Digital video platforms and phylogenetic modelling reveal parrot tool use is not rare

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Abstract

Putatively rare behaviours, such as tool use, are challenging to study because absence of evidence can arise either from a species' inability to produce the behaviour or from insufficient research effort. Here, we tackle this challenge by combining data from digital platforms and phylogenetic modelling to estimate actual rates of tool use in parrots.

Videos on a social media platform revealed novel instances of tool use in 17 parrot species, more than doubling the confirmed number of tool-using parrot species from 11 (3%) to 28 (7%). Phylogenetic modelling ranked additional species that are most likely to be unobserved tool users, suggesting that between 11% and 17% of extant parrot species may be tool users. These discoveries have implications for inferences about the evolutionary drivers and origins of tool use in parrots, revealing associations with relative brain size and feeding generalism and indicating several genera where tool use was likely an ancestral trait. Overall, our findings challenge the assumption that current sampling effort captures the full distribution of putatively rare animal behaviours. Combining our sampling and analysis methods offers a fruitful approach for investigating the distribution, drivers, and origins of other rare behaviours.

Keywords: tool use; technical intelligence; parrots; digital video platforms; phylogenetic modelling

Introduction

Our understanding of the evolution of animal behaviour is built on the assumption
that we have access to sufficient data¹⁻³. However, this is not always the case. Data on
behaviours that are rare, fleeting, or otherwise difficult to observe are likely to be patchy
and incomplete^{4,5}. Among species for which such behaviours have not been observed, it can
be difficult to differentiate between cases in which the species is truly incapable of
producing the behaviour and cases in which the species is capable of producing the
behaviour but the behaviour has not yet been observed. Such a distinction can be critical
for drawing conclusions about the rarity and evolution of the behaviour in question.

Comparative work on the evolution of tool use is a paradigmatic example of this 10 issue. The initial discoveries of tool use in chimpanzees⁶, birds⁷, dolphins⁸, and octopuses⁹ 11 occurred decades after significant advances on other more easily measurable aspects of 12 their biology. Since then, scholars have proposed a clear operational definition of tool use 13 applicable to all species — the manipulation of an unattached object as an extension of the animal's body to achieve a goal¹⁰ — and have used the distribution of species meeting this 15 definition to make various claims about the evolutionary drivers of tool use behaviours. For 16 example, based on the observation that bird species with reported tool use tend to have 17 larger brains, researchers have identified higher relative brain size as a likely precondition 18 for tool using capabilities^{11–14}. These researchers argue that larger brains are better able to 19 integrate visual and somatosensory information when innovating novel behaviours, such as tool use, in changing environments^{15,16} (but see ref¹⁷). Similarly, researchers have used 21 existing reports of tool use in birds to debate the roles of generalist versus specialist feeding strategies in driving the evolution of tool use, with some arguing that feeding generalists require technical innovations to expand their dietary niche^{15,18,19} and others arguing that feeding specialists require technical innovations for extractive foraging of specific foods^{20,21}.

However, before we can make claims of this kind, we need to know whether current

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research effort in the literature is sufficient for robust conclusions to be drawn about the evolution of tool use. In fact, evidence suggests that research effort is often systematically 28 biased towards particular taxonomic groups, parts of the world that are easy to access, and 29 species with life history traits that make them easier to study, such as larger distribution 30 ranges and population sizes²². This is a crucial limitation because insufficient observation 31 may lead researchers to miss true instances of tool behaviours and thus draw premature 32 conclusions about the evolutionary drivers and origins of tool use. Researchers are keenly 33 aware of this problem and have attempted to deal with it in different ways, such as controlling for the number of scientific papers published on different species. But previous 35 work has not yet attempted to quantify and explicitly model the relationship between actual tool-using behaviour and what is reported in the scientific literature. If more 37 tool-using species exist than previously thought, this could have important implications for theories of the evolutionary drivers and origins of tool use and for our understanding of how rare this behaviour actually is.

One potentially powerful method for quantifying actual rates of rare animal
behaviours is by collating evidence of the behaviours from digital video platforms²³. With
millions of videos posted every day, digital platforms like YouTube offer a valuable source
of data on animal behaviour. Digital video platforms have already been successfully used
to uncover a variety of rare animal behaviours, including interspecies play in dogs²³, novel
problem-solving behaviours in horses²⁴, and death-related behaviours in Asian elephants²⁵.

By casting the net wider than the scientific literature, digital video platforms can provide
an indication of the tool-using species that the literature might be missing.

Even after collating evidence from digital video platforms, some tool-users could *still*remain unobserved. One principled framework for identifying these unobserved species is to
specify a causal model of the process that generates the observed data. We propose one
such causal model in Figure 1. In this model, we assume that the presence of tool use in
the scientific literature (or on digital video platforms) is caused by both unobserved tool

use capabilities and the number of published studies (or the number of videos) for any given species. Tool users are more likely to be observed if they are well studied, but understudied tool users may go undetected. Furthermore, based on existing theories of the evolution of tool use^{12–16,18–21}, we propose that the unobserved tool use capabilities are additionally caused by relative brain size, feeding strategy, and shared phylogenetic ancestry. Expressing this causal model as a statistical model can suggest further species which are likely to be unobserved tool-users and, simultaneously, test existing theories of the evolutionary drivers of tool use without incorrectly assuming that absence of evidence is evidence of absence.

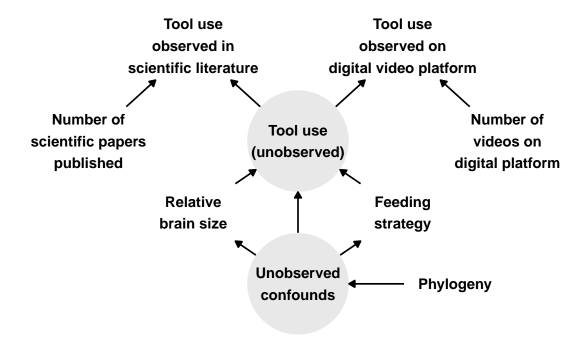


Figure 1. Causal model of observed tool use. Directed acyclic graph of the causal relationships between observed tool use and other variables. Available scientific data on tool use is caused both by unobserved tool use presence and scientific research effort (i.e., number of publications). Available video data on tool use is caused both by unobserved tool use presence and video research effort (i.e., number of videos). According to theory, unobserved tool use presence should be caused by relative brain size (encephalisation quotient) and feeding strategy (generalist vs. specialist). These variables all share unobserved confounds generated by shared phylogenetic history. Grey circles indicate unobserved variables.

Here, we apply these approaches to tool use in the parrot order. We focus on tool use

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in parrots for a number of reasons. First, the scientific literature suggests that only a small proportion of extant parrot species (11 out of 398; 3%) use tools ^{12,26–38}. Parrot tool use thus provides an ideal test case for examining how robust sampling is in the scientific literature. Second, parrots are highly popular as pets. Over 70% of all extant parrot species are bred in the aviculture industry and kept as pets worldwide ^{39–45}, enabling us to leverage the power of digital video platforms to search for evidence of tool use ²³. Third, detailed data on relative brain sizes ^{46–51}, feeding strategies ⁵², and shared ancestry ⁵³ exist for parrots, allowing us to fit the statistical model implied by Figure 1 to the entire parrot order.

We first present the results from our video survey, in which we collated videos of tool
use in parrots from a digital video platform. This survey reveals a number of previously
unidentified tool-using parrot species, which we map onto the phylogeny of the parrot
order. We then describe our statistical model in more detail, and use it to (i) rank further
parrot species that are likely unobserved tool users and (ii) re-examine key hypotheses
regarding the evolutionary drivers and origins of tool use in parrots.

78 Results

9 Digital video platform reveals tool use in additional parrot species

We surveyed the digital media platform YouTube for video evidence of tool use in parrots (see Methods for detailed search criteria). In our search, we used the standard criteria for identifying tool use in the literature, defining "true" tool use behaviour as the manipulation of an unattached object as an extension of the animal's body to achieve a goal¹⁰, while "borderline" tool use involved the use of an object that was still attached to a substrate⁵⁴.

In total, we found 116 videos of 104 individuals from 25 parrot species performing
behaviours that met the definition of either true tool use (100 videos of 89 individuals from
22 species) or borderline tool use (16 videos of 16 individuals from 7 species). All videos

featured pet parrots in captive settings. In 68 of these videos, owners did not appear to
interact with the subjects. In 43 videos, there was potential human interaction, either from
the owners being in close physical contact with the bird (e.g., bird perching on hand),
talking to the bird, or handing it the tool (which occurred in only two videos). We could
not establish the degree of human interaction in the remaining 5 videos, as sound had been
removed or was substituted by music. None of the videos featured owners directly
rewarding tool use behaviours with food. All borderline tool use cases were excluded from
further analyses.

