- 1 Pterosaur integumentary structures with complex feather-like branching
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Pterosaurs were the first vertebrates to achieve true flapping flight, but in the 20 absence of living representatives, many questions concerning their biology and 21 lifestyle remain unresolved. Pycnofibres, the integumentary coverings of 22 pterosaurs, are particularly enigmatic: although many reconstructions depict 23 fur-like coverings composed of pycnofibres, their affinities and function are not 24 25 fully understood. Here we report the preservation in two anurognathid pterosaur specimens of morphologically diverse pycnofibres that show diagnostic features 26 of feathers, including non-vaned grouped filaments and bilaterally branched 27 filaments, hitherto considered unique to maniraptoran dinosaurs, and preserved 28 melanosomes with diverse geometries. These findings could imply that feathers 29 had deep evolutionary origins in ancestral archosaurs, or that these structures 30 arose independently in pterosaurs. The presence of feather-like structures 31 32 suggests that anurognathids, and potentially other pterosaurs, possessed a dense filamentous covering that likely functioned in thermoregulation, tactile sensing, 33 signalling, and aerodynamics. 34

Feathers are the most complex integumentary appendages in vertebrates¹. Most 35 feathers in modern birds possess an axial shaft from which branch lateral barbs and 36 barbules. Much is known about the anatomy, developmental biology, and genomic 37 regulation of these structures, but their deep evolutionary origin is controversial²⁻⁴. 38 Feathers and feather-like integumentary structures have been reported in many 39 theropod dinosaurs (including birds)^{3,5} and ornithischians such as *Psittacosaurus*⁶, 40 Tianyulong⁷, and Kulindadromeus⁸. Feather-like or hair-like structures, termed 41 pycnofibres⁹, have also been reported in several pterosaur specimens⁹⁻¹³, but their 42 nature is not resolved. 43

Here we report remarkably well-preserved pycnofibres in two anurognathid 44 pterosaurs and demonstrate, using evidence from morphology, chemistry and 45 macroevolutionary analyses, that the preserved pycnofibres bear key features of 46 feathers: monofilaments, two types of non-vaned grouped filaments, bilaterally 47 48 branched filaments that were previously considered unique to maniraptoran dinosaurs, and preserved melanosomes with diverse geometries. Both specimens studied are 49 from the Middle-Late Jurassic Yanliao Biota (ca. 165-160 Mya¹⁴). NJU-57003 50 (Nanjing University) is a newly excavated specimen from the Mutoudeng locality and 51 CAGS-Z070 (Institute of Geology, Chinese Academy of Geological Sciences), which 52 has been noted briefly for its feather-like branched pycnofibres¹³, is from the 53 Daohugou locality. Both specimens are near-complete and well-articulated, with 54 extensive soft tissues (Figs. 1 and 2, and Supplementary Figs. 1-5). Both specimens 55 are identified as anurognathids¹⁷ (see Supplementary text for osteological 56 descriptions). 57

Preserved soft tissues include structural fibres (actinofibrils) and pycnofibres. Structural fibres, common in the pterosaur wing membrane^{9,12,18}, are observed only in the posterior portion of the uropatagium in CAGS–Z070 (Fig. 1**o**–**p**). As reported elsewhere, they are parallel to subparallel and closely packed. Individual fibres are 0.08–0.11 mm wide (ca. 5 fibres per mm) and at least 1.9 mm long. Pycnofibres are preserved extensively in both pterosaur specimens (especially CAGS–Z070; Figs. 1 and 2, and Supplementary Figs. 1, 4 and 5) and are discriminated from structural
fibres based on their curved morphology and overlapping arrangement. In the
posterior portion of the uropatagium in CAGS–Z070, pycnofibres co-occur with
structural fibres; oblique intersections reflect superposition of these features during
decay (Fig. 1o-p).

