1 **Comparative lifespan and healthspan of nonhuman primate species common to biomedical research**

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- **Abstract**

There is a critical need to generate age- and sex-specific survival curves to characterize chronological aging consistently across nonhuman primates (NHP) used in biomedical research. Accurate measures of chronological aging are essential for inferences into genetic, demographic, and physiological variables driving differences in NHP lifespan within and between species. Understanding NHP lifespans is relevant to public health because unraveling the demographic, molecular, and clinical bases of health across the life course in translationally relevant NHP species is fundamentally important to the study of human aging. Data from more than 110,000 captive individual NHP were contributed by 15 major research institutions to generate sex-specific Kaplan-Meier survival curves using uniform methods in 12 translational aging models: *Callithrix jacchus* (common marmoset), *Chlorocebus aethiops sabaeus* (vervet/African green), *Macaca fascicularis* (cynomolgus macaque), *M. fuscata* (Japanese macaque), *M. mulatta* (rhesus macaque), *M. nemestrina* (pigtail macaque), *M. radiata* (bonnet macaque), *Pan troglodytes* spp. (chimpanzee), *Papio hamadryas* spp. (baboon), *Plecturocebus cupreus* (coppery titi monkey), *Saguinus oedipus* (cotton-top tamarin), and *Saimiri* spp. (squirrel monkey). After employing strict inclusion criteria, primary analysis results are based on 12,269 NHP that survived to adulthood and died of natural/health-related causes. A secondary analysis was completed for 32,616 NHP that died of any 47 cause. For the primary analyses, we report ages of 25th, 50th, 75th, and 85th percentiles of survival, maximum observed ages, rates of survivorship, and sex-based differences captured by quantile regression models and Kolmogorov-Smirnov tests. Our findings show a pattern of reduced male survival among catarrhines (African and Asian primates), especially macaques, but not platyrrhines (Central and South American primates). For many species, median lifespans were lower than previously reported. An important consideration is that these analyses may offer a better reflection of healthspan than lifespan. Captive NHP used in research are typically euthanized for humane welfare reasons before their natural end of life, often after diagnosis of their first major disease requiring long-term treatment with reduced quality of life (e.g., endometriosis, cancer, osteoarthritis).

Supporting the idea that these data are capturing healthspan, for several species typical age at onset of chronic disease is similar to the median lifespan estimates. This data resource represents the most comprehensive characterization of sex-specific lifespan and age-at-death distributions for 12 biomedically relevant species, to date. The results clarify the relationships among NHP ages and will provide a valuable resource for the aging research community, improving human-NHP age equivalencies, informing investigators of the expected survival rates of NHP assigned to studies, providing a metric for comparisons in future studies, and contributing to our understanding of the factors that drive lifespan differences within and among species.

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Introduction

Nonhuman primates (NHPs) are genetically, physiologically, and behaviorally the best translational models for human aging as their genomes, developmental trajectory, reproductive strategies, and aging-related changes in physical function, cognitive function, and disease development are more similar to humans than those of 67 other mammals.^{1–4} Yet we know little about longevity in the NHPs most commonly used as translational models. Few studies have attempted cross-species comparisons and reports are often contradictory, likely due to the use of different methodological approaches (e.g., inclusion criteria). To determine how NHP ages correspond with human age, it is essential to fully characterize the demography of NHP longevity within each species, rather than focusing on individual reports of maximum longevity. Numerous publications list NHP maximum lifespans in tables that include a variety of other life history features, but few cite primary sources. This leads to overreporting of the same statistics without verifying the validity of the measure or the relevance to animals under study. For example, 37.5 years is often cited as the lifespan of baboons (*Papio hamadryas* spp.).^{5–8} However, tracing citations to the primary source reveals that this statistic comes from a single baboon that died at the Brookfield Zoo in 1972; the birth date is given as June 1, 1935 (one year after the zoo opened), $77\quad$ but it is not documented whether this date is known or estimated. 9 This estimate of maximum longevity in baboons is not particularly useful without additional context indicating how many baboons survive to the 79 maximum or what the median baboon lifespan is. Median captive baboon lifespan has been reported as 21^{10} or 11¹¹ years but the report of maximum longevity is more frequently cited. It is likely that the discrepancy in median baboon lifespan reflects differences in methodological approaches to data analysis. This example in

baboons highlights how differences in analytic approaches across studies make it difficult to compare reports within or across species. The unclear data on NHP lifespan, such as the reporting of maximum longevity to indicate "lifespan," creates confusion in scientific analysis and in the peer review process.

