, Dr. William Mann, who was an outstanding Zoo director for many years.

The welcome given me by Dr. Wetmore in 1926 remains vivid in my memory, and my mother and I visited Washington under his cordial and competent guidance. Later on, we had many opportunities of getting together at meetings and congresses, as we have had many interests in common. We met in Europe and in America frequently, working together for bird preservation since the inception of the International Council for Bird Preservation. We saw even more of each other after 1940, when I came to live in the United States.

We are now among the few ornithologists of our generation still alive. We sadly miss many of our old friends, particularly Frank Chapman, Tom Barbour, Robert Cushman Murphy, James Chapin and T. Gilbert Pearson, to list only a few who worked with us on different projects. It is to me a very special comfort to know that Alex still is here, looking and acting and writing much as he always has, and I wish him all the happiness he deserves. As past Secretary of the Smithsonian Institution he joins the ranks of those others who have seemed over the years almost immortal; thus his continuing research for many years seems assured.
Alexander Wetmore and the Study of Fossil Birds

Storrs L. Olson

In most general discussions of paleontology or ornithology, the subject of fossil birds is almost invariably treated with a predictable uniformity. Mention is made of Archaeopteryx and the Cretaceous toothed birds, and occasionally some of the large Tertiary predators like Diatryma and Phorusrhacos. This is accompanied by a statement explaining that bird bones are fragile and seldom preserved, thus accounting for what is alleged to be a meager and uninformative fossil record for the entire class. Through frequent repetition, this myth has gained such general acceptance that the uninformed find it difficult to conceive of an avian paleontologist being able to find enough to keep himself occupied. Yet for 60 years Alexander Wetmore has produced a steady stream of papers on fossil birds. With over 150 such entries and nearly as many new fossil taxa to his credit, he can without reservation be said to have contributed more to this field than any other single person. One cannot help but be humbled to think that this is but a fraction of his total scientific output.

Bringing together this collection of papers in avian paleontology to honor Alexander Wetmore’s 90th birthday on 18 June 1976 provides not only an opportunity to review his influence on paleornithology over the past six decades, but also offers a chance to begin dispelling the fiction that fossil birds are rare and provide little information on avian evolution.

Wetmore’s most intensive work on fossil birds took place in the period after the waning of excitement over the spectacular 19th century discoveries of Mesozoic birds, but before most of the renewed modern interest in avian paleontology had been sparked. For many years Wetmore was virtually the only person anywhere who was engaged in research on fossil birds, with the notable exception of the California school of Loye and Alden Miller and Hildegarde Howard. Thus it was natural that bird fossils from all parts of the United States and from areas of the world as diverse as Inner Mongolia, Java, St. Helena, Hawaii, and Bermuda, passed through Wetmore’s hands continually. To this day, the cabinets in his office hold a rich trove of undescribed treasures from a wide array of horizons and localities.

For many years, Wetmore has assiduously maintained an extensive card catalog of references from which he prepared three separate editions of a checklist of fossil birds of North America. He also endeavored to keep his colleagues abreast of current developments in avian paleontology through numerous addresses, lectures, and entertaining synoptic papers—all the while maintaining a consistently high level of production of basic detailed descriptions and diagnoses of new forms.

Wetmore’s first paper on fossil birds involved removing the large Miocene bird described by R. W. Shufeldt as Palaeochenoides miocaenus from the Anseriformes to the Pelecaniformes. Shufeldt, whom Wetmore knew well, was in no way
pleased by this, but Wetmore's action was quite correct. Specimens possibly representing two new species of Palaeochenoides have recently come to the National Museum and it now appears that these may provide a breakthrough in our understanding of these huge, enigmatic seabirds. Wetmore's recognition of the true affinities of Palaeochenoides marked the first step toward this understanding. Shufeldt, it might be noted, was a singular eccentric who, although making many contributions to avian paleontology, repeatedly made serious errors in identification. The process of re-evaluating Shufeldt's taxa, begun by Wetmore and others, has continued up to the present, as seen, for example, in the papers on Eocene Piciformes elsewhere in this volume.

The first new bird Wetmore described from osteological remains was a new genus and species of large flightless rail, Nesotrochis debooyi, found in Indian middens in the Virgin Islands. That such deposits may still be of interest to avian paleontologists is clearly demonstrated by Morejohn in the final paper of the present volume. In recent years two new species of Nesotrochis have been described from Cuba and Hispaniola; despite this, the genus remains so distinctive that there is not yet a good clue as to its affinities within the Rallidae.

Wetmore continued to draw notice to the extinct Pleistocene birds of the West Indies, analyzing fossil avifaunas from Puerto Rico, Haiti, Cuba, and the Bahamas. Among the most notable of his discoveries was the giant barn owl, Tyto ostologa, of Haiti, which he correctly diagnosed from a small fragment of tarsometatarsus. He later described a similar species, T. pollens, along with two new large eagles, from the Bahamas. As late as 1959, Brodkorb, in dedicating to him a new fossil species of crow from New Providence Island, remarked that Alexander Wetmore was "responsible for all previous knowledge of fossil birds of the West Indies." Since then, there have been many additional discoveries of avian fossils in the Antilles, the most remarkable of which are certainly the gigantic raptors of Cuba brought to light through the labors of Oscar Arredondo (summarized in this volume). Among the material from the same deposits that yielded Tyto ostologa, a new rail and a new falcon have recently been found. There is every reason to believe that the fossil resources of the Greater Antilles will continue to produce surprises, while as far as avian paleontology is concerned, the Lesser Antilles are terra incognita.

Perhaps the greatest proportion of Wetmore's paleontological efforts concerned the identification and description of Tertiary birds from North America, especially those of the Eocene, Oligocene, and Miocene terrestrial deposits of the western states and the marine Miocene of the east coast. In these areas he has laid the groundwork for all future researches.

Some of the most exciting recent finds of fossil birds are from the extensive lower Eocene deposits of the Green River Formation, for these often yield complete, articulated skeletons, as for example a particularly fine specimen of primitive frigatebird now under study by the writer. Feduccia and Martin in this volume discuss the significance of the Green River Piciformes, which are now coming to light with remarkable rapidity since Brodkorb's recognition of the first species in 1970. But perhaps the most astonishing of developments in Green River paleornithology are the tremendous deposits of flamingo bones discovered by Paul O. McGrew and now under study by him and Alan Feduccia. Here too, Wetmore's past contributions have played a part, for he described this flamingo in 1926 as a new genus of recurvirostrid, Presbyornis. This case of mistaken identity is understandable in view of Feduccia's further investigations, which
have disclosed some extraordinary similarities between the skeletons of recurvirostrids and flamingos, particularly those of the lower Eocene forms. This is further confirmed by an undescribed flamingo of Bridgerian age in the National Museum which is even more similar to recurvirostrids than is *Presbyornis*. These discoveries now appear to be leading to a reappraisal of the affinities of both the flamingos and the shorebirds.

Wetmore's several contributions on Eocene owls resulted in his erecting a new family, the Protostrigidae, the importance of which is only now becoming apparent. The fossil record of owls is particularly good and we now know that the order extends back at least as far as the Paleocene (see Rich and Bohaska's paper in this volume). Much unstudied material of fossil owls is to be found in various museums, which, along with the revision of the many forms already known, should provide an especially fruitful area of inquiry for avian paleontologists in the future. Of Wetmore's Eocene birds, perhaps the most provocative is *Neocathartes grallator*, a long-legged vulture that was based on a nearly complete skeleton.

Wetmore's contributions once provided just about all that was known of the birds from the extensive Oligocene deposits of western North America. These are now producing new and extremely interesting fossil birds almost annually (e.g., Olson's paper in this volume). One of the predominant groups of birds in the North American Oligocene was the gruiform family that Wetmore named the Bathornithidae. Wetmore himself offered more than one interpretation of the possible relationships of this group and Cracraft has recently proposed others. It seems certain that the final word has not been said on this matter, but the importance of the Bathornithidae is undisputed. Once again, it was Wetmore's pioneering work on the group that has made possible all subsequent investigations. It now appears that the Oligocene limpkins (Aramidae) described by Wetmore will soon be augmented by a new genus, known from much of a skeleton collected in Wyoming by Dr. R. J. Emry of the National Museum. Oligocene raptors described by Wetmore include two forms inseparable from the modern genus *Buteo*, and an intriguing species, *Palaeoplancus sternbergi*, which was made the type of a new subfamily of Accipitridae.

For Wetmore, some of the most interesting fossil deposits were those closest to home—the Miocene marine beds of the Chesapeake Group. Most of what we know of the birds of these deposits is to be found in Wetmore's publications, including the description of a diminutive gannet, *Microsula avita*, which is now known to be relatively common in these beds. In the past few years many new specimens, some of them highly significant, have come to the National Museum from this area, although these are as yet undescribed. As abundant as this material is, it is far overshadowed by the tremendous collections of Miocene and Pliocene age that have recently been acquired from a phosphate mine in North Carolina and which this writer has had the privilege of studying in collaboration with Dr. Wetmore. This is probably the largest deposit of Tertiary birds in existence and thousands of fossils of more than 50 species have so far been recovered. These collections, along with those from Bone Valley, Florida, being studied by Brodkorb, and those from the Pacific coast, which are constantly productive (see the contributions by Howard and Wärter in this volume), provide a solid basis for making unprecedented gains in our knowledge of evolution in the Alcidae, Procellariidae, Diomedeidae, Gaviidae, Sulidae, Phalacrocoracidae, and other families of marine birds.
In 1931, Wetmore published a large paper on the Pleistocene avifauna of Florida in which it was shown that several birds, such as the California condor and the huge vulture *Teratornis*, then known only from the west, particularly from the tarpits at Rancho la Brea, were also present in Florida. This opened up a very fertile area of investigation and in succeeding years the studies of Brodkorb and others have continued to be a source of new information on the rich Pleistocene avifauna of Florida (e.g., Storer's paper herein). In his many years of involvement in paleornithology, Wetmore has repeatedly been called upon to identify material from Pleistocene caves and from Indian middens, a task which as often as not holds few rewards but which nevertheless he pursued with alacrity. From such studies he published numerous notes showing that the distribution of many modern North American species was once much different than at present, as indicated, for example, by Canada Jays, Magpies, and Sharp-tailed Grouse in Virginia, and Spruce Grouse in Virginia and Georgia. The sum of these observations has proved to be a significant contribution to our knowledge of the effects of Pleistocene climatic changes on avian distribution.

When the Central Asiatic Expeditions of the American Museum of Natural History discovered fossil birds in the Eocene of Inner Mongolia, it was to Wetmore that the specimens were sent for study. The most abundant material was that of the crane-like bird which Wetmore named *Eogrus aeola*, assigning it to
a new family, Eogruidae. Recently, the significance of these birds as the probable ancestors of the peculiar two-toed running birds of the family Ergilornithidae has been demonstrated (see Kurochkin's paper herein) and provides one of the most interesting examples of an evolutionary lineage in the avian fossil record.

Oceanic islands are of particular interest to the avian paleontologist because of the rapid extinction of species after the introduction of exotic predators by man. Most such introductions occurred before the era of scientific exploration and thus many insular species can be known only from the study of fossil or subfossil remains. Here Wetmore has likewise made numerous contributions. In 1945 he described an extinct goose from the island of Hawaii. This turned out to be but a small indication of what was to come, for in the past few years the Bishop Museum has forwarded to him for examination numerous fossils from Molokai and Maui, which comprise one of the most extraordinary avifaunas ever uncovered, some of the species being so anomalous as to be quite beyond the wildest imaginations of the most whimsical fantasizer. From Pleistocene deposits on Bermuda, Wetmore described a crane and a duck, leaving to Brodkorb the naming of five new rails from these and other deposits on the island (as yet undescribed). From St. Helena, in the South Atlantic Ocean, Wetmore named a new rail to provide a first step in the elucidation of the extensive fossil avifauna of that island, which this writer has recently had the opportunity to expand.

We have touched on but a few of Alexander Wetmore's contributions to avian paleontology and their importance to present and future research. It should by now be clear that, contrary to persistent belief, fossil birds are not uncommon, and in the following pages it should be equally evident that there is much to be gained from their study.

At last there is some light being shed on the study of Cretaceous land birds (see Brodkorb's paper herein), an area that had hitherto been a void. The renowned Pleistocene tar pits at Rancho la Brea, California, long erroneously held to be the only really productive source of avian fossils, now find a rival in similar deposits in South America which portend a new era of discovery on that continent (see Campbell's paper in the present volume). Although these many new finds are of paramount importance, the avian paleontologist has also inherited a rich source of information in the fossils that have been made known previously. Re-examination of the much discussed but widely misunderstood Mesozoic birds, such as the Jurassic Archaeopteryx and the Cretaceous toothed divers, has generated exciting new ideas and controversy, all of which can only lead to a better understanding of avian evolution (see the papers by Ostrom, Gingerich, and Martin and Tate in this volume). Long-neglected fossil birds, such as those from the vast Tertiary collections of France and from the wealth of material in the New Zealand Quaternary, are coming under scrutiny once again, and in the light of modern concepts find a better place in the evolutionary scheme (see papers herein by Collins and Cracraft). It would seem, therefore, that avian paleontology is truly experiencing a renaissance.

In 1932, Joseph Grinnell (Auk, 49:9-13) in pondering the latest edition of the American Ornithologists' Union's Checklist of North American Birds, to which Wetmore contributed the portion on fossils, attempted to make some inferences about future lists and the number of species they might contain. Concerning the fossil list he queried, "And what about the number and relative acumen of future students in avian paleontology: Will they be more numerous and more
alert than heretofore or will the attractions in this field wane in the face of the ascending allurements for bright minds of bio-physics, bio-chemistry and cosmic mathematics? These questions are more or less baffling of answer.” Forty-five years later, the answers are apparent. We offer the present volume as testimony to the fact that avian paleontology has quite enough allure of its own to attract numerous and perspicacious practitioners, and that the materials they study allow of significant advances not only in the knowledge of birds, but of biology and paleontology as a whole. The discipline that Alexander Wetmore nurtured for sixty years is expanding and vigorous and reaping the benefits of his devotion.
Publications in Avian Paleontology
by Alexander Wetmore

1917

1918

1920

1922

1923

1924

1925

1926
12. Descriptions of Additional Fossil Birds from the Miocene of Nebraska. American Museum Novitates, 211:1-5, 6 figures. (11 March)

¹ Exact dates of publication, when known, are included for papers in which new taxa are proposed.


1928


1929


1930


1931


44. The Avifauna of the Pleistocene in Florida. Smithsonian Miscellaneous Collections, 85(2):1–41, 16 figures, 6 plates. (15 May)


1932


1933


60. The Status of Minerva antiqua, Aquila ferox, and Aquila lydekkeri as Fossil Birds. American Museum Novitates, 680:1–4, 1 figure. (4 December)

61. An Oligocene Eagle from Wyoming. Smithsonian Miscellaneous Collections, 87(19):1–9, 19 figures. (26 December)


1934


64. A Fossil Quail from Nebraska. Condor, 36(1):50, figure 5. (15 January)


1935

1936

1937

1938

1939

1940

1941
1942


1943


1944


1945


1948


1949

110. Dittyma. Page 324 in volume 7 of *Encyclopaedia Britannica*.
111. Hesperornis. Pages 530–531 in volume 11 of *Encyclopaedia Britannica*.
112. Ichthyornis. Page 58A in volume 12 of *Encyclopaedia Britannica*.

1950


1951

1952


1955


1956


125. A Fossil Guan from the Oligocene of South Dakota. Condor, 58(3):234–255, 1 figure. (23 May)


1957


1958

129. Miscellaneous Notes on Fossil Birds. Smithsonian Miscellaneous Collections, 135(8):1–11, 5 plates. (26 June)

1959


1960


133. Pleistocene Birds in Bermuda. Smithsonian Miscellaneous Collections, 140(2):1–11, 3 plates. (7 July)


1962


1963

143. An Extinct Rail from the Island of St. Helena. *Ibis*, 103b(3):379–381, plate 9. (1 September)

1964


1965


1967


1968


1972


Index to Fossil Avian Taxa Described
by Alexander Wetmore

The status of a number of these taxa has changed since their original description and therefore only an alphabetical arrangement is attempted here. Species are listed in the genera in which they were originally described. Taxa marked with an asterisk are preoccupied and no longer available. Following each name is the publication number (from the preceding bibliography) and page in which the name was proposed.

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- Cladornithidae, 132:4
- Cyphornithidae, 27:4
- Eleutherornithidae, 118:3
  - *Eocathartidae, 103:69
  - *Eocathartoidea, 103:69
- Eogruidae, 65:30
- Eonessinae, 85:280
- Gaviellinae, 89:30
- Geranoididae, 57:115
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- Palaeospizidae, 10:192

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ABSTRACT
The five known skeletal specimens of *Archaeopteryx* provide the only presently available anatomical evidence pertaining to the earliest stages in the evolution of the avian flight apparatus. This evidence, together with the osteology of modern birds, makes possible the reconstruction of some hypothetical anatomical stages that must have occurred during the course of avian evolution. It is postulated that one of the most critical components of the flight apparatus is the coracoid. Evolutionary changes in coracoid morphology elevated the actions of the principal humeral extensor (M. coracobrachialis) and forearm flexor (M. biceps), and as a consequence, caused deflection of the course of the M. supracoracoideus, converting it from a humeral depressor to a wing elevator. These changes appear to have been related to predation and feeding activities in the earliest birds, rather than to early stages of flight. Subsequently, additional changes in the forelimb components provided for restricted elbow and wrist movements, compact folding of the forelimb, and more stable support of the remiges. These last changes appear to have taken place after the acquisition of incipient flight capability.

Introduction
One of the most remarkable of all animal adaptations is that of flight, which perhaps has reached its zenith among vertebrates in the diverse kinds of flight displayed by modern birds. Strangely enough, there have been only a few investigations or speculations about the origins of avian flight, but perhaps that stems from the clear logic (Bock, 1965, 1969) of the currently favored arboreal theory of flight origins (Marsh, 1880). The purpose of this paper, however, is not to explore that particular question, which I have already reviewed elsewhere (Ostrom, 1974), but rather it is to present purely theoretical reconstructions of some of the anatomical stages that must have occurred during the course of evolution of the avian flight apparatus, and to discuss the implications thereof.

Reconstruction of such hypothetical evolutionary stages is speculative to be sure, but it is a fruitful exercise in this instance because we know the nature of the starting point, the almost non-bird *Archaeopteryx* (Figure 1), as well as the "end point," the highly perfected flight apparatus of modern birds. A few authors (Heptonstall, 1970; Yalden, 1970) have investigated the possible flight capabilities of *Archaeopteryx*, but apparently no one has examined in any detail the anatomical changes that clearly must have occurred in the flight apparatus between the *Archaeopteryx* stage and that of modern birds. In the absence of any recognized intermediate stages within the avian fossil record, consideration of these necessary anatomical changes assumes major significance, since they may very well provide the only possible clues about early selective factors that led to the develop-
Figure 1.—The Berlin specimen of *Archaeopteryx lithographica* found in 1877 near Eichstätt, Germany, in the Late Jurassic Solnhofen Limestones. Preservation of feather impressions, showing remarkably fine structural details, established these as the remains of a true bird, despite the fact that the skeletal anatomy is more like that of theropod dinosaurs than that of modern birds. (The scale is 100 mm long.)
ment of powered avian flight. Conceivably, such considerations might even shed light on the actual beginnings of flight.

A premise that is critical for the remarks that follow is that the several specimens of Archaeopteryx represent an extremely primitive stage in the evolution of birds (Ostrom, 1973, 1975). (I also believe that Archaeopteryx represents a preflight stage [Ostrom, 1974], but not everyone concurs with such an interpretation.) Some authors (de Beer, 1954; Swinton, 1960, 1964) have maintained that Archaeopteryx was not in the main lineage of avian evolution, but so far not one single bit of evidence has been found, either in the known specimens of Archaeopteryx or elsewhere, to support such a contention. Indeed, as Simpson (1946) observed, Archaeopteryx is anatomically intermediate between reptiles and modern birds, and regardless of whether it is directly ancestral to modern carinates, it is entirely reasonable to assume that the early main-line ancestry of birds included an anatomical stage comparable, if not identical, to that of Archaeopteryx. Thus, any consideration of the evolution of avian flight must start with Archaeopteryx.

Acknowledgments.—I gratefully acknowledge the assistance and courtesies of A. J. Charig of the British Museum (Natural History), London; H. Jaeger of the Humboldt Museum fur Naturkunde, East Berlin; T. Kress of the Solenhofen Actien-Verein, Solnhofen, Bavaria; C. O. van Regteren Altena of Teyler’s Stichting, Haarlem; and P. Wellnhofer of Bayerische Staatssammlung fur Paläontologie und historische Geologie, Munich, who granted me the privilege of studying the specimens of Archaeopteryx in their care. I am also indebted to Walter Bock, who read an early version of the manuscript and offered valuable suggestions and criticisms. These studies were funded by grants from the Frank M. Chapman Memorial Fund of the American Museum of Natural History, and the John T. Doneghy Fund of the Yale Peabody Museum.

Flight Apparatus of Modern Birds

By way of introduction to this section, certain generalized comparisons among higher vertebrates may be useful. In modern quadrupedal reptiles, the proximal components of both the fore and hind limbs extend laterally from the hip and shoulder joints (sprawling posture), which are situated well below the level of the vertebral column. In quadrupedal mammals, both appendages are normally positioned in near-parasagittal orientation (upright posture) articulating with hip and shoulder sockets that are close to the level of the vertebral column. In birds, the hip and shoulder sockets are both elevated and lie in or near the plane of the vertebrae. But birds are peculiar in that the hind limb projects downward in a nearly parasagittal orientation, whereas the forelimb extends out laterally from the body. These contrasting limb orientations in birds obviously are correlated with the different limb movements in the two modes of avian locomotion: terrestrial locomotion by means of alternating (or synchronous) longitudinal limb excursion in the hind quarters, and powered flight by means of complex, but chiefly synchronous (nonalternating) dorsoventral transverse movements of the forelimbs.

The avian skeleton includes a number of specializations that are directly or indirectly involved with powered flight: (1) The trunk region is quite rigid due to fusion or restricted articular freedom of the thoracic vertebrae, the solid bony connection between the vertebral column and the sternum, by full ossification of the ventral (sternal) as well as the dorsal ribs, and the development of uncinate processes on the dorsal ribs. (2) Fixation of the shoulder joints by means of elongation of the coracoids which have developed solid bony articulations with a fully ossified sternum; fusion of the clavicles into a single median strut, the furcula, which appears to function as a spring-like spacer maintaining proper transverse spacing of the shoulder joints. (3) Complete ossification and enlargement of the sternum and the development of a deep and robust sternal keel. (4) Modification of the forelimb skeleton into a rigid but collapsible airfoil support in which the shoulder joint permits humeral movements in nearly all directions (including limited long-axis humeral rotation), but the elbow and wrist joints are restricted so as to confine forearm flexion and extension chiefly to the plane of the wing, wrist movements being limited to flexion and extension in the wing plane only; fusion of some carpals and metacarpals to provide a solid platform for the attachment of the primary remiges; and reduction of the manus to
digits I, II and III, with II and III united into a relatively rigid structure. Elsewhere (Ostrom, in press), I have given reasons for discounting the suggestion by some authors (Holmgren, 1955) that the remaining digits of the hand are II, III and IV. (5) The caudal vertebrae are reduced in number and coalesced into a short pygostyle, providing a firmer and more readily controllable base of attachment for the tail feathers. (6) Of particular importance is the great development of the coracoids and furcula, which are constructed so as to prevent the shoulder sockets from being pulled downward or squeezed toward the midline by the powerful contractions of the flight muscles that originate on the sternum.

In addition to these skeletal specializations, the pectoral and forelimb musculature of carinates have also been highly modified from the primitive tetrapod condition, to the extent that in some instances homologies are very much in doubt. Fortunately, the establishment of homologies is not critical for the theoretical reconstructions and interpretations that follow here. The flight musculature of modern carinates has been studied and described by many authorities, among them Stresemann (1933), Sy (1936), Fisher (1946), Hudson and Lanzillotti (1955), Berger (1960), and George and Berger (1966). From these studies, we may classify the flight muscles in six broadly functional categories as follows: (1) those that fix or adjust the pectoral girdle and the shoulder socket; (2) those that power the wing, producing the propulsive down stroke; (3) those producing the recovery stroke of the wing; (4) the flexors, for folding the wing; (5) the extensors, for unfolding the wing, and (6) the muscles that produce minor adjustments of the wing components, including the remiges. Some thoracic and appendicular muscles are involved in two or more of these actions. The following tabulation summarizes the principal muscles in each of these generalized categories. In the discussion that follows, the emphasis will be on those muscles that are concerned with the power and recovery strokes of the wing, not because other muscles are less important, but because these are more conspicuously involved in the evolutionary changes that occurred between Archaeopteryx and later birds.

### Shoulder Joint Fixors and Adjustors
- Rhomboideus superficialis
- Rhomboideus profundus
- Serratus superficialis anterior

### Flight Muscles

#### Pectoralis superficialis

#### Wing Recovery Muscles
- Supracoracoideus
- Coracobrachialis anterior
- Deltoideus major anterior

#### Wing Folders
- Latissimus dorsi anterior
- Latissimus dorsi posterior
- Scapulohumeralis anterior
- Scapulohumeralis posterior
- Coracobrachialis posterior
- Subcoracoideus
- Subscapularis

#### Wing Unfolders
- Coracobrachialis anterior
- Triceps brachii
- Extensor metacarpi radialis
- Extensor digitorum communis

#### Wing Adjustors
- Serratus superficialis
- Pectoralis propatagialis
- Pectoralis propatagialis longus
- Pectoralis propatagialis brevis
- Cucullaris propatagialis
- Propatagialis longus
- Propatagialis brevis
- Expansor secundariorum
- Pronator sublimis
- Pronator profundus
- Entepicondylo-ulnaris
- Flexor carpi ulnaris
- Ulnometacarpalis ventralis
- Extensor metacarpi radialis
- Supinator
- Extensor digitorum communis
- Extensor carpi ulnaris
- Extensor carpi ulnaris

Powered avian flight is produced by synchronous down strokes of the wing caused by contraction of the large ventral muscle complex, the M. pectoralis. This complex usually consists of three or four distinct muscles, the M. pectoralis thoracica, or pectoralis superficialis, being the largest and most important. The other pectoralis muscles typically are small slips that function to tense the protopatagium, thus belonging to the last category listed above. The M. pectoralis superficialis originates extensively on the posterior and lateroventral surfaces of the sternum, the ventral half of the entire length of the carina, the entire posterolateral
surface of the clavicle and the anterior margin of the sterno-coracoclavicular membrane. The pectoralis tendon inserts broadly on the ventral surface over most of the length of the deltopectoral crest (crista lateralis humeri) of the humerus. This muscle provides nearly all the force for flight and is the largest of all avian muscles, averaging more than 15 percent of total body weight among all flying birds (Hartman, 1961; Greenwalt, 1962). Two osseous features reflect the size and functional importance of this muscle: the very large sternum and its carina, and the long and prominent deltopectoral crest of the humerus.

**Wing elevation (recovery stroke)** is accomplished by the combined actions of several muscles: the M. supracoracoideus, M. coracobrachialis anterior and Mm. deltoideus major and minor. Of these, the supracoracoideus is by far the most important. The coracobrachialis, by virtue of its origin on the anterodorsal extremity of the coracoid (the acrocoracoid) anterior and dorsal to the glenoid fossa, provides some lifting of the humerus, but its chief action is to extend or pull the humerus forward, thereby unfolding the wing. Typically, it is the smallest "elevator" muscle. The M. deltoideus major usually consists of a pars anterior and pars posterior. The pars anterior arises from a small area on the dorsal side of the scapula adjacent to the glenoid. The pars posterior originates on the dorsal end of the clavicle and the anterodorsal surface of the scapula. Accordingly, these fibers tend to elevate the humerus and draw it forward. The M. deltoideus minor also originates on the anterodorsal apex of the scapula, above, medial, and slightly anterior to the glenoid, hence also acting to elevate the humerus.

The largest humeral abductor, as noted above, is the M. supracoracoideus, also termed the pectoralis secundus or pectoralis minor (Figure 2). This muscle arises by extensive attachment on the dorsal parts of the sternal carina, the anterolateral surfaces of the sternum, the ventro-anteromedial surface of the coracoid and the lateral part of the coracoclavicular membrane. Its fibers converge dorsally, attaching to a narrow tendon that passes backward through an osseous canal, the foramen tririoso, between the dorsal extremities of the coracoid and clavicle and the anterior extremity of the scapula. From there, the tendon turns downward to insert on the dorsal surface of the humerus between the head and the deltopectoral crest. The fact that the triosseal canal is situated above the insertion point when the humerus is depressed allows this ventrally placed muscle to *elevate* rather than depress the humerus.

**Figures 2 and 3** illustrate the structure of the triosseal canal and its relationship to the supracoracoideus muscle. Of particular importance is the very prominent dorsal process of the coracoid (the acrocoracoid) that extends well above and anterior to the glenoid. The medial side of this process forms the lateral wall of the triosseal canal. A further factor of importance is that two important muscles arise from the upper anterior surface of the acrocoracoid, the M. coracobrachialis anterior and the M. biceps brachii. As noted earlier, the coracobrachialis anterior is a primary extensor of the humerus and the biceps is equally important as the principal flexor of the humerus.
forearm. It is safe to assume that the elevated positions of these origins at the apex of the acrococloid have functional significance.

Without concerning ourselves with homologies, or the proper name for the avian "supracoracoideus," the action of that muscle in modern carinates emerges as extremely important for reconstructing some of the details of avian evolution. By the nature of its location and architecture, it is clear that at some earlier stage in the evolution of birds the antecedent of this muscle must have acted to depress the arm. Therefore, its action has been completely reversed, probably as a consequence of the development of the pulley-like arrangement of the triosseal canal and its interposition between the points of origin and insertion. The avian wing is elevated chiefly by this ventral muscle, rather than by dorsal muscles as we would expect, and as is the case in bats.

The fact that virtually all muscles in all organ-
isms follow the most direct route between the points of origin and insertion argues strongly against the possibility that the insertion of the supracoracoideus gradually migrated to the dorsal side of the humerus, without prior or concurrent deflection of the fibers or tendon leading to that insertion. Even if the insertion had shifted to a dorsal position on the humerus, contraction of the muscle would still depress, as well as rotate, the humerus—unless the fibers approached the humerus from above. Consequently, the most logical explanation of the peculiar organization and action of the modern avian supracoracoideus would seem to be that its path was altered during the course of avian evolution. Modern carinates, together with the specimens of Archaeopteryx, establish that these postulated changes resulted from drastic changes in the shape of the coracoid and that these changes occurred subsequent to the Archaeopteryx stage.

"Flight" Apparatus of Archaeopteryx

The portion of the skeleton of Archaeopteryx that can be equated with the flight apparatus of modern carinates displays a number of important features:

1. There appears to be little or no loss of flexibility in the trunk region, either by vertebral fusion or by restriction of vertebral articular freedom. Although fully ossified gastralia are present, there is no evidence of ossification of either sternal ribs or the sternum. Also, there are no uncinate processes on the dorsal ribs.

2. The pectoral arch does not appear to have been as rigidly fixed as in modern birds. The coracoids are short, subquadrangular, not strut-like, and had only cartilaginous or membranous contact with the sternum. The clavicles, however, were fused and fully ossified into a robust furcula, but the nature of its contacts with the scapulocoracoid are not known.

3. Contrary to de Beer's (1954) interpretation, no sternum is preserved in any of the presently known specimens of Archaeopteryx (Ostrom, in press). This indicates that the sternum was almost certainly cartilaginous and probably lacked a keel. It may even have been membranous. Furthermore, the space anterior to the gastralia is quite short, a clear indication that the sternum, whether ossified or not, could not have been enlarged, as it is in all modern carinates.

4. The forelimb is elongated, but it does not possess any of the skeletal specializations of modern carinates that are usually equated with avian flight. The deltopectoral crest of the humerus is comparable to that of small theropod dinosaurs and is longer and more elevated above the shaft than is typical of most carinates. The elbow and wrist joints are unmodified, the carpals and metacarpals are not fused and digits I, II, and III are separate and unfused. The London and Berlin specimens clearly show that the forelimbs bore large, remex-like feathers, but it is uncertain whether these feathers were attached directly to the forelimb skeleton as in modern birds and as would seem to be required of true "flight" feathers. Despite exceptional preservation of several of the specimens, none shows anything that can reasonably be interpreted as quill nodes on the ulna. This is negative evidence only, but a further indication that the "flight" feathers were not firmly attached to the skeleton is the fact that imprints of the "primaries" of both wings in the London specimen are preserved with only slight disarrangement, yet the left hand is disarticulated and the right hand is missing altogether.

5. The long reptilian tail of Archaeopteryx bore feathers, but there is no indication in any of the specimens that the caudal series was undergoing reduction or fusion into a pygostyle. On account of the feathers, we can conclude that the tail may have functioned as an aerodynamic, rather than an inertial, stabilizer, but this should not be construed as proof of flight capability in Archaeopteryx.

The more important of the above conditions in Archaeopteryx are the nonavian form of the coracoid, the absence of an ossified sternum, the unfused carpometacarpus and the unfused digits of the manus. As Figures 4 and 5 show, the coracoid of Archaeopteryx is not elongated, and clearly did not serve as a strong, anticompressive brace against the sternum. It appears to have been fused with the scapula and its sternal border, although not as robust as the scapular margin, is well defined, but thin. The glenoid segment is stout, a relatively large supracoracoid foramen is present and a very prominent lateral process occurs just anterior to and below the glenoid. This last feature, some-
Figure 4.—Three views of the pectoral girdle of *Archaeopteryx* as reconstructed from the London, Berlin, and Maxberg specimens: a, anterior view of the left coracoid; b, lateral view of the left scapulo-coracoid; c, dorsal view of the left scapulo-coracoid.

Figure 5.—Left coracoid and glenoid of the London specimen of *Archaeopteryx*, as seen in anterior view from the underside of the main slab. (The smallest divisions on the scale equal 0.5 mm.)
times referred to as the biceps tubercle (Walker, 1972), is of special significance because it appears to be the precursor of the avian acrocoracoid. Contrary to Bakker and Galton’s (1974) interpretation, the glenoid does not face downward, but is directed laterally (Figures 4 and 5) more or less as in modern carinates (Figure 3).

The Transition from *Archaeopteryx* to Modern Birds

Changes in the Pectoral Girdle

Before attempting to reconstruct hypothetical transitional stages in the evolution of the pectoral arch between *Archaeopteryx* and modern birds, it may be useful to review certain facts. First, the coracoid of all lower tetrapods, including birds, has certain constant relationships with other elements of the trunk. It occupies a position between the scapula (with which it usually forms the shoulder socket) and the sternum, regardless of whether the latter is ossified or cartilaginous. Thus, at least two regions of the coracoid, the sternal border and the scapular border, are unmistakable reference points no matter what the shape or size of the coracoid. Similarly, the glenoid portion is always recognizable.

The second consideration is the role of the coracoid in forelimb biomechanics of lower tetrapods. Chief among the various muscles that attach to the coracoid (most of which insert on the humerus) is the biceps, the principal flexor of the antebrachium. (A structural and functional analog, the M. coracoradialis proprius, is present in amphibians.) The biceps passes between the coracoid and the internal proximal surfaces of the radius and ulna. Even in mammals, where the coracoid is no longer present as a separate bone, the major forearm flexor (which also happens to be termed the biceps) originates on the presumed relict of the coracoid, the coracoid process of the scapula.

The final consideration is that the location of the flexor origin relative to the glenoid fossa determines the approximate path of forearm flexion. Thus, for any given position of the humerus, the approximate orientation of the plane of forearm flexion can be determined from those two points. For example: the biceps brachii of lizards originates on a small area adjacent to the sternal border of the coracoid, anterior and ventral to the glenoid. With the humerus positioned in a horizontal transverse position, the biceps flexes the forearm anteroventrally toward the midline. But with the humerus extended forward, forearm flexion is down and *backward*. In birds, the site of origin of the biceps on the anterolateral surface of the acrocoracoid is situated in front of and above the glenoid; consequently, forearm flexion is restricted to a forward movement (Figure 6).

![Figure 6.—Dorsal aspect of the wing skeleton and pectoral girdle of *Corvus brachyrhynchos*, showing the location and action of the M. biceps brachii (heavy arrow), the chief flexor of the forearm in modern birds.](image)

If, as seems reasonable, we accept the so-called biceps tubercle of *Archaeopteryx* as the homolog of the acrocoracoid of modern birds and the probable site of origin of the chief flexor of the forearm (whatever we call it), we can reconstruct the general nature of forearm flexion in *Archaeopteryx*. Although the precise orientation of the scapulocoracoid in *Archaeopteryx* cannot be established from any of the presently known specimens, there can be little doubt that the biceps tubercle was situated well below and anterior to the glenoid (Figure 4b). Consequently, there must necessarily have been a major *downward* component in forearm flexion, regardless of whether the humerus was extended, retracted, or even adducted.

Transformation of the avian coracoid from the condition in *Archaeopteryx* to that of modern birds involved two major changes: the dorsoventral elongation of the main body of the coracoid and the raising of the site of origin of the M. biceps brachii by anterodorsal prolongation of the acrocoracoid. Elongation of the coracoid increased the distance between the glenoid and the sternum, pre-
Figure 7.—Hypothetical stages in the evolution of the avian coracoid from the *Archaeopteryx* stage to that of a modern carinate (*Cathartes*). The arrows indicate the hypothesized course of the M. supracoracoideus fibers in each stage and their progressive deflection resulting from evolutionary elevation and expansion of the biceps tubercle (= acrocoracoid). Upper arrows indicate the line of action of the supracoracoideus at each stage. Dashed lines indicate the acromion and adjacent regions of the scapula. All stages are of a left coracoid viewed from the front.

sumably increasing the range of dorsoventral humeral excursion. This in turn may have been correlated with the anteroposterior elongation of the sternum, the development of the sternal keel, and the enlargement of the ventral adductor muscles—the M. pectoralis. Increased force of forelimb adduction, for whatever biological role, required strengthening of the coracoid into a strong, anti-compressive strut between the shoulder socket and the enlarged muscle origins on the sternum.

Because the supracoracoideus of lower tetrapods originates ventral and anterior to the glenoid, and because it also has a ventral origin close to the sternal border in modern birds, the primitive site of origin of this muscle in *Archaeopteryx* probably was in a similar position—ventral and somewhat medial to the biceps tubercle. If so, then any upward expansion of the biceps tubercle would have impinged against the supracoracoideus tendon, gradually deflecting its course medially around the base of the expanding “protoacrocoracoid.” Once the base of this process reached the level of the glenoid, the then-deflected supracoracoideus would have pulled the humerus anteromedially, rather than downward. Continued expansion and elevation of the acrocoracoid would have resulted in further deflection of the supracoracoideus. The action of this muscle almost certainly was not reversed abruptly, but probably changed gradually from that of a humeral adductor, to an anteroventral extensor, to a forward extensor, to an antero-dorsal extensor and finally becoming an abductor of the humerus. Figure 7 illustrates how this transformation may have taken place.

If the above reconstruction is even approximately correct, it is clear that one of the major factors in the evolution of avian flight structures was the upward expansion of the acrocoracoid. This conclusion is established beyond any doubt by the presently reversed action of the supracoracoideus in modern birds. The critical question is: What brought about the upward expansion of the acrocoracoid? There appear to be several possibilities: (1) elevation of the anterior part of the glenoid and rotation of the shoulder socket to face directly laterally, thereby permitting unrestricted transverse (up and down) movements of the forelimb; (2) provision of an enlarged buttress at the level of the glenoid for the furcula to brace against, thereby insuring proper transverse separation of the shoulder sockets; (3) raising of the levels of humeral extension and forearm flexion by elevat-
ing the sites of origin of the coracobrachialis and biceps. In all probability, none of these factors
acted alone, and other less obvious factors may have been involved as well.

Whether enlargement of the pectoral adductor muscles and the elongation of the coracoids into
robust struts occurred before, after, or concurrently with upward expansion of the acrocoracid cannot
be determined in the absence of intermediate stages in the avian fossil record. Whatever the se-
quencing, the upward growth of the acrocoracid would have progressively deflected the action of
the supracoracoideus. It also brought about signifi-
cant changes in other forelimb movements, especially in elevating the range of humeral ex-
tension and increasingly confining it to the cranial sector. As a direct consequence, the level of
forearm flexion was also elevated to a nearly hori-
Zontal fore-aft plane more or less perpendicular to
the transverse, up and down, humeral movements
produced by the enlarged pectoral muscles.

Considering these three possibilities, it appears
that the glenoid in Archaeopteryx already faced
laterally and slightly forward (Figure 4b,c) not
ventrolaterally, as Bakker and Galton (1974)
claim. The coracoid portion of the glenoid also
seems to have been elevated. Yet, the biceps
tubercle was still small and located well below the
glenoid. Also, as was noted earlier, a robust furcula
is present in Archaeopteryx (as seen in the Lon-
don [Figure 8] and Maxberg specimens), and al-
though the nature of its articulations with the
other elements of the pectoral girdle is not clear,
there does not appear to have been any special
structure of the coracoid that might have served to
buttress it, since, as already noted, the biceps tu-
bercle is not elevated. This, of course, raises the
question of the function of the furcula in Archaeo-
pteryx. Did it serve as a transverse spacer between
the shoulder sockets? If so, it would appear to have
been related to some activity other than powered
flight—perhaps predation. Since both the M. cora-
cobrachialis anterior and the M. biceps brachii
arise from the upper anterior surface of the acro-
coracoid in all modern carinates, then by virtue of
their positions above and in front of the glenoid,
these muscles, respectively, pull the humerus for-
ward and up, and flex the forearm forward and in-
ward toward the midline. In Archaeopteryx, the
humerus apparently could not be extended for-
ward and upward above the level of the shoulder
because no part of the coracoid was situated above

Figure 8.—The furcula of Archaeopteryx as preserved in the London specimen. The exposed
surface is probably the anterior surface. (The smallest divisions on the scale equal 0.5 mm.)
and in front of the glenoid (Figures 4 and 9). Thus, of the three possibilities suggested above, the evolutionary expansion of the avian acrocoracoid would seem to have been most critically linked with the actions of the coracobrachialis and biceps muscles. There appears to have been some selective advantage in raising the level or attitude of forelimb extension and forearm flexion.

So far, I have given little attention to the scapula. This is because the scapula of Archaeopteryx already had acquired a form remarkably similar to that of modern birds, being very long, narrow and strap-like. Its principal distinctions from the condition in most modern birds are its fusion to the coracoid, the form of the acromion, and the shape of the distal extremity, which is rectangular or slightly flared in Archaeopteryx rather than tapered. The fact that the acromion is more prominent and robust than in most modern birds suggests that the M. deltoideus was perhaps a more important humeral elevator at the Archaeopteryx stage of avian evolution. This would be consistent with the conclusion reached above that the supracoracoideus of Archaeopteryx could not have elevated the humerus (as was noted by Walker, 1972), but rather must have been a lateral adductor. If the deltoideus, however, was more important as a humeral elevator at the Archaeopteryx stage than it is in modern carinates, then it would appear that the force of the recovery stroke must have continued to decline in birds succeeding Archaeopteryx, until complete deflection of the supracoracoideus was accomplished. This implies that there probably was no tendency at the Archaeopteryx stage, or immediately afterward, toward powered flight.

It should also be noted here that the stout acromion in Archaeopteryx may not have had anything to do with the deltoideus muscles, but might have served as a buttress for the stout furcula. This cannot be established on the basis of present specimens, however.

The narrow form of the scapula, as compared with the broad, triangular form in all other tetrapods except theropod dinosaurs, suggests that the musculature that inserted or originated on the scapular blade—and particularly on its dorsal surface—was greatly reduced. This certainly is true of modern birds in which the M. rhomboideus and M. scapulohumeralis (the largest dorsal shoulder muscles) are of relatively small size. The fact that this narrow scapular form occurs only in obligate bipeds (birds, Archaeopteryx, and theropod dinosaurs), but not in facultative bipeds (such as non-human primates, kangaroos, or ornithopod dinosaurs), or in any quadrupedal animal is highly suggestive. It indicates that strong stabilization of the pectoral arch by muscles connecting the scapular blade with the vertebral column and dorsal ribs, and powerful abduction of the limb by large muscles extending between the humerus and the scapular blade, were unnecessary in obligate bipeds in which the forelimb was no longer involved in weight support.

**Changes in the Forelimb**

Comparison of the forelimb skeleton of Archaeopteryx with that of modern birds reveals several major differences, the most conspicuous of which
Figure 10.—The right manus and carpus of the Berlin specimen of Archaeopteryx, seen in dorsal aspect. Notice the separated fingers and the unfused metacarpus and carpus, as well as the extent of lateral flexion. (The smallest divisions on the scale equal 0.5 mm. Roman numerals identify the digits.)

occur in the hand and wrist. Figure 10 shows the right hand and wrist of the Berlin specimen of Archaeopteryx with its unfused metacarpus and three separated fingers. The same construction is present in the other three specimens in which the hands are preserved. This construction is in sharp contrast to the united metacarpus and manus of modern birds (Figure 11). It is obvious that phalanges have been lost or co-ossified in at least the external finger (digit III) of modern birds, but the most interesting changes have taken place in the metacarpus and wrist. Figure 12 illustrates the carpus and metacarpus as they are preserved in the Berlin (Figure 12a) and Eichstädt (Figure 12b) specimens, compared with the same elements of a modern carinate, Cathartes aura (Figure 12c,d). The first metacarpal is considerably shorter than the other two (Figure 10), as it is in modern forms, but it does not appear to be co-ossified with metacarpal II, nor are the second and third metacarpals fused. The carpus consists of only three elements, a large distal carpal with a distinctive semicircular proximal profile, and two smaller bones, which probably represent the radiale (scapulolunar) and the unare (cuneiform). Although neither of the last two elements resemble modern bird carpals, two features in Archaeopteryx do preview specialized conditions of the modern avian carpometacarpus. These are the large lunate distal carpal that is closely articulated with the first and second metacarpals (Figure 12a,b), and the internal expansion at the base of metacarpal I. There can be little doubt that the lunate carpal of Archaeopteryx, by fusion with the two metacarpals, became the pulley-like trochlea of the carinate carpometacarpus. The proximal internal expansion at the base of the first metacarpal in Archaeopteryx is almost certainly the precursor of the large extensor process (processus metacarpalis I) of the modern carpometacarpus. In Figure 13, I have attempted to show how the modern avian carpometacarpus probably evolved from the condition in Archaeopteryx.

Reconstructing the above intermediate stages is far simpler than trying to account for the conditions that brought about such changes. The second digit clearly was the dominant finger and ultimately became the main supporting structure of
the primary remiges. The third or outermost finger gradually was reduced and metacarpal III was modified into a posterior (external) brace against metacarpal II. These changes could well have come about in connection with flight and the attachment of the primaries to the second metacarpal, presumably bracing it against lift forces that would tend to rotate the second digit and metacarpal upward. Fusion of the lunate carpal to the metacarpus, and its expansion into the pulley-like trochlea, increased the degree of flexion possible at the wrist, but at the same time reduced wrist mobility to the plane of the metacarpus and the wing. The prominent extensor process of the modern carpometacarpus is the point of insertion of the largest muscle of the avian forearm, the M. extensor metacarpus radialis, the action of which extends or unfolds the hand (Figure 14).

In the discussion that follows, it is essential to distinguish between two very different kinds of flexing movements at the wrist: that in which the extremity is flexed toward the ulnar or external side of the forearm (termed lateral flexion here) and that in which the hand and metacarpus are “flexed” inward toward the radial side of the forearm. This last movement might be termed “medial flexion,” but for the sake of clarity it is designated here as “hyperextension.” These terms differ from the usual terminology applied by ornithologists (which by convention is in terms of a laterally extended wing), but hopefully they will be clear to all readers. The term extension is used here in the sense of straightening the wrist, and where necessary for clarity, it will be specified as extension from the laterally flexed or the hyperextended condition.

In the Berlin and Eichstätt specimens of Archaeopteryx, the hands are flexed laterally toward the ulna at about 80 degrees to the radius and ulna. Close examination of the wrist in each case (Figure 12a,b), and especially of the morphology of the lunate carpal and the external aspect of the ulnar extremity, reveals that in both specimens the wrists are fully flexed. Notice that the internal condyle or condylus metacarpalis does not extend proximally along the outer surface of the ulnar shaft as it does in modern birds. For contrast, Figure 12c shows the much greater maximum degree of lateral flexion (hyperflexion) possible in the modern bird wrist. Also conspicuous in modern birds is the elongated extensor process of the carpometacarpus, which greatly increases the leverage of the principal extensor of the hand.

It is tempting to relate these features to some aspect of flight; for example, the need for adjusting or changing the surface area of the airfoil by improved efficiency and precision of extension and flexion at the wrist. Once flight capability had been achieved, increased leverage for the M. extensor metacarpus radialis would reduce the amount of energy required to counteract the force of the airstream that tends to flex or fold the wing extremity laterally. On the other hand, during the power stroke, lift forces tend to open or extend the wing extremities. Another possibility is that the extensor process grew larger in conjunction with the development of wrist hyperflexion, which in turn was made possible by gradual expansion of
Figure 12.—The wrists of Archaeopteryx (a and b) and Cathartes aura (c and d) as viewed from above: a, left wrist of the Berlin specimen; b, right wrist of the recently recognized Eichstätt specimen (b and c are preserved flexed laterally, toward the ulnar side of the forearm, to the maximum degree possible); c, left wrist of Cathartes drawn in the same laterally hyperflexed position to show the greater degree of flexion possible in modern carinates; d, “exploded” dorsal view of the right wrist of Cathartes, flexed to the same degree as b, to show the specialized facets of the wrist elements, arrows indicating complementary articular facets. Notice in particular the lengths of the external portions of the condylus metacarpalis of the ulna and also the trochlea carpalis of the carpometacarpus, as compared with the corresponding regions in Archaeopteryx. Also notice the large extensor process of the carpometacarpus compared with the modest expansion on metacarpal I of Archaeopteryx. The phalanges have been omitted from digit I in a and b. (Roman numerals identify the metacarpals; the horizontal lines equal 10 mm).

the trochlea carpalis of the carpometacarpus and elongation of the condylus metacarpalis of the ulna. A critical point here, however, is that extreme hyperflexion of the manus has no obvious “flight” advantage, but it clearly is advantageous for compact folding of the forelimb extremities to protect the airfoil when not in use. Under these circumstances, it would appear that the increased extension leverage that was provided by a larger extensor process on the carpometacarpus was not related to the first explanation above, but probably was advantageous for quick unfolding of a hyperflexed wing. This interpretation is reinforced when it is considered in conjunction with the unique linkage between the modern avian elbow and wrist that automatically synchronizes flexion (or extension) at those two joints. As first observed by Coues (1871) and Headley (1895), and confirmed by Fisher’s (1957) experiments, the radius of birds functions as a “connecting rod” between...
the elbow and wrist. Because of the greater length of the radial versus the ulnar condyle of the humerus, the radius slides distally along its axis when the elbow is flexed by the M. biceps, thereby pushing against the carpus and metacarpus and forcing the wing extremity to flex. Extension of the wrist and elbow are similarly linked. In fact, because of the "connecting rod" action of the radius, and the increased leverage of the M. extensor metacarpus radialis provided by the enlarged extensor process, it is possible for that muscle to function as the primary unfolder of the entire wing, not just of the hand. Although smaller than the M. triceps brachii, the forearm extensor, the M. extensor metacarpus radialis of most carinates has far better leverage than the triceps (which inserts on the olecranon) for extending the wing extremity.

It is not possible to establish which, if either, of the above possibilities was the decisive factor in the evolution of the modern avian wrist, but the specimens of *Archaeopteryx* seem to provide a clue. All four of the specimens in which the hand is preserved show what appears to be a maximum degree (about 80°) of flexion of the hands toward the ulnar side of the forearm. In other words, the hand could not be hyperflexed or folded back tightly against the forearm as in modern birds. Also, the extensor process is only very weakly developed in these specimens. The nature of the articular surfaces in the wrists of the Berlin and Eichstätt specimens, however, indicates that the hands almost certainly could have been hyperextended medially, or bent toward the radial side of the forearm, to about the same degree that they are preserved flexed laterally toward the ulna, perhaps even more so. This last is important, because medial hyperextension of the hand is not possible in modern birds. In fact, the manus cannot even be fully extended to align parallel with the radius and ulna. From this, the most probable conclusion is that the extensor process is most important for recovery (extension) of the avian manus from a folded or laterally hyperflexed condition. If it had developed for enhancing medial hyperextension it is difficult to understand why this process was retained, even enlarged, while at the same time...
the capacity for medial hyperextension of the hand was being reduced and ultimately eliminated.

The rest of the forelimb appears to have been altered in much less conspicuous ways during the transition from Archaeopteryx to modern birds, yet those changes that can be recognized may have significant implications. In the ulna, the most obvious changes involved the external expansion and elongation of the condylus metacarpalis (Figure 12), the articular facet of which permits hyperflexion of the manus laterally. Less obvious is the apparent lack of direct attachment of the secondary remiges to the ulna, or of the primaries to the metacarpus, as is indicated by the absence of quill nodes. These conditions are lacking in the specimens of Archaeopteryx, but are well developed in a variety of modern carinates.

The humerus of Archaeopteryx, although very bird-like, is much simpler than that of modern carinates. There is a long and well-defined deltopectoral crest, but as can be seen in Figure 15, this crest projects farther from the shaft than is characteristic of most later birds. More important, though, are the features that are missing from the humerus of Archaeopteryx. There is no sign of either the external or internal tuberosity, nor is there a bicipital crest. Distally, the ectepicondyle is also absent, or at least there is no detectable tubercle preserved in presently known specimens. In view of the other related features of the distal segments of the forelimb of Archaeopteryx, the absence of these processes seems to have special significance, because in modern carinates they play a direct part in the compact folding of the wing.

The internal tuberosity (tuberculum mediale) is the site of insertion of the three principal humeral retractors (the M. subcapularis, M. subcoracoideus, and M. coracobrachialis posterior). The external tuberosity (tuberculum laterale) is the site of insertion of the M. supracoracoideus which, in addition to elevating the wing, also rotates the entire folded wing dorsally toward the midline in modern birds. In Archaeopteryx, however, this muscle must have been a humeral depressor, as has been emphasized above. The bicipital crest (crista medialis) of modern birds is the area of insertion of the M. scapulohumeralis posterior, which draws the humerus back against the body. The implications of these conditions are obvious: in the absence of all specialized features of the humerus, ulna, and carpometacarpus that in modern birds are directly related to the folding of the wing, we are forced to conclude that Archaeopteryx was unable to fold the forelimb back against the body as in modern birds. Add to this the absence in Archaeopteryx of an ectepicondyle, which is the site of origin of the M. extensor metacarpus radialis, and also the weak development of the extensor process of metacarpal I, which is the site of insertion of this same muscle in later birds. These conditions indicate that powerful or rapid extension of the manus was unlikely, and probably not necessary, because the wrists of Archaeopteryx clearly show that lateral hyperflexion of the manus was not possible. On the other hand, a high degree of medial hyperextension was retained, perhaps as a critical action for prey catching or feeding activities.

Discussion

It would appear that the acquisition of obligate bipedal posture and locomotion in some pre-Archaeopteryx stage of avian evolution was responsible in large part for the ultimate development of powered avian flight. An early consequence was the narrowing of the scapula. Strong stabiliza-
tion of the scapula and shoulder joint, and powerful abduction of the forelimb became less critical than in obligate or occasional quadrupeds (whether of sprawling or upright posture), where antagonistic and synergistic interaction of dorsal abductors and ventral adductors are necessary for precise dynamic control of limb movements and positions under weight-bearing conditions. By the Archaeopteryx stage, forelimb abduction may have been accomplished solely by the action of a reduced remnant of the M. deltoideus that presumably originated on the prominent acromion.

With the assumption of upright, obligate bipedal posture and the release of the forelimbs from a weight-supporting role, new forelimb functions became possible. At the Archaeopteryx stage these functions apparently involved laterally elevated movements of the forearm (as indicated by the outward facing glenoid), anteroventral extension of the humerus (as indicated by the anteriorly facing surface of the coracoid below the level of the glenoid—the only available site of origin for humeral extensors), and powerful anteroventral flexion of the forearm toward the midline (as indicated by the prominent biceps tubercle below and anterior to the glenoid—the most probable site of origin of the forearm flexor). The hands were capable of nearly 180 degrees of lateral flexion and medial hyperextension, as noted above. The capacity for extreme hyperextension at the wrist (not possible in modern birds), coupled with the evidence for strong flexion of the forearm toward the sagittal plane, appears to be especially significant. Perhaps even more significant is the evidence that the forelimb of Archaeopteryx probably could not have been raised above the level of the glenoid when in the anteriorly extended position, simply because no part of the shoulder girdle was situated above and anterior to the glenoid. The strong anterior extensor (M. coracobrachialis anterior) and forearm flexor (M. biceps brachii) of modern birds have their present actions only because of the elevated positions of their origins on the acrocoracoid. The evolutionary upward expansion of the acrocoracoid would seem to have been linked causally with the actions of those two muscles and most especially with that of the humeral extensor. Selection apparently favored the elevation of forearm and hand activities.

As observed above, it is tempting to equate such changes with some aspect of flight. For example, these changes might permit alterations in the surface area of the "wing" by means of flexion or extension of distal components more or less in the plane of the "wing." Notice, however, that these capabilities apparently were not yet available in Archaeopteryx, where wrist and elbow movements were not restricted. Another possibility is that elevation of forelimb extension and forearm and hand flexion and extension ostensibly might improve the aerodynamic qualities of an incipient "wing" by making possible a positive angle of attack (where the leading edge of the airfoil is above the trailing edge, relative to the airflow or flight path). A positive angle of wing attack is essential for all forms of flight, whether powered or passive, because without it there can be no lift. That being true, then there is a critical flaw in attributing the above anatomical changes between Archaeopteryx and modern birds to aerodynamic adaptations. The flaw is that there can be no lift, and thus no aero-kinetic selective advantage in raising the attitude of a potential airfoil until after the smallest degree of a positive angle of attack has been acquired.

An aerodynamic explanation of the anatomical changes noted above is also weakened by the absence of an ossified sternum in all specimens of Archaeopteryx. The absence of a sternum strongly suggests that the "flight" muscles of Archaeopteryx were not of unusual size, a conclusion that is substantiated by the short space available for the sternum in front of the ossified gastralia, as well as by the short nonstrut-like form of the coracoid. If all these assessments are correct, then some biological role other than flight must have been involved in the initial and early phases of the upward expansion of the biceps tubercle into the future acrocoracoid.

Aside from making lift possible, the only other obvious consequence of raising the level of forelimb extension and flexion is to place the hands and their activities directly in front of and above the animal. Two activities immediately come to mind: climbing and prey-catching. Various authors (Bock, 1965, 1969; de Beer, 1954; Swinton, 1960) have interpreted Archaeopteryx as being an arboreal animal. I have argued that there is no compelling evidence for this (Ostrom, 1974), and instead, the skeletal anatomy of Archaeopteryx appears to have been adapted for ground-dwelling activities.
Even if *Archaeopteryx* were arboreal, however, a possibility that I do not deny, then it acquired its climbing skills *prior* to elevation of the acrocoracoid and the capacity of elevated forelimb extension, and after the acquisition of obligate bipedal posture.

Obviously the same is true of prey-catching and feeding activities of *Archaeopteryx*. If *Archaeopteryx* were insectivorous, as seems almost certain, it clearly must have been proficient at catching insects, whether it did so with its mouth by quick darting movements of the head on the long flexible neck, or by grasping them in the hands or snaring them beneath the forelimb plumage. Considering the general absence of flight-related skeletal structures in the forelimb and pectoral girdle, it does not seem unreasonable to conclude that the forelimbs of this obligatory bipedal predator must have taken part in prey-catching activities.

If the forelimbs of *Archaeopteryx* were used to catch prey, and if the original advantages behind the enlargement of the contour feathers of the forelimb was to enhance insect-catching skills (Ostrom, 1974), there would be very real selective advantages in any changes that increased the scope of forelimb movements, especially if we think in terms of leaping or flying insects.

At this point it is not possible to identify the exact activities or selective advantages that promoted the upward expansion of the acrocoracoid, but it seems clear that these were related to upward extension of the arms and hands. It also appears that flight was not a factor in these first modifications. It was perhaps only coincidental that once a certain degree of upward enlargement of the acrocoracoid had been accomplished, the action of the coracobrachialis anterior would have been supplemented by the newly deflected supra-acrocoraoides acting as an anterodorsal extensor of the humerus.

The various specialized features of the modern avian forelimb skeleton mentioned above (reduced fingers, fused carpometacarpus, novel tubercles and crests on the humerus) seem best explained as flight-related adaptations that appeared subsequent to the dorsal expansion of the acrocoracoid and the resultant ability to raise the attitude of the extended forelimb, thereby achieving at least a minimal positive angle of attack. Fusion of the metacarpus would solidify the structural support of the primary flight feathers and brace the second metacarpal against long-axis rotation resulting from lift forces. Phalangeal reduction may have been correlated with changing the function of the manus to that of an airfoil and the reduction of the primitive prey-grasping role of the long, separated fingers. Fusion of the distal carpal to the metacarpus reduced the amount of abduction-adduction possible at the wrist, but at the same time facilitated precise flexion-extension of the manus in the plane of the wing, essentially perpendicular to the powerful adductive actions of the enlarged pectoral muscles. The capacity for medial hyperextension of the manus was reduced and ultimately lost, presumably as the primitive avian hand became less and less involved with prey-catching and feeding activities and was increasingly adapted for flight-related functions. Later stages presumably involved development of structures related to compact folding of the wing and rapid unfolding—the various tubercles of the humerus noted above, the capacity for lateral hyperflexion of the manus, and the enlarged extensor process of the carpometacarpus.

**Summary**

The existence of several specimens of *Archaeopteryx*, the oldest known fossil remains that are universally accepted as avian, provides important anatomical details of an extremely early stage in bird evolution. Despite impressions of what appear to have been modern-type "flight" feathers attached to the forelimb (but possibly not attached to the forelimb skeleton), the five presently known specimens of *Archaeopteryx* show almost no osteological features that compare with the skeletal adaptations of the modern avian flight apparatus. The only exception is the furcula, preserved in the two largest specimens.

Assuming that *Archaeopteryx* is in fact an ancestral bird, and in the absence of any known intermediate structural stages in the avian fossil record between *Archaeopteryx* (of Late Jurassic age) and the essentially modern birds of Late Cretaceous and Early Tertiary ages, we can postulate only the most obvious structural changes that occurred during the evolution of the avian flight mechanism. From the *Archaeopteryx* stage, the
following sequence of developments seems probable, with the first two possibly taking place concurrently or, less probably, in reversed order: (1) Upward expansion of the biceps tubercle, thereby raising the sites of origin of the M. biceps and M. coracobrachialis and thus the level of humeral extension and forearm flexion—possibly in connection with prey-catching or feeding activities, or perhaps to facilitate quadrupedal climbing. A direct consequence of the expansion of the acrocoracoid was the conversion of the M. supracoracoideus from a humeral adductor to a humeral elevator. (2) Enlargement of the pectoral muscles for more powerful arm adduction, accompanied by enlargement and ossification of the sternum, and elongation and strengthening of the coracoids to immobilize the shoulder joints. (3) Attachment of the remiges to the ulna and the second digit to resist feather deflection during the wing downstroke. (4) Fusion of carpals and metacarpals into a united carpometacarpus for firmer fixation of the primaries, modification of the troclea carpalis to permit only planar flexion and extension at the wrist, and loss of phalanges from all three fingers. (5) Loss of medial hyperextension of the hand and development of the capacity for compact folding of the wing, due to elongation of the condylus metacarpalis of the ulna and various tubercles on the humerus. This was associated with enlargement of the extensor process of the carpometacarpus to provide leverage for rapid unfolding of the wing.

