

Extensive longevity and DNA virus-driven adaptation in nearctic *Myotis* bats

Juan M Vazquez^{1,18}, M. Elise Lauterbur^{2,3,18}, Saba Mottaghinia⁴, Melanie Bucci², Devaughn Fraser^{5,6}, Genavieve Gray-Sandoval², Léa Gaucherand⁷, Zeinab R Haidar^{8,9}, Melissa Han¹⁰, William Kohler¹⁰, Tanya M. Lama¹¹, Amandine Le Corff⁴, Clara Loyer⁴, Sarah Maesen⁴, Dakota McMillan^{1,12}, Stacy Li^{1,13}, Johnathan Lo¹³, Carine Rey⁴, Samantha LR Capel⁶, Michael Singer¹⁴, Kathleen Slocum¹⁵, William Thomas¹⁶, Janet Debelak Tyburec¹⁷, Sarah Villa¹⁴, Richard Miller¹⁰, Michael Buchalski⁵, Jose Pablo Vazquez-Medina¹, Sébastien Pfeffer⁷, Lucie Etienne^{4,19*}, David Enard^{2,19,20**}, Peter H Sudmant^{1,13,19,20,21***}

Affiliations:

1. Department of Integrative Biology, University of California, Berkeley, Berkeley, CA USA
2. Department of Ecology and Evolutionary Biology, University of Arizona, Tucson, AZ USA
3. Current affiliation: Department of Biology, University of Vermont, Burlington, VT USA
4. Centre International de Recherche en Infectiologie (CIRI), Inserm U1111, UCBL1, CNRS UMR5308, Ecole Normale Supérieure ENS de Lyon, Université de Lyon, Lyon, France
5. Wildlife Genetics Research Unit, Wildlife Health Laboratory, California Department of Fish and Wildlife, Sacramento, CA, United States
6. Current affiliation: Wildlife Diversity Program, Wildlife Division, Connecticut Department of Energy and Environmental Protection, Burlington, CT, United States
7. Université de Strasbourg, Architecture et Réactivité de l'ARN, Institut de Biologie Moléculaire et Cellulaire du CNRS, Strasbourg, France.
8. Department of Biology, California State Polytechnic University, Humboldt, Arcata, CA USA
9. Current affiliation: Western EcoSystems Technology Inc, Cheyenne, WY USA
10. Department of Pathology and Clinical Laboratories, University of Michigan, Ann Arbor, MI USA
11. Department of Biological Sciences, Smith College, Northampton, MA USA
12. Department of Science and Biotechnology, Berkeley City College, Berkeley, CA USA
13. Center for Computational Biology, University of California, Berkeley, Berkeley, CA USA
14. Department of Molecular and Cellular Biology, University of California, Berkeley, Berkeley, CA USA
15. Bat Conservation International, Austin, TX USA
16. Department of Ecology and Evolution, Stony Brook University, Stony Brook NY USA
17. Bat Survey Solutions, LLC, Tucson, AZ USA
18. These authors contributed equally
19. Senior author
20. These authors contributed equally
21. Lead contact

* Correspondence: lucie.etienne@ens-lyon.fr

** Correspondence: denard@arizona.edu
 *** Correspondence: psudmant@berkeley.edu

Abstract

The genus *Myotis* is one of the largest clades of bats, and exhibits some of the most extreme variation in lifespans among mammals alongside unique adaptations to viral tolerance and immune defense. To study the evolution of longevity-associated traits and infectious disease, we generated near-complete genome assemblies and cell lines for 8 closely related species of *Myotis*. Using genome-wide screens of positive selection, analyses of structural variation, and functional experiments in primary cell lines, we identify new patterns of adaptation contributing to longevity, cancer resistance, and viral interactions in bats. We find that *Myotis* bats have some of the most significant variation in cancer risk across mammals and demonstrate a unique DNA damage response in primary cells of the long-lived *M. lucifugus*. We also find evidence of abundant adaptation in response to DNA viruses - but not RNA viruses - in *Myotis* and other bats in sharp contrast with other mammals, potentially contributing to the role of bats as reservoirs of zoonoses. Together, our results demonstrate how genomics and primary cells derived from diverse taxa uncover the molecular bases of extreme adaptations in non-model organisms.

Keywords

Aging, Bats, Cancer, Evolutionary Biology, Functional Genomics, Immunity, Infectious Disease

Introduction

Bats (order *Chiroptera*) represent approximately 20% of all known mammalian species and are one of the most phenotypically diverse clades of mammals^{1,2}. Since their emergence 60 million years ago³⁻⁵, many bat lineages have independently evolved a wide variety of life history strategies and phenotypic traits, including exceptional changes in longevity, viral tolerance, and immune defense⁶⁻¹¹. Such systems, in which shared traits have evolved *de novo* multiple times, are powerful resources for dissecting the genetic basis of phenotypes. Rigorous approaches to studying these traits, however, depend on high-quality, well-annotated genomes to test evolutionary and genomic hypotheses, and on experimental functional systems to validate these hypotheses.

The largest genus of bats - *Myotis* - is estimated to have emerged approximately 33 million years ago^{12,13}, and encompasses over 139 described species spanning six continents and a wide range of ecological niches^{1,12-14}. *Myotis* species demonstrate some of the most extreme variation in lifespan amongst mammals^{6,15-18}, including a six-fold difference in lifespan between the longest-lived species (*M. brandtii*, 42 years^{15,19}, **Figure 1A**) and the shortest-lived species (*M. nigricans*, 7 yrs^{15,20}) which diverged approximately 10.6 million years ago^{5,14,21,22}. In addition, *Myotis* species have been used as systems for investigating virus tolerance and other pathogen resistance²³⁻²⁵ associated with the expansion and

contraction of antiviral defenses^{26–29}, which have contributed to bats' ecological role as zoonotic reservoirs^{10,11,30–33}.

The origin, evolution, and functional basis of these phenotypes can be studied experimentally in model organisms as well as via comparative evolutionary methods. The power of comparative evolutionary studies is constrained by several factors including incomplete phylogenetic coverage; poor temporal resolution; the quality and composition of gene annotations; and availability of functional data and tools for validation. Rapidly evolving genes, such as those associated with adaptations to pathogens^{34–36}, present particular challenges for homology and alignment based methods. Similarly, poor phenotypic resolution and long divergence times between study species hinders the power of statistical approaches to identify patterns of selection and diversification^{37–40}. Meanwhile, model organism-based approaches contribute a different, complementary perspective and provide the power of functional analyses; however, these studies can suffer from issues related to the suitability and diversity of the model species' genotype and phenotype.

While studies on the genetic basis of longevity in short-lived model organisms have been crucial for identifying and dissecting several key aging pathways, comparative studies of exceptionally long-lived organisms have uncovered novel genes and pleiotropic effects governing lifespan^{36,41–53}. The comparative approach, however, has historically been hindered by limitations in available genomic resources and genetic tools for study. Similarly, studies of infectious disease response are common and powerful in model organisms, but the lack of diversity and inbred lines limits their scope. Bats in particular present an important case study in, and opportunity to study, variation in virus adaptation strategies due to bats' role as zoonotic reservoirs and their specific resistance to viruses^{36,54}. While previous studies have shown unique infectious disease adaptations in bats, including loss of important inflammatory genes and expansions of and adaptation in some immune gene families^{54–56}, they are typically hampered by the breadth and number of species analyzed, and only rarely functionally validate results from genomic analyses.

Here we combined comparative and functional approaches in *Myotis* to uncover strong genomic and functional evidence of adaptation to both aging-related and infectious diseases. We present for the first time a robust quantification of relative intrinsic cancer risk across mammals, finding that *Myotis* are overrepresented at the extreme of increased cancer risk. Consistent with this observation, we identified pervasive selection of genes in longevity- and cancer-related processes, especially in lineages which have undergone the greatest changes in lifespan. Furthermore, we found strong evidence of adaptation in response to DNA viruses in *Myotis* and other bats. Genome-wide enrichment of adaptation being driven by DNA viruses is unique to bats in comparison with other large groups of mammals. Finally, using near-complete assemblies, we identified structural variations encompassing stress response, immunity, and inflammation genes, including a trans-species copy number polymorphism of protein kinase R (PKR). Together, our results suggest that pleiotropy and co-evolution of traits in *Myotis* has played a key role in the evolution of exceptional longevity and infectious disease resistance.

Results

High quality chromosome-level assemblies of 8 *Myotis* bat species

To study how lifespan and viral response have evolved in *Myotis*, we collected skin punches and derived primary cell lines from several North American (“Nearctic”)²¹ species (**Figure 1A,C**), including from one of the longest-lived mammals, *Myotis lucifugus*¹⁵. Using these cell lines and flash frozen tissues we generated *de novo* haplotype-resolved, chromosome-scale genome assemblies for eight species (**Figure 1A**) using a combination of long-read PacBio HiFi sequencing and HiC scaffolding. These genomes are highly contiguous, with an average of 98.6% of nucleotides assembled into 22-23 syntenic chromosome-scale scaffolds corresponding to the published karyotype⁵⁷ with an average QV of 66. These genomes have among the lowest auNG scores of any *Chiroptera* genome published to date (**Figure 1A, E; Table S1**). Across all 8 genomes, each autosome has been completely assembled telomere-to-telomere (T2T) in at least one species (**Figure 1E**). Within assemblies, 29%-70% of chromosomes are fully assembled with an average of less than one gap per chromosome (**Table S1**). When comparing the assemblies of species generated from tissue samples versus primary cell lines, we found that they were broadly comparable and structurally similar. However, genomes assembled from cell lines had slightly improved statistics likely attributable to the increased quality and molecular weight of extracted DNA (**Figure 1A, D, E; Table S1**).

Genomes were annotated using well-established pipelines³⁶ leveraging multiple lines of evidence, including short-read RNAseq, gene prediction (AUGUSTUS-CGP⁵⁸, GeneMark-ES⁵⁹; gene projections⁶⁰, TOGA⁶¹); and homology (miniprot⁶²). In total, we identified an average of 27,536 protein coding genes per species. We benchmarked our annotations using BUSCO^{63,64} (V5.4.3) mammalian ortholog sets indicating these annotations are 98.2%-98.5% complete (**Figure 1C**). We also annotated a recent assembly of *Myotis yumanensis*⁶⁵ for inclusion in downstream analyses. Overall, these fully annotated genomes represent some of the most contiguous mammalian assemblies to date.