Pycnofibres are categorized here into four types. Type 1 occurs around the head, 69 neck, shoulder, torso, all four limbs and tail of both specimens (Figs. 1c-e, o-p, 2b-c 70 and f). It comprises curved monofilaments that are 3.5-12.8 mm long and 70-430 µm 71 wide. Some short, distally tapering examples discriminate between dark-toned lateral 72 margins and light-toned axial regions, especially near the filament base where the 73 light-toned axis is wider, suggesting a tube-like morphology (Fig. 1c-e). Type 2 is 74 preserved in the neck, proximal forelimb, plantar metatarsus and proximal tail regions 75 76 of CAGS-Z070. It consists of bundles of curved filaments of similar length that appear to form brush-like structures at the distal ends of thicker filaments (2.0-13.8 77 mm long and 80–180 µm wide) (Fig. 1f-h). The latter may represent individual thick 78 filaments or fused proximal regions of thinner distal filaments. Type 3 occurs around 79 80 the head of CAGS-Z070. It comprises straight to slightly curved, distally tapered, central filaments (4.5-7.0 mm long and 50-450 µm wide) with short lateral branches 81 that diverge from the central filament near the midpoint (Fig. 1i-k). There are five 82 Type 3 filaments identified on the head, next to five similar filaments likely of the 83 same nature but obscured by overlapping filaments (Supplementary Fig. 5b). Type 4 84 occurs on the wing membrane of both specimens. It comprises tufts of curved 85 filaments (2.5–8.0 mm long and 70–130 µm wide) that diverge proximally (Figs. 11–n 86 and 2d-e), in contrast to the clear separation between Type 1 filaments (Fig. 1o-p). 87

Filamentous integumentary structures in extant and fossil vertebrates commonly 88 contain melanin-bearing organelles (melanosomes). Scanning electron microscopy 89 (SEM) of the filamentous structures of NJU-57003 reveals densely packed 90 microbodies $0.70 \pm 0.11 \,\mu\text{m}$ long and $0.32 \pm 0.05 \,\mu\text{m}$ wide (Fig. 2g-h, Supplementary 91 Figs. 4a–f, 6 and 7, and Supplementary Table 2). As with most melanosome-rich 92 fossil feathers¹⁹⁻²¹, energy dispersive X-ray spectroscopy (EDS) spectra of the 93 filaments are dominated by a major peak for carbon (Supplementary Fig. 8). These 94 carbonaceous microbodies resemble fossil melanosomes in terms of their geometry, 95 dense packing, parallel alignment relative to the long axis of the integumentary 96 structure (i.e. barbules in Paraves), and preservation within the matrix of the filament 97 (see Supplementary text). Most of the microbodies are oblate and morphologically 98 similar to those that are usually interpreted as phaeomelanosomes in fossils¹⁹ (Fig. 99 2h). Rod-shaped examples, usually interpreted as eumelanosomes in fossils¹⁹ (Fig. 100 2g), are rare. 101

Fourier transform infrared spectroscopy (FTIR) of samples of pterosaur filaments shows four major peaks unique to the filaments (Fig. 2i). These peaks are consistent with the absorption regions of amide I at ca. 1650 cm⁻¹ (principally the C=O asymmetric stretching vibration with some C–N bending), amide II at ca. 1540 cm⁻¹ (a combination of N–H in-plane bending and C–N and C–C stretching as in indole and pyrrole in melanin and amino acids), and aliphatic C–H stretching at 2850 cm⁻¹

and 2918 cm^{-1 22}. These peaks also occur in spectra obtained from extant feathers^{21,23}, 108 fossil feathers of the paravian Anchiornis²⁰, and melanosomes isolated from human 109 hair²⁴. Further, spectra of the pterosaur filaments more closely resemble those of 110 pheomelanin-rich red human hair in the stronger absorption regions at ca. 2850 cm⁻¹ 111 and 2918 cm^{-1} and higher resolution in the region ca. 1500–1700 cm^{-1} than those 112 from eumelanin-rich black human hair and the ink sac of cuttlefish²⁴. This, together 113 with the SEM results, suggests that the densely packed microbodies in the pterosaur 114 filaments are preserved melanosomes. The amide I peak at 1650 cm⁻¹ is more 115 consistent with α -keratin (characteristic of extant mammal hair²⁵) than β -keratin (the 116 primary keratin in extant avian feathers^{22,26}). This signal may be original or 117 diagenetic; the molecular configuration of keratin²⁶ and other proteins²⁷ can alter 118 under mechanical stress and changes in hydration levels. 119