Cross-species comparisons are a major goal of aging research since they can reveal factors contributing to variation in lifespans. Inconsistent lifespan estimates are problematic when looking at a single species, and the problem is compounded by cross-species comparisons. We address this knowledge gap by creating rigorous and reproducible survivorship data, identifying mortality risk and its relationship to biological age at different chronological ages, and examining the shape of mortality and healthspan curves across 12 captive NHP species. The initial dataset, prior to quality control and filtering, included lifespan data from 114,255 animals from 58 species at 15 institutions. We highlight that while maximum age is an easily reported statistic as it is purely observational, calculating median lifespan is more challenging, as methodological decisions about inclusion and exclusion criteria vary among studies, producing substantial discrepancies across cohorts and species. With the data herein, we have the unique ability to calculate survival probabilities using the same criteria for all 12 species, producing the most methodologically consistent cross-species comparison to date. 97 The value of such a large dataset is the ability to filter the data to the most representative sample and retain adequate sample sizes for statistical analyses. In this study, survival curves were generated on animals that survived to at least adulthood (defined in Methods) because, as in most mammals including humans, risk of death in infancy is substantial and strongly biases the median lifespan. Primary results and comparisons by sex are built using data from animals that died of natural causes or were euthanized for clinical/health reasons. This report provides comprehensive data summaries and tools to improve biomedical research involving NHPs within and beyond the field of aging.

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Results

 Primary analyses. Sample counts of primary analysis datasets, featuring natural or health-related deaths only, are shown in **Table 1**. Maximum observed age including all types of deaths (e.g., research-related sacrifice, clinical/health-related euthanasia, and natural), as well as median age at death calculated from only natural

 and clinical deaths, are summarized by sex and species in **Table 2**. **Figure 1** shows a scatter plot of datapoints for natural and clinical deaths, with medians, interquartile ranges, and proportions of data by sex and species. Combined survival curves for all 12 species in males and females are shown in **Figure 2**. To evaluate the rate of decline for the survivorship curves, data from the first and last quartiles of the Kaplan-Meier survivorship function were fit to an exponential model that captures rate of decay (i.e., change in probability of death), and species were then compared within and between sexes. Comparing first and last quartiles illustrated that species predominantly experienced faster rates of death within the first quartile of adulthood. Comparing male and female rates of decline within both quartiles highlighted the faster rates of decline for males within the first quartile. However, in the last quartile, this pattern was nearly reversed; the majority of species (except cotton- top tamarin, vervet/African green monkey, and common marmoset) exhibited slower rates of decline in males compared to females (**Figure 3**).

 For each species, individual survival curves are shown in **Figure 4** and sex-based comparisons in **Table 3**. In most species, males showed reduced survival compared to females. Among vervets, Japanese macaques, 123 and chimpanzees, males showed reduced survival at every age with a different overall distribution of age at death. *C*ynomolgus macaque and baboon males showed reduced survival compared to females at younger 125 ages (25th and 50th percentiles), but there was no difference in survival at later stages of life. Rhesus macaque 126 males showed reduced survival compared to females at the $25th$, 50th, and 75th percentiles, but females had lower age of survival at the 85th percentile. There was a strong difference in the distribution of age at death 128 between males and females (P-value= $2.20x10^{-16}$). Pig-tailed macaque males showed reduced survival 129 compared to females early in life (25%) but the sexes were similar at other ages. In contrast, females showed reduced survival compared to males at every age in common marmosets. Male and female survival was similar at every age with no difference in the distribution of age at death between sexes for cotton-top tamarins and 132 squirrel monkeys. There was also no difference in distributions for coppery titi monkeys and bonnet macaques; 133 however, the modest sample size for the species limits power to detect small differences.