Resolving the phylogeny and the evolution of body size and lifespan in nearctic *Myotis*

The phylogenetic relationships within *Myotis* have been the subject of much debate, with a number of conflicting phylogenies described in the literature based on different choices of genetic markers^{14,66–69}. To resolve the phylogeny of Nearctic *Myotis*, we identified single copy orthologs of 17,509 protein genes present in 536 mammalian genomes resulting in 30.6M aligned nucleotides. These alignments were used to build a maximum likelihood tree of *Eutheria*. The *Chiroptera* sub-clade was then time-calibrated using available fossil-based node calibrations (**Figure 1B; Figure S1; Table S2**). Our results conclusively recapitulate known sister species pairs including *M. lucifugus* and *M. occultus*; *M. yumanensis* and *M. velifer*; and *M. evotis* and *M. thysanodes*. Our proposed phylogeny resolves the complex relationship between these sister taxa, with 100% bootstrap support at all nodes throughout *Chiroptera*.

Using our resolved Nearctic *Myotis* phylogeny, we re-examined the evolution of body size and lifespan in *Chiroptera*. In mammals and other metazoans, there is a strong allometric scaling (positive correlation with body size) of lifespan. Bats have been noted as an exception to this rule: they are exceptionally long-lived for their body size^{17,18,70}, and this exceptional longevity has evolved *de novo* multiple times^{6,70,71}. However, these observations have not been tested using phylogenetically corrected statistics leveraging well-resolved phylogenies. To test the hypothesis of non-allometric scaling of lifespan in bats, we modeled the evolution of body size and lifespan across a supertree of over 1000 placental mammals (*Eutheria*)⁶⁷ (**Figure 2; Table S2**). In agreement with previous studies in vertebrates^{7,17,44,50,72–80}, changes in body size are pervasive across mammals, with extreme changes seen in whales (*Cetacea*)^{78,79}, elephantids (*Proboscidea*)^{42,44,72}, and in sloths and armadillos (*Xenarthra*)^{73,80–82} (**Figure 2A; Table S2**). Within bats, major changes in body size are only observed at the root of the lineage and within *Yinpterochiroptera* (megabats including genera *Pteropus*, *Eidolon*, *Megaderma*, and *Rhinolophus*). Outside of these clades, only minor changes in body size were observed (**Figure 2A**). The evolution of lifespan across mammals mirrors the evolution of body size; branches with large increases in body size (e.g. *Cetacea* ancestor, *Primate* ancestor) have also experienced large increases in lifespan (**Figure 2B**), leading to an overall positive association between lifespan and body size (**Figure S2A**). However, despite little change in body size in bats (**Figure 2A, C**), we observed some of the largest changes in lifespan across mammals towards the tips of the tree (**Figure 2B, D**), consistent with the theory of multiple independent increases in lifespan across bats. This is especially true in *Myotis*, where we saw many of the fastest increases in lifespan, including for *Myotis grisescens* (4.15x increase, 100th percentile), *Myotis brandtii* (2.25x increase, 100th percentile), *Myotis lucifugus* (1.56x, 98th percentile), *Myotis myotis* (1.1x increase, 79nd percentile), and the *Myotis* common ancestor (1.26x increase, 92rd percentile) (**Figure 2D; Figure S2C; Table S2**). We next used phylogenetically-corrected generalized linear models and ANCOVA to study the relationship between body size and lifespan across mammals. While we find that non-bat mammals experience a 0.159% increase in lifespan per 1% increase in body size on average, bats experience a 0.223% increase in lifespan years per 1% increase in body size; these rates were not significantly different, however, suggesting that lifespan allometry is conserved in bats after accounting for phylogeny (**Figure S2E-F**; pANCOVA, p=0.29).

Rapid changes in body size and lifespan can have major implications for the evolution of cancer risk and resistance across mammals. The lifetime cancer risk of an individual is modeled as the product of body size (i.e. the number of cells within an individual), lifespan, and a constant representing the intrinsic cancer risk per cell per unit time. Within species, lifetime cancer risk scales linearly with body size, and with lifespan by a power-law of exponent 6^{83–86}. In contrast to this *within*-species relationship, there is no significant correlation between body size, lifespan, and cancer risk *across* species^{86–89} - a phenomenon known as Peto's Paradox. The observation of similar lifetime cancer incidence rates across mammals^{73,89,90} suggests that species with more cells or longer lifespans have adapted to reduce their cancer risks (i.e. increased cancer resistance) (**Figure 2E**).

We hypothesized that the very rapid evolution of increased lifespan in *Myotis* would thus result in a dramatic increase in their expected cancer risk compared to other mammals. This can be quantified by the Reduced Intrinsic Cancer Risk per cell (RICR) between an extant mammal and its most recent ancestor, calculated as the log₂ ratio of body size and lifespan between the two nodes (**Figure 2E**)^{44,86}. Decreases in RICR correspond to an increase in the expected cancer risk. We used estimates of body size and lifespan across *Eutheria* to quantify changes in (RICR) across placental mammals (**Figure 2F**)⁴⁴.

Bats overall were slightly overrepresented in the bottom 10% of RICR with an odds ratio of 1.15, highlighting the impact of rapid lifespan evolution on cancer risk. The longest-lived *Myotis* (*M. grisescens*, 39 yrs & 1st pct; *M. brandtii*, 42 yrs & 2nd percentile; *M. lucifugus*, 36 yrs & 4th pct) and their most recent common ancestors (*lucifugus-occultus*, ~26 yrs & 8th pct; *Myotis* common ancestor, ~22 yrs & 14th pct) demonstrated some of the most pronounced decreases in RICR among mammals (**Figure 2F**; **Figure S2D**; **Table S2**). Similar to other extreme cases of body size and lifespan in vertebrates^{44,46,50,52,73,91–94}, the pronounced changes in RICR seen in *Myotis* imply an extraordinarily strong selective pressure to evolve cancer resistance mechanisms at multiple points across *Chiroptera* in general, and within *Myotis* in particular.

Evolutionary signatures of cancer resistance in *Myotis*

We next set out to identify genes under positive selection across our phylogeny of Nearctic *Myotis*. We used aBSREL⁹⁵ to test for branch-specific positive selection among 15,734 single-copy orthologous genes identified in 536 mammalian genomes. We found that on average, 22.7% of genes were under selection across the 9 nearctic *Myotis* species and their internal branches after multiple testing correction at FDR≤5%; and 5.23% of genes were significant and had omega values above 1, signaling positive selection (**Table S3**). These genes were enriched for several pathways involved in immunity, cancer, and aging (**Table S3**). Many of these genes lie at the intersection of these two processes, including members of the Cluster of Differentiation (CD) family, Serpin family, insulin signaling pathway, redox repair, and iron storage (**Figure 3A**; **Table S3**), suggesting possible pleiotropic influences on genes under selection.

To test this, we quantified the contribution of genes under selection to pathways associated with the hallmark of cancer^{96–98} by measuring the proportion of cancer-associated pathways overrepresented among genes under selection throughout the phylogeny (**Figure 3A**; **insets**). Many nodes within nearctic *Myotis* were enriched for cancer hallmark pathways, especially at the recent ancestors of the longest-lived species (e.g. *M. lucifugus*, *M. occultus*; **Figure 3A**). Testing the overall contribution of genes that have undergone selection in each species since the common *Myotis* ancestor, we observed significant enrichments in the representation of cancer-associated pathways only in species lineages with reductions in RICR (*M. lucifugus*, *M. occultus*, *M. evotis*, *M. thysanodes*, *M. yumanensis*; **Figure 3B**). This suggests that while genes under selection in nearctic *Myotis* frequently contribute to cancer-associated pathways, cancer resistance has only driven consistent selection in the longest-lived lineages with the greatest increases in cancer risk.

We also observed that many key genes involved in ferroptosis - specifically in iron transport, glutathione metabolism, and lipid peroxidation - were under both positive and negative selection at multiple instances throughout the phylogeny (**Table S3**). Many of these genes were recurrently under selection in each species' lineage, such as with ferritin (both heavy and light chains) at three distinct points in the evolutionary history of *M. yumanensis*. Genes under selection in iron transport are specifically involved in the regulation of free iron in the cell, specifically in the export and reduction of the free radical catalyst Fe²⁺ (ferroportin, *HMOX1*) and the import, storage, and maintenance of Fe³⁺ (ferritin and transferrin receptors 1 and 2). Additionally, we observe selective signatures in glutathione metabolism and oxidative stress response including: *SLC3A2* and *SLC7A11*, a heterodimer pair facilitating cystine import and glutamate export; glutathione synthetase; and glutathione peroxidase 3

(*GPX3*). Finally, we observed a pattern of selection in genes involved in synthesizing and maintaining key polyunsaturated fatty acids involved in ferroptosis, including *LPCAT3*, *ALOX15*, and *PRDX5*.

To test for intensified and relaxed selection in genes in long-lived or short-lived *Myotis*, we ran RELAX⁹⁹ on 12,438 genes present across 11 *Myotis* species, identifying 263 genes under intensified selection ($k > 1$) and 101 genes under relaxed selection ($k \leq 1$) after multiple testing correction ($p_{adj} \leq 0.05$). Among genes of note showing significant intensified selection were *USP9X* (an X-linked ubiquitin protease associated with cancer and T cell development^{100,101}, $k=48.6$); *CDK16* (an oncogenic cyclin-dependent kinase that regulates autophagy^{102,103}, $k=44.9$); and *FGFR2* (a cell growth receptor associated with human cancers that is also a viral interacting protein^{104,105}, $k=26.1$) (**Figure S3B; Table S4**). Performing a gene set enrichment analysis for the 364 significant genes, we find a strong association among genes under intensified selection with FGF2 signaling, chromatin remodeling, and pathways associated with both retroviruses and coronaviruses (**Figure S3C; Table S4**). Finally, using RERConverge¹⁰⁶, we investigated how genes' evolutionary rates correlated with the evolution of body size, lifespan, or the first two principal components of body size and lifespan across *Myotis*, and found a number of genes enriched in pathways associated with innate immunity, gamete production, and various metabolic processes, consistent with our other results (**Figure S3D-E; Table S4**).