120 The ultrastructural and chemical features of the pterosaur filaments confirm that they are hair-like or feather-like integumentary structures. The four types of filaments 121 described here show distinct distributions and morphologies. They are separated 122 clearly from the sedimentary matrix by sharp boundaries (Supplementary Fig. 4g-i). 123 There is no evidence that one or more filament type(s) were generated 124 taphonomically, e.g. through selective degradation or fossilization, or superimposition 125 of filaments. For instance, although Type 1 and 4 filaments occur widely in both 126 specimens, Type 4 occurs only in the wings, while Type 1 occupies the remaining 127 body regions. Type 1 filaments are thus not degraded products of Type 4, and Type 4 128 filaments do not represent superimposed clusters of Type 1 filaments. Filament types 129 2 and 3 occur only in CAGS-Z070. Type 3 occurs only in the facial area and is 130 associated with Type 1, where Types 2 and 4 are not evident. Type 3 filaments are 131 thus not degraded Type 2 or 4 filaments. Central filaments of Type 3 are 132 morphologically identical to the short, distally tapering filaments of Type 1, but the 133 branching filaments are much thinner (< 40 µm (Type 3) versus >70 µm (Type 1) 134 wide) and shorter (< 0.6 mm vs. > 3.5 mm long) than the latter. The branching 135 filaments are thus unlikely to reflect superimposition of clusters of Type 1 filaments. 136 In contrast, the distal ends of Type 2 filaments are similar, and have a similar 137 distribution pattern to, Type 1 filaments. An alternative interpretation, that Type 2 138 filaments might represent superimposition of Type 1 filaments at their proximal ends, 139 is unlikely (see detailed discussion in Supplementary text). Feathers and feather-like 140 integumentary structures have been reported in non-avian dinosaurs, although debate 141 continues about their true nature². These structures have been ascribed to several 142 morphotypes, some absent in living birds^{3,5}, and provide a basis to analyse the 143 evolutionary significance of pterosaur pycnofibres. The pterosaur Type 1 filaments 144 resemble monofilaments in the ornithischian dinosaurs Tianyulong and Psittacosaurus 145 and the coelurosaur Beipiaosaurus: unbranched, cylindrical structures with a midline 146 groove that widens towards the base (presumed in *Beipiaosaurus*)^{3,5}. The pterosaur 147 Type 2 filaments resemble the brush-like bundles of filaments in the coelurosaurs 148 *Epidexipteryx* and *Yi*^{3,5,28}: both comprise parallel filaments that unite proximally. The 149 morphology and circum-cranial distribution of pterosaur Type 3 filaments resemble 150 bristles in modern birds¹, but surprisingly do not correspond to any reported 151

morphotype in non-avian dinosaurs. The Type 3 filaments recall bilaterally branched
filaments in *Sinornithosaurus*, *Anchiornis*, and *Dilong*, but the latter filaments branch
throughout their length rather than halfway along the central filament(s), as in the
pterosaur structure^{3,5}. The pterosaur Type 4 filaments are identical to the radially
branched, downy feather-like morphotype found widely in coelurosaurs such as *Sinornithosaurus*, *Beipiaosaurus*, *Protarchaeopteryx*, *Caudipteryx*, and *Dilong*^{3,5}.