Of the 22 parrot species performing true tool use, 13 were represented in our video survey by two or more individuals over multiple independent observations. True tool use always involved the subject using an object for self-scratching (95 videos involved scratching the head and/or neck). The most common tool (53 videos) was a moulted feather. Human-made objects (e.g., pens, spoons, pieces of wood, cardboard) were also common. In 66 videos, parrots manipulated the tool while keeping their body still, rather than holding the tool still and moving towards it.

According to YouTube video descriptions and owner comments, 45 of the individuals 104 performing true tool use were males and 18 were females. No sex information was provided 105 for the remaining 26 individuals. As owners provided no information on whether sex had 106 been established through genetic testing, and sexual dimorphism in parrots is rare^{55,56}, we 107 could not typically ascertain if descriptions were accurate. It is unclear if the 108 disproportionately large number of males in the sample is a consequence of owners more 109 readily assuming their parrots are male when they have not been genetically tested, owners being more likely to own or film male parrots, or male parrots exhibiting more true tool use 111 behaviours than female parrots. 112

Figure 2 maps the findings from the video survey onto a maximum clade credibility phylogeny for the parrot order, plotted alongside species previously identified in the

scientific literature. Before the video survey, 11 parrot species (3%) had been identified as 115 tool users in the scientific literature. Across our video survey, we observed true tool use in 116 22 species, 5 of which overlapped with the scientific literature and 17 of which were novel 117 species. All of the species identified in the video survey were cockatoos (Cacatuidae), Old 118 World parrots (*Psittacinae*), or neotropical parrots (*Arinae*). The most common species in 119 our survey, accounting for 41 videos from 37 individuals, was the green-cheeked conure 120 (Pyrrhura molinae). In accordance with the scientific literature, the video survey found no 121 evidence of tool use in any species of Psittaculidae, despite this family containing some of 122 the most commonly kept pet species, including lovebirds, lorikeets, and Asian parakeet 123 species. Combining both the video survey and the scientific literature, we can thus identify 124 28 tool-using parrot species overall (7%), compared to the 11 previously reported. 125

The identification of new tool-using species in our video survey increases the extent 126 to which phylogeny can explain the distribution of tool use in the parrot order. We 127 estimated phylogenetic signal (Pagel's λ) of tool use using both the pre-video-survey and 128 post-video-survey data. Pagel's λ varies between 0 and 1, where 0 implies that the 129 distribution of a trait across species is unexplained by phylogenetic relatedness and 1 130 implies that the distribution of a trait across species is fully explained by phylogeny. Using 131 the evidence of tool use from the scientific literature alone (pre-video-survey data; 11 132 tool-using species), we estimated an average posterior Pagel's λ of 0.60 (95% credible 133 interval [0.00 0.90]; total n = 174 species). This estimate was moderate-to-strong, but 134 highly uncertain. In comparison, combining the evidence from both the literature and the 135 video survey (post-video-survey data; 28 tool-using species) resulted in a stronger and more 136 certain estimate of phylogenetic signal. With these data, we estimated Pagel's $\lambda = 0.65$ 137 $(95\% \text{ CI } [0.50 \ 0.77]; \text{ total } n = 174 \text{ species})$. Thus, the results of our video survey increase 138 the extent to which the distribution of tool use across parrot species can be explained by 139 shared phylogenetic ancestry. This suggests that we can potentially use phylogenetic information, along with other variables, to identify further tool-using parrot species that

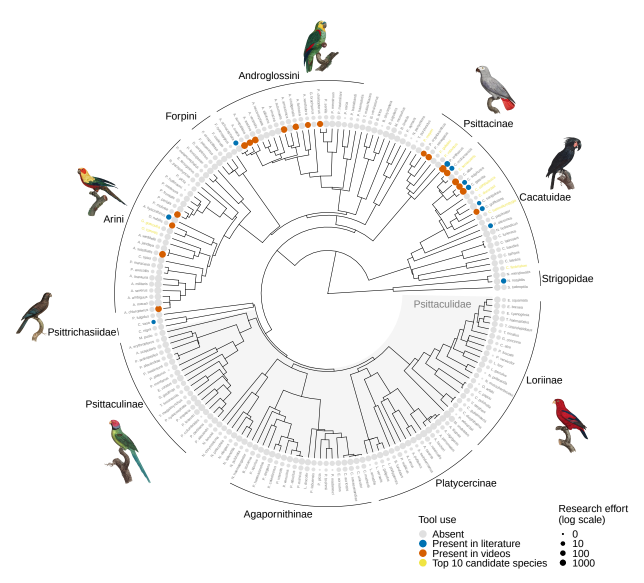


Figure 2. Results of video survey and phylogenetic survival cure modelling mapped onto a maximum clade credibility phylogeny of the parrot order. Orange points in the inner ring indicate species observed in the video survey, with point size scaled by the number of videos for each species (note that three species observed only in the video survey are not present in the phylogeny due to a lack of genomic data: Psittacara erythrogenys, Psittacus timneh, and Aratinga nenday). Blue points in the outer ring indicate species observed in the scientific literature, with point size scaled by the number of papers published on each species. Yellow species names indicate the top ten most likely tool-using species from our phylogenetic survival cure model which were not observed in the scientific literature or the video survey. Total n = 174 species.

may remain undetected.

Phylogenetic survival cure modelling identifies further candidate tool users

In addition to the 28 tool-using species identified in the literature and our video survey, we fitted a Bayesian phylogenetic survival cure model to rank further species that are likely to be undetected tool-users (i.e., tool-using species with no tool use reported in the literature or in videos).

Survival cure models⁵⁷, also known as split population models⁵⁸, are used to analyse 148 the time to some event of interest with the added assumption that a certain proportion of 149 the population will never experience the event, no matter how long they are measured for. 150 These models have been used to analyse a variety of right-censored outcomes, from cancer 151 relapse⁵⁷ to criminal recidivism⁵⁸. The data are right-censored because some individuals 152 will have experienced the event when they are measured (e.g., disease onset, return to 153 prison) while others will have not experienced the event. For those who have not 154 experienced the event, this may be because (i) the event has not happened to them yet or 155 (ii) the event will never happen to them. Survival cure models treat these two processes 156 separately. 157

Our tool use problem has the same features. We are modelling a time-to-event;
specifically, the amount of "time" (i.e., observation opportunities measured as the number
of published papers or videos) until tool use is identified. This is right-censored data
because many species will not have had tool use identified when we measure them.
Moreover, we can assume that a certain proportion of the population will never experience
the event – that is, they are a non-tool-user and so we will never identify tool use no
matter how long we measure them for.

In our model, we infer the tool-using status of each species by allowing each species to have their own probability of being a non-tool-user. Following our causal model (Figure

167 1), we predict these probabilities based on feeding strategy, encephalisation quotient, and
168 phylogenetic history (see Methods for full model). The model additionally takes research
169 effort into account by allowing that, among species for which tool use is unobserved, all else
170 being equal those with fewer published papers and fewer video search hits have a higher
171 probability of being undetected tool users (Figure S1).

We found that this phylogenetic survival cure model was able to adequately 172 distinguish between species with and without evidence for tool use, with an 173 area-under-the-curve classification statistic of 0.95 (Figure S2). To further estimate the 174 accuracy of the model's predictions, we also used a leave-one-species-out approach with known tool users. For each of the 25 tool-using species that were represented on the phylogeny and for which we had brain size and genomic data (we lacked data for three 177 tool-using species), we fitted the model to a modified dataset which set tool use to be 178 absent for the target species in both the scientific literature and the video survey. Across 179 25 cross-validation models, 18 models (72%) continued to predict the target species as 180 having a median posterior probability of tool use that was within the range of all other tool 181 users. This classification rate was greater than the baseline classification rate of 26% for 182 species without evidence of tool use in the full model (38 of 149 species without evidence of 183 tool use had a median posterior probability of tool use that was within the range of the 184 tool-using species). Together, the area-under-the-curve statistic and the 185 leave-one-species-out approach suggest that the model is able to adequately classify known 186 tool users, with some error. 187

Figure 3 visualises the ranked posterior probabilities of tool use from the phylogenetic survival cure model for all parrot species. As expected, the known tool users are ranked towards the top of this list. However, several "tool use absent" species also rank highly on the list, despite not being identified as tool users in the scientific literature or in our video survey. In fact, according to the model, the most likely tool user is a species for which tool use is unobserved in our data: the blue-eyed cockatoo (*Cacatua ophthalmica*). This species

is endemic to Papua New Guinea and is relatively understudied, with only 6 published 194 papers and 596 video search hits, which is fewer than the model expects are necessary to 195 discover tool use when it is present (Figure 4). This species is also found in the Cacatua 196 genus, a clade containing several known tool users. This prediction makes sense given the 197 high phylogenetic signal for tool use reported by the model (Figures S3 and S4). Beyond 198 the blue-eyed cockatoo, other highly ranked species without observed evidence of tool use 199 are the Meyer's parrot (Poicephalus meyeri), the golden parakeet (Guaruba quarouba), the 200 long-billed corella (Cacatua tenuirostris), the Solomons cockatoo (Cacatua ducorpsii), the 201 red-fronted parrot (*Poicephalus gulielmi*), the Cape parrot (*Poicephalus robustus*), the 202 yellow-eared parrot (Ognorhynchus icterotis), the red-vented cockatoo (Cacatua 203 haematuropygia), and the gang-gang cockatoo (Callocephalon fimbriatum). Figure 2 plots 204 these species on the parrot phylogeny, using the top ten highest ranked species without observed evidence of tool use as an arbitrary cutoff for visualisation purposes.