 Secondary analyses. *C*ensored data (deaths due to research sacrifice and colony management) were biased by sex (**Extended Data Figure 1**) and prevented statistical comparisons between males and females when 137 including censored data.¹² However, as a secondary analysis, survival curves that include censored events are presented as extended data (**Extended Data Figure 2**) for reference. Across species, inclusion of additional datapoints from censored events increased median lifespan estimates. We note that the high proportion of censored events (**Extended Data Figure 1**), especially in some species (i.e., greater than 50% of deaths in baboons, cynomolgus, pigtails, rhesus, squirrel monkeys, and vervets), yielded survivorship functions that never reach zero, limiting utility and inference for the full lifespan.

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Discussion

 Lifespan vs healthspan. A major consideration of note for this study is that few research NHPs live until natural death. Most are humanely euthanized due to study protocols or clinical determinations based on quality 147 of life. The issues considered by veterinarians in making euthanasia decisions vary by facility and study protocol, but a common approach is to euthanize at the first diagnosis of major disease or injury requiring long- term treatment with reduced quality of life. Reasons for humane euthanasia may include such diverse conditions as advanced spinal or knee osteoarthritis, endometriosis, broken limbs, tumors, and meningitis – not all of which are the result of aging-related diseases. Therefore, we posit that these findings may be measuring healthspan rather than lifespan in NHP cohorts housed at research facilities. For our survival analyses, this potential limitation is partially mediated by our very large database, which enabled analyses even after removing experimental and other non-clinical deaths.

 Supporting the idea that we are measuring healthspan rather than lifespan, for several species, typical age at onset of chronic disease is similar to the median lifespan estimates. Among baboons, age-related diseases are apparent around 9 years old (e.g., edema, kyphosis, prolapse, myocarditis), and by 12 years many more are 159 evident (e.g., pancreatitis, stricture, lymphosarcoma).¹³ Median baboon lifespan in this report is 10.1 years for males and 11.1 years for females. Marmoset age-related diseases tend to emerge in animals >6 years old, 161 including cardiovascular disease, diabetes, and neoplasias.¹⁴ Median marmoset lifespan in our study is 5.5

 years in males and 5.0 years in females. Rhesus macaques are on average diagnosed with the first chronic 163 condition at age 9.0 years and the second at age 10.7 years.¹⁵ Median rhesus lifespan in our study is 9.1 years in males and 10.6 years in females. Differences in veterinary care for these conditions mean that some 165 pathologies in some species may be treated medically, whereas others proceed to veterinarian-suggested euthanasia. We speculate that zoo NHPs may be treated for more chronic conditions than research NHPs and would make a useful lifespan and healthspan comparison to humans.

 The ability to make more accurate comparisons between NHP age and the human equivalent was a primary 170 goal of the current analyses. Since the NHP estimates herein may be closer to healthspan than lifespan, it is useful to consider them in relation to human healthspan. The most frequently studied measures of human healthspan are deficit accumulation indices, which measure accumulation of health deficits and decline in 173 physical function or frailty.^{16–20} In one study of 66,589 Canadians in the National Population Health Survey, accumulation of health deficits was gradual before age 46 years, with 40% of 45-50 year-olds having a frailty 175 index score of 0 (no health deficits); starting at age 46, deficit accumulation was much more rapid, and at age 176 80, only 5% still had a score of 0.^{20,21} Among 73,396 people from the Longitudinal Ageing Study in India, 177 average age of onset of any chronic disease was 53 years.²² We speculate that our NHP median lifespan estimates may align better with human onset and accumulation of health deficits, rather than human lifespan. However, our analysis does not address onset of health deficits, and we are unable to distinguish between which NHPs died at the end of their lifespan versus those which died at the end of their healthspan. Therefore, we are unable to make specific comparisons between human and NHP healthspans.