The occurrence of other changes in the musculoskeletal system that affected the flight apparatus cannot be determined in the above sequence, but powered flight, as opposed to either gliding or flapping leaps, almost certainly could not have occurred before the first three of the above stages had been completed.

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Evolutionary Significance of the Mesozoic Toothed Birds

Philip D. Gingerich

ABSTRACT

Well-preserved fossils of the Mesozoic toothed birds Archaeopteryx, Hesperornis, and Ichthyornis, and of the bird-like dinosaur Compsognathus, discovered in the 19th century, indicated to early evolutionary biologists that dinosaurs and birds were closely related, and that birds in all probability evolved from a dinosaur similar to Compsognathus. The modern ratites, sharing some distinctive similarities with Hesperornis, were regarded as survivors of a primitive initial radiation of birds. Several workers have subsequently challenged the idea that the Cretaceous birds Ichthyornis and Hesperornis had teeth or that they bore any similarity to the ratites. After careful study of the actual fossil specimens of Hesperornis, it is clear that this Cretaceous bird had toothed jaws and a palaeognathous palate, the latter condition being shared with ratites and certain dinosaurs. These and other characters place Hesperornis, like Archaeopteryx, in a position morphologically, as well as temporally, intermediate between dinosaurs and typical birds. The few significant features unifying the living ratites and tinamous all appear to be primitive characteristics, suggesting that ratites and tinamous are either survivors of an early radiation of birds, or are possibly a more recently derived artificial group in which primitive characters have reappeared secondarily through neoteny.

Introduction

The discovery of fossil birds with teeth was one of the most dramatic events in 19th century paleontology. In 1861 a partial skeleton of the feathered Archaeopteryx was discovered in the Jurassic deposits of Bavaria. In the next 16 years, skeletons of Ichthyornis and Hesperornis were discovered in the Cretaceous of North America and a more complete skeleton of Archaeopteryx was found in Germany. Surprisingly, the jaws of each of these birds bore reptile-like teeth. Being discovered only a few years after publication of The Origin of Species, toothed birds were much discussed in connection with Darwin’s evolutionary hypothesis.

As spectacular as the original discoveries were, it is remarkable in retrospect how little detailed study was made of the actual specimens until relatively recently. The history of the original discoveries of toothed birds, the initial recognition of their evolutionary significance, and their subsequent fate are reviewed here. The whole provides an interesting historical comment on the treatment of intermediate forms that do not conform to preconceived archetypical categorizations.

Acknowledgments.—I should like to acknowledge here the encouragement Dr. Wetmore gave to continued study of the Yale collection of Mesozoic birds when work was initiated on Hesperornis several years ago. My study of Hesperornis began, curiously enough, as a tutorial with K. S. Thomson on kinesis and jaw mechanics in fishes. Expanding the range of comparisons, the Mesozoic bird material at Yale was examined to determine the form of kinesis of primitive birds. When no simple answer was forthcoming, J. H. Ostrom authorized Peter Whybrow to undertake further preparation of the original specimens. Thus I am particularly indebted to Professors Thomson and Ostrom and to Mr. Whybrow for their assistance and encouragement.
In addition, Drs. C. C. Black, T. H. Eaton, and L. D. Martin permitted an extended loan of the University of Kansas skull of Hesperornis. Drs. C. E. Ray, Nicholas Hotton III, and Mr. Robert Purdy allowed me to borrow the cranial material of Hesperornis in the National Museum of Natural History. Drs. W. J. Bock, Pierce Brodkorb, Peter Dodson, J. T. Gregory, Hildegard Howard, P. V. Rich, and M. V. Walker have all assisted one way or another as well.

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Mesozoic Birds with Teeth

It is now generally admitted by biologists who have made a study of the vertebrates, that Birds have come down to us through the Dinosaurs, and the close affinity of the latter with recent Struthious Birds will hardly be questioned. The case amounts almost to a demonstration, if we compare, with Dinosaurs, their contemporaries, the Mesozoic Birds. The classes of Birds and Reptiles, as now living, are separated by a gulf so profound that a few years since it was cited by the opponents of evolution as the most important break in the animal series, and one which that doctrine could not bridge over. Since then, as Huxley has clearly shown, this gap has been virtually filled by the discovery of bird-like Reptiles and reptilian Birds. Compsognathus and Archaeopteryx of the Old World, and Icthyornis and Hesperornis of the New, are the stepping stones by which the evolutionist of to-day leads the doubting brother across the shallow remnant of the gulf, once thought impassable. (O. C. Marsh, 1877:352).

In 1859, perhaps the gravest deficiency of Darwin’s hypothesis of evolutionary descent was the rarity of intermediate forms in the geological record. Intermediate forms linking species into graded chains or linking major groups of animals to a common ancestor were at that time poorly known. Evidence remedying this deficiency was supplied in a most spectacular way by the discovery of several intermediate forms linking birds to a reptilian origin. Interestingly, each discovery of itself was insufficient to overcome archetypical categorizations of birds and reptiles, and a truly evolutionary view of both classes was necessary in order to interpret literally the clear evidence for bird-reptile relationships offered by the skeletons of Compsognathus, Archaeopteryx, and Icthyornis.

J. A. Wagner (1861) described a remarkably complete skeleton of a very small new dinosaur, Compsognathus longipes, from the Jurassic lithographic limestone of Solenhofen, Germany. In the same year H. von Meyer (1861) first published a notice on the skeleton of a bird from the same deposit, which he named Archaeopteryx lithographica. Having a dinosaurian skeleton, Compsognathus was clearly a variant of the “Bird type.” On the other hand, Archaeopteryx, with its distinct impressions of feathers, was from the beginning regarded as a variant of the “Bird type.” Influenced at least in part by Darwin’s dynamic view of evolution, T. H. Huxley was able to overcome his contemporaries’ fixed categorizations, even of groups as large as reptiles and birds, and he found in Compsognathus a bird-like dinosaur, and in Archaeopteryx the most reptilian of birds. Thus, Huxley (1868) confirmed the Darwinian expectation of intermediate forms linking birds and reptiles in the fossil record. Although the actual common ancestor of living reptiles and birds had not been found, Huxley judged from their morphology that late Jurassic birds and reptiles were clearly much more closely related than their living descendants seemed to suggest. This closer similarity of the early forms was itself strong evidence favoring Darwin’s dynamic view of evolutionary descent, as opposed to the then-prevailing view that living “reptiles” and “birds” were static groups persisting through time within some predetermined bounds.

There was, however, a limit to the intermediate position even Huxley would accept for Archaeopteryx. Thus, of the single skeleton of Archaeopteryx then known, he wrote “unfortunately the skull is lost” (Huxley, 1868:70), making no mention of an earlier paper by Sir John Evans (1865) describing a premaxilla with four teeth preserved among the other bones of the specimen. Evans’ note (1865:421) quotes a letter from von Meyer himself concerning the apparent association of a toothed premaxilla with Archaeopteryx:

Teeth of this sort I do not know in the lithographic stone .... From this it would appear that the jaw really belongs to the Archaeopteryx. An arming of the jaw with teeth would contradict the view of the Archaeopteryx being a bird or an embryonic form of bird. But after all, I do not believe that God formed his creatures after the systems devised by our philosophical wisdom. Of the classes of birds and reptiles as we define them, the Creator knows nothing, and just as little
of a prototype, or of a constant embryonic condition of the bird, which might be recognized in the Archaeopteryx. The Archaeopteryx is of its kind just as perfect a creature as other creatures, and if we are not able to include this fossil animal in our system, our short-sightedness is alone to blame.

The presence of teeth in the bird Archaeopteryx was apparently too reptilian a characteristic for even Huxley to accept.

O. C. Marsh was the first to discover the unequivocal presence of teeth in primitive birds, though he too was at the outset apparently unable to accept the evidence. In September 1872, Professor Mudge of Kansas presented Marsh with some fossils from the Cretaceous Niobrara Chalk, the formation from which Marsh had earlier described the headless skeleton of a large, flightless, diving bird as Hesperornis regalis. Marsh studied Mudge's new fossils, and in October published a note describing the postcrania of a new form of smaller volant bird, Ichthyornis dispar (Marsh, 1872a). A month later he published another note (Marsh, 1872b) on the jaws of a new small "reptile," Colosaurus mudgei, found in association with the remains of Ichthyornis. In the same month that Colosaurus was described (November, 1872), Marsh's assistant T. H. Russell discovered a nearly perfect skeleton of Hesperornis, again in the Niobrara Chalk. This new skeleton included a skull with associated toothed jaws (Figure 1). Immediately after the discovery of this skeleton of Hesperornis, Marsh published a short paper in February 1873 stating that the toothed jaws of "Colosaurus" actually belonged to Ichthyornis. Of Ichthyornis dispar, Marsh (1873: 162) wrote:

When the remains of this species were first described, the portions of lower jaws found with them were regarded by the writer as reptilian; the possibility of their forming part of the same skeleton, although considered at the time, was not deemed sufficiently strong to be placed on record. On subsequently removing the surrounding shale, the skull and additional portions of both jaws were brought to light, so that there cannot now be a reasonable doubt that all are parts of the same bird.

Although no mention was then made of the toothed jaws of Hesperornis, that discovery probably provided Marsh with the necessary corroboration for him to accept the previously evident association of toothed jaws with Ichthyornis. Two years after the toothed jaws of Hesperornis were first described by Marsh (1875), the Berlin specimen of Archaeopteryx was found about 10 miles from the original Solenhofen discovery, and its feathers, reptilian skeleton, and toothed jaws left no doubt about the reptilian ancestry of birds.

Beyond their importance in dramatically filling a gap in the fossil evidence of evolution originally available to Darwin, the three early avian fossils Archaeopteryx, Ichthyornis, and Hesperornis are of interest for another reason. Huxley (1868:74) originally interpreted the great similarity of Compsognathus as indicating a dinosaurian (more specifically, coelurosaurian) origin of birds:

Surely there is nothing very wild or illegitimate in the hypothesis that the phylum of the Class Aves has its foot in the Dinosaurian reptiles—that these, passing through a series of such modifications as are exhibited in one of their phases by Compsognathus, have given rise to the Ratitae—while the Carinatae are still further modifications and differentiations of these last, . . .

Similarly, Marsh (1880:189) saw in the skull of Hesperornis certain resemblances to the "Ratitae," a group he regarded as being survivors of an evolutionary stage intermediate between reptiles and the true "ornithic type."

Three principal ideas have come out of the early work of Huxley and Marsh: (1) that ratites are survivors of a primitive stock of birds, (2) that Hesperornis was similar to ratites, and (3) that Hesperornis and Ichthyornis actually possessed jaws with teeth. All three of these views have been challenged in the century since their first publication by Huxley and Marsh. Disagreement with these ideas has come in part from authorities urging caution in attempting any interpretation at all, but in most cases a strong contrary interpretation has been offered, usually without critical examination of even the evidence available to Huxley and Marsh. Advocating ratites as a derived group of birds, reconstructing Hesperornis with a "neognathous" skull, and denying the presence of teeth in Ichthyornis or Hesperornis have a common effect—to deny the primitiveness and the reptilian characters of the best known Cretaceous birds and to maintain a wide gulf between birds and reptiles. This common effect of so many studies by post-Darwinian evolutionary biologists can only be ascribed to a deep-seated typological conception of "birds" and "reptiles"—an interesting comment on the pervasiveness of typological thinking.
The Skull of Hesperornis

Our knowledge of the structure of the skull in Hesperornis is based almost entirely on three specimens: (1) that found by Marsh and Russell in 1872, now in the Yale Peabody Museum (YPM 1206) (Figure 1); (2) the premaxillae and mandibles of a skull in the National Museum of Natural History, Smithsonian Institution (USNM 4978); and (3) a nearly complete but crushed skull in the collections of the University of Kansas (KU 2287) (Figures 2 and 3). The first of these skulls was described and illustrated in some detail by Marsh (1880:5-12, plates 1,2), and a brief description of the last two was published by Lucas (1903), who illustrated the quadrate and pterygoid of the Kansas specimen and the lacrimal of the National Museum specimen. The Yale and National Museum specimens are very nearly the same size and both have been identified as Hesperornis regalis by virtually all workers. The Kansas specimen, on the other hand, is slightly smaller than the other two and was placed by Lucas (1903) in a new genus, Hargeria, having as its type the species Hesperornis gracilis Marsh. After extensive comparison of the three skulls, I agree with Gregory (1952) that all three are of the same genus, Hesperornis. It remains an open question whether more than a single species should be recognized.

The Yale skull was only partly removed from the enclosing rock by Marsh, and those portions that were freed for study were subsequently remounted on the original slab for display purposes. Consequently, the specimen was not really available for examination until relatively recently, when it was removed from public exhibition.

The Yale skull is in many respects the best one for study, because its components were scattered before fossilization and are now disarticulated and very little crushed (except for the braincase). The major portion of the Yale skull is illustrated here as it was mounted for exhibition (Figure 1). The braincase and some smaller fragments were completely removed from the rock by Marsh and it is not certain that their positions as shown in Figure 1 are those in which they were found. The pre-

Figure 1.—The Yale skull of Hesperornis regalis Marsh (YPM 1206), showing the individual disarticulated bones well preserved. Premaxilla, nasal, maxilla, and vomers are illustrated in the position in which they were found—all have subsequently been removed and cleaned for study. (d = dentary, f = frontal, l = lacrimal, m = maxilla, n = nasal, pl = palatine, pm = premaxilla, q = quadrate, t = tooth, v = vomer.) Note presence of teeth in dentary, as illustrated by Marsh (1880, pl. 1). (Approximately one-half natural size.)
maxilla, maxilla, nasal, vomers, and palatine, however, were never removed and thus retain their original orientation as buried. It should be noted that Marsh had the nasal and maxilla exposed from both sides of the slab, but they were never completely removed. All of the important pieces of the Yale skull were carefully removed from their matrix in 1971 by Mr. Peter Whybrow, and they can now be studied freely and articulated.

The cranium of the University of Kansas skull of *Hesperornis* is also in a slab of Niobrara chalk, but unlike the Yale specimen, it was preserved in articulation and both the braincase and the maxillary portion of the skull have suffered considerable crushing. Furthermore, Lucas (1903) reported that the specimen was preserved with the skull doubled backwards against the pelvis, and that portions of both the dorsal and the sternal ribs were crushed into the palate. It is possible to identify most of the bones preserved in this specimen, but the maxillae are conspicuously lacking—whether they are crushed beyond recognition into the palate or lost entirely cannot be determined.

In addition to the portions illustrated in Figure 2, the Kansas specimen includes most of the lower jaws, a complete left quadrate, and a complete left pterygoid, which have been fully prepared and can be articulated with each other and also with the left palatine preserved with the main part of the cranium. The quadrate and pterygoid were illustrated by Lucas (1903, figs. 1, 2; the left pterygoid is incorrectly identified as a right pterygoid), and they are illustrated here in articulation (Figure 3). The complicated S-shaped surface of the left pterygoid (“Apl” in Figure 3) articulates with the S-shaped proximal end of the palatine (“Apt” in Figure 2).

The principal contribution of the USNM specimen to our understanding of the skull morphology of *Hesperornis* is furnished by the nearly complete left lacrimal (illustrated by Lucas, 1903, fig. 3).

By studying all three specimens it is possible to reconstruct the major features of the morphology of the rostrum, the palate, and the mandible (Figure 4). The reconstruction has been discussed elsewhere (Gingerich, 1973), but some additional notes are added here. These notes and the illustrations of the Yale and Kansas specimens (Figures 1–3) are preliminary to a more definitive description of this important material. They are intended to provide additional documentation of the remarkable completeness of the preserved specimens and to answer, in part, some questions raised by several skeptical colleagues.

The length of the reconstructed skull was determined from the Yale specimen (YPM 1206). The dorsal surface of the braincase in this specimen is crushed forward, but without affecting the length from the occipital condyle to the anterior end of the frontals. The overlapping articulation between the nasal and the frontal is outlined on the surface of the frontal, and the two can be fitted together as in life. The nasal-premaxillary articulation is preserved in both of the elements and these too can be fitted together accurately. As neither frontals, nasals, premaxillae, nor the base of the braincase appear in any way distorted in length, a total length of 26–27 cm is estimated for this skull.

Regarding the possibilities of cranial kinesis, little can be added to my previous discussion (Gingerich, 1973) except perhaps to add a more cautionary note. Rhynchokinesis in *Hesperornis* is almost certainly ruled out by the complete ring of bone formed by the premaxillae and nasals around the external narial opening. Some slight prokinetic movement might have been possible if the premaxillae and nasals were capable of being lifted off the frontals, although I know of no modern bird with such thick bone in the region of bending, and the complex interdigitation of the nasal and lacrimal in *Hesperornis* would likewise limit prokinetic movement. The quadrates were clearly streptostylic, which appears to have been correlated with a unique form of maxillolokinesis whereby the maxillae were able to slide anteroposteriorly on rails formed by the nasal-premaxillary subnarial bars (Gingerich, 1973). While I am reluctant to postulate a form of kinetic motion so distinctive from that of any other animal, the preserved osteology of the rostrum in *Hesperornis* is unique and its adaptations were clearly different from those of any known vertebrate. Maxillolokinesis appears to explain several unique features of the known fossil material.

One of the most curious features of the upper jaw of *Hesperornis* is the fact that the premaxilla bore a horny sheath as in modern birds (indicated by the vascular nature of the underlying bone), while the teeth were confined to the maxillae.
Figure 2.—The Kansas skull of *Hesperornis* (KU 2287), ventral view as preserved, articulated on a slab of Niobrara Chalk. Note particularly the little-disturbed contact between the premaxillae and nasals, while the maxillae are completely missing. (Apt = pterygoid articulation of palatine, Aq = quadrate articulation of squamosal, n = nasal, pl = palatine, pm = premaxilla, v = vomer; approximately two-thirds natural size.)
Figure 3.—Articulated left pterygoid (pt) and quadrate (q) of Kansas specimen of *Hesperornis* (KU 2287): *a*, medial view; *b*, lateral view. Note particularly the complicated articulation between quadrate and pterygoid, the broad basisphenoid articulation of the pterygoid, and the complicated S-shaped articulation of the pterygoid with the palatine. (Abs = basisphenoid articulation of pterygoid, Am = mandibular articulation of quadrate, Apl = palatine articulation of pterygoid, Aqj = quadratojugal articulation of quadrate, Asq = squamosal articulation of quadrate; twice natural size.)

The lower jaw bore teeth throughout the length of the dentary. Secondly, in both the Yale and Kansas specimens, the maxillae have clearly separated from the nasal-premaxillary subnarial bars while, at least in the Kansas specimen, the subnarial bars were little disturbed by crushing. It should be noted also that the anterior end of each maxilla was grooved to fit over anteroposteriorly aligned keys or ridges of bone on the ventral surface of the premaxilla. This system of locking would keep the anterior ends of the maxillae from dropping away from the subnarial bars, while permitting anteroposterior motion of the maxillae relative to the subnarial bars. Finally, it now seems unlikely that the left and right vomers were fused to each other at their anterior ends. Such fusion would have prevented independent motion of the left and right maxillary segments of the palate relative to each other. The only possible functional advantage of having the kind of maxillary kinesis postulated here would be in moving each side independently. As evidenced by the unfused mandibular symphysis, such independent movement of the lower jaws was clearly possible. Independent movement of the maxillae would further expand the range of possible movements used in ingesting prey, which in this case was almost certainly fish.

A new specimen of *Archaeopteryx*, described recently by Wellnhofer (1974), fortunately has a relatively well-preserved skull. Wellnhofer (1974: 185) interprets the skull as being definitely kinetic, but in *Archaeopteryx*, as in *Hesperornis*, it is difficult to see where bending that would lift a significant portion of the rostrum could have taken place. Wellnhofer favors bending in the dorsal processes of the premaxillae, but at most this would lift only the tip of the upper jaw. Kinesis approaching that of modern birds seems not to have been present in either *Archaeopteryx* or *Hesperornis*.

The present evidence bearing on Huxley's and Marsh's conclusions regarding the evolutionary position of the ratites, the relationship of *Hesperornis* to the ratites, and the presence of teeth in *Hesperornis* and *Ichthyornis* can now be considered. The skeleton of *Archaeopteryx* is more reptilian than avian, and the uncontested fact that its jaws bear teeth is easy to believe. The skeletons of *Hesperornis* and *Ichthyornis*, on the other hand, are more typically avian. That a bird with an avian postcranial skeleton should have jaws with
the quadrate is not preserved in the original specimen of *Ichthyornis* and the toothed jaws that Marsh found associated with this skeleton thus cannot be articulated with the remainder of the cranium. The articular regions of the original jaws are also badly distorted. Gregory (1952) made a careful study of the lower jaws of *Ichthyornis* and concluded that they belonged to a small mosasaur. Therefore *Hesperornis* alone was left with the combination of toothed jaws and a nearly typically avian skeleton. Inevitably, the association of teeth with the skull of *Hesperornis* was also questioned. Bock (1969) claimed that the teeth found with *Hesperornis* were not in place in the jaws, but scattered and cemented with matrix onto the skull. However, one need only examine the Yale specimen to see that teeth are preserved in the jaws as well as being scattered through the matrix (Figure 1). Discovery of a new, uncrushed posterior portion of a mandible of *Ichthyornis* (Gingerich, 1972), and its comparison with the mandibles of the original specimen and with those of *Hesperornis* and modern birds, leaves little doubt that Marsh was correct in associating toothed jaws with *Ichthyornis*.

Interpretation of the structure of the palate in *Hesperornis* has had an interesting history. Marsh (1880:6) originally determined that the palate resembled most closely that of "Struthious" birds, but he confused the vomers with the palatines of his specimen of *Hesperornis* and gave no figure or reconstruction of the palate. Thompson (1890), followed by Lucas (1903), Shufeldt (1915), and Heilmann (1926), challenged Marsh's interpretation of *Hesperornis* as indicating any relationship to the ratites. In the course of the 36 years from 1890 to 1926, the palatal structure of *Hesperornis* "evolved" rapidly in the literature, ultimately "converging" toward the neognathous palatal type of the modern loon (*Gavia*), a fish-eating, diving bird with certain similar locomotor adaptations.

Fortunately, the Yale and Kansas specimens of *Hesperornis* (Figures 1-3) preserve virtually intact at least one example of each of the palatal bones. The quadrate and pterygoid are complete in the Kansas specimen, portions of both vomers are present in the Yale specimen (Figure 1), a crushed left vomer remains in the Kansas specimen (Figure 2), and virtually complete palatines are preserved in both. About midway along their length, a rounded surface is present on the medial side of the vomers, which apparently articulated with the parasphenoid rostrum. The left maxilla is preserved in the Yale specimen (Figure 1) and it fits together with, and is overlapped by, the left vomer, as shown in Figure 4. There appears to be an articular facet on a ventrolateral expansion of the vomer for the narrow anterior end of the palate (Figure 4). It is possible, but unlikely, that the palatines articulated directly with posterior projections of the maxillae (not preserved) rather than with the vomers. As noted above, the maxillae articulated with the subnarial bars formed by the premaxillae and nasals. Returning to the pterygoid-quadrate complex, it should be noted that each pterygoid bears a large, round, flat surface that articulates with a "basipterygoid" process of the basiphenoid (Figure 3, "Abs").

The entire reassembled palate is illustrated in Figure 4c. Compared with that of living ratites, the palate of *Hesperornis* is obviously different from an emu or an ostrich in being much longer and narrower. This lengthening has clearly been accomplished by elongation of the premaxillae, maxillae, vomers, and palatines relative to the more posterior elements of the skull. Although having adaptations quite different from those of any living palaeognathous bird, *Hesperornis* shares with palaeognathous birds all essential palatal characters that distinguish them from neognathous birds: (1) a relatively large vomer, (2) a firm pterygoid-palatine connection, (3) palatines widely separated from the sphenoid rostrum by the pterygoids, (4) strong basipterygoid processes of the sphenoid articulating with the pterygoids, and (5) a complex pterygoid-quadrate articulation including portions of the orbital process of the quadrate (Figure 3).

The structure of the palate is still unknown in *Archaeopteryx*, but the presence of a palaеognathous palate in *Hesperornis* would appear to be strong evidence favoring the view that the palaеognathous conformation is primitive in birds. Additional evidence bearing on the primitive structure of the palate of birds is offered by this structure in theropod dinosaurs. Ostrom (1973) has compared the skeleton of *Archaeopteryx* with that of reptiles and concluded that birds originated from theropod dinosaurs, more specifically, from a coelurosaurian stock of theropods. The palatal
structure is not known in any coelurosaur, but it is completely preserved in the large carnivorous Tyrannosaurus (Osborn, 1912) and less well preserved in the smaller Dromaeosaurus (Colbert and Russell, 1969) and Deinonychus (Ostrom, 1969). The structure of each of these skulls appears to meet all of the criteria listed above for the palaeognathous palate. Osborn (1912:11) noted this "analogy" implicitly in comparing the palate of Tyrannosaurus with that of a cassowary. The presence of a palaeognathous palate in Mesozoic theropods, the "sister group" of birds, together with the palaeognathous palate of the Cretaceous bird Hesperornis, should leave little doubt that this palatal conformation is truly primitive in birds.

I emphasize the strength of the evidence in this case because Cracraft (1974) has proposed that the living ratite birds are cladistically a "strictly monophyletic" group on the basis of their "derived" palaeognathous palate, their unique rhamphothecal structure, and their large ilioischiatric fenestra. Cracraft asserts that the palaeognathous palate is a derived state in birds, not a primitive one, because "it is restricted to a small number of species within this large class" (Cracraft, 1974:497). This specious reasoning would lead one to assume that teeth in Mesozoic birds are a derived condition also, an unlikely hypothesis.

The unique rhamphothecal structure and other resemblances of ratites and tinamous were interpreted by Parkes and Clark (1966) rather less stringently than Cracraft now proposes. They (1966:469) noted that "resemblances are to be attributed to parallel evolution from a common stock . . . rather than to convergence from unrelated stocks, and thus, employing Simpson's concepts, the group may be considered monophyletic." The resemblance in rhamphothecal structure of
ratites and tinamous provides no evidence that this group is strictly monophyletic in Cracraft’s sense rather than monophyletic in G. G. Simpson’s sense (i.e., possibly paraphyletic, if indeed the unique rhamphothecal structure is a derived state at all—it may very well be primitive).

The third character Cracraft (1974:505) cites as evidence that ratites and tinamous are “each other’s closest relatives” is their possession of a large ilioischiatic fenestra. *Archaeopteryx* has long been known to have a large ilioischiatic fenestra (see for example Petronievics and Smith Woodward, 1917), and Cracraft (1974:503) himself notes that this is the condition in *Hesperornis* and *Ichthyornis*. In short, of the three “derived” characters cited by Cracraft (1974), the first and third are almost certainly primitive and the second may be primitive as well.

Evidence that ratites are strictly monophyletic remains to be discovered and it is possible, even probable, that the groups of living ratites and the tinamous are paraphyletic. Huxley (1867:419) envisioned the living palaeognathous ratites as “waifs and strays” of an early radiation of birds, the neognathous types representing a subsequent radiation. Judging from the fossil record, successive adaptive radiations replacing older stocks by newer ones are common in vertebrate evolution, and the class Aves is no exception. Although they are sometimes highly modified from the ancestral stock, we are fortunate to have in many groups of vertebrates surviving “waifs and strays,” and still more fortunate to have well-preserved archaic fossil forms. In the absence of a more complete fossil record, some question must remain as to whether the modern ratites and tinamous are in fact survivors of a primitive radiation of birds, or whether their primitive characteristics are neotenic solutions to particular adaptive problems, since both the palaeognathous palate and the open ilioischiatic fenestra appear to be present in the developmental stages of modern nonratite birds (Jollie, 1958; Olson, 1973:35–36). To explain away the primitive morphology of *Hesperornis* and ally it with modern loons and grebes (Cracraft, 1974:497, 503), however, illustrates on the one hand the arbitrary nature of the cladistic method of reconstructing a phylogeny, and on the other hand exemplifies another typological attempt to force an archaic bird into a modern morphological category. To paraphrase von Meyer (1861), if *Hesperornis* does not fit our philosophical wisdom and if we are not able to include this fossil in our system, our shortsightedness is alone to blame.

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The Skeleton of *Baptornis advenus*  
(Aves: Hesperornithiformes)  

Larry D. Martin and James Tate, Jr.

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**ABSTRACT**

*Baptornis advenus* is a foot-propelled diving bird from the Late Cretaceous of Kansas. It was slightly larger than the largest living loon and had an unusually long neck. The feet were large, with only slight modifications for toe-rotation. In this and many other respects, *Baptornis* was a less specialized diving bird than the contemporaneous *Hesperornis*. However, examination of almost the entire skeleton shows that *Baptornis* is more closely related to *Hesperornis* than to any living diving bird and should be included in the order Hesperornithiformes. It should not be regarded as the earliest record of the Podicipediformes. We maintain *Baptornis* in a family Baptornithidae separate from Hesperornithidae. Both *Hesperornis* and *Baptornis* are in many respects very primitive birds, which in some characters appear to be little modified from *Archaeopteryx*.

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**Introduction**

In 1964 we discovered a previously unrecognized partial skeleton of *Baptornis advenus* in the collections of the University of Nebraska State Museum. This specimen was more complete and better preserved than any other known example of the species and it encouraged us to review all known specimens of the form. In other institutions we found numerous examples of *Baptornis*. We were able to study all of these except a tarsometatarsus reported by Lambrecht (1933) to be in Germany. We thus were able to reconstruct the skeleton of *Baptornis advenus*, apart from most of the skull and jaws.

After *Hesperornis* and *Ichthyornis*, *Baptornis* is now the best known Cretaceous bird. It was first described by Marsh (1877) as a new swimming bird allied to *Hesperornis*, but readily separable from that genus in having the third and fourth trochleae of the tarsometatarsus of about equal size. He did not illustrate the type, but did give detailed measurements and commented that the bird was about the size of a loon and may have had similar habits. In 1880, Marsh illustrated the holotype tarsometatarsus and mentioned a referred femur and tibiotarsus. Lucas (1903:553-555) described a partial skeleton housed at the University of Kansas, and illustrated the coracoid, humerus, radius, ulna, and patella. Shufeldt (1915:9-11, figs. 1-6) published photographs of the holotype and discussed it in great detail. Finally, Lambrecht (1933:258-260) summarized what was then known about the anatomy of *Baptornis*.

After Lambrecht, relatively little has been published on *Baptornis*, although Swinton (1965) mentioned the genus briefly in his semipopular book *Fossil Birds*, and suggested that the Cretaceous diving bird *Lonchodytes* might be related. Storer (1958), in his discussion of evolution in diving birds, made some remarks concerning the evolutionary position of *Baptornis*, although he mistakenly spelled *Baptornis* and Baptornithidae as "Bathornis" and "Bathornithidae". Walker (1967) reviewed some of the material and previous work, but erroneously assigned an alleged large humerus to *Baptornis*. Other than the aforementioned works, the published references to *Baptornis* are restricted to checklists and catalogs
A close relationship between \textit{Baptornis} and \textit{Hesperornis} was first suggested by Marsh (1877) and went essentially unquestioned until Brodkorb (1963b) placed the former in the Podicipediformes. Brodkorb’s arrangement has been challenged by Storer (1971) and by Martin and Tate (1969) on the basis of the new material discussed in detail below. Relationships also have been suggested between \textit{Baptornis} and numerous other Mesozoic and Tertiary foot-propelled diving birds including \textit{Enaliornis}, \textit{Lonchodytes}, \textit{Hesperornis}, \textit{Neogaeornis}, and \textit{Eupterornis}. Those with \textit{Enaliornis} and \textit{Hesperornis} seem best established, as \textit{Baptornis} seems to differ from these two genera only at the familial level, whereas it seems to have no affinities with the Podicipediformes.

\textbf{Distribution.}—\textit{Baptornis} is known only from Kansas (Gove, Logan, and Wallace counties) and only from the Smoky Hill (upper) Member of the Niobrara Chalk Formation (Coniacian). Other North American Cretaceous birds (\textit{Ichthyornis} and \textit{Hesperornis}) are known from the present Gulf of Mexico to above the Arctic Circle in Canada (Russell, 1966), and range in age from the Turonian, Greenhorn Formation, to the Campanian, Pierre Shale (Walker, 1967; Martin and Tate, 1967). \textit{Baptornis} is known only from marine deposits. A specimen from Logan County (KUVP 16112) and one from Wallace County (YPM 5768) are from very immature individuals, suggesting that \textit{Baptornis} may have bred in the vicinity. Young specimens of \textit{Hesperornis} are uncommon, although Russell (1967) reported several examples of subadult \textit{Hesperornis} from a bituminous marine shale along the Anderson River in Canada (latitude 69° N) and suggested that a nesting colony might have existed nearby.

The Niobrara Chalk is a carbonate deposit with no evidence of associated continental sediments. The absence of a shoreline would imply that \textit{Pteranodon}, \textit{Ichthyornis}, \textit{Hesperornis}, and \textit{Baptornis} were accustomed to venturing many hundreds of miles into open sea. Evidence for a nearby shoreline in Kansas in the Coniacian is somewhat sketchy and not completely convincing. The Niobrara deposits do seem to thin, in an easterly direction, toward areas that are known to have been continental in the Early Cretaceous. It also has produced a few examples of dinosaurs which are presumed to have floated out to sea after death. How far these dinosaurs may have floated is presently unknown, but one wonders if it could have been hundreds of miles in such a scavenger-rich sea.

\textbf{Material Examined.}—All available specimens were studied. These included partial skeletons at the University of Nebraska State Museum, UNSM 20030; Field Museum of Natural History, FMNH 395; American Museum of Natural History, AMNH 5101; University of Kansas Museum of Natural History, KUVP 2290 and 16112; Yale Peabody Museum, YPM 1465 and 5768. We were also permitted to examine some additional material at the Fick Fossil Museum, Oakley, Kansas. Most of the available hesperornithid specimens were also examined, and the extensive skeletal collection of recent birds at the University of Kansas Museum of Natural History was used for comparisons.

\textbf{Acknowledgments.}—The authors are deeply indebted to the following persons for permission to use specimens in their care: C. B. Schulz, University of Nebraska State Museum; R. Zangerl, Field Museum of Natural History; E. Simons and J. Ostrom, Peabody Museum of Natural History, Yale University; M. Walker, Sternberg Memorial Museum; R. Shaeffer, American Museum of Natural History. We have also benefitted from conversations with P. Brodkorb, R. W. Storer, and J. Cracraft. M. A. Jenkinson read the manuscript and offered many helpful suggestions. The skeletal restoration was prepared by M. Tanner, the life restoration by B. Dalzell, and the other drawings by M. Tanner, D. K. Bennett, and D. Brennfoerder.

\textbf{Class AVES}

\textbf{Subclass ODONTOHOLCAE} Stejneger, 1885

\textbf{Order HESPERORNITHIFORMES} (Fürbringer), 1888

\textbf{Amended Diagnosis.}—Foot-propelled diving birds with teeth; skull paleognathous (at least in \textit{Hesperornis}); mandibular symphysis not fused; reduced wings; coracoid with glenoid facet on tip of scapular end; clavicles unfused; sternum flat; patella large and perforated for tendon of ambiens muscle; posterior extremities of ilium, ischium, and
pubis separate; tibiotarsus lacking supratendinal bridge; tarsometatarsus lacking hypotarsal grooves and proximal foramina.

**Family BAPTORNITHIDAE American Ornithologist’s Union, 1910**

**Included Genera.** — *Baptornis* and *Neogaeornis*.

**Amended Diagnosis.** — Foot-propelled diving birds with fully heterocoelous vertebrae; uncinate processes turned dorsally (straight in *Hesperornis*); coracoid more slender than in hesperornithids; pelvis with the preacetabular portion of the ilium relatively longer than in hesperornithids; patella pyramidal in shape (much more laterally compressed in *Hesperornis*); intracentral bones not fused to caudal vertebrae; pygostyle long and laterally compressed; outer trochlea of tarsometatarsus not enlarged; distal foramen on tarsometatarsus an open groove; toe-rotation not well developed.

**Genus *Baptornis* Marsh, 1877**

**Type-Species.** — *Baptornis advenus*.

**Included Species.** — Type-species only.

**Amended Diagnosis.** — Foot-propelled diving bird about the size of a Yellow-billed Loon (*Gavia adamsii*); neck elongate; wing greatly reduced, but radius and ulna present; tarsometatarsus not as compressed as in *Neogaeornis*.

*Baptornis advenus* Marsh, 1877

**Lectotype.** — Distal end of tarsometatarsus, YPM 1465.

**Horizon.** — Smoky Hill Member of Niobrara Chalk, Late Cretaceous.

**Type-Locality.** — Wallace County, Kansas.

**Diagnosis.** — Same as for genus.

**Morphology**

**Skull.** — The skull of *Baptornis* is known from only a few fragments. If teeth were present they were restricted to the maxillae and dentaries, as in *Hesperornis*. KUVP 16112 includes a fragment of the bill just anterior to the nasal openings. There are fairly large, distinct, triangular grooves on the lateral sides of the fragment, extending towards the tip. The sides of the bill are quite thick and vaulted. There is no evidence that the bill came to a point to form a spear as in the grebe *Aechmophorus*. A fragment of the left side of the bill (AMNH 5101) is probably from a position just posterior to that of KUVP 16112. This fragment shows a single large, elongate, longitudinal nutrient foramen. In KUVP 16112 a large number of smaller foramina extend over the dorsal surface of the bill as in the top of the bill in loons. Apparently the tip of the bill was relatively short and broad, being shaped like that of *Hesperornis* (Marsh, 1880, pl. 1). It was probably covered by a horny sheath.

AMNH 5101 includes a fragment of the frontal bone which is very difficult to orient (Figure 1a). The dorsal surface bears a faint scroll-like pattern similar to that on the frontal bone of grebes. However, the frontal-parietal suture is present and the sagittal groove extends up to it, thus extending further posteriorly than in the grebes and indicating that the top of the skull was more similar to that of loons. The cerebral hemispheres of the brain seem to have been expanded, as in modern birds, but better material is needed to confirm this.

The ventral half of a right quadrate (Figure 1c–e) is preserved with AMNH 5101. The shaft is broad but not as massive as in *Gavia*. The orbital process has been broken off, but it had a triangular base and originated very low on the quadrate. A prominent pit is present on the dorsomedial margin of the base of the orbital foramen, but the quadrate was not pneumatic. There is a small, rectangular facet for the articulation of the pterygoid. This facet is elongated dorsoventrally. Lateral to the pterygoid articulation is a shallow, rectangular depression from the middle of which a low ridge connects to the socket for the quadratojugal. The socket for the quadratojugal is large, shallow, and somewhat triangular in shape (rounded and very deep in loons and grebes). In loons and grebes, the mandibular articulation is divided into two parts by a groove, and the medial facet runs anteroposteriorly. *Baptornis* resembles *Hesperornis* in having a single comma-shaped facet running from the pterygoid articulation.

The few fragments preserved are suggestive of a skull close to that of *Hesperornis*. *Hesperornis* and *Baptornis* have very similar quadrates, and both differ from loons and grebes in most details of that bone.
Mandibular Ramus.—FMNH 395 includes the posterior portion of the left mandibular ramus with the articular cotyla (Figure 1b). It is slightly crushed dorsoventrally. The only other Cretaceous birds for which any part of the jaw is known are Hesperornis and Ichthyornis (excluding Caenagnathus, which is not a bird).

In Baptornis, the surangular is tightly fused to the articular, with a groove near the ventrolateral border of the specimen along the suture between
these two bones. The articular cotyla is divided into anterolateral and posteromedial sections, which are separated by a low ridge. A groove, which includes the articular foramen, lies on the margin of the posterior cotyla and runs under the internal articular process. Anterior to the posterior articular surface is a small depression and a wide, deep groove, which forms at the junction of the surangular and the articular.

The articular surfaces are of about the same shape and occupy approximately the same position as those on the ramus of Hesperornis (Gregory, 1952, fig. 7). They differ from the articular cotylae of the mosasaurs Clidastes and Platycarpus in having a concave, oblique, and elongate posterior articular surface and in the fusion of the jaw elements. Like Hesperornis, Baptornis lacks a depression for the condyle of the quadrate. This depression is present on the jaws of modern diving birds. In the Double-crested Cormorant (Phalacrocorax auritus), for example, it is centrally located and occupies nearly the entire posterior articulation. A posterior articular foramen is similarly located in Hesperornis and modern species of birds, but is absent in the mosasaurs. The posterior portion of the ramus of Baptornis is remarkably similar to that illustrated for Ichthyornis (Gingerich, 1973; fig. 2).

The mandibular articulation of Baptornis is very similar to that of both Hesperornis and Ichthyornis (Figure 2.) The jaws of Cretaceous toothed birds are more similar to one another than is generally supposed. Hesperornis shares with Ichthyornis the following features: teeth flattened from side to side, with expanded bases; anterior end of dentary blunt and symphysis not fused; teeth restricted to dentaries and maxillae; quadrate articulation double. Gingerich (1972:472) lists several additional features shared by Ichthyornis and Hesperornis in the posterior portion of the ramus. In so far as we can tell, Baptornis also shares these characters, and although it cannot yet be proven, we suspect that it was toothed.

Presacral Vertebrae (Table 1).—AMNH 5101 contains the most complete vertebral series of Baptornis, with at least 22 presacral vertebrae present. UNSM 20030 includes the first eight vertebrae anterior to the sacrum, and at least one of these is absent from AMNH 5101 suggesting that the total number of presacral vertebrae in Baptornis may have been 23, the same as in Hesperornis. Most of the vertebrae with AMNH 5101 are fragmentary, however, which may cause errors in interpreting their number and position. It is possible that Baptornis may have had 24 or 25 vertebrae, but the number could not have been less than 23. There are four thoracic and parts of six cervical vertebrae with KUVP 16112. KUVP 2290 also contains four thoracic and six cervical vertebrae. FMNH 395 includes the last unfused thoracic vertebra. Three thoracic and two cervical vertebrae in AMNH 5101 and three cervicals with KUVP 2290 are preserved in articulation. These are the only natural associations available. The relative positions of the other vertebrae have been determined by comparisons with Hesperornis.

Only the cervicals reported on by Lucas (1903: 553) have previously been described. He pointed
out that these were somewhat more elongate than the comparable vertebrae in *Hesperornis*.

The atlas (lacking the dorsal arch) and the crushed anterior portion of the axis (Figure 3a) are present in AMNH 5101. The atlas is similar to the atlas of *Hesperornis* illustrated by Marsh (1880:196, fig. 40). In *Hesperornis*, loons, and grebes, however, the ventral border of the atlas is directed posteriorly to form a shelf. In *Baptornis* the ventral border of the centrum is directed anteriorly and does not form a shelf. The hypapophysis is short and broad so that it is not well separated from the rest of the centrum. On either side of the hypapophysis is a short posteriorly directed process. These are very small in *Hesperornis* and absent in the loons and grebes. Short, blunt, transverse processes are also present. The odontoid process of the axis is short and broad. The axis has a large, deep, anteriorly directed pit on its anterolateral side and a pair of small pits on each side of the midline on the anteroventral surface. Its centrum is very narrow, with a thin blade-like hypapophysis. The anterior cervicals are all narrow and elongate (Figure 3c-e). They become much shorter and more massive posteriorly. The first eight vertebrae after the axis seem modified for downward flexion (Figure 3) and the articular facets of the prezygapophyses curve ventrally. The anapophyses are short and somewhat dorsally directed. The sublateral crests form a triangle with the anteroventral margin of the centrum on most of these vertebrae. In *Hesperornis*, the sublateral crests on the anterior cervicals are nearly parallel and the centra are much broader ventrally. What appears to be the tenth vertebra is very narrow posterovertrally, with its posteroventral border directed downward. Articulating posteriorly with this vertebra is a short, broad eleventh cervical with the zygapophyses tilted laterally and not directed downwards. The ventral surfaces of this and the four following vertebrae are broad and short and bear low parallel sublateral processes that do not join each other ventrally as they do in similar vertebrae in grebes (Zusi and Storer, 1969, fig. 10). Vertebrae 14–16 bear very large, deep, lateral depressions in their centra. The ventral border of the centrum of the 16th vertebra is directed ventrally and forms a distinct, thick, ventral process. In *Baptornis* there is no evidence that large vertebralarterial canals were present in as many vertebrae as they are in *Hesperornis*, loons, and grebes; however, the vertebrae which should show these canals are all badly damaged in the material of *Baptornis*.

The eighth vertebra anterior to the synsacrum (Figure 4a–d) corresponds to the 16th cervical vertebra of *Hesperornis*. Its dorsal spine is short, thick, and broad at its base. The tubercle is not bifurcated; the post- and prezygapophyses are set at a low angle and are widely spaced; the neural canal is circular in outline; the anterior articulation of the centrum is short dorsoventrally, but very wide; the transverse processes are wide, swing slightly upwards, and are directed posteriorly; the posterior border is thickened while the anterior portion nearest the centrum is slightly concave; and there is no vertebralarterial canal. On this ver-

| Table 1.—Measurements (mm) of the thoracic vertebrae of *Baptornis* numbered from the synsacrum forward |
| --- | --- | --- | --- | --- | --- | --- |
| Character | **FIRST** | **SECOND** | **THIRD** | **FOURTH** | **FIFTH** | **SIXTH** |
| **UNSM** | **FMNH** | **UNSM** | **KUVP** | **UNSM** | **KUVP** | **UNSM** | **UNSM** |
| Length centrum | 19.5 | 18.7 | 22 | 22 | (22) | 21.5 | 21 | 21.5 | (20) | 21 |
| Width anterior articulation | 18.3 | 17.6 | – | 18 | (14.5) | 16 | – | 15 | 13 | 14 | 13.5 |
| Height anterior articulation | 9 | 8.8 | 9 | 8.5 | 9 | 8.5 | 8.7 | 9 | 7.7 | 9 | 10 |
| Width posterior articulation | 14 | 12.8 | 15 | – | 13 | 15.5 | 12 | 14 | 10.5 | – | 11 |
| Height posterior articulation | 9.5 | 8.5 | 9.5 | 10.5 | 9.5 | 9 | 9.5 | 9 | – | 9 |
| Diameter of rib articulations | * | * | – | – | 4.7 | – | 4.7 | – | 5.5 | 5 | 4.5 |
| Head | – | – | – | – | – | – | – | – | 11 | – | 8.5 |
| Tuberculum | – | – | – | – | – | – | – | – | – | – | 3.5 |
| Height neural spine | 14 | – | – | – | – | – | – | 11 | – | 8.5 |
| Length base neural spine | 12 | 12 | – | 12 | – | 13 | – | 13 | – | 13 |
| Width across transverse process | 23 | 23.5 | – | – | – | – | – | – | – | 36 |

* Ribs do not occur on first vertebra; ( ) = measurements from crushed specimen.
Figure 3.—Stereophotographs of vertebrae of Baptoptis advenus: a, atlas and axis vertebrae (AMNH 5101), X 4; b, cervical vertebra (AMNH 5101), anterior view, X 1; c, same, ventral view, X 1; d, three partial cervical vertebrae (KUVP 2290), dorsal view, X 1; e, same, lateral view, X 1; f, thoracic vertebra (KUVP 2290), posterior view, X 1.

tebra the articulation for the tuberculum of the rib is shallow, circular in outline, and slightly recessed from the outermost margin of the diapophyses. The articulation for the head of the rib is deep, circular, and placed anteriorly on the centrum. The centrum itself is short and thick, with a strong indentation on either side. The hypapophysis is extremely short, stout, and directed posteriorly, practically merging in with the rest of the centrum. The distal hypapophysis is terminated by two stubby horizontal wings.

The seventh vertebra anterior to the sacrum corresponds to the last cervical of Hesperornis and resembles the previous vertebra. However, the anterior articulation is not as wide and is higher dorsoventrally; the neural canal is not as rounded; the prezygapophyses are closer together and at a more acute angle; the dorsal spine is wider and thicker; there is a concavity on the anteroventral margin of the transverse process above the articulation for the head of the rib; and the transverse processes are shorter and do not swing back pos-
The articulation for the tuberculum of the rib is nearer the lateral edge of the diapophysis than in the 16th vertebra. The hypapophysis is thin and posteriorly directed.

The sixth vertebra anterior to the sacrum resembles the seventh except that the anterior articulation is not quite as wide and is higher dorsoventrally; the dorsal spine is of about the same height but is thinner; the transverse processes curve posteriorly; the pit above the articulation for the rib is deeper, and the hypapophysis is thin and directed anteriorly.

The fifth vertebra anterior to the sacrum has a narrow, higher anterior articulation than the sixth; the centrum itself is not as indented and the prezygapophyses are closer together and at a more acute angle; the dorsal spine is thin, high, and directed anteriorly; the pit above the articulation for the head of the rib is deep, as in the preceding vertebrae; the diapophysis and hypapophysis are
missing, but enough remains of the hypapophysis to show that it was thin and large.

The fourth vertebra anterior to the sacrum similarly has a high, narrow anterior articulation; however, the centrum is less deeply indented; a small pit forms on the dorsal anterior margin of both of the postzygapophyses; the pit above the articulation for the head of the rib has become shallow and elongate. The dorsal spine, transverse processes, right prezygapophysis, and most of the hypapophysis are broken off, but enough remains to show that the hypapophysis was thin, posteriorly directed, and probably short.

The third vertebra anterior to the sacrum has a lower, but wider, anterior articulation; the centrum is less indented; the pit above the articulation for the head of the rib is absent; and the hypapophysis is very short, thin, and triangular. The dorsal spine and diapophysis are missing from the available specimens.

The second vertebra anterior to the sacrum has a wider anterior articulation than the preceding vertebra. The top of the vertebra and the transverse processes are missing in the one available specimen. The articulation for the head of the rib is shallower than in the preceding vertebra, and there is a shallow pit just above it. The hypapophysis is represented by a small knob on the ventral border of the vertebra. The centrum of this specimen had been broken through the middle and repaired with glue. We separated the two halves with acetone but found no medullary cavity.

The first vertebra anterior to the synsacrum (Figure 4e–h) is distinctive. The anterior articulation is wider than in the preceding vertebra. The dorsal spine is high and thin, and the posterior margin is bordered by two small grooves lying on top of the postzygapophyses. The diapophyses are short, thick, and directed posteriorly, their ends being flattened to buttress against the inner sides of the ilia. There is a small pit lying directly anterior to the diapophysis. The centrum is slightly indented, the ventral border is flat, and there are no rib articulations.

The vertebrae of Baptornis are nonpneumatic and heterocoelous (amphicoelous vertebrae occur in Archaeopteryx, Ichthyornis, and Enaliornis). Compared to Hesperornis, the cervical vertebrae are more elongate and not as deep, the anapophysies are less developed, and the sublateral processes tend to converge more posteriorly. At least the first five postaxial vertebrae are modified for downward flexion, as are the presumed 14th and 15th vertebrae. The specimens of intervening vertebrae (7–13) are too fragmentary to be certain of their adaptations but some must have been modified for upward flexion.

The description of the thoracic vertebrae is based almost entirely on the beautifully preserved series with UNSM 20080. These all have good heterocoelous articular surfaces, although circular pits (Figure 3f) occur in the centra of some specimens. They are not fused as in cormorants, or fused and further immobilized by ossified dorsal tendons, as in grebes. They also lack the very deep lateral excavations found in the thoracic vertebrae of Ichthyornis and can best be compared to the thoracic vertebrae of Hesperornis. As Lucas (1903) indicated, the hypapophyses are more anteriorly situated, not as well developed as in Hesperornis, and not bifurcated as in most modern diving birds.

**Synsacrum.—** The synsacrum (Figure 5b–d) is represented in KUVP 2290, FMNH 395, AMNH 5101, and a few fragments from KUVP 16112. It is nonpneumatic and extremely narrow. Lucas (1903) reported that the synsacrum of KUVP 2290 contains 10 fused vertebrae and that the first bore a rib. Although he was correct in the number of vertebrae (10 also being the number found in FMNH 395) he was mistaken about the presence of a rib with the first fused sacral. Rib facets do not occur on any of the synsacra available nor even on the last unfused thoracic vertebra.

The anterior sacral vertebra has a high, posteriorly sloping neural spine. The transverse processes are tilted and flattened laterally to form a broad contact with the ilium. The second sacral is sutured dorsally to the first. It has a large, laterally flattened, pointed transverse process projecting anteriorly (Figure 5b) that also abuts the ilium. There are three vertebrae from the acetabulum forward, and these may represent fused lumbar with the rest of the sacrum being composed of seven fused caudals (urosacrals). The neural spines of the synsacral vertebrae are low and form a median ridge bounded by projections on either side. The posterior urosacrals lack the ventral keel found in Gavia, and the posterior central articulations of the last two urosacral vertebrae are divided.
Caudal Vertebrae.—UNSM 20030 has five caudal vertebrae and a pygostyle, as does FMNH 395. AMNH 5101 also contains five caudals but includes four anterior ones that are not represented in either UNSM 20030 or FMNH 395. The first caudal has already been discussed in the section on the sacrum. The following three vertebrae have low neural spines, flat ventral borders and widen posteriorly, assuming a triangular shape. All of these vertebrae are flanked by the pelvis and their transverse processes abut the ilia. The vertebrae posterior to the pelvis have high, straight, triangular neural spines with rounded tips that become

Figure 5.—Pelvis of Baptornis advenus: a, right innominate bone (UNSM 20030), lateral view. Synsacra of B. advenus: b, dorsal view (KUVP 2290); c, lateral view (FMNH 395); d, ventral view (AMNH 5101). All × 1.
progressively lower posteriorly. The caudals also have large, posteriorly directed and ventrally depressed transverse processes that, along with the centra, become smaller posteriorly. The vertebral centra are either amphicoelous or amphiplatyan and the centra themselves are rounded ventrally and bear ventrolateral depressions or pits. The pygostyle is elongate and laterally flattened. It includes five fused centra. The general configuration of the posterior two-thirds of the tail (Figure 6) is roughly similar to that in Gavia and quite different from Hesperornis, which has dorsoventrally flattened caudals with wide, flat transverse processes and shelf-like, fused intracentral bones. Hesperornis has only two fused centra in its pygostyle. Intracentral bones are present in many living diving birds, but they have not been found in Baptornis. In Baptornis, the posteroventral margin of the second fused centrum of the pygostyle bears a distinct projection, which probably served as a muscle insertion.

RIBS AND UNCINATE PROCESSES.—Some fragments of ribs are present with all of the specimens of Baptornis. The best material (Figure 7) is preserved in situ on slabs of matrix with UNSM 20030. This material includes at least four different dorsal ribs and four different sternal ribs. These are flattened and shaped about as in Hesperornis, suggesting a narrow rib cage. Six uncinate processes representing five pairs are present in UNSM 20030. The uncinate processes of Hesperornis and Baptornis do not fuse to the ribs—a condition similar to that seen in grebes and loons, as well as certain other modern birds. In Hesperornis, the uncinate...
processes are broad and straight. In *Baptornis*, they are narrow, and bend upward at an acute angle, thus resembling the uncinate processes of grebes (Figure 8). Because there is no association between ribs and uncinate processes on the slab, none can be placed in order with certainty. The ribs of *Baptornis* are heavier than those of grebes and loons and do not expand as much ventrally. The tuberculum is not separated as far from the head as it is in *Hesperornis* or in the modern foot-propelled divers.

**Sternum.**—The anterior and lateral margins of the sternum (Figure 9a) are preserved with UNSM 20030, the entire central portion and the posterior margin having been destroyed.

In the area of the dorsal manubrial spine there is only a smooth, thickened border from which a shallow sulcus flares ventrally into a flattened, rounded ventral manubrial spine. The coracoidal sulci are deep, closely spaced, and lie at an angle of 22° from a line perpendicular to the midline of the sternum. The sternocoracoidal processes are large, rounded, flare laterally, and are not strongly curved. The costal margins are short, and bear five costal ridges.

The sternum is most similar to that found in *Hesperornis*. It appears to bear a shallow rectangular depression on its anteroventral surface which does not occur in *Hesperornis*. The sternum also appears relatively smaller than in *Hesperornis* and consequently the body of *Baptornis* may have been somewhat narrower. The width of the anterior end of the sternum is 53 mm and the length of the costal margin is 27 mm.

**Coracoid** (Table 2).—UNSM 20030 includes the scapular end and a fragment of the sternal end of the left coracoid. KUVP 2290 includes most of the right coracoid (Figure 9c). This specimen was illustrated by Lucas (1903, fig. 6).

The head of the coracoid is small and turned towards the procoracoid. The glenoid facet is large, elliptical, and shallow. The furcular facet is low, narrow, not undercut, and set almost directly on the scapular end of the shaft so that it is completely visible when the scapular end of the coracoid is viewed from above (Figure 10a, 11a). The surface of the scapular facet is rough and bears two or more distinct pits on its anterior end. The procoracoid is short and recurved towards the shaft to form part of the triosseal canal. Just below the procoracoid is a foramen leading into the shaft of the bone, which is probably the procoracoid foramen. In KUVP 2290 this foramen penetrates from the anterior to the posterior surface, as well as branching into the shaft. Lucas (1903:553) incorrectly states that the procoracoid process and foramen are absent. The shaft is long and narrow. The posterior surface is slightly concave, while the anterior surface is curved and convex. This gives the coracoid the appearance of a shallow spoon with a square end. The sternal end is wide, thin, and lacks definite facets.

The coracoid of *Hesperornis* is fundamentally

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**Figure 8.**—Ribs and uncinate processes: a, grebe, *Aechmophorus occidentalis*; b, *Baptornis advenus*; c, *Hesperornis regalis*. (Not to scale.)

**Figure 9.**—Pectoral and wing elements of *Baptornis advenus*: a (top to bottom), anterior, ventral, and left lateral views of sternum (UNSM 20030) × 1; b, external and internal views of left humerus, X 1; c, ventral, dorsal, and external views of right coracoid (KUVP 2290), X 1; d, distal end of right humerus, external view, X 5.

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**Table 2.**—Measurements (mm) of the coracoid of *Baptornis*

<table>
<thead>
<tr>
<th>Character</th>
<th>UNSM 20030</th>
<th>KUVP 2290</th>
</tr>
</thead>
<tbody>
<tr>
<td>Approximate length</td>
<td>– 53</td>
<td></td>
</tr>
<tr>
<td>Width glenoid facet</td>
<td>4</td>
<td>3.5</td>
</tr>
<tr>
<td>Length glenoid facet</td>
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<tr>
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<tr>
<td>Length scapular facet</td>
<td>7</td>
<td>7.5</td>
</tr>
<tr>
<td>Width sternal end</td>
<td>– 23</td>
<td></td>
</tr>
</tbody>
</table>
similar to that of *Archeopteryx*. In both genera the coracoid is short and broad and the scapula and humerus have articulations on the tip of the scapular end. The coracoid of *Baptornis* resembles *Hesperornis* in these features, as well as in having the sternocoracoidal process above the midpoint of the bone. The main longitudinal axis of the shaft of the coracoid is perpendicular to the edge of the sternal facet in the Hesperornithiformes and inclined in all other foot-propelled diving birds for which the coracoid is known (Figure 10). The coracoid of *Baptornis* is more elongate than in *Hesperornis* and the procoracoid process is smaller and differently shaped. In *Baptornis*, the internal edge of the coracoid is nearly straight, while in *Hesperornis* there is a large, square, internal projection (Figure 10c). It should be noted that Lucas (1903, fig. 6) has illustrated the posterior view of the sternal fragment of the coracoid with the anterior view of the scapular portion (compare his figure 6 with figure 10 of this paper).

A small area for the attachment of the clavicle seems to be present although no clavicles are known.

**Scapula.**—KUVP 2290 includes the articular end of the left scapula (Figure 11b–d), which was described and figured by Lucas (1903:553–554). The coracoidal articulation is long, narrow, and slightly curved to fit the contour of the scapular facet on the coracoid. It bears two large pits similar to those found on that facet. The anterodorsal margin of the proximal end does not show any articulation for the furcula. This margin slopes at about 47 degrees to the main axis of the shaft, and is terminated dorsally by a small projection. The ventral border bears a long, shallow groove. The measurements (in mm) of the scapula are: width of neck, 7.5; depth of neck, 4.0; width of proximal end, 10.5; length of glenoid facet, 9.4.

The neck of the scapula is wide and thick, which led Lucas to suggest that it may have been expanded posteriorly as in penguins. The scapula of *Hesperornis* is similarly thickened, but does not expand posteriorly (Marsh, 1880:58). Therefore, the posterior portion of the scapula of *Baptornis* probably did not differ much from that of *Hesperornis*.

**Humerus.**—KUVP 2290 includes the distal end and a portion of the shaft of the right humerus. This specimen was described by Lucas (1903:554), who reported it as being a left humerus, but the curvature of the shaft matches that of Marsh's illustration of the right humerus of *Hesperornis regalis* (Marsh, 1880, pl. 8: fig. 1). UNSM 20030 includes a slightly abraded proximal end and the greater portion of the shaft of the left humerus. We have examined the large alleged humeri of *Baptornis* reported by Walker (1967) and have determined that they are fragmentary shafts of the tibiotarsus.

The proximal end of the humerus is simplified as compared with that of modern birds (Figure 9b). The shaft curves downward, then expands noticeably and is twisted postero-medially 75 mm from its proximal end. There is a nutrient foramen situated in a groove 48.5 mm from the proximal end in UNSM 20030. This foramen is likewise present in KUVP 2290, which also has a
second groove and a foramen 17.5 mm distal to the previously mentioned one.

A small, rounded prominence set off from the internal condyle by a shallow groove may represent the distal external condyle. The internal condyle is not of the form usually found in birds and is only slightly delineated from the rest of the distal end. It bears a small foramen on its articular surface. There is no distinct olecranal fossa or prominent grooving. The shaft of the University of Nebraska
specimen was x-rayed and the broken ends of KUVP 2290 and UNSM 20030 were examined. Neither are pneumatic. By transposing the humerus of UNSM 20030 over that of KUVP 2290 so that the curvatures and foramina are in alignment (Figure 9b), a reconstructed length of 118 mm is obtained. This is only 18 mm more than Lucas's (1903:553) estimate of 100 mm based on KUVP 2290 alone. The width of the proximal end is 8.9 mm and of the distal end 6.8 mm. The greatest diameter of the shaft is 6.5 mm.

The humerus of Baptornis is relatively long, curved, and rounded. It is not flattened and straight as in the wing-propelled auks and penguins. Lucas (1903:554) suggests that the wing may have been used in conjunction with the feet for locomotion. This seems unlikely, although we have shown it as having a stabilizing function (Figure 20). The humerus is larger than that of Hesperornis, which is a much bigger bird. In Hesperornis, the distal end is also much more reduced and the radius and ulna may not have been present.