The longest-lived bat in our study, *M. lucifugus*, had an overrepresentation of pathways specifically associated with DNA double-strand break (DSB) repair when looking at both lineage-wide and node-specific enrichments in positive selection using the Reactome database¹⁰⁷ (**Figure 3C; Table S3**). This includes 35 out of 65 genes in the high-fidelity Homologous Recombination Repair pathway, and 21/37 members of the Homology-Directed Repair via Single Strand Annealing (**Figure 3C; Table S3**). These results suggest that *M. lucifugus* might have an enhanced response to DNA DSBs relative to other bats. To test this hypothesis, we assessed the tolerance of *M. lucifugus* to neocarzinostatin, a potent radiomimetic agent that induces DNA double-strand breaks (**Figure 3D**), compared to *M. evotis*, three non-*Myotis* bats (*Eidolon helvum*, *Pteropus rodrigensis*, and *Rousettus lanosus*), and humans. At low doses of neocarzinostatin, *M. lucifugus* was the only species tested showing sensitivity to neocarzinostatin after 24 hours, with a drop in viability and concomitant increase in apoptosis. At high doses, *M. lucifugus* had the highest level of apoptosis and the greatest drop in viability of all the bats tested, although all bats were more resistant to DNA damage than humans. This is consistent with other long-lived species, including elephants^{42,43,90}, naked mole rats⁵¹, and bowhead whales^{46,108}, where longevity and RICR are associated with an increased ability to clear out damaged cells. Together, these results support the hypothesis that *M. lucifugus* has evolved an enhanced DNA double-strand break response as predicted by genes exhibiting signatures of positive selection in this species.

Adaptation to DNA viruses

Amongst genes under selection, a substantial portion were involved with immunity, including members of the immunoglobulin and Cluster of Differentiation gene families. These genes exhibited some of the highest evolutionary rates (ω) in our dataset, suggesting that they are under strong selection in *Myotis* (**Table S3; Table S4**). Because immune pathways are only one aspect of host viral adaptation¹⁰⁹, we tested for adaptive signatures in virus-interacting proteins (VIPs) in *Myotis* and other bats. VIPs are host proteins that physically interact with viral proteins (e.g. *CD45*, **Figure 4A**), and can be proviral (contributing to viral

infection, e.g. viral receptors), antiviral (protective against viral infection, e.g. interferons), or both depending on infection stage and virus type. Previous studies investigating positive selection across mammals have found an enrichment for adaptation among a set of 5,528 manually curated VIPs, defined as host proteins that have at least one experimentally verified physical interaction with a viral protein, RNA, or DNA¹⁰⁹.

By calculating an enrichment score from the ratio of positive selection in VIPs compared to their matched control genes using BUSTED-MH¹¹⁰, we found that, like other mammals, *Myotis* show an enrichment for adaptation at VIPs (**Figure 4B; Table S5**). Physical host-virus interactions may not always result in fitness effects in the host. We therefore repeated our analysis using a gene set restricted to VIPs with experimental evidence of specific pro- or anti-viral effects, and thus with a stronger expectation of fitness effects. We observed an even stronger significant elevation in the ratio of positive selection in these proviral and antiviral VIPs (**Figure 4C; Table S5**), but no elevation in this ratio in other VIPs (**Figure 4D; Table S5**). This is consistent with the expectation of viral interaction as the cause of enrichment of positive selection in VIPs in bats¹¹¹. We repeated this analysis using a dataset of 47 publicly-available non-*Myotis* bat genomes, and confirmed these same patterns across bats more broadly, even when excluding *Myotis* genomes (**Figure 4B inset**).

Previous work has suggested that bats may have different physiological responses to DNA and RNA viruses¹¹². To determine if this was reflected in genomic VIP adaptation, we compared the enrichment of positive selection in VIPs that interact only with DNA viruses (DNA VIPs) to those that interact only with RNA viruses (RNA VIPs). Remarkably, we found that VIP adaptation in *Myotis* and other bats is driven by selection in DNA VIPs (**Figure 4E and inset**). This is in marked contrast to the observed pattern in RNA VIPs, which show no evidence of enrichment in adaptation (**Figure 4F and inset**). Note that this difference between DNA and RNA VIPs cannot be explained by a difference in the conservation of VIP status between the two. The vast majority of VIPs were discovered between human proteins and viruses that infect humans¹¹¹, and a concern could then be that those proteins that are RNA VIPs in humans have evolved faster than DNA VIPs in bats, ultimately resulting in the more frequent loss of their VIP status in bats. We can however exclude this possibility, since DNA and RNA VIPs have very similar average dN/dS ratios (*Myotis*, 0.2 vs. 0.18 respectively; non-*Myotis* bats, 0.163 vs. 0.153 respectively).

In contrast to what we observe in bats, VIP adaptation in humans is driven by positive selection in RNA - and not DNA - VIPs^{109,113}. To investigate if DNA VIP-driven adaptation in bats is exceptional among mammals, we replicated these analyses across four other large mammalian orders that are well represented among publicly-available mammalian genomes: *Primates*, *Glires*, *Eeungulata*, and *Carnivora*. We found that while other mammalian orders show a mix of adaptation enrichments in both RNA and DNA VIPs, none show an absence of genome-wide enrichment of adaptation in RNA VIPs as observed in bats (**Figure S4**). These results highlight that bats, including *Myotis*, may have faced greater selective pressures from DNA viruses than from RNA viruses, in contrast to other mammals.

Evolution of structural variation within constrained karyotypes

With only six known exceptions, all *Myotis* species with cytological data have a conserved karyotype (60+ *Myotis* spp.: 2n = 44^{114–118}; *M. annectans*: 2n = 46¹¹⁶; *M. laniger*: 2n = 48¹¹⁷; *M. bechsteinii*:

2n = 42¹¹⁹; *M. daubentoni*: 2n = 42¹²⁰; *M. davidii*: 2n = 46¹²¹; *M. macrodactylus*: 2n = 44/45^{122,123}). This conserved *Myotis* karyotype, shared among species spread across six continents^{1,2}, consists of three large autosomes and one small metacentric autosome; 17 small telocentric autosomes; and metacentric X and Y chromosomes^{57,124}. Consistent with this broad cytological conservation, we find large scale synteny across the Nearctic *Myotis* in this study. However, structural variants (SVs) including inversions, duplications, and translocations are relatively common within chromosomes, especially in putative centromeric regions (**Figure 5A, B**).

We used SyRI¹²⁵ to identify SVs across pairwise alignments of Nearctic *Myotis* genomes relative to the outgroup *M. myotis* and identified 6,813 - 8,013 SVs per genome. Most of these events were small, with 97 - 99% of events under 10Kb. In the three large autosomes, which constitute ~30% of each genome, we cataloged an average of 509 SVs (**Table S6**). In contrast, in the small autosomes, constituting ~65% of each genome, we observed an average of 316 events, highlighting the distinct structural evolution between these chromosome types (**Table S6**). However, large (≥10Kb) duplications, large inverted duplications, and large inverted translocations were more common on small autosomes compared to the large autosomes (**Table S6**).

We also quantified the distribution of transposable elements (TEs) across chromosomes. Surprisingly, LINE elements were significantly enriched around the centromeres of all chromosomes, both metacentric and telocentric (**Figure 5B**); while this is rare in mammals, it has been recently described as a feature of Phyllostomid genomes¹²⁶. In many cases, particularly in the 3 large metacentric chromosomes, LINE elements appear to have displaced other TEs. Rolling circle and SINE elements were particularly depleted concomitant with LINE enrichment. In contrast, SINE elements were enriched at telomeres. The concentration of segmental duplications is significantly correlated with TE density in each species (linear regression, $p < 0.01$; **Figure 5B**; **Figure S5J**) highlighting the possible importance of TEs in facilitating structural evolution.

One particularly striking example of structural evolution we identified is a ~20-Mb block at the subtelomeric end of chromosome V15 undergoing frequent and recurrent inversions and translocations in nearctic *Myotis* (**Figure 5A**). This region spans several immune-related genes including multiple members of interleukin signaling pathways, including IL-1 and IL-36. A 10Mb portion of this block was recently identified as a potential target of recent selection by adaptive introgression⁶⁹. We identified between 2-3 major (8+ kb) blocks in this region exhibiting inversions between Nearctic *Myotis*, which correspond to similarly sized regions in the outgroup *M. myotis* (**Figure 5A**; **Table S5**). Additionally, we noted a depletion of DNA transposable elements at the boundaries of each inversion (**Figure 5B**), particularly for rolling circle (RC) and SINE elements. Both of these elements can catalyze large-scale structural rearrangements via DNA damage repair and homologous recombination, respectively¹²⁷⁻¹³¹.

Gene duplications and losses can be drivers of evolution via dosage modification^{132,133}, sub- and neofunctionalization^{134,135}, regulatory network remodeling¹³⁶, and other processes¹³². We quantified gene gains and losses across *Myotis* relative to their single-copy human orthologs. Using CAFE¹³⁷, we found 38 gene families underwent significant expansions or contractions in at least one nearctic *Myotis* species (**Figure 5C**). However, gain and loss rates varied substantially across branches of the *Myotis* phylogeny. The terminal *M. auricolus* and *M. velifer* branches had ~4-fold more significant gene family expansions (37 and 35 families, respectively; **Figure 5C**) than other *Myotis* branches. In contrast, the terminal *M.*

californicus and *M. yumanensis* branches had ~2-fold more significant contractions (24 and 23 families, respectively; **Figure 5C**) than other *Myotis* branches. We observe significant overrepresentation of pathways at FDR≤10% in only 4 gene sets: gene families that underwent significant expansions in *M. auriculus*, *M. velifer*, and *M. volans*; and genes that underwent significant contractions in *M. lucifugus* (**Figure S5A-H**). Many of these pathways were shared between all sets, including pathways involved in translation regulation; ROBO receptors and neuronal development; selenoprotein and selenocystine metabolism; and influenza life cycle (**Figure S5A-H**).

Given that many of the genes in these pathways are VIPs, we used the method of Huang et al (2023)⁴⁸ to test if VIP genes in particular underwent significant copy number changes relative to non-VIP genes. We found that while the birth-death rate of VIP genes is similar to that of other genes ($p = 0.071$), together VIP genes are significantly more likely to have undergone expansions and/or contractions on at least one branch of the *Myotis* family ($p < 0.001$; **Figure S5I-J**). This suggests that there is variation in gene family birth rates across species, but that VIPs are more dynamic across the Nearctic *Myotis* as a whole than other types of genes.