The filamentous integumentary structures in our anurognathid pterosaurs are thus 158 remarkably similar to feathers and feather-like structures in non-avian dinosaurs. 159 Intriguingly, cylindrical (Type 1), radially symmetrical branched (Types 2 and 4) and 160 bilaterally symmetrical branched (Type 3) filaments clearly coexisted in individual 161 animals; these structures may represent transitional forms in the evolution of feathers, 162 as revealed by developmental studies^{3,5}. These new findings warrant revision of the 163 origin of complex feather-like branching integumentary structures from Dinosauria to 164 Avemetatarsalia, the wider clade that includes dinosaurs, pterosaurs, and close 165 relatives^{4,29}. The early evolutionary history of bird feathers and homologous structures 166 in dinosaurs, and the multiple complex pycnofibres of pterosaurs, is enigmatic. A 167 previous study concluded that the common ancestor of these clades bore scales and 168 not filamentous integumentary appendages², but this result emerged only when the 169 filaments of pterosaurs were coded as non-homologous with those of dinosaurs. There 170 are no morphological criteria, however, for such a determination. The presence of 171 multiple pycnofibre types and their morphological, ultrastructural and chemical 172 similarity to feathers and feather-like structures in various dinosaurian clades, 173 confirms their likely homology with filamentous structures in non-avian dinosaurs 174 and birds. Comparative phylogenetic analysis produces equivocal results: maximum 175 likelihood modelling of plausible ancestral states, against various combinations of 176 branch length and character transition models (Supplementary text and 177 Supplementary Fig. 9, Table 3), reveals various potential solutions. The statistically 178 most likely result (Fig. 3 and Supplementary Table 3, highest log-likelihood value) 179 shows that the avemetatarsalian ancestors of dinosaurs and pterosaurs possessed 180 integumentary filaments, with highest likelihood of possessing monofilaments; tufts 181 of filaments, and, especially, brush-type filaments, are less likely ancestral states. This 182 confirms that feather-like structures arose in the Early or Middle Triassic. The 183 alternative tree for Dinosauria, with Ornithischia and Theropoda paired as 184 Ornithoscelida³⁰, produces an identical result. 185

We present these modelling data with caution, however, for two reasons: (1) the tree rooting method can influence the result (Supplementary Table 3), favouring results in which either scales are the basal condition or where non-theropod featherlike structures and feathers evolved independently (Supplementary Figure 9, Table 3), and (2) there is no adequate way to model probabilities of evolution of all six feather types, or to model probabilities of transitions between the six different feather types.

192 The discovery of multiple types of feather-like structures in pterosaurs has broad 193 implications for our understanding of pterosaur biology and the functional origin of 194 feather-like structures in Avemetatarsalia^{31,32}. Potential functions of these structures 195 include insulation, tactile sensing, streamlining and coloration (primarily for

camouflage and signalling), as for bristles, down feathers and mammalian hairs³¹⁻³⁴. 196 Type 1, 2 and 4 filaments could shape a filamentous covering around the body and 197 wings (Fig. 4) that might have functioned in streamlining the body surface in order to 198 reduce drag during flight, as for modern bat fur or avian covert feathers^{33,35}. Type 1 199 and 2 filaments occur in considerably high densities, particularly around the neck, 200 201 shoulder, hindlimb and tail regions where the high degree of superposition prevents easy discrimination of adjacent fibres. This, along with the wide distribution and 202 frayed appearance, resembles mammalian underfur adapted for thermal insulation^{36,35}. 203 Despite the less dense packing of Type 4 filaments on the wings, the morphology of 204 the structures is consistent with a thermoregulatory function: down feathers can 205 achieve similar insulation as mammalian hair with only about half the mass, due to 206 their air-trapping properties and high mechanical resilience, effective in retaining an 207 insulating layer of still air³⁸. This may optimize the encumbrance of the large wing 208 area to wing locomotion¹⁸. Type 3 filaments around the jaw (Fig. 4) may have had 209 tactile functions in e.g. prey handling, information gathering during flight, navigating 210 in nest cavities and on the ground at night, similar to bristles in birds³⁹. 211

213 Methods

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Sampling. The specimen NJU-57003 is represented by two fragmented slabs, both 214 containing original bone, fossilized soft tissues, and natural moulds of bones. Each 215 slab was glued together along the fissures by fossil dealers with the fossil on the 216 surfaces untouched. The specimen CAGS-Z070 is represented by a single unbroken 217 slab. Small flakes (1-3 mm wide) of samples with preserved integument and/or 218 enclosing sediments were carefully removed from the inferred integumentary 219 filaments from different parts of NJU–57003 (Supplementary Figs. 1a and 4a–c) 220 using a dissecting scalpel. This method was used to avoid sampling from degraded 221 products of other tissues, such as dermis, epidermis, or even internal organs. Most 222 samples were not treated further; the remainder were sputter-coated with Au to 223 224 enhance SEM resolution (Fig. 2g-h and Supplementary Figs. 4a-f and 6). All 225 experiments described below were repeated in order to validate the results.