 Sources of variation within and between species. Our findings show great variation in adult life expectancy among all 12 species, in contrast to a prior cross-species analysis of six primate species that found little 185 variation in adult survival.²³ Many factors contribute to variation in adult survival. Some may assume that in captive research populations, quality of veterinary care is a major driving force. While this may have been 187 important in the early years of NHP research, most species have been in captivity for decades and quality care is well defined. Institutional management practices are important factors, such as how decisions are made

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199 Within species, life history features can influence lifespan. It has been proposed that reproductive strategies 200 play an evolutionary role in regulating lifespan, since there may be tradeoffs between female fertility, $201\quad$ investment in offspring, and longevity, 26 although this long-held view has been challenged since the 202 Felationships between reproduction and longevity are not consistent across species.^{27,28} Adult body size also 203 factors into survival because a longer period of growth will likely result in later reproductive maturity and a 204 greater need for investment in offspring. In our data, common marmosets have the shortest maximum and 205 median lifespan of all 12 species. Marmosets are also the smallest species (average weight 350-400 g), reach 206 adulthood at the youngest age (1.5 years), and usually give birth to twins.^{14,29} However, cotton-top tamarins, 207 the other small (average weight in captivity 565.7 g), quickly maturing (2.5 years at adulthood), twinning 208 callitrichine³⁰ in this study, has maximum and median lifespan resembling that of several larger bodied, slower 209 maturing species that give birth to singletons, including squirrel monkeys, baboons, vervets, and macaques. It 210 is unclear to what extent these patterns are driven by inherent species characteristics versus institutional 211 practices, but it would be advantageous to explore this question in future studies.

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213 **Sex-based differences.** Among primates, males have been shown to have higher age-specific mortality than 214 females throughout adulthood.³¹ We see this in some species included in the current study. One pattern is 215 shorter lifespan among macaque males. Five macaque species (*Macaca* spp.) are reported here. In three

216 species males have shorter median lifespan than females (cynomolgus, Japanese, and rhesus macaques). In 217 pigtails, males have lower survival probability in early adulthood (25%) but similar survival probability at older 218 ages, and in bonnet macaques male lifespan appears shorter in the curves and estimates, but sample size 219 may be too small to detect a difference (female n=43, male n=19). This pattern seems to extend to all of the 220 parvorder Catarrhini (Old World monkeys- Cercopithecoidea and apes- Hominoidea). Vervets have the largest 221 sex-based differential with median age of 8.3 years for males and 17.9 years for females. For baboons, males 222 show borderline lower survival probability at the 25th and 75th percentiles. Male chimpanzees also have lower 223 survival probability relative to females at every life stage.

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225 In contrast, in the parvorder Platyrrhini (Central and South American monkeys), there is generally no difference 226 between males and females in survival estimates. For context, a phylogenetic tree for the 12 species in this study is shown in **Extended Data Figure 3**. ³² 227 The exception is the common marmoset, with lower female 228 survival at every age, replicating the findings of another marmoset report.¹⁴ The relatively short female 229 marmoset lifespan is related to their high fertility rates.^{26,29} There are no differences in survival between males 230 and females in coppery titi monkeys, squirrel monkeys, or cotton-top tamarins. A prior primate lifespan 231 comparison that suggested female primates have longer lifespan than males included several catarrhine 232 species but few data from platyrrhine species.³¹A recent study of coppery titi monkey lifespan showed a trend 233 toward longer lifespan in males relative to females using the same population of monkeys in the current study 234 but with different inclusion criteria. 33

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236 It is difficult to know if the observed sex-based differences between catarrhine versus platyrrhine species are due to inherent species characteristics, institutional practices, or their interactions. For example, in catarrhine monkeys, it is common to house a single breeding or vasectomized male with multiple females. Fewer males than females are needed for breeding programs because males will mate with multiple females. In some 240 species, especially baboons, males are much larger than females, requiring more space and resources. These factors and more mean males and females are not equally distributed and are subject to different animal selection practices in research institutions. The difference is also evident in the sample size. Before data

 filtering, the sample size included 44,704 females and 43,413 males. After data filtering, there were 8,296 females and 3,973 males. A larger proportion of the males were filtered out of the analyses because of research-related endpoints or humane euthanasia for management reasons, reflecting bias in how sexes are 246 deployed in research.