To further explore the functional impact of gene duplications we ranked genes by their maximum copy number across all genomes. We found that the gene families with the highest copy numbers were concentrated in pathways associated with cancer, aging, immunity, and olfaction (**Figure 5D**). One striking case is *FBXO31*, with ~2.4x more copies on average than the next most duplicated gene in *Myotis* (20-48 copies). *FBXO31* is a SCF (SKP1-cullin-F-box) protein ligase involved in cell cycle regulation and DNA damage response, consisting of two functional domains: a F-Box domain and a CDK binding domain¹³⁸, and has previously been speculated as a driver of longevity in *Myotis*⁹³. Quantifying *FBXO31* copy number across over 500 mammals using reciprocal best-hit BLAT, we found that this gene was more highly duplicated in *Myotis* than in any other mammal genome (**Figure 5E**). Furthermore, while there were additional partial matches of non-canonical copies of *FBXO31* in non-*Myotis* species, all copies identified in *Myotis* are full-length genes with functional domains. To model the evolution of gene copy number, we used GeneRax¹³⁹ to reconcile the gene tree and species tree. GeneRax infers a gene family tree under scenarios of gene duplication and loss, taking into account the species tree. We found support for an original 14 duplications in the common ancestor of Nearctic *Myotis*, with subsequent gains and losses in each lineage (**Figure 5F**). These results highlight a massively expanded gene family in *Myotis* with potential consequences for the regulation of stress response and other processes.

An actively segregating, trans-species copy number polymorphism of the antiviral factor Protein Kinase R, PKR

Our highly contiguous genome assemblies provide a unique opportunity to understand the evolutionary and functional dynamics of structural variation in adaptation. To illustrate this, we explored the antiviral innate immune Protein Kinase R (*PKR/EIF2AK2*), an interferon-stimulated gene with adaptive duplications unique to *Myotis*²⁸. Among our Nearctic *Myotis* genome assemblies, we resolved the structure of the two known structural haplotypes: H1, containing a single copy of *PKR* (*PKR2*); and H2, containing two tandemly duplicated copies of *PKR* (*PKR1* and *PKR2*; **Figure 6A**). We also identified a third haplotype - H3 - with three tandem duplicates of *PKR* (*PKR1*, *PKR2*, and a third copy). While 7 out of 9 *Myotis* species carried duplicated haplotypes (H2 in 6 species, H3 in *M. californicus*), to our surprise,

5 of these cases were heterozygous for the duplicated haplotype: (i.e. H1/H2 or H1/H3; **Figure 6B**). Furthermore, two *Myotis* individuals (*lucifugus* and *evotis*) only encoded for *PKR1* (i.e. H1/H1; **Figure 6B**). To determine the evolutionary history of the duplicates, we used GeneRax¹³⁹ to construct a tree from alignments of all *PKR* gene copies across Nearctic *Myotis*, using *Pipistrellus pygmaeus* as a non-*Myotis* outgroup (**Figure 6C**). Our results suggest that *PKR2* is the ancestral copy of *PKR*, and that *PKR1* originated from a single duplication event at the root of *Myotis*. Intriguingly, we observed that in the heterozygous species, both *PKR1* and *PKR2* on the duplicated haplotype clustered with other duplicated haplotypes, resulting in species tree violations for the ancestral copy, *PKR2* (**Figure 6C**). These results highlight that both the duplicated and unduplicated haplotypes have likely been segregating for over 30 million years, representing an ancient trans-species polymorphism.

PKR is a stress response and innate immune factor that interacts with viral or inverted Alu repeats dsRNAs via its dsRNA binding motifs (dsRBMs), leading to *PKR* auto-phosphorylation and dimerization^{140,141}. Upon activation, *PKR* can then phosphorylate various molecules leading to protein translation shutdown and restriction of viral replication^{140,141}. While the independent functional impacts of *PKR1* and *PKR2* were previously investigated²⁸, the effects of co-expressing both copies remain unknown. This is important because their final effects may be additive, synergistic or dominant negative, providing clues into why the *PKR* duplication is polymorphic both within and between *Myotis* species. We therefore investigated the functional impact of the duplicates' co-expression on steady state protein levels, homo/hetero-dimer formation, cell viability, protein translation shutdown and antiviral restriction (**Figure 6D-G**). We used *PKR*-KO Hela cells transfected with either *Myotis myotis* or *Myotis velifer* *PKR1*, *PKR2*, and *PKR1+2*. We found that the coexpression of *Myotis* Flag-*PKR1* and Flag-*PKR2* did not affect their protein expression levels (**Figure S6A**). Interestingly, co-immunoprecipitation (coIP) experiments show that *Myotis myotis* *PKR1* and *PKR2* do not interact (i.e. no heterodimers), even though *Myotis myotis* *PKR1* can dimerize (**Figure 6D, Figure S6B**). Furthermore, coexpression of *PKR1* and *PKR2* led to a simple additive effect in their translation shutdown activity (**Figure 6E**), suggesting that neither copy is dominant negative. Using non-toxic doses of *Myotis* *PKRs* in the context of VSV-GFP (Vesicular stomatitis virus encoding a GFP reporter¹⁴²) infections, we found that, although *PKR1* and *PKR2* are both antiviral²⁸, the coexpression of *PKR1* and *PKR2* is not beneficial against VSV (**Figure 6F**). Similar results were found with an unrelated virus, SINV-GFP (Sindbis virus encoding a GFP reporter) (**Figure S6C**). Finally, because duplicated haplotypes may lead to increased doses of *PKR* in *Myotis* cells, we tested *PKR* impact on cell viability. We found that at low doses none of the *Myotis* *PKRs* affected cell viability. However, higher doses of *PKRs* led to more cell toxicity, potentially resulting in a tradeoff (**Figure 6G**). Altogether, this may explain why *PKR* is rarely duplicated in mammals, and why both single- and duplicate haplotypes of the loci are segregating across several *Myotis* species. These genomic and functional results highlight the impact of an unfixed gene duplicate which may play a role in adaptation to viral infections.

Discussion

A functionally empowered approach to comparative genomics

Bats are widely known for their long lifespan, cancer resistance, and viral tolerance^{6,10,11,36,70,89,143–145}. As highly complex and pleiotropic processes, the genes and mechanisms underlying these phenotypes can be challenging to identify. Comparative approaches to identify the genetic bases of these traits are constrained by the availability of high-quality genomes, annotations, and functional resources for validation. These challenges are exacerbated in the case of rapidly-evolving phenotypes, such as host-pathogen interactions.

Here we outline an approach that enables functional comparative biology by generating cell lines from wing punches of wild caught bats for genome assembly, comparative genomics, and functional follow up. Cell lines are generated from minimally-invasive biopsies collected in the field thus avoiding disturbing natural populations. Given the high density of bat species concentrated at single locations world-wide^{146,147} it is feasible to collect wing punches from a large number of individuals across a wide phylogenetic range; these wing punches can be used to generate cell lines and sequencing libraries for reference genomes in a matter of weeks. This is an important advance, not only for efforts to expand genetic resources across the tree of life^{148–150}, but for conservation genomics. As our approach can generate genomic resources from minimal material gathered via non-lethal sampling, it is well-suited for the study of rare or endangered species for which acquiring sufficient amounts of material can be challenging.

Evolution of lifespan and cancer risk in a new phylogenetic context

The evolution of body size and lifespan across mammals - and the rapid evolution of lifespan in *Yinpterochiroptera* in particular - has major implications for the co-evolution of cancer risk and resistance. While models of body size evolution are well-studied in mammals^{7,44,72,74} the evolution of lifespan is less well understood. By explicitly modeling the evolution of lifespan separately from body size, we recapitulate the extant relationship between body size and lifespan across mammals in evolutionary time. Contrary to prior work, we show that bats exhibit relaxed allometric scaling of lifespan comparable to other mammals. However, *Myotis* demonstrates an increased rate of change in lifespan given body size compared to other mammals. This altered scaling of longevity in *Myotis* has dramatic consequences for their intrinsic, per-cell cancer risk and for the evolution of tumor-suppressor genes and pathways. While cancer risk scales linearly with body size, it scales over time as a power law of $6^{83,86,87}$. Meanwhile, while mammalian body sizes span a 10^6 range of masses, they only span a 10^2 range of lifespans^{16,151}. Unlike other systems where the evolution of cancer resistance has been driven by rapid changes in body size^{42–44,50,91,94}, the body size of *Myotis* has not significantly changed since their common ancestor. Instead, the rapid and repeated changes in lifespan across an order of magnitude in *Myotis* lead to some of the most significant changes in intrinsic cancer risk seen across mammals.

We found a number of genes under selection across multiple longevity-associated pathways, consistent with the pleiotropic nature of the aging process. These include members of canonical longevity pathways such as mTOR-IGF signaling, DNA damage repair, oxidative stress, and the senescence-

associated secretory phenotype. We additionally identified selection in various pathways that have likely emerged as a result of the unique biology of bats, including genes at the intersection of immunity and senescence, such as Serpin-family genes; genes in metabolic pathways including amino acid metabolism; and pervasive selection observed in the ferroptosis pathway, which sits at the intersection of bats' extreme oxidative challenges, metabolic demands, immune function, and cancer resistance. By quantifying the relative contributions of genes under selection to cancer-related pathways at each node, we found significant enrichment of these processes across the phylogeny, especially at nodes undergoing the greatest changes in lifespan and cancer risk.

While the implications of an increased cancer risk are clear, the implications of decreases in relative cancer risk are less so. As expected by Peto's Paradox, we observe an overrepresentation of cancer-related pathways among genes under selection at nodes experiencing high increases in relative cancer risk, consistent with patterns observed in other vertebrates^{44,46,50,52,73,91–94}. However, we also observed an enrichment in cancer-related pathway representation among genes under selection in nodes with significant decreases in cancer risk (e.g. *M. thysanodes*, *M. velifer*). This combination of low intrinsic cancer risk alongside the persistence of cancer-related adaptations, has been observed previously in sloths and armadillos⁷³. Intriguingly, these species demonstrate some of the lowest known rates of cancer among mammals. While no reports or studies of neoplasia rates have been published in *Myotis*, the use of *in vitro* models of carcinogenesis provides a promising avenue for comparative studies of cancer resistance under controlled conditions. In agreement with our results, *in vitro* and xenograft transplant models have shown that cells of long-lived bats, including *M. lucifugus*, are more resistant to carcinogenesis than shorter-lived bats and other mammals¹⁴⁵. Such studies provide a reliable route for the experimental validation of the evolution of cancer resistance in species where *in vivo* work would otherwise prove ethically or practically intractable.