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SEM. Samples were examined using a JEOL 8530F Hyperprobe at the School of
Earth Sciences, University of Bristol, and a LEO 1530VP scanning electron
microscope at the Technical Services Centre, Nanjing Institute of Geology and
Palaeontology, Chinese Academy of Sciences. Both instruments were equipped with a
secondary electron (SE) detector, a back-scattered electron (BSE) detector and an
energy dispersive X-ray spectrometer (EDS).

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Measurements of melanosomes. The geometry of melanosomes was measured from
SEM images using the image-processing program ImageJ (available for download at
http://rsbweb.nih.gov/ij/). We measured maximum short and long axis length of
melanosomes that were oriented perpendicular to line of sight, and from these data we
calculated mean and coefficient of variation (CV) of the long and short axis, and mean
aspect ratio (long:short axis). Based on the proposed taphonomic alteration of fossil

- 240 melanosome size (shrinkage up to $\sim 20\%$ in both length and diameter)^{40,41}, we 241 modelled potential diagenetic alteration by enlarging original measurements by 20%. 242
- FTIR microspectroscopy. Samples of the filamentous tissues and the associated 243 sediments were removed separately from NJU-57003 and placed on a BaF₂ plate 244 245 without further treatment. The IR absorbance spectra were collected using a Thermo iN10MX infrared microscope with a cooled MCT detector, at the School of Earth 246 Sciences, University of Bristol. The microscope was operated in transmission mode 247 with a 15x15 micron aperture. 10 spectra were obtained from the filamentous tissues. 248 The spectra show consistent results and the example presented in Fig. 2 shows the 249 highest signal to noise ratio and was obtained with 2 cm⁻¹ resolution and 2000 scans. 250 251
- Fluorescence microscopy. Selected areas with extensive soft tissue preservation in
 NJU–57003 were investigated and photographed using a Zeiss Axio Imager Z2
 microscope with a digital camera (AxioCam HRc) and a fluorescence illuminator
- 255 (514 nm LED) attached, at the Technical Services Centre, Nanjing Institute of
- 256 Geology and Palaeontology, Chinese Academy of Sciences.
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Laser-stimulated fluorescence (LSF) imaging and data reduction protocol. LSF 258 images were collected using the protocol of Kaye et al. ^{15,16}. NJU-57003 was imaged 259 with a 405 nm 500 mw laser that was projected into a vertical line by a Laserline 260 Optics Canada lens. The laser line was swept repeatedly over the specimen during the 261 exposure time for each image in a dark room. Images were captured with a Nikon 262 D610 DSLR camera fitted with an appropriate long pass blocking filter in front of the 263 lens to prevent image saturation by the laser. Standard laser safety protocols were 264 followed during laser usage. The images were post processed in Photoshop CS6 for 265 sharpness, colour balance and saturation. 266

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Phylogenetic macroevolutionary analysis. In order to analyse the evolution of 268 feather characters, data were compiled on known integumentary characters across 269 dinosaurs and pterosaurs. The basic data were taken from the Supplementary data of 270 Barrett et al.², comprising 74 dinosaurs (33 ornithischians, seven sauropods and 44 271 theropods (including four Mesozoic birds)); to this dataset we added four pterosaurs. 272 Barrett et al.² scored taxa for three integumentary states (scales, filaments, feathers) 273 in their macroevolutionary analyses. We checked and followed these basic categories 274 and added three more; we then cross-referenced these six categories against the 275 feather morphotypes defined by Xu et al. ⁴². The categories used herein are: scales (1; 276 not included in Xu et al.⁴²), monofilaments (2; morphotypes 1 and 2 in Xu et al.⁴²), 277 brush-like filaments associated with a planar basal feature (3; morphotypes 4 and 6 in 278 Xu et al. ⁴²), tufts of filaments joined basally (4; morphotype 3 in Xu et al. ⁴²), open 279 pennaceous vane, lacking secondary branching (5; morphotype 5 in Xu et al., ⁴²), and 280 closed pennaceous feathers comprising a rachis-like structure associated with lateral 281 branches (barbs and barbules) (6). There was some uncertainty over feathers coded 282 herein as type 3, which could correspond to morphotype 6, or morphotypes 4 and 6 in 283