Radius.—A description of the left radius (Figure 12b) KUVP 2290, appears in Lucas (1903:554). The humeral cotyla is large and oval, with the bicipital tuberosity situated along its rim. The shaft expands markedly at about the midpoint. In this area there is a nutrient foramen on the palmar side and also a faint intermuscular line. The lunar depression is shallow. The scapholunar facet is long and narrow and the distal ligamental process is relatively large. The radius is 20.5 mm long, the proximal end is 3.0 mm wide, and the distal end is 3.5 mm wide.

Ulna.—Lucas (1903:554) briefly describes the short, robust, left ulna, KUVP 2290 (Figure 12a). The olecranon process is short, massive, and not noticeably twisted as in many flying birds. The internal and external cotylae on the proximal end are only slightly separated by an intercotyla area and almost form a single articular surface. The proximal radial depression is slightly discernable. The impression of the brachialis anticus is large and oblong, and crosses the palmar surface of the bone diagonally. A faint intermuscular line stretches for 2–3 mm below the distal end of the impression for the brachialis anticus. No nutrient foramen can be discerned. The distal radial depression is small and quite shallow. The internal condyle on the distal end is a small projection. The external condyle is not a distinct ridge but bears a large, flat articular surface which tilts toward the anconal and ventral margins. Although we examined the bone carefully, we could see no scars for feather attachment, Lucas' (1903:554) assertion of their presence notwithstanding. Measurements (in mm) of the ulna are: length, 21.6; width proximal end, 4.2; width distal end, 2.9; depth distal end, 3.5.

The radius and ulna of Baptornis are short, stout bones resembling in general form their counterparts in the extinct diving goose Chendytes milleri (Howard, 1955). The reduction is much greater in Baptornis, however. In C. milleri the ulna is 43 percent as long as the humerus (Howard, 1955:142), while in Baptornis it is only about 19 percent of the restored length (118 mm) of the humerus. The general form of the ulna is similar to that found in theropod dinosaurs (Figure 13) and it seems possible that the Hesperornithiformes may have split off from the line leading to modern flying birds before the wing had developed the adaptations seen in modern birds. The presence of well-formed articulations on the distal ends of the radius and ulna show that some sort of carpal was present, although these may not have been fused into a true carpometacarpus.

Pelvis.—All the specimens of Baptornis we studied included at least some fragments of the pelvis. Most of the right side is preserved with UNSM 20030, and our description of the pelvis is based on this specimen (Figure 5a). The synsacrum is in place only in AMNH 5101, in which the acetabulum is opposite the third fused sacral.

The pelvis is similar to that of Hesperornis, but differs in having a much longer preacetabular portion of the ilium. The postacetabular part of the ilium is quite long, as it is in all of the foot-propelled diving birds. The middorsal and the posteriormost portions of the ilium are not known, but were probably similar in shape to the same region in Hesperornis, although the whole pelvis is somewhat narrower proportionally. The pectoral process is broad and blunt, as it is in Hesperornis, and the acetabulum is partially closed. The acetabula of the loons and grebes have vertical sides and are completely open. The antitrochanter of Baptornis is large and rectangular, resembling that of Hesperornis. As in Hesperornis, the ante-
rior end of the ischium sweeps up to form the posteroverentral border of the antitrochanter, and there is a prominent suture here in both *Baptornis* and *Hesperornis*. On the pelvis of the Common Loon, *Gavia immer*, just anterior to the antitrochanter, there is a small scar for the gluteus medius and minimus muscles. This scar is absent in the pelvis of grebes, *Baptornis*, and *Hesperornis*. The ilium and ischium of *Baptornis* are separate throughout their length, as they are in *Hesperor-
nis. The ischium is long, thin, flattened internally, and rounded externally, so that in cross-section it appears bell-shaped. The pubis is long, heavy, and flattened dorsoventrally. It bears a shallow groove along the anterior quarter of its dorsal surface.

The measurements (in mm) of the pelvis are: total length (estimated), 179; length of preacetabular ilium, 54; depth of ilium anterior to pectinal process, 19; depth of ilium at pectinal process 27; greatest diameter of acetabulum, 14; height of antitrochanter, 13; width of antitrochanter, 11; length of free ischium, 103; length of free pubis, 109.

Loons and grebes resemble each other and differ from Baptornis and Hesperornis in having the preacetabular portion of the pelvis narrower, twisted laterally, and spread apart anteriorly. Baptornis also differs from the modern foot-propelled diving birds in having the ventral margin of the ilium turned medially (Figure 5). The origins of the iliotrochantericus medius and antitrochantericus muscles are ventral in Baptornis, but are dorsal in loons and grebes. The dorsal surface of the preacetabular ilium is turned medially in loons and grebes and laterally in Baptornis. The postacetabular portion of the pelvis of Baptornis is, on a whole, narrower than in other diving birds except Hesperornis.

Femur (Table 3).—Both femora are represented in FMNH 395 and KUVP 2290. UNSM 20030 has the right femur present (Figure 14b) and KUVP 16112 includes fragments of both femora.

The femur of Baptornis is proportionately more elongate than that of Hesperornis; the neck is more constricted; the insertion of the round ligament is smaller; and the trochanter rises slightly above the head, whereas both are of about the same height in Hesperornis. When the femur is articulated with the pelvis its position is more inclined than in Hesperornis, but the difference is probably not as great as Lucas (1903:554) suggested. The iliac facet occupies about the same shape and area as it does in Hesperornis, and the obturator ridge is very similar in form. Baptornis resembles loons in having the lateral margin of the trochanter close to and parallel with the axis of the shaft, whereas the lateral margin of the trochanter extends a considerable distance away from the axis of the shaft in Hesperornis. The anterior intermuscular line sweeps down in a low arc from the trochanteric ridge to the external condyle. The posterior intermuscular line is not well defined, but runs down the medial surface of the bone from just below the head to the internal condyle. The trochanteric ridge is proportionately larger and heavier than it is in Hesperornis. There are no large nutrient foramina evident on the shaft. The popliteal area is broad and shallow. The fibular condyle sends off a distinct wing in Baptornis and Hesperornis and there is a thick connection between the internal and external condyles (this connection is thick in cormorants and grebes and thin in loons). The fibular groove is broad and shallow in Baptornis and Hesperornis. The internal condyles are at about the same level in Hesperornis.

<table>
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<tbody>
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<td>Length</td>
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<td>72</td>
<td>75</td>
</tr>
<tr>
<td>Diameter head</td>
<td>10</td>
<td>10</td>
<td>11.5</td>
</tr>
<tr>
<td>Diameter distal articulation</td>
<td>25</td>
<td>24</td>
<td>26</td>
</tr>
<tr>
<td>Diameter proximal end</td>
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<td>24.5</td>
<td>28</td>
</tr>
<tr>
<td>Antero-posterior diameter midshaft</td>
<td>11.5</td>
<td>12</td>
<td>12.5</td>
</tr>
<tr>
<td>Transverse diameter midshaft</td>
<td>10</td>
<td>10.5</td>
<td>11</td>
</tr>
</tbody>
</table>
Figure 14.—Hindlimb elements of Baptornis advenus: a, distal end of left tibiotarsus (UNSM 20030); b (top to bottom and left to right), proximal, distal, anterior, lateral, posterior, and medial views of right femur (UNSM 20030); c (top to bottom and left to right), proximal, posterior, and anterior views of right fibula (UNSM 20030); d (top to bottom), anterior, posterior, and distal views of right patella (UNSM 20030). (All × 1.)
perornis while the external condyle is much more distal than the internal condyle in Baptornis.

A discussion of the muscle scars on the femur would be valuable but the homologies are difficult to ascertain. Those of the following scars seem fairly certain. Along the posterior intermuscular line, just below the head, is a small raised triangular area which may correspond to the insertion of the M. iliacus. On the lateral surface of the femur, along the trochanteric ridge, is a complex region of muscle attachments resembling the same area in Hesperornis. The tubercle for M. piriformis is not as prominent as it is in Hesperornis and is situated about half way up the shaft.

The proportions of the femur of Baptornis suggest that the legs were not bound so closely to the pelvis as in Hesperornis and may have had slightly greater freedom of movement. In Hesperornis, the femora may have been permanently held in the extended position illustrated by Heilmann (1927, fig. 34), while Baptornis may have been able to rotate the legs ventrally for paddling and then abduct them for diving as do some diving ducks (Raikow, 1970:6).

Patella (Table 4).—The patella of KUVP 2290 was described and illustrated by Lucas (1903). There is also an excellently preserved patella with UNSM 20030 (Figure 14d). This is a short trihedral bone resembling in some respects the patella of a cormorant. The foramen for the tendon of the ambiens muscle is large and perforate. The articular surface on the base of the bone is double, indicating that it probably articulated with both the internal and external condyles of the femur. In Hesperornis there is a single concave surface, which articulated with the external condyle of the femur. Therefore, the patella of Hesperornis would have been lateral to the main axis of the tibiotalus, while that of Baptornis would almost have been centered on it (contrary to Lucas, 1903:554). Cormorants have the patella placed as in Baptornis. The patella in grebes is shaped as in Hesperornis and articulates on the external condyle of the femur. However, grebes lack the ambiens muscle and therefore no foramen is present in the patella.

Tibiotalus (Table 5).—Both right and left tibiotali occur in UNSM 20030 and in FMNH 395 and parts of both are present in KUVP 2290 and 16112.

The tibiotalus (Figure 15) is like that of Hesperornis in being elongate and nonpneumatic (like Hesperornis, it has a large medullary cavity). The proximal end flares out as in Hesperornis due to the lateral expansion of the outer cnemial crest. This region is not as expanded in loons and grebes. As in Hesperornis, the inner cnemial crest is low, so that the groove between the two crests is broad and shallow. In loons and grebes, the inner cnemial crest is high and the surface between the two crests is narrow and deeply excavated. The rotular process is lower than in loons or grebes and is similar to Hesperornis. The external articular surface is small and slopes ventrally. It is not as rounded as in loons or grebes, nor is it set apart anteriorly and posteriorly from the inner articular facet by grooves (Figure 16a) as in Hesperornis. The inner and outer articular facets are about equal in size and are separated by a groove in the interarticular area in grebes. The inner articular facet is flat, oval, and directed posteromedially in Baptornis, and just below the inner articular facet is a deep roughened pit for the origin of M. plantaris which appears to be divided into dorsal and ventral parts. The fibular crest extends about half way down the shaft and is deeply grooved along its

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**Table 4.** Measurements (mm) of the patella of Baptornis

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<tr>
<td>Length</td>
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<td>Distal antero-posterior diameter</td>
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<td>Distal transverse diameter</td>
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<td>16.5</td>
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<tr>
<td>Diameter ambiens foramen</td>
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<td>7</td>
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**Table 5.** Measurements (mm) of the tibiotalus of Baptornis

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<td>191</td>
<td>-</td>
</tr>
<tr>
<td>Elevation cnemial process</td>
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<td>14</td>
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<tr>
<td>Diameter proximal articulation</td>
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<td>17</td>
<td>18.5</td>
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<tr>
<td>Antero-posterior diameter shaft*</td>
<td>9</td>
<td>8.5</td>
<td>-</td>
</tr>
<tr>
<td>Transverse diameter shaft*</td>
<td>12</td>
<td>11.5</td>
<td>-</td>
</tr>
<tr>
<td>Diameter distal end</td>
<td>19</td>
<td>18</td>
<td>-</td>
</tr>
</tbody>
</table>

* Below fibular ridge.

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**Figure 15.** Left tibiotalus of Baptornis advenus: a, posterior, b, lateral, c, anterior, and d, medial views, (× 1).
posterior margin, said groove crossing the outer border of the shaft just below the fibular crest. The foramen for the medullary artery lies in this groove. The distal attachment for the spine of the fibula is much smaller than in *Hesperornis*.

The distal end of the tibiotarsus (Figures 14a, 15, 16b) is slightly inflected medially, but not as much as in *Gavia*. In *Hesperornis regalis* and the grebes, the distal end is almost centered on the shaft. The internal and external condyles of the tibiotarsus are spread farther apart than in *Hesperornis* and the posterior crests are not as high. The tendinal groove is broad and terminates in a deep lateral pocket. The internal and external condyles are roughly parallel to each other and the anterior intercondylar sulcus is shallow (more so than in *Hesperornis*). Both the external and internal ligamentary prominences are very low. There is no supratendinal bridge, although a large ligamental attachment occurs above the medial side of the external condyle. In FMNH 395 the high ascending process of the astragulus is still clearly discernable and the fusion of this tarsal element to the tibia evidently took place later in ontogeny than in modern birds.

**Fibula** (Table 6).—The fibula of *Baptornis* (Figure 14c) is most similar to that of *Hesperornis*. The head is large and rectangular. The shaft has two distinct ridges on its posterior proximal surface and the bicipital tubercle is large and elongate. The head is as in *Hesperornis* and is not undercut posteriorly as much as in loons. There is no tubercle for *M. flexor perforatus digiti III* as there is in loons, but instead there is a large triangular roughened area as in *Hesperornis* and grebes.

**Tarsometatarsus and Toes** (Tables 7,8).—As Shufeldt (1915:9) noted, the holotype tarsometatarsus designated by Marsh consists of two portions that are quite probably from different individuals, as indicated by the facts that the fracture lines of the two halves do not coincide and the proximal portion is from a juvenile, while the distal portion appears to be from an adult. If the two pieces had been from one individual, Shufeldt estimated that as much as a third of the shaft must be missing. At our request, the curators of the Division of Vertebrate Paleontology of the Yale Peabody Museum have agreed to retain YPM 1465 for the distal portion of this specimen, which we here designate as the lectotype. The proximal portion has been renumbered as YPM 5768.

KUVP 2290 represents a well-ossified individual in which the proximal and distal ends of a left tarsometatarsus are uncrushed, but in which the middle of the shaft is missing. YPM 5768 is from a young bird, and along with FMNH 395 (Figure 16f,g) and KUVP 16112 shows the lines of fusion between the metatarsal bones. The tarsometatarsus of UNSM 20030 is mature, but crushed.

The tarsometatarsus of *Baptornis* is compressed laterally. The external cotyla is slightly larger than the internal one (Figure 16e) and both tilt slightly anteriorly; the intercotylar prominence is low and blunt as compared to *Hesperornis*; calcaneal ridges are absent, and there are no proximal foramina; the tubercle for tibialis anticus is situated high on the shaft; the anteroproximal face of the shaft is deeply excavated; the attachment for the external ligament is not clearly developed; the outer extensor groove is long and shallow and leads to the region of the distal foramen, with the anterior metatarsal groove running parallel to it; the inner metatarsal groove crosses the medial side about one-third the way down the shaft; the intertrochlear notch between the third and fourth trochleae narrows and then widens so that the dis-

### Table 7.—Measurements (mm) of the tarsometatarsus of *Baptornis*

<table>
<thead>
<tr>
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<th>KUVP 2290</th>
</tr>
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<tbody>
<tr>
<td>Length</td>
<td>84</td>
<td>83</td>
<td>83a</td>
</tr>
<tr>
<td>Proximal antero-posterior diameter</td>
<td>(10)</td>
<td>13.8</td>
<td>13.5</td>
</tr>
<tr>
<td>Proximal width</td>
<td>(17)</td>
<td>17.6</td>
<td>18</td>
</tr>
<tr>
<td>Distal antero-posterior diameter</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Distal width</td>
<td>15</td>
<td>15.1</td>
<td>16</td>
</tr>
<tr>
<td>Tip trochlea II to distal end</td>
<td>11.5</td>
<td>11.5</td>
<td>12</td>
</tr>
</tbody>
</table>

a = restored length; ( ) = measurements from crushed specimens.

### Table 6.—Measurements (mm) of the fibula of *Baptornis*

<table>
<thead>
<tr>
<th>Character</th>
<th>UNSM 20030</th>
<th>FMNH 395</th>
<th>KUVP 2290</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antero-posterior diameter proximal end</td>
<td>7</td>
<td>6</td>
<td>8.5</td>
</tr>
<tr>
<td>Transverse diameter proximal end</td>
<td>11</td>
<td>9</td>
<td>11</td>
</tr>
<tr>
<td>Greatest transverse shaft diameter</td>
<td>7.5</td>
<td>7</td>
<td>–</td>
</tr>
<tr>
<td>Antero-posterior diameter at this point</td>
<td>5</td>
<td>4</td>
<td>–</td>
</tr>
</tbody>
</table>
Figure 16.—Stereophotographs of hindlimb elements of Baptornis advenus: a, proximal end of right tibiotarsus (UNSM 20030); b, distal end of right tibiotarsus (UNSM 20030); c, distal end of left tarsometatarsus (KUVP 2290); d, left tarsometatarsus (UNSM 20030), posterior view; e, proximal end of left tarsometatarsus (KUVP 2290) (not stereo); f, left tarsometatarsus (FMNH 395), anterior view; g, same, posterior view. (All × 1, except c and e, which are slightly enlarged.)
tal foramen communicates with the notch and the metatarsal facet is high and posterior. In UNSM 20030 (Figure 16d) the inner trochlea is set about 7 mm proximal to the middle trochlea. The outer trochlea is in line with the major longitudinal axis of the bone and is approximately equal to the middle trochlea in size. In mature specimens the medial rim of the inner trochlea develops a prominent posterior flange.

There is some ontogenetic variation in the trochleae. Those of YPM 1465 and FMNH 395 have shallow tendinal grooves, are less expanded, and lack the small flange on the inner trochlea that is present in UNSM 20030 and KUVP 2290. The lateral rim of this trochlea extends farther distally than the medial rim.

The outer trochlea extends only slightly farther distally than the middle trochlea and both are about equal in size. In both, the tendinal groove is shallow and continuous along both the plantar and the acrotarsal faces of the middle trochlea, but it is missing from the acrotarsal face of the outer trochlea. The medial rim of the outer trochlea is slightly larger than the lateral rim. The lateral rim extends into a posterior flange in mature specimens. The trochleae are compressed laterally and the extensor grooves run up onto the anterior face as in loons and Hesperornis. In grebes the trochleae are less compressed and tend to be almost smooth on their anterior face. In both grebes and Hesperornis, the inner trochlea has rotated posteriorly, while in loons and Baptornis it is more nearly parallel to the main axis of the shaft.

The tarsometatarsus of Baptornis is like that of Hesperornis in that it is laterally compressed, lacks calcaneal ridges on the hypotarsus, lacks proximal foramina, and the outer trochlea is in line with the longitudinal axis of the bone. Baptornis differs from Hesperornis in that the necks of the trochleae are longer and more delicate.

In Hesperornis the outer extensor groove is very large and deep, and the anterior metatarsal groove is quite prominent, although it is only about one-third the width of the outer extensor groove. There is a distinct groove originating near the distal foramen and extending from the medial rim of the outer trochlea over the trochlea to its lateral rim. This groove is much fainter in Baptornis. In both Baptornis and Hesperornis the distal foramen is well within the intertrochlear notch, between the middle and outer trochleae. Its position in Baptornis is delineated by indentations in the necks of the trochleae, but the distal margin is not closed off to form a real foramen. In Hesperornis extensions of the sides of the trochleae meet to form the distal margin of the foramen, and a suture line is usually visible where they meet.

FMNH 395 includes metatarsal I. It is not as short and flat as in Hesperornis. The bone diverges from the shaft of the tarsometatarsus at a fairly steep angle and terminates in a rounded knob. The phalanx of the first digit is a thin, elongate, highly curved bone. None of the toe bones were articulated, so their identification is somewhat subjective. Fourteen phalanges, counting 2 unguals, are preserved with FMNH 395, 6 with

<table>
<thead>
<tr>
<th>Character</th>
<th>Metatarsal I</th>
<th>Digit II</th>
<th>Digit III</th>
<th>Digit IV</th>
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<tr>
<td></td>
<td>FMNH 395</td>
<td>FMNH 395</td>
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<tr>
<td></td>
<td>1</td>
<td>2</td>
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<tr>
<td>Length</td>
<td>14</td>
<td>22</td>
<td>31.7</td>
<td>37.0</td>
</tr>
<tr>
<td>Proximal vertical diameter</td>
<td>3.8</td>
<td>4.4</td>
<td>11.5</td>
<td>8.8</td>
</tr>
<tr>
<td>Proximal transverse diameter</td>
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<td>2.8</td>
<td>8.3</td>
<td>5.8</td>
</tr>
<tr>
<td>Distal vertical diameter</td>
<td>-</td>
<td>3.5</td>
<td>2.6</td>
<td>8.5</td>
</tr>
<tr>
<td>Distal transverse diameter</td>
<td>-</td>
<td>4</td>
<td>4.7</td>
<td>5</td>
</tr>
</tbody>
</table>

( ) = Measurements from crushed specimens.
Figure 17.—Left foot of *Baptornis advenus*: a, anterior view; b, lateral view.

UNSM 20030, and 3 fragments with KUVP 16112. Phalanx 1 of digit II appeared to be absent from our sample. Phalanx 2 of digit II is flattened dorso-ventrally, with a broad proximal end, which rapidly narrows anteriorly. It has a simple distal articulation for the claw. Phalanx 1 of digit III is the largest of the foot. It is flattened laterally, with a deep proximal articulation. The distal articulation consists of two parallel ridges, which would permit mainly anterior or posterior movements. Phalanx 2 of digit III is a smaller version of the first. Phalanx 3 of digit IV is about the same length as that of digit III but neither articulates as in loons or grebes. That of digit IV is narrower and rounder than in the other toes, and the distal end has the medial ridge of the articulation inclined, suggesting that some degree of toe-rotation was possible. Phalanges 2 and 3 of digit IV are much shorter and deeper than the first phalanx, while the fourth is elongate, laterally compressed, and terminates in an articulation for the claw.

Although phalanx 1 of digit IV is more elongate than in *Hesperornis*, nevertheless the third and fourth toes of *Baptornis* are of about the same size and length (Figure 17), whereas in *Hesperornis* the fourth toe is much the longest. The phalanges of the fourth toe also lack the deep ventral grooving found in *Hesperornis*. The claws of *Baptornis* are somewhat flattened, gently curved and pointed. They are not the broad nail-like structures that grebes have.

Grebes have lobed feet while loons have webbed feet. When swimming, both loons and grebes have their toes spread for the power stroke. On the recovery stroke, loons fold their toes posteriorly while grebes rotate their foot 90° (Storer, 1958) so that the edges of the lobes cut through the water. These actions are reflected in the morphology of the phalanges of digit IV. In loons the distal articular ridges are parallel to each other and are about the same size. This is the condition found in most birds. In grebes the medial ridge is enlarged and the lateral ridge is small and rounded (Figure 18c). In *Hesperornis* the distal articulations of the phalanges for digit IV are even more specialized, with the medial ridge extending over the small, rounded lateral ridge (Figure 18d) suggesting highly developed toe-rotation and lobed feet. *Baptornis* differs from the loons in having the medial ridges enlarged and inclined on the distal articulations of the phalanges for digit IV. However, it is not nearly as specialized as either grebes or hesperornithids (Figure 18b).

**HABITS OF *Baptornis***

From the skeletal remains now available, it appears that adults of *Baptornis* were about one meter long from the tip of the tail to the tip of the
Figure 19.—Skeleton of *Baptornis advenus* (stippled areas restored after *Hesperornis*).
Figure 20.—Restoration of Baptoptis advenus. (The feet probably were not totipalmate, as shown here, and may even have been lobed.)
bill. We have restored the bird with teeth (Figures 19, 20) because of its relationship to the toothed bird *Hesperornis*. All Mesozoic birds for which dentaries are known (*Archaeopteryx*, *Hesperornis*, and *Ichthyornis*) had teeth.

The skeletons of both *Hesperornis* and *Baptornis* are nonpneumatic and are composed of relatively heavy, compact bone. Bones of *Hesperornis* are often recognized in the field because they appear more solid than most of the other small bones in the Niobrara Chalk. This is hardly surprising, for additional weight is actually of an advantage for a diving animal. The added weight, along with the long necks and fusiform bodies of the Cretaceous divers, suggest that they might have swum partially submerged, like modern anhingas. However, neither *Baptornis* nor *Hesperornis* appears to have had the neck modified for stabbing. This is further

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**Figure 21.**—Pelvis and right hindlimb of *Baptornis advenus*, dorsal view.
emphasized by their relatively broad, blunt bills. In both Baptornis and Hesperornis the wing is greatly reduced. This is often the case in marine foot-propelled divers; various adaptations for flight would make them too buoyant for efficient diving, while the loss of flight is not too serious a hardship for them. On the other hand, foot-propelled divers that occupy disjunct bodies of fresh water need to be able to fly in order to disperse. Lonchodytes is the only known nonmarine foot-propelled diver from the Mesozoic. It is from freshwater river deposits, and appears to have been volant (Brodkorb, 1963a). The tiny wing of Baptornis could scarcely have been used for locomotion, but it may have had a stabilizing and steering function, much like the pectoral fins in fishes. The sternum in Baptornis is unusually small, even for a flightless bird, and the body must have been very long and narrow. The tibiotarsus was bound to the pelvis as it is in loons, but it was probably not as closely appressed to the postacetabular region as it was in Hesperornis, and the bone itself extended somewhat past the posterior margin of the pelvis (Figures 19, 21). In Hesperornis the distal end of the tibiotarsus is about even with the posterior margin of the pelvis. The above features, along with characters of the acetabulum and femur, suggest that Baptornis had some capacity to rotate its feet under its body, probably for swimming on the surface. When diving, the feet were certainly held out to the side. It seems almost certain that Baptornis could not walk upright on land and in fact must have pushed itself along on its stomach like a seal or a loon. The feet could have been webbed or lobed. They do show some evidence of inefficient toe-rotation and might well have been lobed, although our restoration shows them webbed. Hesperornis almost certainly had lobed feet. The feet of Baptornis are exceptionally large for the size of the bird. The tail is quite long, with a long, laterally compressed pygostyle that probably acted as a rudder. Loons appear to have similar, but shorter tails, while in Hesperornis the tail is long but is dorsoventrally compressed and has a short pygostyle.

Included with UNSM 20030 are eight coprolites, two of which show small fish jaw and other bones. Most are round or elliptical in cross-section and are elongate, except for the two containing the fish material. None shows spiral grooving or surface impressions. George Sternberg, the collector, in a 1937 communication preserved in the records of the University of Nebraska State Museum, makes the following reference to the association of these coprolites with the Baptornis skeleton: "There are 7 or 8 coprolites; . . . . 2 show small fish bones. These are small compared to other coprolites I have seen and were found mingled with the bones." It seems likely that these coprolites are correctly associated with the Baptornis skeleton; if so, they are the only ones known for a Cretaceous bird. However, several of them fit together to form a long rounded structure that might be better interpreted as an intestinal cast (Figure 22a). A fish jaw in one coprolite was identified by Orville W. Bonner of the University of Kansas (pers. comm., 1972) as Enchodus cf. parvus Stewart. Enchodus was a very common genus of fish in the Niobrara seas. It included some very small species, which probably formed a significant item in the diet of the Late Cretaceous toothed birds. Hesperornis had scutellate-rieticulate tarsi (Williston, 1898). We have a fragment of matrix associated with UNSM 20030 showing what may possibly be the impression of scutes (Figure 22b), suggesting that the tarsometatarsus of Baptornis might also have been scutellate-rieticulate (grebes have reticulate tarsi and loons have scutellate-rieticulate tarsi).

Hesperornis and Baptornis form a natural group of specialized divers, separate from, and more specialized than any of the other foot-propelled divers. Their advanced degree of specialization is not surprising, as the Mesozoic birds occupied a fairly stable environment for the entire Cretaceous and must have been well adapted to it.

Figure 22.—Coprolites (a) and possible scute impressions from the foot (b) of Baptornis advenus (UNSM 20030). (Reduced.)
Baptornis was probably able to range far from shore, as most of the finds indicate. At the present, all of the known records of Baptornis are from the Smoky Hill Member of the Niobrara Chalk Formation, Upper Cretaceous of Central Kansas. The immature specimens suggest that Baptornis may have nested in this region.

Afferities of Baptornis

Baptornis and Hesperornis share a number of primitive features with Archaeopteryx. These include: (1) the nature of shoulder girdle, and in particular the shape of the coracoid, with the glenoid and scapular facets near the tip of the scapular end; (2) the character of the distal articulation of the ulna in Baptornis (the ulna of Hesperornis is unknown), which makes the presence of a fused carpometacarpus doubtful; (3) the separation of the distal segments of the ilium, ischium, and pubis; (4) the absence of a supratendinal bridge on the tibiotarsus; (5) the prominence and fusion late in ontogeny of the ascending process of the astragalus on the tibiotarsus; (6) the absence of hypotarsal canals, a closed distal foramen, or proximal foramina in the tarsometatarsus (proximal foramina are present in all modern birds we have examined). To the above we can probably add the various primitive features of the skull, including the shape of the quadrate and the presence of teeth.

The absence of a supratendinal bridge, hypotarsus, and proximal foramina indicates that these are not necessary for foot-propelled diving. Further, the presence of these characters in modern foot-propelled divers suggests that these birds may have been derived from ancestors that were not foot-propelled divers.

The supratendinal bridge is well developed in loons and cormorants, and this may be related to the fact that the distal end of the tibiotarsus in these forms is inflected medially. The supratendinal bridge is not well developed in some grebes which have (as do hesperornithiform birds) a straight articulation between the tarsometatarsus and tibia, causing the foot to be brought more directly out to the side of the body.

Baptornis and Hesperornis also share a large suite of derived characters. Many of these are directly related to foot-propelled diving and can be found in one state or another in all birds that share this form of locomotion. Some of the characters that tend to unite Baptornis and Hesperornis, in addition to the shape and reduction of the shoulder girdle and the wing, are: (1) the broad shallow fibular groove on the femur; (2) the large trihedral patella with a foramen for the tendon of the ambiens muscle; (3) the large triangular pronemial crest on the tibiotarsus; and (4) the compressed tarsometatarsus with the anterolateral edge developed as a high thin ridge. No particular relationship is indicated between Baptornis and any living group of foot-propelled divers. The foot is more similar to that of loons than of grebes, but the coracoid is totally unlike that of any living bird. Throughout, the skeleton is most similar to that of the Hesperornithidae.

Marsh did not attempt to classify Baptornis, but clearly regarded it as a member of his order Odontolcae, which included Hesperornis. His student Williston (1898) included it in this order when he listed the Cretaceous birds of Kansas. Lucas (1903) later suggested that it was separable from Hesperornis at the familial and perhaps the ordinal level. Lambrecht (1933) thought Baptornis was related to the loons and grebes and placed it in the family Enaliornithidae, in which he also included Enaliornis, Neogaeornis, and possibly Eupterornis of the French Paleocene. Romer, in earlier editions of his text Vertebrate Paleontology, placed Eupterornis questionably in the Baptornithidae, but later (1966) followed Brodkorb (1963b) in placing Eupterornis in the Gaviiformes and Baptornis in the Podicipediformes. Wetmore (1956) used the superorder Odontognathae for the New World toothed birds and placed the Baptornithidae in the order Hesperornithiformes. Brodkorb (1963b) treated the Baptornithidae as an ancient family of grebes, which included two genera, Baptornis and Neogaeornis. At the same time, he put Enaliornis and Eupterornis in the Gaviiformes (loons), along with the volant Mesozoic foot-propelled divers of the family Lonchodytidae. Brodkorb (1971:89) has since reaffirmed his opinion regarding Baptornis in stating that “the proper position of the family Baptornithidae is surely in the order Podicipediformes.” This position follows closely a statement by Lucas (1903:555): “In the slender cervical, arrangement of the tibia and patella, and
general structures of the leg, Baptornis is more like a grebe than is the contemporary Hesperornis, and if, with the small amount of material available, it is deemed essential to establish any connection between groups of existing and fossil birds, it is suggested that the ancestors of Baptornis are much more likely to have been also the progenitors of the Colymbine group [= grebes] than are those of Hesperornis." However, examination of much better material of both Hesperornis and Baptornis than was available to Lucas has failed to substantiate his statement. Actually, Hesperornis is more grebe-like than is Baptornis, which tends more to resemble the loons. Our studies here confirm that Baptornis and Hesperornis resemble each other more than they do any modern birds and that both are far removed from, and unrelated to, either loons or grebes. This removes the Podicipediformes from the Cretaceous record, their next earliest occurrence being in the lower Miocene (Arikareean) of Oregon (Brodkorb, 1963b:227).

Other than Baptornis the only genus we presently include in the Baptornithidae is Neogaeornis from the Late Cretaceous of Chile. We have not examined specimens of Neogaeornis, but the highly compressed tarsometatarsus with the outer trochlea slightly more distal than the middle one (Lambrecht, 1933, fig. 100) suggests affiliation with the Hesperornithiformes. The fact that the outer and middle trochleae are of about equal size supports an allocation to the Baptornithidae rather than the Hesperornithidae.

Enaliornis, from the Lower Cretaceous of Britain, shares many features with the Hesperornithiformes, including a broad shallow fibular groove on the femur, a broad triangular procnemial process and the absence of the supratendinal bridge of the tibiotarsus, and the outer trochlea of the tarsometatarsus slightly more distal than the middle trochlea. Enaliornis shares many features with Baptornis, from which it differs in having amphicoelous dorsal vertebrae. We presently maintain the Baptornithidae as a family separate from the Enaliornithidae. Brodkorb (1963a:60) placed Enaliornis in the Gaviiformes stating that "although currently placed in the order Hesperornithiformes, these birds are slightly less specialized than modern loons of the family Gaviidae and should be transferred to the order Gaviiformes." However, we think Enaliornis is more similar to Baptornis than to any other known bird and regard it as the primitive basal stalk of the Hesperornithiformes. This leaves Lonchodytes as the only Mesozoic bird with any possible affinities with loons.

We agree with Baird's (1967) suggestion that the order Hesperornithiformes probably represents an archaic group that became extinct along with the great sea-lizards at the close of the Cretaceous.

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Walker, M. V.

Wetmore, A.

Williston, S. W.

Zusi, R. L., and R. W. Storer
Discovery of a Cretaceous Bird, Apparently Ancestral to the Orders Coraciiformes and Piciformes (Aves: Carinatae)

Pierce Brodkorb

ABSTRACT

Alexornis antecedens, new genus and species, is described from the Bocana Roja Formation, Upper Cretaceous (Campanian age), near El Rosario, Baja California, Mexico. The humerus, ulna, scapula, coracoid, femur, and tibiotarsus are represented. The fossil is referred to a new family, Alexornithidae, and a new order, Alexornithiformes, thought to be ancestral to the Tertiary and Recent orders Coraciiformes and Piciformes. Since Caenagnathus collinsi Sternberg and C. sternbergi Cracraft are reptiles, and Gobipteryx minuta Elzanowski appears to be reptilian also, Alexornis is the only certain land bird known from the Cretaceous.

Introduction

The major deficiency in our knowledge of the history of birds is their inadequate Mesozoic record. The only known Jurassic genus is Archaeopteryx, which was certainly a land bird, whether it was arboreal (Brodkorb, 1971b) or a ground-dweller (Ostrom, 1974). In the Cretaceous period we have records of about 37 species of water birds, distributed among 18 genera, 12 families, and 7 orders. Because many groups of land birds occur early in the Tertiary, they must also have been present and undergoing radiation during the Cretaceous. However, none of the hitherto known alleged birds from terrestrial Cretaceous deposits is certainly referable to the class Aves.

Three species of supposed land birds have been described from the late Cretaceous. These are: Caenagnathus collinsi Sternberg (1940), Caenagnathus sternbergi Cracraft (1971), and Gobipteryx minuta Elzanowski (1974). Each was described from a single specimen of jaw or skull. Although not comparable with any living or fossil bird, the describers of Caenagnathus and Gobipteryx made them the types of new families and orders of Aves. Sternberg (1940) based Caenagnathus collinsi on a mandible from the Belly River Series of Alberta, and regarded it as representing a new order of birds. Wetmore (1960) transferred it to the theropod dinosaurs, near Ornithomimidae, an assignment in which Romer (1966) concurred. Cracraft (1971) returned the genus to Aves and founded a second species, C. sternbergi, on a scrap of a lower jaw from the same formation as C. collinsi. White (1973) included Caenognathus [sic] and Caenognathidae [sic] as a valid genus and family of coelurosaurian theropods. Dale A. Russell of the National Museum of Canada (pers. comm.) informs me that new material of Caenagnathus, plus specimens of related forms from Mongolia, confirms that the Caenognathidae are theropod dinosaurs having affinities with Oviraptor of the Ornithomimidae, thus vindicating Wetmore.

Gobipteryx minuta was described on a small, poorly preserved, flattened skull and mandible from the Barun Goyot Formation in Mongolia. Neither the specimen, which I have seen, nor the published illustrations, are convincingly avian.
Elzanowski (1974) placed Gobipteryx in the “Superorder Palaeognathae” (i.e., Ratitae), but this is certainly wrong. The specimen is quite small, and the palate is unlike that of ratites or any other bird. Better material is needed before it can be assigned confidently to either Aves or Reptilia.

With the relegation of the above forms to Reptilia or taxa incertae sedis, there are no land birds known from the 72-million-year span of the Cretaceous period. It was therefore with great interest that I agreed to study some possibly avian remains from a terrestrial deposit of Late Cretaceous age in Baja California, Mexico. The best preserved specimens so far recovered are from a small land bird—represented by elements of the pectoral girdle, wings, and legs—the subject of the present paper.

Stratigraphy.—The Rosario Group is composed of three formations, each separated by an unconformity. In descending order these are the Rosario, “El Gallo,” and “La Bocana Roja.” According to Morris (1972) the last two formations were defined by Kilmer in his doctoral dissertation (1963).

The Rosario Formation is a marine deposit whose invertebrate fauna was assigned an early Maestrichtian or late Campanian age by Durham and Allison (1960). The nannofossils indicate that it is situated close to the Maestrichtian-Campanian boundary (Morris, 1973).

The Gallo Formation has a thickness of more than 150 m. A potassium/argon (K/Ar) date of 73 ± 2 million years is available for the middle third of the formation (Morris, 1972, 1973). This would make it of late Campanian age. It contains the remains of hadrosaurian dinosaurs (cf. Lambeosaurus), smaller reptiles, amphibians, mammals, and large tree trunks with well-developed root systems. This assemblage is the only extensive Late Cretaceous terrestrial vertebrate fauna from the Pacific margin of North or Middle America.

The Bocana Roja Formation contained the remains of the bird described here. The type-specimen of the theropod dinosaur Labocanania anomalula Molnar (1974) also came from this formation, along with hadrosaur ribs and numerous small vertebrate fossils, as yet unstudied. The age of the formation is Campanian or earlier.

During the Cretaceous period the arrangement of the continents was different from that of today, Mexico being in proximity to North Africa and bordering the western part of the Tethys Sea (Deitz and Holden, 1970).

Acknowledgments.—Field work was supported by the National Geographic Society through grants to Dr. William J. Morris of Occidental College and was sponsored by the Instituto de Geología of the Universidad Nacional Autónoma de México and by the Natural History Museum of Los Angeles County. Over several years the field parties included, in addition to Dr. Morris, Dr. Ismael Ferrusquia V. of the Instituto de Geología and Harley Garbani, J. Loewe, and Robert M. McKenzie of the Natural History Museum of Los Angeles County (LACM). I am greatly indebted to Dr. Hildegarde Howard for suggesting that the fossils be sent to me for study. The beautiful photographs of the tiny specimens are by Victor E. Krantz of the Smithsonian Institution. Donald Baird, Nicholas Hotton III, John H. Ostrom, and Dale A. Russell made suggestions or furnished information about Caenagnathus and Gobipteryx.

Alexornis, new genus

Type of Genus.—Alexornis antecedens, new species.

Diagnosis.—As for the type and only known species.

Etymology.—From Greek aléxō (I defend) and ornis (common gender, masculine selected here) bird. On the occasion of his ninetieth birthday this genus is dedicated to my friend Alexander Wetmore, who, in addition to his many other accomplishments, has done more to foster palornithology and has described more species of fossil birds than any other author.

Alexornis antecedens, new species

Holotype.—Distal 10 mm of right humerus, LACM 33213 (Figure 1a,b). From LACM locality 7256, 6 miles southwest of El Rosario, Baja California del Norte, Mexico. Bocana Roja Formation, Upper Cretaceous, Campanian age. Collected by H. Garbani and J. Loewe, 16 July 1971.

Paratype.—Distal 10 mm of left humerus, collected in association with the holotype and cataloged with the same number.
Figure 1.—Alexornis antecedens, new genus and species, Bocana Roja Formation, Campanian age, near Rosario, Baja California, Mexico (LACM 33212): a, holotype right humerus, palmar view; b, same, anconal view; c, left scapula, ventral view; d, same, dorsal view; e, left coracoid, anterior view; f, same, posterior view; g, right ulna, internal view; h, same, external view; i, right tibiotarsus, anterior view; j, same, posterior view; k, left femur, anterior view; l, same, posterior view; m, same, lateral view. (All X 5.)
Hypodigm.—The holotype, paratype, and the following referred material, collected in association with the types and cataloged under the same number: proximal 4 mm of left scapula (Figure 1c,d), upper 4.4 mm of left coracoid (Figure 1e,f), proximal 3.5 mm of right ulna (Figure 1g,h), distal 6 mm of left femur (Figure 1k-m), and proximal 16.3 mm of right tibiotarsus (Figure 1i,j). More than 20 other fragments were also collected with the types and cataloged under the same number, but they are left unidentified at this time.

Diagnosis.—Comparison of the hypodigm with the homologous skeletal elements of the known orders of birds shows that the resemblances of Alexornis are closest to certain members of the Piciformes and Coraciiformes. Within those two orders the piciform family Bucconidae and the coraciiform family Momotidae have the most similarity to the fossil. The fossil shares certain characters with both Bucconidae and Momotidae, some with Bucconidae alone, and some with Momotidae alone; but more of its characters are unique. In size the fossil falls between the bucconid Malacoptila panamensis and the motmot Hylomanes momotula (Table 1).

Etymology.—Latin antecedens, going before in rank or time, ancestral, in reference to the supposed ancestry of this bird to the orders Piciformes and Coraciiformes.

Description.—Humerus: (1) Olecranal fossa deep (shallow in Bucconidae and Momotidae), and (2) wide, extending toward entepicondylar area (as in Momotidae; in Bucconidae less extended toward entepicondyle). (3) Entepicondyle produced distally, resembling Bucconidae (in Momotidae less produced; in Passeriformes much more produced). (4) External condyle bulbous as in both Bucconidae and Momotidae, but (5) oriented transversely at an angle of about 60 degrees to shaft (in Momotidae angle to shaft is about 45 degrees; in Bucconidae condyle is more upright at angle of 30 degrees to shaft). (6) Internal condyle lies inclined toward entepicondyle, resembling condition in Momotidae (in Bucconidae condyle is more transverse), (7) with a strong facet for medial cotyla of ulna, resembling both Momotidae and Bucconidae. (8) Ectepicondylar prominence large and rounded, resembling Bucconidae (less developed in Momotidae), (9) with a transverse ridge across anconal surface proximal to its base (ridge absent in Bucconidae and Momotidae).

Insofar as preserved, the humerus of Alexornis has three unique characteristics (numbers 1, 5, and 9 above), shares two with Bucconidae alone (3 and 8), shares two with Momotidae alone (2 and 6), and is similar to both Bucconidae and Momotidae in two others (4 and 7).

Ulna: (1) Olecranon straight, short, stout (as in Bucconidae and Momotidae), (2) with a pit in the tip (no pit in Bucconidae and Momotidae). (3) External cotyla large and (4) strongly convex (of moderate size and moderately convex in Bucconidae, and (5) with a transverse ridge across anconal surface proximal to its base (ridge absent in Bucconidae and Momotidae).

Table 1.—Measurements (mm) of skeletal elements

<table>
<thead>
<tr>
<th>Character</th>
<th>Alexornis antecedens</th>
<th>Hylomanes momotula</th>
<th>Malacoptila panamensis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HUMERUS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Distal width</td>
<td>5.6, 5.7</td>
<td>4.4</td>
<td>4.8</td>
</tr>
<tr>
<td>Shaft width</td>
<td>2.2, 2.3</td>
<td>1.7</td>
<td>2.4</td>
</tr>
<tr>
<td>Shaft depth</td>
<td>1.8, 1.8</td>
<td>1.6</td>
<td>1.8</td>
</tr>
<tr>
<td>Least depth through brachial depression</td>
<td>1.1, 1.3</td>
<td>1.1</td>
<td>1.7</td>
</tr>
<tr>
<td><strong>SCAPULA</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diagonal width across acromion and glenoid</td>
<td>4.2</td>
<td>3.5</td>
<td>4.6</td>
</tr>
<tr>
<td>Shaft width</td>
<td>1.7</td>
<td>1.6</td>
<td>1.5</td>
</tr>
<tr>
<td><strong>FEMUR</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Distal width</td>
<td>[3.1]</td>
<td>3.4</td>
<td>3.7</td>
</tr>
<tr>
<td>Shaft width</td>
<td>2</td>
<td>1.7</td>
<td>2</td>
</tr>
<tr>
<td>External condyle depth</td>
<td>2.35</td>
<td>2.5</td>
<td>2.5</td>
</tr>
<tr>
<td>Fibular condyle depth</td>
<td>1.6</td>
<td>2.2</td>
<td>2.2</td>
</tr>
<tr>
<td>Internal condyle depth</td>
<td>2.6</td>
<td>2.0</td>
<td>2.1</td>
</tr>
<tr>
<td>External condyle height</td>
<td>2.5</td>
<td>1.5</td>
<td>1.6</td>
</tr>
<tr>
<td>Fibular condyle height</td>
<td>1.6</td>
<td>1.3</td>
<td>1.4</td>
</tr>
<tr>
<td>Internal condyle height</td>
<td>2</td>
<td>1.7</td>
<td>1.3</td>
</tr>
<tr>
<td>Shaft depth</td>
<td>1.8</td>
<td>1.4</td>
<td>1.7</td>
</tr>
<tr>
<td><strong>TIBIOTARSUS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Proximal width</td>
<td>3.7</td>
<td>2.6</td>
<td>2.9</td>
</tr>
<tr>
<td>Shaft width</td>
<td>1.9</td>
<td>1.3</td>
<td>1.6</td>
</tr>
<tr>
<td>Shaft depth</td>
<td>1.6</td>
<td>1.2</td>
<td>1.6</td>
</tr>
<tr>
<td>Width through internal cnemial crest</td>
<td>4.1</td>
<td>3.1</td>
<td>4</td>
</tr>
<tr>
<td>Length through fibular ridge</td>
<td>10.5</td>
<td>6.7</td>
<td>8.1</td>
</tr>
</tbody>
</table>

*The first measurement of the humerus of Alexornis is of the holotype, the second of the paratype. Measurement in brackets estimated.
dae; small and concave in Momotidae), (5) separated from olecranon by a deep groove (groove absent in Bucconidae and Momotidae); (6) medial rim of external cotyla thick [lateral portion of cotyla missing] (resembling Bucconidae; edge very thin in Momotidae). (7) Internal cotyla small (as in Bucconidae; large in Momotidae), (8) with surface flat (moderately concave in Bucconidae; strongly concave in Momotidae). (9) Proximal radial depression deeply undercuts entire width of rim of internal cotyla (in Bucconidae the depression falls far short of rim of cotyla; in Momotidae the depression extends to the medial edge of cotyla but fails to undercut it).

Insofar as preserved, the ulna of Alexornis has six unique characteristics (numbers 2, 3, 4, 5, 8, and 9 above), shares two with Bucconidae alone (6 and 7), shares none with Momotidae alone, and is similar to both Bucconidae and Momotidae in one characteristic (1).

Scapula: (1) Acromion rather short (long in Bucconidae and Momotidae), with tip slightly damaged, but (2) apparently blunt (as in Bucconidae; tip forms a recurved hook in Momotidae and Passeriformes). (3) Glenoid facet flat (cup-like in Bucconidae and Momotidae).

Thus the scapula, so far as preserved, has two unique characteristics (numbers 1 and 3), shares one with Bucconidae (2), and none with Momotidae.

Coracoid: (1) Brachial tuberosity with a recurved hook directed toward area where procoracoid process would be if preserved (slightly hooked in Bucconidae; hook absent in Momotidae). (2) Triosseal canal very deep (very shallow in Bucconidae; flat in Momotidae). (3) Scapular facet convex (as in Bucconidae and Momotidae), but (4) very broad (very narrow in Bucconidae and Momotidae).

The coracoid has three unique features (numbers 1, 2, and 4), none is shared with Bucconidae alone or with Momotidae alone, and one is shared with both Bucconidae and Momotidae (3).

Femur: The specimen shows evidence of some postmortem compression and distortion. (1) Shaft stout (resembling Bucconidae; slender in Momotidae). (2) Fibular condyle with only very slight indication of a transverse shelf on posterior surface (resembling Bucconidae and Eurylaimidae; shelf very prominent in Momotidae). (3) Fibular condyle small (resembling Momotidae and Eurylaimidae; very stout in Bucconidae). (6) Popliteal area deeply excavated (resembling Momotidae; area nearly flat in Bucconidae and Eurylaimidae). (7) Rotular groove shallow (well developed in Bucconidae and Momotidae).

The femur has three unique features (numbers 2, 3, and 7), three are shared with Bucconidae alone (1, 4, and 5), one with Momotidae alone (6), and none are held in common with both Bucconidae and Momotidae.

Tibiotarsus: The cnemial crests are broken off, but their bases are preserved. (1) Shaft stout (resembling Bucconidae; slender in Momotidae). (2) Fibular crest wide (rudimentary in Bucconidae and Momotidae), (3) extending proximally all the way up shaft (as in Momotidae; in Bucconidae falling far short of proximal end of shaft). (4) Distal end of fibular crest merges gently with shaft (as in Momotidae; in Bucconidae distal end joins shaft at a rather abrupt angle). (5) Anterior and posterior surfaces of fibular crest concave, with an anterior and a posterior groove running along junction with shaft (a slight anterior and posterior groove in Bucconidae; in Momotidae anterior and posterior surfaces of crest flat and ungrooved). (6) Proximal internal articular surface swollen and convex (in Momotidae slightly swollen; surface more concave in Bucconidae). (7) Without posterior overhang (with slight overhang in Bucconidae; in Momotidae a lip overhangs shaft posteriorly). (8) Outer cnemial crest short (as in Bucconidae and Momotidae). (9) Inner cnemial crest short, although considerably longer than outer crest (resembling Bucconidae and Momotidae).

The tibiotarsus has three unique characteristics (numbers 2, 5, and 7), shares one with Bucconidae alone (1), shares three with Momotidae alone (3, 4, and 6), and agrees with both Bucconidae and Momotidae in two features (8 and 9).

Familial Position of Alexornis

The characteristics described above for Alexor-
nis are grouped in Table 2 to show the number of features confined to a single taxon, those shared by two taxa, and those common to all three taxa. By far the strongest grouping of characteristics is of those confined to a single taxon—49 percent in *Alexornis*, 44 percent in Momotidae, and 37 percent in Bucconidae. I interpret this as indicating that the three taxa are of equal taxonomic rank and, therefore, propose *Alexornis* as the type of a new family.

Table 2.—Summary of shared characteristics of *Alexornis*, Momotidae, and Bucconidae (+ = similar to; — = different from)

<table>
<thead>
<tr>
<th>Wing</th>
<th>Pectoral</th>
<th>Leg</th>
<th>Total</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>girdle</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(18)</td>
<td>(7)</td>
<td>(16)</td>
<td>(41)</td>
<td></td>
</tr>
</tbody>
</table>

1. *Alexornis* + Momotidae + Bucconidae
2. *Alexornis* — Momotidae + Bucconidae
3. *Alexornis* + Bucconidae — Momotidae
4. Bucconidae + Momotidae — *Alexornis*
5. Bucconidae — Momotidae — *Alexornis*
6. Momotidae — Bucconidae — *Alexornis*
7. *Alexornis* — Momotidae — Bucconidae

Numbers in parentheses represent number of characters considered.

**ALEXORNITHIDAE**, new family

**Diagnosis.**—Humerus with olecranal fossa deep; entepicondylar area much produced distally; external condyle oriented transversely at an angle of about 60 degrees to shaft; ectepicondylar promi-

nence with a transverse ridge across anconal surface proximal to its base. Ulna with a pit at the tip of olecranon; external cotyla large, strongly convex, and separated from olecranon by a deep groove; surface of internal cotyla flat; proximal radial depression deeply undercutting the entire width of lip of internal cotyla. Scapula with acromion rather short and glenoid facet flat. Coracoid with brachial tuberosity hooked; triosseal canal very deep; scapular facet very broad. Femur with external condyle very long, extending both proximally and distally far beyond both internal and fibular condyles; rotular groove obsolete. Tibiotarsus with its proximal articular surface not overhanging shaft posteriorly; fibular crest wide, with both its surfaces concave and separated from shaft by an anterior and a posterior groove.

**Ordinal Position of Alexornis**

Although possessing a large number of unique features, Alexornithidae shares some characters with the order Coraciiformes as exemplified by the Momotidae, others with the order Piciformes as exemplified by the Bucconidae, and still others with both of those orders. These similarities are summarized in Table 2, and several different hypotheses at the ordinal level could be formed from these data.

The three taxa might be combined in a single order, for which the name Piciformes would have priority (for order-group synonymies see Brodkorb, 1971a:248, 256). But of the 41 characters analyzed, only 6 are shared by the 3 families (Table 2, line 1). Such a small proportion of common characteristics militates against merging the taxa in a single order.

Alexornithidae might be referred to Coraciiformes, but such an arrangement is also supported by 6 characters only (Table 2, line 2), and this hypothesis is likewise discarded.

Alexornithidae might be placed in Piciformes, as the order is currently understood. Nine characters support this combination (Table 2, line 3), but line 4 of Table 2 argues against it, as Piciformes and Coraciiformes share more characters than any other combination of the taxa under consideration.

The conclusion thus derived from Table 2 is that the three taxa represent separate but related
orders. I therefore propose Alexornis as the type-genus of a new order.

ALEXORNITHIFORMES, new order

Diagnosis.—Same as for the only known family, Alexornithidae.

Remarks.—The age of Alexornis is about 81 million years BP, much earlier than the earliest known occurrence of either the Coraciiformes and Piciformes. The earliest record of the Piciformes is early Eocene, about 51 million years BP, when bucconid-like forms appear in Wyoming (Brodkorb, 1970; Feduccia and Martin, herein). If Harrison and Walker (1972) are correct in assigning the British Halcyornis to the Coraciiformes, the earliest record of that order is also early Eocene. Undoubted members of the Coraciiformes occur in European deposits of middle and late Eocene age (Brodkorb, 1971a). Both morphology and the temporal sequence thus suggest Alexornis as the presumptive ancestor of the orders Coraciiformes and Piciformes.

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Romer, A. S.

Sternberg, R. M.

Wetmore, A.

White, T. E.
A Survey of the Paleogene Birds of Asia

E. N. Kurochkin

ABSTRACT

In the past few years, much new material of Paleogene birds has been collected in Mongolia and Kazakhstan, the fossil collections now comprising 350 satisfactory specimens of postcranial elements. The first Paleocene birds known from Asia were obtained in Mongolia, where remains referable to several avian orders were recovered. Fossils from a new middle Eocene site at Khaichin Ula 2, also situated in the South Gobi of Mongolia, were identified as various waterbirds and galliforms. A variety of birds is represented in early Oligocene material from Mongolia and a wealth of new avian material of middle Oligocene age has been obtained from the so-called Indricotherium beds of Central Kazakhstan. Paleoecological conditions in the Asian Paleogene are assessed on the basis of avian fossils, and the sketchy picture of the possible interrelationships of Paleogene birds from Asia, Europe, and North America is outlined.

Abundant new fossil material has made possible a deeper insight into the composition, characteristics, and relationships of the gruiform families Eogruidae and Ergilornithidae, and proves the existence of a phylogenetic continuity between the Eocene genus Eogrus, the Oligocene genera Ergilornis and Proergilornis, and the Pliocene genus Urmiornis. A fragment of humerus assigned to the Ergilornithidae shows that these birds were flightless.

Introduction

Until recently, knowledge of the Paleogene birds of Asia has been derived mainly from casual discoveries, with few attempts at a purposeful gathering of avian remains and their subsequent study being made. Publications on the subject have generally been concerned with systematic descriptions of individual specimens collected in association with Paleogene mammals from isolated Asian localities. The list of these publications is deplorably short, consisting of 15 titles, in which 21 forms of birds are described.

Discoveries of Paleogene birds in Asia have been confined largely to the following three territories: Central Kazakhstan east of the Aral Sea; East Kazakhstan east of Lake Zaisan; and the southern part of Mongolia together with adjoining provinces of China (Figure 1). Avian fossils from outside this area consist of individual specimens from the early middle Eocene of Fergana in Kirgizia (Eobalearica tugarinovi Gureev, 1949), the middle Eocene of Sumatra (Protoplotus beauforti Lambrecht, 1931), and Gruidae genus indet. from the Oligocene of Ordos, China (Teilhard de Chardin, 1926).

Three upper Eocene birds, two of which were described as new, were reported by Wetmore (1934) from China. Lower Oligocene ergilornithids from East Gobi, Mongolia, were studied by Kozlova (1960). Several species of birds of prey, Anseriformes and rails were described by Kurochkin (1968a, 1968b) from the Oligocene of Kazakhstan and Mongolia. Other specimens of Paleogene birds from Asia were discussed by Tugarinov (1940) and Bendukidze (1971). A record of Aquila sp. from middle Oligocene deposits at Min Eske Suek, Kazakhstan (Aubekerova, 1965) has not been confirmed. Surveys of the Tertiary birds from the U.S.S.R, including several Paleogene discoveries, were published by Burchak-Abramovich (1958) and the present writer (Kurochkin, 1974).

In the past few years, great quantities of new avian material have been obtained from the Oligo-
Figure 1.—Schematic map showing collection localities for Paleogene birds in Asia. **Paleocene:** 1, Naran Bulak and Tsagan Khushu. **Middle Eocene:** 2, Khaichin Ula 2; 3, Kolobolchi; 4, Kalmakpai; 5, Andarak (in Fergana). **Late Eocene:** 6, Andreevka; 7, Irdin Manha and Ulan Shireh; 8, Iren Dabasu; 9, Tung Gur (Miocene); 10, (inset) Sipang, Sumatra. **Early Oligocene:** 11, Zhongiz Shoki; 12, Ergelyeen Dzo (Ardyn Obo, Ergil Obo); 13, Khoer Dzan. **Middle Oligocene:** 14, Kur Sai and Min Sai (Tchelkar Nura); 15, Donguz Tau; 16, Kyzyl Kak; 17, Kusto; 18, Kusto Kysylkain; 19, Tatal Gol (Shand Gol, Loh, Tatshin Gol); 20, Sen Zhak (Ordos, China); 21. Shunkht. **Late Oligocene:** 22. Agispe.

cene of Kazakhstan and in various other Paleogene localities. This material was gathered by expeditions of the Paleontological Institute of the U.S.S.R. Academy of Sciences, and also by a joint Soviet-Mongolian paleontological expedition. The tentative studies made so far have shown that a small part of this material belongs to previously known taxa, whereas the greater part represents completely new forms.

In all, about 240 fossils of Paleogene birds have been found in Mongolia. These include the first Asian Paleocene birds known, found in deposits at Naran Bulak and Tsagan Khushu, as well as interesting discoveries made in the Eocene deposits at Khaichin Ula 2 and Kolobolchi, and a large collection from the Oligocene localities at Ergelyeen Dzo, Khoer Dzan, Tatsin Gol, Tatul Gol, and others. New material from the Oligocene sites at Kur Sai and Min Sai, Donguz Tau and Kyzyl Kak in Central Kazakhstan and several Oligocene localities in the Zaisan basin, East Kazakhstan, includes 80 fragments of avian bones, most of which can be identified to the species level.

**Acknowledgments.**—I want to thank many persons for their valuable assistance in preparing this survey and in gathering the fossil birds reported in this work: Dr. N. Shevyreva, Paleontological Institute of the U.S.S.R. Academy of Sciences (PIN), who provided me with avian material from Zaisan basin; V. Y. Reshetov, also of the Paleontological Institute, for his valuable contributions to the discovery of fossil birds in Naran Bulak, Tsagan Khushu, and Khaichin Ula 2; Dr. D. Dashzeveg, Geological Institute of the Mongolian Academy of Sciences, who provided me with avian remains from Tsagan Khushu and Khoer Dzan; Dr. Malcolm C. McKenna, American Museum of Natural History, for his help in obtaining casts of the original material of *Eogrus* and for editorial assistance; Dr. K. Yudin and Dr. B. Nekrasov, Zoological Institute of the U.S.S.R. Academy of Science, for permitting me to study a skeleton of *Strigops*;
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The Paleocene

Birds have not previously been known from the Paleocene of Asia. As a result of field work by the Soviet-Mongolian paleontological expedition, the first such avian fossils were obtained from Naran Bulak and Tsagan Khushu in South Gobi, in the western part of the Nemeget basin. The geology of these localities, which are situated about 6 km from one another, has been described by Novozhilov (1954), Gradziński, et al. (1968), and Shishkin (1975). The mammalian assemblage from these sites is generally referred to as the Naran Bulak fauna (Belaeva, et al., 1974). Although immediately after its discovery this fauna was believed to be early Eocene in age (Efremov, 1954; Novizhilov, 1954), it is presently placed in the late Paleocene (Gradziński, et al., 1968; Szalay and McKenna, 1971; Belaeva, et al., 1974). The Naran Bulak mammalian fauna consists of insectivores, multituberculates, lagomorphs, condylarths, dinoceratids, pantodontids, and notoungulates. It has many species in common with the Gashato fauna collected by the Central Asiatic expedition of the American Museum of Natural History (Belaeva, et al., 1974).

In washing thousands of mammalian bones at Tsagan Khushu, only 12 avian bones were found. Half of these were coracoids and scapulae belonging to small anseriforms possessing specific structural details not found in Recent ducks. Also included were remains of a peculiar new type of heron similar to the Ardeidae, and shorebirds having definite affinities with the Scolopacidae.

The Eocene

Gureev (1949) described Eobalearica tugarinovi from a distal end of tibiotarsus from Andarak in the Fergana valley, a site which was then considered to be late Eocene. New stratigraphic correlations have shown it to belong to the early middle Eocene (Hekker, et al., 1962). Recently, bird remains have been reported from the Zaisan basin, where sediments bearing the so-called Obailin fauna are rather common. Various authors date this fauna as middle to late Eocene. Bendukidze (1971) described Progrus turanicus from a distal end of tibiotarsus from the Kalmakpai River, without giving an exact indication of locality. He placed this bird in the family Geranoididae, but this allocation would appear to require further scrutiny. P. A. Aubekerova of the Institute of Zoology of the Kazakh S.S.R. Academy of Sciences obtained unidentifiable fragmentary avian remains from the late Eocene localities near Andreevka on the Chinzhili River in Southeast Kazakhstan. Another isolated discovery was that of Protoplotus beauforti Lambrecht (1931), an upper Eocene anhinga from Sipang, in the western part of Sumatra, which was described from an almost complete skeletal impression.