The posterior probabilities shown in Figure 3 are estimated with uncertainty, so it is difficult to "identify" any particular species as an undetected tool user. Nevertheless, taking the sum of all the posterior probabilities for the 149 species without observed evidence of tool use, we can estimate that around 26 of those species are likely to be undetected tool users (median sum of probabilities = 25.68, 95% CI [15.15 41.33]). When combined with the species known to use tools, this implies that between 11% and 17% of extant parrot species may be tool users.

214 Implications for the evolutionary drivers and origins of tool use

The predicted probabilities from our phylogenetic survival cure model have implications for inferences about the evolutionary drivers and origins of tool use in the parrot order. Regarding the drivers of tool use hypothesised in Figure 1, the phylogenetic survival cure model revealed that encephalisation quotient strongly positively predicted the probability of tool use (median posterior log odds slope = 1.12, 95% CI [0.39 2.00]; total n

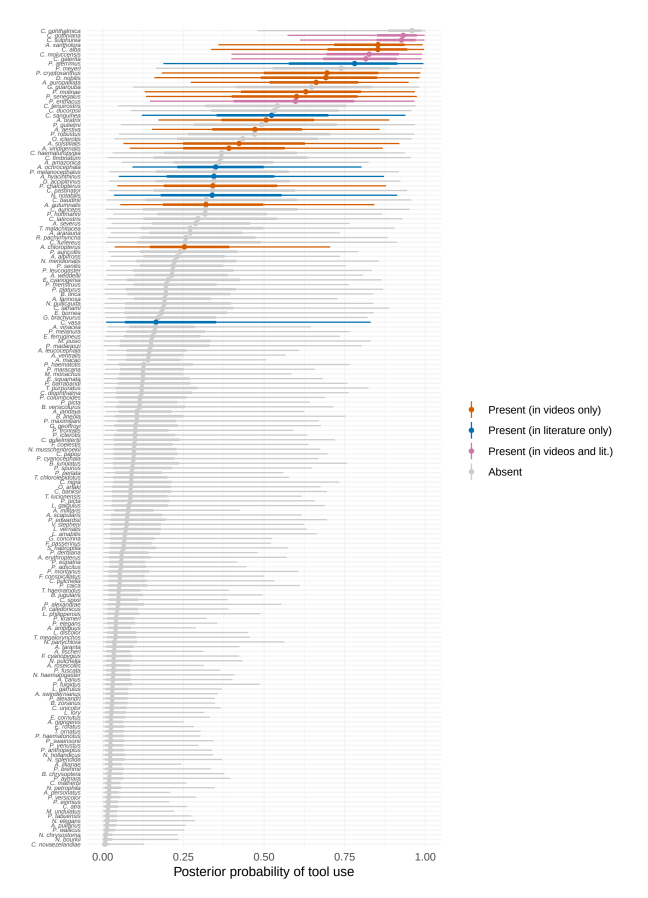


Figure 3. Posterior predicted probabilities of tool use for each species from our phylogenetic survival cure model. Points are posterior medians and lines are 50% and 95% credible intervals. Total n=174 species.

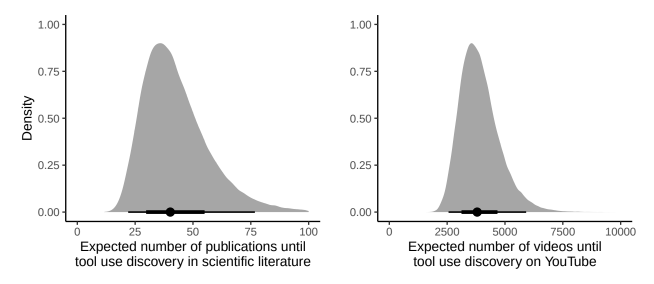


Figure 4. Expected number of published papers and videos until tool use discovery, according to the survival component of the phylogenetic survival cure model. Densities are full posterior distributions, points are posterior medians, and lines are 50% and 95% credible intervals.

= 174 species; Figure 5). This helps explain the ranking in Figure 3: the blue-eyed 220 cockatoo has the largest relative brain size in the dataset. We also found that feeding 221 generalist species were slightly more likely to be tool users, though the posterior difference 222 between generalists and specialists was quite uncertain (median posterior log odds 223 difference = 0.33, 95\% CI [-1.13 1.76]; total n = 174 species). These results from the 224 survival cure model differed from the results of models fitted to pre-video-survey and 225 post-video-survey data without the survival cure component, which found no effect of 226 relative brain size and no difference between feeding strategies, respectively (Figure S5). 227

Regarding the origins of tool use, we fitted exploratory ancestral state reconstruction 228 models to the pre-video-survey data, the post-video-survey data, and the predicted probabilities from the phylogenetic survival cure model. The discoveries from our video 230 survey and from our phylogenetic modelling increased the likelihood that the capacity for tool use was present in the most recent common ancestors for several parrot genera. These 232 include the most recent common ancestor of amazon parrots native to the Americas (Amazona), the most recent common ancestor of the true white cockatoos and corellas found in South East Asia and Australasia (Cacatua), the most recent common ancestor of

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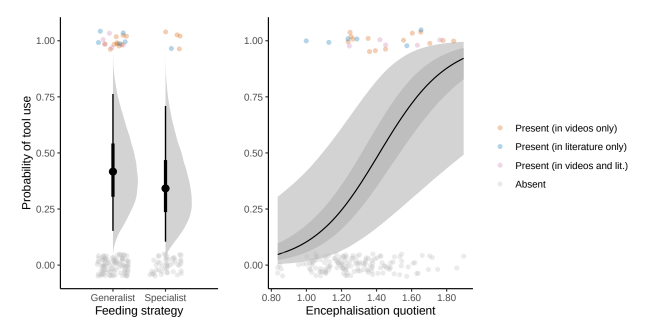


Figure 5. Posterior predictions for the effects of feeding strategy and encephalisation quotient on the probability of tool use from the phylogenetic survival cure model. In the left plot, points and lines represent posterior medians and 50% and 95% credible intervals, with densities representing full posterior distributions. In the right plot, the line and shaded areas represent the posterior median regression line with 50% and 95% credible intervals. In both plots, individual species are coloured according to the presence / absence of tool use in the video survey and the scientific literature. Total n=174 species, generalist n=121 species, specialist n=53 species.

the kea and the kākā from New Zealand (*Nestor*), and the most recent common ancestor for the *Poicephalus* genus native to Africa (Table 1; Figures S6 - S8).

238 Discussion

Since the earliest anecdotes of parrots using tools by Wallace in the 1880s⁵⁹ and more systematic anecdotal reports in the 1970s⁶⁰, only 11 parrot species (3% of all extant parrots) have been documented as tool users in the scientific literature. Our study collated data from a digital video platform to identify 17 additional tool-using parrot species that are new to science, more than doubling the overall count to 28 species (7%). These species consisted of cockatoos (*Cacatuidae*), Old World parrots (*Psittacinae*), and neotropical parrots (*Arinae*). Beyond the video survey, the strong phylogenetic signal in our dataset allowed us to use phylogenetic information, along with other variables, to infer the

Table 1 Estimated probabilities of tool use for most recent common ancestors of several parrot genera. Probabilities estimated using exploratory ancestral state reconstruction models fitted to the pre-video-survey data, post-video-survey data, and predicted probabilities from the phylogenetic survival cure model. Total n = 174 species.

Genus	Pre-video-survey	Post-video-survey	Survival cure probabilities
Amazona	0.06, 95% CI [0.01 0.17]	0.21, 95% CI [0.04 0.66]	0.48, 95% CI [0.05 0.84]
Cacatua	0.08, 95% CI [0.03 0.20]	0.17, 95% CI [0.03 0.45]	0.81, 95% CI [0.23 0.99]
Nestor	0.11, 95% CI [0.05 0.29]	0.21, 95% CI [0.12 0.37]	0.53, 95% CI [0.29 0.66]
Poice phalus	0.06, 95% CI [0.02 0.11]	$0.19, 95\% \text{ CI } [0.12 \ 0.41]$	$0.72, 95\% \text{ CI } [0.16 \ 0.90]$

unobserved probabilities of tool use across the parrot order. Our phylogenetic survival cure
model incorporated information on phylogenetic history, research effort, relative brain size,
and feeding specialisation to rank parrot species that were most likely to be undetected
tool users. The sum of probabilities from this model implied that between 15 and 41 of the
species without observed evidence of tool use are likely to be undetected tool users,
suggesting that the true proportion of tool users may be as high as 17%.