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248 **Comparison with prior reports of captive NHP lifespan.** As mentioned in the introduction, captive baboon 249 $\,$ maximum lifespan has been reported as 37.5 years, $^{5\textrm{--}8}$ and median lifespan as 21 10 or 11 11 years. Our median 250 lifespan findings align with the lowest of those estimates, and close inspection of the methods used to arrive at 251 that estimate reveals that the study employed similar inclusion and exclusion criteria as the current study.¹¹ 252 The 37.5 year estimate is based on a single zoo baboon⁹ and is a rare case of extreme maximum longevity. 253 The 21-year baboon lifespan estimate uses different methods from the current study, such as inclusion of live animals as right censored datapoints.10 254 In another report that includes 4,480 zoo baboons, male *P. hamadryas* 255 were estimated to live 13.2 years and females 17.1 years from birth.²³ We expect that this difference is due to 256 both methodological differences in calculating median lifespan and differences in the veterinary care for the 257 small numbers of baboons in zoo settings, e.g., they frequently receive long-term treatment for chronic 258 diseases. It may also be due to differences between hamadryas and the mixed baboons in our study. Prior 259 reports of lifespan of rhesus macaques have hovered around a median lifespan of 25 years and maximum 40 260 years, but again, these studies employed right censored data approaches.^{24,34–36} In contrast, our median 261 lifespan estimate for rhesus is 7.9 years in males and 10.3 years in females using data only from animals with 262 known ages at death, rather than including ages from still living animals with a right censored approach. To 263 highlight this methodological difference, we provide survivorship probabilities with censored data for reference 264 (**Extended Data Figure 2**). A prior study of common marmosets at a single institution estimated median 265 lifespan of 6.5 years in animals that survived to at least two years (compared with our starting age of 1.5 266 years).¹⁴ Another marmoset study from a different institution estimated median lifespan at four years in 267 marmosets that survived for 60 days; the same study reported cotton-top tamarin median life expectancy of 7.2 268 years.³⁷ Our estimates from marmosets at 4 different institutions are 5.3 years in females and 6.0 years in 269 males. For cotton-top tamarins, our estimates of median lifespan (from animals living at one institution) are 9.6

270 years for males and 8.9 years for females. Chimpanzee median survival in a biomedical research population 271 has been reported as 31.0 years in males and 38.8 years in females among individuals who reached 1 year of 272 $\,$ age.³⁸ In a zoo population, male chimpanzees lived a median of 26.0 years and females 30.5 years from 273 birth.²³ Our estimates are 33.0 years in males and 44.0 years in females among individuals who reached ten 274 years of age and are therefore fairly consistent with previous reports. For coppery titi monkeys, median 275 lifespan has been reported as 14.9 years in males and 11.4 years in females among individuals surviving to 31 276 days,³³ compared with our estimates of 8.6 years for males and 9.2 years for females. Once again, the 277 differences between estimates in our studies and prior reports likely arise methodologically, such as choices 278 made about age of inclusion and use of a right censored approach to include individuals still alive and/or those 279 euthanized for research-related endpoints. A major strength of the current study is the use of uniform methods 280 across 12 different NHP species.

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282 **Importance of data filtering.** This study highlights the necessity of thorough methodological documentation in 283 NHP lifespan studies. As illustrated with our primary and secondary analyses, filtering and methodological 284 decisions impact the results and interpretation. The simplest example is the minimum age threshold for 285 computing the survivorship functions. Including juveniles dramatically lowers median lifespan due to high rates 286 of juvenile mortality among primates. Additionally, by including only animals that were born and died at the 287 same institute, it sometimes eliminated the oldest known individuals from the dataset, such as two 19-year-old 288 SNPRC marmosets; however, these instances were rare in our very large sample. Decisions that greatly 289 reduced our analysis sample size, such as date-of-birth (DOB) cutoffs, are a privilege of a large initial (pre-290 filtered) dataset. So, while the DOB cutoffs greatly reduced our final sample size, it removed bias associated 291 with very early deaths (since our dataset did not include currently alive animals). Overall, given the impact of 292 filtering decisions, we emphasize the need for robust reporting of the decision criteria in NHP survival studies. 293 We encourage authors to follow the ARRIVE guidelines (Animal Research: Reporting of In Vivo Experiments; 294 https://arriveguidelines.org/), a checklist for full and transparent reporting aimed at improving rigor, 295 transparency, and reproducibility in animal research.³⁹ In longevity research, it is particularly crucial to report 296 inclusion and exclusion criteria in addition to the details of statistical approaches.