Xu et al. ⁴². However, the only taxa coded with these as the most derived feather type are *Sordes pilosus* and *Beipiaosaurus inexpectus*. These taxa belong to separate clades and thus the calculation of ancestral states is not affected by how our feather type 3 is coded (i.e. whether treating morphotypes 4 and 6 of Xu et al. ⁴² in combination or separately).

As in previous studies², we used maximum-likelihood (ML) approaches to 289 explore trait evolution. There are many methods to estimate ancestral states for 290 continuous characters, but choices are more limited for discrete characters, such as 291 here, where only ML estimation of ancestral states is appropriate⁴³. We calculated ML 292 reconstructions of ancestral character states using the 'ace' function of the ape R 293 package⁴⁴, with tree branch lengths estimated in terms of time, derived using the 294 'timePaleoPhy' function in the paleotree package⁴⁵ and the 'DatePhylo' function in 295 the strap R package⁴⁶. These enabled us to assess results according to three methods 296 of estimating branch lengths, the 'basic' method, which makes each internal node in a 297 tree the age of its oldest descendant, the 'equal branch length' (equal) method, which 298 adds a pre-determined branch length (often 1 Myr) to the tree root and then evenly 299 distributes zero-length branches at the base of the tree, and the 'minimum branch 300 301 length' (mbl) method, which minimizes inferred branching times and closely resembles the raw, time-calibrated tree. A problem with the 'basic' branch length 302 303 estimation is that it results in many branch lengths of length zero, in cases where many related taxa are of the same age; in these cases, we added a line of code to make 304 such zero branch lengths equal to 1/1000000 of the total tree length. A criticism of the 305 mbl method is that it tends to extend terminal branching events back in time, 306 especially when internal ghost lineages are extensive², but this is not the case here, 307 and the base of the tree barely extends to the Triassic / Jurassic boundary. 308

We ran our analyses using three evolutionary models with different rates of transition between the specified number of character states (six here), namely "ER", an equal-rates model, "ARD", an all-rates-different model and "SYM", a symmetrical model. These were calculated using the 'ace' function in ape² and the 'add.simmap.legend' function of the R package 'phytools' ⁴⁷.

In a further series of analyses, we attempted to model the macroevolution of all 314 traits, as coded (see Supplementary results), so coding multiple trait values for taxa 315 that preserve multiple feather types. This did not shed much light on patterns of 316 evolution of feather types because the multiple trait codings (e.g. 1,2 or 2,5,6) were 317 each made into a new state, making 14 in all, and these were not linked. Therefore, 318 319 the six multiply coded taxa that each had feather type 6 were represented as six independent states and their evolution tracked in those terms. Further, we attempted to 320 separate the six characters, so they would track through the tree, whether recorded as 321 singles or multiples in different taxa; however, we did not have the information to 322 enable us to do this with confidence because of gaps in coding. In terms of reality, 323 these multiply coded taxa still represent an incomplete sample of the true presence 324 and absence of character states - by chance, many coelurosaurs are not coded for 325 scales (1) or monofilaments (1), and yet it is likely they all had these epidermal 326 appendages. Therefore, attempting to run such multiple codings, with characters 327

328	either as groups or coded	independently, encounters	s so many gaps that the result is
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- hard to interpret. Our approach is to code the most derived feather in each taxon, and
- that too is incomplete because of fossilization gaps, but at least it represents a
- minimal, or conservative, approach to trait coding and hence to the discoveries of
- macroevolutionary patterns of feather evolution; complete fossil data might show
- wider distributions of each feather type and hence deeper hypothesized points of
- origin. Complete coding of feather types would of course allow each trait to be
- tracked in a multiple-traits analysis.
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337 Data availability

The data that support the findings of this study are available from the correspondingauthors upon reasonable request.