These findings have a number of important implications. First, our findings show 253 that current research effort in the scientific literature is insufficient to capture the real 254 world occurrence of parrot tool use. If the scientific literature had sampled the natural 255 world sufficiently, we would expect to see close correspondence between those species 256 reported as tool users in the literature and those species the public have filmed performing 257 tool use. Instead, we discovered a large discrepancy between these two data sources, both 258 in the prevalence of tool use and the species identified, likely due to the numerous 259 difficulties inherent to conducting observations in the wild. This raises the possibility that 260 other rare behaviours remain yet to be discovered in the scientific literature. 261

Second, in terms of the evolution of tool use in parrots, our study challenges a key assumption made in the literature to date: that only a minority of parrots are tool

users^{30,31,34,37,38}. The paucity of evidence for tool use across parrots in the literature 264 initially implied that the capacity for tool use may have evolved independently in different 265 parrot species. Our discovery of the widespread distribution of tool use across the parrot 266 phylogeny, along with the strong phylogenetic signal in this expanded dataset, challenges 267 this and suggests that, at least for some parrot clades, the capacity for tool use might be a 268 homologous trait that has been evolutionarily conserved. Our exploratory ancestral state 260 reconstruction analysis provides preliminary support for this hypothesis, revealing an 270 increased probability of the capacity for tool use among the most recent common ancestors 271 for the Amazona, Cacatua, Nestor, and Poicephalus genera. The complexity of tool use 272 behaviour that such a basic capacity might afford, and the conditions under which it 273 emerges in the wild, are yet to be established. However, even at this preliminary stage, our 274 analysis raises an alternative hypothesis for the observed tool use in Cacatua^{30,38} and $Nestor^{26-29,31,34}$, namely that tool behaviours have arisen due to the common ancestor 276 having the capacity to use tools, rather than from independent evolution or behavioural innovation within species.

Third, our results support existing theories of the drivers of tool use. We found that 279 encephalisation was strongly positively related to the probability of tool use in our 280 phylogenetic model, supporting previous theories linking relative brain size to increased 281 tool innovation in birds¹²⁻¹⁴ and primates⁶¹. We acknowledge that encephalisation quotient 282 is not a perfect measure due to measurement error and challenges with interpretation^{3,4}. 283 However, encephalisation quotient has much greater coverage across the parrot phylogeny 284 than more fine-grained measures like whole neuron count⁶² and there is no reason to think 285 that measurement error would produce the consistent patterns across our study and prior 286 work. To understand the causal mechanisms responsible for these relationships, we 287 encourage further work on the specific neural correlates of technical intelligence in parrots, 288 e.g., 63. In our phylogenetic model, we also found that tool use was somewhat more likely 289 among feeding generalists compared to feeding specialists, although this difference was

uncertain. This trend supports previous suggestions that increased cognitive abilities and technical innovation rates are required to expand a generalist species' dietary niche^{18,20,64}. However, the trend contradicts theories linking tool use to dietary specialisation, whereby species eating specific foods that require extractive foraging have higher cognitive ability and are especially prone to using tools²⁰.

All instances of tool use in our video survey met the established criteria for tool use 296 in the literature: the manipulation of an unattached object as an extension of the animal's 297 body to achieve a goal¹⁰. However, a striking feature of the dataset is that the 17 additional species identified in the video survey were exclusively observed using tools for 299 self-scratching. While parrots in the wild use tools to achieve a variety of goals, including extractive foraging and preening^{34,38}, parrots in captivity only require tools to achieve the 301 latter goal. It could be argued that self-scratching is a more cognitively simplistic form of 302 tool use than extractive foraging as it is egocentric, meaning that the tool is directed 303 towards oneself rather than an external object. Nevertheless, self-scratching meets the 304 definition for a more complex type of embodied tool use known as 'tooling', the deliberate 305 generation of a mechanical interface by using an object to manipulate another target or 306 surface⁶⁵. In our dataset, most self-scratching tooling instances involved manipulating the 307 tool while keeping the body still, suggesting that these were goal-directed, deliberate 308 movements by individuals. This makes it difficult to dismiss self-scratching as 309 unimportant: to do so would require a redefinition of tool-use and tooling in non-human 310 animals. Moreover, the fact that we found strong phylogenetic clustering of self-scratching 311 tool use from the video survey with other examples of tool use from the literature supports 312 a common underlying cognitive mechanism. In line with this, some of the species of parrots 313 that use tools for self-scratching in captivity also use tools for other purposes in the 314 wild 30,31,34,38 . 315

A potential concern with data originating from digital video platforms is its reliability^{23,66-70}. In particular, the self-care tool use instances detected in our video survey

might be attributable to training or unintentional cueing by the birds' owners. However, 318 there was little evidence to suggest that the observations of self-scratching tool use in the 319 video survey were merely unintentional accidents or explicitly trained behaviours. 320 Individual parrots often used tools slowly and repetitively over long periods of time, even 321 across multiple different videos, suggesting that their behaviour was not random or 322 accidental⁶⁹. Parrots preferentially employed self-scratching tools on areas of their body 323 that were otherwise inaccessible, with 96\% of all instances involving scratching of the head 324 or neck, suggesting intentional tool use. For 60% of the species in the video survey, we 325 found two or more videos of repetitive and sustained scratching by different individuals of 326 the same species in separate households, suggesting that the manipulations were 327 intentional and recurring events that did not represent unusual stereotypies of any single 328 individual. Regarding the possibility of training or cueing from owners, over half of the videos contained no evidence of human interaction aside from filming the behaviour. 330 Humans only handed parrots their tools in two of the videos, and none of the videos featured owners directly rewarding tool use behaviours with food. It is unlikely that owners 332 would specifically train parrots to scratch themselves with tools. Finally, the high levels of 333 phylogenetic signal in our data provide strong evidence that the observations from our 334 video survey reflect biologically-endowed capacities for tool use rather than accidental or 335 trained behaviours, which would likely appear uniformly across the phylogeny. 336 Nevertheless, future work using data such as this should choose target behaviours carefully 337 and examine videos for evidence of training or cueing. 338

Our data reveals that a wide range of parrot species exhibit self-scratching tool use
behaviours in captive environments. It is possible that these self-care behaviours emerge
more often in captivity because pet parrots are kept with few or no conspecifics, and
therefore must innovate tool use to preen inaccessible body areas, such as their heads and
necks, which would have been otherwise preened by flock members. This may indicate that
human interaction and allopreening in the absence of a conspecific is insufficient for pet

parrots, as suggested by studies on welfare and chronic stress in lone captive parrots^{71–74}. 345 Beyond social isolation in captivity, other factors that reduce preening efficacy by an 346 individual might also lead to the innovation of self-care tooling: for example, a captive kea 347 housed with conspecifics in a naturalistic environment was shown to have innovated the use 348 of a pebble tool for self-care likely as a consequence of a missing upper mandible³¹. These 349 behaviours might therefore be innovated by individuals when ecologically necessary. 350 Critically, however, a cognitive capacity to innovate a behaviour such as tool use must exist 351 in a species in order to manifest in any environment, whether that be captive or in the 352 wild. As such, despite differences in individuals' life experiences in captive and wild 353 settings, such behaviour emerging in a captive setting at the very least indicates an 354 underlying cognitive capacity to innovate that same behaviour in the wild. 355

While missing data imputation is becoming more common in phylogenetic analyses⁷⁵. 356 the important distinction between absence of evidence and evidence of absence has not 357 been given as much attention. Our phylogenetic analysis provides one approach to this 358 problem by distinguishing between true absences of tool use and absences of tool use due 359 to a lack of research effort in the scientific literature or in videos. To achieve this, we 360 explicitly modelled the measurement of the outcome variable along a research effort time 361 series, such that species with lower research effort in the literature or in videos were likely 362 to be censored. In line with our causal model, we also included relative brain size, feeding 363 strategy, and phylogenetic history as predictors of unobserved tool use. We encourage 364 researchers to test this model by directing future study efforts towards the parrot species 365 with the highest probabilities of being undetected tool-users. Future work should also 366 refine the causal model in Figure 1 to provide more certain estimates of tool use 367 probabilities, either by including additional predictor variables or modelling further causes 368 of measurement error in the taxonomic record, such as species abundance and geographic accessibility 22 . 370

In conclusion, we have shown that the scientific literature has insufficiently captured

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the full distribution of tool use in the parrot order. Our digital video survey more than 372 doubled the number of known tool using parrot species from 11 to 28, and our phylogenetic 373 model suggested that the true proportion of parrot tool users could be as high as 17% of all 374 species in this order. These discoveries have implications for theories of the evolutionary 375 drivers and origins of tool use in parrots. Beyond parrot tool use, the methods used in this 376 study have the potential to be applied to other rarely observed behaviours, including tool 377 use in other taxa⁷⁶, rhythmic entrainment in birds^{77–80}, teasing behaviours in primates⁸¹, 378 and tactical deception across all animals^{82–84}. We hope that these methods will continue to 379 uncover a diverse array of ephemeral behaviours that have as yet gone undetected in the 380 scientific literature.