Viral adaptation and immunity

The nature of viral tolerance and infectious disease adaptation in bats has major implications for understanding their role as zoonotic reservoirs and mechanisms of infectious disease adaptation. Here we focus on Virus Interacting Proteins (VIPs) that influence viral response and contain vital information about the nature of host adaptation to viruses¹⁰⁹. By integrating comparative analyses of VIP adaptation, VIP and immune gene family expansion and contraction, and functional experiments, we show that virus adaptation in bats is mostly driven by DNA viruses, as opposed to RNA viruses; we recapitulate and expand on previous results related to positive selection in immune genes and immune gene family expansion, contraction, and loss; and demonstrate complex patterns of structural variation, including a segregating duplication of protein kinase R (PKR), a major protein involved in the antiviral innate immune system, that has functional relevance in its activity against viruses.

The remarkable dominance of adaptation in response to DNA viruses in bats is in contrast with viral adaptation in humans and other primates, which is driven by RNA viruses^{113,152}; and in other mammals, in which virus adaptation is driven by a combination of DNA and RNA viruses. Most zoonoses, including those hosted by bats, are RNA viruses¹⁰, making this especially important in understanding the dynamics of emerging infectious diseases. This novel finding complements previous observations that bats are more likely than other mammals to asymptotically harbor RNA viruses, while being more susceptible themselves to other pathogens, such as fungi¹¹². This suggests multiple, non-exclusive,

possibilities. First, bats may have some other form of response to RNA viruses that sufficiently reduces the fitness effect of these viruses such that the associated VIPs did not adapt as strongly. Second, our result does not imply that bats have not adapted to RNA viruses, rather that adaptation to RNA viruses does not exceed the genomic baseline adaptation, while adaptation to DNA viruses does. Indeed, bats are known to mount adaptive immune responses to some RNA viruses and the strength of their immune response can have complex interactions with hibernation and reproduction¹⁰. It has been previously suggested that bats may rely more strongly on adaptive immunity in response to RNA viruses than to other pathogens¹¹², though evolutionary functional analyses have also found evidence of innate immune adaptation to RNA viruses, including *RTP4* to flaviviruses¹⁵³ and *OAS1* to SARS-COVs¹⁵⁴. This is consistent with our findings of positive selection and gene family expansion in adaptive immune proteins.

While previous work has shown associations between gene family size and certain phenotypic traits in bats^{36,54,155,156}, confirmation of functional effects of copy number is rare. By resolving individual haplotypes in these nine *Myotis* species, we were able to confirm a single duplication event at the origin of *Myotis* *PKR1* and *PKR2*. We further demonstrate functional implications of copy number variation in Protein kinase R, as previously shown in functional evolutionary studies (eg. Jacquet et al. 2022). These results are especially interesting in the light of other studies that have found trans-species polymorphisms related to immune genes¹⁵⁷. This further illustrates the importance of high-quality genome assemblies and annotations, to distinguish copy number variation between haplotypes, as well as between functional copies and pseudogenes¹⁵⁸.

The role of agonistic pleiotropy in driving adaptations in bats

Multiple hypotheses have been proposed to connect the unique physiology and ecology of bats with the evolution of remarkable adaptations such as viral infection tolerance, stress tolerance, and exceptional longevity¹⁴³. Hypothesized drivers of disease resistance and longevity evolution in bats include the evolution of flight (e.g. “flight as fever” hypothesis¹⁵⁹, though this hypothesis has recently been critiqued¹⁶⁰), the disposable soma hypothesis¹⁶¹; metabolic state¹⁶²; torpor⁶; and other adaptations to specific environments^{9,156,163,164}. Additionally, many studies have highlighted the intersection of one or more of these traits, including a relationship between hibernation and both longevity⁶ and disease resistance¹¹². Our results are consistent with an *agonistic* pleiotropy hypothesis, wherein genetic adaptations for many specific traits (e.g. physiological stress to flight, hibernation, DNA virus innate immunity) may prove beneficial to other seemingly-unrelated traits (e.g. cancer resistance, cellular homeostasis, longevity).

Consistent with this, many of the genes and pathways highlighted in this study have been found to play vital roles across physiological traits in bats and other species. For example, two genes under selection in nearctic *Myotis* - *FTH1* and *IGFN1* - have been implicated in functional studies as key hibernation genes^{165–167}, viral interacting proteins^{168–171}, and as pro-longevity genes^{172–174}. Similarly, many DNA VIPs such as *BRCA1/2* and *POLG* represent core DNA maintenance genes essential for cancer resistance and longevity^{51,175–181}; the existence of active DNA transposable elements such as *Helitron* in *Myotis* may provide another selective pressure on DNA repair genes¹⁸². Beyond individual genes, many of the overarching pathways under selection in *Myotis*, such as those associated with inflammation, senescence, and ferroptosis lie directly at the intersection of aging-related immune processes^{36,54,56,75,167,172,183–188}. While these results suggest the possibility that traits such as cancer risk,

cellular homeostasis, and antiviral response have evolved in tandem due to pleiotropic selection at overlapping points in bats' evolutionary histories, further functional validation will be required to disentangle the functional impacts of these genetic changes and disambiguate the drivers of selection.

Acknowledgements

PacBio HiFi sequencing was done by the DNA Technologies and Expression Analysis Cores at the UC Davis Genome Center, supported by NIH Shared Instrumentation Grant 1S10OD010786-01. This material is based in part upon High Performance Computing (HPC) resources supported by the University of Arizona TRIF, UITS, and Research, Innovation, and Impact (RII) and maintained by the UArizona Research Technologies department. This research used the Savio computational cluster resource provided by the Berkeley Research Computing program at the University of California, Berkeley (supported by the UC Berkeley Chancellor, Vice Chancellor for Research, and Chief Information Officer). Computations were performed, in part, on the Vermont Advanced Computing Core. We thank the members of the LP2L team (CIRI) for helpful discussions.

MEL was funded by NSF PRFB #2010884, and a Dovetail Tree of Life Grant. JMV was funded by NSF PRFB #2109915 and NIH NIA T32AG000266. DE was funded by NIH NIGMS MIRA grant 5R35GM142677. PHS was funded by NIH NIGMS grant R35GM142916 and by the Vallee Scholars Award. LE was funded by a grant from the Agence Nationale de la Recherche (ANR, #ANR-202-CE15-0020-01 to LE) and by the CNRS. LE and DE were further funded by a grant from the Joint Call for Proposals between the CNRS and the University of Arizona (International Research Center, IRC, 2021-2024). LE, DE, PHS and SP were further funded by the International Research Project (IRP) RAPIDvBAT from the CNRS, the University of Arizona and the University of California, Berkeley.

Image attributions for the bat photos featured in Figure 1: J. Scott Altenbach (*M. thysanodes*, *M. velifer*); Rick & Nora Bowers (*M. occultus*); U.S. National Park Service (*M. auriculus*); Frank Carey (*M. californicus*); Michael Durham (*M. evotis*); J. N. Stuart (*M. volans*); and SMBishop & the Wikimedia Foundation (*M. lucifugus*).

Author Contributions:

Conceptualization, J.M.V., M.E.L., D.E., P.H.S., L.E., D.F., M.B.; Methodology, J.M.V., M.E.L., D.E., P.H.S., L.E., G.G.S.; Software, J.M.V., M.E.L., D.E., P.H.S., L.E., T.M.L.; Validation, J.M.V., M.E.L., D.E., P.H.S., L.E., J.P.V.M.; Formal Analysis, J.M.V., M.E.L., D.E., P.H.S., L.E., S.M., A.L.C., D.M., S.V.; Investigation, J.M.V., M.E.L., S.M., M.B., D.F., G.G.S., L.G., Z.R.H., M.H., W.K., T.M.L., A.L.C., C.L. S.M., D.M., S.L., J.L., C.R., S.L.R.C., M.S., K.S., W.T., J.D.T., S.V., R.M., M.B., J.P.V.M., S.P., L.E., D.E., P.H.S.; Resources, J.M.V., M.E.L., D.E., P.H.S., L.E., M.B., D.F., G.G.S., L.G., Z.R.H., M.H., W.K., T.M.L., A.L.C., S.M., D.M., S.L., J.L., C.R., S.L.R.C., M.S., K.S., W.T., J.D.T., S.V., R.M., M.B.; Data Curation, J.M.V., M.E.L., S.M., M.B., D.F., G.G.S., L.G., Z.R.H., M.H., W.K., T.M.L., A.L.C., S.M., D.M., S.L., J.L., C.R., S.L.R.C., M.S., K.S., W.T., J.D.T., S.V., R.M., M.B., J.P.V.M., S.P., L.E., D.E., P.H.S.; Writing - Original Draft, J.M.V., M.E.L., D.E., P.H.S., L.E., S.M., J.L.; Writing - Review & Editing, J.M.V., M.E.L.,

S.M., M.B., D.F., G.G.S., L.G., Z.R.H., M.H., W.K., T.M.L., A.L.C., S.M., D.M., S.L., J.L., C.R., S.L.R.C., M.S., K.S., W.T., J.D.T., S.V., R.M., M.B., J.P.V.M., S.P., L.E., D.E., P.H.S.; Visualization, J.M.V., M.E.L., D.E., P.H.S., L.E., S.M., A.L.C.; Supervision, J.M.V., M.E.L., D.E., P.H.S., L.E.; Project Administration, J.M.V., M.E.L., D.E., P.H.S., L.E.; Funding Acquisition, J.M.V., M.E.L., D.E., P.H.S., L.E.

Declaration of Interests:

The authors declare no competing interests.

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1353 Materials and Methods

1354 Data availability

1355 All sequencing data and genomes generated in this study are available on NCBI under Bioprojects
1356 PRJNA973719 and PRJNA1035541. Annotations generated in this study are available at
1357 <https://github.com/docmanny/myotis-gene-annotations>. All other code is available at
1358 <https://github.com/sudmantlab/MyotisGenomeAssembly>.