 Limitations. One limitation of the study is that the stringent inclusion criteria reduced our starting sample size by 86%. This was necessary to ensure appropriate comparisons across institutions and species. For example, some species (cynomolgus, pigtails, baboons) have a very high percentage of deaths by research sacrifice, 301 rather than by natural or health-related causes. Including research-related deaths as right censored data results in highly skewed models with limited utility for these species (e.g., survival curves for female baboons do not converge past the median survivorship when including censored data). Further, censoring was biased by sex because of the differences in research utilization and breeding needs, statistically hindering the 305 possibility of comparisons between males and females. Therefore, primary analyses were limited to data from natural or clinical deaths, eliminating the need for right censoring. Another constraint of the study is our limited knowledge of specific cause of death. Differences in institutional death coding systems make it difficult to easily 308 determine cause of death, since some record systems group many types of deaths, while others have more granular codes to distinguish among death types.

Conclusions. The need for comparative analyses of lifespans across species has been widely

312 acknowledged.⁴⁰ Investigators need access to reliable lifespan tables, survivorship graphs, and maximum

313 lifespan measurements to conduct relevant translational aging studies. Here we provide the largest dataset yet

assembled from captive research NHPs. These data provide a valuable comparative resource for translational

NHP research, primary data on multispecies NHP lifespan in captivity, and context for consideration of

morbidity and mortality in the study of diverse diseases.

Methods

Species

 Twelve NHP species for analyses are shown in **Table 1**. We are considering all members of the genus *Papio* a single species and considering Indian- and Chinese-origin rhesus macaques together, as captive research 322 baboons have a high degree of morphotype mixing^{41,42} and captive rhesus are similarly highly admixed from

323 these geographic source populations.⁴³ We included chimpanzees (*Pan troglodytes* spp.), but it must be noted

 that biomedical research with great apes is heavily restricted across the world. Still, many retired chimpanzees reside at research facilities and they provide a valuable comparison since their estimated lifespan is between 326 that of humans and the monkey species commonly found at biomedical research facilities. Similarly, while cotton-top tamarins (*Saguinus oedipus*) were at one time biomedical research models, they have not been used for that purpose since 2008 when deforestation resulted in animals being listed as critically endangered.

Participating institutions

 Data from eight United States National Primate Research Centers (NPRCs) are included: California (CNPRC), Emory (ENPRC), New England (NEPRC; this center is no longer open but we obtained archival data), Oregon (ONPRC), Southwest (SNPRC), Tulane (TNPRC), Washington (WaNPRC), and Wisconsin (WNPRC). Data also originated from Primate Research Center IPB University in Indonesia, Keeling Center for Comparative Medicine and Research at The University of Texas MD Anderson Cancer Center, National Institute on Aging Intramural Research Program, Sam and Ann Barshop Institute for Longevity and Aging Studies at UT Health San Antonio, Vervet Research Colony at Wake Forest University, and Yale University. **Table S1** shows species sample sizes contributed by each institute. A data extraction standard operating protocol (SOP) was developed to ensure consistency among institutions. The SOP requested data from all NHPs that were born and died at the same institute going back through all historical records, along with sex, species, date of birth, date of death, and disposition (i.e., death) code and description. We received data from 27 species categories 342 at the Duke Lemur Center, but ultimately did not include these data herein because they did not meet stage 1 343 filtering requirements of this study. We also note that life history profiles for these animals are published⁴⁴ and the data are available for public download (https://lemur.duke.edu/duke-lemur-center-database/).