Wetmore (1934) examined the paleornithological material collected by the Central Asiatic expedition at Ulan Shireh (upper Eocene of Irdin Manha) in China. He referred a coracoid to the Falconiformes and also described a femur as a new genus and species of Rallidae, Telecrex grangeri. Recently, Olson (1974) has provided convincing evidence that Telecrex grangeri is a representative
of the family Numididae, showing some similarities to the modern genus *Phasidus*. This is the earliest evidence of the presence of this group in Asia. From the same locality, Wetmore (1934) described numerous specimens of a crane as a new genus and species, *Eogrus aeola*, and reported specimens thought to be of the same species from another upper Eocene locality in Inner Mongolia in the vicinity of Iren Dabasu. Material referable to *Eogrus aeola* was obtained in 1959 by the Soviet-Chinese paleontological expedition at the same locality, Irdin Manha, in the Shara Murun region of China. This includes 10 distal ends of tibiotarsi and tarsometatarsi (PIN 2197-185–194), which confirm the original characterization of *Eogrus* and provide a solid basis for a comparative study of new specimens of Eogruidae and Ergilornithidae from the lower Oligocene.

Of interest are recent discoveries from Khaichin Ula 2 on the southern slope of Bugin Tsav in southern Gobi. In 1971–1973, 27 avian postcranial bones were found in these sediments, which are middle upper Eocene in age (Shuvalov, et al., 1974). Most of the bones are well preserved and can be studied in detail. Tentative identifications of this material have shown that the majority of the bones belong to shorebirds of two groups. The lesser part of these are clearly referable to the Scopacidae, whereas the remainder may be attributed to small birds of the suborder Charadriidea having no parallel among the recent families of this group and represented by species of different sizes. Coracoids and scapulae were especially numerous and a radius and tarsometatarsus were also found. From the same locality are remains belonging to the Ardeidae, Rallidae, and the suborder Lari. Diurnal birds of prey are represented by several rather large femora resembling the recent Aegypiinae but differing from *Gyps*, *Aegypius*, and *Gypaetus* in a number of important characteristics. A tarsometatarsus of a smaller species of Accipitridae was also found. Ducks, swans, rails, and herons from this locality are fairly comparable to recent families, although a fragment of skull belonging to a large representative of the suborder Lari has no exact parallels among the Recent families of that group.

Most of the avian bones from Khoer Dzan belong to the two gruiform families Eogruidae and Ergilornithidae. Remains of both groups were also found at Ergelyeen Dzo, and the more diagnostic material of the former confirms Wetmore's (1934) assertion that *Eogrus* may be traced to deposits of lower Oligocene age. These two families are discussed in greater detail in a following section.

The Early Oligocene

Information on the early Oligocene of Asia, especially of birds, is, in general, extremely scanty (Flerov, et al., 1974). *Cygnavus formosus* was described from the Aksirskaya site in East Kazakhstan (Kurochkin, 1968b). Kozlova (1960) described two peculiar, almost didactyloous species of Gruiformes, *Proergilornis minor* and *Ergilornis rapidus*, from Ergelyeen Dzo in Mongolia. Wetmore (1934) referred to a specimen from the same locality as "Eogrus sp." In 1970–1971, large scale paleontological field work was conducted at Ergelyeen Dzo (Ardyn Obo, Ergil Obo) and in other lower Oligocene localities in East Gobi, Mongolia, by Mongolian and Soviet paleontologists. Numerous avian remains were found, especially at Khoer Dzan, situated 70 km northwest of the railway station at Zamin Ud, at a locality that we called Shuvalov's Hills, after a student from Saratov State University who first discovered them in 1971. In all, about 200 avian bones were found at Khoer Dzan, 150 of which were in good or satisfactory condition and can be studied in detail. Several avian bones were also found in the eastern part of the classical locality at Ergelyeen Dzo.

The fossil material from Khoer Dzan contains specimens referable to the Accipitridae, Anatidae (including Cygninae), Rallidae, Ardeidae, and the suborder Lari. Diurnal birds of prey are represented by several rather large femora resembling the recent Aegypiinae but differing from *Gyps*, *Aegypius*, and *Gypaetus* in a number of important characteristics. A tarsometatarsus of a smaller species of Accipitridae was also found. Ducks, swans, rails, and herons from this locality are fairly comparable to recent families, although a fragment of skull belonging to a large representative of the suborder Lari has no exact parallels among the Recent families of that group.

Most of the avian bones from Khoer Dzan belong to the two gruiform families Eogruidae and Ergilornithidae. Remains of both groups were also found at Ergelyeen Dzo, and the more diagnostic material of the former confirms Wetmore's (1934) assertion that *Eogrus* may be traced to deposits of lower Oligocene age. These two families are discussed in greater detail in a following section.

The Middle Oligocene

Information on birds from the middle Oligocene of Asia is relatively rich. In Central Kazakhstan avian remains were found in the classical Indricotherium localities in the vicinity of the solonchak Tchelkar Tengiz. *Agnopterus turgaiensis*
was described from this region by Tugarinov (1940), and Cygnopterus lambrechti, Somateria sp., Limicorallus saiensis, and Megagallinula havundinea were later described by the present author (Kurochkin, 1968b). Much new material was also collected in this region in 1968. Among the fossils from the sites at Kur Sai and Min Sai in Tchelkar Nura, north of Tchelkar Tengiz, were remains identified as belonging to the Pelecaniformes, Anatidae (including Cygninae), Accipitridae, Galliformes, Gruidae, Rallidae, and Otididae. Collections made at Donguz Tau, another locality situated on the northeastern part of Tchelkar Tengiz, included avian remains referable to the Gaviiformes, Pocipediformes, Ciconiiformes (including the Threskiornithidae), Anatidae (including members of the Cygninae and Anserinae), Rallidae, Otididae, and Charadriformes.

The fossil site at Kyzyl Kak, on the northern slope of the basin of the same name in Central Kazakhstan, south of the town of Dzhezkazgan, was discovered and excavated in the past few years. This locality was thought to be middle Oligocene in age (Klebanova, 1965), but the mammalian material obtained there in 1968 casts doubt on this dating and most probably indicates a greater age. Several scraps of avian bones were reported from Kyzyl Kak, and were determined as belonging to the Podicipediformes, Aquilavus sp., Gruidae, and Otididae.

In East Kazakhstan, 60 km south of Lake Zaisan, new avian material was obtained from Kusto and Kusto Kysylkain, the sediments of which were dated by comparison with the mammalian fauna found at the middle Oligocene site at Kustovskaya. Flamingos, ducks, swans, Cygnopterus sp. and Eogrus sp. were identified from here. In the past few years several different birds were also reported from Bulkair, Tchaibulak, Akzhar, and Bobrovaya Struya, located 90 km north of Lake Zaisan. All the material was collected by Dr. N. Shevyreva of the Paleontological Institute of the U.S.S.R. Academy of Sciences.

Middle Oligocene sediments are very common on the northern slope of the Lakes Valley at the foot of the highest peaks of the Mongolian Altai, Ikhe Bogdo and Baga Bogdo. The first of these localities, Hsanda Gol and Loh (sediments of the Hsanda Gol Formation), were discovered by the Central Asiatic expedition in the 1920s. Small birds of prey and rails were described from the avian material collected in Tatal Gol in 1947 by the Mongolian paleontological expedition of the U.S.S.R. Academy of Sciences (Kurochkin, 1968a; 1969). Two of these birds of prey belong to the new genera Gobihierax and Venerator. The third species was placed in the recent genus Buteo, the fossil record of which extends back to the middle Oligocene of North America. This material was not included in Mellett's survey of the fauna of the Hsanda Gol Formation (Mellett, 1968). New material from the middle Oligocene sediments of the Lakes Valley was obtained in 1972. The complete tarsometatarsus of a small owl, differing markedly from all recent forms of Strigidae, is the most interesting of these discoveries. In another region of Mongolia, an unidentifiable bird bone was recovered from the middle Oligocene site at Shunkht, in Middle Gobi Aimak, east of Manlai Somon.

The Late Oligocene

Birds in the late Oligocene deposits at Agispe, on the northwest shore of the Aral Sea, are represented so far only by Anas oligocaena and several other species of Anatidae (Tugarinov, 1940, and more recent unpublished data). A specimen from this locality was identified by the present author as being galliform.

Paleoecology and Paleozoogeography of Asian Paleogene Birds

We can make some judgment of the life and environment of fossil birds by analogy with Recent birds belonging to allied groups, for in many cases, such as with cormorants, ducks, or bustards, there is no reason to suspect that there would have been significant differences in the habits of the Paleogene forms. However, in studying Paleogene birds with no analogs among the recent fauna, we can only make certain assumptions from their osteology as to their way of life.

The birds from Naran Bulak and Tsagan Khu-shu provide but scanty evidence of the environment of this region in the Paleocene. The discovery here of Anseriformes, Charadrii, and Ardeidae tells us no more than that there were moist areas present. The distinctive morphology of these birds as compared to Recent forms, makes
Along with *Eoegrus*, the lower Oligocene localities at Ergelyeen Dzo and Khoer Dzan in southeastern Mongolia contain the remains of two gruiform birds, *Proergilornis minor* and *Ergilornis rapidus*. In the Paleogene ergilornithids, and in their Pliocene descendent, *Urmiornis*, the inner toe was vestigial or absent, and the birds had lost the ability to fly, indicating that they were adapted for running in open areas. In this respect their ecological niche was probably perfectly analogous to that of the modern African ostrich (*Struthio*) and of the Asian ostrich of the Pliocene (Kurochkin and Lungu, 1970). The presence of ergilornithids and *Eoegrus* in the faunas of Ergelyeen Dzo and Khoer Dzan is strong evidence that sparsely vegetated open spaces were present in this area in the early Oligocene.

The middle Oligocene avian material from Tatal Gol in central Mongolia does little to illuminate our knowledge of the ecological conditions that existed at that time. The three birds of prey (*Accipitridae*) found there could have lived either in forests or in open areas, since Recent species of raptors can live in extremely varied environments. Fossil remains of an owl are similarly uninformative, as this species might well have lived under any of several ecological conditions. The rail described from Tatal Gol appears to have been more terrestrially than aquatically adapted.

We can hypothesize with much more confidence about the habitat found in the middle Oligocene of Central Kazakhstan, near the present solonchak of Tchelkar Tengiz. The specimens from the sites at Tchelkar Nura and those from Donguz Tau may be treated together, as these sites are separated by a mere 30 km and no great difference in their past contemporaneous environments is likely. It is possible, however, that certain taphonomic factors could have affected the species composition of the birds collected from these two localities. Waterbirds are numerous and diverse at both sites, particularly various anatids and loons. In addition, birds intimately associated with shoals and with marshes overgrown with vegetation are also abundantly represented. These include various rails, grebes, ciconiiforms, pelecaniforms, flamingos, shorebirds, and probably cranes. Certain of these, such as the Limicolae and ciconiiforms, would have required shorelines as feeding sites, while others, such as loons, swans, and other anatids, indicate the former presence of large areas of open water. Although aquatic species constitute a large proportion of the birds found at Donguz Tau and Tchelkar Nura, this does not necessarily reflect the actual situation that existed, as waterbirds are more likely to have been preserved than terrestrial ones. Birds of open dry habitats are also represented here, viz. Galliformes and Otididae. There is much evidence to suggest that in the middle Oligocene in the area of the northern coast of the present Tchalkar Tengiz there were vast basins with stretches of open water, alternating with shoals and marshes and open drier savannas.

The birds of the Kyzyl Kak locality are not numerous but are distinctive in their ecological preferences. The presence of small forms of bustards and cranes suggests open country, perhaps with low dense vegetation, while grebes indicate the presence of some nearby water. The birds from the middle Oligocene deposits of East Kazakhstan at Kusto and Kyzylkain are almost all waterbirds, such as ducks, swans, and flamingos, which certainly indicate the existence of aquatic habitats.

The paleornithological data from the early Oligocene of East Gobi (Ergelyeen Dzo and Khoer Dzan), and the middle Oligocene of Central Kazakhstan (Donguz Tau, Tchelkar Nura, and Kyzyl Kak) and Mongolia in the Lakes Valley (Tatal Gol) confirms the environmental picture derived from an analysis of the mammalian fauna of the same age (Flerov, 1961). On the whole, this suggests a more arid environment for Mongolia as compared with Kazakhstan. This is evidenced not only by the predominance of predatory and crane-like birds in Mongolia, but also by the absence of Anseriformes, which are customarily encountered in localities of this type in other areas. A variety of aquatic and marsh birds prevails in the middle Oligocene of Kazakhstan, whereas terrestrial birds are almost lacking, the few bustard-like birds probably being associated with watering places.

It is appropriate here to consider the possible role of the Turgai area in the evolution of Paleartic waterbirds in the latter half of the Paleogene. Notwithstanding the general paucity of material, the discoveries made so far seem to suggest that in the late Paleogene a number of groups of waterfowl may have originated on the coasts of the Turgai strait and later spread from there to
other areas. Further studies are needed to confirm or refute this surmise.

For a number of reasons it is difficult to determine the relationships of the Paleogene birds of Kazakhstan and Mongolia to those species from other areas. In the first place, the material thus far described and studied in detail is limited. As can be seen from the preceding discussion, most of it was obtained very recently and so far has been studied but tentatively. The character of much of the paleornithological material also impedes comparisons, since fossil forms of the same systematic group from different geographical areas may be represented by different parts of the skeleton and cannot be compared directly. Generally, the Paleogene birds of Western Europe and North America are incomparably better known than those of Asia. We can cite here only a few examples of Asian Paleogene birds with apparent relatives from other areas.

_Eogrus_, from the upper Eocene and lower Oligocene of Asia, appears to bear some resemblance to the genus _Palaeogrus_ from the middle Eocene to early Miocene of Europe. _Cygnatus formosus_ from the lower Oligocene of eastern Kazakhstan is comparable to _C. senckenbergi_ Lambrecht from the lower Miocene of West Germany. The middle Oligocene forms of _Cygnopterus_ from central and eastern Kazakhstan are close to _Cygnopterus affinis_ (Van Beneden) from the middle Oligocene of Belgium. The genus _Agnopterus_, known from the middle Oligocene of Central Kazakhstan (Tugarinov, 1940) is represented in the upper Eocene of England and France by two other species. _Aquilavus_, represented by new material from the middle Oligocene of Kazakhstan, is widely represented by several species in Western Europe that range from upper Eocene to lower Miocene. An undescribed middle Oligocene loon from Central Kazakhstan may prove to be referable to the genus _Colymboides_, the two other species of which are known in the upper Eocene of England and the lower Miocene of France.

The Families Eogruidae and Ergilornithidae in Asia

Of special interest is the evolutionary history of the gruiform families Eogruidae and Ergilornithidae in the Tertiary of Asia. The collections

Figure 2.—Left tarsometatarsus (PIN 3109-125) of _Proergilornis minor_ Kozlova, 1960; lower Oligocene, Ergelyeem Dzo: anterior, medial, and distal views. (Reduced.)
from the early Oligocene site at Khoer Dzan are of particular importance in this regard.

A complete tarsometatarsus of *Proergilornis minor* Kozlova (1960) from Ergelyeen Dzo (Figure 2) was used for comparison with 15 fragments of tarsometatarsi of Eogruidae (PIN 3110-59, 3110-101–113, 3110-172) and 22 fragments of tarsometatarsi of Ergilornithidae (PIN 3110-54, 3110-55, 3110-70–76, 3110-88-100) from Khoer Dzan. These collections also include 9 distal ends of tibiotarsi of Eogruidae (PIN 3110-77–85) and 9 of Ergilornithidae (PIN 3110-64–67) (Figure 3). Unfortunately, no proximal ends of this bone were recovered.

All of the eogruid hindlimb elements from Khoer Dzan are readily attributable to the genus *Eogrus*, although they exhibit several characters distinguishing them from *Eogrus aeola* of the late Eocene of Inner Mongolia and *E. wetmorei* from the Miocene deposits of Tung Gur. It should be noted that the remains of *Eogrus* from Khoer Dzan are easily separable into two size groups. Quite arbitrarily, we identified several other bones (phalanges 1 and 2 of pedal digit III and a fragment of femur) as belonging to *Eogrus*.

The remains of both *Proergilornis minor* and *Ergilornis* sp. from Khoer Dzan also fall readily into two groups differing in size. As with *Eogrus*, these differences are detectable in the tibiotarsi, tarsometatarsi, and phalanges. In the absence of any structural differences between the corresponding bones of the two size groups, we have every reason to believe that there were significant sexual differences in size in the species of Ergilornithidae.

In examining numerous limb bones belonging to Ergilornithidae and Eogruidae we experienced considerable difficulty in distinguishing specimens of one family from those of the other. It seems desirable therefore to detail here the structural differences in the distal end of the tibiotarsus and in the tarsometatarsus that separate the two families (Table 1). In making our comparisons we used Kozlova's original specimens of *Proergilornis* and *Ergilornis* in the collection of the Paleontological Institute of the U.S.S.R. Academy of Sciences, casts of the type-material of *Eogrus aeola* Wetmore (1934) and *E. wetmorei* Brodkorb (1967), obtained from the American Museum of Natural History, and the referred material of *Eogrus aeola* from Irdin Manha, also in the collection of the Paleontological Institute.
Table 1.—Comparisons of the hindlimb of the *Eogruidae* and *Ergilornithidae*

<table>
<thead>
<tr>
<th></th>
<th><em>Eogruidae</em></th>
<th><em>Ergilornithidae</em></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DISTAL END OF TIBIOTARSUS</strong></td>
<td>Posterior edge of internal condyle on the same level with anterior edge</td>
<td>Posterior edge of internal condyle slopes distally</td>
</tr>
<tr>
<td></td>
<td>Inner tuberculum fixatum closer to outer edge</td>
<td>Inner tuberculum fixatum on mid-line</td>
</tr>
<tr>
<td></td>
<td>Anterior part of external condyle flattened laterally</td>
<td>Anterior part of external condyle with lateral swelling</td>
</tr>
<tr>
<td></td>
<td>Internal ligamental prominence small</td>
<td>Internal ligamental prominence large</td>
</tr>
<tr>
<td></td>
<td>Distal end narrow relative to shaft</td>
<td>Distal end wide relative to shaft</td>
</tr>
<tr>
<td><strong>PROXIMAL END OF TARSOMETATARSUS</strong></td>
<td>External cotyla round</td>
<td>External cotyla oval</td>
</tr>
<tr>
<td></td>
<td>Hypotarsus with broad plantar surface and three tendinal grooves</td>
<td>Hypotarsus with narrow plantar surface and no traces of tendinal grooves</td>
</tr>
<tr>
<td><strong>DISTAL END OF TARSOMETATARSUS</strong></td>
<td>Middle trochlea relatively large, with parallel lateral surfaces</td>
<td>Middle trochlea relatively small with lateral surfaces widening distally</td>
</tr>
<tr>
<td></td>
<td>Outer trochlea closer to middle trochlea</td>
<td>Outer trochlea set farther laterally</td>
</tr>
<tr>
<td></td>
<td>In posterior view, groove between inner and outer trochlea located more proximally in relation to groove between middle and outer trochlea</td>
<td>Groove between inner and middle trochlea on same level or ending farther distally in relation to groove between middle and outer trochlea</td>
</tr>
</tbody>
</table>

In the development of the plantar crest on the shaft of the tarsometatarsus, the specimens of *Eogrus* from Khoer Dzan are intermediate between *Eogrus aeola* and the Ergilornithidae, thus tending to bridge the differences between the two groups.

Apart from tibiotarsi and tarsometatarsi, the collections from Khoer Dzan include 28 pedal phalanges belonging to the Ergilornithidae, although it has not been possible to assign these phalanges to a particular genus. These phalanges are short and flat and are virtually identical to those found in association with a tarsometatarsus of the Pliocene genus *Urmiornis*. Figure 4 shows the basal phalanges of digit III as reconstructed from elements belonging to different individuals. The phalanges of digit IV are also present and are easily distinguishable by their shorter length, while certain others we have rather arbitrarily identified as those of digit II. It appears that digit I had been completely lost in the Ergilornithidae.

Because the ergilornithids were large and had massive hindlimbs with reduced toes and flattened phalanges well adapted for running, it was natural to suggest that they might have been flightless. This supposition was borne out when a proximal end of humerus (PIN 3110–60) of a bird in which the locomotor function of the wing had obviously been lost (Figure 5) was found at Khoer Dzan. This specimen represents the second fragment of...
its kind, the first, which was found a day earlier, being lost soon after its discovery. This humerus has been so modified through structural degeneration that there is little basis for comparing it with Recent Gruiformes. Nevertheless, we suggest that this fragment is referable to the Ergilornithidae. It is characterized by general dorsoventral flattening, a markedly reduced and distally sloping head, the complete absence of a deltoid crest and ligamental furrow, and a weakly developed bicipital surface.

We can compare this humeral fragment with the humerus of *Strigops habroptilus* (Psittacidae), a completely unrelated bird whose ability to fly has been completely lost. Parallelism in the process of reduction is quite evident in both. The humerus of *Strigops* is similarly characterized by the elongated and flattened proximal end and reduced head and ligamental groove. In the ergilornithid humerus from Khoer Dzan the process of reduction was much farther advanced, however.

The collection from Khoer Dzan also contains a phalanx 1 of digit I of the wing (PIN 3110-163) which is characterized by its swollen structure and its shortened and rounded distal apex. We suggest that it also belongs to the Ergilornithidae.

The question of the nature of the relationship between *Proergilornis* and *Ergilornis* naturally arises. The present author is inclined to share Kozlova's (1960) assertion that there is considerable reason to regard these two genera as independent. The complete tarsometatarsus from Ergelyeen Dzo (PIN 3109–125) belongs to *Proergilornis minor* (Figure 2). The tarsometatarsi from Khoer Dzan also belong to this species, the only exception being PIN 3110–55, which belongs to *Ergilornis rapidus*. The fossils were obtained from sediments belonging to two different levels formed at different periods of time, though their accumulation took place within the framework of one and the same sedimentary cycle of the early Oligocene. Presently these two levels are separated by a section 10–15 m thick. In Shuvalov's Hills, most of the fossils were collected from the lower level, which was composed of a white mass of oligomict sand and clay. Only the tarsometatarsi of *Proergilornis minor* were represented in this layer. The upper level, an ochrous mass of sands and gravels of polymict composition, contained only five avian bones: a tarsometatarsus of *Ergilornis rapidus*, a tarsometatarsus of *Proergilornis minor*, two phalanges referable to the Ergilornithidae and Eogruidae, and a gruiform cervical vertebra. No definite inferences could be made from this material. Nevertheless, it ought to be kept in mind that the numerous bones from the lower level represented a single genus only, whereas the few specimens from the upper level represent two genera.

The question of the possible ancestor of the Ergilornithidae now arises. Our comparative studies suggest that this ancestor may be found in the Eocene forms of the Eogruidae. The structure of the hindlimb elements of the early Oligocene representatives of the two groups exhibit a rather close resemblance, both in overall appearance and in details. If we take into consideration that in the Eocene the Eogruidae exhibit a tendency toward reduction of the inner toe (first noted by Wetmore, 1934), as well as toward development of a sagittal crest on the tarsometatarsus and a medial fixing nodule on the distal end of tibiotarsus, we may suggest that the Eocene Eogruidae were the original members of the lineage leading to the Ergilornithidae. At a later stage these two families were developing in parallel, as is evident from their mutual occurrence in the lower Oligocene of Khoer Dzan, as well as from the presence of *Eogrus wetmorei* in the Miocene at Tung Gur (Brodtkorb, 1967).

The appearance of the flightless didactylous Ergilornithidae in early Oligocene time correlates well with the general trends observed in the to-
ography, climate, and vertebrate fauna in the transition from the late Eocene to the early Oligocene (Flerov, et al., 1974). This period was characterized by a complete disappearance of contacts between the faunas of Asia and America, an intensive development of open landscapes, and the advent of various groups of Artiodactyla and Perissodactyla that later underwent further development on the vast arid open areas of Asia, Europe, and Africa. Under such circumstances the appearance of the flightless cursorial Ergilornithidae may be considered as a natural product of gruiform evolution.

The Neogene descendant of the Paleogene ergilornithids is Urmiornis, which was described for the first time from a lower Pliocene site at Maragha, Iran (Mecquenem, 1925). Later, remains of representatives of this genus were found in the U.S.S.R. in the southern part of the Ukraine and Moldavia (Burchak-Abramovich, 1951). Additional specimens have been reported from other localities in the Ukraine and Caucasus and also from middle Pliocene deposits (Bendukidze, 1972; Umanskaja, 1973). The collections of the Palaeontological Institute of the U.S.S.R. Academy of Science also contain representative material of Urmiornis from the middle Pliocene at Kalmakpai in the Zaisan basin, East Kazakhstan, and several fragments from the middle Pliocene locality at Khirgis Nur in the Great Lakes Valley, Mongolia. Evidently, Urmiornis was very common in both the lower and middle Pliocene, throughout the entire range of the Hipparion Fauna. Comparisons of the original material of the Ergilornithidae with Urmiornis leaves no doubt as to the affinity between these didactylous gruiform birds, as the structure of their limb bones is very similar. Brodkorb (1967) was quite correct in placing Urmiornis in the family Ergilornithidae. A direct phylogenetic link between the Oligocene ergilornithids and the Pliocene Urmiornis can no longer be doubted.

Thus, a distinct phylogenetic line of Asian gruiforms that survived at least up to the middle Pliocene may be traced through the Tertiary as follows: Eogrus (Late Eocene), Ergilornis and Proergilornis (early Oligocene), Urmiornis (lower and middle Pliocene).

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The World's Oldest Owl: A New Strigiform from the Paleocene of Southwestern Colorado

Pat Vickers Rich and David J. Bohaska

ABSTRACT

Among the fossils recovered from a small, mid-Paleocene fissure filling in southwestern Colorado is the oldest known owl, Ogygoptynx wetmorei, new genus and species. This form, represented by a single tarsometatarsus, does not clearly belong in any of the known families of Strigiformes and may represent a new higher category of owls that provides a link between the Strigidae and the Tytonidae.

Introduction

In 1916, Walter Granger, following up the exploratory work of J. W. Gidley, examined sediments around Tiffany in southwestern Colorado and located several fossil-bearing areas that produced a moderately diverse mammalian fauna (Simpson, 1935). This assemblage became important in the definition of a North American land mammal age, the Tiffanian (Wood, et al., 1941). Among the bones found by Granger was a small avian tarsometatarsus. The following paper describes this fossil and considers its relationships.

Acknowledgments.—Thanks are due Mr. Earl Manning, who first recognized the avian affinities of Ogygoptynx, to Dr. Malcolm C. McKenna, American Museum of Natural History (AMNH), for allowing us to borrow the specimen for study, and Ms. Charlotte Holton for arranging the loan. Comparative material of the living genus Phodilus was kindly furnished by Dr. Ned K. Johnson, Museum of Vertebrate Zoology, University of California, Berkeley. The photographs and chart were prepared by Lloyd Logan and Robert Suddartha. Sincere thanks are also due Dr. Don E. Russell, Institut de Paléontologie, Museum National d'Histoire Naturelle, Paris (PM) for lending the European Paleogene owls briefly surveyed in this paper and presently under more extensive study. The manuscript was read by Hildegarde Howard and typed by Carol Baldwin.

Order STRIGIFORMES

Family incertae sedis

Ogygoptynx, new genus

Type-Species.—Ogygoptynx wetmorei, new species.

Distribution.—Known only from the Tiffanian (mid-Paleocene) of Colorado.

Diagnosis.—See Table 1.

Characters differing from the Strigidae but similar to the Tytonidae (including the Phodilinae): tarsometatarsus lacking a supratendinal bridge on the proximal end (three strigids also lack this bridge; Ford, 1967); posterior metatarsal groove not separated from the proximal articular surface by a marked bony ledge; anterior metatarsal groove deep across the entire width of the shaft at the proximal end.

Characters differing from the Phodilinae and Strigidae but resembling the Tytoninae: tarsometatarsus elongate and slender; posterior metatarsal groove only slightly excavated, lacking a lateral wall near the proximal end.

Characters resembling the Phodilinae and inter-
### Table 1.—Comparison of the tarsometatarsus in *Ogygoptynx* with several families and subfamilies of owls

<table>
<thead>
<tr>
<th>Character</th>
<th>Protostrigidae</th>
<th>Strigidae</th>
<th>Tytoninae</th>
<th>Phodilinae, <em>Phodilus badius</em></th>
<th><em>Ogygoptynx wetmorei</em></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Proximal End</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intercotylar prominence</td>
<td>–</td>
<td>Set back from anterior margin of proximal end, located midway between anterior and posterior margins</td>
<td>Occupies most anterior part of proximal end; protrudes anterior to cotyla</td>
<td>Occupies most anterior part of proximal end; protrudes anterior to cotyla</td>
<td>Occupies anterior margin of proximal end; not protruding anterior to cotyla</td>
</tr>
<tr>
<td>Anteroposterior width across intercotylar area</td>
<td>–</td>
<td>Narrow</td>
<td>Deep</td>
<td>Deep</td>
<td>Narrow</td>
</tr>
<tr>
<td>Groove lateral to intercotylar prominence</td>
<td>–</td>
<td>Distinct</td>
<td>Indistinct</td>
<td>Indicated slightly</td>
<td>Indicated moderately, intermediate between Strigidae and Phodilinae</td>
</tr>
<tr>
<td>Supratendinal bridge</td>
<td>–</td>
<td>Ossified in all but 3 species (see Ford, 1967)</td>
<td>Not ossified</td>
<td>Not ossified</td>
<td>Not ossified</td>
</tr>
<tr>
<td>Width of calcaneal canal</td>
<td>–</td>
<td>Wide</td>
<td>Narrow</td>
<td>Narrow</td>
<td>Wide?</td>
</tr>
<tr>
<td>Orientation of external cotyla</td>
<td>–</td>
<td>Directed laterally</td>
<td>Directed posteriorly</td>
<td>Directed posteriorly</td>
<td>Directed laterally</td>
</tr>
<tr>
<td>Degree of excavation of posterior metatarsal groove</td>
<td>–</td>
<td>Highly excavated</td>
<td>Slightly excavated</td>
<td>Highly excavated</td>
<td>Slightly excavated</td>
</tr>
<tr>
<td>Degree of separation of posterior metatarsal groove and proximal articular surface</td>
<td>–</td>
<td>Bony ledge separating groove from intercotylar area</td>
<td>No bony ledge present</td>
<td>No bony ledge present</td>
<td>No bony ledge present</td>
</tr>
<tr>
<td>Depth of anterior metatarsal groove near proximal end of tarsometatarsus</td>
<td>–</td>
<td>Deep only at inner part of supratendinal bridge</td>
<td>Deep across full width of proximal end, undercuts cotylar area</td>
<td>Deep across full width of proximal end, undercuts cotylar area</td>
<td>Deep across full width of proximal end, undercuts cotylar area</td>
</tr>
<tr>
<td>Lateral wall of posterior metatarsal groove</td>
<td>–</td>
<td>Present</td>
<td>Absent</td>
<td>Present</td>
<td>Absent</td>
</tr>
<tr>
<td><strong>Distal End</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Relative distal extension of trochleae</td>
<td>Middle trochlea extends farther distal than inner</td>
<td>Middle trochlea extends farther distal than inner</td>
<td>Inner trochlea extends slightly more distal than middle</td>
<td>Middle trochlea extends slightly farther distal than inner</td>
<td>Inner trochlea extends much farther distal than middle</td>
</tr>
<tr>
<td>Groove on middle trochlea</td>
<td>Anterior part not noticeably grooved, blends smoothly into shaft</td>
<td>Anterior part not noticeably grooved</td>
<td>Anterior part not noticeably grooved, blends smoothly into shaft</td>
<td>Anterior part with shallow groove</td>
<td>Anterior part not noticeably grooved, blends smoothly into shaft</td>
</tr>
</tbody>
</table>
mediate between the Strigidae and Tytoninae: shallow groove slightly lateral to the intercotylar prominence.

Characters differing from the Tytonidae but resembling the Strigidae: intercotylar area shallow anteroposteriorly, due in part to the intercotylar prominence not protruding anterior to cotylar margins; distal margin of outer trochlea in lateral view flattened and only slightly rounded, not grooved; outer trochlea in lateral view only slightly curved.

Characters differing from the Protostrigidae: distal margin of outer trochlea in lateral view flattened and only slightly, rather than greatly, rounded.

Characters within the Strigiformes unique to *Ogygoptynx*: proximal end in lateral view shaped like a parallelogram with unequal angles, not rectangular; outer trochlea in distal view not smoothly rounded but slightly grooved laterally, not tapering to a point but broadened posteriorly; anterior portion of shaft just proximal to middle trochlea elevated farther anteriorly than the remainder of the shaft, but distal portion of shaft medial to this region markedly planar and not rounded; inner trochlea decidedly more elongate than middle trochlea.

**Etymology.**—From Greek, *Ogyges*, mythical king of Thebes, suggesting ancient or primeval, and *ptynx*, an owl.

**Ogygoptynx wetmorei**, new species

**Holotype.**—Nearly complete right tarsometatarsus, AMNH 2653, missing only a small part of the calcaneal ridge and possibly some of the midshaft.

**Type-Locality and Stratigraphic Position.**—Mason Pocket near old Mason schoolhouse, northern drainage slope of the San Juan River, 6.5 to 8 km north of Tiffany, Section 20, T33N, R6W, La Plata County, southwestern Colorado. Found in “a small pocket of gray shale imbedded in a stratum of mottled purplish and brownish clay,” which has been thought by some workers to be a fissure filling of some type. “Tiffany Beds,” Tiffanian, mid-Paleocene in age (Simpson, 1935).

**Diagnosis.**—As for the genus.

**Measurements.**—Maximum width of proximal end, 9.0 mm; maximum depth of external cotyla, 4.9; maximum depth of internal cotyla, 5.1; maximum length from intercotylar prominence to distal end of attachment of tibialis anticus, 13.8; maximum width of distal end, 9.4; maximum width of inner trochlea, 4.2; maximum width of middle trochlea, 3.7; depth of internal border of inner trochlea, 3.2; depth of external border of inner trochlea, 3.6; depth of internal border of middle trochlea, 3.5; depth of external border of middle
Figure 2.—Ogygoptynx wetmorei, new genus and species, stereo pairs of holotype right tarsometatarsus (AMNH 2653): a, proximal view; b, distal view. (X 3.)

trochlea, 3.9; depth of external border of outer trochlea, 4.0; overall length from intercotylar prominence to distal end of inner trochlea, at least 48.2.

Etymology.—Named in honor of Dr. Alexander Wetmore, who has done much to further the knowledge of fossil owls as well as other birds.

Comparison with Modern Strigiformes

Ogygoptynx, known only from a tarsometatarsus, clearly belongs in the order Strigiformes as indicated by the broadly and deeply excavated anterior metatarsal groove; the single, slender calcaneal ridge; the shallow, narrow middle trochlea relative to the inner and outer trochleae; and the inner trochlea extending distally nearly as far as, or farther than, the middle trochlea.

Within the order, Ogygoptynx appears to be intermediate between the Tytonidae (including the Phodilinae; Ford, 1967) and the Strigidae, since it possesses a mosaic of character states of each of these families (Table 1).

One modern owl, Phodilus (subfamily Phodilinae), presently allocated to the Tytonidae (Ford, 1967), is to some degree a strigid-tytonid mosaic, but unlike Ogygoptynx, it is clearly most closely related to the Tytonidae. Strigid features seen in Phodilus include the deep proximal excavation of the posterior metatarsal groove, development of a lateral wall on the metatarsal groove near the proximal end, the moderately arched distal end, and the overall short, stout configuration of the tarsometatarsus. In most of its characters, however, Phodilus resembles the Tytonidae (Table 1), and thus, at least as evidenced by the tarsometatarsus, the allocation of Phodilus to the Tytonidae appears to be correct.

Because the tarsometatarsus of Ogygoptynx has such a mingling of strigid and tytonid character states, it cannot clearly be referred to any of the higher categories of modern Strigiformes.

Comparison with Fossil Strigiformes

The mid-Paleocene Ogygoptynx is the earliest known owl (Figure 3) and it is certain that by the early Eocene a number of other strigiforms had made their appearance in North America. Protostrix and Eostrix, in the family Protostrigidae, are known from a number of localities in the American West (Wetmore, 1958; Brodkorb, 1971; Martin and Black, 1972; Rich, unpublished data). The distal end of the tarsometatarsus is known for both genera and thus they may be compared directly with Ogygoptynx. Unfortunately, the proximal end of the tarsometatarsus is not known in any of the protostrigids. The smallest genus, Eostrix, closely approximates Ogygoptynx in size, but along with Protostrix it differs in the relative distal extension of the inner and middle trochleae, in the shape of the outer trochlea, and in lacking those character states unique to Ogygoptynx (see "Diagnosis"). It is difficult to assess the value of many of these unique character states due to the partial crushing of the type of O. wetmorei, as well as to our incomplete knowledge of the tarsometatarsus in the Protostrigidae.

A number of other fossil owls, previously classified in the Strigidae (Brodkorb, 1971), show a mixture of strigid and tytonid character states suggesting that they may bear some relationship to Phodilus and Ogygoptynx. Brodkorb (1970) erected the genus Paratyto for Bubo arvernensis Milne-Edwards, from the lower Miocene of France, placing it in the "Phodilidae" on the basis of its
lacking an ossified supratendinal bridge on the tarsometatarsus (as in the Tytoninae and Phodi-linae) and on the stoutness of the tarsometatarsus (as in the Strigidae and Phodilinae). He did not mention, however, that three species of strigids also lack an ossified supratendinal bridge, nor did he discuss other characters useful in separating the Strigidae from the Tytonidae (including the Phodilinae). Thus, reexamination of Paratyto would be useful for determining whether it is a strigid or a phodiline.

At least four other European Tertiary owls (Figure 3), Necrobyas harpax, N. rossignoli, "Otus" henrici, and "Otus" sp. (PM 3120) (Milne-Edwards, 1892), are based entirely or in part on tarsometatarsi that are stout and also lack an ossified supratendinal bridge. The tarsometatarsus of Ogygoptynx is narrower than in these owls, with N. harpax exhibiting proportions most similar to those of the new Paleocene form. Another tytonid character seen in the four European owls and Ogygoptynx is the lack of a ledge proximal to the posterior metatarsal groove on the tarsometatarsus. The calcaneal groove is narrow in the four European owls, as in the tytonids, but is probably wide in Ogygoptynx, as in the strigids. In all of the above four species, the external cotyla points posteriorly, as in the Tytonidae, except in two specimens assigned to "Otus" henrici (PM 3117 and PM 3118) in which the cotyla points laterally as in the Strigidae and Ogygoptynx. The position of the intercotylar prominence in all four species is intermediate between that in the tytonids and strigids and very similar to that in Ogygoptynx. A fifth

<table>
<thead>
<tr>
<th>m.y.</th>
<th>Stage/Age</th>
<th>Epoch</th>
<th>NORTH AMERICA</th>
<th>EUROPE</th>
<th>m.y.</th>
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<tr>
<td>30</td>
<td>WHITNEYAN/ORELLAN</td>
<td>OLIGOCENE</td>
<td></td>
<td>Necrobyas Strigogyps</td>
<td></td>
</tr>
<tr>
<td>35</td>
<td>CHADRONIAN</td>
<td></td>
<td></td>
<td>&quot;Otus&quot; henrici</td>
<td>35</td>
</tr>
<tr>
<td>40</td>
<td>DUCHESNEAN</td>
<td></td>
<td></td>
<td>&quot;Otus&quot;, species &quot;Bubo&quot; incertis</td>
<td>40</td>
</tr>
<tr>
<td>45</td>
<td>UINTAN</td>
<td></td>
<td></td>
<td>Prostrix</td>
<td>45</td>
</tr>
<tr>
<td>50</td>
<td>BRIDGERIAN</td>
<td></td>
<td></td>
<td>Eostrix</td>
<td>50</td>
</tr>
<tr>
<td>55</td>
<td>WASATCHIAN</td>
<td></td>
<td></td>
<td>Ogygoptynx, new genus</td>
<td>55</td>
</tr>
<tr>
<td>60</td>
<td>CLARKFORKIAN</td>
<td></td>
<td></td>
<td></td>
<td>60</td>
</tr>
<tr>
<td>65</td>
<td>TIFFIAN</td>
<td></td>
<td></td>
<td></td>
<td>65</td>
</tr>
<tr>
<td></td>
<td>TORRJONIAN</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>PUERCAN/DRAGONIAN</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Figure 3.**—Temporal distribution of early Tertiary owls.
owl, *Necrobyas edwardsi* Gaillard (1908, 1939), from the Tertiary of France, also based on a tarsometatarsus, was unavailable for study.

Other possibly intermediate genera of owls include *Prosybris* Brodkorb (1970) and *Lechusa* Miller (1956). *Prosybris*, from the lower Miocene of France, is placed with the Tytonidae but has a somewhat stouter tarsometatarsus than *Tyto*; like *Ogygoptynx*, its external trochlea is elongate. *Lechusa stirtoni*, based on a coracoid from the Pliocene of San Diego, California, was classified by Miller (1956) as a tytonid, although both he and Wetmore (in Miller, 1956:620) noted several strigid characters in this form. Miller stated, however, that *Lechusa* was definitely not a phodiline. Direct comparison between *Lechusa* and *Ogygoptynx* is impossible because of the lack of corresponding elements.

An extensive review of the above-mentioned fossil forms and some other European fossil owls is needed before the interrelationships of these, as well as modern forms, can be interpreted. *Ogygoptynx*, the oldest known owl, does not clearly belong in any of the presently established families within the order Strigiformes (*Protostrigidae*, *Tytonidae*, *Strigidae*) and may well be distinct enough to rate separate familial rank. Until a thorough revision of the early Tertiary European owls is completed, however, we will refrain from making any such designation and only point out the mosaic nature of this early owl.

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Simpson, G. G.

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Neanis schucherti Restudied:
Another Eocene Piciform Bird

Alan Feduccia

ABSTRACT

Neanis schucherti, an avian fossil from the lower Eocene Green River Formation of Wyoming, was described in 1913 under the genus Hebe by Shufeldt as the earliest representative of the order Passeriformes. The name Neanis has since been substituted for Hebe for reasons of priority. The specimen is here restudied and is found to be of piciform, not passeriform, affinity. It is assigned to the extinct Eocene family Primobucconidae. Of the two other previously described lower Eocene Piciformes, Primobucco kistneri Feduccia 1973 is reassigned to Neanis, while P. mcgrewi Brodkorb 1970 is retained in Primobucco, which genus is recognized by its larger size.

Introduction

The oldest North American avian fossil referred to the order Passeriformes and the only one assigned to the Neotropical family Rhinocryptidae (tapaculos) is Neanis schucherti (Shufeldt, 1913) from the lower Eocene Green River Formation of Wyoming. This fossil was first described under the generic name Hebe. Brodkorb (1965) substituted Neanis for Shufeldt’s genus, which was preoccupied by Hebe Risso 1826 (Crustacea). The type of Neanis schucherti consists of a small slab and counterslab (YPM 1233) containing impressions of bone and feathers and some poorly preserved pieces of bone (Figures 1, 2). These were collected in 1874 by F. A. C. Richardson of the Powell Expedition. The original label reads: “Found five miles west of Green River City, Wyoming. In the fish cut of the R.R. . . . Associated with insects described by Scudder.” On the back of the label is the following inscription: “Compare Pteroptochidae; see Ibis, 1874, p. 191 (July), for sternum with 2 emarginations in sternum.” The citation is to Sclater’s (1874) paper on the tapaculos and it was no doubt the above inscription that led Shufeldt to place the fossil in the family Pteroptochidae (= Rhinocryptidae). He based this conclusion almost entirely on the presence of a four-notched sternum (i.e., with “2 emarginations” on each side), and on an alleged “large manubrium which is bifurcated anteriorly” (Shufeldt, 1913:647). I have recently had the opportunity to examine the type of Neanis schucherti, and I present here my conclusion that the affinities of Neanis are with the Piciformes rather than the Passeriformes.

Acknowledgments.—The loan of the type of Neanis schucherti was arranged through the kindness of Drs. Charles G. Sibley and John H. Ostrom of the Yale University Peabody Museum of Natural History (YPM). Dr. Paul O. McGrew of the University of Wyoming Department of Geology kindly lent the types of Primobucco mcgrewi and Primobucco kistneri. Skeletons of modern species were made available through the courtesy of Dr. Pierce Brodkorb (collection of Pierce Brodkorb), Dr. R. L. Storer (University of Michigan Museum of Zoology), and Dr. R. L. Zusi (National Museum of Natural History, Smithsonian Institution). I wish to thank Drs. Pierce Brodkorb, Larry D. Martin, and Helmut C. Mueller for their critical review of the manuscript. This work was supported by a grant from the University of North Carolina Research Council.
Description and Comparisons

Two facts render the assignment of *Neanis* to the Rhinocryptidae questionable or suspect. First, the characters used in the identification, namely the presence of a four-notched sternum and a bifurcate manubrium (= forked spina externa of the sternum), are unique neither to the Rhinocryptidae nor the Passeriformes. Second, in recent years several primitive piciform birds have been described from the lower Eocene Green River Formation of Wyoming, the same horizon and locality from which *Neanis* was recovered. Piciform birds typically have a four-notched sternum and a forked spina externa. The Green River species *Primobucco mcgregori* Brodkorb (1970) and *Primobucco kistneri* Feduccia (1973)* provide the earliest records of the order Piciformes. Among the structurally primitive piciform families, the Bucconidae has been used in the past to accommodate these lower Eocene zygodactyl birds, but they are now considered to merit their own family, the Primo-bucconidae, which also includes three middle Eocene genera (Feduccia and Martin, p. 101, herein).

The presence of a four-notched sternum is an unreliable taxonomic character even at the ordinal level. Both two- and four-notched sterna occur in many orders not related to the passerines and their allies (e.g., Giconiformes and Charadriiformes), and four-notched sterna are commonly found in certain coraciiform birds (e.g., rollers and kingfishers). All members of the Piciformes possess a four-notched sternum (Feduccia, 1972), and within the Passeriformes the four-notched sternum is found within the Formicariidae and Rhinocryptidae (Heimerdinger and Ames, 1967). I am able to confirm that *Neanis schucherti* does indeed possess a four-notched sternum although to determine this requires “very close and careful examination with a high-power lens” (Shufeldt, 1913:646). Shufeldt (1913:646) goes on to point out that “the sternum of [Neanis] differed in this particular from all typical existing passerines as they now occur in North America, at least north of Costa Rica.”

In birds, the spina externa of the sternum may be forked (as in most passerines) or a simple rod (as in most nonpasserines). Olson (1971) has examined the spina externa of the sternum and found it to be highly variable. It is typically forked (bifurcate) in the Passeriformes (exclusively so in the suborder Passeres) but within the “suboscines” an unforked spina externa occurs within the Eurylaimidae, Cotingidae, Philepittidae (Ames, 1971), and Dendrocolaptidae (*Xiphocolaptes promeropirhynchos*, personal observation). However, *Smithornis* of the Eurylaimidae has a forked spina externa and within the Cotingidae the character is intragenerically variable in *Procynias* (Olson, 1971). Olson (1971:509) also points out that, “the forked versus simple spina externa is variable in other orders as well. In the Piciformes either conformation may be found in the Picidae, Capitonidae, Bucconidae and Galbulidae. The character is also variable in the Coraciformes and Trogoniformes.” Thus, perhaps the only taxonomic use of the nature of the spina externa would be to exclude forms with the simple, unforked spina externa from the suborder Passeres. In any case, I am unable to confirm the presence of a forked spina externa in *Neanis schucherti*. I have found the structure Shufeldt evidently intended (Figure 1), but the area is so crushed that some imagination is necessary to envision it as a bifurcate spina externa.

The two slabs containing the type of *Neanis schucherti* are illustrated in Figures 1 and 2. Figure 1 depicts the slab containing the actual bones; Figure 2 shows the counterslab with bone and feather impressions. The bones present on the slab are as follows: sternum (left side: ventral aspect); both coracoids (left, 10.8 mm); furcula (12.1 mm from furcular process to scapular tuberosity); left scapula; left humerus (proximal portion as preserved, 13.3 mm; width of proximal end, 5.4 mm; least width of shaft, 1.8 mm); right ulna (19.1 mm); right radius (approximately 18.4 mm); bones of manus (very faint). Unfortunately, no useful ratios could be obtained. Furthermore, few useful osteological characters are exhibited in the bones exposed, except for the form of the proximal end of the humerus, which is preserved in palmar view. It is in the humerus that one finds characters that ally *Neanis* with the structurally primitive families of the Piciformes, such as the modern family Bucconidae. *Neanis* conforms with the characters

* Misspelled “kisterni” in two places in the original reference.
that Brodkorb (1970:13) used to place *Primobucco mcgrewi* in the Piciformes and Bucconidae: "(1) proximal end inflected, so that entire caput humeri is medial to inner border of shaft (head of humerus more nearly in line with shaft in other families of Piciformes); (2) shaft more curved than in other piciform families; (3) deltoid crest long, bent near its mid-length at an angle of about 150 degrees (deltoid crest nearly parallel with shaft in other piciform families)." The other characters outlined by Brodkorb are not clearly visible in *Neanis*; however, it is clear from the palmar view of the hu-
merus (Figure 1) that Neanis is not passerine, but piciform, and of living families is most similar to the Bucconidae, as are the two other species of Piciformes from the lower Eocene Green River Formation.

The length of the humerus of *Primobucco mcgrewi* is 26.7 mm, and that of *Primobucco kistneri* was accurately estimated at 18-19 mm, as the entire outline of the bone was extant. In outline, the humerus of *Neanis schucherti* is somewhat similar to that of *Primobucco kistneri*, but the bones of the latter are so crushed that the comparison is unsatisfactory; however, the two forms were of the same general size, which, as I stated of *Primobucco kistneri* (Feduccia, 1973:503), "would probably best approximate . . . some of the modern African barbets (Capitonidae) of the genus *Pogoniulus* (including *Viridobucco*), which are approximately 4-5 inches in total length." In the absence of the evidence to the contrary, it seems best for the present to regard *Neanis schucherti* and *Primobucco kistneri* as distinct species; however, because of their general similarity in size, I recommend that *P. kistneri* be included in the genus *Neanis*, which has priority over *Primobucco*. Because *Primobucco mcgrewi* is considerably larger than either *Neanis schucherti* or *Neanis kistneri*, I strongly recommend the retention of the genus *Primobucco* to represent the large lower Eocene piciform birds from the Green River Formation.
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Shufeldt, R. W.
The Eocene Zygodactyl Birds of North America
(Aves: Piciformes)

Alan Feduccia and Larry D. Martin

ABSTRACT

Recent discoveries of zygodactyl birds in the Eocene of Wyoming, along with reinterpretation of previously described taxa, show that these forms belong to an extinct family, affiliated with the Bucconidae, for which we here propose the name Primobucconidae. The genera Primobucco, Neanis, Uintornis, Botauroides, and a new genus, Eobucco, are assigned to this family, and three new species, Primobucco olsoni, Uintornis marionae, and Eobucco brodkorbi are described. Primobucconids appear to have been the dominant small perching birds of the Eocene of North America.

Introduction

Recent discoveries of piciform zygodactyl birds from the lower Eocene Green River Formation of Wyoming brought to light an entirely new element in the avifauna of the North American Tertiary. Brodkorb (1970a) described the first of these forms as a new genus and species of the Bucconidae, which is structurally the most primitive family of the Piciformes. The genera Primobucco, Neanis, Uintornis, Botauroides, and a new genus, Eobucco, are assigned to this family, and three new species, Primobucco olsoni, Uintornis marionae, and Eobucco brodkorbi are described. Primobucconids appear to have been the dominant small perching birds of the Eocene of North America.

...
ically that this assignment was erroneous, but left the question of the true affinities of the genus unsettled. Cracraft examined the type and agreed with Shufeldt that it was not from a woodpecker, suggesting instead that it belonged with the Cuculiformes (Cracraft and Morony, 1969:6). On Cracraft’s suggestion, Brodkorb (1970b; 1971) placed Uintornis in the Cuculidae. Our present studies of the type of Uintornis lucaris show that it is not a cuckulid and that Brodkorb’s (1970a) original assignment of it to the Bucconidae was more nearly correct. We found three other tarsometatarsi, also from middle Eocene deposits in North America, to be affiliated with Uintornis at the family level. One of these was originally described by Shufeldt (1915) as a new genus and species of heron, Botauroides parvus; the second represents an undescribed species of Uintornis; while the last represents a new genus.

Two species of piciform birds from the Miocene of Europe have been placed in an extinct family Zygodactylidae (Brodkorb, 1971), based on the genus Zygodactylus (Ballman, 1969a; 1969b). These forms are distinctly different from the above species and further study of them will be necessary in order to clarify their affinities with other groups of modern and Tertiary zygodactyl birds. Meanwhile, the morphology of the Eocene forms precludes their assignment to any known family of the Piciformes and the erection of a new family is therefore made necessary.

Acknowledgments.—We are greatly indebted to S. W. Shannon of the Geological Survey of Alabama for bringing to our attention the type of Primobucco olsoni and placing it at our disposal for study. P. O. McGrew of the University of Wyoming, Department of Geology, kindly lent the types of Primobucco mcgrewi and Neanis kistneri. Skeletons of modern species were made available through the courtesy of Pierce Brodkorb (University of Florida), R. W. Storer (University of Michigan Museum of Zoology), and R. L. Zusi (National Museum of Natural History, Smithsonian Institution). C. B. Schultz made the University of Nebraska specimen available. D. Adams and D. Bennett rendered the illustrations. Abbreviations are as follows: Geological Survey of Alabama Type Collection (GSATC), University of Kansas Museum of Natural History (KUVP), University of Nebraska State Museum (UNSM), University of Wyoming Geological Museum (UWGM), and Yale Peabody Museum (YPM).

Order PICIFORMES

Suborder GALBULAE

PRIMOBUCCONIDAE, new family

Included Genera.—Primobucco, Neanis, Uintornis, Botauroides, Eobucco.

Diagnosis.—Small perching birds with the following combination of characters: (1) humerus (Figure 1a) with shaft curved, the head inflected medially, and the deltoid crest low, slightly rounded and proximally located; (2) radius and ulna (Figure 1a) slender and elongate; (3) metacarpals II

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Figure 1.—a, Restoration of the right wing of Primobucco olsoni, palmar view. Palmar views of right humeri: b, Cheli-doptera tenebrosa (Bucconidae); c, Megalaima lineata (Capitonidae); d, Tauraco sp. (Musophagidae); e, Tapera naevia (Cuculidae).
and III (Figure 1a) nearly parallel to each other, with only a narrow intermetacarpal space; (4) phalanx I digit II of manus (Figure 1a) broader proximally than in most other Piciformes; (5) tarsometatarsus (Figure 2e, f) relatively short, broad, and flat; (6) hypotarsus with a square lateral block of bone and a low ridge leading distally from it (Figure 6), probably with only a single tendinal canal (not clearly determinable in any known specimen); (7) papilla for tibialis anticus on the extreme internal margin of the tarsometatarsus (Figure 2e, f); (8) tarsometatarsus with a single proximal foramen (Figure 6); (9) distal end of tarsometatarsus flared, with large intertrocLEAR spaces and trochleae lying in the same anterior-posterior plane (Figures 2e,f,k; 5; 6); (10) middle trochlea the most distal, with the inner and outer trochleae about subequal (Figures 2e,f; 5; 6); (11) facet for metatarsal I entirely medial (Figures 5, 6); (12) inner trochlea grooved distally and posteriorly, middle trochlea deeply grooved (Figures 5, 6); (13) outer trochlea inflected inwards, with-

Figure 2.—Proximal views of left tarsometatarsi: a, Notarchus macrorhynchos (Bucconidae); b, Megalaima lineata (Capitonidae); c, Tauraco sp. (Musophagidae); d, Piaya cayana (Cuculidae). Restoration of the distal end of the left tibiotarsus and tarsometatarsus: e, Primobucco olsoni. Anterior (f–j) and distal (k–o) views of left tarsometatarsi: f, k, Eobucco brodkorbi (Primobucco conidae); g, l, Notarchus macrorhynchos (Bucconidae); h, m, Megalaima lineata (Capitonidae); i, n, Tauraco sp. (Musophagidae); j, o, Piaya cayana (Cuculidae).
out a well-developed sehnenhalter, but with a distinct groove separating the posterior portion of the trochlea from the remainder.

Remarks.—Characters 1, 6, and 13 of the diagnosis eliminate all families of the Piciformes except the Bucconidae. In the other piciformes, the bill is long (except in indicatorids and capitonids), the shaft of the humerus tends to be straight with the deltoid crest parallel to it (Figure 1c), there are two or more hypotarsal canals arranged one behind the other (Figure 2b), and the sehnenhalter is well developed (Figure 2m). The fossils tend to be bucconid-like in characters 1, 2, 3, 5, 6, 7, 8, and 13. They differ from the Bucconidae, however, in having a shorter mandibular symphysis, and in characters 4, and 9 through 12. The bucconids and the other piciforms differ from the Primobucconidae in that phalanx 1 digit II of the manus is narrower, especially at its proximal end; the inner and middle trochleae of the tarsometatarsus are placed closer together and the outer trochlea is elevated; the anterior face of the middle trochlea is not deeply grooved; and the facet for the first metatarsal is posteriorly situated.

Because Uintornis was suggested as being culculeform (Cracraft and Morony, 1969), we have compared all members of the Primobucconidae with nine genera of Cuculidae and three genera of Musophagidae. The Primobucconidae differ from both of these families in all but character 10 in the above diagnosis. The musophagids and culcules differ from the Primobucconidae in that the ulna is short, strongly curved, and has large feather papillae; metacarpal III is strongly curved so that there is a large intermetacarpal space; the supratendinal bridge is much above the proximal margin of the condyles of the tibiotarsus; the tarsometatarsus is elongate (Figure 2i,j); the inner and middle trochleae are close to each other, and the outer trochlea is elevated and inflected inwards, with no separation of the posterior portion from the rest of the trochlea (Figure 2n,o); the facet for metatarsal I is located posteriorly and the inner trochlea is not prominently grooved. The anterior face of the tarsometatarsus is often deeply excavated in culculiforms and there are usually two proximal foramina. The culcules always have two large enclosed hypotarsal canals (Figure 2d), a feature that is certainly absent in the only primobucconid in which it can be checked. Also, musophagids have a very triangular and distally situated deltoid crest (Figure 1d).

Clearly the Primobucconidae belong in the Piciformes and are most closely related to the Bucconidae. We can find no substantive evidence for a relationship between Uintornis (or any of the other forms of Primobucconidae) and any group of the Cuculiformes.

The family Primobucconidae includes five genera. The species of two of these genera, Primobucco and Neanis, are preserved as crushed specimens on slabs and represent medium-sized and small forms, respectively, of lower Eocene age. Three genera, Uintornis, Botauroides, and Eobucco, are medium to large in size and are middle Eocene in age; all are represented by fairly well-preserved tarsometatarsi only. Obviously, distinguishing the lower from the middle Eocene forms is difficult because the material is not strictly comparable. Nevertheless, size differences and such characters of the tarsometatarsus as can be discerned in the lower Eocene genera will permit them to be distinguished from most, if not all, of the middle Eocene forms. For this reason, and because of the time element involved, we believe it is best to recognize five genera in the Primobucconidae.

Primobucco Brodkorb, 1970a

Type-Species.—Primobucco mcgregori Brodkorb, 1970a.

Included Species.—P. mcgregori, P. olsoni.

Amended Diagnosis.—Medium-sized primobucconids, larger than Neanis and probably smaller than Uintornis, Botauroides, or Eobucco.

Primobucco mcgregori Brodkorb, 1970a

Holotype.—Right wing, UWGM 3255.

Type-Locality and Horizon.—From fish quarries in SE ¼ of Sec 18, T21N, R117W, near Fossil, Lincoln County, Wyoming; lower beds of Green River Formation, lower Eocene (Late Wasatchian) (Brodkorb, 1970a).

Primobucco olsoni, new species

Figures 1a, 3, 4

Holotype.—Two slabs containing a nearly complete skeletal impression and counterimpression,
Geological Survey of Alabama Type Collection, GSATC 217 (Figures 3, 4).

Type-Locality and Horizon.—The "first bluff" north of US Highway 30 north, across from Nug¬get, Lincoln County, Wyoming; Green River For¬mation, lower Eocene. Collected by Mr. George Moravec.

Diagnosis.—Smaller than *Primobucco mcgrewi* but larger than *Neanis*. Wing more slender and humerus longer than in *Primobucco mcgrewi*. Middle trochlea of tarsometatarsus extending farther distally than the other trochleae.

Description.—Skeleton preserved on two slabs with numerous feather impressions; mandible broad with a short symphysis; humerus with a low, gently curved deltoid crest almost parallel to shaft (Figure 1a); head of humerus inflected medially; radius and ulna slender, straight, and elongate, the ulna with no evidence of feather papil¬lae; carpometacarpus long and slender with a large, straight process for metacarpal I and a narrow intermetacarpal space; phalanx I digit II of the manus broad (narrower proximally in most bucconids); tibiotarsus short and robust with the supratendinal bridge straight, lying just above the condyles; tarsometatarsus broad, short, and flat with a high intercotylar prominence, anterior face of shaft with a low medial ridge, a single medial

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**Figure 3.**—Holotype slab of *Primobucco olsoni*, new species (GSATC 217), viewed ventrally. (a.t. = anterior toes, co = coracoid, cm = carpometacarpus, d = digit II, h = humerus, m = mandibular ramus, p.t. = posterior toes, r.u. = radius and ulna, tm = tarsometatarsus, tt = tibiotarsus)
proximal foramen, the papilla for tibialis anticus along the medial edge, the facet for metatarsal I situated medially, the middle trochlea situated distally, and the trochleae deeply grooved. Compared to *Primobucco mcgrewi* the humerus of *P. olsoni* is longer (28.8 vs. 26.7 mm), the shaft more slender (least width c. 2.2 vs. 3.0 mm), the proximal width greater (c. 7.5 vs. 6.7 mm), the ulna shorter (c. 32.0 vs. 34.2 mm), and phalanx I digit II of manus shorter (c 6.7 vs. 7.0 mm). The approximate length of the coracoid is c. 15.0 mm and that of the mandibular ramus c. 34.5 mm. The estimated toe lengths measured (in mm) through the arc are: digit I, 11.0; digit II, 16.4; digit III, 18.8; and digit IV, 13.3. Equivalent estimates of the toe arcs for *Neanis kistneri* are 6.8, 7.4, 8.6, and 8.2, respectively. The chord of the left wing of *P. olsoni* is estimated at 92 mm; of the species of Bucconidae listed in Ridgway (1914), the Barred Puffbird, *Nystalus radiatus*, is the nearest in size, with the wing chord of males averaging about 92 mm.

**Etymology.**—The specific name is in honor of Storrs L. Olson for his contributions to avian paleontology.

**Remarks.**—No accurate measurements of the tibiotarsus were possible but by comparing the two sides we were able to estimate the total length of the tibiotarsus very roughly as 28.5 (± 3) mm. The tarsometatarsus measures c. 15.5 mm, giving a ratio of tarsometatarsus to tibiotarsus of approximately 0.54. This rules out an affinity with the
Picidae (average ratio of 8 species, 0.67) or the Cuculidae (average of 7 species, 0.69). The same ratio for Neanis kistneri is 0.56; the average for 6 species of Bucconidae, 0.58; 3 species of Galbulidae, 0.51; five species of Capitonidae, 0.58; and 2 species of Indicatoridae, 0.59 (Feduccia, 1973). These ratios indicate only that the fossil does not belong to either the Cuculidae or the Picidae, but is a “perching” piciform bird. In woodpeckers, the different ratio results from the tibiotarsus being proportionately reduced. Comparative measurements of bucconids are given in Brodkorb (1970a).