1359 Sample collection and cell line derivation

1360 All bats sampled for this study were wild caught under scientific collection permits for California
1361 and Arizona (see Supplemental Table 1). Bats were sampled using standard mist-netting procedures,
1362 including taking standard body measurements, following USGS recommendations for White-Nose
1363 Syndrome and COVID-19 prevention^{189,190}.

For *M. lucifugus*, the donor individual was field-caught in California and transported to the Genetics Laboratory of the California Department of Fish and Wildlife, where they were euthanized via isoflurane. The *M. velifer* individual was caught in Arizona and euthanized in the field via isoflurane. For both *M. lucifugus* and *M. velifer*, tissues were collected and preserved via flash-freezing in liquid nitrogen.

For *M. volans*, *M. occultus*, *M. auriculus*, and *M. californicus*, two 3-mm wing punch biopsies were taken from the left and right plagiopatagium of each donor individual and placed in a live cell collection media consisting of DMEM/F12 (Gibco) supplemented with 15mM HEPES (Gibco), 20% FBS (Gibco), and 0.2% Primocin (Invivogen) [yohe2019; curty2023; capel2023]. Wing punches were then brought back to a cell culture facility in Berkeley, where they were used to generate cell lines as previously described[yohe2019; curty2023; capel2023]. Additional cell lines for *M. lucifugus*, *M. velifer*, *M. yumanensis*, *M. evotis*, and *M. thysanodes* were similarly collected and generated.

Cell lines for the *M. evotis* and *M. thysanodes* genomes were generously provided by Richard Miller. Cell lines for functional work in *Rousettus langosus*, *Pteropus rodrigensis*, and *Eidolon helvum* were provided by the San Diego Frozen Zoo.

Sequencing and assembly

For 6 genomes (*M. evotis*, *M. thysanodes*, *M. volans*, *M. occultus*, *M. auriculus*, and *M. californicus*) DNA was extracted from primary cell lines expanded from 3M cells at Passage 2-4 to approximately 40M cells per line using a Circulomics BigDNA CCB kit following the UHMW protocol for cells. DNA from *M. lucifugus* was extracted from flash-frozen tissue by the Genetics Lab of the California Department of Fish and Wildlife. PacBio HiFi libraries were generated and sequenced on a Sequel II (PacBio) by the Functional Genomics Core at the University of California, Berkeley. For cell-line-derived genomes, Hi-C libraries for these genomes were generated from 1M cells at Passage 3 using the OmniC for Illumina kit (Dovetail genomics); libraries were submitted for quality control and sequencing on the Illumina NovaSeq platform (Novogene). For the *M. velifer* genomes, DNA was extracted from flash-frozen tissues, and all DNA extraction, library prep, and sequencing was completed by Dovetail Genomics following standard protocols. For *M. lucifugus*, a previously published Hi-C dataset from 4 pooled individuals was used for scaffolding^{191,192}.

The PacBio reads were processed using SMRTTools (v6.0.0-1, PacBio) to generate the circular consensus sequences using the settings --minPasses=3 --minRQ=0.99. Hi-C reads were processed using trimmomatic¹⁹³ (v0.35-6) to remove adapter sequences and low-quality bases using the settings ILLUMINACLIP:data/trimmomatic-adapters/TruSeq3-PE-2.fa:2:40:15 SLIDINGWINDOW:5:20. To generate the primary contig assemblies, we used hifiasm^{194,195} (v0.14-hd174df1_0) in Hi-C mode, providing both the CCS reads and the trimmed Hi-C reads as input, and purging duplicates using the -l2 option. For our reference genomes, we proceeded with the primary contig assembly (*.asm.hic.p_ctg.gfa).

All reference genomes were scaffolded with YAHS¹⁹⁶ (v1.1a.1s) and the Hi-C datasets. Dovetail Omni-C data were processed and mapped to the genome following the manufacturer's instructions using bwa^{197,198} (v0.7.17-h5bf99c6_8), pairtools¹⁹⁹ (v0.3.0-py37hb9c2fc3_5), and samtools²⁰⁰ (v1.12-

h9aed4be_1). YAHS was run using both default settings as well as with --no-contig-ec; after comparing the outputs, we proceeded with the --no-contig-ec version for our final assemblies.

To finalize the assemblies, we performed manual curation using PreTextView²⁰¹ and the Rapid Curation toolkit²⁰² (version ff964069). The X chromosomes were identified based on size, synteny across genomes, and half-coverage observed in XY genomes; putative Y chromosomes were similarly identified in XY genomes. Mitochondrial genomes were identified and removed from the final assembly by running mitohifi²⁰³ (v3.0) in contig mode on the assembly and removing all scaffolds identified as mitogenomes. The consensus mitogenome from mitohifi was designated as the representative mitogenome for the assembly after manual curation. Finally, to eliminate spurious duplicates, we used FunAnnotate²⁰⁴ (v1.8.15) and the “clean” function to identify and remove any remaining scaffolds with 90% identical to a larger scaffold.

Identification and annotation of repetitive elements

We used RepeatMasker²⁰⁵ (version 4.0.7-open) to annotate repetitive elements in our genomes. We first ran RepeatMasker using a curated database of transposable elements from 249 mammalian species^{36,206} (David Ray, pers. comm.) and the settings “-engine ncbi -s -noisy -xsmall” followed by a second run using RepeatModeler²⁰⁷ and RepeatMasker to identify *de novo* repeats missing from the curated database. All repeats were then soft-masked in all genomes. To assess the repeat landscape, we calculated the summary of divergence from the repeat alignments and created the repeat landscape using auxiliary RepeatMasker scripts (calcDivergenceFromAlign.pl & createRepeatLandscape.pl).

Structural variation

To understand the distribution of structural variants, including segmental duplication events, we used SyRI (Synteny and Rearrangement Identifier¹²⁵) and BISER (Brisk Inference of Segmental duplication Evolutionary stRucture²⁰⁸). We first masked telomere regions using TIDK (Telomere Identification toolKit²⁰⁹), and mapped the primary 22 scaffolds of the nearctic *Myotis* genomes to each other with minimap2²¹⁰. The scaffold corresponding to the X chromosome was omitted because there is no corresponding scaffold in the *M. yumanensis* assembly. To verify homologous chromosomes and fix strand orientation, we used *fixchr* from the SyRI package and manually renamed scaffolds accordingly, then re-mapped with minimap2. We ran SyRI on the resulting files and plotted the results with plotsr²¹¹. We ran BISER on the primary 22 scaffolds of the nearctic *Myotis* genomes with --keep-contigs and default settings to generate bed files with the inferred segmental duplication regions.

RNA-seq

To assist our annotation efforts, we generated mRNA-seq for 7 of the species sequenced *de novo* in this study. For *M. velifer*, samples of heart, brain, kidneys, lungs, pancreas, and testis collected from the donor individual were provided to Dovetail Genomics (CA, USA) for mRNA-seq library preparation and sequencing. Using the same cell lines used for the genomes of *M. occultus*, *M. thysanodes*, *M.*

evotis, *M. volans*, *M. auriculus*, and *M. californicus*, we generated rRNA-depleted total RNA-seq libraries using the NEBNext rRNA Depletion Kit v2 and Ultra II Directional RNA Library Prep Kits. RNA and libraries were quality controlled on an Agilent Bioanalyzer using the RNA 6000 Nano and DNA High Sensitivity assays, respectively. Samples were sequenced on to 100M 150PE reads per sample using the Novoseq platform (Novogene). For *M. lucifugus*, we used the following published RNA-seq data on NCBI SRA generated using poly-A selection and paired-end sequencing: SRR6793287, SRR6793288, SRR6793289, SRR6793290, SRR6793291, SRR6793292, SRR6793293, SRR6793294, SRR6793295, SRR6793296, SRR6793297, SRR6793298, SRR6793299, SRR6793300, SRR6793301, SRR7064951, SRR10512805, SRR10512806, SRR10512807, SRR10512808, SRR10512809, SRR10512818, SRR10512829, SRR10512840, SRR10512845, SRR10512846, SRR10512847, SRR10512848, SRR10512849, SRR10512850, SRR10512851, SRR10512852, SRR10083333, SRR10083334, SRR10083335, SRR10083336, SRR10083337, SRR10083338, SRR10083339, SRR10083340, SRR10083351, SRR10083352, SRR1916825, SRR1916826, SRR1916827, SRR1916830, SRR1916832, SRR1916834, SRR1916836, SRR1916839, SRR1916841, SRR1916842, SRR18761564, SRR18761566, SRR18761568, SRR18761571, SRR18761573, SRR18761563, SRR18761565, SRR18761567, SRR18761569, SRR18761570, SRR18761572, SRR18761574, SRR1270869, SRR1270914, SRR1270919, SRR1270921, SRR1270922, SRR1270923, SRR4249979, SRR4249988, SRR5676382, SRR5676383, SRR5676395, SRR5676396, SRR5676402, SRR1869462, and SRR1013468.

Gene annotation and alignment

Gene predictions

To create optimal gene annotations, we combined *ab initio* gene predictions; orthology inferences; and transcriptomic evidence for a total-evidence dataset facilitated using FunAnnotate^{204,212} with manual interventions. To generate high-quality orthology-based evidence, we downloaded the UNIPARC database²¹³ of genes present in all Chiropteran genomes and mapped these proteins to our genomes using miniprot⁶² (v 0.6-r194-dirty). We assembled our transcriptome data into transcripts using TRINITY²¹⁴ (v 2.13.2), and mapped these transcripts to our genomes using minimap2²¹⁰ (v 2.24).

Next, we ran BUSCO^{63,64} (version 5.4.3) using the “eutheria_odb10” database and AUGUSTUS⁵⁸ to identify BUSCO orthologs in our genomes. GFFs describing the gene structure of single-copy BUSCO orthologs was then used by FunAnnotate to train SNAP²¹⁵ and GlimmerHMM²¹⁶ (v 3.0.4) prior to gene prediction. GeneMark-ES⁵⁹ (v 4.72) was run using its self-trained model. AUGUSTUS²¹⁷ (v 3.4) was run using a previously-generated model jointly trained on 6 high-quality bat genome assemblies³⁶ and supplemented with protein and transcriptome hints generated by FunAnnotate from the UNIPARC and Trinity datasets.