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Author Contributions

All authors contributed to the conceptualisation of the paper. A.P.M.B., X.J.N., and A.H.T. developed the video search methodology. S.C., D.W., and Q.A.D. developed the statistical models and analysed the data. All the authors wrote the manuscript and approved the final version for submission.

Declaration of Interests

The authors declare no competing interests.

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STAR Methods

Resource Availability

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Lead contact. Further information about resources should be directed to Scott

Claessens (scott.claessens@gmail.com).

Materials availability. This study did not generate new unique materials.

Data and code availability. All data have been deposited on GitHub and are publicly available as of the date of publication. DOIs are listed in the key resources table.

All original code has been deposited on GitHub and is publicly available as of the date of publication. DOIs are listed in the key resources table.

91 Method Details

Video searches and coding. Our video search was conducted on YouTube in 392 July 2020. Searches were conducted manually by the first author over a month-long period, 393 using the same IP address and not logged in with a YouTube user account. Search terms 394 included "parrot using tool" and variants (e.g., "macaw using tool", "lorikeet using tool", 395 "parakeet using tool"), "tool use in parrot", "parrot tool use", "parrot scratching itself" 396 (included after we found several videos demonstrating self-care tool use in previous 397 searches) and equivalent terms (e.g., "parrot preening itself", "parrot grooming itself", 398 "parrot scratching"). For all species that did not display results including object 390 manipulation or scratching behaviours, we also searched the species' common name(s) + 400 "tool use", as well as the species' common name(s) + "scratching". We also searched for 401 translations of the terms "parrot tool use" and "parrot scratching" in languages for all countries where bird ownership was reported as $>5\%^{85}$, namely, Turkish, Czech, Polish, French, Italian, Dutch, German, Russian, Spanish, Portuguese, and Mandarin. Browser search histories were not cleared between searches.

When we found a relevant video, we also searched for similar content uploaded by the
same person/channel. For each YouTube search conducted, we watched all relevant videos
until we reached five consecutive videos that did not feature any parrots. At this point, we
ended that search and initiated the next search. In line with previous recommendations²³,
we planned to exclude any videos that consisted of four or more shots edited together so as
to ensure the behaviours being observed were not edited or manipulated, but none of the
videos obtained qualified for exclusion.

All videos featuring parrots manipulating objects were investigated for potential tool 413 use or borderline tool use. We defined tool use as the manipulation of an unattached object as an extension of the beak or foot to achieve a goal towards another object, individual, or 415 oneself¹⁰. Borderline tool use was similarly defined, except that it involved the use of an 416 object that was still attached to a substrate⁵⁴. For example, if individuals used a fallen 417 feather or stick for self-scratching this was defined as tool use, but using one's currently 418 attached tail feathers or cage furnishings for the same purpose was defined as borderline 419 tool use. Self-scratching had to involve slow and repeated movements of touching an object 420 to one's body (or, in the case of borderline tool use, rubbing repetitively against an 421 attached object⁶⁹). 422

All relevant videos were coded for video length, species, tool use presence

(yes/borderline), tool use type (e.g., scratching, feeding), the object being used (e.g.,

feather, stick), tool use target, human interaction (talking or handing object to parrot,

holding parrot), and the number of shots within each video. Our complete dataset also

includes the name for each video, link, subject name, sex (as declared by owner, as most

parrot species are not sexually dimorphic), publishing date, and dates found and coded.

Since YouTube is constantly growing and evolving in both its uploaded content and its proprietary recommendation algorithms, the same search strategy today would not return exactly the same videos as our search in July 2020. It is possible that some videos

have since been deleted or more videos of parrot tool use have since been uploaded.

However, this does not affect our conclusions about the number of parrot species found to

use tools in our initial search.

Data for parrot species. We collected data for 194 parrot species (Figures S9 -435 S11). We gathered feeding strategy data as a dichotomous variable ("generalist" or 436 "specialist") from the EltonTraits ecological database⁵². As per the database, specialists 437 were defined as species whose diet comprised at least 70% of a single food source. To 438 calculate relative brain size, we collated data from the literature for all known body mass 430 (g) and brain mass (g) values across parrots^{46–51}. For all species for which we obtained 440 body and brain mass data, we calculated the encephalisation quotient (EQ) using the 441 following formula⁸⁶: $BrainWeight/(0.12*BodyWeight(\frac{2}{3}))$. We found body mass and 442 brain mass data for a total of 194 parrot species. This included all tool-using species in our 443 video dataset, with the exception of three species: Diopsittaca nobilis, Psittacara 444 erythrogenys, and Coracopsis vasa. For the latter, we used values for the closely related 445 Coracopsis nigra. The other two species were excluded from the final dataset. 446

For modelling purposes, we coded research effort in both the scientific literature and 447 the YouTube videos. For the scientific literature, we operationalised research effort as the 448 number of papers published for each species' Latin name up to and including the first 449 paper containing tool use for that species. If no tool use had been identified in the 450 scientific literature for a species, then we coded the total number of papers published to date. We used the scientific database Scopus for coding the number of published papers. 452 For the videos, we coded research effort as the number of search hits for each species on YouTube. If tool use had been identified on YouTube, we estimated the number of search hits when the first video of tool use was published on YouTube, assuming linear growth of 455 search hits since the inception of YouTube. If tool use had not been identified, we used the 456 current number of search hits. 457

For phylogenetic data, we used the phylogenetic tool at www.birdtree.org 53 to

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compile 1000 posterior draws of phylogenetic trees for 174 of the 194 parrot species for which both EQ and genomic data exist. A single maximum clade credibility tree was generated from these posterior draws for visualisation purposes. In our analyses, we iterated over posterior draws of the phylogeny to account for phylogenetic uncertainty.

463 Quantification and Statistical Analysis

Phylogenetic signal. We used the *fitDiscrete* function in the *ape* R package⁸⁷ to calculate phylogenetic signal, for both the pre-survey and post-survey tool use data. We iterated the model over 100 posterior parrot phylogenies to incorporate phylogenetic uncertainty.

To infer unobserved probabilities of tool use across Causal model of tool use. 468 parrots, we proposed a causal model of observed tool use (Figure 1). We assumed that observed tool use in the scientific literature and in the videos is caused by both the 470 unobserved presence or absence of tool use and research effort, proxied by the number of 471 papers published on a species and the number of videos published on a species. Tool users 472 are more likely to be observed if they are well studied, but understudied tool users may go undetected. In addition, based on theory, we also assumed that unobserved tool use is caused by feeding strategy and relative brain size^{12–15,18–20}. Finally, we assumed that 475 shared phylogenetic history causes unobserved confounding and non-independence in 476 unobserved tool use, feeding strategy, and relative brain size across the parrot phylogeny. 477

Bayesian phylogenetic survival cure model. Given our proposed causal model,
we constructed a statistical model to impute unobserved probabilities of tool use and test
existing theories of the evolution of tool use in parrots. To understand the model, suppose
that we have the following observed variables for parrot species i. For the scientific
literature, we declare $N_{\text{Lit},i}$ as the number of papers published before and up to tool use
identification for species i (or, if tool use has not been identified, the total number of
papers published for species i) and $T_{\text{Lit},i}$ as a binary variable stating whether (1) or not (0)

tool use has yet been observed in the scientific literature for species i. For the videos, we declare $N_{Vid,i}$ as the number of videos published before and up to tool use identification for species i (or, if tool use has not been identified, the total number of videos published for species i) and $T_{Vid,i}$ as a binary variable stating whether (1) or not (0) tool use has yet been observed in the videos for species i. Additionally, F_i and EQ_i are feeding strategy and encephalisation quotient values for species i and we have a phylogenetic distance matrix D that describes the patristic distances between all parrot species.