Data Filtering and Quality Control

Received data were first processed via a series of quality control checks for non-NHP species labels,

348 inconsistent or undefined codes, and duplicated records (e.g., ensuring one observation (date of birth and

death) per animal in data). We attempted to resolve inconsistencies or undefined codes via follow-up with the

350 original data source. Records that were unable to be resolved were removed from subsequent analyses. The

 resulting data were then parsed through a two-stage filtering process. Stage One filtering retained records with: 1) sex classified as male or female, 2) known date of birth (not estimated), and 3) survived at least 30 days (removing neonatal deaths). Species were then filtered to only include those which retained at least 150 animals. These Stage One filtered data yielded over 77,000 animals across 12 species. Stage Two filtering retained 1) animals that survived to adulthood using the National Institutes of Health Nonhuman Primate 356 Evaluation and Analysis table of NHP life stages (Table 1).⁴⁵ The earliest age listed as adult for each species 357 was used, supplemented by additional references for two species not present in the table, chimpanzees⁴⁶ and $\,$ coppery titi monkeys. 33 Stage Two filtering also implemented a date of birth (DOB) cutoff. This step was critical for survival analyses and lifespan inference as received data did not include records on alive animals. Removing later (more recent) births avoided skewing results towards earlier deaths, and inference was thus based on the dataset of animals that had greatest opportunity to live to their maximum ages (**Figure S1**). The DOB threshold was implemented by retaining animals born before 2023 minus the number of years 363 corresponding to the initial assessment of the 85th percentile of lifespan for that species (combined sexes; non-natural deaths as censored events). In total, this filtering stage yielded a dataset of 32,616 animals, across 12 species.

 Defining censored events by death types. Given that these data did not include alive animals, for survival analyses, censored events were based on death type, as follows: 1) death types pertaining to research sacrifice and colony management were categorized as right censored events; 2) death types pertaining to natural causes or humane euthanasia for health reasons were coded as un-censored events. Right censoring is a statistical approach in survival analysis that enables inclusion of the knowledge that the subject survived at 372 least to that point.¹² Treating deaths related to research sacrifice and colony management as right-censored 373 events enabled animals to contribute to the survivorship model up until age of censoring. That is, this accounts for the lack of knowledge of how long the animal would have lived until a natural or health-related death. The 375 final Stage Two filtered dataset was comprised of 12,269 events and 20,347 censored events.

Statistical analyses

 We computed the Kaplan-Meier estimator⁴⁷ of the survivorship function for each species and sex, using the 379 ggsurvfit package⁴⁸ in R version 4.1.2. Survival curves and median lifespan estimates were calculated for both including and excluding censored (research sacrifice; colony management death types) data. A critical analytic consideration was that censoring was greatly biased by sex. Thus, the primary analyses presented with comparisons by sex were limited to natural/health-related deaths only (no censored data). For many species, proportional hazards assumptions were violated (preventing usage of the cox-proportional hazards model), but since the primary analysis datasets were absent of censored events, analyses were not restricted to methods 385 for censored data. The analysis plan followed one that was applicable across all twelve species of various 386 sample sizes. For each species, maximum ages were compared between males and females using two analytic approaches. First, quantile regression models were analyzed in SAS version 9.2 using the 388 QUANTREG procedure at the 25th, 50th, 75th, and 85th maximum age percentiles with sex as the predictor and primate center was included as a covariate. Effects of sex at each percentile were tested using the Wald statistic and standard errors for regression coefficients were computed using resampling method (seed=12333). For each species, we also tested for differences in the maximum age distributions by sex using the nonparametric two-sample Kolmogorov-Smirnov test (ks.test function in R version 4.1.2), two-sided test p-393 values are reported.⁴⁷ Finally, to evaluate the uniformity of the rate of decline across survivorship curves, we fit 394 an exponential model (e^{$β$}), separately, to the first and last quartiles of the Kaplan-Meier survival curves using the nonlinear least squares function in R (version 4.1.2), shown in **Figure S3**. As β captures the function's rate 396 of decay, we illustrated trends across species, by sex, by plotting the magnitude of β for these two quartiles. Computations were performed using the Wake Forest University (WFU) High Performance Computing 398 Facility. 49