The holotype of Primobucco olsoni is especially important because it permits us to associate the skeleton of Primobucco with those primobucconids known only from the tarsometatarsus. Features in which Primobucco resembles Eobucco and Uintornis are the short, broad tarsometatarsus with a high intercotylar prominence, the single lateral proximal foramen, the small and very medially situated tubercle for the tibialis anticus, and the medially situated facet for metatarsal I.

Neanis Brodkorb, 1965

**Synonym.**—Hebe Shufeldt, 1913 (preoccupied).
**Type-Species.**—Hebe schucherti Shufeldt, 1913.
**Included Species.**—N. schucherti, N. kistneri.
**Amended Diagnosis.**—Wing relatively shorter than in Primobucco; tarsometatarsus with large hypotarsus.

**Remarks.**—Better preserved material is needed for a full diagnosis of the very small zygodactyl birds referred to Neanis.

*Neanis schucherti* (Shufeldt, 1913)

**Holotype.**—YPM 1233, partial skeleton on a slab and impression on counterslab.

**Type-Locality and Horizon.**—Fish cut of the railroad, 8 km west of Green River City, Wyoming; Green River Formation, lower Eocene.

*Neanis kistneri* (Feduccia, 1973)

**Holotype.**—UWGM 3196, partial skeleton on a slab.

**Type-Locality and Horizon.**—N ½, NW ¼, Sec 6, T23N, R104W, Sweetwater County, Wyoming; Tipton Tongue Member of the Green River Formation, lower Eocene (Wasatchian).

**Amended Diagnosis.**—Smaller than *N. schucherti* or any of the other primobucconids.

*Uintornis* Marsh, 1872

**Type-Species.**—*Uintornis lucaris* Marsh, 1872.
**Included Species.**—*U. lucaris, U. marionae.*

**Amended Diagnosis.**—Zygodactyl birds with the outer trochlea of the tarsometatarsus not as strongly rotated as in other primobucconids; middle trochlea situated distally; distal foramen located above the outer trochlea.

*Uintornis lucaris* Marsh, 1872

**Holotype.**—YPM 617, distal end of right tarsometatarsus (Figure 5f,g).
Type-Locality and horizon.—Henry’s Fork, Sweetwater County, Wyoming; Bridger Formation, middle Eocene.

Amended Diagnosis.—Outer trochlea of tarsometatarsus inflected inwards but with inner and outer trochlear ridges not widely separated; middle trochlea located much farther distally than the others and distal foramen located more proximad.

Description.—Shaft of tarsometatarsus broad and flat; inner extensor grooves deep; distal foramen more proximal than the outer trochlea; distal end expanded with wide intertrochlear notches; trochlea deeply grooved; inner trochlea small, smooth anteriorly and grooved posteriorly, with a small medially directed posterior flange; middle trochlea deeply grooved, with high, thick, trochlear ridges, the inner trochlear ridge distal to the outer and with a short, thick neck; outer trochlea large, at the same level as the inner trochlea and inflected medially; outer trochlea grooved, with the outer ridge produced posteriorly as a flange or incipient sehnenhalter; facet for metatarsal I medially situated. Measurements (in mm) of type: width of distal end, 4.77; width of shaft at distal foramen, 4.18; depth and width of inner trochlea, 1.83 and 1.34; depth and width of middle trochlea, 2.25 and 2.02.

Etymology.—Named for Marion A. Jenkinson, who has often assisted us in our work on fossil birds.

Remarks.—The size difference and various qualitative features of the specimen separate it from Uintornis lucaris. After Eobucco, described later in this paper, it is the largest known member of the Primobucconidae. Uintornis seems to be the least specialized genus of the family in terms of toe rotation.

Botauroides Shufeldt, 1915

Type-Species.—Botauroides parvus Shufeldt, 1915.

Included Species.—B. parvus.

Amended Diagnosis.—Botauroides differs from Uintornis in having a proportionately wider shaft, the notch for the facet of metatarsal I shallower and slightly more posterior, the inner trochlea at about the same level as the middle trochlea, and outer trochlea not as elevated. The inner ridge of the outer trochlea is slightly more rotated and does not project as far anteriorly as it does in Uintornis. The outer trochlea is proportionately smaller and the medial ridge of the middle trochlea swings farther medially. Uintornis has a shallow groove just proximal to the outer trochlea, and the inner trochlea is relatively smaller than in Botauroides.

Uintornis marionae, new species

Figure 5a–d

Holotype.—KUVP 26906, distal end of right tarsometatarsus lacking outer trochlea (Figure 5a–d).

Type-Locality and horizon.—Sage Creek, Sweetwater County, Wyoming; Bridger Formation, middle Eocene.

Diagnosis.—Resembles Uintornis and differs from Botauroides in having a more slender shaft, a small inner trochlea, and a distinct groove proximal to the outer trochlea. Differs from Uintornis lucaris in being about 25 percent larger, and in having the facet for metatarsal I less deeply impressed and the outer ridge of the outer trochlea more medially inclined and elongated.

Description.—Tarsometatarsus broad and flat; distal foramen lying proximal to outer trochlea in anterior view and not visible in posterior view; facet for metatarsal I situated laterally; inner trochlea grooved posteriorly and smooth anteriorly; intertrochlear spaces wide; middle trochlea deeply grooved and situated distally; trochleae not arched. Measurements (in mm) of type: width of shaft at distal foramen, 4.18; depth and width of inner trochlea, 1.83 and 1.34; depth and width of middle trochlea, 2.25 and 2.02.

Botauroides parvus Shufeldt, 1915

Holotype.—YPM 1030, distal end of left tarso-metatarsus (Figure 5e).

Type-Locality and Horizon.—Spanish John Meadow, Wyoming; Bridger Formation, middle Eocene.

Diagnosis.—As for the genus.

Description.—Shaft very broad and flat, not expanded distally; distal foramen situated far proximally; inner trochlea larger and grooved posteriorly; inner and middle trochleae at same level;
outer trochlea relatively small, rotated medially and not very elevated; trochleae not arched.

Remarks.—Cracraft (pers. comm.) directed our attention to the similarities between Botauroides and Uintornis. Shufeldt (1915), with his uncanny ability to err, had referred it to the Ardeidae, where it appears in Brodkorb’s (1963) catalog.

**Eobucco, new genus**

**Type-Species.**—*Eobucco brodkorbi*, new species.

**Diagnosis.**—Largest known primobucconid; resembles *Uintornis* and differs from *Botauroides* in having the facet for metatarsal I deeply impressed and the middle trochlea located farther distally. Differs from *Uintornis* in having the ridges of the middle trochlea not extending as far proximally on the anterior side; in having the outer ridge of the outer trochlea inclined more medially and elongated until it extends past the outer ridge of the middle trochlea (it does not reach this trochlea in *Uintornis*); in having the outer ridge of the inner trochlea extending posteriorly as a distinct flange; and in the far distal position of the distal foramen.

**Eobucco brodkorbi, new species**

**Figure 6**

**Holotype.**—UNSM 20046 (Figure 6), left tarsometatarsus.

**Type-Locality and Horizon.**—56 km north of Green River, Sweetwater County, Wyoming; Bridger Formation, middle Eocene.

**Diagnosis.**—As for the genus.

**Description.**—Shaft of tarsometatarsus short, broad, and flattened; intercotylar prominence high and large; hypotarsus damaged but with a large lateral square of bone, which may have included a closed canal (the two closed canals, such as found in cuculids, could not have been present); a shallow groove rather than a ridge leads distally from the hypotarsus; anterior face of shaft grooved, with a high lateral ridge present; medial edge of shaft thin; a single large proximal foramen present near the midline; tubercle for tibialis anticus very small and distally situated along the medial edge of the shaft; facet for metatarsal I high and deeply impressed into the medial side of the shaft; distal foramen small, level with the outer trochlea; trochleae widely spread, not arched; inner trochlea relatively large, anterior face smooth, posterior face grooved; proximal part of the outer rim of the outer trochlea produced into a large posterior flange; middle trochlea large with high trochlear ridges diverging posteriorly; middle trochlea more distal than other trochleae; large outer trochlea at the same level as the inner trochlea; outer trochlea rotated medially with its elongate outer ridge extending medially past the outer rim of the middle trochlea; outer trochlea grooved. Measurements (in mm) of type: total length, 26.95; width of proximal end, 6.75; width of distal end, 6.00; depth and width of the inner trochlea, 1.36 and 1.27; depth and width of middle trochlea, 2.00 and 1.90; depth and width of outer trochlea, 2.90 and 1.80.

**Etymology.**—Named for Pierce Brodkorb, who...
was the first to recognize the true affinity of the primobucconids.

Remarks.—This is the largest and most advanced of the known primobucconids. It shares with Uintornis the distal position of the middle trochlea but has the outer trochlea much larger and more medially rotated. The single proximal foramen is remarkable for its large size and central position, thereby resembling the proximal foramen in Galbula. The medial side of the shaft is deeply indented under the internal cotyla where it is reduced to a thin blade. The tubercle for the tibialis anticus is extremely reduced. Eobucco possesses a combination of primitive and specialized characters that probably preclude its having given rise to any modern group of piciforms.

Conclusion

The allocation of Neanis schucherti, Uintornis lucaris, and Botauroides parvus to the Piciformes, and the description of the new forms Primobucco olsoni, Uintornis marionae, and Eobucco brodkorbi, brings the total number of species of North American Eocene zygodactyl birds to eight: two very small forms, Neanis schucherti and Neanis kistneri; two medium-size forms, Primobucco mekgrewi and Primobucco olsoni; two large forms, Eobucco brodkorbi and Uintornis marionae, and two others, Uintornis lucaris and Botauroides parvus, slightly smaller than the last two.

The order Piciformes probably arose in the New World and its forms occupied the “perching” arboreal adaptive zone in the early Tertiary of North America when tropical and subtropical climates predominated. Later, the order spread to the Old World where the Miocene Zygodactylidae and the modern families Picidae, Capitonidae, and Indicatoridae are represented, the last named being the only modern piciform family not found in the New World. Probably through climatic change and competition with more advanced land birds, e.g., passerines, the piciforms retreated mainly to tropical zones of the New and Old Worlds. The most primitive living piciform families, the Bucconidae and Galbulidae, are presently confined to the New World tropics. The Bucconidae are structurally closest to the Eocene family Primobucconidae.

The preponderance of evidence now indicates that the typical “perching” birds of the early Tertiary of North America were primitive piciforms, rather than passerine birds. Thus, it was probably not until the mid-Tertiary that the passerines took over in North America as the predominant “perching” group.

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Oligocene Fossils Bearing on the Origins of the Todidae and the Momotidae (Aves: Coraciiformes)

Storrs L. Olson

ABSTRACT

A new genus and species of tody, Palaeotodus emryi, is described from the “middle” Oligocene (Orellan land mammal stage) of Wyoming, providing the first record of the modern family Todidae outside the West Indies. The fossil bird Protornis glarniensis from the lower Oligocene of Switzerland is removed from the Alcedinidae to the Momotidae to provide the first occurrence of the latter family outside the New World. The Todidae and Momotidae appear to be more closely related to each other than either is to any other family of Coraciiformes. The Momotidae were evidently derived from the Old World. The Todidae appear to have been derived from a momotid-like ancestor in the Oligocene or earlier. The present distribution of these two families in the New World tropics is relictual. The Coraciiformes appear to have been one of the prevalent groups of small land birds in the Oligocene.

Introduction

The five modern species of todies (Todidae), endemic to the Greater Antilles, are among the most intriguing birds of the West Indies. The Momotidae of Central and South America and the Todidae are the only families of Coraciiformes confined to the New World. Apart from late Pleistocene remains of modern species, there has hitherto been no fossil record of either family.

Now, a recently discovered fossil and a new interpretation of one of the first fossil birds to be described, provide us with increased information bearing on the evolution and geographic origins of both the Todidae and the Momotidae.

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An Oligocene Tody from Wyoming

In June of 1972, Dr. Robert J. Emry of the Smithsonian Institution collected several blocks of matrix containing great concentrations of bones of small vertebrates from a deposit of Orellan age (“middle” Oligocene) in east-central Wyoming. Present in these samples are the abundant remains of at least two species of squirrels, various smaller rodents, small marsupials, and insectivores. That this great concentration of bone may be attributed to the work of owls is virtually certain since the blocks also contain the beautifully preserved skeletons of at least four small owls, possibly of two species.
On the surface of one of these blocks, Dr. Emry noticed the bill of a small bird and later found a crushed avian cranium that fit perfectly with the bill. After the two portions had been reunited, an incomplete avian humerus was found attached to the lower surface of the skull. There is no reason to believe that the humerus and skull are not from the same individual. Were they not, the owl that cast the pellet containing these remains would have had to eat two different birds at the same time, since it is rare for elements from a single prey item to be found in two successive pellets (Raczyński and Ruprecht, 1974). This would seem unlikely, particularly in view of the absence of birds other than owls in the remainder of the samples.

The distinctively shaped, flattened bill of the fossil was immediately reminiscent of the Todidae, but since bills of similar shape have evolved independently in several groups of birds, many of them passerines, identification was made cautiously. After careful comparisons, I concluded that this specimen is indeed referable to the family Todidae.

**Palaeotodus, new genus**

**Type.** *Palaeotodus emryi*, new species.

**Diagnosis.**—Similar to modern *Todus* but with different proportions, the wing apparently being better developed. Bill proportionately shorter and broader, not as pointed as in *Todus*. Mandibular rami not as flattened, the anterior portions grooved, so as to form a distinct dorsal shelf. Three ridges on the ventral side of the interorbital bridge separate, rather than coalesced as in *Todus*.

**Etymology.**—Greek, *palaeos*, ancient, plus *Todus*, the genus of modern todies. See Newton (1896:970, footnote) for the etymology of *Todus*.

*Palaeotodus emryi*, new species

**Figure 1**

**Holotype.**—Incomplete and partially crushed skull with most of the anterior portions of the rostrum intact, including the mandibular symphysis and parts of both rami; crushed posterior portion of cranium with ventral surface of interorbital bridge well preserved; much of the rest of the skull crushed, jumbled, and displaced ventroanteriorly. Vertebrate Paleontological Collections of the National Museum of Natural History, Smithsonian Institution, USNM 205608. Collected in NW 1/4, SE 1/4, Sec. 27, T32N, R71W, about 5.6 km SSE of Douglas, Converse County, Wyoming (42°42/55′N; 105°21′15/W) on 12 June 1972 by Robert J. Emry (Field No. WYO. 72-246) and Leroy Glenn.

**Horizon.**—Brule Formation, Orellan land-mammal stage, “middle” Oligocene.

**Paratype.**—Somewhat distorted right humerus with the shaft crushed and lacking the distal end; same number and data as the holotype.

**Measurements.**—Overall length of skull as preserved 34.5 mm, length of bill from anterior of nostril 10.0, length of mandibular symphysis 7.8, width of mandible at beginning of symphysis 5.6, width of mandibular ramus 1.7, proximal width of humerus 6.7.

**Etymology.**—After Dr. Robert J. Emry, the collector, in recognition of his significant contributions to our knowledge of the Oligocene fauna of North America.

**Description.**—Bill flat, broad, nearly spatulate, with a broader more rounded tip than in *Todus*. Internarial bar long, slender, terete, and somewhat heavier than in *Todus*, continued out the rostrum as a slightly elevated ridge. Mandible flattened, the symphysis shorter and broader than in *Todus*; mandibular rami deeper than in *Todus*, grooved anteriorly to form a dorsal shelf. Ventral surface of interorbital bridge with three ridges, the middle one terminating in a pointed process (tip broken off in the type), the outer ones flaring laterally to form the edges of the huge anterior cranial fenestra. The condition in *Todus* is essentially similar, but the three ridges are not as distinct anteriorly and coalesce to form a narrower, deeper interorbital bridge. The middle process in *Todus* is a...
point of attachment for a narrow ligament that joins with similar ligaments from the parapersphenoid rostrum and the lateral edges of the cranial fenestra to form a tenuous cross holding the anterior part of the brain in place. The humerus of *Palaeotodus* is similar to that of *Todus*, but is much larger; the internal tuberosity is heavier and much less lateromedially elongate; and the ligamental furrow and the scar on the external tuberosity are both considerably deeper.

**Remarks.**—Few nonpasserine landbirds have the bill anywhere near as flattened as in *Todus* and *Palaeotodus*. Those with the most flattened bills are *Electron*, *Hylomanes*, and *Aspatha* (Momotidae), *Jacamarops* (Galbulidae), and *Myioceyx* (Alcedinidae). In all of these forms the bill is deeper and the internarial bar broader and shorter than in either of the two genera of Todidae. The bony structure of the bill in passerines, as for example in *Todirostrum* (Tyrannidae), is quite different from that of *Todus* and *Palaeotodus*, being deeper and more arched, with a more trough-like mandible and a shorter, wider, and less terete internarial bar.

*Palaeotodus* agrees with *Todus* and differs conspicuously from the Momotidae, Alcedinidae, Galbulidae, and indeed from all of the rest of the Coraciiformes and Piciformes, in having the anterior wall of the cranium and the interorbital septum unossified. In the Momotidae, Alcedinidae, and Galbulidae the interorbital septum is partially or very heavily ossified. The anterior wall of the cranium is ossified in all of these families, whereas in *Todus* and *Palaeotodus* there is a great, open fenestra.

The humerus of *Palaeotodus* is broken and distorted, with the head crushed down distally relative to the internal and external tuberosities. Although in *Todus* there is slightly more of a depression beneath the head than in other Coraciiformes, the humerus can in no sense be regarded as having a double fossa, as stated by Bock (1962), and has a single pneumatic opening beneath the internal tuberosity. *Palaeotodus* is similar to *Todus* in this respect. The humerus of *Palaeotodus* differs from that of the Passeriformes in the less bulbous head, the much lesser development of the bicipital crest, the higher and more pronounced external tuberosity, and the narrower, more ventrally projecting internal tuberosity. In the Piciformes the deltoid crest is much more expanded, the head more bulbous, the capital groove much deeper, and the internal tuberosity less perpendicular to the shaft than in *Palaeotodus*, although these differences are considerably less pronounced in the Galbulidae and Buccoidea. Within the Coraciiformes the humeri of the Upupidae, Phoeniculidae, and Meropidae have larger, more triangular deltoid crests than in *Palaeotodus*, while in the Coraciidae the bicipital crest is more extensive. The humerus in the Alcedinidae has the head more bulbous and situated much higher above the external tuberosity, the shaft straighter, the internal tuberosity much heavier, and the bicipital surface much less produced than in *Palaeotodus*. The humerus of *Palaeotodus* is most similar to that of the Todidae and the Momotidae. In the conformation of the internal tuberosity it is more similar to the Momotidae, whereas in the greater excavation of the external tuberosity and ligamental furrow it more closely resembles the Todidae.

The manner in which the skull was crushed in the type of *Palaeotodus* makes it appear wider, while the distortion of the humerus is such as to make it appear wider and thus larger. Nevertheless, it is quite evident that the proportions of *Palaeotodus* are different from those of *Todus*, the wing being much larger in relation to the head.

This difference in proportions may be due at least in part to the small size of *Todus* being secondarily derived, since the species of this genus are the smallest members of the order Coraciiformes. In the evolution of vertebrates, body size usually changes at a more rapid rate than head size, so that small forms derived from larger ones tend to have proportionately larger heads, and vice versa. Many authors have remarked on the large-headed appearance of *Todus* in life. In the Oligocene, the Todidae were possibly more diverse than at present and probably included larger, more actively flying forms with better developed wings than the strictly sedentary modern todies.

**An Oligocene Motmot from Switzerland**

In 1889, von Meyer called attention to the remains of what he thought to be a passerine bird from slate deposits (Glarner Fischschiefer) in Switzerland, then considered to be of Cretaceous age. In a subsequent publication he named this specimen *Protornis glarniensis* (von Meyer, 1844).
Later (von Meyer, 1856), he emended the name to *P. glaronensis* and this spelling was in general use until Brodkorb (1971), whom I have followed, revived the original orthography. Lambrecht (1933) maintained Protornis as a genus incertae sedis in the Passeriformes. At that time the deposits from which the type of *P. glarniensis* was derived were regarded as upper Eocene in age. Subsequent studies have shown them to be of lower Oligocene age (Peyer, 1957).

The type of *P. glarniensis* consists of a slab containing the bones of all four limbs, the pectoral girdle, a complete mandible, the quadrate, and a few vertebrae and ribs. These were insufficiently exposed when von Meyer studied them, but Peyer (1957) undertook further preparation of the type, illustrating his results with numerous photographs and x-radiographs. The fossil is slightly distorted from stresses imposed on the rock after deposition; Stüssi (1958) and Baumann (1958) have offered mathematical and optical corrections, respectively, for this distortion. Another less complete specimen was referred to *Protornis*, possibly *glarniensis*, by Peyer (1957). I have had access to casts of both these specimens, as well as to Peyer’s excellent illustrations. The casts were made by a copper electrophate process and appear to be very accurate representations of the original specimens.

After his study of the type of *P. glarniensis*, Peyer (1957) concluded that it belonged with the Alcedinidae (kingfishers) and more particularly that it was nearest to *Dacelo*. I agree with the assignment of this form to the Coraciiformes, but numerous characters of its skeleton show conclusively that *Protornis* cannot be a kingfisher.

*Protornis glarniensis* is a small bird, slightly smaller than the modern motmot *Hylomanes momotula*. As detailed by Peyer, many aspects of its structure demonstrate that it does not belong with the Passeriformes and the clearly anisodactyl feet eliminate the Piciformes from consideration. The mandibular symphysis is broad, flattened and somewhat spatulate, differing from most non-passerine landbirds except the Momotidae and Todidae. The overall conformation of the mandible is in fact, markedly similar to that of the motmot genera *Electron* and *Hylomanes* and is quite distinct from that of the kingfishers, including the flat-billed genus *Myioceyx* (Figure 2). The symphysis is broader than in the Todidae and somewhat shorter than in the modern genera of Momotidae, being most similar in this respect to *Hylomanes*, which genus is generally conceded to be the most primitive of living motmots.

In the mandibular articulation of *Protornis* the internal process is a long, thin splint set off from a well-developed retroarticular process by a distinct notch, with the actual articulating surface for the quadrate much reduced. This is exactly the condition seen in the Momotidae and Todidae. In the Alcedinidae the articular cup for the quadrate is large and deep, the retroarticular process virtually absent, and the internal process wide, heavy, and triangular, quite unlike *Protornis* or the motmots and todies. Bee-eaters, Meropidae, have a fairly long, slender internal process, but it is not set off from the retroarticular by a notch, and the articular cup is deep, as in kingfishers. Furthermore, the bill shape of *Protornis* is not at all like that of the Meropidae.

The shape of the hypotarsus in *Protornis* is exactly as in motmots and differs from that of the kingfishers, in which it projects above the cotyae in a distinct point. The tarsometatarsus of *Protornis* is only slightly shorter than the middle toe with claw, as in the motmots. In todies the tarsus is longer than the middle toe with claw, whereas in the kingfishers and bee-eaters the tarsus is squat and much shorter than the middle toe. The procoracoidal process appears to be nearly absent in *Protornis*, as in motmots and todies, whereas it is better developed in kingfishers.

From the evidence detailed above it is clear that *Protornis* does not belong with the Alcedinidae, where Peyer (1957) placed it. The proportions of the bill and of the hindlimb and toes preclude its assignment to the Todidae. In all of its important features it agrees with the Momotidae. It differs from the modern forms of the family mainly in the shorter mandibular symphysis and the higher, more expanded sternocoracoidal process of the coracoid. *Protornis glarniensis* should, therefore, be assigned to the family Momotidae.

A second fossil from the Glarner Fischschiefer, consisting of a slab with both hindlimbs, the right wing, some ribs, and portions of the pelvis superimposed on the sternum, was assigned to the genus *Protornis* by Peyer (1957), who suggested that it might be referable to the species *P. glarniensis*. This is plainly impossible, for the second specimen...
is much too large to belong to the same species as *P. glarniensis*; the carpometacarpus, for example, is twice as long. Furthermore, in the right foot of this specimen it can be clearly seen that both digits I and II are reversed (Figure 3)—a condition found only in the Trogonidae among modern birds. The rest of the skeleton of this specimen is generally similar to that of modern trogons, although differing in some details. This specimen deserves a great deal more attention since it provides the
earliest evidence of the occurrence of the heterodactyl foot. It obviously cannot be assigned to Protornis or the Momotidae and for the present should be regarded as belonging to the Trogonidae. Four fossil species of trogons are known—all from France (Brodkorb, 1971). Three of these, in the genus Archaeotrogon, are from the Phosphorites du Quercy, which range in age from upper Eocene to lower Oligocene, and are thus possibly contemporaneous with the Swiss specimen. The fourth species, Paratrogon gallicus, is from lower Miocene (Aquitanian) deposits at Langy.

A second species of Protornis, P. blumeri, was described from the Glarner Fischschiefer in 1865 by Heer (1876). The type appears to have been poorly preserved and has not been restudied, its whereabouts being unknown. Brodkorb (1971) placed this species, along with P. glarniensis, in the Alcedinidae, but considered its position uncertain. From the original illustration one cannot even ascertain that the specimen was avian. In view of this, and since more than one family of birds occurs in the Glarner deposits, P. blumeri should be relegated to the category of Aves incertae sedis.

Discussion

The ten families of the order Coraciiformes fall into several diverse groups whose relationships within and without the order are as yet uncertain. Sibley and Ahlquist (1972:230) maintained that, “no compelling evidence exists to ally any group of the Coraciiformes more closely to a non-coraciiform than to other members of the Coraciiformes.” On the basis of biochemical analysis of egg-white proteins, Sibley and Ahlquist concluded, as have other taxonomists in the past, that within the Coraciiformes, the Alcedinidae, Todidae, Momotidae, and Meropidae appear to form a natural but distantly interrelated group. Feduccia (1975) discovered that these families possess a highly derived stapes, which is shared only with the Trogonidae, and concluded that all five families are closely related. Contrary to most earlier opinions, Sibley and Ahlquist (1972:230) suggested that the Todidae are more closely allied to the Alcedinidae than to the Momotidae. The osteology of these families does not support this contention, and along with their distributional history strongly indicates that a fairly close affinity exists between todies and motmots and that these families differ considerably from the kingfishers.

Seven of the ten families of Coraciiformes are confined to the Old World. Of approximately 89 Recent species of Alcedinidae, only six, in two genera, are found in the New World. Of these, two are in the genus Ceryle, which also contains two Old World species, while the genus Chloroceryle, which is only weakly differentiated from Ceryle, contains four species endemic to the New World. Clearly the kingfishers are an Old World family that has only recently invaded the Americas. Thus, the Todidae and Momotidae are the only truly New World families of modern Coraciiformes.

Recent motmots are neotropical in distribution, ranging from southern Mexico south through Brazil. In a classical exercise in zoogeography, Chapman (1923) analyzed the distribution of the genera, species, and subspecies of motmots, concluding that they had originated in Central America, the few South American forms having been derived from the north. Lönnberg (1927), noting that Central and North America probably presented a more or less continuous tropical environment in the Tertiary, felt that the motmots could as easily be con-
sidered North American in origin. A North or Central American origin of the modern members of the family, as opposed to a South American one, has properly gained general acceptance. Nevertheless, this leaves unanswered the question of the origins of ancestral motmots.

The modern todies, confined to the West Indies, are also thought of as being North American in origin. This is due in part to their presumed relationship with the motmots and in part to the North American derivation of most of the rest of the Antillean avifauna (Bond, 1966). It might then be asked whether the todies evolved their distinctive characteristics from some momotid-like ancestor after arriving in the Greater Antilles, or had already assumed these characteristics before colonizing the islands.

Because of the small size and feeble flight of modern todies, Bond (1966) expressed reservations about their ability to cross even narrow water gaps and suggested that they might have been rafted to the West Indies from Central America. However, because the five species of modern todies are very similar in plumage and morphology, one must assume that members of the genus Todus have been able to cross the water barriers between the Greater Antilles within relatively recent geologic time. Moreover, since there are two species of Todus on Hispaniola, one must either assume sympatric speciation or a double invasion of the island. Bond's own remarks (1974) on the greater similarity of the voice of Hispaniolan T. angustirostris to that of Cuban T. multicolor than to that of the other Hispaniolan species, T. subulatus, suggest a double invasion. Thus, if todies were able to cross the water barriers between the islands of the Greater Antilles they might as easily have crossed from the mainland. Furthermore, the evidence provided by Palaeotodus shows that in the Oligocene, todies were larger and had proportions suggestive of greater powers of flight. It therefore seems possible that todies might have colonized the West Indies over water as easily as, say, kingfishers, and it is not necessary to invoke rafting to explain their present distribution.

Without doubt, the order Coraciiformes, as traditionally conceived, arose in the Old World. The existence of Protornis in the lower Oligocene of Switzerland now provides evidence that the family Momotidae, presently confined to the New World, actually had its origins in the Old World. The place and time of origin of the Todidae are uncertain. The Orellan land-mammal stage represents a geologically very short span of time following the much longer Chadronian stage and preceding the Whitneyan stage, the latter also representing a very short span of time. The deposits from which Palaeotodus was recovered are about 30 million years old and have traditionally been regarded as middle Oligocene, although there is as yet no paleontological correlation between the North American terrestrial deposits of so-called Oligocene age and those of Europe. By the reduced ossification of the skull, Palaeotodus certainly seems to be referable to the Todidae rather than the Momotidae, but perhaps with material from earlier in the Oligocene it would not be possible to distinguish the two families, the family Todidae having assumed its characteristics since that time.

Probably through a combination of climatic change and competition with more advanced land birds, the motmot-tody group was entirely supplanted in the Old World. The deterioration of tropical conditions in North America in the late Tertiary left motmots only in Central America, from whence they have spread into South America since the closing of the Panamanian seaway in the late Pliocene. Similar factors affected the North American todies and only the isolated West Indian relicts of the genus Todus have survived up to the present.

Feduccia and Martin (p. 110, herein) have shown that the predominant order of small land birds of the Eocene in North America was the Piciformes. It is now becoming evident that the Oligocene was similarly important in the evolution of the Coraciiformes. Although the evidence is far from conclusive, if the Coraciiformes (including the Trogonidae) were not the predominant perching land birds of the Oligocene, they were certainly much more prevalent than today. Recently I have examined a number of fragments of small land birds of Chadronian and Orellan age from the western United States. All of these appear to be referable either to the Coraciiformes or Piciformes and definitely are not passerine. Thus, it would appear that the Passeriformes may not have gained a strong foothold in North America until the Miocene.
Baumann, E.

Bock, W. J.

Bond, J.

Brodkorb, P.

Chapman, F. M.

Feduccia, A.

Heer, O.

Lambrecht, K.

Lönnberg, E.

von Meyer, H.

Newton, A.

Peyer, B.

Raczynski, J., and A. L. Ruprecht

Sibley, C. G., and J. E. Ahlquist

Stiissi, F.
Two New Species of *Aegialornis* from France, with Comments on the Ordinal Affinities of the Aegialornithidae

Charles T. Collins

**ABSTRACT**

Collections from the upper Eocene–lower Oligocene phosphorite deposits of Quercy, France, include numerous fossil elements attributed to two species of *Aegialornis*. An examination of the humeri in this series disclosed the presence of two unrecognized species, which are newly described here as *Aegialornis wetmorei* and *A. broweri*. Preliminary study of the other skeletal elements previously assigned to *Aegialornis* indicates that at least some of them are probably referable to the Charadriiformes and the Coraciiformes. The humeri of *Aegialornis* show closer similarity to *Chordeiles* than to any members of the Hemiprocidae or Apodidae, and, therefore, the Aegialornithidae is removed from the Apodiformes and placed in the Caprimulgiformes near the Caprimulgidae.

**Introduction**

In the Museum National D'Histoire Naturelle, Paris, and the British Museum (Natural History), are extensive collections of bird fossils from the upper Eocene to lower Oligocene phosphorite deposits of Quercy, France. These include numerous distinctive humeri and some additional material referred to two species in the genus *Aegialornis*: *A. gallicus* Lydekker and *A. leenhardti* Gaillard. An additional form, *Primapus lacki*, was later described from the lower Eocene London Clay of Britain (Harrison and Walker, 1975). Further study of the Quercy material indicates the presence of two additional species of *Aegialornis*. This genus has been placed in a distinct family, the Aegialornithidae, the taxonomic history of which has been summarized by Harrison (1975). The family was first proposed by Lydekker (1891) who treated it as incertae sedis near the Laridae. The subsequent view of Milne-Edwards (1892) and Gaillard (1908) that *Aegialornis* is more properly included in the Apodiformes has been widely, though seemingly uncritically, accepted. Brodkorb (1971), on the basis of the evidence now presented here, included the Aegialornithidae in the Caprimulgiformes, a placement recently disputed by Harrison (1975). It is the purpose of this paper to review the species of *Aegialornis* and to comment on the possible affinities of the Aegialornithidae.

A wide array of fossil and recent material was examined in this study. Included were the types of *Aegialornis gallicus*, *A. leenhardti*, and *Tachyornis hirundo*, and much of the additional material referred to these species. Recent skeletons examined included many genera of Caprimulgiformes, particularly *Chordeiles*, *Caprimulgus*, and *Phalaenoptilus*, and from one to several species of swifts and crested swifts in the genera *Hemiprocne*, *Cypseloides*, *Streptoprocne*, *Apus*, *Aeronautes*, *Reinarda*, *Hirundapus*, and *Chaetura*.

Acknowledgments.—The study of these fossils, part of a wider study of the biology and evolution of swifts, has been generously supported by research grants and a postdoctoral fellowship from the Frank M. Chapman Memorial Fund of the American Museum of Natural History. Fossil material was obtained on loan from the Institut de Paléontologie, Museum National D'Histoire Naturelle, Paris (PM); the Department of Paleontology, British Museum (Natural History) (BMNH); and the Musée D'Histoire Naturelle de Montauban.
Recent comparative material of *Hemiproene* was obtained from the National Museum of Natural History, Smithsonian Institution. I am indebted to the curators of these institutions for their kindness. C. J. O. Harrison and C. A. Walker generously provided advance copies of their papers and photographs of *Primapus*. John Auth provided essential photographic assistance.

**AEGIALORNITHIDAE** Lydekker, 1891

*Aegialornis* Lydekker, 1891

**Synonyms.** — *Tachyornis* Milne-Edwards, 1892; *Belornis* Milne-Edwards, 1893.

*Aegialornis gallicus* Lydekker, 1891

*Figures 1, 4b*

**Type-Localities.** — *Aegialornis gallicus* near Lalbenque, France.


This species is represented by at least 20 humeri: the type-series of 13 and 2 additional specimens in the British Museum, and 5 specimens, including the type of *Tachyornis hirundo*, in the Paris Museum. Referred material includes 2 coracoids, 3 ulnae, 11 carpometacarpi, and 3 proximal phalanges of digit II (BMNH); and 26 tarsometatarsi (PM). As noted by Lydekker (1891) and Harrison and Walker (1975), the humerus is short and stout with a long, prominent, angular deltoid crest; deep ligamental furrow; large, flattened ectepicondylar process; laterally compressed head; broad bicipital surface and bicipital crest; and deep brachial depression. The humeri of *A. gallicus* are

![Figure 1](image-url)
smaller than those of *A. leenhardti* (Table 1), but larger than those of other species of *Aegialornis* or *Primapus*. The referred elements, not all of which appear to be properly assigned to *Aegialornis*, are discussed below.

**Aegialornis leenhardti** Gaillard, 1908

**Figures 2, 4a**

**Synonym.**—Originally proposed as *Aegialornis leenhardtii* Gaillard, 1908; spelling emended to *leenhardti* by Brodkorb, 1971:233.

**Type-Locality.**—Phosphorites du Quercy, France.

The holotype right humerus (Musee D'Histoire Naturelle de Montauban) and a previously unrecognized left humerus (PM) agree in being larger than *A. gallicus* and in having a larger and more distally located ectepicondylar process (Table 1). Additional material of this species (not examined in this study) is present in other museum collections (Gaillard, 1908; P. Ballmann, pers. comm.). A left tarsometatarsus from Caylux (Museum de Lyon) figured by Gaillard (1908) appears similar to the numerous tarsometatarsi he referred to *A. gallicus* and which were examined in this study. These specimens are, in my opinion, from an undescribed species possibly belonging in the Coraciiformes.
**Aegialornis wetmorei**, new species

**Figures 3a, 4c**

**Holotype.**—Complete right humerus (PM 15478) from the upper Eocene-lower Oligocene Phosphorites du Quercy, France.

**Measurements of Holotype.**—Overall length from head to internal condyle 21.8 mm, width and thickness of shaft at midpoint 2.8 × 2.1 mm, greatest width of distal end 5.0 mm, thickness through internal condyle 3.4 mm, height of proximal edge of ectepicondylar process above distal edge of ectepicondyle 6.3 mm.

**Paratypes.**—Two nearly complete left humeri (PM 15479 and 15480) slightly abraded on deltoid crest, bicipital crest, and internal tuberosity; from the same deposits as the type.

**Etymology.**—This species is named after Dr. Alexander Wetmore on the occasion of his ninetieth birthday, in recognition of his many contributions to the field of paleornithology.

**Diagnosis and Description.**—These humeri are appreciably smaller and more slender than the smallest humerus of *A. gallicus* (the type of *Tachyornis hirundo*) or that of the still larger species *A. leenhardti*. The ectepicondylar process is less robust than in *A. leenhardti* or *A. gallicus* and is located farther proximally, being well above the proximal edge of the brachial depression, whereas the ectepicondylar process is located at the level of the proximal end of the brachial depression in *A. gallicus* and *A. broweri* and is somewhat more distal in *A. leenhardti* (Table 1).

**Aegialornis broweri**, new species

**Figures 3b, 4d**

**Holotype.**—Nearly complete right humerus (PM 15481) from the upper Eocene-lower Oligocene Phosphorites du Quercy, France.

**Measurements of Holotype.**—Overall length from head to internal condyle 19.1 mm, width and thickness of shaft at midpoint 2.3 × 1.7 mm, greatest width of distal end 4.3 mm, thickness of distal end through external condyle 2.6 mm, thickness through internal condyle 2.7 mm, height of proximal edge of ectepicondylar process above distal edge of ectepicondyle 4.8 mm.

**Etymology.**—This species is named after Dr. Lincoln P. Brower in recognition of his contributions to other fields of biology and also for instilling in me a way of thinking I have tried to follow throughout my career.

**Diagnosis and Description.**—The single known humerus of *A. broweri* differs from *A. leenhardti*, *A. gallicus*, and *A. wetmorei* in being smaller, with a proportionately more slender shaft. The ectepicondylar process is more proximally located than in either *A. leenhardti* or *A. gallicus*, but is not as far proximal as in *A. wetmorei*. The brachial depression is less excavated and the muscle attachments of the proximal end are less well defined than in the other species of the genus. The type shows no signs of immaturity and must therefore pertain to an additional small species of *Aegialornis* in this fauna. The lower Eocene species *Primaapus lacki* is still smaller, the humerus being little more than two-thirds the length of that of *A. broweri*.

**Discussion**

It is perhaps surprising that there should be four
such closely related species of *Aegialornis* (Figure 4) in the same fauna. The differences in the position of the ectepicondylar process in these forms, however, make it unlikely that the apparent species limits are simply breaks in a continuum of one or two highly variable or sexually dimorphic species. *Primapus lacki* from the lower Eocene of Britain differs from the four species of *Aegialornis* in being much smaller and in having a slightly bilobed appearance to the bicipital crest and the entepicondyle projecting distally beyond the internal condyle (Harrison and Walker, 1975). The putative swift, *Cypselavus gallicus* Gaillard, from the upper Eocene-lower Oligocene Phosphorites du Quercy, was not examined in this study, but as noted elsewhere (p. 131, herein), it appears from the published illustrations that it may be a small member of the Aegialornithidae, about the same size as *Primapus lacki*. The earliest known modern swift (Apodidae) is *Cypseloides ignotus* (Milne-Edwards) from the lower Miocene (Aquitanian) of France.

The affinity of the Aegialornithidae to the Apodidae and Hemiprocnidae of the suborder Apodi, has been accepted, largely uncritically, since the early suggestions of Milne-Edwards (1892) and Gaillard (1908). This action has recently been endorsed by Harrison (1975) on the basis of a review of the humeri and other referred elements of *Aegialornis gallicus* and *Primapus lacki*. From my study of the referred material of *A. gallicus* I am convinced that the coracoids, the proximal phalanges of digit 2, and the tarsometatarsi belong to species in the orders Charadriiformes and Coraciiformes, and thus cannot be used to elucidate the ordinal affinities of *Aegialornis*. The similarity of some of these elements to those of the Laridae was noted by Lydekker (1891) in the original description of *Aegialornis*. Until it is possible to restudy all of the referred material, it seems wisest to confine discussion of the possible affinities of *Aegialornis* to characters of the humerus, the type-element in all the species of the Aegialornithidae.

The original allocation of *Aegialornis* to a fam-

![Figure 4](https://example.com/figure4.png)

**Figure 4**.—Size comparison of palmar views of right humeri of the four species of *Aegialornis*: *a*, *A. leenhardtii*, holotype; *b*, *A. gallicus*; *c*, *A. wetmorei*, new species, holotype; *d*, *A. broweri*, new species, holotype. (× 3.)
families. I feel within the Apodi seems to have been based principally on the superficial resemblance of the short and stout humeri to those of the Hemiprocnoiidae and Apodidae. Harrison (1975) also cites the prominent ectepicondylar process, shorter and more abruptly projecting deltoid crest, less proximally deflected internal tuberosity, and the presence of the distinct flange on the bicipital crest as characteristics shared with the Apodi. Although there are some definite similarities between the humeri of *Aegialornis* and those of the Apodi, particularly *Hemiprocone*, I feel there is a greater resemblance between *Aegialornis* and some members of the Caprimulgidae, particularly *Chordeiles* and related genera (the Chordeilidae of Oberholser, 1914).

In *Aegialornis* the head is deflected anconally and the distal end is directed palmarly. This condition, somewhat accentuated by the longer shaft, is also found in *Chordeiles*. The humerus is notably straight in *Hemiprocone* and the primitive swifts of the subfamily Cypseloidinae; only a slight anconal deflection of the head is present in the Apodinae and Chaeturinae. The deep ligamental furrow in *Aegialornis* and *Chordeiles* extends well out onto the internal tuberosity, where it curves to approach the distal margin. In the Apodi the ligamental furrow is shorter and straighter, ending near the proximal base of the internal tuberosity. In *Chordeiles* the internal tuberosity is deflected slightly more proximally at the tip than in *Aegialornis*, and the pneumatic fossa is more exposed. In the Apodi the internal tuberosity projects laterally or is deflected distally and bears little resemblance to that of *Aegialornis*. A distinct similarity exists between *Aegialornis* and *Chordeiles* in the thickened median crest and broadly excavated capital groove proximal to it. A slight projecting flange on the bicipital crest of *Aegialornis* can also be noted in some specimens of *Chordeiles*. Although this flange is usually much more highly developed in *Aegialornis*, considerable variation is shown in the specimens examined in this study, with some individuals showing only slight development of this feature. The shape of the deltoid crest is very similar in *Chordeiles* and *Aegialornis* and lacks the more abrupt taper and concave proximal edge of the Apodi. The deltoid crest is appreciably different in other genera of the Caprimulgidae (e.g., *Caprimulgus* and *Phalaenoptilus*), which have a more flattened lateral edge and a longer, more gradual slope to the distal edge. Thus, only some of the genera of Caprimulgidae have the “longer and more smoothly rounded” profile of the deltoid crest incorrectly attributed to the entire family by Harrison (1975).

The distal end of the humerus shows many similarities between *Aegialornis*, *Hemiprocone*, and to a lesser extent *Streptoprocne*, in the flared external tricipital groove and medially expanded entepicondyle. *Aegialornis* and *Chordeiles* are alike in having a broader and more protruding attachment for the anterior articular ligament and a deeper intercondylar groove. A large peglike ectepicondylar process, the single most distinctive characteristic of the humeri of swifts and *Aegialornis*, is accompanied by a distinct, raised lateral muscle scar at its base in *Aegialornis* which is lacking in the Apodi. A small ectepicondylar process and associated muscle scar is also present in *Chordeiles* and, as noted by Harrison (1975), in *Podager*. In the Apodi the ectepicondylar process is never as thickened as in *Aegialornis* and is always substantially more proximally located. In some of the Apodi there is also a secondary process located distal to the ectepicondylar process. This is particularly well developed in the Hemiprocnoiidae and Cypseloidinae but completely absent in *Aegialornis*. Within the Caprimulgidae there is considerable difference in the appearance of the distal portion of the humerus, as well as in the development of the ectepicondylar process. A strong resemblance to *Aegialornis* can be found in *Chordeiles* and related genera, but not in *Caprimulgus*, *Phalaenoptilus*, and *Eurostopodus*.

Although the Aegialornithidae show some similarities to the Hemiprocnoiidae, I feel that the majority of the characters of the humeri indicate a closer relationship with the *Chordeiles* group of the Caprimulgidae. I therefore place the Aegialornithidae as a family within the Caprimulgiformes, possibly allied to the Caprimulgidae.

With the tentative removal of *Cypselavanus galli¬cus* from the Apodidae to the Aegialornithidae, the earliest fossil swifts appear in the lower and middle Miocene deposits of France (p. 131, herein). There is thus no longer any evidence to support the earlier notion that the Apodidae and Aegialornithidae were contemporaneous during the late Eocene or early Oligocene. Therefore, the possi-
bility exists that the Aegialornithidae are representatives of a caprimulgiform lineage that later gave rise to the swifts and crested swifts. Although a close relationship between the Caprimulgiformes and the Apodi is not supported by presently available biochemical information (Sibley and Ahlquist, 1972), neither does this information provide any conclusive evidence of the affinities of swifts to other groups. A caprimulgiform-apodiform relationship should be reviewed further when additional fossil elements are found that can definitely be assigned to the Aegialornithidae.

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Sibley, C. G., and J. E. Ahlquist
A Review of the Lower Miocene Swifts
(Aves: Apodidae)

Charles T. Collins

ABSTRACT

Three nominal species of swifts have been described from lower Miocene (Aquitanian) deposits of France. Re-examination of these forms, Cypselus [≡ Apus] ignotus Milne-Edwards, Collocalia incerta Milne-Edwards, and Cypselavus intermedius Gaillard, indicates that they are attributable to a single species, ignotus, referable to the modern genus Cypseloides. This provides the first occurrence of the Cypseloidinae in the fossil record and indicates a possible origin in the Old World for this primitive group of swifts, presently restricted to the New World.

Introduction

Up to now, five species of fossil swifts have been described, all coming from Tertiary deposits in France. The present paper is aimed at reviewing the three nominal species from deposits of early Miocene age, with comparisons being made with a much wider array of skeletal material of modern swifts than were available to the original describers of the fossil forms. Recent swifts examined in this study included Cypseloides rutilus, C. cherriei, C. niger, and Streptoprocne zonaris in the Cypseloidinae; from one to several species in the genera Apus, Aeronautes, Cypsiurus, Tachornis, Reinarda, and Panyptila in the Apodinae; and Chaetura, Collocalia, and Hirundapus in the Chaeturinae.

Acknowledgments.—I thank the Frank M. Chapman Memorial Fund of the American Museum of Natural History for support and I am also indebted to Dr. J. P. Lehman of the Institut de Paléontologie, Museum National D'Histoire Naturelle, Paris, for lending the types of Cypselus ignotus and Collocalia incerta.

Family APODIDAE

Subfamily CYPSELOIDINAE

Cypseloides ignotus (Milne-Edwards, 1871)


From the Aquitanian deposits at St.-Gerand-le-Puy, Departement de Allier, France, Milne-Edwards (1871) described a new species of swift, Cypselus [≡ Apus] ignotus. This was based on a complete right carpometacarpus and a left ulna with the proximal end badly chipped. When I examined these specimens, a second left ulna, excellently preserved, had somehow been associated with the two syntypes. This is identical to the first ulna and I therefore refer it to the species ignotus also. From the same deposits, Milne-Edwards (1871) named a second species of swift, Collocalia incerta, based on a single well-preserved left tibiotarsus. This was characterized as being much too small to have come from the same species as the wing elements assigned to Apus ignotus. Considerably later, Gaillard (1939) reported a left humerus of a swift from Aquitanian deposits at Chavroches, also in the Departement de Allier, which he described as a new species, intermedius, in the
Eocene-Oligocene genus *Cypselavus* (Gaillard, 1908).

I have studied the original material of *Apus ignotus* and *Collocalia incerta* as well as Gaillard's (1939) description and illustrations of *Cypselavus intermedius*. While all these fossils clearly belong to the Apodidae, it is equally apparent that none is referable either to *Apus* or to *Collocalia*.

In size and conformation, the two ulnae of *ignotus* (Figure 1a) are very similar to those of some of the smaller species of the genus *Cypseloides*. They lack the well-developed olecran process found in the subfamilies Apodinae and Chaeturinae. This condition is typical of the Cypseloidinae. The fossil ulnae are slightly longer and stockier than the ulnae of *Cypseloides rutilus* or *C. cherriei*, and the prominence for the anterior articular ligament is more shelf-like. Also, the external cotyla appears slightly more undercut at its palmar edge and the distal ligamental attachment of the carpal tuberosity is less laterally flared.

The carpometacarpus of swifts shows less marked distinctions between the subfamilies than does the ulna. That of *ignotus* (Figure 1b) is somewhat longer and stockier than in *Cypseloides rutilus* or *C. cherriei*, but it does have the more widely flared pollical facet of metacarpal I and the more pronounced tendinal groove on metacarpal II typical of the Cypseloidinae. The fossil also has a broader and more flared tuberosity of metacarpal II, providing a wider articular facet for digit II, which is typical of the two smaller species of *Cypseloides* studied, but not of swifts of other subfamilies.

As noted by Milne-Edwards (1871) the tibiotarsus of *Collocalia incerta* (Figure 1c,d) is indeed more delicate than would be expected for any member of the Apodinae or Chaeturinae of the size of *Apus ignotus*. However, the tibiotarsus in the Cypseloidinae is proportionately more slender than in the other subfamilies of swifts, particularly the Apodinae. There is, in fact, a very close agreement in overall size and morphology between the type of *Collocalia incerta* and Recent specimens of *Cypseloides rutilus*. The posterior intercondylar groove of *incerta* is not deeply excavated as it is in members of the Apodinae and Chaeturinae, including *Apus* and *Collocalia*. The proximal portion of the shaft is straight, as in *Cypseloides*, and not distinctly bent laterally as typical of many other swifts. The fossil element is slightly smaller and stockier than in *C. rutilus* (*C. cherriei* has a much longer tibiotarsus than *C. rutilus* in spite of its having wing elements similar in size to both *C. rutilus* and *A. ignotus*), and the internal ligamental prominence is less developed but more excavated under the lip of the rotular crest.

The wing elements of *ignotus* are clearly those of a small swift belonging to the genus *Cypseloides*. The tibiotarsus of *incerta* similarly shows affinities to *Cypseloides* particularly to *C. rutilus*. Contrary to Milne-Edwards (1871), it is entirely probable that these fossils, which are from the same locality and horizon, come from the same species. This species should now be known as *Cypseloides ignotus* (Milne-Edwards) with *incerta* becoming a junior synonym, *ignotus* being chosen on the basis of line priority.

In the referred ulna of *Cypseloides ignotus*, the
maximum length is 17.9 mm, distal width 3.2 mm, proximal width 3.7 mm, and shaft width 1.6 mm. No accurate length could be determined for the chipped ulna in which the distal width is 3.2 mm, proximal width 3.7 mm, and shaft width 1.8 mm. The single carpometacarpus measures 16.4 mm in total length, proximal height 5.2 mm, proximal width 2.35 mm, and distal width 3.95 mm. The tibiotarsus has a total length of 21.1 mm, width across condyles 2.2 mm, width across proximal articular surfaces 2.5 mm, and shaft dimensions of 0.9 \times 1.0 \text{ mm} at the narrowest point and 1.0 \times 1.5 \text{ mm} at the middle of the fibular crest.

The type-humerus of *Cypselavus intermedius* Gaillard (1939) was not examined in this study, but from the original figures it appears to have the distinctively longer and narrow proportions characteristic of the species of *Cypseloides*. As was noted by Lowe (1939:324), the ectepicondylar process of *intermedius* is much more distally positioned than in any of the modern forms of the Apodinae or Chaeturinae but is only slightly more distal than in *Cypseloides*, a genus that was not compared by earlier workers. The measurements of the type of *C. intermedius* as given by Gaillard (1939:43) are: total length 11 mm, proximal width 4.5 mm, distal width 3 mm. Thus, this specimen agrees closely in size with specimens of modern *Cypseloides rutilus* and *C. cherriei*, and it would therefore also be of the same approximate size as *C. ignotus*. The type of *Cypselavus intermedius* comes from the same horizon and from a locality close to that of *Cypseloides ignotus*. Since it also appears to belong to the genus *Cypseloides* and is of the same size as *C. ignotus*, I feel that *Cypselavus intermedius* should also be synonymized with *Cypseloides* ignotus. As a result, the genus *Cypselavus* Gaillard is reduced to a single species, *C. gallicus*, from the upper Eocene or lower Oligocene (Phosphorites du Quercy) of France; the genus *Collocalia* is deleted from the fossil record; and the earliest fossil possibly attributable to *Apus* now becomes *Apus gaillardi* (Ennouchi) from the upper middle Miocene (Tortonian) of France (Brodkorb, 1971).

Although the specimens of *Cypselavus gallicus* and *Apus gaillardi* were not examined in this study, the published illustrations are sufficient to determine that neither species shows any similarities to *Cypseloides ignotus* or the modern *Cypseloidinae*. In fact, *Cypselavus gallicus* shows a distinctly closer resemblance to the Aegialornithidae, the humerus agreeing in size with the newly described small aegialornithid *Primapus lacki*, from the lower Eocene of Britain (Harrison and Walker, 1975). In the published illustrations (Gaillard, 1908), the humerus of *Cypselavus gallicus* appears to lack the prominent ectepicondylar process seen in the Aegialornithidae, but this could well be the result of damage. The illustrations of the humerus of *Apus gaillardi* (Ennouchi, 1930) show it to have the general proportions of the modern Apodidae and Chaeturinae. This species, and an additional swift from the upper Miocene of Italy, are currently under review elsewhere (P. Ballmann, pers. comm.).

The *Cypseloidinae* (see Brooke, 1970:14–15 for use of this term) appears to be the most primitive subfamily of the Apodidae. It is therefore not unexpected that an extinct species of *Cypseloides* be among the earliest known swifts. Like the vultures of the family Cathartidae [= Vulturidae], the modern species of *Cypseloidinae* are confined to the New World; but also like the Cathartidae (Cracraft and Rich, 1972), they can now be shown to have had a past distribution and possible origin in the Old World. Further elucidation of the origin and evolution of the Apodidae will have to await a review of additional modern forms and the remaining fossil swifts, as well as the swift-like members of the Aegialornithidae (see Harrison and Walker, 1975; Harrison, 1975, Collins, pp. 121–127, herein).

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Paris, P.
A New Osprey from the Miocene of California
(Falconiformes: Pandionidae)

Stuart L. Warter

ABSTRACT

Two nearly complete humeri and two partial ulnae from Barstovian age Miocene deposits at Sharktooth Hill, near Bakersfield, Kern County, California, are described as the first known extinct species of the modern genus Pandion. Possible functional implications of the morphological differences observed between the fossil species and modern P. haliaetus are discussed and the fossil record of the Pandionidae is reviewed.

Introduction

In 1973 an avian fossil owned by a private collector was brought for identification to the Natural History Museum of Los Angeles County (LACM) by Mr. Raj Guruswami-Naidu. The specimen, from the Miocene Sharktooth Hill beds, was identified by Dr. Hildegarde Howard and me as a right humerus closely resembling that of a modern osprey, Pandion haliaetus. The specimen was cast and returned, subsequently to be obtained anew by the collector and original owner, Mr. William Hawes, who donated it to the LACM, along with portions of a left humerus and parts of right and left ulnae that were found associated with it. Through the courtesy of Dr. Howard, Dr. Lawrence Barnes and Dr. David Whistler, all of the Department of Vertebrate Paleontology, LACM, the specimens were made available to me for study.

Upon detailed comparison, the bones, which bear a remarkable resemblance to those of modern Pandion haliaetus, were found to differ from that species in a number of subtle, but apparently significant features. These were considered sufficiently important to warrant recognition of a new species, thus extending the history of the genus Pandion back as far as the Miocene.

The terminology used follows that of Howard (1929) and Fisher (1946). Twelve specimens of modern P. haliaetus were examined, four at the LACM and eight at the University of California, Los Angeles. Appreciation is expressed to the curators of these collections for their cooperation. Detailed comparisons are based on skeleton LACM Bi 268, which is typical of larger specimens of P. haliaetus.

Pandion homalopteron, new species

Figures 1-3

Holotype.—Nearly complete left and right humeri and proximal portions of left and right ulnae, all associated; LACM 42815; collected by Mr. William Hawes of Bakersfield, California. Right humerus entire, but shattered and filled in two places with plaster; surfaces of head and internal tuberosity sufficiently intact to permit reasonably accurate total measurement (151 mm), but other contours of both ends badly eroded; deltoid crest missing. Left humerus consisting of three pieces, plus fragments; shaft and distal end joined by actual contact along external surface, proximal portion joined by comparison with companion right humerus and with left humerus of recent Pandion. Head of left humerus entire, but other contours of proximal end severely eroded; all articular surfaces of distal end intact; only a small portion of

Stuart L. Warter, Department of Biology, California State University, Long Beach, California 90840, and Research Associate, Natural History Museum of Los Angeles County, Los Angeles, California 90007.
base of deltoid crest remaining. Left ulna badly shattered, but nearly complete (180 mm), lacking approximately 20 mm or less of the distal end; proximal articular surfaces largely intact, tip of olecranon and tip of external cotyla broken. Right ulna less complete (120 mm) with proximal articular surface largely intact, but olecranon, tip of internal cotyla, and edge of external cotyla missing; distal 48 mm of shaft lacking the anconal surface, last 32 mm displaced to the palmar side and joined only by matrix. Colors variable: right humerus light tan; right ulnar fragment brown; left hu-
Figure 2.—Holotype humeri of *Pandion homalopteron*, new species (LACM 42815): *a*, external view and *b*, internal view of left and right elements. Natural size. (Courtesy of Natural History Museum of Los Angeles County)

merus with proximal segment light tan, shaft and distal segment brown; left ulna brown proximally, fading to light tan distally.

Locality and Age.—From the Sharktooth Hill bone bed, middle Miocene (Barstovian age, Savage and Barnes, 1972:133). Round Mountain Silt, Sharktooth Hill, near Bakersfield, Kern County, California; LACM locality 3205.

Measurements of Holotype.—See Table 1. The brachialis scar of the right humerus is $13 \pm 2.5$ mm long by $6 \pm 1$ mm wide (margin indistinct) and the length of the brachialis scar of the right
Table 1.—Measurements (mm) of the holotype humeri (R = right; L = left) of Pandion homalopteron, new species, compared with modern P. haliaetus

<table>
<thead>
<tr>
<th>Character</th>
<th>P. homalopteron</th>
<th>P. haliaetus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total length</td>
<td>151 (R)</td>
<td>145.9</td>
</tr>
<tr>
<td>Width of head from the external to the internal tuberosity</td>
<td>27.5 (R)</td>
<td>27.1</td>
</tr>
<tr>
<td>Length from head to distal margin of deltoid crest</td>
<td>63 (R &amp; L)</td>
<td>63.3</td>
</tr>
<tr>
<td>Ratio of above measurement to total length (= % of total length)</td>
<td>41</td>
<td>42.9</td>
</tr>
<tr>
<td>Width of shaft at distal end of deltoid crest</td>
<td>12 (R &amp; L)</td>
<td>11.5</td>
</tr>
<tr>
<td>Length of ectepicondyle from proximal margin of extensor metacarpi radialis to distal margin of flexor metacarpi radialis scar</td>
<td>14.5 (R &amp; L)</td>
<td>15.8</td>
</tr>
<tr>
<td>Length of entepicondyle from proximal margin of pronator brevis scar to distal margin of flexor carpi ulnaris scar</td>
<td>12.5 (L)</td>
<td>14.6</td>
</tr>
<tr>
<td>Width of distal end</td>
<td>24.5 (L)</td>
<td>23.8</td>
</tr>
</tbody>
</table>

ulna is 30 mm. These measurements in P. haliaetus are variable and may differ between the right and left sides of the same individual: the brachialis scar of the humerus ranges from 15 to 17.5 mm in length and 7 to 8.5 mm in width (n = 10); that of the ulna ranges from 28 to 36 mm in length (n = 10).

Diagnosis.—Pandion with humerus and ulna resembling those of large individuals of modern P. haliaetus, but showing evidence of weaker musculature and other osteological features that probably permitted less extension at the elbow and less rotation at the shoulder.

Etymology.—Greek homalos, even, level; and pteron, wing; referring to the more level configuration of the wing that would result from a reduced ability to raise the wrist during soaring, thereby reducing or eliminating the “kinked-wing” appearance often presented in flight by members of the modern species.

Description.—Humerus with head more triangular, less rounded than in the modern form; capital groove and ligamental furrow shallower, less deeply excavated; anconal surface of internal tuberosity in internal view less tapering, more nearly perpendicular to main axis of shaft; capital groove and median crest not extending below pneumatic foramen as they do in P. haliaetus.

Distal end of humerus with internal condyle higher, more rounded than in P. haliaetus; olecranal fossa in palmar-distal view shallower and wider; border of fossa in anconal-distal view less rounded, more triangular; brachial depression noticeably smaller and less excavated, its external margin situated more externally; external condyle in palmar view rotated, its long axis at a greater angle from the axis of the shaft; viewed from the external side the external condyle rounder, less squared, and less deep than in the modern form. Ectepicondylar and entepicondylar prominences shorter, the scars for M. extensor metacarpi radialis and M. pronator brevis closer to the distal end of the bone; facet of anterior articular ligament wider and shorter, its surface flat to concave, this facet in P. haliaetus being longer, narrower, and convex.

Proximal end of ulna with surface of internal cotyla shallower, its lip (palmar surface) more extensively flared; palmar lip of radial depression less enlarged; surface of external cotyla less angled from the axis of the shaft, more steeply inclined from the surface of the internal cotyla; scar for the insertion of M. brachialis (= M. brachialis anticus) shorter than in most modern specimens of equivalent size; prominence for anterior articular ligament with shorter, wider facet; proximal half of radial surface of ulna convex in cross-section, whereas flattened or concave in the modern form; olecranon apparently less robust.

Discussion

The modern Osprey, Pandion haliaetus, is a highly specialized fish hunter. It is capable of hovering over one spot with rapidly beating wings held high over the back. Also, it is capable of increasing
Figure 3.—Holotype ulnae and humerus of *Pandion homalopteron*, new species (LACM 42815):  
*a*, dorsal view and *b*, palmar view of left and right ulnae; *c*, distal end of left humerus in distopalmar view. Natural size. (Courtesy of Natural History Museum of Los Angeles County)
the angle of attack of the wings in level soaring flight by raising the wrists through rotation of the long outstretched forelimbs. In this position the wings present the "kinked" or "crooked" appearance for which it is so well known.