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467 Author Contributions

B.Y.J. and M.J.B. designed the research, Z.X.Y., B.Y.J. and X.X. systematically
studied the specimens, Z.X.Y., S.L.K., M.E.M, and P.J.O. did the SEM analysis,
Z.X.Y. and B.Y.J. did the FTIR analysis, M.P. and T.G.K. did the LSF imaging, data
reduction and interpretation, M.J.B. did the maximum likelihood analyses, and
Z.X.Y., B.Y.J., M.J.B., M.E.M, X.X. and P.J.O. wrote the paper; all authors approved
the final draft of the paper.

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- 480

Figure 1 | Integumentary filamentous structures in CAGS–Z070. a, Overview
shows extensive preservation of soft tissues. b–p, Details of the integumentary
filaments in the regions indicated in a on the head and neck (b–d, i–j), forelimb (f–g),
wing (l–m) and tail (o–p), and illustrated reconstructions of the filaments (e: Type 1
filament; h: Type 2 filament; k: Type 3 filament; n: Type 4 filament). Scale bars: 20
mm in a; 10 mm in b; 500 µm in c and i; 100 µm in d; 1 mm in f, l, m and p; 200 µm
in g and j; 5 mm in o.

488

489 Figure 2 | Preservation, microstructure and chemistry of the integumentary

490 **filamentous structures in NJU–57003. a**, Laser-stimulated fluorescence^{6,15,16} image 491 highlights extensive preservation of soft tissues (black areas). **b–f**, Details of the

- 492 integumentary filaments in the regions indicated in A on the head and neck $(\mathbf{b}-\mathbf{c})$,
- 493 wing $(\mathbf{d}-\mathbf{e})$ and tail (f). g-h, Scanning electron micrographs of the monofilaments on
- the neck and hindlimb of NJU–57003 (samples 10 and 39, respectively,
- 495 Supplementary Fig. 1a) show densely packed, elongate and oblate melanosomes. i,
- 496 FTIR absorbance spectra of the monofilaments, monofilaments with sediment matrix,
- 497 and sediment matrix in NJU–57003 (Sample 15, Supplementary Fig. 1a) compared
- with spectra from a feather of *Anchiornis* (from ref. ²⁰), extant Marabou stork feather
 (from ref. ²¹) and black and red human hair melanosomes (from ref. ²⁴). Scale bars: 20
- 500 mm in \mathbf{a} ; 1 mm in \mathbf{b} , \mathbf{c} and \mathbf{e} ; 5 mm in \mathbf{d} and \mathbf{f} ; 1 μ m in \mathbf{g} and \mathbf{h} .
- 501

502 Figure 3 | Phylogenetic comparative analysis of integumentary filament and

503 feather evolution in pterosaurs and archosaurs. The phylogeny is scaled to

- geological time, with recorded terminal character states for each species, and 504 estimated ancestral character states at the lower nodes. The model is the most likely of 505 the maximum likelihood models, based on minimum-branch lengths (mbl) and 506 transitions occurring as all-rates-different (ARD), but other results with lower 507 likelihoods show scales as ancestral. The ancestral state reconstruction shows a 508 combination of monofilaments, tuft-like filaments, and brush-type filaments as the 509 ancestral state for Avemetatarsalia and for Dinosauria. The estimated ancestral state 510 for Theropoda comprises all five feather states. Numbered small vertical arrows 511 indicate earliest occurrences of feather types 2-6. Two hypotheses for timing of avian 512 feather origins are indicated: A, early origin, at the base of Avemetatarsalia in the 513 Early Triassic, or B, late origin, at the base of Maniraptora in the Early-Middle 514 Jurassic. 515 516
- 517 Figure 4 | Reconstruction of one of the studied anurognathid pterosaurs, exhibiting
- 518 diverse types of pycnofibres distributed in different body parts.