Data Availability

 Raw, de-identified data are available via the password-protected database MIDAS (Monkey Inventory and DAta management of Samples), request for access available from [https://midas.wakehealth.edu/MIDAS.](https://midas.wakehealth.edu/MIDAS) The MIDAS database will include the same information provided in the manuscript and extended tables, and

- provide tools for species comparisons, which will make this a user-friendly resource accessible to researchers.
- Data sharing will be limited to scientific uses.
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Code Availability

- Analyses and summaries were computed using functions and libraries, as described in methods, in
- accordance with standard practices and their vignettes. Custom Code for fitting exponential curves to survival
- data is available in Supplementary Information and is available via MIDAS as described in Data Availability.
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Comparative lifespan and healthspan of nonhuman primate species common to biomedical research.

Primary Display Items and Extended Data Figures

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Primary Figures

Figure 1. Scatter plot of data points for natural and health-related euthanasia deaths by species.

Boxplot overlay depicts median and interquartile range by species and sex. Proportion of data by sex and species also shown. The vertical dashed line denotes equal counts of males and females by species.

Figure 2. Combined survival curves for females (A) and males (B) of all 12 species.

Data shown are for animals with deaths resulting from natural causes or humane euthanasia for health-related reasons.

Figure 3. Comparison of rate of survivorship decline by quartile and sex.

Rates of decline were calculated from fitting an exponential model to the first and last quartiles of the sex-specific Kaplan-Meier survival curves. Males and females are compared by quartile. Rate of decline was generally faster in males within the first quartile with the pattern nearly reversed by sex in the last quartile.

Figure 4. Kaplan-Meier survival curves by sex and species for natural deaths or humane euthanasia for health-related reasons.

For each plot, the X-axis scaling (maximum age) is species-specific.

Primary Tables

Table 1. Sample sizes of primary analysis datasets and species-specific age categories*.*

For each species, age categories and estimated age ranges are shown. 33,44,45

**Natural or Health-related deaths only*

Table 2. Maximum and median age at death by sex and species

**Median age at death is calculated from natural and clinical deaths only; maximum observed age includes animals with any type of death. Maximum ages are from the current dataset only; there are known older animals of some of these species at research institutes, such as a 29-year-old titi monkey male at CNPRC and two 19-year-old male marmosets at SNPRC.*

Table 3. Sex-based comparisons of age by species.

Quantile regression for 25^{th} , 50^{th} , 75^{th} , and 85^{th} percentiles. Regression models adjusted for primate location (data source). Distribution of ages by sex were assessed using the Kolmogorov Smirnov test. Complete data used for analyses (natural or clinical deaths) with no censoring.

Extended Data Figures (Active Links within Article)

Extended Data Figure 1: Distribution and proportions of sex and censored data points for filtered data (n=32,616).

Deaths due to natural causes or health-related reasons are labeled as 'natural.' Deaths due to research sacrifice or colony management are labeled as 'censored.' Multiple species had very high proportions of censored events.

Extended Data Figure 2: Kaplan-Meier survival curves for 12 species, by sex, including censored events.

Graphs of Kaplan-Meier estimates of the survivorship function for all data passing quality control filtering are provided. Presented are survival curves where deaths from research sacrifice and colony management are included as censored events; censoring was biased by sex, so sex-specific comparisons were not computed for these data and were instead limited to data containing natural deaths, only. For many species (e.g., baboons), the high proportion of censored events yielded unstable survival probability estimates for latter ages. For each plot, the X-axis scaling (maximum age) is species-specific.

Extended Data Figure 3: Phylogenetic tree of 12 species analyzed in study.

This tree was generated with the 10kTrees Project and modified to match taxonomic names with those used in our study and to simplify the presentation.³² Only the 12 species studied herein are represented in the tree; there are many other species of primates in these clades not pictured.