The distinctive shape of the deltoid crest of *P. haliaetus* is undoubtedly related to the aforementioned abilities. It is a large, triangular plate, deflexed from the external surface in a palmar direction and beginning abruptly at the lesser trochanters. The deltoid crest is missing from both humeri of *P. homalopteron*, sufficient base of the crest of the left humerus to tell that the entire process was similarly positioned, enlarged and deflexed (Figure 1).

A large deltoid crest usually is indicative of strong pectoral musculature and well-developed powers of flight. Perhaps paradoxically, such a crest occurs in *Pandion* along with a rounded sternal carina, a feature that in soaring birds like cather-tid vultures often is associated with a small deltoid crest and a relatively weak flight mechanism. In *Coragyps* (Fisher, 1946:603), the palmar surface of the deltoid crest provides the area of insertion of the superficial layer of M. pectoralis (= pectoralis superficialis), and on the anconal surface of the crest, M. deltoideus minor and M. deltoideus major originate on the crest and on extensive areas of the shaft of the humerus proximal, distal, and posterior to the deltoid crest. Judging from muscle scars on the humerus of *P. haliaetus*, however, the enlarged deltoid crest serves as the area of insertion for much, if not all, of the large M. deltoideus major, whereas M. deltoideus minor is small and inserts anconally along the shaft anterior to the crest.

In *Coragyps*, the anterior portion of M. deltoideus major "is more important in elevating the leading edge of the wing since the posterior extension of the deltoid crest provides a longer lever arm" (Fisher, 1946:590). An important function of the expanded, deflexed, deltoid crest in *Pandion*, then, is to provide a lever arm for increasing the upward rotational ability of the humerus through the action of M. deltoideus major. A similar increase in downward rotational ability would probably be conferred to the humerus by M. pectoralis.

The humerus of *Pandion* also has an enlarged internal tuberosity, the function of which is to increase the lever arm for several small muscles inserting upon it which, in vultures (Fisher, 1946:603), serve to depress the trailing edge of the wing, thereby raising the leading edge. The described differences in the humeral head, capital groove, ligamental furrow, and internal tuberosity of *P. homalopteron* may indicate a lesser degree of muscular development and rotational ability in the shoulder than in *P. haliaetus*.

The differences in the morphology of the elbow joint provide additional evidence of some degree of functional dissimilarity between the two species of *Pandion*. The configuration of the joint surfaces would seem to indicate a lesser degree of extension at the elbow in *P. homalopteron*. This appears to be borne out by mechanical manipulation of the bones. The forearm of *P. haliaetus* exhibits a much greater degree of extension at the elbow than does that of *Buteo*. The robust olecranon of *P. haliaetus* fits closely into its corresponding depression on the humerus, possibly serving as a bony stop against further extension. The degree of this extension in *P. homalopteron* is also greater than in *Buteo*, but less than in *P. haliaetus*. In spite of the olecranon being incomplete, the ulna of *P. homalopteron* could not be extended to the same degree as that of *P. haliaetus* without partially disarticulating the joint. The observed differences in the size and configuration of the attachment for the anterior articulat ligament also might be related to the decreased ability to extend the elbow.

In the fossil form, the convex radial surface of the ulna, the smaller M. brachialis scars, and possibly the more distal origins of Mm. pronator brevis and extensor metacarpi radialis, may indicate weaker intrinsic musculature. All three of the above muscles are involved in flexion of the forearm, while Mm. brachialis and pronator brevis also are involved in supination and pronation, respectively, of the manus (Fisher, 1946:591-594).

Intrinsic rotational movements of the hand and forearm in birds are limited (Bellairs and Jenkin, 1960:258), and the degree to which they occur has not been determined (George and Berger, 1966:14). However, *P. haliaetus* may have greater abilities to raise the wrist through intrinsic rotation, as well as by rotation at the shoulder of an entire, more extended wing, than did *P. homalopteron*.

There is no reason to assume that *P. homalopteron* was any less variable in its dimensions than is modern *P. haliaetus*. To the extent that the
single available specimen can be considered typical of the Miocene population, the species *P. homalopteron* appears to have been larger in absolute gross skeletal dimensions than an average-sized modern Osprey, but was smaller than average in other measurable features. Several of these features may indicate a lighter wing musculature relative to bone size than is found in *P. haliaeetus*. This, along with the seemingly lesser powers of extension and rotation of the wing, presents a picture of a bird similar to the modern Osprey in size, but one with a more level wing and less refined powers of soaring and hovering. Such a bird could be ancestral to *P. haliaeetus*.

**The Fossil Record of the Pandionidae**

*Pandion homalopteron* provides the only Tertiary record of the family Pandionidae founded on adequate and diagnostic material. Brunet (1970) has placed the species *Palaeocircus cuvieri* Milne-Edwards, based primarily on an incomplete carpo-metacarpus from the upper Eocene of France, in the Pandionidae, stating that the type, while specifically distinct, is scarcely separable from *Pandion* at the generic level. Storrs Olson (pers. comm.) believes that Brunet's illustrations of the specimen indicate to the contrary, however, since both the proximal and distal symphyses between metacarpals II and III are longer than in *Pandion*. The assignment of *Palaeocircus* to the Pandionidae should be regarded with caution, particularly since the family is at present monotypic and the addition of another genus would require redefinition of the family.

A record of *Pandion* from the middle Pliocene Bone Valley Formation in Central Florida (Brodkorb, 1972) is based on a single claw (Storrs Olson, pers. comm.). Another claw, kindly lent to me by the National Museum of Natural History, Smithsonian Institution, (USNM 192193), comes from the Lee Creek phosphate mine near Aurora, Beaufort County, North Carolina. Middle Miocene and Pliocene fossiliferous deposits are exposed there, and in this case it is not certain from which level the specimen was derived (Storrs Olson, pers. comm.). This claw is from digit III of the right foot, but has the tip broken so that an accurate measurement of the chord is not possible. It is referable to the genus *Pandion* and is of a size appropriate for either *P. homalopteron* or *P. haliaeetus*, but since this element is not diagnostic and its age is uncertain, no specific identification can be made.

The only other fossil records for the Pandionidae are Pleistocene remains of the modern species *Pandion haliaeetus*. To the various localities listed in Brodkorb (1964:260) may be added a previously unreported left tarsometatarsus (LACM 27082) from Pleistocene deposits at Kelly Springs, Kelly Park, Orange County, Florida (LACM locality 7119).

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George, J. C., and A. J. Berger  

Howard, H.  

Savage, D. E., and L. G. Barnes  
A New Species of Flightless Auk from the Miocene of California (Alcidae: Mancallinae)

Hildegarde Howard

ABSTRACT

Praemancalla wetmorei is described from the late Miocene of Orange County, California, with humerus and ulna as holotype and paratype, and radius, carpometacarpus, and coracoid referred. The species, although less specialized as a flightless diver than the geologically younger genus Mancalla, appears to be more advanced than Praemancalla lagunensis, which is believed to be derived from slightly older deposits.

Introduction

Since Lucas (1901) described the first humerus of Mancalla, knowledge of the flightless mancalline alcids (Mancallinae) has increased to include nearly all skeletal elements and to involve five species and two genera.

The type-genus, Mancalla, is known from four species—M. californiensis Lucas (1901), M. diegense (Miller, 1937), M. milleri Howard (1970) and M. cedrosensis Howard (1971)—and is recorded from Humboldt County in northern California to Cedros Island, Mexico. The Humboldt County site, with a single humerus assigned to M. diegense (Howard, 1970), is believed by Kohl (1974:217) to be Pleistocene in age. The other records are middle to late Pliocene. Praemancalla is known from the single species, P. lagunensis Howard (1966), described from a late Miocene deposit in Leisure World, Laguna Hills, Orange County, California.

Recently, three other Miocene sites in Orange County have yielded mancalline bones. These sites are in Laguna Niguel, approximately 5 km south of the Laguna Hills locality. The specimens from these sites are in the collections of the Natural History Museum of Los Angeles County. The catalog and locality numbers are listed under Los Angeles County Museum (LACM).

The associated avifauna from these sites includes the same families recorded at the type-locality of Praemancalla lagunensis (LACM Loc. 1945), but the species represented are not identical. None of the species described as new from locality 1945 has appeared in the Laguna Niguel localities. On the basis of associated marine mammals, it is suggested that these sites may represent a later subdivision of the late Miocene than locality 1945 (Barnes, et al., in prep.).

The mancalline skeletal elements from Laguna Niguel include humerus, ulna, radius, carpometacarpus, and coracoid, all of which have been previously described for Mancalla. Only for the carpometacarpus, coracoid, and distal end of the humerus is there comparable material of Praemancalla. The newly found specimens suggest a generally larger form than any previously described mancalline species. Qualitative characters show distinction from comparable elements of all species of Mancalla. Distinctions are also apparent with respect to Praemancalla lagunensis, but the degree of adaptation towards restriction of the wings for swimming is closer to Praemancalla than to Mancalla. Possibly a third genus is indicated. At the
present state of knowledge, however, it seems wiser to assign the specimens to the genus *Praemancalla* under a new species name.

**Acknowledgments.**—I am grateful to the Earth Sciences Division of the Natural History Museum of Los Angeles County for placing the museum's collections at my disposal and for the many courtesies of the staff members. I particularly appreciate the assistance of Dr. Lawrence Barnes in discussion of matters pertaining to the field work in Orange County. The photographs were taken by Lawrence Reynolds, museum photographer. Special thanks go to my husband, Henry Anson Wylde, for the art work in connection with the preparation of the plate.

*Praemancalla Howard*

In describing *Praemancalla lagunensis*, the specific diagnosis of the holotype humerus and paratype carpometacarpus served also as the generic diagnosis. In the specimens from Laguna Niguel now at hand, the following characters are in agreement with that diagnosis:

Humerus with groove separating base of ectepicondyle from external condyle, brachial impression faint and running diagonally from ectepicondylar prominence to a point slightly proximal to attachment of articular ligament, with no papilla present above condyles. The tricipital grooves and ridges are broken in the humerus from Laguna Niguel, so the characters of this area set forth in the original diagnosis cannot be assessed.

Carpometacarpus with distinct, blunt pisiform process, trochlear area having narrow, deep groove between internal and external crests posteriorly, metacarpal II relatively broad with more rounded anterior contour and more angular internal contour than in *Mancalla*, and process of metacarpal I relatively shorter.

The following additional characters observed in the specimens now at hand are considered to be of generic value when compared with *Mancalla*: humerus with head only slightly extended over capital groove, deltoid crest weakly developed, area of anterior articular ligament slightly swollen; ulna with prominent olecranon process; radius lacking prominent crest on convex contour; coracoid with scapular facet facing dorsally, coracohumeral attachment flat and angular in outline.

**Praemancalla wetmorei, new species**

**Figure 1a, b, e-g, i-k**

**Holotype.**—Humerus, LACM 42653, complete except for tricipital area of distal end (Figure 1i,j).

**Type-Locality.**—LACM Loc. 6906, site of excavation for North American Rockwell building (now U.S. General Services Administration building) on El Lazo Road, Laguna Niguel, Orange County, California; 914 m north of junction of Aliso Creek and Sulphur Creek, in yellow sands and laminated gray shale. Latitude 33°33’43” N, longitude 117°42’44” W. In the NE 1/4 NE 1/4 SE 1/4 of unsurveyed Sec 16, T7S, R8W, San Juan Capistrano quadrangle, USGS 1948, 1:24000.

**Formation and Age.**—Monterey Formation, late Miocene.

**Paratype.**—Proximal end of ulna LACM 32429 from type-locality (Figure 1e).

**Diagnosis.**—Humerus broad proximally; medial profile of capital groove a wide open curve; depth through deltoid crest only 5 percent greater than depth of shaft above distal end; ectepicondylar prominence notably protuberant at its proximal tip and slightly lateral in position with respect to palmar surface of shaft; groove between external condyle and base of ectepicondyle more constricted and less distal in extent than in *P. lagunensis*; shaft breadth above ectepicondylar prominence 55 percent of shaft depth at same point; shaft depth 113 percent of breadth of distal end.

Ulna laterally compressed, with short brachial impression partially palmar in position and bordered palmar by heavy ridge; olecranon blunt but protruding proximally beyond cotylae and distinctly set off from cotylae by lateral depression both externally and internally.

**Measurements.**—Humerus: length to external condyle 82.7 mm, greatest proximal breadth from pectoral to bicipital crests 22.2 mm, breadth across head 19.6 mm, breadth through distal condyles 8.5 mm, breadth and depth of shaft above ectepicondylar prominence 5.1 mm and 9.6 mm, respectively, height of ectepicondylar prominence above distal end 16.9 mm, greatest depth through deltoid crest 10.1 mm, breadth of shaft at same point 5.5 mm. Ulna: proximal breadth across cotylae 7.5 mm, proximal depth through olecranon 11.3 mm,
Figure 1.—Skeletal elements of Praemancalla and Mancalla: a, b, coracoid (LACM 37637) of P. wetmorei, new species, medial and dorsal views; c, coracoid (LACM 15289) of P. lagunensis, dorsal view; d, coracoid (LACM 2310) of M. diegense, dorsal view; e, paratype ulna (LACM 32429) of P. wetmorei, internal view; f, referred carpometacarpus (LACM 52216) of P. wetmorei, internal view; g, radius (LACM 53907) of P. wetmorei, palmar view, h, humerus (LACM 15367) of M. cedrosensis, palmar view; i, j, holotype humerus (LACM 42653) of P. wetmorei, palmar and anconal views; k, referred humerus (LACM 32432) of P. wetmorei, anconal view; l, humerus (LACM 2331) of M. diegense, anconal view. (Approximately natural size.)
breadth and depth at middle of shaft 4.2 mm and 6.2 mm respectively.

Referred Material.—From Laguna Niguel, Orange County, California, late Miocene, Monterey Formation. Proximal end of humerus LACM 32432 (Figure 1k) from type-locality (LACM Loc. 6906). Complete radius LACM 53907 (Figure 1g) and scapular end of coracid LACM 37637 (Figure 1a,b) from LACM locality 6902 at northwest end of El Lazo Road, 365 m northwest of type-locality (LACM Loc. 6906). Complete radius LACM 53907 from LACM locality 6902 at northwest end of El Lazo Road, 365 m northwest of type-locality and 244 m east of Aliso Creek, in basal 0.5–1.5 m of coarse yellow sand directly overlying laminated gray shale. Proximal 3/4 of carpometacarpus LACM 52216 (Figure 1f) from LACM locality 3185, in tributary gully west of Aliso Creek in coarse yellow sand.

Etymology.—I take pleasure in naming this new species in honor of Dr. Alexander Wetmore, who has done so much to further the science of paleornithology and who has generously given advice and counsel to me throughout my years of study in this field.

Description.—Compared with Mancalla, the humerus of P. wetmorei is relatively, as well as actually, broader both proximally and distally (ratio of greatest proximal breadth to length 26.8 percent in P. wetmorei; 23–25 percent in Mancalla; ratio of distal breadth to length 10.2 percent in P. wetmorei, 8.2–9.6 percent in Mancalla) and exceeds in length all but one specimen of Mancalla (the maximum of M. diegense). It is, however, 8 percent longer than the average for M. diegense and 12 percent longer than the average for M. cedrosensis (Table 1), and 32 percent longer than the much smaller M. milleri.

The lesser protrusion of the head over the capital groove is reflected in the wide, open curve between the head and internal tuberosity as seen in palmar and anconal views; this condition contrasts with the narrow, U-shaped curve found in Mancalla (Figure 1k–l). Further distinction from Mancalla is seen in the deltoid crest which, in P. wetmorei, describes a low, even arc and is not expanded towards its distal termination. Distally, the greater breadth of the humerus is observed not only in the width through the condyles but also in a slight expansion in the region of the attachment of the anterior ligament. In this character, as well as in the lateral slant of the brachial impression and absence of a prominent papilla above the condyles, P. wetmorei resembles Praemancalla lagunensis. It is distinguished from that species in the greater projection of the ectepicondylar prominence from the shaft, narrower groove between the base of the ectepicondyle and external condyle, and relatively narrower and deeper shaft (relative breadth to depth of shaft 53 percent in P. wetmorei, 66 percent in P. lagunensis). In depth of shaft relative to breadth of distal end, P. wetmorei is intermediate between Praemancalla lagunensis and the several species of Mancalla (99 percent in P. lagunensis, 115 percent in P. wetmorei, 126–140 percent in Mancalla).

The prominence of the olecranon immediately distinguishes the ulna of P. wetmorei from all species of Mancalla, but the palmar position of the brachial impression and the shortened lip of the external cotyla assign the element to the subfamily Mancallinae rather than the typical alcids.

The radius (LACM 53907), although short and laterally compressed as in Mancalla, is less blade-like and lacks the prominent crest on its convex contour. The ulnar depression is broader and deeper than in Mancalla. Neither the ulna nor the radius is known for Praemancalla lagunensis. Those assigned to P. wetmorei both show less modification towards a flipper-like wing than in Mancalla, and in this regard are in keeping with the character of the other elements known for Praemancalla.

The radius is 12 percent longer than the maximum known for any species of Mancalla (Table 1). Using the radius as a guide, and comparing the relative size of ulna to radius in the type of Mancalla cedrosensis (associated skeletal elements of one individual), it is suggested that the ulna of P. wetmorei attained a length of 36.5 mm.

Carpometacarpus LACM 52216 differs from that of Mancalla and resembles Praemancalla in the presence of a distinct, blunt pisiform process, rounded anterior contour of shaft of metacarpal II and deep narrow groove between the internal and external crests of the trochlea posteriorly. It is distinguished from P. lagunensis by the narrower shaft and the relatively longer process of metacarpal I, with more than half its length distal to the level of the metacarpal symphysis; also, the trochlea extends higher above that process and the lateral surface of the internal crest of the trochlea is more broadly and less deeply depressed.

In coracid LACM 37637 the furcular facet is
Table 1.—Skeletal measurements (mm) of *Praemancalla wetmorei* compared with *P. lagunensis*, *Mancalla diegense*, *M. cedrosensis*, *M. californiensis*

<table>
<thead>
<tr>
<th>Character</th>
<th>P. wetmorei</th>
<th>P. lagunensis</th>
<th>M. diegense min. mean max.</th>
<th>M. cedrosensis min. mean max.</th>
<th>M. californiensis*</th>
</tr>
</thead>
<tbody>
<tr>
<td>HUMERUS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length</td>
<td>82.7</td>
<td>-</td>
<td>71</td>
<td>76.5</td>
<td>85.2</td>
</tr>
<tr>
<td>Greatest proximal breadth</td>
<td>22.2</td>
<td>-</td>
<td>17.3</td>
<td>18.7</td>
<td>20.3</td>
</tr>
<tr>
<td>Distal breadth</td>
<td>8.5</td>
<td>7.8</td>
<td>6.4</td>
<td>6.7</td>
<td>8</td>
</tr>
<tr>
<td>ULNA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Proximal depth</td>
<td>11.3</td>
<td>-</td>
<td>9†</td>
<td>9.3</td>
<td>8.8</td>
</tr>
<tr>
<td>Proximal breadth</td>
<td>7.5</td>
<td>-</td>
<td>5.9</td>
<td>6.4</td>
<td>6.6</td>
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<tr>
<td>RADIUS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Greatest length</td>
<td>35.8</td>
<td>-</td>
<td>29.6</td>
<td>30.9</td>
<td>31.8</td>
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<tr>
<td>Greatest shaft depth</td>
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<td>-</td>
<td>6.3</td>
<td>6.25</td>
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<tr>
<td>Shaft breadth</td>
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<td>-</td>
<td>2.3</td>
<td>2.5</td>
<td>2.7</td>
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<tr>
<td>CARPOMETACARPUS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length process</td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>Metacarpal I</td>
<td>15.7</td>
<td>14</td>
<td>15.2</td>
<td>15.3</td>
<td>15.5</td>
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<tr>
<td>Proximal depth</td>
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<td>11.9</td>
<td>9.7</td>
<td>10.3</td>
<td>11</td>
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<tr>
<td>Shaft breadth</td>
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<td>4.5</td>
<td>3.1</td>
<td>3.4</td>
<td>3.7</td>
</tr>
<tr>
<td>CORACOID</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length from below scapular facet to head</td>
<td>20.9</td>
<td>18.8</td>
<td>15.4</td>
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<td>19.5</td>
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<tr>
<td>Breadth below furcular facet</td>
<td>5.8</td>
<td>6</td>
<td>5.4</td>
<td>5.8</td>
<td>6.3</td>
</tr>
<tr>
<td>Breadth furcular facet</td>
<td>10.3</td>
<td>-</td>
<td>7.3</td>
<td>7.8</td>
<td>8.6</td>
</tr>
</tbody>
</table>

* Only one specimen of each element of *M. californiensis*, except carpometacarpus (average of four).
† Only two specimens measurable for this dimension.

broad and deep; it extends ventrally well beyond the triosseal canal, is strongly thrust mediad above the canal, and is markedly undercut. Below the facet the bone narrows and the ventral border of the triosseal canal is sharply angular. The species of *Mancalla* vary in development of the furcular facet and the bordering of the triosseal canal. The greatest overhang of the facet and the least angular border of the triosseal canal are found in *M. cedrosensis*; the least overhang and most angular border of the canal occur in *M. californiensis*. In no specimen of *Mancalla* is the furcular facet as ventrally extended as in *P. wetmorei*. This facet is broken ventrally in the single known coracoid of *P. lagunensis*, but the portion that remains is deep and has a strong overhang; below the facet, however, the area is broader and more rounded than in *P. wetmorei*. In direct dorsal view (with dorsal surface of shaft held horizontally) the scapular facet in *P. wetmorei* is more dorsally and less laterally directed than in *Mancalla*, and the triosseal canal faces more mediad. Resemblance is closer to *Praemancalla lagunensis*, although the canal is even more mediad directed in the latter species. The attachment of the coracohumeral muscle in *P. wetmorei* is broad, flat, and angular in outline at its anterior end, as in *P. lagunensis*, but is relatively longer, and narrows near the glenoid facet. In *Mancalla* the attachment is narrow and rounded.
Conclusions

Four wing elements and a coracoid from three correlative localities of late Miocene age in Laguna Niguel, Orange County, California, represent a new species, *Praemancalla wetmorei*, in the alcid subfamily Mancallinae. The degree of specialization towards a flipper-like wing is less than in the Pliocene genus *Mancalla*, and reflects a stage of development closer to the Miocene genus *Praemancalla*. Differences noted with respect to *Praemancalla lagunensis*, however, suggest a slight advance towards the more specialized wing of *Mancalla*. The humerus is more compressed, the trissoseal canal of the coracoid more dorsally rotated and narrower, and the process of metacarpal I of the carpometacarpus longer.

This suggested evolutionary trend is in keeping with the evidence presented by the associated faunas of the localities involved, which indicates a slightly greater age for the type-locality of *P. lagunensis* than for the Laguna Niguel sites.

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Howard, Hildegarde

Kohl, Roy F.

Lucas, Frederic A.

Miller, Loye
The Pleistocene Pied-billed Grebes
(Aves: Podicipedidae)

Robert W. Storer

ABSTRACT

Pleistocene specimens of pied-billed grebes (Podilymbus) were compared with a series of skeletons of the modern North American form, Podilymbus podiceps podiceps. Most of the fossils agreed closely with this form and are allocated to it. The co-types of Podilymbus magnus Shufeldt also fall within the range of variation of this form, hence P. magnus becomes a synonym of P. podiceps. A new species, Podilymbus wetmorei, characterized by a wide tarsometatarsus and a heavy femur, is described from the Pleistocene of Florida.

Introduction

The Pied-billed Grebe (Podilymbus podiceps) is widely distributed in the New World from Canada to southern South America. The only other living species of the genus, the Atitlán or Giant Pied-billed Grebe (P. gigas), is confined to Lake Atitlán, Guatemala. The genus is represented in upper Pliocene deposits of Idaho by a large species, P. majuscus (Murray, 1967), and in numerous Pleistocene deposits. Most of the Pleistocene specimens have been assigned to the living species, P. podiceps, but a few have been referred to an allegedly larger extinct species, P. magnus. The latter was first described by Shufeldt (1913:136–137) on the basis of two tarsometatarsi and a coracoid from Fossil Lake, Oregon. Later, Wetmore (1937:198–199) synonymized P. magnus with P. podiceps, pointing out that there is considerable sexual dimorphism in the genus and that Shufeldt had only one skeleton (a female) of the living species with which to compare his fossil material. Wetmore found that the larger of the tarsometatarsi described by Shufeldt was only slightly larger than those of two males of the living North American subspecies (P. p. podiceps) and was matched by an example of the slightly larger South American race (P. p. antarcticus). More recently, Brodkorb (1959:273–274) revived the name P. magnus for twelve bones from Arredondo, Florida, using eight skeletons of the living North American form for comparison. He (1963a:113) also referred material from the Santa Fe River, Florida, to P. magnus.

McCoy (1963:337) in his report on the fossil avifauna of the Ichetucknee River, a tributary of the Santa Fe, referred two tarsometatarsi to P. magnus and 47 other bones (including two other tarsometatarsi) to P. podiceps. Subsequently, Brodkorb (1963b:230) wrote that “specimens from Fossil Lake and some of the Floridian localities average large and are perhaps recognizable as a temporal subspecies, Podilymbus podiceps magnus Shufeldt.”

The availability of a series of 39 skeletons of the modern North American form (Podilymbus p. podiceps) from Michigan and Wisconsin has permitted a better estimate of variation within a living population of this species than was heretofore possible, as well as providing a comparison of skeletal elements of this population with a large number of fossil elements from late Pleistocene deposits. The following fossil material has been examined: California: McKittrick, 1 tarsometatarsus; Rancho La Brea, 1 femur; Florida: Reddick, 3 coracoids, 1 humerus, 1 tibiotarsus, 2 tarsometatarsi; Haile, 1 coracoid, 1 ulna, 1 tibiotarsus; Arredondo, 2 coracoids, 3 humeri, 1 ulna, 3 carpometacarpi, 1 femur,
### Table 1.—Measurements (mm) of modern and late Pleistocene Pied-billed Grebe bones

<table>
<thead>
<tr>
<th>Character</th>
<th>MODERN</th>
<th>Fossil</th>
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<th>FEMALES</th>
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<tr>
<td></td>
<td>n</td>
<td>max.</td>
<td>min.</td>
<td>mean±σm</td>
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<tr>
<td>Length</td>
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<td>33.9</td>
<td>30.1</td>
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<tr>
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<td>2.7</td>
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<tr>
<td>Depth head</td>
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<td>4.6</td>
<td>5.17±0.05</td>
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<td><strong>HUMERUS</strong></td>
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<tr>
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<td><strong>ULNA</strong></td>
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<td></td>
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<tr>
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<td>69.1</td>
<td>71.87±0.04</td>
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* Calculated by subtracting the length measured from the articulation from the total length.

3 tarsometatarsi; Catalina Lake, 1 coracoid; Hornsby Spring, 1 femur, 1 tibiotarsus; Itchucknee River, 28 coracoids, 58 humeri, 7 ulnae, 7 carpo-
metacarpi, 18 femora, 23 tibiotarsi, 20 tarsometatarsi; Jenny Spring, 1 humerus; Lake Monroe, 2
humeri, 1 femur, 1 tibiotarsus; Rock Spring, 2
coracoids, 5 humeri, 1 ulna, 6 tibiotarsi; St. John’s
Lock, 3 humeri, 1 ulna, 1 tibiotarsus, 1 tarsomet-
tarsus; Vero Beach, 1 humerus. Nevada: Smith
Creek Cave, 1 coracoid. Oregon: Fossil Lake, 4
tarsometatarsi (including the types of P. magnus).
Total: 220 bones.

Using dial calipers reading to one tenth of a
millimeter, 33 measurements were taken from the
fossil and modern bones (Table 1). Each fossil
specimen was “sexed” by comparing it with the
largest and smallest element for each sex in the
modern sample. There is very little or no overlap
between the sexes in the total lengths of the elements measured, males being larger. In instances in which a fossil was within a zone of overlap in length, it was almost always outside a zone of overlap in another measurement. Therefore, I believe that few, if any, of the fossils were assigned to the wrong sex and only a few fragmentary specimens could not be "sexed" in the above manner. Variability within the sexes of the modern material and the fossils "sexed" in this way proved similar, a further indication that most, if not all of the fossils were assigned to the correct sex.

The means and, where practical, the standard errors of the means and standard deviations of the measurements for these bones were calculated and compared with those of the modern material (Table 1).

Acknowledgments.—I am indebted to Pierce Brodkorb of the University of Florida (PB) and to the curators of the American Museum of Natural History (AMNH), the Natural History Museum of Los Angeles County (LACM), the Museum of Comparative Zoology (MCZ), the University of Florida (UF), and the University of Michigan Museum of Paleontology for permission to use material under their care. The figure was prepared by Mark Orsen. Part of this work was subsidized by the National Science Foundation through Grant GB-8269.

Discussion

The late Pleistocene material is discussed below by skeletal element. Unless otherwise indicated, all specimens in the following sections are from the Ichathucknee River.

Coracoid.—In both sexes, the means of all measurements of the modern and Pleistocene coracoids are very close. One specimen (PB1209 "♂") falls outside the expected range (i.e., mean ± 2σ) of the modern sample in the width of the head (2.5 mm as opposed to 2.6 mm in the smallest modern specimen).

Humerus.—With five exceptions, the fossil humeri agree with the modern series in size. A distal portion (UF 15309) measures 3.0 mm in shaft width and 6.5 mm in distal width, the former measurement being near the mean for males and the latter near the mean for females in the modern series. (This bone was not included in the calculations of the fossil sample because it could not be assigned with confidence to either sex.) Humerus PB 8005 has a shaft width of 3.4 mm, which is approximately 2.5 standard deviations above the mean for males, whereas its other measurements are well within the range of modern males. (It was not included in the calculations of the fossil sample because its identification was not certain; it might belong to the new species described herein.) UF 15297 has a shaft width of 2.4 mm, three standard deviations below the mean for modern females, and a distal width of 6.3 mm, which is within the range of modern females. UF 15280 and UF 15307 have distal widths of 7.2 and 7.0 mm, respectively, which are somewhat outside the expected range of the modern sample.

Ulna.—One partial ulna (PB 7687 from Rock Spring) has a distal width of 4.6 mm or 2.35 standard deviations above the mean for modern males. In other measurements, it is within the expected range of modern specimens, as are the other fossil ulnae examined.

Carpometacarpus.—The fossil carpometacarpi are all within the expected range of variation of the modern sample.

Femur.—With two exceptions (UF 15214 and UF 15220), all the fossil femora fall within the expected range of modern specimens. The two exceptions are referred to the new species and are not included in the calculations of the fossil sample.

Tibiotarsus.—Four fossils are outside the range of the modern sample. UF 15251 "♂" is more than 2 standard deviations above the means for males in distal width (8.2 mm) and least width of shaft (4.4 mm). Two modern specimens are 8.0 mm in distal width and one is 4.3 mm in shaft width.) UF 15254 is 68.7 mm in length, measured from the articulation, which is between the ranges of the two sexes but slightly nearer the mean of females, 3.7 mm in width of shaft, within the range of either sex, and 2.4 mm in depth of shaft, within the range of modern males. MCZ 2606 "♀" and PB 1851 "♀" have cnemial crests well below the range expected for their assigned sexes. In other measurements, they are within the range of modern specimens or differ by only a tenth of a millimeter. These four fossils were not included in the calculations of means for the fossil samples.

Tarsometatarsus.—All but three fossils of this element fall within the expected range of the mod-
ern sample. Two of these (UF 15223 and PB 1762 from Reddick) are very broad and are referred to the new species. The third (PB 1854) measures 44.8 mm in length and 3.5 in least width of shaft. The former measurement is 2.2 and the latter 1.2 standard deviations above the mean for the modern series. These three fossils are not included in the calculations of the fossil sample.

Thus, most of the Pleistocene fossils agree well with the sample of modern specimens. These fossils include the type tarsometatarsi of *P. magnus*, which measure 43.6 and 42.9 mm in length and 3.5 and 3.2 mm in least width. I have not examined the coracoid Shufeldt (1913, pl. 38:fig. 449) tentatively assigned to *P. magnus*, but his photograph of it agrees better with the modern form than two other coracoids (his figures 461, 462) he did assign to *P. podiceps*. Wetmore (1937:199) referred this specimen to the modern form. The 12 fossils from Arredondo referred to *P. magnus* by Brodkorb (1959:273–274) fall within the expected range of variation of the modern form. I have not seen the fossil (or fossils) from the Santa Fe River referred without comment to *P. magnus* by Brodkorb (1963a:115). The tarsometatarsi from the Itchtucknee River referred to *P. magnus* by McCoy (1963:337) also fall within the expected range of the modern form, although one (PB 1854) is 0.6 mm longer than the longest modern specimen measured. Both measure 3.5 mm in least width and are equaled or surpassed by both fossil and modern specimens in this dimension. Thus, I can find no evidence that the late Pleistocene birds were significantly larger than the modern ones. Because the types of *P. magnus* are indistinguishable from the modern form, *Podilymbus magnus* Shufeldt must be considered a synonym of *Podilymbus podiceps* (Linnaeus).

Of the few fossils falling outside the range of the modern series, several are heavier than the corresponding elements of *P. p. podiceps* and are established here as a new species.

**Podilymbus wetmorei**, new species

**Figure 1b,d**

**Holotype.**–Nearly complete left tarsometatarsus in the collection of Pierce Brodkorb (PB 1762) from the Dixie Lime Products quarry, locality IA, 1.6 km south of Reddick, Marion County, Florida. Collected by Pierce Brodkorb, 2 March 1957.

**Age.**–Late Pleistocene (Rancholabrean) fide Webb (1974:13).

**Range.**–Known so far only from two localities in peninsular Florida.

**Diagnosis.**–Similar in length and general configuration to the tarsometatarsus of *Podilymbus podiceps*, the shaft much heavier (Figure 1; Table 2). Differs from *Podilymbus majuscus* Murray (1967), from the upper Pliocene of Idaho, in being shorter, comparatively wider in the shaft, and in having the ridge along the external side of the anterior surface of the bone much reduced for its distal third. Shorter and relatively heavier than the tarsometatarsus of *P. gigas*.

**Referred Material.**–One tarsometatarsus (UF 15223) and two femora (UF 15214 and UF 15220) all from the Itchtucknee River, Columbia County, Florida. Like the type, the second tarsometatarsus has a notably heavy shaft; the femora have thick shafts and wider heads and distal ends than *P. podiceps* (Table 2). Two exceptional humeri mentioned above (UF 15309 and PB 8005 from the Itchtucknee River) have wide shafts and may likewise represent *P. wetmorei*. Two tibiotarsi (MCZ 2606 and PB 1851 from the Itchtucknee River) have short cnemial crests. The former is wide at both articulations and has a thick (deep) shaft, suggesting that it may also belong to the new form. However, the second bone is small in all these dimensions and probably is an aberrant or worn example of *P. podiceps*. The placement of the other fossils that fall outside the range of the modern form is uncertain, and will probably remain so until more material becomes available.

**Remarks.**–Having both a thick tarsometatarsus

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**Table 2.—Measurements (mm) of Podilymbus wetmorei**
and a heavy femur is paradoxical in a foot-propelled diving bird, because the complex of structural modifications which appear adaptive for rapid locomotion under water includes a short, heavy femur, a long cnemial crest, and a narrow tarsometatarsus. In the case of *P. wetmorei*, a thickened but not shortened femur might have provided a greater area for the attachment of muscles used in swimming (notably the gastrocnemius), which could compensate for the area lost through the shorter cnemial crest. Modern pied-billed grebes inhabit bodies of water with considerable emergent vegetation and have wider tarsometatarsi than grebes inhabiting more open water. Presumably this is an adaptation for diving almost directly downward, breast first, rather than springing forward like most other grebes. The even wider tarsometatarsi of *P. wetmorei* probably represent a further extension of this adaptive trend already evident in modern species of *Podilymbus*.

The presence of two species of *Podilymbus* in the same Rancholabrean deposits in Florida suggests that one (*P. wetmorei*) may have been a resident form that differentiated from the widespread *P. podiceps*, which latter may only have wintered there. The situation between the fossil form known as *Gallinula brodkorbi*, also from the Itchucknee, and *G. chloropus* (Olson, 1974:174) suggests an interesting parallel and might be accounted for similarly.

**Etymology.**—I take great pleasure in naming this species in honor of Alexander Wetmore.
**Literature Cited**

Brodkorb, P.


McCoy, J. J.

Murray, B. G., Jr.

Olson, S. L.

Shufeldt, R. W.

Webb, S. D.

Wetmore, A.
The Late Pleistocene Avifauna of La Carolina, Southwestern Ecuador

Kenneth E. Campbell, Jr.

ABSTRACT

A collection of fossils from the late Pleistocene site known as La Carolina, located on the arid Santa Elena Peninsula of southwestern Ecuador, contains 53 species of birds, representing 16 families and 42 genera, including 7 extinct species previously recorded only from the Talara Tar Seeps of northwestern Peru. New species of *Buteo* and *Oreopholus* are described. The genus *Protoconurus* Spillman is synonymized with *Aratinga*.

Seventy-two percent of the species recorded from La Carolina were also recorded from the Talara Tar Seeps. The resemblance between the two avifaunas suggests a similarity in age, habitat, and climatological conditions at the two sites at the time of deposition. Evidence suggests that during glacial periods the currently arid Santa Elena Peninsula was part of a broad, forested coastal savanna extending from central Ecuador south to northern Peru.

Introduction

While on a visit to Quito, Ecuador, in early 1970, in connection with my research on the late Pleistocene avifauna of the Talara Tar Seeps of northwestern Peru (Campbell, in press), I had the pleasure of meeting and talking with Professor Gustav Orcés V. of the Escuela Politécnica Nacional. During the course of our conversations I learned that he had in storage at the Escuela a collection of fossil birds from a site known as La Carolina, located on the Santa Elena Peninsula. The collection was made by Dr. Robert Hoffstetter in the course of his work on the Pleistocene mammals of Ecuador (Hoffstetter, 1952). I was most graciously granted permission to take the fossil birds back to the University of Florida where I could work on them in conjunction with the Talara avifauna.

The La Carolina site is located on the north side of the Santa Elena Peninsula near the town of La Libertad (Figure 1). Hoffstetter (1952) and Edmund (1965) have described the location and geological history of this and other nearby sites. Spillman (1942) also contributed to our knowledge of the paleontology and geological history of the Santa Elena Peninsula, including descriptions of what he regarded as two extinct genera and species of birds.

I have not visited the site personally and must therefore refer the reader to the description published by Hoffstetter (1952). The fossil beds consist of fine-grained aeolian and fluvialite sands that were deposited in a broad, shallow estuarine environment. Certain zones of the deposit are impregnated with pitch and some evidence suggests that this occurred subsequent to the deposition of the sands (Edmund, 1965). Preservation of the fossil material is generally excellent, indicating rapid burial with little disturbance. In addition to the specimens recorded below, there are a large number of small, delicate specimens of various species of Passeriformes that are not sufficiently prepared to include in this report.

Species distributions were taken from Chapman (1926), Marchant (1958), and Meyer de Schauensee (1966). For osteological characters of the various species the reader is referred to a pending paper on the paleoavifauna of the Talara Tar Seeps (Campbell, in press). The La Carolina specimens

Kenneth E. Campbell, Jr., Department of Zoology, University of Florida, Gainesville, Florida 32611.
are currently uncataloged and will be returned to Professor Orcés in the near future.

Acknowledgments.—I wish to thank Professor Gustav Orcés V. for his kind permission to study the avian material from La Carolina. Special thanks also go to Pierce Brodkorb for his assistance in the preparation of this manuscript and for access to his skeletal collection. I am very grateful to the Frank M. Chapman Fund of the American Museum of Natural History, the Society of Sigma Xi, and the Organization for Tropical Studies, Inc. (Pilot Research Proposal No. 69-23) for providing the crucial financial assistance that made possible my field studies in Ecuador and Peru. The National Museum of Natural History and the American Museum of Natural History kindly loaned specimens that were critical for identification of some of the fossil material reported on here. A. Gordon Edmund of the Royal Ontario Museum provided numerous useful comments.

ARDEIDAE

Nycticorax nycticorax (Linnaeus)

Material.—One complete right coracoid, humeral end of 1 left coracoid, 1 complete left carpometacarpus, proximal ends of 1 right and 1 left femur.

Remarks.—N. nycticorax is resident in Ecuador.

ANATIDAE

Dendrocygna autumnalis (Linnaeus)

Material.—One left scapula, 1 complete right coracoid, proximal end of 1 right ulna, distal end of 1 left tibiotarsus.

Remarks.—D. autumnalis is resident in Ecuador.

Anatidae genus and species

Material.—Two complete right and 2 complete left coracoids, humeral end of 1 right coracoid, 1 complete left ulna, 1 complete left radius, 1 complete right carpometacarpus, proximal ends of 1 right and 1 left carpometacarpus.

Remarks.—This new genus and species of Anatidae is being described as a form of Tadorninae from the Talara Tar Seeps of northwestern Peru (Campbell, in press). The presence of this and the other extinct species listed below indicate that paleoclimatic conditions at La Carolina were similar to those at the Talara Tar Seeps, much as similar conditions exist at the two sites today.

Anas species 1

Material.—One complete right coracoid, proximal end of 1 right radius, distal end of 1 right tibiotarsus, 1 complete left tarsometatarsus.

Remarks.—The proximal end of the radius is not included in the description of Anas species 1, from the Talara Tar Seeps (Campbell, in press). The present specimen is referred to this species on the basis of size (proximal width, 4.4 mm), and because it differs from the radius being referred to Anas species 2 (Campbell, in press) by having: (1) ligamental prominence high, (2) ulnar depression deep, (3) scapholunar facet broad and flat.

Anas species 3

Material.—One left scapula, 1 complete left humerus (holotype), 2 complete left carpometacarpi.
Remarks.—The complete humerus from La Carolina is being used as the holotype in the description of *Anas* species 3 (Campbell, in press), because it is in better condition than any of the material from the Talara Tar Seeps. Elements of this species not found in the Talara Tar Seeps include the scapula and carpometacarpus. The small size of these specimens indicates that they may be safely referred to *Anas* species 3. In addition to its small size, the scapula is characterized by having: (1) acromion of moderate length and width, with anteroventral corner a prominent projection; (3) attachment of Lig. furculoscapulare dorsale oval in shape, located on midline of shaft. Measurements of the scapula are: proximal height, 7.7 mm; proximal width, 2.4 mm.

The carpometacarpus is characterized by having: (1) process of metacarpal I moderately high and long; (2) proximal edge of metacarpal I sloping slightly proximad; (3) proximal metacarpal synphysis of moderate length, distal synphysis long; (4) internal face of external rim of carpal trochlea angular proximally; (6) area covered by external cuneiform ligament moderately convex; (7) posterior carpal fossa of moderate width for its length; (8) surface of carpal trochlea extending deep into posterior carpal fossa; (9) internal rim of carpal trochlea with posterior edge bowing externally. Measurements (in mm) of the two carpometacarpi are as follows: length, 35.0 and 35.2; height through metacarpal I, 8.1 and broken; proximal width, 3.9 and 3.9; least width of shaft, 2.5 and 2.7; length of distal forniax, 5.0 and 4.9.

The numbered characters listed above correspond to those being used in the descriptions of *Anas* species 1, and *Anas* species 2 (Campbell, in press). The absence of *Anas* species 2, from La Carolina may be a result of the small sample size, although it was more common at the Talara Tar Seeps than *Anas* species 3.

*Anas bahamensis* Linnaeus

Material.—Portions of 12 crania, 9 right and 4 left scapulae, 23 complete right and 27 complete left coracoids, humeral ends of 5 right and 7 left coracoids, 4 complete right and 2 complete left humeri, proximal ends of 1 right and 2 left humeri, distal ends of 1 right and 4 left humeri, 16 complete right and 18 complete left ulnae, proximal ends of 3 right and 3 left ulnae, distal ends of 2 right and 3 left ulnae, 11 complete right and 3 complete left radii, proximal ends of 7 right and 4 left radii, distal ends of 3 right and 5 left radii, 28 complete right and 54 complete left carpometacarpi, proximal ends of 3 left carpometacarpi, 5 complete right and 8 complete left femora, proximal ends of 2 left femora, distal ends of 3 left femora, 3 complete right and 1 complete left tibiotarsus, 1 almost complete right and 1 almost complete left tibiotarsus, proximal ends of 1 right and 1 left tibiotarsus, distal ends of 4 right and 5 left tibiotarsi, 19 complete right and 24 complete left tarsometatarsi, proximal ends of 2 right and 4 left tarsometatarsi.

Remarks.—In terms of numbers of specimens and of individuals, *A. bahamensis* is the second most abundant species in the La Carolina deposits. This species is currently found in large numbers in southwestern Ecuador when standing bodies of fresh water occur (Marchant, 1958).

I am elsewhere (Campbell, in press) expressing my belief that it was material of this species that Spillman (1942) named *Archeoquerquedula lambrechti*. The absence of any large, extinct duck from the La Carolina deposits upholds the view that *Archeoquerquedula lambrechti* should be synonymized with *Anas bahamensis*.

**VULTURIDAE**

**Vulturidae genus and species indeterminate**

Material.—Proximal end of 1 right tarsometatarsus.

Remarks.—This specimen is too broken to be identified further. It represents a condor the size of a large individual of *Vultur gryphus* Linnaeus. The intercotylar prominence is very broad and low, unlike that found in either *Vultur* Linnaeus, *Gymnogyps* Lesson, or the new genus of condor being described from the Talara Tar Seeps (Campbell, in press). I am convinced the specimen represents a new genus and species, but consider the specimen too fragmentary to describe.

**Coragyps cf. atratus** (Bechstein)

Material.—Distal ends of 1 right and 1 left carpometacarpus.
Remarks.—These specimens resemble those from the Talara Tar Seeps in differing slightly from the subspecies of *C. atratus* currently resident in southwestern Ecuador and northwestern Peru.

**Cathartes aura** (Linnaeus)

Material.—One right scapula, proximal end of 1 left radius, 1 complete left tarsometatarsus, distal ends of 2 left tarsometatarsi.

Remarks.—*C. aura* is resident in southwestern Ecuador.

**ACCIPITRIDAE**

**Accipitridae genus and species**

Material.—Proximal end of 1 left femur.

Remarks.—This extinct new genus and species of large eagle is being described from the Talara Tar Seeps (Campbell, in press). Hoffstetter (1952: 40) mentioned an enormous eagle from La Carolina. The specimens he was referring to probably belong to this species.

**Geranoaetus melanoleucus** (Vieillot)

Material.—Proximal end of 1 right carpometacarpus, distal end of 1 left tibiotarsus.

Remarks.—*G. melanoleucus* is resident in western Ecuador today.

**Buteo polyosoma** (Quoy and Gaimard)

Material.—Proximal ends of 1 right and 1 left humerus, proximal end of 1 right radius, 1 complete left carpometacarpus, distal end of 1 right tibiotarsus, 2 complete left tarsometatarsi.

Remarks.—*B. polyosoma* is a common resident in southwestern Ecuador.

**Buteo hoffstetteri**, new species

Figure 2

Holotype.—Right tarsometatarsus lacking medial portion of proximal end. Uncataloged.

Paratype.—Almost complete left tarsometatarsus.

Diagnosis.—Tarsometatarsus agrees with that of *Buteo* Lacépède and differs from that of all other South American genera of Accipitridae by having those characters of the genus *Buteo* as listed by Campbell (in press).

Tarsometatarsus characterized by having: (1) internal cotylna very concave (moderately concave in *B. lineatus*, very concave in *B. polyosoma*); (2) anterior metatarsal groove very deep immediately distal to intercotylar prominence and distal to tubercle for tibialis anticus (moderately deep in both areas in *B. lineatus* and *B. polyosoma*); (3) shaft narrow abruptly distal to tubercle for tibialis anticus (narrowly gradually in *B. lineatus*, abruptly in *B. polyosoma*); (4) shaft with posterior metatarsal groove moderately deep (very deep in *B. lineatus*, moderately deep in *B. polyosoma*); (5) intertrochlear notches wide (narrow in *B. lineatus* and *B. polyosoma*); (6) internal trochlea with very prominent angular proximolateral projection (not as prominent in *B. lineatus*, very prominent in *B. polyosoma*); (7) external trochlea narrow, short, and projecting posterodistad at approximately 45° to axis of shaft (wide, long, and at approximately 85° to axis of shaft in *B. lineatus* and *B. polyosoma*); (8) shaft very slender (similar in *B. lineatus*, wider in *B. polyosoma*).

Measurements (in mm) of the holotype and paratype (in parentheses) are as follows: length, 74.6 (76.3); proximal width, broken (11.1 ± 1.0); distal width, 10.7 (12.2 ± 0.5); least width of shaft, 4.0 (4.2).

Referred Material.—Distal end of 1 left and 1 right tibiotarsus.

Characters.—Tibiotarsus characterized by having: (1) shaft only slightly concave at postero-proximal end of internal condyle (moderately to deeply concave in *B. lineatus* and *B. polyosoma*); (2) internal condyle short (long in *B. lineatus*, of moderate length in *B. polyosoma*); (3) external condyle merging gradually with shaft antero-proximally, not forming a lip (merges abruptly, forming a lip in *B. lineatus* and *B. polyosoma*). The only measurement that can be taken is the distal width (10.6 mm) of one specimen.

Etymology.—This species is named for Dr. Robert Hoffstetter of the Muséum National d'His-toire Naturelle, Paris, France, in recognition of his paleontological work in Ecuador and his role in the preservation of the La Carolina fossils.

Remarks.—Although *Buteo lineatus* does not oc-
Figure 2.—Holotype right tarsometatarsus of *Buteo hoffstetteri*, new species (uncataloged), in anterior, external, and posterior view, × 1.

Cur in South America, it was used in the diagnosis because its tarsometatarsus resembles that of *B. hoffstetteri* in being long and slender rather than short and stout. While the tarsometatarsus of *B. polysoma* is the same length as that of *B. lineatus*, it is much more robust. In all other species of *Buteo* with tarsometatarsi of similar length, the bone is even more robust than in *B. polysoma* and these species must therefore also differ from *B. hoffstetteri*.

**Circus cinereus** Vieillot

**Material.**—Proximal end of 1 left carpometacarpus, 1 complete right femur, distal end of 1 left tibiotarsus.

**Remarks.**—*C. cinereus* is resident in western Ecuador, but has not been recorded from the Santa Elena Peninsula.

**FALCONIDAE**

**Falco peregrinus** Tunstall

**Material.**—Two left scapulae, sternal end of 1 right coracid, proximal end of 2 left humeri, distal end of 1 right humerus, 1 complete left ulna, proximal end of 1 right ulna, distal ends of 2 left ulnae, proximal end of 1 right radius, 2 complete right carpometacarpi, proximal ends of 1 right and 1 left carpometacarpus, distal end of 1 right carpometacarpus, 1 complete left femur, distal end of 1 right tibiotarsus, 1 complete right tarsometatarsus, proximal ends of 2 left tarsometatarsi.

**Remarks.**—*F. peregrinus* occurs in Ecuador both as a northern and a southern migrant.

**Falco femoralis** Temminck

**Material.**—One complete left coracid, proximal end of 1 left tibiotarsus.

**Remarks.**—*F. femoralis* is resident in western Ecuador.

**Polyborus plancus** (Miller)

**Material.**—Five right and 4 left scapulae, 2 complete right and 2 complete left coracoids, humeral ends of 4 right and 1 left coracid, proximal ends of 3 right and 1 left humerus, distal ends of 1 right and 1 left humerus, 1 complete right ulna, proximal ends of 2 left ulnae, distal ends of 3 right and 2 left ulnae, 1 complete right and 3 complete left carpometacarpi, proximal ends of 2 left carpometacarpus, proximal ends of 2 left femora, distal end of 1 right femur, distal ends of 2 right and 1 left tibiotarsus, proximal end of 1 left tarsometatarsus, distal ends of 3 right and 4 left tarsometatarsi.

**Remarks.**—These specimens differ from the subspecies presently found in southwestern Ecuador and northwestern Peru in the same way as do the specimens from the Talara Tar Seeps. The complete right carpometacarpus is considerably lighter in build than the rest of the carpometacarpi from this site as well as those from the Talara Tar Seeps and is perhaps sufficiently different to warrant consideration as a distinct species. However, the great variation found in *Polyborus* makes it unwise to describe a new species without a larger sample. *P. plancus* is common in southwestern Ecuador.

**Milvago species**

**Material.**—Two left scapulae, 1 complete left carpometacarpus.

**Remarks.**—This species is being described from
the Talara Tar Seeps (Campbell, in press). No species of Milvago presently occurs west of the Andes Mountains in Peru or Ecuador.

CHARADRIIDAE

**Pluvialis dominica** (Statius Muller)

**Material.**—One complete right and 3 complete left coracoids, 8 complete right and 4 complete left humeri, distal ends of 1 right and 2 left humeri, 5 complete right and 5 complete left carpometacarpi, 2 complete left femora, proximal ends of 2 right tibiotarsi, distal ends of 3 right and 1 left tibiotarsus, 1 complete right and 5 complete left carpometacarpi, proximal ends of 1 right and 1 left tarsometatarsi, distal ends of 2 right tarsometatarsi.

**Remarks.**—As North American migrants, *P. dominica* and the following 3 species of plovers occur as seasonal visitors or residents along the coast of Ecuador.

**Pluvialis squatarola** (Linnaeus)

**Material.**—Proximal end of 1 right humerus.

**Charadrius vociferus** Linnaeus

**Material.**—Proximal end of 1 left tibiotarsus.

**Charadrius semipalmatus** Bonaparte

**Material.**—One complete left humerus, distal end of 1 left tibiotarsus.

**Oreopholus orcesi, new species**

**Figure 3**

**Holotype.**—Complete left femur. Uncataloged.

**Diagnosis.**—Femur agrees with that of *Oreopholus* and differs from that of all other genera of South American plovers (with possible exception of Zonibyx Reichenbach, Pluvianellus Gray, and Phegornis Gray, which were not available for comparison) by having: (1) head disc-shaped, projecting mediad, perpendicular to shaft, without protruding dorsad above level of flat iliac facet; (2) attachment of M. flexor ischiofemoralis elevated along anterior edge; (3) shaft with proximal two-thirds straight, and marked anteroposterior flexure in distal third, in lateral view; (4) shaft with distal end turned sharply mediad posterior to anteroproximal termination of internal condyle, in anterior view; (5) internal condyle with posteroproximal corner projecting proximad more than external condyle; (6) shaft very wide immediately proximal to condyles.

Femur differs from that of *Oreopholus ruficollis* (Wagler) by having: (1) head much smaller; (2) attachment of M. iliacus marked by sharp, high ridge along corner of shaft (ridge absent and attachment much larger in area in *O. ruficollis*); (3) shaft with greater anteroposterior flexure; (4) shaft with marked concavity at point of flexure medial to internal condyle, resulting in internal condyle terminating proximally in high, narrow ridge (not as excavated in *O. ruficollis*, but with internal condyle more rounded); (5) internal condyle deeper anteroposteriad, with posterior articular surface flat (convexity present on surface in *O. ruficollis*); (6) external condyle projecting less proximad, and at a greater angle to axis of shaft, in posterior view; (7) attachment of M. gastrocnemius, pars externa, smaller in area, more elevated, and positioned more posteriad; (8) rotular groove narrower and deeper.

Measurements (in mm) of the holotype, with
those of one specimen of *O. ruficollis* (in parentheses), are as follows: length from external condyle to iliac facet, 29.7 (30.0); width of proximal end, 5.9 (6.2); width of distal end, 6.0 (6.1); least width of shaft, 2.5 (2.4).

**Etymology.**—This species is named for Professor Gustav Orcés V., of the Escuela Politécnica Nacional, Quito, Ecuador, for his pioneering work in the natural history of Ecuador.

**Remarks.**—Although only one specimen of *O. ruficollis* was available for comparison, the majority of the characters cited above are considered unlikely to vary with a larger sample. The species of *Zonibyx*, *Pluvianellus*, and *Phegornis* are all much smaller than *O. orcesi*. In the event that the species described here does not belong in *Oreopholus*, but in one of the three genera listed above, it would still represent a new species, if based solely on size.

**SCOLOPACIDAE**

**Tringa solitaria** Wilson

**Material.**—Proximal end of 1 left tibiotarsus, proximal end of 1 left tarsometatarsus, distal end of 1 left tarsometatarsus.

**Remarks.**—As North American migrants, *T. solitaria* and the following 12 species of the family Scolopacidae occur as seasonal visitors or residents along the coast of Ecuador.

**Totanus flavipes** (Gmelin)

**Material.**—One complete right coracoid, 1 complete right and 2 complete left humeri, 1 complete right and 1 complete left carpometacarpus, shaft and proximal end of 1 right femur, distal ends of 3 right and 1 left tarsometatarsus.

**Remarks.**—I consider the osteological differences between *Tringa* and *Totanus* as sufficient to separate them at the generic level.

**Totanus melanoleucus** (Gmelin)

**Material.**—Humeral end of 1 right coracoid, 1 complete left humerus, proximal end of 1 right humerus, distal end of 1 left humerus, 1 complete left carpometacarpus.

**Catoptrophorus semipalmatus** (Gmelin)

**Material.**—Distal ends of 2 right and 1 left humerus, 3 complete right and 3 complete left carpometacarpi, proximal end of 1 left carpometacarpus, shaft and proximal end of 1 right femur, distal end of 1 right tibiotarsus, proximal end of 1 right tarsometatarsus.

**Calidris canutus** (Linnaeus)

**Material.**—One complete left coracoid, 6 complete right and 6 complete left humeri, 5 almost complete right and 5 almost complete left humeri, proximal ends of 2 right and 3 left humeri, distal ends of 4 right and 5 left humeri, 5 complete right and 3 complete left carpometacarpi, proximal ends of 2 right carpometacarpi, distal ends of 7 right and 3 left tibiotarsi.

**Erolia melanotos** (Vieillot)

**Material.**—Three complete right and 8 complete left humeri, 1 almost complete right humerus, proximal ends of 1 right and 1 left humerus, distal end of 1 left humerus, 1 complete right and 2 complete left carpometacarpi, 1 complete right femur, distal end of 1 left femur, distal end of 1 right tibiotarsus, 1 complete left tarsometatarsus.

**Remarks.**—I consider the osteological differences between *Calidris*, *Erolia*, and *Ereunetes* as sufficient to separate them as three distinct genera.

**Ereunetes pusillus** (Linnaeus)

**Material.**—One complete right and 1 complete left humerus, distal end of 1 right humerus.

**Ereunetes mauri** (Cabanis)

**Material.**—Two complete right and 4 complete left humeri, distal ends of 1 right and 3 left humeri.

**Micropalama himantopus** (Bonaparte)

**Material.**—One complete right femur.

**Bartramia longicauda** (Bechstein)

**Material.**—One complete right humerus, distal
end of 1 right humerus, 1 complete right and 1 complete left carpometacarpus, distal ends of 2 left tibiotarsi, proximal end of 1 left tarsometatarsus, distal end of 1 left tarsometatarsus.

**Numenius phaeopus** (Linnaeus)

**Material.**—Two complete left coracoids, 1 complete right carpometacarpus, distal end of 1 left tibiotarsus.

**Limosa fedoa** (Linnaeus)

**Material.**—One complete right and 1 fragmentary left carpometacarpus.

**Arenaria interpres** (Linnaeus)

**Material.**—Two complete left and 1 complete right coracoid, humeral end of 1 right coracoid, proximal end of 1 left carpometacarpus, 1 complete left femur, 1 complete left tarsometatarsus.

**Scolopacidae genus and species**

**Material.**—One complete left coracoid, 1 complete right femur.

**Remarks.**—A new genus and species of Scolopacidae is being described from the Talara Tar Seeps (Campbell, in press) on the basis of a coracoid and a tarsometatarsus. The coracoid from La Carolina agrees in all characters with the holotype coracoid, differing only in being more robust. Measurements (in mm) of the La Carolina coracoid are as follows: length, 17.3; head to scapular facet, 6.7; depth of proximal end, 4.9; least width of shaft, 2.4; length of sternal facet, 5.8.

The above femur agrees with the other elements assigned to the new genus in superficially resembling *Limnodromus* more than any other genus of scolopacid. The femur differs from that of *L. griseus* by having: (1) neck longer and much more constricted; (2) head directed more anteriorly; (3) shaft flexed anteriad at level of attachment of M. iliaca, in lateral view (straight in *L. griseus*); (4) shaft curved gradually posteriad in distal half, in lateral view (curvature more localized in *L. griseus*); (5) attachment of M. gastrocnemius, pars externa, more elevated, facing more posteriad, but lying more on corner of shaft, farther from base of external condyle; (6) external condyle more distinctly set off at base posteriorly. The internal condyle is too broken to yield any characters.

Measurements (in mm) of the femur are as follows: length from iliac facet to external condyle, 28.9; width of proximal end, 5.3; width of distal end, 4.8; least width of shaft, 2.2; depth of proximal end, 4.0.

**RECURVIROSTRIDAE**

**Himantopus mexicanus** (Statius Müller)

**Material.**—One complete right coracoid.

**Remarks.**—*H. mexicanus* is generally distributed along the coast of Ecuador.

**PHALAROPODIDAE**

**Phalaropus fulicarius** (Linnaeus)

**Material.**—One complete right humerus, shaft and distal ends of 2 right humeri, 2 complete left carpometacarpi.

**Remarks.**—As North American migrants, *P. fulicarius* and the following two species of phalaropes occur in western Ecuador as seasonal visitors or residents.

**Lobipes lobatus** (Linnaeus)

**Material.**—Two complete right and 6 complete left humeri, 1 almost complete left humerus, 1 complete right and 4 complete left coracoids, proximal ends of 1 right and 1 left humerus, distal ends of 1 right and 2 left humeri, 2 complete right and 2 complete left carpometacarpi, proximal end of 1 right carpometacarpus, 1 complete right and 3 complete left tarsometatarsi.

**Steganopus tricolor** Vieillot

**Material.**—Five complete right and 4 complete left coracoids, 19 complete right and 20 complete left humeri, 3 almost complete right and 2 almost complete left humeri, proximal ends of 1 right and 2 left humeri, distal ends of 3 right and 7 left humeri, 5 complete right and 9 complete left carpo-
metacarpi, 2 complete right and 2 complete left femora, distal ends of 6 right and 5 left tibiotarsi, 2 complete right tarsometatarsi, 1 almost complete right and 1 almost complete left tarsometatarsus, proximal ends of 1 right and 1 left tarsometatarsus, distal end of 1 right tarsometatarsus.

**Steganopus species**

**Material.**—One complete right and 1 complete left humerus, distal end of 1 left humerus, 1 complete right tarsometatarsus.

**Remarks.**—This species is being described from the Talara Tar Seeps on the basis of a complete femur (Campbell, in press). As the above elements differ significantly from *S. tricolor* they are here referred to the new species.