We assume that species i is a non-tool-user with some probability p_i . We also assume that tool use is identified in the scientific literature and the videos at constant rates λ_{Lit} and λ_{Vid} following exponential survival functions. Given these assumptions, we can then describe the different ways in which variables N_{Lit} and N_{Vid} can be distributed. Focusing on the scientific literature, if tool use has been observed ($T_{\text{Lit},i} = 1$), then the likelihood for $N_{\text{Lit},i}$ is:

$$\Pr(N_{\text{Lit},i}|T_{\text{Lit},i} = 1, p_i, \lambda_{\text{Lit}}) = (1 - p_i) \cdot \text{Exponential}(N_{\text{Lit},i}|\lambda_{\text{Lit}})$$
(1)

On the other hand, if tool use has not yet been observed ($T_{Lit,i} = 0$), there are two ways that the outcome variable could have been realised. First, the species could be a non-tool-user with probability p_i . Second, the species could be a tool-user with probability (1 - p_i) that has been censored and has not had its tool use measured yet. Together, then, the likelihood for $N_{Lit,i}$ is:

$$Pr(N_{\text{Lit},i}|T_{\text{Lit},i} = 0, p_i, \lambda_{\text{Lit}}) = p_i + ((1 - p_i) \cdot \text{Exponential-CCDF}(N_{\text{Lit},i}|\lambda_{\text{Lit}}))$$
(2)

The Exponential-CCDF function allows for the censored nature of the data. The

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same data generating process is assumed to underlie the videos.

We define the mixture likelihood SurvivalCure as the distribution above, with parameters p (the probability of being a non-tool-user) and λ (the rate of the exponential distribution). We use an Ornstein-Uhlenbeck Gaussian process⁸⁸ to model phylogenetic covariance. Below, we specify the full model with priors:

$$N_{\text{Lit},i} \sim \text{SurvivalCure}(\lambda_{\text{Lit},i}, p_i)$$

$$N_{\text{Vid},i} \sim \text{SurvivalCure}(\lambda_{\text{Vid},i}, p_i)$$

$$\lambda_{\text{Lit},i} = 1/\exp(\gamma_{\text{Lit}})$$

$$\lambda_{\text{Vid},i} = 1/\exp(\gamma_{\text{Vid}})$$

$$\log \text{it}(p_i) = \alpha_{\text{FEEDING}[i]} + \beta \text{EQ}_i + k_{\text{SPECIES}[i]}$$

$$\begin{pmatrix} k_1 \\ k_2 \\ \dots \\ k_n \end{pmatrix} \sim \text{MVNormal} \begin{pmatrix} 0 \\ 0 \\ \dots \\ 0 \end{pmatrix}, \mathbf{K}$$

$$\begin{pmatrix} \mathbf{K}_{\text{ij}} \\ \mathbf{K}_{\text{ij$$

The priors for this model produce reasonable prior predictions of the probabilities of tool use for each parrot species (Figure S12), but a sensitivity analysis revealed that the ranking and posterior probabilities reported in the main text were robust to modifying these priors (Figure S13). We estimated the posterior distribution of this model using Hamiltonian Monte Carlo as implemented in Stan version 2.26.1⁸⁹. We ran the model for 4000 samples, with 2000 warmup samples, and iterated the model over 100 posterior parrot

phylogenies to incorporate phylogenetic uncertainty. R-hat values and effective sample sizes suggested that the model converged normally. Trace plots are reported in Figure S14. We report equal-tailed credible intervals to describe the posterior distribution of this model in the main text.

To validate our method, we fitted the model to 100 simulated datasets with known parameters. The model was able to successfully recover those parameters (Figure S15). We also ran a leave-one-species-out exercise to ensure that we could accurately predict known tool users. We repeated this approach for each known tool user by setting observed tool use to zero. Cross-validation results are reported in the main text.

Ancestral state reconstruction. To determine whether the identification of novel 524 tool-using species has implications for our understanding of the evolutionary origins of tool 525 use in parrots, we fitted three exploratory ancestral state reconstruction models. We used 526 the ancThresh function from the phytools R package⁹⁰, iterating the function over 100 527 posterior parrot phylogenies. This function estimates discrete ancestral states by assuming 528 the evolution of a latent continuous variable following an Ornstein-Uhlenbeck process. We 529 fitted this model to three different outcome variables: (i) presence vs. absence of tool use 530 in scientific literature only, (ii) presence vs. absence of tool use in literature and/or videos, 531 and (iii) the median predicted probabilities of tool use from the phylogenetic survival cure 532 model. 533

Reproducibility. All analyses were conducted in R v4.2.1.⁹¹. Visualisations were produced using the *ggtree*⁹², *ggplot2*⁹³, and *cowplot*⁹⁴ packages. The manuscript was reproducibly generated using the *targets*⁹⁵ and *papaja*⁹⁶ packages. Code to reproduce all analyses and figures can be found here: https://github.com/ScottClaessens/phyloParrot

Supplemental Information

Digital video platforms and phylogenetic modelling reveal parrot tool use is not rare Amalia P. M. Bastos^{1,2}, Scott Claessens², Ximena J. Nelson³, David Welch⁴, Quentin D. Atkinson², Alex H. Taylor^{2,3,5,6}

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Supplemental Figures

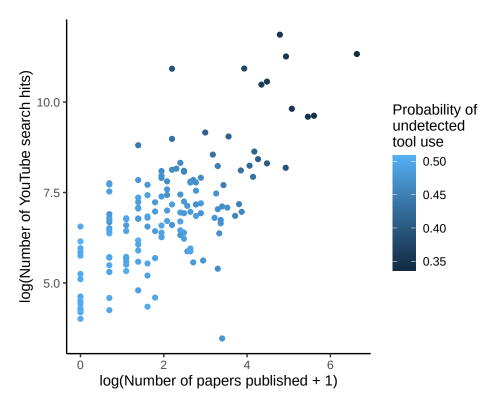


Figure S1. Median posterior probabilities of undetected tool use for each parrot species without observed evidence of tool use from reduced model. This reduced version of the phylogenetic survival cure model does not contain relative brain size or feeding strategy as predictors, nor does it contain any phylogenetic covariance. The only information included in the model is the number of papers published and the number of YouTube search hits for each species. Each point is a parrot species without observed evidence of tool use, and the colour of the points scales with the probability of undetected tool use. All else being equal, those species with fewer published papers and fewer YouTube search hits have a higher probability of being undetected tool users.

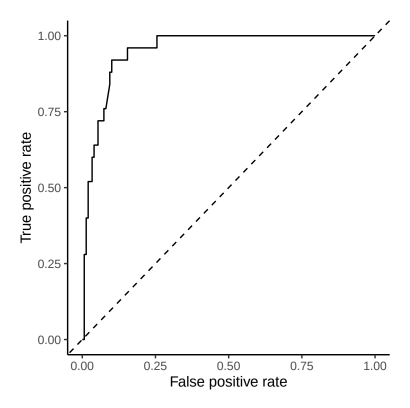


Figure S2. Receiver operating characteristic (ROC) curve for the phylogenetic survival cure model. The area-under-the-curve in this plot is 0.95, suggesting that the model is able to adequately classify observed tool users and non-tool users.

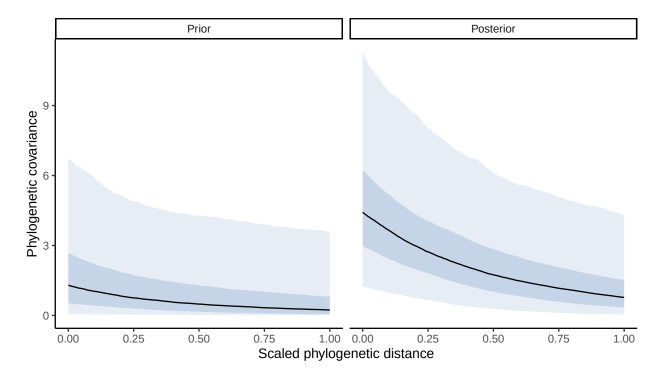


Figure S3. Prior and posterior phylogenetic covariance functions from the Bayesian survival cure model fitted to the full dataset. Lines are median posterior functions and shaded areas are 50% and 95% credible intervals.

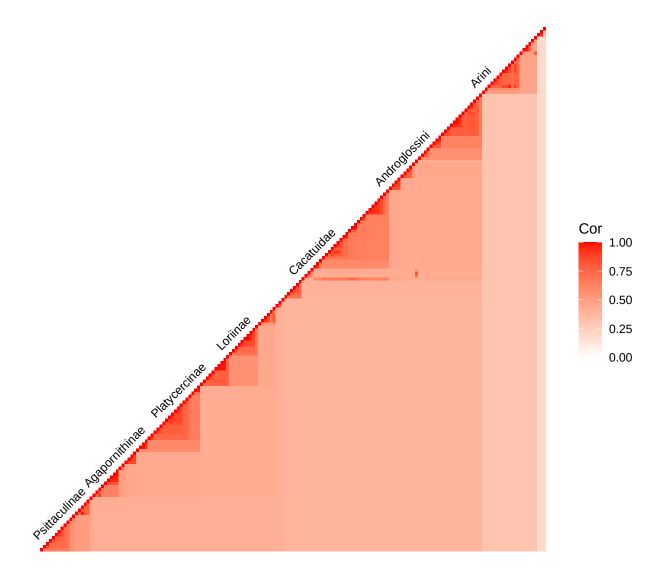


Figure S4. Between-species correlation matrix implied by the posterior phylogenetic covariance function from the Bayesian survival cure model. Correlations are median posterior estimates. Individual species names omitted for space reasons.