To leverage high-quality annotations from other genomes, we used TOGA⁶¹ (version 1.0.1) to generate gene annotations for each of our species, using inference from hg38 annotations. TOGA outputs a table of genes (“reg” genes) associated with the projected transcripts from the reference genomes, and a BED file describing the CDS structure of these projected transcripts. To generate a final

GFF file summarizing these data, we converted the original BED file to a GFF file using [program]; removed the erroneous “Gene” level attributes; and added in new “Gene” entries describing the TOGA-designated genes, modifying the “Parent” attributes of the mRNAs to refer to the correct parent gene. Transcript projections that were not associated with a TOGA gene designation were then dropped.

Finally, we used LiftOff⁶⁰ (v1.6.3) to lift over annotations from the *Myotis myotis* genome (mMyoMyo1.0_primary³⁶). Using BUSCO and manual curation, we assessed both the original GenBank (GCF_014108235.1) and NCBI RefSeq (GCA_014108235.1) annotations, and selected the NCBI RefSeq annotation, as it had slightly improved BUSCO scoring and less erroneous intron-exon junctions at select genes. We removed all non-protein-coding genes from the initial GFF file, then ran LiftOff using the settings “-exclude_partial -polish -cds”.

We evaluated each line of evidence by assessing their completeness using BUSCO and comparing the completeness score to the total number of predicted genes. We found that SNAP and GLIMMERHMM performed the poorest for gene annotations, with both the lowest BUSCO scores and the highest number of low-quality predictions. The miniprot-UniParc and TOGA-hg38 datasets generated the highest quality gene prediction datasets, with near-complete BUSCO scores and reduced low-quality protein predictions.

Gene prediction curation

We used EvidenceModeler²¹⁸ (version 2.0) to generate an initial consensus gene set using only the best lines of evidence (AUGUSTUS, weight 2; high quality AUGUSTUS, weight 5; TOGA-hg38, weight 12; miniprot-UniParc, weight 5; and LiftOff-mMyoMyo1, weight 5) with hints from protein orthology (miniprot-UniParc, weight 6) and RNA-seq (TRINITY, weight 5) for alternative splicing. By default, EvidenceModeler does not consider genes that are located within intronic regions of other genes. To restore these genes, we intersected the EvidenceModeler consensus gene GFF with the TOGA-hg38 GFF to identify which genes were present in intronic regions and omitted from EvidenceModeler; these genes were then added back to the EvidenceModeler gene set.

To eliminate remaining spurious predictions, we cross-referenced our gene annotations against the SwissProt²¹⁹ database using DIAMOND²²⁰ (v. 2.1.4) with settings “--ultra-sensitive --outfmt 6 qseqid bitscore sseqid pident length mismatch gapopen qlen qstart qend slen sstart send ppos evalve --max-target-seqs 1 --evalve 1e-10”. We kept all genes that matched a protein on SwissProt with at least 80% identity, matched over 50% of the target sequence, and coded for at least 50 amino acids. Of the remaining genes, we kept them only if they contained both a start and stop codon with no internal stop codons.

Finally, we further curated our annotations by putting the EVM and TOGA gene predictions in competition with each other when they both annotated the same locus, but with different overlapping or neighboring annotations. In such cases, one of the gene annotations is likely closer to the truth. To determine which, we compared EVM and TOGA gene models with their closest human gene BLAST hits. Only proteins with a BLAST match to a human Ensembl v99 annotation with the lowest E-values below 0.001 were considered. These human homologs were used as a reference for curation as they are well-defined and characterized. We observed that occasionally, either the EVM or TOGA model predicted a

transcript much longer than their human closest homolog. Closer inspection revealed that such cases represent artifactual mergers of neighboring genes during the annotation process, clearly visible from the fact that they map to two distinct human homologs in succession. Such cases were resolved by choosing the annotations (between EVM and TOGA) that were not affected by the artificial merger. We further observed a specific class of mergers between neighboring, segmentally duplicated genes, with the resulting annotations representing chimeric mixes of exons from the duplicates. In such cases we selected the annotations that clearly stayed within the boundaries of the separate duplicates, as identified by mapping to the closest human homolog. For the remaining annotations where both TOGA and EVM both mapped to a single human homolog throughout their entire length, we selected the most complete annotation that was closest in length to the human homolog.

Orthologous Gene Alignments

Phylogeny and selection analyses described in this manuscript rely on high-quality alignments of bat orthologous coding sequences. To first find and align orthologous *Myotis* genes to the greatest extent possible, we first complemented the gene annotations described above with likely missing annotations that could still be found through BLAT homology searches. Missing gene annotations are always expected in non-model species genomes and reflect a feature of annotation pipelines in general, not an artifactual issue. For example if the first coding exon of a gene falls into a small local assembly gap, the lack of a start codon may prevent the trigger of a CDS annotation, or may lead to the clearly incomplete CDS being subsequently filtered out. Similarly, erroneous indels representing sequencing errors may interrupt coding reading frames. Genes with missing annotations can still be detected in assemblies through classic BLAST or BLAT homology searches, and then aligned with their annotated orthologs from other species. To align orthologous *Myotis* genes from ten species (those sequenced here plus *Myotis myotis* and *M. yumanensis*), we first decided to use *Myotis velifer* as the *Myotis* species of reference, since the RNA-seq data we used was generated with *M. velifer* tissues.

We first looked for missing homologs of *M. velifer* genes in the other *Myotis* genomes by blatting *M. velifer* CDS to the other *Myotis* assemblies (BLAT command line including non-default options -q=dnax -t=dnax -fine) to find matches outside of already annotated genomic segments. When multiple velifer CDS matched to the same locus with multiple overlapping homologous BLAT matches, we selected the match with the highest number of identical nucleotides. The remaining matching BLAT sequences were further considered if they spanned at least 50% of the velifer CDS, and included 100 codons or more. BLAT matches including stop codons were removed. This process added 1,837 putative CDS to consider for orthologous alignments for *M. auriculus*, 1,785 for *M. californicus*, 1,796 for *M. evotis*, 1,505 for *M. lucifugus*, 3,234 for *M. myotis*, 1,826 for *M. occultus*, 1,822 for *M. thysanodes*, 1,800 for *M. volans* and 1,729 for *M. yumanensis*. The correct reading frames for these putative CDS were then determined by aligning to the velifer CDS that generated the initial match with MACSE v2. MACSE has the crucial advantage over other aligners of being able to repair broken reading frames due to sequencing indel errors or erroneous gene annotations. At this stage, we restricted any further analysis to those velifer CDS with human homologs (BLASTP E-value<0.001 with at least one human canonical protein-coding gene from Ensembl). One-to-one orthologs with the 23,030 remaining velifer CDS in other *Myotis* species were then determined using Orthofinder v2.5.4²²¹. The sequences of each group of ortholog were then aligned with MACSE v2²²² with default settings. The resulting CDS with potentially repaired reading

frames were then checked with PREQUAL²²³ to exclude sequencing errors and erroneous inclusion of non-homologous segments in annotations. The remaining parts of orthologous sequences that passed PREQUAL filtering were then aligned again using MACSE v2 with default settings. The first round of alignment with MACSE ensures that we do not remove portions of CDS that look like they have no homology and would thus be removed by PREQUAL, just because of frameshifts that are easy to repair first with MACSE. The second round of MACSE is to align the remaining codons once PREQUAL has removed erroneous portions of CDS that could have otherwise disturbed the alignment process. We further masked (i.e. replaced with indels) codons near indels with putative alignment errors as described in Bowman et al.²²⁴. Of the 23,030 initial *M. velifer* CDSs, this process resulted in 21,756 alignments with at least one ortholog in another *Myotis* species.

We also aligned pan-Chiroptera orthologs from 47 non-*Myotis* genomes publicly available on NCBI at the time of analysis, to test the generality of our observations to all bats. We used the same strategy described above to complement *Myotis* gene annotations with BLAT matches, but this time blatting velifer CDS on non-*Myotis* assemblies (with -q=dnax -t=dnax -fine again) to find all the potential orthologs in the non-*Myotis* assemblies. We previously found that because BLAT represents a first filter to include only portions of homologous CDS with good local similarity, using BLAT matches results in higher quality alignments of orthologs than using existing gene annotations of disparate qualities that too often include non-homologous portions of introns among other issues^{224,225}. As before with only *Myotis* species, we recovered putative one-to-one orthologs with Orthofinder. This process resulted in the alignment (as previously described with two rounds of MACSE and PREQUAL in the middle) of 19,009 orthologous CDS with at least one non-*Myotis* orthologous CDS.

To test whether the patterns of virus-driven adaptation observed in bats are unique across mammals, we also prepared four more datasets of 70 primate orthologous CDS alignments, 138 euungulate alignments, 127 glires alignments, and 82 carnivora alignments (see supplementary files XY for the species and their respective assemblies used). We used the same pipeline as the one used to align 47 pan-chiroptera species as described above, except that instead of starting from velifer CDS, we started from human Ensembl v109²²⁶ CDS (the longest isoform available in each case) for primates, *Mus musculus* Ensembl v109 longest CDS for glires, *Canis familiaris* Ensembl v109 longest CDS for carnivores, and *Bos taurus* Ensembl v109 longest CDS for euungulates. These species were chosen for the very high quality of their gene annotations.

Gene Trees & Phylogeny

A phylogeny of all 536 mammals in our alignments was generated using IQTREE²²⁷ (version 2.3.1) using all gene alignments with the settings “-B 1000 -m GTR+F3x4+R6.” To generate gene trees, we first filtered our gene alignments to exclude alignments with over 50% gaps in the sequence and less than 4 species. With the remaining alignments, we used IQTREE to find the best-fitting substitution model and tree using settings “--wbtl --bnni --alrt 1000 -B 1000 --safe”. The best substitution models for each gene were saved as a NEXUS file. To generate the phylogeny of bats, we first concatenated all gene alignments using *catfasta2phym* (<https://github.com/nylander/catfasta2phym>) to concatenate our individual gene alignments into species-level alignments, filling in missing species in each sub-alignment

with gap symbols to preserve the alignment structure. Furthermore, we generated a partition file describing the region of each gene sub-alignment within the concatenated alignment.