The humerus of the new *Steganopus* differs from that of *S. tricolor* by having: (1) median crest larger, less excavated; (2) internal tuberosity larger, more rounded, directed less dorsad, and projecting less anconally; (3) head not undercut by capital shaft groove; (4) attachment of M. procapulohumeralis positioned closer to base of median crest; (5) attachment of M. triceps, external head, larger dorsoventrad and bordered ventrally by distinct lip of bicipital crest, resulting in a larger bicipital crest; (6) deltoid crest longer, thicker, and higher; (7) impression of M. brachialis anticus shallower, not as deeply inset at the edges; (8) attachment of anterior articular liga¬ment shorter and higher; (9) attachment of M. pronator brevis positioned more distad; (10) internal and external condyles longer; (11) shaft much larger.

The tarsometatarsus of the new species differs from that of *S. tricolor* by having: (1) internal trochlea larger, but projecting less posteriorly; (2) internal trochlea less excavated adjacent to middle trochlea; (3) middle trochlea more rounded posteriorly in medial view; (4) internal trochlea positioned close to, and directed more toward, midline of shaft. The proximal ends of the tarsometatarsi of *Steganopus* apparently do not ossify completely except in older individuals. Although the present specimen showed only a slight indication of pitting, no reliable characters could be obtained from the proximal end.

For measurements of the humerus and tarsometatarsus of the two species of *Steganopus*, see Table 1.

**BURHINIDAE**

**Burhinus superciliaris** (Tschudi)

**Material.**—Proximal end of 1 left humerus, distal ends of 1 right and 3 left humeri, 1 complete right and 3 complete left coracoids, proximal end of 1 left coracoid, distal end of 1 right scapula, distal end of 1 right ulna, 1 complete right and 2

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<th>Character and measurement</th>
<th>Steganopus species</th>
<th>Steganopus tricolor Recent</th>
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<td>7.3</td>
<td>7.4</td>
<td>7.3</td>
</tr>
<tr>
<td>N</td>
<td>2</td>
<td>8</td>
<td>35</td>
</tr>
<tr>
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<td>5.1</td>
<td>4.5-5.6</td>
<td>4.7-5.6</td>
</tr>
<tr>
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<td>5.1</td>
</tr>
<tr>
<td>N</td>
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<td>8</td>
<td>35</td>
</tr>
<tr>
<td>Least width of shaft</td>
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<td>2-2.3</td>
<td>2-2.6</td>
</tr>
<tr>
<td>M</td>
<td>2.4</td>
<td>2.2</td>
<td>2.3</td>
</tr>
<tr>
<td>N</td>
<td>2</td>
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</tr>
<tr>
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<td></td>
<td></td>
</tr>
<tr>
<td>Length</td>
<td>30.6</td>
<td>31-34.7</td>
<td>32.8-34.5</td>
</tr>
<tr>
<td>OR</td>
<td>32</td>
<td>32.2</td>
<td>33.6</td>
</tr>
<tr>
<td>M</td>
<td>32</td>
<td>32.2</td>
<td>33.6</td>
</tr>
<tr>
<td>N</td>
<td>1</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>Width of proximal end</td>
<td>3.5</td>
<td>3.2-3.7</td>
<td>3.5-3.8</td>
</tr>
<tr>
<td>M</td>
<td>3.5</td>
<td>3.4</td>
<td>3.6</td>
</tr>
<tr>
<td>N</td>
<td>1</td>
<td>8</td>
<td>4</td>
</tr>
<tr>
<td>Width of distal end</td>
<td>3.2</td>
<td>3-3.3</td>
<td>3.3-3.5</td>
</tr>
<tr>
<td>M</td>
<td>3.2</td>
<td>3.2</td>
<td>3.4</td>
</tr>
<tr>
<td>N</td>
<td>1</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>Least width of shaft</td>
<td>1.2</td>
<td>1.2-1.3</td>
<td>1.3</td>
</tr>
<tr>
<td>M</td>
<td>1.2</td>
<td>1.2</td>
<td>1.3</td>
</tr>
<tr>
<td>N</td>
<td>1</td>
<td>8</td>
<td>4</td>
</tr>
</tbody>
</table>
complete left carpometacarpi, 1 complete left femur, distal end of 1 left tibiotarsus, distal end of 1 right tarsometatarsus.

Remarks.—B. superciliaris is common in the semi-arid region of Ecuador.

THINOCORIDAE

Thinocorus rumicivorus Eschscholtz

Material.—One complete right and 1 complete left humerus, 1 complete right and 1 complete left carpometacarpus.

Remarks.—T. rumicivorus is resident in the arid regions of Ecuador.

LARIDAE

Larus pipixcan Wagler

Material.—One complete right carpometacarpus, proximal end of 1 right carpometacarpus.

Remarks.—A North American migrant, L. pipixcan is a seasonal resident of Ecuador.

Larosterna inca (Lesson)

Material.—Humeral end of 1 left coracoid.

Remarks.—A small, shallow attachment of Lig. humerocoracoideum anterius superius, located on the anteroventral corner of the head of the coracoid, is characteristic of Larosterna. This is the first fossil record of the genus. L. inca is found along the coast of southern Ecuador.

COLUMBIDAE

Zenaida auriculata (Des Murs)

Material.—Two right and 2 left scapulae, 1 complete right and 8 complete left coracoids, humeral ends of 1 right and 1 left coracoid, 1 complete right and 6 complete left humeri, proximal ends of 2 right and 3 left humeri, distal ends of 1 right and 1 left humerus, 4 complete right and 5 complete left ulnae, 2 complete right and 2 complete left radii, 5 complete right and 6 complete left carpometacarpi, 1 complete right and 1 complete left femur, 1 complete right and 1 complete left tibiotarsus, distal end of 1 right tibiotarsus, 4 complete right and 1 complete left tarsometatarsus, shaft and proximal end of 1 left tarsometatarsus.

Remarks.—Z. auriculata is resident in western Ecuador.

Zenaida asiatica (Linnaeus)

Material.—Proximal end of 1 left carpometacarpus.

Remarks.—Z. asiatica is resident in southwestern Ecuador.

Columbina cruziana (Knip and Prevost)

Material.—One complete right humerus.

Remarks.—C. cruziana is common in southwestern Ecuador.

PSITTACIDAE

Aratinga roosevelti (Spillman)

Material.—Portions of 11 crania, 12 maxillaries, 14 mandibles, 24 right and 17 left scapulae, 35 complete right and 46 complete left coracoids, humeral ends of 2 left coracoids, 25 complete right and 24 complete left humeri, proximal end of 1 right humerus, distal ends of 3 left humeri, 55 complete right and 56 complete left ulnae, proximal ends of 1 right and 1 left ulna, distal end of 1 left ulna, 30 complete right and 28 complete left radii, proximal ends of 5 right and 1 left radius, distal ends of 2 right and 2 left radii, 85 complete right and 71 complete left carpometacarpi, proximal ends of 3 right and 2 left carpometacarpi, shafts of 2 left carpometacarpi, 29 complete right and 22 complete left femora, distal ends of 1 right and 1 left femur, 36 complete right and 24 complete left tibiotarsus, proximal ends of 1 right and 1 left tibiotarsus, distal ends of 4 right and 1 left tibiotarsus, 21 complete right and 19 complete left tarsometatarsus, proximal ends of 2 left tarsometatarsi, distal ends of 3 left tarsometatarsi.

Remarks.—Spillman (1942) reported a large collection of parrot fossils from a site on the Santa Elena Peninsula near La Carolina. He placed this material in a new genus and species, Protoconurus roosevelti. From his descriptions and illustrations
it is reasonably certain that the present material belongs to the same species. I could not detect sufficient differences between this fossil material and Recent specimens of the genus *Aratinga* to warrant the placement of the former in a different genus. Therefore, I suggest that *Protoconurus* be synonymized with *Aratinga*.

It is not possible at this time to say whether the fossil material represents an extinct species, as Spillman believed, or an extant species. This cannot be determined until a satisfactory series of comparative material is available. The fossil material is slightly larger than the one available specimen of *A. erythrogenys* from southwestern Ecuador and differs from that species in many ways.

One coracoid of those listed above is significantly smaller than the others and may represent a different species.

*Aratinga* species

Material.—One complete left coracoid, distal end of 1 right humerus.

Remarks.—These two specimens appear to represent a species of *Aratinga*, but may possibly represent another genus, as not all genera of Ecuadorian parrots were available for comparison. They are considerably larger than the corresponding elements of *A. erythrogenys*.

**TYTONIDAE**

*Tyto alba* (Scopoli)

Material.—One complete right coracoid.

Remarks.—*T. alba* is resident in western Ecuador.

**STRIGIDAE**

*Speotyto cunicularia* Gloger

Material.—One complete right and 1 complete left carpometacarpus, distal ends of 2 right tibiotarsi, distal end of 1 left tarsometatarsus.

Remarks.—*S. cunicularia* is resident in western Ecuador.

Discussion

Ecological Considerations.—La Carolina is located in the most arid region of Ecuador. Vegetation is very sparse, except in the larger dry river valleys. Ecological and climatological conditions at the site are very similar to those found at the Talara Tar Seeps, except that La Carolina is much closer to the coastline and receives slightly more rainfall. The major difference between the arid region of Ecuador and that of coastal Peru is in the greater area of the latter. Whereas the Peruvian coastal desert is thousands of kilometers long, the Ecuadorian desert is essentially limited to the Santa Elena Peninsula (Figure 1). The transition from barren desert to heavy forest is very abrupt, in some places occurring over only a few kilometers.

The principal references concerning the modern avifauna of southwestern Ecuador are the works of Chapman (1926) and Marchant (1958, 1959, 1960). A point stressed by both authors is the rapidity with which the arid and semi-arid regions are colonized by birds from the adjacent forests following the rare periods of rainfall. Marchant also documents how large numbers of birds are attracted to standing bodies of fresh water on the peninsula, such as an artificial lake.

The importance of the La Carolina avifauna in interpreting the paleoecology of southwestern Ecuador is difficult to assess. This is because the desert region is so small, the transition to forest so sharp, and the speed of colonization under the proper conditions so rapid, that only a few years of heavy rainfall are needed to change the desert into a lush savanna with seasonally flowing rivers. Such rivers would in turn form wide deltas in the low-lying, flat coastal zone. This type of environment undoubtedly attracts birds from the surrounding areas and provides excellent stop-over points for the North American migrants that make up over 43 percent of the paleoavifauna.

A climatological phenomenon known as El Niño (Chapman, 1926; Murphy, 1926) periodically brings heavy rains to southwestern Ecuador and northern coastal Peru. This supposedly occurs every seven years, but in reality the phenomenon may take place in any year. When rains do come to southwestern Ecuador they occur during the northern hemisphere winter. This increases the probability of migrant species encountering favor-
able conditions during passage to and from their wintering grounds.

If the avifauna of the Talara Tar Seeps were not known, one would probably interpret the avifauna of La Carolina, and possibly the mammalian fauna as well, as having been deposited during El Niño years. However, the Talara avifauna strongly suggests that the entire climatological regime of northern coastal Peru, and thus by inference that of southwestern Ecuador, was quite different from that found today. Instead of desert, the area was probably a lush savanna or, in the case of the Santa Elena Peninsula, more probably a tropical dry deciduous forest. Archeological work on the Santa Elena Peninsula also indicates a moister climatic regime in the past (Meggers, Evans, and Estrada, 1965; Richardson, 1973). The causes of the present dry climate, as opposed to the wetter climate present during glacial periods, are the subject of another paper currently in preparation.

Comparison with the Avifauna of the Talara Tar Seeps.—Of the 53 species from La Carolina, 38 (72%) were also found at the Talara Tar Seeps (Table 2). This includes 7 extinct species common to both sites that were first recorded from the Talara Tar Seeps. The high degree of similarity between the paleoavifaunas of the two localities, as well as that between the modern avifaunas, confirms that there must have been a great deal of interchange between coastal Ecuador and coastal Peru. A similar situation is observed between the mammalian faunas of the two sites (Lemon and Churcher, 1961).

The straight line distance between La Carolina and the Talara Tar Seeps is approximately 273 kilometers. At the present time the Gulf of Guayaquil separates the arid region of Ecuador from that of northern Peru. During glacial periods, however, the lowered sea level exposed as dry land the area currently covered by the Gulf of Guayaquil, allowing easy movement between the two areas. The formation of the Gulf of Guayaquil at the end of the Pleistocene resulted in the isolation of the avifauna of the arid Santa Elena Peninsula from that of coastal Peru. Consequently, a great deal of subspeciation has occurred between the two areas (Chapman, 1926).

The differences observed in the composition of the two paleoavifaunas can be attributed to two major causes—the means of entrapment and the location relative to the coastline. The active tar seeps at Talara attracted large numbers of scavenger species that came to feed upon other animals, including large mammals, trapped in the tar. The drawing power of La Carolina in this regard must have been weak. As the Talara site is located approximately 32 kilometers inland, it has a higher percentage of terrestrial species and fewer strictly marine species that do not venture inland. The opposite is true of La Carolina which was on, or very near, the coastline.

The similarities between the avian and mammalian faunas of La Carolina and the Talara Tar Seeps, particularly in the extinct species, suggest that material was being deposited at the two sites contemporaneously. The Talara site has been dated at c. 13,900 B.P. (Churcher, 1966). If the age of La Carolina differs significantly from that of the Talara Tar Seeps, I believe it will prove to be younger rather than older, for climatological reasons to be discussed in a later publication.

Entrapment of the Birds at La Carolina.—How birds became trapped, or their remains concentrated, at La Carolina is not clear. The undisturbed matrix available with the collection was extremely rich in avian specimens, and contained only a few fragmentary mammalian bones. At other nearby sites, however, Edmund (pers. comm.) found only mammalian and no avian material.

None of the fossil material I studied shows any large concentration of pitch, although there are some indications of its presence in the matrix surrounding some specimens. The absence of heavy pitch concentrations on the bones indicates that the birds were not trapped in active tar seeps, as at the Talara Tar Seeps and Rancho La Brea. It is quite possible, however, that oil seeps may have coated the water with a surface layer of oil and that upon landing in this the feathers of the birds would become oil-soaked, preventing further flight and resulting in their drowning. As the bodies decomposed, the bones would be freed from the oil and drop to settle in the soft mud. Such a film of oil would also help to explain the lack of mammalian specimens in the concentration of avian remains. Mammals are better able to sense and avoid oil-covered water than are birds alighting from flight. For larger mammals, passage through oil covered water would be no more than an unpleasant inconvenience.
Table 2.—Fifty-three nonpasserine species from La Carolina, Ecuador

<table>
<thead>
<tr>
<th>Species</th>
<th>No. of Specimens</th>
<th>Minimum No. of Individuals</th>
<th>Species</th>
<th>No. of Specimens</th>
<th>Minimum No. of Individuals</th>
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<td>1</td>
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<tr>
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<td>Bartramia longicauda</td>
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<td>2</td>
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<td>1</td>
<td>Limosa feda</td>
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<td><em>Arenaria interpres</em></td>
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<tr>
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<td>3</td>
<td><em>Scolopacidae</em> genus and species</td>
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<tr>
<td><em>Accipitridae</em> genus and species</td>
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<td>1</td>
<td><em>Himantopus mexicanus</em></td>
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<td>1</td>
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<tr>
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<td>1</td>
<td><em>Phalaropus fulicarius</em></td>
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<td>3</td>
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<tr>
<td><em>Buteo polyosoma</em></td>
<td>7</td>
<td>2</td>
<td><em>Lobipes lobatus</em></td>
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<td>8</td>
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<tr>
<td>Buteo hoffstetteri, new species</td>
<td>4</td>
<td>1</td>
<td><em>Steaganopus</em> species</td>
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<td>Steganopus tricolor</td>
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<td><em>Burhinus supercilialis</em></td>
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<td><em>Polyborus plancus</em></td>
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<td>1</td>
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<td>Aratinga species</td>
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<tr>
<td><em>Tringa solitaria</em></td>
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<td><em>Tyto alba</em></td>
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<td><em>Catoptrophorus semipalmatus</em></td>
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<td>Calidris canutus</td>
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| Total | | | Total | | |
| 1611 | 284 |

*Also recorded from the Talara Tar Seeps, Peru.

Perhaps the most remarkable aspect of the avifauna from La Carolina is the tremendous number of specimens of parrots, making up approximately 44 percent of the entire collection. Parrots must have been attracted to the site in large flocks, perhaps for water, and subsequently trapped by the oil. From such large flocks of parrots one could reason either that water was scarce and La Carolina was the only watering site available, the parrots thus having to travel long distances to reach it, or water may have been abundant and La Carolina close to the roosting or feeding areas of the parrots. If the latter were true, it would indicate the presence of forest on the presently barren peninsula.

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Murphy, R. C.

Richardson, J. B.

Spillman, F.
The Great Predatory Birds of the Pleistocene of Cuba

Oscar Arredondo
translated and amended by Storrs L. Olson

ABSTRACT

Recent paleontological investigations in Cuba have shown that the island was formerly inhabited by large populations of rodents and edentates. Based on discoveries of giant fossil raptorial birds (Strigiformes and Accipitriformes) in Cuba and other Antillean islands, the theory is put forth here that these predators were the principal agents in controlling the abundance of these native mammals. Details are given of the discovery in Cuba of the gigantic owl *Ornimegalonyx*, two species of giant barn owls (*Tyto*), a giant species of eagle (*Aquila borrasi*), and a vulture (*Antillovultur*) similar in size to the Andean Condor (*Vultur gryphus*). The relationships and possible origins of gigantism in these birds are discussed. Descriptions, geographic distribution, tables of measurements, and a list of the extinct faunas found in each of the type-localities are offered for each species.

Se expone la tesis, basada en los últimos descubrimientos realizados en Cuba y otras Antillas sobre aves fósiles gigantes de los órdenes Strigiformes y Accipitriformes, de que dichos taxones constituían el elemento faunístico primordial que controlara con su acción predatora a roedores y desdentados que en estas islas existían, manteniendo así el equilibrio biológico necesario. Se ofrecen datos concretos probatorios de la extremada abundancia en las Antillas de poblaciones de roedores y desdentados. Se dan detalles del descubrimiento en Cuba del gigantesco bubo *Ornimegalonyx*, de dos especies gigantes de lechuzas del género *Tyto*, de una especie de águila gigante y de un buitre similar en talla a *Vultur gryphus*. Se plantea una discusión sobre el motivo que pudo originar el gigantismo en estas aves. Se ofrecen descripciones específicas de todas estas grandes aves predatoras, incluyéndose, distribución geográfica, tablas de medidas y una relación de la fauna extinguida hallada asociada en las localidades típicas.

Introduction

Recent studies of fossil material from numerous caves in Cuba have convincingly demonstrated the former extraordinary abundance of endemic mammals there. These include rodents of the genera *Heteropsomys*, *Capromys* (including *Geocapromys*), and *Macrocopromys*; ground sloths (*Edeontata*) of the genera *Cubanocnus*, *Miocnus*, *Mesocnus*, and *Megalocnus*; and insectivores of the genera *Solenodon* and *Nesophontes*. These genera were represented throughout the island by species very numerous in terms of individuals. From such caves as Paredones and El Túnel in the province of Habana it is no exaggeration to say that tens of thousands of mandibles of *Capromys pleistocenicus* can be extracted from a single small chamber, although most of these specimens are poorly preserved. An examination of only a part of the fossil material taken from a small cave in Cayo Salinas, a few miles east-southeast of Caibarién, Las Villas, yielded the remains of over 200 individuals of the edentate genus *Mesocnus* (Acevedo, Arredondo, and Gonzalez, 1975). Further confirming the former abundance of native mammals is the widespread occurrence in Cuba of remains of blood-eating vampire bats of the genus *Desmodus* (Koopman, 1958; Arredondo, 1958b; Woloszyn and Mayo, 1974). These bats would necessarily have required numerous, large, warm-blooded mammals to sustain them. Similar abundant faunas of large rodents and ground sloths are also known from cave deposits in Hispaniola and Puerto Rico.
Obviously, some natural regulator must have acted to maintain a biological equilibrium between these prolific herbivorous mammals and the vegetation on which they fed. In the Antilles, however, there are virtually no native carnivorous mammals. Exceptions are *Cubacyon transversidens* (Arredondo and Varona, 1974), a canid known from a single fragment of maxilla found in association with extinct vertebrates in a cave in the province of Habana, and the extant populations of *Procyon* known from the Bahamas, Barbados, and Guadeloupe, some or all of which may have been introduced to these islands by man (Varona, 1974; Arredondo and Varona, 1974). A mandible found in an Indian midden in Camagiiey, Cuba, and attributed to *Procyon lotor* by the naturalist Andres Poey (Harrington, 1935), may similarly have pertained to an introduced animal; the record is dubious in any case, since the specimen can no longer be found. Regardless, small procyonids could not have been an influential factor in the control of the endemic Antillean rodents and edentates.

The numerous and dense deposits of smaller vertebrates (Table 1) frequently found in Cuban caves abound with remains of the rodents *Capromys pleistocenicus, Capromys nanus, Heteropsemys torrei,* and *H. offella;* the insectivores *Nesonophontes micrus* and *N. major;* several genera of bats; passeriform birds, particularly of the genera *Mimocichla, Quiscalus* and *Dives;* as well as non-passerines such as *Crotophaga* and *Glaucidium.* These remains are certainly attributable to the actions of medium-sized owls still existing in the island such as *Tyto alba* and *Asio stygius,* and perhaps to others now extinct, such as *Pulsatrix arredondoi* (Brodkorb, 1969).

The most outstanding accumulations of bones, however, are those of the larger mammals (Table 1) such as *Capromys pilorides, C. columbianus, Macrocapromys acevedo,* and occasional examples of *Cubanocnus gliriformis,* as well as large birds of various orders. Such remains suggest that many larger predators were once active in Cuba.

Much of the accumulation of vertebrate remains in cave deposits in Haiti was attributed to the depredations of the extinct giant barn owl *Tyto ostologa,* first described by Wetmore (1922). Another giant barn owl, *Tyto pollens* Wetmore (1937), first reported from Great Exuma and later from New Providence Island (Brodkorb, 1959), must have been an effective predator of *Capromys* in the Bahamas, as no doubt were the extinct diurnal raptores (Accipitridae) of the genera *Calohierax* and *Titanohierax* (Wetmore, 1937). The presence of such large raptores elsewhere in the Antilles suggested that the Cuban cave deposits might be attributable to similar avian predators. This was vividly confirmed by the discovery in Cuba of remains of five species of predatory birds of truly tremendous proportions. These birds occupied the niches which in continental areas are usually filled by various groups of carnivorous mammals. The absence of mammalian competitors, combined with a superabundance of large prey, are no doubt the principal factors contributing to the gigantism observed in these birds. The larger forms may have functioned particularly to keep the populations of edentates in check.

**Brief History of Certain Discoveries**

On 2 January 1954, the remains of a gigantic owl, the largest known, were discovered for the first time in the depths of a large cavern known as Pio Domingo Cave, located in the Sierra de Sumidero, opposite Pica-Pica Valley in Pinar del Río. These bones (Figure 5) were found in place, fastened to the calcareous surface of the floor by travertine, just as were those of the various edentates encountered in the immediate vicinity. This owl was described under the name *Ornimegalonyx oteroi* (Arredondo, 1958a) and was at first erroneously regarded as a member of the Phorusrhacidae, an extinct family of flightless South American carnivorous birds. Dr. Byran Patterson, who studied some of these remains, afterwards informed me (pers. comm.) of his belief that they actually pertained to a great owl, two times larger in linear dimensions than *Tyto ostologa* of Haiti. Brodkorb (1961), recognizing the validity of the nomenclature proposed in 1958, established that the species belonged in the family Strigidae, where he maintained it in later publications (Brodkorb, 1969, 1971). Addi-
Table 1.—Associated fauna found in the type-localities of the large extinct species of Cuban birds of prey

<table>
<thead>
<tr>
<th>Species</th>
<th>Ornimegalonyx oteroi</th>
<th>Tyto riveroi</th>
<th>Tyto noeli and Aquila borrasii</th>
<th>Antillovultur varonai</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cueva de Pío Domingo, Pinar del Río</td>
<td>Cueva de Bellamar, Matanzas</td>
<td>Cueva del Túnel, Habana</td>
<td>Cueva de Paredones, Habana</td>
</tr>
<tr>
<td>AMPHIBIA</td>
<td>Bufo sp.</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td>REPTILIA</td>
<td>Iguanidae gen. and sp. indet.</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Epicrates cf. angulifer</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Geochelone cubensis</td>
<td>-</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Crocodylus sp.</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td>AVES</td>
<td>Cathartes aura</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Antillovultur varonai</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Vulturidae gen. and sp. indet.</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Aquila borrasii</td>
<td>?</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Grus cubensis</td>
<td>?</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Ornimegalonyx oteroi</td>
<td>X</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Ornimegalonyx sp.</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Pulsatrix arredondoi</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Gymnoglaux sp.</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Tyto alba</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Tyto noeli</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Tyto riveroi</td>
<td>-</td>
<td>X</td>
<td>-</td>
</tr>
<tr>
<td>MAMMALIA</td>
<td>Solenodon cf. cubanus</td>
<td>X</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Nesophontes micrus</td>
<td>X</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Nesophontes major</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Phyllonycteris poeyi</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Natalus lepidus</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Cubacyon transversidens</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Acratocnus sp.</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Micnus antillensis</td>
<td>X</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Cubanocnus gliriformis</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Mesocnus torrei</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Mesocnus browni</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Neomesocnus brevirostris</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Megalocnus rodens</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Megalocnus sp.</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Capromys pilorides</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Capromys frechensilis</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Capromys nanus</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Capromys columbianus</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Capromys pleistocenicus</td>
<td>-</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Capromys sp. A</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Capromys sp. B</td>
<td>X</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Capromys sp. C</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Capromys sp. D</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Macrocapromys acevedo</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Heteropsomys torrei</td>
<td>X</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>Heteropsomys offella</td>
<td>X</td>
<td>-</td>
<td>X</td>
</tr>
</tbody>
</table>

X = present; ? = uncertain; - = not present.
tional fossils of this giant owl were later found in various caves in the provinces of Habana, Matanzas, Las Villas, and Camagüey, as well as in the Isle of Pines. It is possible that two unrecognized species of Ornimegalonyx may exist among these remains; one of these seems to be somewhat smaller than O. oteroi, while the other is larger. Considering the enormous size of this owl, particularly of its claws, it is quite conceivable that it could have made victims of juvenile edentates, notwithstanding the fact that Ornimegalonyx appears to have been incapable of flight.

In July 1954, explorations in the Cueva de Paredones in San Antonio de los Baños, revealed for the first time the fossil bones of an eagle larger than any of the living species of the family Accipitridae. Other bones of it were found a few years later in the Cueva del Túnel, La Salud, Habana. This eagle was named Aquila borrasii (Arredondo, 1970). According to its tarsometatarsus, it was very similar in size and morphology to the recently described species Garganoaetus freudenthali from the upper Miocene of Italy (Ballmann, 1973).

A fossil vulture from Cueva de Paredones was recently described as a new genus and species, Antillovultur varonai Arredondo (1971), and was the size of an Andean Condor. Various bones found in a cave in Habana and now under study, possibly indicate another large species of vulture.

Two species of giant barn owls of the genus Tyto were discovered a little later (Arredondo, 1972a, 1972b). One of these, Tyto noeli, founded on abundant bones from two caves in Habana, was similar in size to Tyto ostologa of Haiti. The other species, Tyto riveroi, based on the distal portion of a tarsometatarsus from Cueva de Bellamar, Matanzas, was truly gigantic, being larger than any of the fossil or living species of the genus. It is the one strigiform that most closely approaches the size of Ornimegalonyx oteroi, and like that species could also have captured small edentates. The eminent paleornithologist Alexander Wetmore (1959) was the first to report fossil remains of large barn owls from Cuba, but these were not named.

The discovery of three species of tytonids in the upper Miocene of Italy (Ballmann, 1973), one of which, Tyto robusta, is equal in size to T. noeli and T. ostologa, and the other, Tyto gigantea, being only slightly smaller than T. riveroi, partly contradicts the hypothesis that gigantism in Antillean barn owls is attributable to insular evolution from smaller species that responded to the great abundance of food and the lack of competition from carnivorous mammals. The genus Tyto evidently had already evolved giant species in Europe, millions of years before the beginning of the Pleistocene. The following conclusions could therefore be drawn: (1) either the giant Antillean barn owls evolved in parallel with those of Europe, arriving through convergence at species of approximately the same size, or (2) the Antillean forms are descended from Tertiary European forms that established themselves in North America and colonized the Antilles before or during the Pleistocene. Against this last suggestion is the absence on the American continent of giant species of Tyto. Ornimegalonyx is truly exceptional for its extremely large size. It appears to have evolved its gigantism in Cuba from some remote smaller ancestor. An affinity of Ornimegalonyx with any of the living genera of large continental owls is not clearly evident and its relationships may lie closer to some extinct form rather than with any presently living.

The study of the origins, evolution, and paleoecology of the giant raptorial birds of the Antilles is of great interest and significance to our understanding of the environment and evolution of many of the terrestrial vertebrates of those islands. It is hoped that this summary of what is known of the Cuban birds will aid in that understanding.

Abbreviations used are as follows: Academia de Ciencias de Cuba (ACC), Departamento de Paleontologia de la Universidad de la Habana (DPUH), Museo del Grupo de Exploraciones Científicas "Pedro Borrás Astorga" (GEC), Museo de Comparative Zoology (MCZ), Museo Felipe Poy de la Academia de Ciencias de Cuba (MFP), Museo Montané de la Universidad de la Habana (MMUH), personal collection of Oscar Arredondo (OA), Sociedad Espeleológica de Cuba (SEC).
Order ACCIPITRIFORMES

Family VULTURIDAE

Genus Antillovultur Arredondo, 1971

Antillovultur varonai Arredondo, 1971

Holotype.—Proximal portion of left tarsometatarsus, DPUH 1254.

Type-Locality.—Cueva de Paredones, San Antonio de los Baños, Habana, Cuba.

Age.—Late Pleistocene.

Other Material.—GEC (unnumbered), distal portion of left humerus; OA 847, external trochlea of left tarsometatarsus; OA 848, body of seventh cervical vertebra. All specimens from the type-locality.

Description.—The type (Figure 1a, b) is a proximal portion of a tarsometatarsus, 42.5 mm in length, lacking the proximal articulating surface, hypotarsus, and slightly more than half the distal portion of the bone. The estimated total length is 141 mm, or about equal to that of Vultur gryphus and longer and slightly more robust than in Gymnogyps californianus (Figure 2, Table 2). From these two species and Cathartes aura it differs in having the groove in the anterior face of the bone narrower and deeper and the internal tubercle for M. tibialis anticus more expanded. In anterior view, the surface of the shaft between the internal border and the groove is notably thick and rounded, whereas in Vultur, Cathartes, and Teratornis it is narrower and sharp-edged. The surface of the shaft delimited by the external border and the groove is more slender than the internal ridge, contrary to the condition in the other genera mentioned. The shaft in medial view is proportionately more slender than in Vultur or Cathartes. These characters are considered to be of generic value.

The distance between the proximal border of the larger tubercle for M. tibialis anticus and the proximal extremity of the anterior groove is greater than in Vultur or Teratornis and equal to that in Gymnogyps. Antillovultur has an additional two proximal foraminae situated above the usual two. Vultur similarly possesses another aperture above the lateral proximal foramen but lacks the medial proximal foramen of Antillovultur.

Figure 1.—Specimens of Antillovultur varonai, Cueva de Paredones: a, holotype fragmentary proximal end of left tarsometatarsus (DPUH 1254), anterior view; b, same, posterior view; c, left humerus lacking proximal end (GEC unnumbered), palmar view. (Natural size.)
Table 2.—Measurements (mm) of the tarsometatarsus of *Antillovultur varonai* compared with other large New World vultures

<table>
<thead>
<tr>
<th>Character</th>
<th><em>Antillovultur varonai</em></th>
<th><em>Vultur gryphus</em></th>
<th><em>Gymnogyps californianus</em></th>
<th><em>Teratornis merriami</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Maximum length</td>
<td>141</td>
<td>141</td>
<td>124.2</td>
<td>132</td>
</tr>
<tr>
<td>Greatest proximal width</td>
<td>32.2</td>
<td>30.2</td>
<td>27</td>
<td>28</td>
</tr>
<tr>
<td>Least width of shaft (at break)</td>
<td>17</td>
<td>17</td>
<td>16.8</td>
<td>15</td>
</tr>
<tr>
<td>Length of outer trochlea from the angle of the middle trochlea</td>
<td>9</td>
<td>9.5</td>
<td>6</td>
<td>9</td>
</tr>
<tr>
<td>Greatest width of outer trochlea</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Greatest width across trochlea</td>
<td>35*</td>
<td>35.4</td>
<td>30</td>
<td>29.5</td>
</tr>
</tbody>
</table>

* Estimated.

*Cathartes aura* has only two proximal foramina. *Teratornis* differs in having three great united foramina arranged so as to form a kind of circle. In *Antillovultur* the medial tubercle for M. tibialis anticus is better developed than in either *Vultur, Gymnogyps,* or *Cathartes,* and is situated over a slight protruberance on the internal border of the anterior groove. In posterior view the surface of the bone from the lateral foramen to the external border is flat and does not slope downward as in *Vultur.* On the opposite side, the surface from the medial foramen to the external border is very depressed compared to that in *Vultur.* An external trochlea from a left tarsometatarsus of this species is similar in size and shape to that of *Vultur.*

**Figure 2.**—Comparison of tarsometatarsi of New World vultures: *a,* *Gymnogyps californianus; b,* *Vultur gryphus; c,* *Antillovultur varonai; d,* *Teratornis merriami* (based on Wetmore, 1931). (Half natural size.)

A seventh cervical vertebra attributed to *Antillovultur* consists of the complete body lacking all of the processes. It is similar to vertebrae of *Vultur* in size and morphology.

A distal portion of a left humerus (Figure 1c) is from a specimen very similar in size to *Vultur.* This fragment has a length of 184 mm from the distal end to a point a little beyond the protruberance at the distal extremity of the deltoid crest. Taking this protruberance as a point of reference, the complete length of the bone can be estimated as 265 mm, which is about 20 mm less than in the specimens of *Vultur* compared. The least width of the shaft is 20 mm (21.7 in *Vultur*); the maximum distal width is 45.8 mm (54.3 in *Vultur*). The bone, although almost as large as that of *Vultur,* is more slender. The distal protruberance of the deltoid crest is at the same level as in *Vultur.* *Antillovultur* differs from *Vultur* in having the ectepicondylar prominence less pronounced, recalling that of *Cathartes.* The internal and external condyles are slightly smaller than in *Vultur* but with the distal borders more prominent. The internal condyle has a smooth and extensive depression on the entepicondylar side, which is present in *Cathartes* but absent in *Vultur.* The attachment of the anterior articular ligament is as large as in *Vultur,* but situated closer to the internal condyle, the space between them being less than in *Vultur* and more similar to *Cathartes.* The foramen located in this space is isolated from the brachial depression by a ridge that is absent in *Vultur* but somewhat evident in *Cathartes.* The brachial depression is deeper and more pronounced than in *Vultur,* particularly in the proximal region, but is notably less expanded than in either *Vultur* or *Cathartes.* The olecranal fossa is similar to that of *Vultur* and
less dilated than in Cathartes. The tricipital grooves are similar to those of Vultur but the external one is somewhat deeper and appears like that of Cathartes.

**Family ACCIPITRIDAE**

**Genus Aquila Brisson**

*Aquila borrasi* Arredondo, 1970

**Holotype.**—Left tarsometatarsus lacking trochleae, DPUH 1250.

**Type-Locality.**—Cueva del Túnel, La Salud, Habana, Cuba.

**Other Localities.**—Cueva de Paredones, San Antonio de los Baños, Habana. Cueva de Pio Domingo, Sumidero, Pinar del Río.

**Age.**—Late Pleistocene.

**Other Material.**—Right femur lacking condyles, SEC P-26; ungual phalanges, SEC P-31, P-32, P-35, P-1147, and ACC 1000a; subterminal phalanx, ACC 1000b; distal end of tarsometatarsus, SEC P-40.

**Description.**—Tarsometatarsus (Figure 3) generally similar to that of *Aquila chrysaetos* but notably longer, since even without the distal end it measures 97.7 mm. The estimated total length of the bone is 130 mm, or 34 mm longer than that of *Aquila chrysaetos* and larger than that of any living species of eagle (Table 3). The proximal articular region is similar to that in *Aquila* but with the proximal foramina located only 5 mm from the internal cotyla, whereas in *A. chrysaetos* this distance is nearly twice as great. That which remains of the base of the hypotarsus in the type indicates that this process was probably similar to that of *A. chrysaetos*.

The tarsometatarsi of the fossil species *Buteo typhoicus* and *B. contortus*, from the upper Miocene of Nebraska, and of *B. conterminus*, from the upper Pliocene of Nebraska (Wetmore, 1923), are larger than those of Recent species of *Aquila* and *Haliaeetus*, but are more slender proximally, the articular region in proximal view being of a different shape than in *Aquila* and also differing in the form and position of the middle trochlea. The tarsometatarsus of *B. contortus* measures 113 mm in length, which is 17 mm less than in *Aquila borrasi*. Although the Cuban species has certain similarities to *B. contortus* in the general structure of the tarsometatarsus, it is larger and more robust than that species and appears more like *A. chrysaetos*. According to the published figures, the recently described species *Garganoaetus freudenthali* Ballmann (1973), from the upper Miocene of Italy, is similar morphologically to *B. borrasi*, although its tarsometatarsus is slightly more robust. *Aquila borrasi* was a gigantic form within its genus, the only other known fossil forms of which are *Aquila delphinensis* and *A. pennatoides*, described by Gaillard (1939) from tarsometatarsi from the upper Miocene of France.

The femur of *Aquila borrasi* is larger and more robust than that of any living eagle. Although the one known specimen is incomplete, its maximum length is estimated at about 155 mm, as opposed to 125 mm in *Aquila chrysaetos*, 114 mm in *Haliaeetus leucocephalus*, and 96 mm in *Spizaetus ornatus*. This is even larger than in the two immense living eagles *Harpia harpyja* (131 mm)

**Figure 3.**—Holotype left tarsometatarsus of *Aquila borrasi* (DPUH 1250), Cueva del Túnel. (Anterior view at natural size.)
Table 3.—Measurements (mm) of the tarsometatarsus and claws of *Aquila borrasi* compared with other species of living eagles

<table>
<thead>
<tr>
<th>Character</th>
<th><em>Aquila borrasi</em></th>
<th><em>Pithecophaga jefferyi</em></th>
<th><em>Harpia harpyja</em></th>
<th><em>Aquila chrysaetos</em></th>
<th><em>Haliaeetus leucocephalus</em></th>
<th><em>Spizaetus ornatus</em></th>
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</thead>
<tbody>
<tr>
<td>Tarsometatarsus</td>
<td>Type, DPUH 1250</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length from the upper border of the internal cotyla to the beginning of the scar for the hallux</td>
<td>97.7</td>
<td>80</td>
<td>70</td>
<td>64.5</td>
<td>55</td>
<td>68.2</td>
</tr>
<tr>
<td>Proximal width</td>
<td>22.4</td>
<td>24</td>
<td>36.6</td>
<td>21.5</td>
<td>20.5</td>
<td>18.5</td>
</tr>
<tr>
<td>Least width of shaft</td>
<td>13.2</td>
<td>13.2</td>
<td>19.3</td>
<td>10</td>
<td>10</td>
<td>9.6</td>
</tr>
<tr>
<td>Ungual Phalanx of Digit I</td>
<td>CEC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dorsal arc</td>
<td>52.6</td>
<td>51.2</td>
<td>47</td>
<td>64</td>
<td>38.6</td>
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<tr>
<td>Ventral arc</td>
<td>46.7</td>
<td>47.2</td>
<td>39</td>
<td>51.5</td>
<td>33.5</td>
<td>31.2</td>
</tr>
</tbody>
</table>

* Measurements from specimens in the USNM collections. It can be seen that the tarsometatarsus of *Aquila borrasi*, while decidedly longer, is proportionately much more gracile than in either *Pithecophaga* or *Harpia*—Ed.

and *Pithecophaga jefferyi* (130 mm). Other measurements (in mm) of this specimen are as follows: total length as preserved, 140, proximal width 45.4, vertical diameter of head 16.3, width of neck 17.2, maximum width of pneumatic foramen 8.0, length of pneumatic foramen 14.0, least width of shaft 19.8.

The femur differs from that of allied genera by the lesser projection of the trochanter above the head and by the greater width between the anterior border of the head and the apex of the trochanter (twice that of *Aquila chrysaetos* or *Haliaeetus leucocephalus*). The pneumatic opening is roughly oval in shape but wider distally. It is located at the base of the trochanter as in *Aquila* but differs in being not perfectly oval and in being oriented semi-obliquely towards the external border of the trochanter. The foramina of the upper region of the trochanter are larger and deeper than in the other species compared. The rugose intermuscular line on the anterior face of the bone angles below and near the pneumatic opening along the external border of the shaft almost to its midpoint. In contrast, this line in *A. chrysaetos* originates farther above the upper border of the pneumatic opening and descends straight to the midpoint of the shaft. The head is massive and the neck is thick and oriented slightly upwards.

The ungual phalanges are very well developed (Figure 4), being almost two times larger than those of *A. chrysaetos*. They resemble those of *Harpia harpyja* in having the same degree of curvature. The ungual phalanx of digit IV is larger, while that of digit I is smaller than in *Harpia* (Table 3). The shape of the articular region and the ventral process of the first ungual phalanx of *A. borrasi* more closely resembles that of *Aquila* than *Haliaeetus, Spizaetus*, or *Buteo*. The ungual phalanges of digits II and IV are likewise similar to those of *Aquila* and differ from those of the other genera examined. The ungual phalanx of digit IV measures 33 mm through the ventral arc and 35 mm through the dorsal arc.

Order STRIGIFORMES

Family STRIGIDAE

Genus *Ornimegalonyx* Arredondo

*Ornimegalonyx oteroi* Arredondo, 1958a

**Synonym.**—*Ornimegalonyx arredondoi* Arredondo, 1958.

**Lectotype.**—Left tarsometatarsus SEC P-383E. Lectotype designated by Brodkorb (1961); deposited in the Museum of Comparative Zoology, Harvard University.

**Type-Locality.**—Caverna de Pio Domingo, Si-

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3 Measurements supplied from specimens in the collections of the National Museum of Natural History, Smithsonian Institution.—Ed.
Other Localities.—Sierra de Anafe, Guanajay, Habana; Cueva de Paredones, San Antonio de los Baños, Habana; Cueva del Túnel, La Salud, Habana; Cueva de Isla, Punta del Este, Isla de Pinos; Cueva de Quinto, Boca de Camarioca, Mantanzas; Canteras de los Hornos de Cal, Sancti Spiritus, Las Villas; Sierra de Cubitas, Camagüey.

Age.—Late Pleistocene.

Other Material.—Caverna de Pio Domingo, SEC P-383E, various pieces from the same individual as the lectotype as follows: left tarsometatarsus and fragments of the right, fragments of both tibiotarsi, left femur and fragments of the right, articular portions of the mandible, fragments of the sternum, scapula, carpometacarpus, and ungual and subterminal phalanges. From Paredones, El Túnel, Quinto, and Isla caves there are unnumbered pieces of tarsometatarsi, tibiotarsi, femora, phalanges, synsacra and other bones, all in private collections. From the collection of the author there are three subterminal phalanges from El Túnel and Paredones (OA 671, 676, 800).

Description.—Eight tarsometatarsi of Ornimegalonyx have come to light so far: the lectotype from Pio Domingo; one from Cueva de Quinto; two proximal halves from Paredones; a distal half from Cueva del Túnel and a proximal half from the same locality. Ultimately, two complete specimens from the same individual were found in a cave in the Sierra de Cubitas, Camagüey, along with other elements.

The dimensions of these bones indicate the enormous size that this bird had in relation to all other known Strigiformes, living or fossil (Figures 5–9, Tables 4–6). The tarsometatarsus is almost double the length of that of Bubo bubo, or more than double if one considers the specimen from Cueva de Quinto (GEC unnumbered) (Figure 7a) or that from Paredones. These bones are more than three times the size of the corresponding element of Nyctea scandiaca, more than four times that of Asio otus, and eight and a half times the size of Glaucidium siju.

In spite of its gigantic size, the tarsometatarsus of Ornimegalonyx is proportionately less robust than that of Bubo, Nyctea, or Pulsatrix, the difference being due to the relative lengthening of the shaft in Ornimegalonyx. If the tarsometatarsus of Ornimegalonyx were reduced to the length of that of Asio otus, the two elements would be seen to be very similar in proportion, whereas if one magnified the tarsometatarsi of Bubo, Nyctea, or Pulsatrix to the size of that of Ornimegalonyx, they would appear much stronger, wider, and more robust.

In Ornimegalonyx the internal trochlea is proportionately shorter and wider than in Bubo, Nyctea, or Asio, being more similar to Pulsatrix. The middle trochlea is narrow and placed very close to the outer trochlea. The distal foramen is somewhat lower than in Bubo, Nyctea, or Asio, and the ossified bridge on the anteroproximal region of the bone recalls that of Bubo and differs from Nyctea in that it is stronger and more circular. The internal cotyla is similar to that in Bubo and Asio, but somewhat lower than in Nyctea. The wide, deep groove on the posterior face of the bone is more pronounced than in Bubo, Nyctea, or Asio.

The tibiotarsi of the type individual are fractured into proximal and distal portions and shafts (Figure 5, Table 5). In a complete state they would have measured some 250 mm in length. The tibio-
Table 4.—Measurements (mm) of the tarsometatarsus of *Ornimegalonyx oteroi* compared with other owls

<table>
<thead>
<tr>
<th>Character</th>
<th><em>Ornimegalonyx oteroi</em></th>
<th><em>Bubo bubo</em></th>
<th><em>Nyctea scandiaca</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>SEC P-383.E Lectotype</td>
<td>147*</td>
<td>177</td>
<td>178</td>
</tr>
<tr>
<td>SEC P-59</td>
<td>32</td>
<td>36</td>
<td>35</td>
</tr>
<tr>
<td>GEC (un-numbered)</td>
<td>c. 34</td>
<td>33</td>
<td>32</td>
</tr>
<tr>
<td>MFP (un-numbered)</td>
<td>c. 15</td>
<td>c. 16</td>
<td>15</td>
</tr>
<tr>
<td>Total length</td>
<td>c. 147*</td>
<td>c. 177</td>
<td>c. 178</td>
</tr>
<tr>
<td>Proximal width</td>
<td>c. 32</td>
<td>36</td>
<td>35</td>
</tr>
<tr>
<td>Distal width across trochleae</td>
<td>c. 34</td>
<td>33</td>
<td>32</td>
</tr>
<tr>
<td>Least width of shaft</td>
<td>c. 15</td>
<td>c. 16</td>
<td>15</td>
</tr>
<tr>
<td>Length of middle trochlea from angle with inner trochlea</td>
<td>c. 6</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Least width of middle trochlea</td>
<td>c. 9</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>Length of outer trochlea from angle with middle trochlea</td>
<td>c. 9</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>Width of outer trochlea</td>
<td>c. 6</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>Length of inner trochlea from angle with middle trochlea</td>
<td>c. 6</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>Width of inner trochlea</td>
<td>c. 6</td>
<td>9</td>
<td>9</td>
</tr>
</tbody>
</table>

* Estimated.

tarsus from Cueva de Quinto has a length of 272 mm, which is almost twice that of *Bubo bubo* or *Nyctea scandiaca*. Its proximal width of 40 mm is likewise twice that of those species. The tibiotarsus of the lectotype individual appears to have been straight, while that from Cueva de Quinto is slightly curved and subtly twisted. Compared to the modern genera examined, the tibiotarsus of *Ornimegalonyx* (Figure 6a) has the fibular articulation more pronounced, the rotular crest, and the cnemial crest more elevated, and the fossa proximal to the condyles on the anterior face deeper.

Eight femora of *Ornimegalonyx* have been found so far. The left femur of the type individual is in the MCZ. Of the right, only a part of the proximal end remains in Cuba. The largest femur, represented by the proximal end only (Figure 7d), was found to the east of Sancti Spiritus and is deposited in the Museo Montané of the University of Havana (MMUH 3072). Two incomplete femora from Paredones (formerly SEC P-37 and SEC P-38, but now in the MCZ) are smaller than those of the type and for now are perhaps best referred to as *Ornimegalonyx* sp.

The femur of *Ornimegalonyx* differs principally from that of other owls in being much larger and notably more robust (Table 6). That of the type individual is almost one and a half times larger than the femur of *Bubo bubo*, while the large femur (MMUH 3072) from Sancti Spiritus is almost double the size of that of *Nyctea scandiaca*. It is curious that when reduced, the tarsometatarsi of *Ornimegalonyx* are less robust than in *Bubo* or *Nyctea*, while in the femora the opposite occurs.

Fragments of the sternum, as well as parts of the scapulae, ribs, vertebrae and carpometacarpi, were associated with the type individual in Pio Domingo cave. The most important sternal fragments are an anterior portion with the articulations for the coracoids, and another fragment of the left side containing four costal facets (the fifth having been fractured off). The costal facets vary slightly in size, the largest being 6 X 6 mm. Through these fragments it has been possible to reconstruct the sternum (Figure 8) as being wide, almost flat (both dorsally and ventrally), with a vestigial keel, which indicates that the bird was hardly able to fly. Its estimated length is 120 mm (vs. 47 mm in *Tyto alba*), the estimated width 75 mm (30 mm in *T. alba*); and the height at the keel some 30 mm (25 mm in *T. alba*). The similarity of this last measurement in two species which otherwise differ so greatly in size is a further indication of the great extent of the atrophy of the keel in *Ornimegalonyx*.

The carpometacarpus of *Ornimegalonyx* is small in proportion to the enormous size of the body. Its total length is estimated at about 90 mm whereas in *Bubo bubo*, a smaller volant species, it is 85 mm.
As with the other bones, the phalanges of *Ornimegalonyx* stand apart from those of other owls by their great size. The length of the ungual phalanges of digits II and III, measured through the dorsal arc, range from 37 to 39 mm, while that of digit I measures 40 mm. The proximal height of these phalanges ranges from 15–17 mm.

It was not until the beginning of 1969 that fragments of mandibles were found among the remains of the type individual of *Ornimegalonyx oteroi*. These consist of the two articular portions of both rami. These fragments permit for the first time a very approximate estimate of the size of the mandible and ultimately of the whole skull (Figure 9a). These mandibles are very similar in overall morphology to those of the diminutive genus *Glauccidium*. They differ from *Bubo bubo* in that the internal angular process is greatly lengthened. The posterior angular process, in comparison to that of *Bubo*, is notably more robust, and the portions that remain of the surangular and angular are greatly thickened at the point of the fracture. Judging by comparison with recent owls, the length of the entire mandible of *Ornimegalonyx oteroi* would have been some 115 mm and the distance between the external borders of the articulations was approximately 100 mm, or about two times larger than in *Bubo bubo*.

A portion of a cranium of *Ornimegalonyx* was...
Table 5.—Measurements (mm) of the tibiotarsus of *Ornimegalonyx oteroi* compared with other owls

<table>
<thead>
<tr>
<th>Character</th>
<th><em>Ornimegalonyx oteroi</em></th>
<th><em>Ornimegalonyx oteroi</em></th>
<th><em>Bubo bubo</em></th>
<th><em>Nyctea scandiaca</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SEC P-383.E</td>
<td>SEC P-28</td>
<td>MFP (un-numbered)</td>
<td>GEC (un-numbered)</td>
</tr>
<tr>
<td>Total length</td>
<td>250*</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Proximal width</td>
<td>c. 36</td>
<td>39</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Least width of shaft</td>
<td>c. 13</td>
<td>14</td>
<td>15*</td>
<td>13*</td>
</tr>
<tr>
<td>Distal width across trochlea</td>
<td>30</td>
<td>-</td>
<td>31</td>
<td>c. 30</td>
</tr>
</tbody>
</table>

* Estimated.

Table 6.—Measurements (mm) of the femur of *Ornimegalonyx oteroi* compared with other owls

<table>
<thead>
<tr>
<th>Character</th>
<th><em>Ornimegalonyx oteroi</em></th>
<th><em>Ornimegalonyx oteroi</em></th>
<th><em>Bubo bubo</em></th>
<th><em>Nyctea scandiaca</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SEC P-383.E</td>
<td>MFP (un-numbered)</td>
<td>MMUH 3072</td>
<td>108</td>
</tr>
<tr>
<td>Maximum length</td>
<td>154</td>
<td>160</td>
<td>168*</td>
<td></td>
</tr>
<tr>
<td>Proximal width between antero-external border of head and postero-external border of trochanter</td>
<td>42</td>
<td>45*</td>
<td>46</td>
<td>23.7</td>
</tr>
<tr>
<td>Antero-posterior diameter of head</td>
<td>c. 11</td>
<td>12*</td>
<td>c. 12</td>
<td>7.5</td>
</tr>
<tr>
<td>Vertical diameter of head</td>
<td>15</td>
<td>-</td>
<td>c. 15</td>
<td>9</td>
</tr>
<tr>
<td>Least width of shaft</td>
<td>16</td>
<td>16*</td>
<td>20*</td>
<td>9.6</td>
</tr>
<tr>
<td>Distal width through condyles</td>
<td>35*</td>
<td>38*</td>
<td>-</td>
<td>22.5</td>
</tr>
</tbody>
</table>

* Estimated.

Figure 8.—Ventral and lateral views of the sternum of *Ornimegalonyx oteroi*, as reconstructed from two fragments from Pío Domingo and Paredones caves. (Natural size.)
found in Cueva de Paredones in 1959 (Figure 9b,c). For the time being it is best referred to only as *Ornimegalonyx* sp., for it is apparent that it does not correspond to the species *O. oteroi*. Rather, it appears to belong to a smaller form, as also evidenced by the femora found in the same locality. The specimen consists of the posterior portion of a cranium from the postorbital arc to the occiput and including the basisphenoid, the foramen magnum, and the occipital condyle. Viewed from the front, the great thickness of the walls of the cranium are seen in the region of the break. Compared to *Bubo bubo* it is larger, and in ventral view it has the postorbital process better developed. The opisthotic process is rather prominent and its extremity bends notably, hanging in the form of an ear. The basipterygoid processes are well developed. The foramen magnum is somewhat higher than wide, as opposed to *Glaucidium* and *Tyto* in which it is wider than high. As in *Nyctea*, the occipital condyle is very well developed in relation to the foramen magnum, whereas it is proportionately much smaller in *Glaucidium* and *Tyto*.

From the actual and estimated measurements of the various bones of *Ornimegalonyx* it can be established that this great owl stood some 1100 mm high in life. Although the general aspect of its skeleton is similar to that of living owls, it is distinguished from them by the long and robust hindlimbs, provided with long, heavy toes armed with the most powerful claws possessed by any strigiform bird. Although the sternum is larger than in any living owl, it is actually small in proportion to the rest of the bones of the skeleton. Its semiflat structure and reduced keel show that *Ornimegalonyx* was little or not at all capable of flight. In accordance with this, the bones of the wing are poorly developed, particularly the carpometacarpus.

Some of the differences in morphology and size that are observed between individuals of *Ornimegalonyx* are probably attributable to sexual dimorphism, since in other owls the females are larger than the males. Nevertheless, other bones that are either larger or smaller than those of the type individual and have distinct differences from it, probably indicate additional species—the cranium and femora from Paredones and the femur from Sancti Spiritus being examples.

*Ornimegalonyx* had to have been the scourge and terror of most of the larger mammals of the Pleistocene of Cuba and the claws and mandibles of this bird would have constituted a terrible combination of superior destructive power.

Family **TYTONIDAE**

Genus *Tyto* Billberg

*Tyto noeli* Arredondo, 1972a

**Holotype.**—Right tarsometatarsus, DPUH 1251.

**Type-Locality.**—Cueva del Túnel, La Salud, Habana, Cuba.

**Other Localities.**—Cueva de Paredones, San Antonio de los Baños, Habana; Cueva del Indio, Reparto El Globo, Calabazar, Habana; quarries near Sancti Spiritus, Las Villas.

**Age.**—Late Pleistocene.

**Other Material.**—Cueva del Túnel: OA 818, right femur; OA 812, distal portion of left tibiotarsus; OA 804, distal portion of right humerus; OA 806, proximal fragment of right humerus; OA 822, shaft of right tibiotarsus; OA 815, distal portion of right ulna. Cueva de Paredones: OA 828, proximal portion of right tarsometatarsus; OA 827, proximal portion of right tibiotarsus; OA 889, right coracoid. Cueva del Indio: OA 1027, right femur.

**Description.**—Similar to the living species *Tyto alba* in its general skeletal configuration, but much larger (Figures 10, 11, Tables 7 and 8), equaling in size the extinct species *Tyto ostologa* of Haiti and *T. pollens* of the Bahamas. The tarsometatarsus was between 90 and 100 mm long and was similar to that of *T. pollens*, but more slender, even in specimens that are longer than in *T. pollens*. The tibiotarsus is likewise similar to that of *T. pollens* but is less robust. This slenderness of the hindlimb is the most notable difference between the two species.

In the femur, humerus, ulna, coracoid, and claws, the only pronounced difference from *Tyto*...
alba is in size. Likewise, a fragment of the anterior portion of a sternum of T. noeli from quarries near Sancti Spiritus has the same conformation as that of T. alba but is larger. One might expect to find greater distinctions in the skull, but so far only fragments of the skull of T. noeli have been found.

Brodkorb (1959:357) suggested that T. pollens may possibly be differentiated from T. ostologa only at the subspecific level. The same could be suggested for T. noeli. From the upper Miocene of Italy, a new species of giant barn owl, Tyto robusta Ballmann (1973), has been described that is near the size of T. noeli. The Cuban species is somewhat larger and heavier, however.
Table 7.—Measurements of limb bones of *Tyto noeli* compared with *Tyto alba*

<table>
<thead>
<tr>
<th>Character</th>
<th><em>Tyto noeli</em></th>
<th><em>Tyto alba furcata</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>OA 818</td>
<td>OA 834</td>
</tr>
<tr>
<td>Total length</td>
<td>74.3</td>
<td>73*</td>
</tr>
<tr>
<td>Proximal width</td>
<td>14.4</td>
<td>13.1</td>
</tr>
<tr>
<td>Antero-posterior diameter of head</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Vertical diameter of head</td>
<td>5.8</td>
<td>5</td>
</tr>
<tr>
<td>Breadth through trochanter</td>
<td>9.2</td>
<td>8.2</td>
</tr>
<tr>
<td>Least width of shaft</td>
<td>6.4</td>
<td>6</td>
</tr>
<tr>
<td>Distal width</td>
<td>14.8</td>
<td>14</td>
</tr>
</tbody>
</table>

| Tibiotarsus                      | OA 827       | OA 831              |
| Total length                     | 147*         | –                   | 108                 |
| Proximal width                   | 16           | –                   | 10.9                |
| Least width of shaft             | 6.4          | –                   | 5.5                 |
| Distal width                     | –            | 15                  | 11.3                |

| Humerus                          | OA 804       | OA 826              |
| Total length                     | 137*         | –                   | 100.1               |
| Proximal width                   | –            | 23                  | 16.9                |
| Width of shaft                   | 8.5          | –                   | 6.2                 |
| Distal width                     | 20.5         | –                   | 15.7                |

* Estimated.

Table 8.—Measurements (mm) of the tarsometatarsi of the three Cuban species of *Tyto*

<table>
<thead>
<tr>
<th>Character</th>
<th><em>Tyto riveroi</em></th>
<th><em>Tyto noeli</em></th>
<th><em>Tyto alba furcata</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DPUH 1252 Type</td>
<td>DPUH 1251 OA 828 Type</td>
<td></td>
</tr>
<tr>
<td>Maximum length</td>
<td>125*</td>
<td>91.7</td>
<td>100*</td>
</tr>
<tr>
<td>Proximal width</td>
<td>22*</td>
<td>14.6</td>
<td>16</td>
</tr>
<tr>
<td>Distal width</td>
<td>22</td>
<td>17.4</td>
<td>–</td>
</tr>
<tr>
<td>Least width of shaft</td>
<td>9</td>
<td>6.7</td>
<td>7.3</td>
</tr>
<tr>
<td>Length of middle trochlea from</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>angle of internal trochlea</td>
<td>6.8</td>
<td>5</td>
<td>–</td>
</tr>
<tr>
<td>Width of middle trochlea</td>
<td>8.5</td>
<td>6.4</td>
<td>–</td>
</tr>
<tr>
<td>Length of outer trochlea from</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>angle of middle trochlea</td>
<td>11.5</td>
<td>c.4</td>
<td>–</td>
</tr>
<tr>
<td>Width of outer trochlea</td>
<td>12.3</td>
<td>4.8</td>
<td>–</td>
</tr>
<tr>
<td>Length of inner trochlea from</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>angle of middle trochlea</td>
<td>6.5</td>
<td>4.4</td>
<td>–</td>
</tr>
<tr>
<td>Width of inner trochlea</td>
<td>9.8</td>
<td>7.5</td>
<td>–</td>
</tr>
</tbody>
</table>

* Estimated.

*Tyto riveroi* Arredondo, 1972b

**Holotype.**—Distal portion of a left tarsometatarsus, DPUH 1252.

**Type-Locality.**—Cueva de Bellamar, Mantanzas, Cuba.

**Age.**—Late Pleistocene.

**Description.**—Tarsometatarsus with the general aspect of that of *Tyto alba* and still more similar to that of *T. noeli*. Except for size, significant morphological differences from the above species are almost absent; however, the measurements of the type of *T. riveroi* notably exceed the limits of either (Figure 11, Table 8). The estimated total length of this bone would be approximately 125
mm. The following slight morphological distinctions are also noted: greater separation of the internal and external trochlea from the middle trochlea, the intertrocchlear spaces being narrower in *T* alba and *T* noeli; shaft proportionately wider and thicker. Compared with *T.* pollens, the same slight differences are apparent.

*Tyto gigantea*, recently described from the upper Miocene of Gargano, Italy (Ballmann, 1973), was an enormous barn owl, equal in size to *T.* riveroi. According to the published figures, the Italian species has the distal foramen somewhat more elevated and the middle trochlea lower and more elongate than in the Antillean species.

![Figure 11](image)

**Figure 11.**—Comparison of the tarsometatarsi of the three Cuban species of *Tyto*: *a*, Recent *Tyto alba furcata*; *b*, *Tyto noeli*, holotype (DPUH 1251), Cueva del Túnel; *c*-*e*, *Tyto riveroi*, holotype (DPUH 1252), Cueva de Bellamar, anterior, posterior, and lateral views. (Natural size.)

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Varona, Luis S.

Wetmore, A.

Wołoszyn, B. W., and N. A. Mayo
The Species of Moas (Aves: Dinornithidae)

Joel Cracraft

ABSTRACT

In order to provide a basis for future studies on the functional morphology and evolutionary history of the New Zealand moas, the species-level systematics of the family is reviewed. Based on a study of museum collections and an analysis of intraspecific variability, only 13 species are considered to be valid: Anomalopteryx didiformis, A. oweni, Megalapteryx didinus, M. benhami, Pachyornis elephanotus, P. mappini, Euryapteryx curts, E. geranoides, Emeus crassus, Dinornis struthoides, D. torosus, D. novaezelandiae, and D. giganteus. The taxa accepted as valid in this paper probably fit the biological species concept more closely than do those of previous classifications, in which as many as 29 species have been recognized. The "species-pairs," Pachyornis mappini–P septentrionalis, Euryapteryx curts–E. exilis, E. geranoides–E. gravis, and Emeus crassus–E. huttonii probably represent examples of sexual size dimorphism.