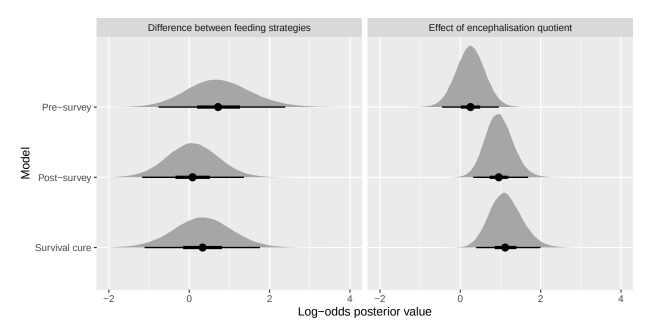


Figure S5. Comparing results between the survival cure model and models fitted to the pre-survey and post-survey data without any survival cure component. Densities are full posterior distributions from three separate models iterated over 100 posterior parrot phylogenies. Points represent posterior medians, and lines represent 50% and 95% credible intervals.

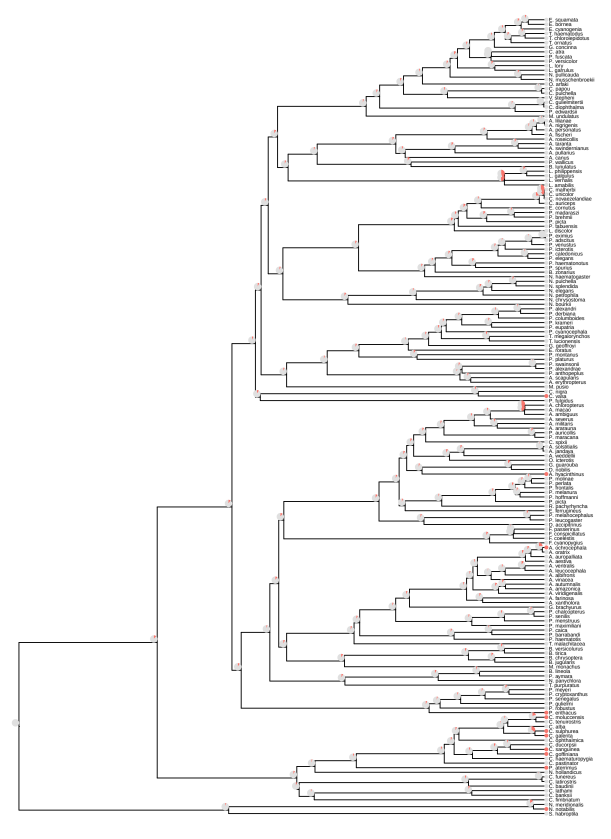


Figure S6. Results of exploratory ancestral state reconstruction analysis fitted to pre-video-survey data, represented on a maximum clade credibility tree. Tip nodes represent the presence (red) or absence (grey) of observed tool use in the scientific literature. Pie charts represent the posterior probability of tool use presence at each ancestral node.

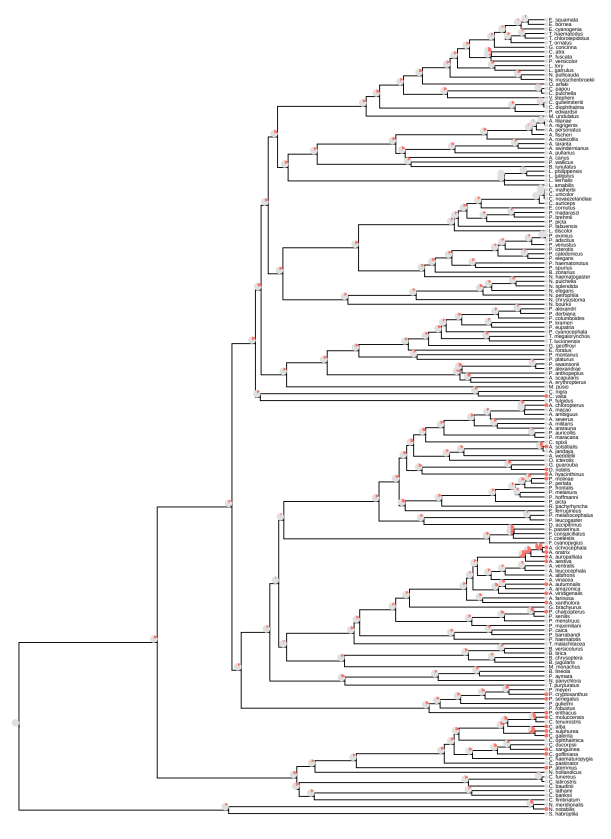


Figure S7. Results of exploratory ancestral state reconstruction analysis fitted to post-video-survey data, represented on a maximum clade credibility tree. Tip nodes represent the presence (red) or absence (grey) of observed tool use in the scientific literature and the video survey. Pie charts represent the posterior probability of tool use presence at each ancestral node.

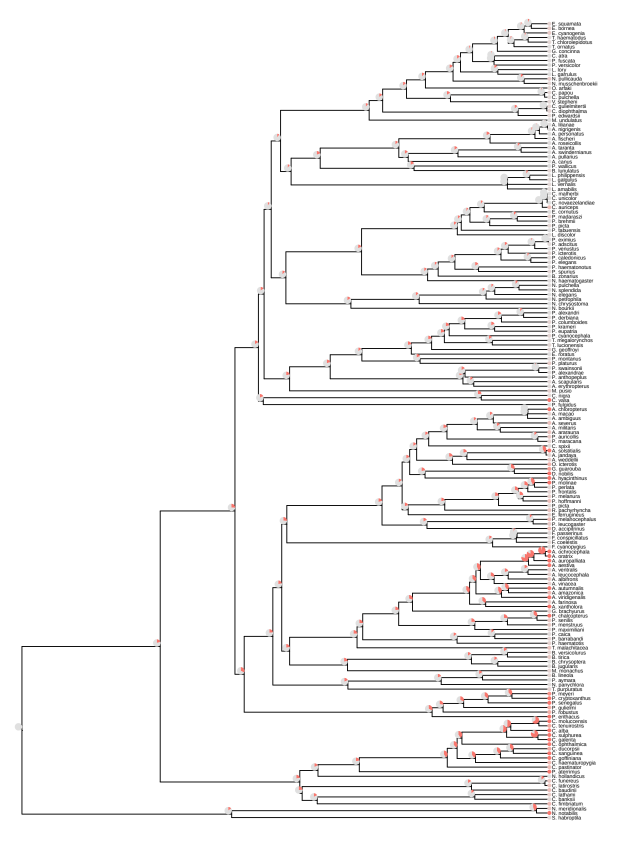


Figure S8. Results of exploratory ancestral state reconstruction analysis fitted to predicted probabilities from the phylogenetic survival cure model, represented on a maximum clade credibility tree. Tip nodes represent the median posterior predicted probabilities of tool use from the phylogenetic survival cure model, with more red indicating an increasing probability of tool use presence and more grey indicating a decreasing probability of tool use presence. Pie charts represent the posterior probability of tool use presence at each ancestral node.

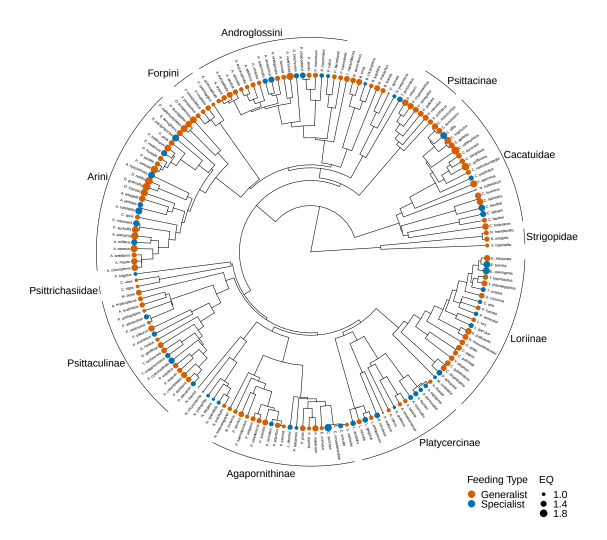


Figure S9. Data on encephalisation quotient and feeding strategy for all parrots, presented on a maximum clade credibility tree. Tip points are coloured according to feeding generalism (orange) and specialism (blue), and scaled according to encephalisation quotient (EQ).