Introduction

Birds provide a number of classic examples of insular adaptive radiations, the best known being the Hawaiian honeycreepers (Drepanididae) and the Galapagos finches (Geospizinae). A unique radiation of this sort is also found in the New Zealand moas (Dinornithidae)—unique because a spectacular radiation took place not only in feeding mechanisms, but also in body size and proportions, in contrast to the well-known passerine radiations. This occurred within a group that was completely flightless, and indeed it is probable that being flightless enhanced selective pressures toward divergence in cranial morphology and body size.

Moas are primitive ratite birds whose closest living relatives are probably the kiwis (Cracraft, 1974). I have postulated elsewhere that the common ancestor of the kiwis and moas probably had a distribution in the Cretaceous that included western Antarctica and parts of South America and that moas and kiwis were isolated on New Zealand following northward continental drift of that landmass beginning in the Late Cretaceous (Cracraft, 1973a; 1974). If this is so, then the radiation of moas probably began in the Tertiary, but it is my opinion that the taxonomic and morphological diversity observed in the Holocene assemblages of moas is the result of a relatively recent, probably Pleistocene, episode of speciation (Cracraft, in prep.). Thus, species formation within moas probably involved repeated cycles of isolation and sympathy among populations of the two major islands and among populations that very likely were isolated in forest refugia at glacial maxima. It is within such a model of their evolution that I have considered the species-level systematics of moas.

Despite the fact that hundreds of papers have been written about moas (see summaries in Lambrecht, 1933; Oliver, 1949), there is comparatively little known about their functional morphology or evolutionary history. Upon initiating such studies, it quickly became clear that the complexities and confusion of presently accepted species-level systematics would hinder any advances in morphological or evolutionary investigations. It thus became necessary to review the status of the various species of moas, and this paper presents the results of that study.

The species-level systematics of moas has suffered from a century of typological thinking and a lack of application of modern concepts of population biology. No less than 60 specific names have been applied to a group that almost certainly con-
tains fewer than 20 biological species (see Brodkorb, 1963, for citations to the taxonomic literature). Each variant, each newly discovered bone in some cases, was frequently given a new specific name. Unfortunately, the taxonomic philosophy of certain influential recent workers, rather than clarifying the systematics of these birds, merely confounded matters further. Oliver (1949:132-134), for example, believed that “it is the work of the systematist . to define the units that make up [a series of continuous sizes between extreme forms],” and that “in dealing with fossil species we should not hesitate to give specific names to forms that differ only slightly from one another . . . .” (1949:164). It was this philosophical approach that enabled Oliver as late as 1949 to describe one new genus and six new species of moas, all of which were based on a small number of isolated bones. As will be seen, none of these taxa appears to be valid.

It is my purpose here to propose species-limits within moas which it is hoped will reflect the biological structure of those species. I have attempted to obtain some estimate of intraspecific variability and to apply this to recognizing species-limits. Species based on isolated bones or on inadequate material are critically evaluated for validity; likewise, geographic representatives on separate islands, previously given separate species names, are here considered conspecific unless there is good evidence to the contrary. Species-limits of moas undoubtedly will remain a matter of personal opinion for some time to come, and I make no pretense at having arrived at a definitive picture. There is a need for additional studies, and the systematics of these birds would benefit especially from a comprehensive numerical-multivariate approach. I do believe, however, that the species-limits proposed in this paper considerably clarify moa taxonomy and facilitate more interesting studies of their evolution.

Materials and Methods.—I have studied material of moas in the following museums (abbreviations used in the text follow in parentheses): British Museum (Natural History) (BM); American Museum of Natural History; Field Museum of Natural History; Canterbury Museum, Christchurch; Otago Museum, Dunedin; and National Museum of New Zealand (formerly Dominion Museum) (DM), Wellington. During this study the only major collection I was unable to examine was that at the Auckland Museum (AM), Auckland. Fortunately, Archey’s (1941) valuable monograph on the moas is based almost entirely on the Auckland collection and I was, therefore, able to incorporate much information on that material into this study.

Of the species recognized by Oliver (1949) I have examined material of all except Pachyornis murihiku, Anomalopteryx antiquus, Megalapteryx hectori, and M. benhami. All of these were described from isolated bones and it is probable that none represents a valid species, with the possible exception of M. benhami. Thus, I was able to study the majority of taxa in need of critical evaluation.

In addition to using standard univariate statistical procedures, I have employed several multivariate morphometric techniques in order to characterize patterns of intra- and interspecific variability in more detail. The theory and methodology of multivariate approaches and their application to biological problems are discussed by Blackith and Reyment (1971) and Oxnard (1973). Basically, these techniques describe patterns of variation or degrees of similarity (or difference) for many variables taken simultaneously over many taxa. I have used two techniques: (1) principal components analysis (BMD01M; Dixon, 1970) in order to examine the structure of variation within groups, primarily to investigate problems of sexual dimorphism in size and shape within a species; (2) discriminant function-canonical analysis (BMD07M; Dixon, 1970), in order to examine the patterns of variation among groups that are defined prior to the analysis. I employed this approach to examine the nature of the separations among the presumed species of a genus to evaluate species distinctness and the presence of sexual dimorphism. Part of the output of BMD07M is a posterior probability classification which allows one to discover whether individuals assigned to one group prior to the analysis are in fact closer to the means of another group.

In all examples employing multivariate techniques, I undertook the analysis of each hindlimb element based either on my own data or that in Archey (1941) and Oliver (1949) using the following four variables: bone length, breadth of proximal end, breadth of shaft at midpoint, and breadth of distal end.
Acknowledgments.—It is with pleasure that I dedicate this paper to Dr. Alexander Wetmore in honor of his ninetieth birthday, his many contributions to avian paleontology and ornithology, and especially for the affection he has shown us all.

I want to thank the following institutions and individuals for making their collections available to me and for making my visits enjoyable and productive: American Museum of Natural History (Wesley Lanyon, Malcolm C. McKenna); British Museum (Natural History) (Alan J. Charig, Cyril A. Walker); Canterbury Museum (Roger S. Duff, Ronald J. Scarlett, Michael Trotter); Field Museum of Natural History (John Bolt, Melvin A. Traylor); National Museum of New Zealand (John C. Yaldwyn, R. K. Dell); and Otago Museum (Roy Forster, J. Darby). I am grateful to Hildegard Howard and Pat V. Rich, John C. Yaldwyn, and Sir Robert Falla for providing helpful comments on an early draft of this paper. My trip to New Zealand and subsequent data analysis were made possible by financial support from the Australian Academy of Science (through H. J. Frith), the Chapman Fund of the American Museum of Natural History, and the National Science Foundation (grant GB-41089).

Intraspecific Variability

The relative variability of one species of kiwi and four species of moas, all of which are believed to represent “good” biological species, are examined here. By gaining some understanding of the degree of variability within species recognized to be valid by nearly all previous workers, a basis of comparison can thereby be provided for assessing species-limits among the more controversial taxa considered in the following section. For reasons to be discussed below, these assessments of variability within “good” species cannot be used as absolute limits or criteria of species variability, but they can serve as guidelines.

Causes of Variability in Moas

It has been known for 100 years or more that moas are highly variable. Moreover, some workers have seemed to appreciate the fact—although they seldom stated so explicitly—that this variation is complex in nature and cannot be attributed to any single factor. The main obstacle to understanding this variation is that of dealing with fossil (perhaps more correctly, subfossil) populations and their well-known problems of sampling in space and time. Some workers have sought to solve these problems either by ignoring them or by naming new species for each variant and thereby eliminating the necessity of delimiting or explaining intraspecific variability (note comments by Oliver cited above).

At this time it is not possible to make quantitative estimates of the separate factors contributing to patterns of variability within species of moas. Sample sizes for some species over their entire range, or for local populations of most species, are usually too small; stratigraphic control is lacking for all but a few of the moa specimens collected so far. Consequently, one is forced to estimate intuitively the relative importance of temporal, geographic, and individual components of variation. Within most fossil populations, particularly of moas, the following five factors seem important.

Intrapopulational Variation.—This is the variation observed in individuals of the same local interbreeding populations. We might expect the amount of such variation to be relatively low compared to those samples in which the influences of geographic or temporal variation are also present. Most of the samples discussed in this section do not represent single local populations; such samples probably do exist for certain species, but restrictions of time while in New Zealand did not permit me to study this aspect of variation in detail. Certain samples of Apteryx australis and Euryapteryx curtus (including E. exilis) might provide examples of this type of variation, and these are discussed below.

Sexual Dimorphism.—Since sexual size dimorphism is common in other ratites (with either males or females being larger), one would expect moas to show sexual size dimorphism also. Unfortunately, sample sizes are usually inadequate to lend support to this, although evidence is presented below of four probable examples of sexual size dimorphism in moas. The question of how much size difference to expect between sexes must be considered when dealing with fossil populations: if there were too great a difference between two particular samples, then we might be inclined to recognize two species rather than two sexes. But
what is “too great” a difference? Almost no quantitative information has been published about dimorphism in other ratites. In one study of the Emu (Dromiceius novaehollandiae), data gathered by Long (1965) show that coefficients of variation (CV) of combined male-female samples range from 3.5 for bill length to 19.14 for body weight. Coefficients of variation for lengths of a few skeletal elements range from 4.77 for the tarsometatarsus to 5.28 for the tibiotarsus. These CVs for the skeletal elements are generally less than those calculated for the expanded species of moas discussed in the next section.

**Geographic Variation.**—The major factor in this type of variation in moas is probably inter-island differentiation. It has previously been noted that the bones of North Island forms tend to be shorter and less stout than those of comparable taxa from the South Island (Archey, 1941:62, 71; Scarlett, 1972:20; Oliver, 1949:164). In addition, some intra-island differentiation may have occurred, but samples are too limited to confirm this. I believe geographic differentiation contributes greatly to the large variability in the samples of certain species discussed below.

**Temporal Variation.**—It is difficult to assess the importance of temporal variation in affecting variability within moas. The chronology of natural moa deposits is not well known. One can be reasonably certain that these assemblages are no older than 7000–8000 years, and most are undoubtedly much younger, apparently less than 4000 years old (Fleming, 1962). Hence, it may be that temporal variation contributes relatively little to the variability of the available samples of moas.

**Variation and Recency of Sympatry.**—If speciation in moas has resulted from isolation in forest refugia during glacial maxima, then it is reasonable to assume that variability would have increased as a result of this isolation, thus providing an example of intra-island geographic variation. Many samples of moas probably are composed of different populations that had come in contact following the last glaciation. Thus, we may be sampling birds that had recently diverged morphologically, and some of the variation observed may be the result of recent character displacement in size following this contact.

### Analysis of Species

Basic statistical data for the femora, tibiotarsi, and tarsometatarsi of *Apteryx australis* and four species of moas are given in Table 1. Of particular importance for the discussions that follow are the coefficients of variation (CV), which are measures of relative variability independent of size.

*Apteryx australis*: This sample (housed in the National Museum of New Zealand) of 24–32 individuals referable to the modern Brown Kiwi, comes from the Castle Rocks cave deposit on the South Island. It is evident from Table 1 that *A. australis*, with CVs ranging from 4.68 to 7.6, exhibits less variability than any of the moas. I attribute this to the relative lack of geographic and temporal influences since the sample comes from a single locality and was probably deposited over a relatively short span of time. Of the species studied, this sample of *A. australis* possibly comes closest to representing only intrapopulational variation. The degree of variability in this sample is similar to that shown by fossil populations of some species of gruiforms (Cracraft, 1973b). In *A. australis*, measurements of length are less variable than those of breadth.

*Megalapteryx didinus*: This sample is taken from many localities in the South Island (data from Archey, 1941; Oliver, 1949). In a later section I synonymize *M. hectori* with *M. didinus*, but specimens assigned to the former are not included in this sample. This species may be one of the more variable of moas in that the lowest CV is 5.79 while the highest is 12.04. Most of this variability is probably attributable to geographic variation. As with *A. australis*, measurements of length are less variable than those of breadth.

*Anomalopteryx didiformis*: The analysis of this species is based on a large sample collected from sites on both islands. Included in this series by Archey (1941, table A) are a few specimens that might be assigned to Oliver’s (1949) species *A. parvus*. Few workers accept *A. parvus* as a distinct species (see below) and my analysis is based on all the specimens listed by Archey. This species also shows a fairly high degree of variability, with CVs ranging from 6.79 to 9.3. Again, lengths are less variable than other measurements. The variability of this sample is probably influenced substantially by geographic and intrapopulational variation.
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Dinornis torosus: The data for this species are based on a small sample collected from many sites in the South Island (Archey, 1941; Oliver, 1949). Variability is fairly high, with CVs ranging from 5.73 to 9.99. This is the only species in which lengths tend to be more variable than breadths. Variability in this sample is probably affected by geographic and temporal components.

Pachyornis elephantopus: This sample is from various sites on the South Island (Archey, 1941; Oliver, 1949; and measurements by the author). Coefficients of variation are comparable to those of A. didiformis and D. torosus, ranging from 5.03 to 9.68. Once again, length measurements are less variable than those of breadth. Intra-island components probably contribute substantially to variability in this species.

Summary

Variability within individual species of moas appears somewhat higher than is found in most other birds so far studied. For example, in six fossil species of gruiforms, CVs for hindlimb elements averaged 6–7 (Cracraft, 1973b:97–107). Simpson (1946) presented data for the King Penguin (Aptenodytes patagonicus) and Goodge (1951) analyzed variability in the Common Murre (Uria aalge); both workers found low CVs, averaging 2–4, in these extant species.

Data for A. didiformis, D. torosus, and P. elephantopus would seem to suggest that in moas CVs generally fall in the range of 6–10, but if the sample of M. didinus were used as a standard, then CVs as high as 10–12 might be expected within a single species. Indeed, estimates based on the above samples may be conservative in that they possibly include only individuals of a single sex, specimens of the other sex previously having been considered as forming a distinct species.

During growth, the length of long bones apparently reaches its maximum value before maximum body weight is obtained (Cock, 1963). Bone breadths of adults, on the other hand, probably reflect final body weight since they have a mechanical relationship to the amount of weight that can be supported. It can thus be expected that breadth measurements will tend to be more variable than those for length, since the former depend upon variation in body weight at the termination of growth. Estimates of variation in length might be preferred over those for breadth as a more precise, and more conservative, measure of intraspecific variability. For this reason the comparisons in the following section will be based on measurements of length.

Systematics

In this section I have attempted to formulate species limits based on information from univariate and multivariate analyses of variation, personal examination and comparison of many complete and partial skeletons, and an evaluation of previously published opinions on moa systematics. Detailed discussions of morphology (particularly at the generic level and higher), natural history, and taxonomic synonymies can be found in Archey (1941), Oliver (1949), Brodkorb (1963), or papers cited therein, and are not included here unless they bear directly on the subject of species limits.

Family DINORNITHIDAE

Subfamily ANOMALOPTERYGINAE

Anomalopteryx Reichenbach, 1852

Summary.—Two species of Anomalopteryx are tentatively admitted here—a larger form, A. didiformis, found on both North and South islands, and a smaller, less common form, A. oweni, found only on the North Island.

Anomalopteryx didiformis (Owen, 1844)

Synonyms.—Anomalopteryx parvus (Owen, 1883), Anomalopteryx antiquus Hutton, 1892.

Of the two species of the genus recognized here, this was the more common and occurred on both the North and South islands. Archey’s (1941:14–29) discussion, although not employing statistical methods, amply demonstrated the great variation present within this species.

Anomalopteryx parvus, based on fairly extensive material from both islands, is included in this species, following Archey (1941), Brodkorb (1963), and Scarlett (1972). This is contrary to Oliver (1949:138), who maintained A. parvus as distinct. Oliver (1949:144–145, figs. 115–116) pictured bones.
of the two species for comparative purposes, and indeed these appear quite different in size. Oliver's method of analysis and argumentation, however, was to compare the extremes in size or shape, thus magnifying the differences. There is a continuity in size and morphology in specimens assigned to these two species (Archey, 1941:18); thus, unlike examples to be described below in other genera, it is not possible to recognize two size groups within _A. didiformis_ that could represent sexual size dimorphism.

According to Scarlett (1972:22), _A. antiquus_ is now considered to come from lower Pleistocene deposits rather than being Miocene or Pliocene in age as previously thought. Scarlett further indicated that _A. antiquus_ is "doubtfully distinct from _didiformis_" and that there are no significant morphological differences between them. Likewise, Archey (1941:29) noted close similarities between the two species. I did not locate the type during my stay at the Canterbury Museum. At present I believe that compelling evidence is lacking to maintain _A. antiquus_ as a separate species.

**Anomalopteryx oweni (Haast, 1885)**

There has been some controversy about the generic assignment of this species. As did all other early workers, Haast (1885, 1886) placed this species in _Dinornis_. Shortly thereafter, Lydekker (1891:280) put _oweni_ in _Anomalopteryx_ on the basis of its skull morphology. Archey (1941:44) transferred the species to _Pachyornis_ without comment; Brodkorb (1963:211) followed Archey. Oliver (1949:134-135) returned _oweni_ to _Anomalopteryx_, claiming that the type cranium illustrated by Haast (1886) shows the diagnostic features of _Anomalopteryx_, as does the associated premaxilla, and that most of the skeletal material assigned to _oweni_ by Archey belongs to _Pachyornis septentrionalis_ (= _P. mappini_ of this paper).

Although I was unable to examine the type-material of _A. oweni_ in the Auckland Museum, several comments on the species can still be made. The cranium and premaxilla illustrated by Haast do appear to be more similar to _Anomalopteryx_ than to _Pachyornis_, although the rounded anterior border of the temporal fossa in dorsal view and the markedly sloping nasal region in lateral view do not resemble species of either genus. Some of the hindlimb elements listed by Archey as belonging to _oweni_ are within the size range of _Pachyornis mappini_ (including _P. septentrionalis_), whereas others appear to be too small to be referred to that species. Thus, until the systematics of _oweni_ can be clarified by restudy of the type and comparison with other material, I tentatively include it as a valid species in the genus _Anomalopteryx_.

**Megalapteryx Haast, 1886**

**Summary.**—Two species of _Megalapteryx_ are accepted here—a small one, _M. didinus_, and a larger one, _M. benhami_. Both are known from the South Island, the alleged presence of _M. didinus_ in the North Island being doubtful.

**Megalapteryx didinus (Owen, 1883)**

**Synonym.**—_Megalapteryx hectori_ Haast, 1886. _Megalapteryx didinus_ is known from a moderate number of bones from the South Island, few of which were found in association. The presence of this species on the North Island is suspect (Oliver, 1949:151-152).

Archey (1941) synonymized _M. hectori_ with _M. didinus_ without comment, but Oliver (1949:149) maintained the species. Brodkorb (1963) and Scarlett (1972) followed Archey. I also believe there is insufficient evidence to justify recognition of _M. hectori_. Only a few limb bones are presumably applicable to this species (I did not examine the type, which is in the Nelson Museum), and their size, although somewhat smaller than typical bones of _M. didinus_, probably falls within the limits of variability for that species. For example, CVs for the lengths of the hindlimb elements for the combined sample of _M. didinus_ and _M. hectori_ are comparable to CVs of the other moas listed in Table 1: e.g., femur, 8.45; tibiotarsus, 6.76; and tarsometatarsus, 7.40. I therefore follow the authors cited above in synonymizing _hectori_ with _didinus_.

**Megalapteryx benhami Archey, 1941**

This species was described on the basis of a femur and a tibiotarsus, not positively associated, from the Mt. Arthur region, South Island. Oliver (1949) also lists a femur from Wairanga, South Island.
The bones were placed in *Megalapteryx* on the basis of several morphological characters associated with the rotular groove and muscle scars (Archev, 1941:35), and it would be important for future workers to confirm the validity of these characters in distinguishing genera. As many authors have noted, there is great variability in the form of the femur and it is frequently difficult to identify this element to genus unless found in association with more diagnostic bones. As it now stands, the species *M. benhami* can be tentatively accepted, as its larger size (mean femur length of 296.5 mm; length of tibiotarsus, 454 mm) is very probably outside the range of variation for *M. didinus*. Future workers need to restudy the material of this species in order to verify its validity and generic assignment.

**Pachyornis Lydekker, 1891**

**Summary.**—Two species of *Pachyornis* are recognized here. The larger, *P. elephantopus*, is known only from the South Island. A smaller species, *P. mappini*, showing what appears to be pronounced sexual dimorphism in size, was restricted to the North Island.

**Pachyornis elephantopus** (Owen, 1856)

**Synonyms.**—*Pachyornis murihiku* Oliver, 1949; *Pachyornis australis* Oliver, 1949.

*Pachyornis elephantopus* was restricted to the South Island and was the larger of the two species of the genus recognized here. It was also one of the more common species of moas and is represented by a number of complete skeletons from the Pyramid Valley Swamp.

Oliver (1949:67) described *P. murihiku* for a single skeleton said to be from a “full-grown but not quite mature” individual from Southland, South Island. The type was supposedly in the Southland Museum, Invercargill, but according to Scarlett (1972:21) it cannot now be found. Both Brodkorb (1963) and Scarlett (1972) accepted *P. australis* as a distinct species, although Scarlett indicated that it may be a variant of *P. elephantopus*. I have compared the type (DM 26) with a large series of skulls of *P. elephantopus* and can find no significant differences that can be regarded as being of specific value. Even though the skull of *P. australis* is at the lower end of the size range for *P. elephantopus*, several skulls assigned to the latter species by Oliver himself (1949:84) are of comparable size (e.g., DM 95, DM 198, DM 333). Cranial of *P. elephantopus* exhibit considerable variability in shape and in the development of processes and muscle scars. For example, within a series of skulls of *P. elephantopus* in the National Museum of New Zealand it was possible to find the following characters attributed to “*P. australis*” by Oliver (1949:70–72): (1) skull wider in proportion to length, (2) evenly rounded cranium, (3) wide space between lambdoidal and temporal ridges, and (4) narrow temporal fossa. Furthermore, there are similarities in rostral shape and considerable variation in the development of the transverse process of the basisphenoid rostrum. Therefore, it does not seem prudent to accept *P. australis* as a valid species.
**Pachyornis mappini** Archey, 1941

SYNONYM.—*Pachyornis septentrionalis* Oliver, 1949.

Archey (1941:41) proposed this species for a small North Island form of *Pachyornis*. The type (AM 124) is an almost complete skeleton. Most of the remaining material assigned to *P. mappini* consists of isolated elements and many of those in the National Museum of New Zealand have been acquired recently and are as yet undescribed.

Oliver (1949:61) described a new species, *P. septentrionalis*, for a partial skeleton (DM 129), also from the North Island. He included in this species those bones from the lower end of the series that Archey (1941) placed in *P. mappini*. Oliver (1949:61) stated that bones of *P. septentrionalis*, in addition to being smaller, are also more slender than those of *P. mappini*. Brodkorb (1963) accepted both species, while Scarlett (1972) suggested that the two might be conspecific.

An analysis of the skeletal measurements of these nominal species (Archey, 1941:139; Oliver, 1949: 86; measurements by the author) indicates that two separable populations do appear to exist, the major difference between them being in size (Table 2). Using principal components analysis of each element of the hindlimb (Figure 1), a moderately well-defined separation between the two forms can be demonstrated along the first principal component, which in this case is a size axis. The second component is a shape axis, mainly defining relative robustness, and it is evident that the two populations do not differ significantly in this respect. Thus, Oliver’s (1949:61) claim that bones of *P. septentrionalis* are more slender than those of *P.

![Figure 1.—Principal components analyses of logarithmically transformed measurements of the femur (a), tibiotarsus (b), and tarsometatarsus (c) of *Pachyornis septentrionalis* (dark circles) and *P. mappini* (open squares). The first principal component of each analysis is graphed along the abscissa and is a size axis; the second principal component is graphed along the ordinate and is a shape axis, primarily indicating relative robustness. (Note that the two taxa are separated by size but not by shape. See text for details.)](image)

<table>
<thead>
<tr>
<th>Table 2.—Statistics for <em>Pachyornis mappini</em> and <em>P. septentrionalis</em> (measurements in mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Character</td>
</tr>
<tr>
<td>Length of femur</td>
</tr>
<tr>
<td>n</td>
</tr>
<tr>
<td>$\bar{X}$</td>
</tr>
<tr>
<td>SD</td>
</tr>
<tr>
<td>CV</td>
</tr>
<tr>
<td>Length of tibiotarsus</td>
</tr>
<tr>
<td>n</td>
</tr>
<tr>
<td>$\bar{X}$</td>
</tr>
<tr>
<td>SD</td>
</tr>
<tr>
<td>CV</td>
</tr>
<tr>
<td>Length of tarsometatarsus</td>
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<tr>
<td>n</td>
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<tr>
<td>$\bar{X}$</td>
</tr>
<tr>
<td>SD</td>
</tr>
<tr>
<td>CV</td>
</tr>
</tbody>
</table>
Table 3.—Posterior probability classification of stepwise discriminate function analysis for bones assigned to Pachyornis mappini and P. septentrionalis

<table>
<thead>
<tr>
<th>Character</th>
<th>P. mappini</th>
<th>P. septentrionalis</th>
<th>n</th>
<th>Percent misclassified</th>
</tr>
</thead>
<tbody>
<tr>
<td>Femur</td>
<td>11</td>
<td>1</td>
<td>12</td>
<td>8.3</td>
</tr>
<tr>
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<td>1</td>
<td>13</td>
<td>14</td>
<td>7.1</td>
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<td>Tibiotarsus</td>
<td>12</td>
<td>0</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>P. septentrionalis</td>
<td>0</td>
<td>12</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Tarsometatarsus</td>
<td>7</td>
<td>0</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>P. mappini</td>
<td>0</td>
<td>10</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>P. septentrionalis</td>
<td>0</td>
<td>10</td>
<td>10</td>
<td>0</td>
</tr>
</tbody>
</table>

mappini is not substantiated. In order to examine further the distinction between the two forms, I analyzed the data using a step-wise discriminate function-canonical analysis. The distinctness of the two groups was further verified and few of the elements were misclassified (Table 3).

What is the meaning of these differences? If, in fact, the two taxa do not represent distinct species, then very likely we are dealing with sexual size differences. Table 2 presents some basic statistical data for lengths of the hindlimb bones. The combined sample of measurements for the lengths of the femur and tarsometatarsus do not show CVs much higher than those for P. mappini alone, or for those of other moas (Table 1). It can be noted also that the CV of 2.79 for the tarsometatarsus length of P. septentrionalis (Table 2) is suspiciously low compared to CVs of other moas. It is my belief that these two skeletal populations probably represent different sexes and that Oliver's (1949) description of a new species was unwarranted. I therefore synonymize septentrionalis with mappini until firm evidence can be offered that they are distinct.

**Euryapteryx Haast, 1874**

**Synonym.**—Zelornis Oliver, 1949.

Oliver (1949:117–128) created the genus Zelornis for the species Euryapteryx exilis Hutton (the genotype) and Emeus haasti Rothschild. Archey (1941) considered the former to be a valid species in the genus Euryapteryx, while the latter he treated as a synonym of Euryapteryx gravis (p. 54). The diagnostic feature separating Zelornis from Euryapteryx was said to be the high arched culmen. Oliver (1949:110) admitted that there were no differences in the shape of the postcranial elements. The type-specimen of Z. exilis is a skeleton from Wangaehu in the Wanganui Museum. The skull was figured by Oliver (1949, figs. 92–94) where it is readily apparent that the premaxilla is considerably broken and abraded. The premaxilla of Z. exilis does not appear to differ in shape from those referred to Z. haasti. I have examined nearly all of the cranial material assigned to Zelornis haasti by Oliver (1949:127) and can find no important differences in size or shape that will distinguish it at the generic level from Euryapteryx. To my knowledge Scarlett (1972) is the only recent author to synonymize Zelornis with Euryapteryx, and I concur with his decision.

**Summary.**—Two sexually dimorphic species are recognized here—a moderately large form, E. gera- noides, present on both North and South islands, and a small species, E. curtus, confined to North Island.

**Euryapteryx curtus** (Owen, 1846)

**Synonyms.**—Euryapteryx exilis Hutton, 1897; Euryapteryx tane Oliver, 1949.

Euryapteryx curtus was a small species of moa, apparently confined to the North Island. The only morphological difference between E. curtus and E. exilis is in size (Archey, 1941:60), E. exilis being slightly larger (Table 4). In order to assess the morphological similarities in size and shape in Euryapteryx, I analyzed the measurements of the femur, tibiotarsus, and tarsometatarsus given by Archey (1941) and Oliver (1949), using principal components and canonical analyses. Figure 2 plots...
Table 4.—Statistics for species of *Euryapteryx* (measurements in mm)

<table>
<thead>
<tr>
<th>Character</th>
<th>E. tane</th>
<th>E. curtus</th>
<th>E. exilis</th>
<th>E. exilis + E. curtus</th>
<th>E. gravis</th>
<th>E. gravis + E. geranoides</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length of Femur</td>
<td>n</td>
<td>x</td>
<td>SD</td>
<td>CV</td>
<td>n</td>
<td>x</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>190.67</td>
<td>9.29</td>
<td>4.87</td>
<td>15</td>
<td>167.6</td>
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<tr>
<td></td>
<td>196.17</td>
<td>184.27</td>
<td>256.63</td>
<td>280.15</td>
<td>263.57</td>
<td>297.68</td>
</tr>
<tr>
<td></td>
<td>186.24</td>
<td>11.5</td>
<td>5.3</td>
<td>3.89</td>
<td>21.57</td>
<td>17.99</td>
</tr>
<tr>
<td>Length of Tibiotarsus</td>
<td>n</td>
<td>x</td>
<td>SD</td>
<td>CV</td>
<td>n</td>
<td>x</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>149</td>
<td>7.55</td>
<td>5.07</td>
<td>19</td>
<td>124.74</td>
</tr>
<tr>
<td></td>
<td>135.12</td>
<td>135.12</td>
<td>12.67</td>
<td>10.09</td>
<td>17</td>
<td>124.74</td>
</tr>
<tr>
<td></td>
<td>175.12</td>
<td>175.12</td>
<td>12.67</td>
<td>10.09</td>
<td>17</td>
<td>124.74</td>
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<tr>
<td></td>
<td>174.78</td>
<td>174.78</td>
<td>12.67</td>
<td>10.09</td>
<td>17</td>
<td>124.74</td>
</tr>
<tr>
<td></td>
<td>173.8</td>
<td>173.8</td>
<td>12.67</td>
<td>10.09</td>
<td>17</td>
<td>124.74</td>
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<td>12.67</td>
<td>10.09</td>
<td>17</td>
<td>124.74</td>
</tr>
</tbody>
</table>

group centroids for the first two canonical axes. The multivariate analysis substantiates the separation of *E. exilis* and *E. curtus*, but the centroids themselves as projected onto the first axis are only about two and a half SD units from each other (except for the tibiotarsus where they are almost four units apart). If one examines the statistics of the combined sample (Table 4), the population *E. exilis + E. curtus* has CVs of about 10–12 for bone lengths. This variability is comparable to that of *Pachyornis mappini*, and I believe it is likely that the difference between *E. exilis* and *E. curtus* is one of sexual size dimorphism. Most of the specimens in Archey's series of these two forms (1941, tables H, I) come from Doubtless Bay, North Island.

Oliver (1949:105) described a separate species, *E. tane*, for a small number of specimens that were larger than those of *E. curtus*. He (1949:123) noted that some of the leg bones of *E. tane* possibly belonged to what he called *Zelornis exilis*. It is readily apparent that the samples of *E. tane* and *E. exilis* are virtually identical (Table 4; Figure 2). There is little question, therefore, that *E. tane* should be synonymized with *E. curtus* as defined here.

**Euryapteryx geranoides** (Owen, 1848)

**Synonyms.**—*Euryapteryx geranoides* (Owen, 1870), *Zelornis haasti* (Rothschild, 1907).

The type-material of *E. geranoides* consists of a cranium, premaxilla, and mandible collected at Te Rangatapu, North Island, and housed in the British Museum (Natural History). According to Oliver (1949:106) the mandible belongs to *Anomalopteryx didiformis*, and the cranium and premaxilla, which may or may not be associated, belong to *Euryapteryx*. To my knowledge a lectotype has not yet been designated, but the cranium (BM 21687) would be a logical choice. The type cranium and premaxilla are smaller than most of the material attributed to *E. gravis* but larger than in *E. curtus*. Therefore, *E. geranoides* has been accepted as a distinct species of intermediate size by Archey (1941), Oliver, (1949), and Brodkorb (1963). It should be pointed out that there is apparently no postcranial material directly associated with cranial material attributable to *E. geranoides* (sensu stricto).

Material assigned to the large form known as *E. gravis* is abundant in South Island localities (the type-skeleton in the British Museum is from Kakani) but very rare on the North Island (Archey, 1941:54–56; Oliver, 1949:108–112). Statistics presented in Table 4 and the results of the canonical analyses shown in Figure 2 confirm the intermediate position of *E. geranoides* between "*E. exilis*" (*=E. curtus*) and *E. gravis*. The question is whether *E. geranoides* is a distinct species, and if not, to which species—the larger *E. gravis* or the smaller *E. curtus*—this skeletal population belongs. Recently, Scarlett (1972:21) suggested that *E. gera-
noides might be united with *E. curtus* once the gaps were eliminated.

I would like to suggest here that *E. geranoides* is conspecific with *E. gravis* and that these forms represent another case of sexual size dimorphism. The evidence is two-fold. First, measurements of the combined sample exhibit CVs very similar to those seen in *P. mappini*-*P. septentrionalis* and *E. curtus*-*E. exilis*, the two other presumed examples of sexual size dimorphism (Table 4). Secondly, bones attributed to *E. geranoides* and *E. gravis* occur on both North and South islands. If *E. geranoides* were conspecific with *E. curtus* (or with *E. exilis*, if this form were distinct from *E. curtus*), then the absence of *E. curtus* from the South Island is unexplained. If *E. geranoides* and *E. exilis* represented different sexes, then both should be present on the South Island. Present evidence, therefore, is more consistent with the hypothesis that *E. curtus*-*E. exilis* constitute one sexually dimorphic species and *E. geranoides*-*E. gravis* another. In the case of the latter, the older name, *geranoides*, has priority.

As noted above, the cranial material of *Zelornis haasti* is very similar to that of *E. geranoides*. The femur and tibiotarsus of the one skeleton of *Z. haasti* are somewhat larger than typical “*E. gravis*” (Oliver, 1949:128; see also Table 4), but the associated tarsometatarsus is easily within the size range of that form. Hence, it is likely that the few bones assigned to *Z. haasti* are large, perhaps aberrant bones of *E. geranoides*, and I follow Archey (1941) in synonymizing *haasti*.

**Emeus Reichenbach, 1852**

**Emeus crassus** (Owen, 1846)

*Synonym.*—*E. huttonii* (Owen, 1879).

There has been little difference of opinion about species-limits within *Emeus*. Most recent authors (Archey, 1941; Oliver, 1949; Brodkorb, 1963) have accepted two species, the large *crassus* and the smaller *huttonii*. Only Scarlett (1972:22) has combined the two species, stating that several specimens from Pyramid Valley are intermediate in
Table 5.—Statistics for species of *Emeus* (measurements in mm)

<table>
<thead>
<tr>
<th>Character</th>
<th><em>E. crassus</em></th>
<th><em>E. huttonii</em></th>
<th><em>E. crassus</em> + <em>E. huttonii</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Length of femur</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>21</td>
<td>10</td>
<td>31</td>
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<tr>
<td>( \bar{x} )</td>
<td>210.81</td>
<td>198.84</td>
<td>209.71</td>
</tr>
<tr>
<td>SD</td>
<td>12.23</td>
<td>11.92</td>
<td>20.24</td>
</tr>
<tr>
<td>CV</td>
<td>4.48</td>
<td>5</td>
<td>7.73</td>
</tr>
<tr>
<td>Length of tibiotarsus</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>21</td>
<td>11</td>
<td>32</td>
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<tr>
<td>( \bar{x} )</td>
<td>214.24</td>
<td>198.64</td>
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<tr>
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<td></td>
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</tr>
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<td>n</td>
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<tr>
<td>CV</td>
<td>5.42</td>
<td>7.3</td>
<td>8.91</td>
</tr>
</tbody>
</table>

Both forms occurred in the South Island and are known primarily from the Canterbury and Otago districts. Although according to Archey (1941:51) several bones referable to *Emeus* have allegedly been found at Martinborough and Te Aute on the North Island, Yaldwyn (1956) does not list *Emeus* from the Martinborough Caves, nor did Oliver (1949) make note of *Emeus* on the North Island. On the South Island *E. crassus* has been reported as moderately common, whereas *E. huttonii* was apparently less so.

Statistics presented in Table 5 show that the bones assigned to *E. crassus* and *E. huttonii* by Archey (1941) and Oliver (1949) comprise two distinct populations. Multivariate analysis of the hindlimb measurements verifies the univariate results, and in the posterior probability classification of the discriminate function analysis, few bones are misclassified (Table 6).

The two forms of *Emeus* could represent two species, or alternatively, they might represent sexual dimorphism within a single species. It is my opinion that the latter hypothesis is more probable. First, the combined sample of the two populations does not exhibit variability that could be considered outside the limits for a single species of moa, all CVs being fairly low (Table 5: femur length, 7.73; tibiotarsus length, 9.86; tarsometatarsus length, 8.91). Secondly, the two forms appear to have been broadly sympatric, both occurring together in the larger fossil deposits. To my knowledge there is no good evidence that one had a distribution exclusive of the other. If the two forms represent dimorphic sexes, then the larger form, *crassus*, would appear to have been the female, as an egg was found preserved in association with a skeleton of this form at Pyramid Valley (Falla, 1941). I therefore follow Scarlett (1972) in tentatively synonymizing *E. huttonii* with *E. crassus*.

Subfamily DINORNITHINAE

*Dinornis* Owen, 1843

**Summary.**—Four species of *Dinornis* are recognized here. The smallest, *D. struthoides*, was confined to the North Island. A second species, *D. torosus*, was only slightly larger than *D. struthoides* and was restricted to the South Island. A larger form, *D. novaeezalandiae*, and the largest species of moa, *D. giganteus*, were both found on the North and South islands.

Table 6.—Posterior probability classification of stepwise discriminate function analysis for bones assigned to *Emeus crassus* and *E. huttonii*

<table>
<thead>
<tr>
<th></th>
<th><em>E. crassus</em></th>
<th><em>E. huttonii</em></th>
<th>n</th>
<th>Percent misclassified</th>
</tr>
</thead>
<tbody>
<tr>
<td>Femur</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td><em>E. crassus</em></td>
<td></td>
<td>18</td>
<td>1</td>
<td>19</td>
</tr>
<tr>
<td><em>E. huttonii</em></td>
<td></td>
<td>0</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Tibiotarsus</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>E. crassus</em></td>
<td></td>
<td>20</td>
<td>0</td>
<td>20</td>
</tr>
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<td><em>E. huttonii</em></td>
<td></td>
<td>0</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Tarsometatarsus</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>E. crassus</em></td>
<td></td>
<td>21</td>
<td>1</td>
<td>22</td>
</tr>
<tr>
<td><em>E. huttonii</em></td>
<td></td>
<td>1</td>
<td>10</td>
<td>11</td>
</tr>
</tbody>
</table>
**Dinornis struthoides** Owen, 1844

**Synonym.**—*Dinornis gazella* Oliver, 1949.

Because of a decision by the International Commission of Zoological Nomenclature (Hemming, 1954), the name *D. struthoides* refers to all those specimens included under the name *D. novaezealandiae* in Archey (1941), Oliver (1949), and Brodkorb (1963). This species was the smallest of the genus and was found on the North Island. Oliver (1949:170) recorded two bones from the South Island which he assigned to this species, noting that some of these bones were “indistinguishable in size and proportions” from those of the North Island. Neither Archey (1941), Brodkorb (1963), nor Scarlett (1972) listed this species from the South Island and it may be that Oliver’s specimens are referable to small individuals of *D. torosus*. Thus, the presence of *D. struthoides* on the South Island needs to be verified (unfortunately, I did not examine the relevant material while in New Zealand). In any case, if *D. struthoides* were present on the South Island, it was evidently uncommon.

Oliver (1949:166) described a new species from the North Island, *D. gazella*, based on a pelvis and some referred bones that are smaller than those typical of *D. struthoides* (Table 7). The size differences are slight, however, and the variation exhibited by the combined sample of *struthoides* and *gazella* is easily within the range of a single species (Table 8). I compared a series of bones (DM 108: 5 tarsometatarsi, 3 tibiotarsi from Te Aute) referred by Oliver to *D. gazella* with bones of *D. struthoides* and found that those of the former had thinner shafts although the two samples were nearly equal in length. The differences appear to be entirely related to age, as the bones referred to *D. gazella* are those of immature individuals. Therefore, I suggest that *D. gazella* be merged with *D. struthoides*.

**Dinornis torosus** Hutton, 1891

This species is only slightly larger than *D. struthoides* and has been considered the South Island counterpart of that species (e.g., Archey, 1941:62; Oliver, 1930). Indeed, I would be inclined to merge *D. torosus* with *D. struthoides* if it were not for some significant differences that apparently exist in cranial structure.

As I will detail in subsequent papers on evolutionary relationships and cranial morphology, *D. torosus* appears to be more advanced in cranial structure than *D. struthoides* but in certain other respects is more primitive than the two larger species *D. novaezealandiae* and *D. giganteus*. The skull of *D. torosus* differs from that of *D. struthoides* in having the basisphenoid rostrum moderately inflated, and the mandible stouter and less deflected ventrally. The most important and consistent difference seems to be in the nature of the

### Table 7.—Statistics for species of Dinornis (measurements in mm)

<table>
<thead>
<tr>
<th>Character</th>
<th><em>D. struthoides</em></th>
<th><em>D. gazella</em></th>
<th><em>D. torosus</em></th>
<th><em>D. novaezealandiae</em></th>
<th><em>D. robustus</em></th>
<th><em>D. hercules</em></th>
<th><em>D. maximus</em></th>
<th><em>D. giganteus</em></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Length of femur</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>n</td>
<td>12</td>
<td>1</td>
<td>15</td>
<td>15</td>
<td>12</td>
<td>1</td>
<td>23</td>
<td>8</td>
</tr>
<tr>
<td>(\bar{x})</td>
<td>265</td>
<td>231</td>
<td>295.33</td>
<td>340.53</td>
<td>354.5</td>
<td>353</td>
<td>406.13</td>
<td>390.88</td>
</tr>
<tr>
<td>SD</td>
<td>17.41</td>
<td>--</td>
<td>18.35</td>
<td>15.14</td>
<td>15.47</td>
<td>--</td>
<td>20.73</td>
<td>22.2</td>
</tr>
<tr>
<td>CV</td>
<td>6.57</td>
<td>--</td>
<td>6.21</td>
<td>3.86</td>
<td>4.36</td>
<td>--</td>
<td>5.1</td>
<td>5.68</td>
</tr>
<tr>
<td><strong>Length of tibiotarsus</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<td>n</td>
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<td>3</td>
<td>14</td>
<td>21</td>
<td>15</td>
<td>5</td>
<td>32</td>
<td>10</td>
</tr>
<tr>
<td>(\bar{x})</td>
<td>520.67</td>
<td>469.67</td>
<td>588.36</td>
<td>701.67</td>
<td>718.6</td>
<td>779.8</td>
<td>866.63</td>
<td>875.2</td>
</tr>
<tr>
<td>SD</td>
<td>29.08</td>
<td>--</td>
<td>36.77</td>
<td>40.87</td>
<td>22.8</td>
<td>37.99</td>
<td>46.54</td>
<td>71.57</td>
</tr>
<tr>
<td>CV</td>
<td>5.59</td>
<td>--</td>
<td>6.25</td>
<td>5.82</td>
<td>3.17</td>
<td>4.87</td>
<td>5.37</td>
<td>8.18</td>
</tr>
<tr>
<td><strong>Length of tarsometatarsus</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
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<td>n</td>
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<td>5</td>
<td>16</td>
<td>13</td>
<td>14</td>
<td>3</td>
<td>29</td>
<td>10</td>
</tr>
<tr>
<td>(\bar{x})</td>
<td>282.2</td>
<td>259</td>
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<td>368.08</td>
<td>382.64</td>
<td>420.33</td>
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<tr>
<td>SD</td>
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<td>22.75</td>
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<td>17.72</td>
<td>--</td>
<td>38.78</td>
<td>30.08</td>
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<tr>
<td>CV</td>
<td>6.83</td>
<td>4.8</td>
<td>7.51</td>
<td>8.3</td>
<td>4.63</td>
<td>--</td>
<td>8.37</td>
<td>6.19</td>
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</tbody>
</table>
Table 8.—Statistics for combined species of Dinornis (measurements in mm)

<table>
<thead>
<tr>
<th>Character</th>
<th>$D. struthoides + D. gazella$</th>
<th>$D. novaezealandiae + D. robustus + D. hercules$</th>
<th>$D. giganteus + D. maximus$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length of femur</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>13</td>
<td>28</td>
<td>31</td>
</tr>
<tr>
<td>$\bar{x}$</td>
<td>262.38</td>
<td>346.96</td>
<td>402.19</td>
</tr>
<tr>
<td>SD</td>
<td>19.15</td>
<td>15.38</td>
<td>21.82</td>
</tr>
<tr>
<td>CV</td>
<td>7.3</td>
<td>4.43</td>
<td>5.42</td>
</tr>
<tr>
<td>Length of tibiotarsus</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>12</td>
<td>41</td>
<td>43</td>
</tr>
<tr>
<td>$\bar{x}$</td>
<td>507.92</td>
<td>717.39</td>
<td>866.91</td>
</tr>
<tr>
<td>SD</td>
<td>34.43</td>
<td>42.17</td>
<td>53.32</td>
</tr>
<tr>
<td>CV</td>
<td>6.78</td>
<td>5.88</td>
<td>6.15</td>
</tr>
<tr>
<td>Length of tarsometatarsus</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>20</td>
<td>30</td>
<td>39</td>
</tr>
<tr>
<td>$\bar{x}$</td>
<td>276.4</td>
<td>380.1</td>
<td>469.23</td>
</tr>
<tr>
<td>SD</td>
<td>20.3</td>
<td>27.79</td>
<td>37.75</td>
</tr>
<tr>
<td>CV</td>
<td>7.35</td>
<td>7.31</td>
<td>8.04</td>
</tr>
</tbody>
</table>

basisphenoid rostrum. In $D. giganteus$, and to a lesser extent in $D. novaezealandiae$, there is a marked inflation of the basisphenoid rostrum as compared to the condition seen in $D. torosus$. Based on the comparative series I was able to examine, these differences are consistent between species.

**Dinornis novaezealandiae** Owen, 1843

**Synonyms.**—*Dinornis ingens* Owen, 1844; *D. robustus* Owen, 1846; *D. hercules* Oliver, 1949.

As a result of the above-mentioned decision of the International Commission of Zoological Nomenclature (Hemming, 1954), *D. ingens* now becomes a synonym of *D. novaezealandiae*. Consequently, the latter name belongs to those forms of *Dinornis* that were larger than *D. struthoides* and *D. torosus* but smaller than *D. giganteus*. Bones referred to *D. novaezealandiae* (=*D. ingens* of Archey, Oliver, and Brodkorb) are known from both the North and South islands, although they were apparently less common in the latter. I include *D. robustus* in this species because it appears to be little more than the South Island representative of *D. novaezealandiae*. Bones referred to *D. novaezealandiae* (=*D. ingens* of Archey, Oliver, and Brodkorb) are known from both the North and South islands, although they were apparently less common in the latter. I include *D. robustus* in this species because it appears to be little more than the South Island representative of *D. novaezealandiae*. Bones referred to the two species overlap considerably in length (Table 7), but those included under the name *D. robustus* are somewhat stouter than those assigned to *D. novaezealandiae* (Archey, 1941:71, Oliver, 1949: 171).

Oliver (1949:174) described an additional species, *D. hercules*, from a few limb bones from the North Island. The type tibiotarsus (DM 217) is about the same length as some tibiotarsi of *D. novaezealandiae* but has the shaft more curved; in general, bones assigned to *D. hercules* by Oliver are slightly larger than those of *D. novaezealandiae* (Table 7). Scarlett (1972:21) suggested that *D. hercules* may be a "bow-legged variant" of *D. giganteus*, but I believe most of the specimens assigned to *D. hercules* by Oliver are closer to *D. novaezealandiae* in size and I here include it with that species. I was able to compare the type of *D. hercules* with other species of *Dinornis* and in my opinion the differences in stoutness and the curvature of the shaft of the type are attributable to individual variation.

The combined sample of bones of *D. novaezealandiae*, *D. robustus*, and *D. hercules* exhibits very little variation, all CVs being less than 7.50 (Table 8). This variability is well within that for a single species of moa.

**Dinornis giganteus** Owen, 1844

**Synonym.**—*Dinornis maximus* Owen, 1867.

*Dinornis giganteus* is the North Island representative of the largest species of moa and *D. maximus* is its South Island form. Bones of the latter are somewhat stouter, but measurements of the two overlap greatly (Table 7). I can see little
value in considering these minor variations to be indicative of species differences. The combined samples show a relatively small amount of variability, certainly within the limits of a single species (Table 8). Judging from the available collections, the North Island form was less common than the South Island form.

Conclusions

In contrast to the 20 species recognized by Archey (1941) and 29 species recognized by Oliver (1949), I here accept only 13 species as being valid. The present arrangement is actually fairly similar in parts to Archey's, but combines several North and South islands counterparts, while several “species-pairs” are regarded as examples of sexual dimorphism. Further study may show that Anomalopteryx oweni and Megalapteryx benhami, which are based on somewhat dubious material, perhaps do not deserve recognition. Certainly the acceptance of the large number of species advocated by Oliver is untenable.

The systematic results of this paper can be summarized by the following classification:

**Family Dinornithidae**

**Subfamily Anomalopteryginae**

- Genus *Anomalopteryx* Reichenbach, 1852
  - *A. didiformis* (Owen, 1844)
  - *A. oweni* (Haast, 1885)

- Genus *Megalapteryx* Haast, 1886
  - *M. didinus* (Owen, 1883)
  - *M. benhami* Archey, 1941

**Subfamily Dinornithinae**

- Genus *Pachyornis* Lydekker, 1891
  - *P. elephantopus* (Owen, 1856)
  - *P. mapipini* Archey, 1941

- Genus *Euryapteryx* Haast, 1874
  - *E. curtus* (Owen, 1846)
  - *E. geranoides* (Owen, 1848)

- Genus *Emeus* Reichenbach, 1852
  - *E. crassus* (Owen, 1846)

- Genus *Dinornis* Owen, 1843
  - *D. struthoides* Owen, 1844
  - *D. torosus* Hutton, 1891
  - *D. novaezealandiae* Owen, 1843
  - *D. giganteus* Owen, 1844

The 13 species recognized here appear to have been distributed as follows:

<table>
<thead>
<tr>
<th>North Island</th>
<th>South Island</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Anomalopteryx didiformis</em></td>
<td><em>Anomalopteryx didiformis</em></td>
</tr>
<tr>
<td><em>A. oweni</em></td>
<td><em>Megalapteryx didinus</em></td>
</tr>
<tr>
<td><em>Pachyornis mappini</em></td>
<td><em>Pachyornis elephantopus</em></td>
</tr>
<tr>
<td><em>Euryapteryx geranoides</em></td>
<td><em>Euryapteryx geranoides</em></td>
</tr>
<tr>
<td><em>Emeus crassus</em> (uncertain)</td>
<td><em>Emeus crassus</em></td>
</tr>
<tr>
<td><em>Dinornis struthoides</em></td>
<td><em>Dinornis struthoides</em> (uncertain)</td>
</tr>
<tr>
<td><em>D. novaezealandiae</em></td>
<td><em>D. novaezealandiae</em></td>
</tr>
<tr>
<td><em>D. giganteus</em></td>
<td><em>D. torosus</em></td>
</tr>
</tbody>
</table>

It is of interest to note that the smaller and topographically less diverse North Island had fewer species than the South Island. Also, many of the North Island forms appear to have been smaller than their South Island counterparts.

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Goode, W.

Haast, J. von

Hemming, F., editor

Lambrecht, K.

Long, J. L.

Lydekker, R.

Oliver, W. R. B.

Oxnard, C. E.

Scarlett, R. J.

Simpson, G. G.

Yaldwyn, J. C.
Evidence of the Survival to Recent Times of the Extinct Flightless Duck *Chendytes lawi* Miller

G. Victor Morejohn

**ABSTRACT**

Skeletal remains of the extinct late Pleistocene flightless diving duck, *Chendytes lawi*, were discovered at two northern California Indian midden sites. Carbon 14 dates of midden shell and aspartic acid racemization of the *Chendytes* bones showed that this species lived into the Holocene and became extinct sometime after 3780 years BP. The remains from these northern California middens and a newly discovered Pleistocene tibiotarsus from the Port Orford Formation of Oregon extend the known range of *Chendytes laid* from the Channel Islands of southern California northward some 720 kilometers.

**Introduction**

Study of avian skeletal remains from an Indian midden on the south shore of Laguna Creek, Santa Cruz County, California, has revealed many elements which unquestionably belong to the extinct, flightless diving duck, *Chendytes lawi* Miller. Two bones of this species were also found in another midden on Año Nuevo Point, about 29 km north of the city of Santa Cruz. Previously, all certain records of *Chendytes* were from Pleistocene deposits. The present account documents its persistence into the Holocene and its contemporaneity with aboriginal man.

**Acknowledgments.**—I wish to thank Kenneth R. Lajoie and Edward Helly of the U.S. Geological Survey for providing Carbon 14 dates and assistance in field work. Jeffery L. Bada of Scripps Institution of Oceanography kindly determined the age of bone fragments through aspartic acid racemization. Burton L. Gordon of San Francisco State University provided many bone fragments found at the Laguna Creek midden, some of which belonged to *Chendytes*. Hildegarde Howard assisted in the original identifications of bones and supplied useful comments on the study. Eric Anderson, Janice Cross, Howard Hutchison, Diana Matthiesen, and Bruce Welton were helpful in the field or assisted me in the museum collections.

**Description and Age of the Sites**

The Laguna Creek midden (State of California Archeological site designation—Santa Cruz 7, hereafter abbreviated as CA-SCr-7) is situated atop a large sand dune (Sandhill Bluff) on a marine terrace some 25 meters above sea level at latitude 36°58'30" N and longitude 122°09'10" W, approximately 12 km north of Santa Cruz, Santa Cruz County, California, and about 230 meters west of U.S. Highway No. 1. The dune occupies an area of about 915 m², with the midden capping the highest part and occupying an area of about 120 m². The windswept seaward side of the midden is greatly eroded, in places to a depth such that the underlying dune substrate is visible. Most of the bones were collected from this exposed face. Bones of *Chendytes lawi* were found from 15 cm below the surface to the basal layer of the midden, and several others were found scattered on the surface of the dune below the exposed face.

Most of the midden consists of broken mollusk and barnacle shells tightly packed to a depth varying between 1.5 and 1.75 m. Early reports (Stearns, 1873–1874:157) indicate that this midden was
originally some 6.5 m high, but recent local use for shell to feed poultry has greatly reduced the size of the midden and removed the younger upper layers. Samples of mollusk and barnacle shells from the upper and lower midden levels were used for Carbon 14 dating. The uppermost layer was aged at 3780 ± 95 years BP and the basal layer was aged at 5390 ± 100 years BP. Two bones of *Chendytes* from the basal layer were determined by aspartic acid racemization (Bada, et al., 1974) to be approximately 6000 years old, which corresponds closely with the Carbon 14 dates of other material from the same layer. Fragments of a human femur and an incomplete skull lacking the rostrum and lower jaw were also found in the basal layer, but the aspartic acid age of these fragments was approximately 4000 years BP, obviously indicating secondary introduction through burial. The above datings establish CA-SCr-7 as the oldest known site of Indian occupation on the central California coast (M. J. Moratto, in litt., 19 December 1974). The Indian midden on Ano Nuevo Point, San Mateo County, California, has not been studied. Archeologists have estimated its age at about 2500 years BP, but it is probably older. Persistent seasonal winds expose part of the midden or cover it with sand. Only a few vertebrate fragments have so far been collected.

The following bones of *Chendytes* were recovered from the Laguna Creek and Ano Nuevo middens: humeri, 1 left, 2 right proximal ends; femora, 8 left, 7 right nearly complete, 4 left, 5 right incomplete; tibiotarsi, 1 left, 5 right proximal ends, 3 left, 3 right distal ends, 10 partial shafts; tarsometatarsus, 1 left incomplete; synsacrum, 1 incomplete; cervical vertebra, 1 incomplete.

**Discussion**

Because comparisons were initially made with reference specimens of living taxa, the elements of *Chendytes* collected at CA-SCr-7 at first proved impossible to identify. Later, while studying anseriform fossils with Dr. Hildegarde Howard at the Natural History Museum of Los Angeles County, I examined specimens of *Chendytes* that I then recognized as being similar to the unidentified bones from CA-SCr-7. In size (Table 1) and morphology, the midden bones were found to conform with those previously described for *Chendytes lawi* by Miller (1925; 1930), Miller, et al. (1961), and Howard (1947, 1949, 1955). I later noted several characteristics of the tibiotarsi and femora that had not previously been considered by these authors.

Miller (1925) first recognized the affinities of *Chendytes* with the diving ducks, particularly the scoters. Compared to *Melanitta fusca*, the largest of the scoters, I found great reduction in the marrow cavities of the tibiotarsi and femora, with a concomitant increase in the thickness of the peripheral compact bone in *C. lawi* (Figure 1a). In fact, certain cross-sections of the femora of *Chendytes* bear a similarity to those of mammalian
femora. The density of the bone in *Chendytes* is probably an adaptation for diving. Cross-sections of the tibiotarsus proximal to the fibular crest show *M. fusca* to have a pronounced posterior ridge, whereas in *Chendytes* this ridge is low and rounded (Figure 1c). Cross-sections through the fibular crest show *M. fusca* and *Chendytes* to have a flattened anterior and a ridged posterior surface, but in *Chendytes* the inner cnemial crest extends distally to about the middle of the fibular crest and angles anteriorly (Figure 1b). The outer cnemial crest in anterior view is also markedly different in the two species (Figure 1d).

Howard (1955) reported a total of 156 fossil skeletal remains of *Chendytes* (*C. lawi*, 89; *C. milleri*, 67) from eleven localities in southern California. Since then, another 130 specimens have been recovered from Anacapa Island (Howard, 1964). While most of the bones were clearly from Pleistocene localities, fifteen were reportedly collected from two Indian middens (one at Malaga Cove in the Palos Verdes area of Los Angeles County, the other on San Nicolas Island). However, these bones are well mineralized, unlike those typically found in middens, and they were considered by Howard (1955) to have been secondarily associated with the Indian midden material. In contrast, bones of *C. lawi* from the Laguna Creek and Año Nuevo Point Indian middens in northern California were not mineralized.

Although bones of a variety of birds have been recovered from other Indian middens along the California coast (Howard, 1929; Howard and Dodson, 1933; and D. M. Howard and Cook, 1971) and demonstrate that a number of avian species were eaten by Indians (Table 2), no other elements of *Chendytes* have thus far been found. Howard (1929), for lack of comparative anseriform skeletal material, chose not to attempt identification of waterfowl in her study of the avian remains from the Emeryville shellmound. I recently reexamined the anseriform material from this site in an attempt to find elements of *Chendytes*, but met with no success.

The Emeryville site was estimated to be no younger than 2310 ± 6 years (Hubbs, et al., 1962). *Chendytes* may have been extinct by this time, or its absence from the Emeryville site may have been due to the lack of appropriate habitat. The site is located on an alluvial plane (Howard, 1929) with-

| Table 1.—Comparison of measurements (mm) of bones of *Chendytes lawi* from northern and southern California localities |
|---|---|---|---|
| Character | Northern California | Southern California* |
| | | range | range |
| **HUMERUS** | | | |
| Breadth of proximal end from external to internal tuberosity | 3 | 13.3 | 13-13.5 | 14.3 |
| Breadth of shaft below external tuberosity | 3 | 4.3 | 4-4.5 | 4.8 |
| **FEMUR** | | | |
| Length | 5 | 73.2 | 69.6-77.5 | 65.7-76.4 |
| Breadth of proximal end | 9 | 16.1 | 15.2-17.8 | 14-17 |
| Breadth of distal end | 6 | 19 | 17.7-20.4 | 16.3-18.2 |
| **TIBIOTARSUS** | | | |
| Breadth of proximal end | 3 | 15 | 14.4-15.7 | 14.8-15.5 |
| Breadth of distal end | 3 | 13.7 | 13.3-14.5 | 13.3-14.5 |

* From Howard, 1955.

| Table 2.—Relative abundance of avian species found associated with remains of *Chendytes lawi* at the Laguna Creek Indian midden |
|---|---|
| Species | Number of specimens |
| Gavia stellata | 1 |
| Puffinus griseus | 6 |
| Fulmarus glacialis | 2 |
| Unidentified albatross | 23 |
| Diomedea albatrus | 10 |
| Unidentified cormorant | 2 |
| Phalacrocorax penicillatus | 3 |
| Phalacrocorax pelagicus | 1 |
| Unidentified anseriform | 12 |
| Unidentified duck | 6 |
| Melanitta fusca | 2 |
| Chendytes lawi | 46 |
| Unidentified falconiform | 1 |
| Buteo jamaicensis | 1 |
| Unidentified galliform | 1 |
| Unidentified charadriiform | 1 |
| Uria aalge | 29 |
| Cepphus columba | 6 |
| Pychoramphus aleutica | 1 |
| Unidentified gull | 26 |
| Unidentified bird | 99 |
out nearby reefs or rocky shorelines such as were probably required by *Chendytes* (Miller, et al., 1961). The elements of *Chendytes lawi* from CA-SCR-7, which may be at least as young as 3780 years, provide the latest known occurrence of the genus. At some time after deposition of these bones, perhaps 2500 to 3000 years ago, *Chendytes* became extinct.

The known fossil localities of *C. lawi* are concentrated in southern California and the range of the species was formerly considered by Miller, et al. (1961:10) to be "south of Point Conception, from Ventura County to Orange County." Although the material from CA-SCR-7 extends this range some 320 km farther northward, a hitherto unreported, nearly complete, fossilized tibiotarsus of *C. lawi* (University of California Museum of Paleontology No. 112026) collected by David Taylor from the lower Pleistocene Port Orford Formation in Curry County, Oregon, near Cape Blanco (UCMP vertebrate locality V-74042) extends the range of this species some 720 km north of the southern California sites (Figure 2).

Evolution of flightlessness in smaller birds is usually associated with insular distribution (Olson, 1973:31–36). Being flightless, the breeding sites of *Chendytes* almost certainly had to be restricted to offshore islands, and it may be assumed that the Channel Islands were of particular importance in this regard. There is no geological evidence of former islands in the immediate vicinity of the Laguna Creek area. Therefore, *C. lawi* was probably present in this area as a nonbreeding migrant.

The frequency of occurrence of bones of *Chendytes lawi* at CA-SCR-7 may indicate selective predation by Indians there. The bird was flightless, was of small size, relatively easy to capture, and had large goose-sized legs which would have made it a preferred food item. The preponderance of leg bones at CA-SCR-7 supports this contention. Prob-

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