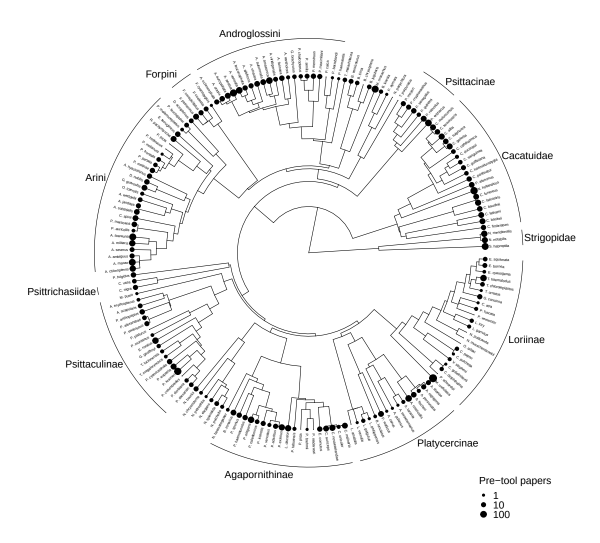


Figure S10. Data on number of scientific publications until tool use discovery for all parrots, presented on a maximum clade credibility tree. Tip points are scaled according to the number of published papers up until tool use discovery (or, if tool use has not been observed, the current number of published papers).



Figure S11. Data on number of video search hits until tool use discovery for all parrots, presented on a maximum clade credibility tree. Tip points are scaled according to the estimated number of video search hits up until tool use discovery (or, if tool use has not been observed, the current number of video search hits).

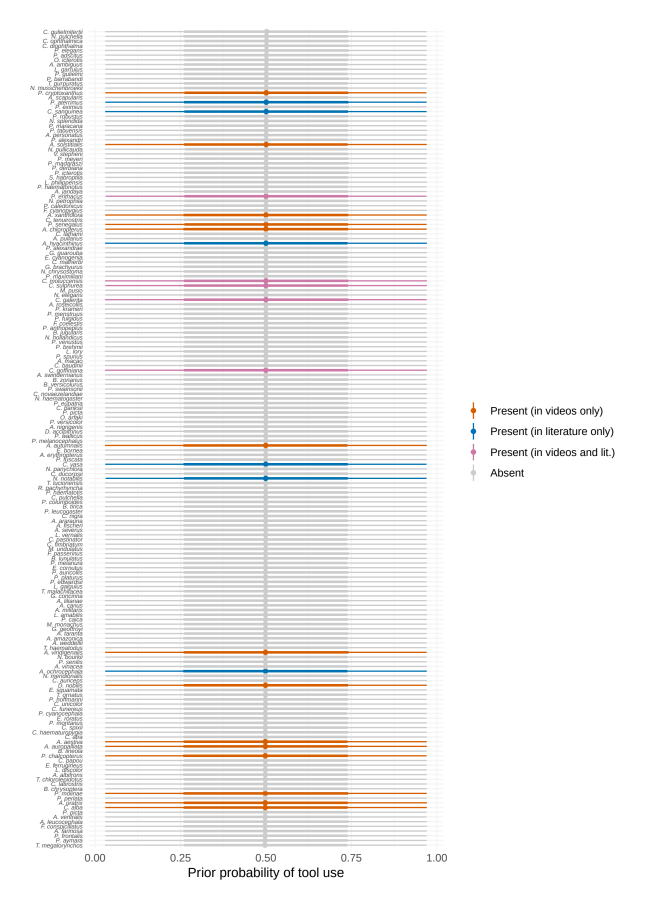


Figure S12. Prior predicted probabilities of tool use for each species from our phylogenetic survival cure model. Points are prior medians and lines are 50% and 95% credible intervals.

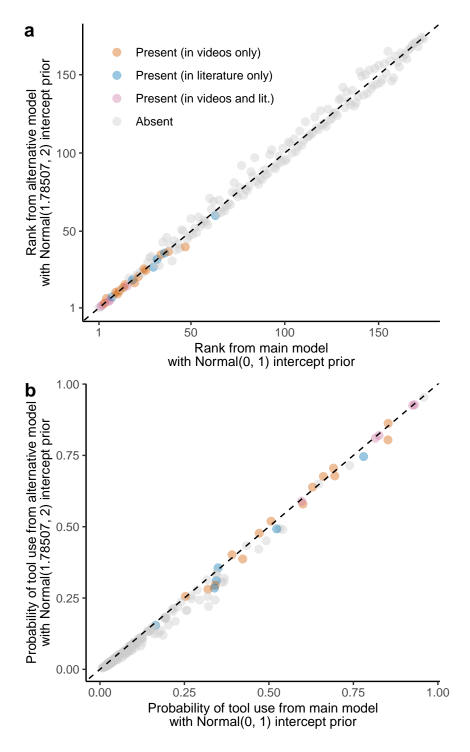


Figure S13. Results of sensitivity analysis. The phylogenetic survival cure model was fitted with either a standard Normal(0, 1) prior on the intercept or an alternative Normal(1.78507, 2) prior on the intercept. This latter prior is wider on the logit scale and roughly converts to a 0.86 prior probability of non-tool-use (or a 0.14 prior probability of tool-use, which is the proportion of tool users in the dataset). The sensitivity analysis showed that changing this intercept prior did not have a marked impact on (a) the posterior rankings of parrot species from 1st to 174th or (b) the median posterior probabilities of tool use for parrot species.

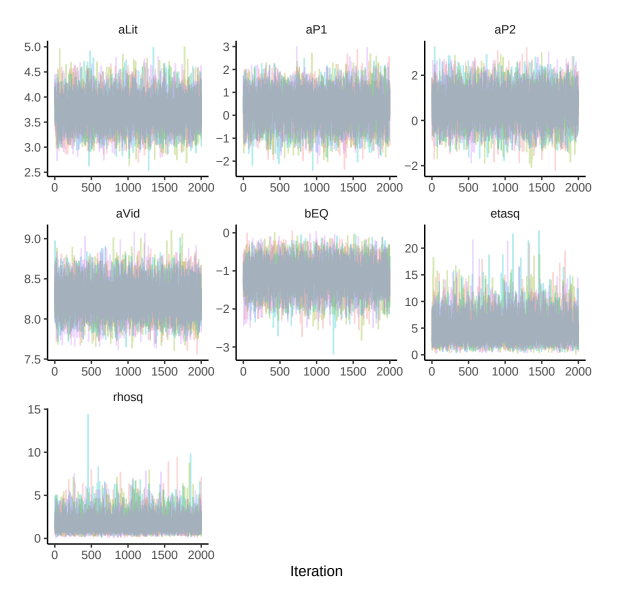


Figure S14. Trace plots for the Bayesian phylogenetic survival cure model. Only four chains are shown for ease of presentation.

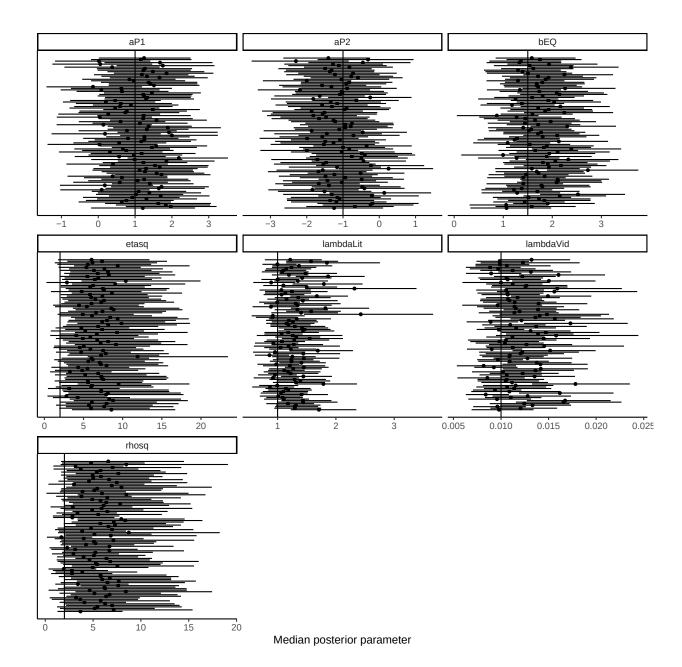


Figure S15. Posterior estimates from Bayesian survival cure models fitted to 100 datasets simulated with known parameters. Each dataset consisted of 100 species. Known parameters are presented as solid vertical lines, whereas points and horizontal lines represent posterior medians and 95% credible intervals. The models were successfully able to recapture the parameters from the simulated datasets.