Time-calibration of 59 bat genomes

Using our codon alignments of 59 bat genomes, we generated a time-calibrated phylogeny using *mcmctree*²²⁸ and PAML²²⁹ (v. 4.10.0) using an approximate likelihood method. Using the pan-bat codon alignments and our phylogeny as input, with fossil calibrations based on previously published work^{4,36,230–237}, we ran *mcmctree* twice to generate the Hessian matrix and confirm convergence. This was followed by 10 independent chains using the “out.BV” file from the first run. Finally, the output files of all 10 chains were combined to compute final divergence time estimates (see Table S2).

Ancestral Body Size, Lifespan, and Cancer Risk reconstruction

To explore how body size and lifespan have evolved over time in mammals, we used a super-phylogeny of mammal species⁶⁷ subsampled to only contain species with extant body size and lifespan data collected from AnAge¹⁵ and PanTHERIA¹⁶. Ancestral body sizes and lifespans were simulated separately using StableTraits²³⁸.

To estimate ancestral longevity quotients (AncLQs), we followed the method of Austad and Fisher¹⁸ and used a linear model of lifespan given body size trained on non-flying mammals to predict the lifespans at each ancestral node given median estimates of body size. AncLQs were then estimated from the ratio of observed lifespan versus predicted lifespan for each node.

Relative Incidence of Cancer Risk (RICR) was calculated across our mammalian phylogeny following the method of Vazquez and Lynch (2021)⁴⁴. The cancer risk K at a given node was calculated using the log of the median predicted body size and lifespan. An organism's lifetime risk of cancer K is proportional to Dt^6 , where D is the body size and T is the maximum lifespan. RICR is then calculated as the $\log_2$ ratio of the cancer risk between a node and its direct ancestor.

Selection Scans & Evolutionary Rates

aBSREL

To conservatively test for branch-specific selection, we used aBSREL^{95,239} (version 2.5.48) to test for selection at each branch within the Nearctic *Myotis* clade for 15,734 gene alignments spanning 536 mammals. These genes were identified as 1:1 orthologs across the full alignment, with no more than 50% sequence gaps and at least 4 species present in the alignment. We defined genes under selection as those with an FDR-corrected p-value of less than 0.05; genes were specifically identified as under positive selection if $\omega > 1$.

BUSTED

To quantify the total amount of positive selection across the *Myotis* tree or the different species trees used in this manuscript, we used an improved version of the BUSTED^{110,239} test called BUSTED-MH. The original BUSTED test estimates for a given gene the proportion of codons that have evolved under positive selection, with $dN/dS > 1$, summed over all the branches of a given tree, regardless of the branch and regardless of the codons in a multi-species alignment. The version of BUSTED we used, BUSTED-MH, includes two crucial improvements over the original BUSTED that make it much less likely to generate false positive inferences of positive selection, albeit at the cost of becoming a very conservative test. First, BUSTED-MH takes synonymous substitution rate variation into account, which prevents mistaking cases where dN/dS is greater than one just because dS is low, with cases where dN/dS is greater than one because positive selection actually increased dN . Second, BUSTED-MH takes complex substitutions that simultaneously involve more than one nucleotide into account in its likelihood models. This prevents attributing positive selection to cases where dN/dS is greater than one where instead a complex substitution changed multiple amino acids in a single event. BUSTED-MH has been shown to strongly reduce the rate of false positives that typically plague dN/dS -based tests of positive selection²⁴⁰.

We applied BUSTED-MH to 19,646 *Myotis* orthologous CDS alignments with at least five orthologs. These orthologs are cases where the Orthofinder gene trees coincide with the species tree. This effectively removes issues regarding whether we should use the gene or the species tree, at the cost of removing 2,110 genes from the *Myotis* selection analysis. Similarly, we applied BUSTED-MH to 17,469 non-*Myotis* bat alignments with at least five orthologs. This includes a subset of 14,091 alignments with orthologs in two thirds of the non-*Myotis* bat species that we specifically used to show that patterns of virus-driven adaptation are representative of all, and not just a limited subset of bats. We also tested 17,890 primate alignments with at least five orthologs with BUSTED-MH, as well as 19,311 glire, 18,000 carnivora and 18,504 ungulate alignments.

RERConverge

Between-species life history diversity may be undergirded by significant evolutionary rate shifts in important genes, where evolutionary change across the gene tree correlates either positively or negatively with changes in a particular life history trait across the trait tree. In *Myotis*, we were interested specifically in testing whether or not longevity-related metrics could be correlated with evolutionary rate shifts for particular genes, and if, among those, we could identify types of genes (gene ontologies) that were enriched.

To answer this question, we used RERconverge¹⁰⁶, an R package which uses gene trees to compute relative evolutionary rates (RERs), then tests for correlations between RERs and trait changes between species. 40 bat genomes were aligned to produce MSAs, which were then split into three groups to be tested independently: all bats ($n=59$), non-*Myotis* bats ($n=29$), and *Myotis* ($n=11$). Gene trees were constructed under the GTR+G model with the same topology as determined in our phylogenetic analysis, across all 39 available bat species. After concatenating the gene trees, RERs were calculated in RERconverge. Trait correlation analysis was performed by regressing these RERs against 4 distinct trait axes. Two of the axes were maximum longevity and size, which were obtained from AnAge¹⁵¹ and PanTHERIA¹⁶; an additional two axes were obtained by plotting species along the first 2 principal

components of size and maximum longevity. Since size generally correlates with longevity, even within *Myotis*, PCA allows us to describe species using orthogonal trait axes that roughly correspond to size-independent longevity and longevity-independent size. Using a Wilcoxon rank-sum test, we then tested for enrichment in correlation significance amongst different gene sets.

RELAX

The evolution of life history diversity across a clade may also manifest in differential selection regimes across relevant genes or types of genes. Specifically, the evolution of a particular life history may be driven by either relaxation or intensification of selection in different genes. In *Myotis*, we were again interested in whether we could identify genes and gene sets related to increased longevity within the clade.

RELAX⁹⁹ is used to identify genes under either relaxation or intensification of selection across groups of species within a clade using MSAs and a labeled species tree. MSAs for 11 available *Myotis* species across ~19,000 shared genes were fit using the BS-REL framework to a branch-site model, using the species tree determined from our phylogenetic analysis. 4 longer-lived species, *Myotis lucifugus*, *M. occultus*, *M. evotis*, and *M. myotis* were set as the foreground class with the remaining species set as the background class. RELAX then used these branch classes to estimate a distribution of ω (dN/dS) for each branch class, constrained by the relaxation factor k . An LRT is performed for $k \neq 1$ against $k = 1$, with $k > 1$ implying relaxation of selection and $k < 1$ implying intensification of selection. The results from this test were then used to perform a Wilcoxon rank-sum test to identify enrichment in the significance of the k -parameter amongst different gene sets.

VIPs

To determine if *Myotis* and other bats are enriched for adaptation at Virus Interacting Proteins (VIPs), we conducted a test comparing levels of adaptation, inferred by BUSTED, in sets of VIP genes compared to matched control genes. Sets of control genes were resampled in a bootstrap procedure (https://github.com/DavidPierreEnard/Gene_Set_Enrichment_Pipeline) to generate 95% confidence intervals for sets of genes at progressively smaller BUSTED p-value thresholds^{109,111,113,152}. When VIPs are subject to greater levels of positive selection than expected relative to the sets of matched control genes, we expect a pattern in which the high p-value thresholds show weaker enrichment but smaller confidence intervals, because more genes are used in these calculations. As the p-value threshold gets smaller, the signal of enrichment is expected to get stronger but at the expense of larger confidence intervals.

We generated five sets of VIP genes: A set of all VIP genes with aligned orthologs from at least five species in the tested clade (Nearctic *Myotis* or pan-Chiroptera without *Myotis*); a set of VIP genes with known pro- and/or anti-viral activity; a set of VIP genes with no known pro- and/or anti-viral activity; a set of VIP genes that interact only with DNA viruses; and a set of VIP genes that interact only with RNA viruses. Because both the number of species and genes included, as well as their level of homology, influences the power of these tests we also tested the influence of the stringency of gene choice by generating a separate set of genes for the pan-Chiroptera analyses that included only genes with aligned orthologs in at least two thirds of the non-*Myotis* species. Analyses using this more limited set of genes

show the same result in terms of enrichment of adaptation in VIP genes and comparing DNA VIPs and RNA VIPs, showing that the observed patterns are valid across bats regardless of the stringency of homology.

The bootstrap procedure matches a tested gene set of interest such as VIPs with sets of control genes (non-VIPs when testing VIPs) that have the same average values as the set of interest for multiple potential confounding factors that could explain differences in adaptation instead of interactions with viruses. For example, if the level of gene mRNA expression has an influence on the rate of adaptation, we then need to match VIPs with control sets of non-VIPs that collectively have the same average expression as VIPs. For each group of tested VIPs we build 1,000 control sets with randomly sampled non-VIPs according to a matching procedure described in Enard & Petrov 2020^{152,241}. We match the 27 following factors between VIPs and non-VIPs, for all tested groups of species:

- the length of the aligned CDS.
- the overall CDS GC content in each orthologous alignment.
- the GC content at aligned codons' position 1, 2 and 3 separately.
- the number of species with a one-to-one ortholog out of all the species included in an alignment, where species with no ortholog are represented by gaps the whole length of the alignment.
- the number of species with an ortholog at least 90% of the length of the species of reference (*Myotis velifer* in bats, human in primates, etc; see above).
- the overall proportion of each orthologous alignment made of indels.
- the three synonymous rates of evolution estimated by the likelihood model of HYPHY Busted.
- the proportions of codons that fall in the three latter synonymous rates.
- average human mRNA expression in 53 GTEx v7 tissues²⁴², in log₂ of Transcripts Per Million (TPM).
- lymphocyte human mRNA expression from GTEx v7, in log₂ of TPM.
- testis human mRNA expression from GTEx v7, in log₂ of TPM.
- mRNA expression in log₂ of TPM for six separate *Myotis velifer* tissues: heart, brain, kidneys, lungs, pancreas, and testis.
- the number in log₂ of protein-protein interactions (PPIs) in the human protein interaction network²⁴³.
- the proportion of genes that are immune genes according to Gene Ontology annotations of the closest human homolog including Gene Ontology terms GO:0002376 (immune system process), GO:0006952 (defense response), and/or GO:0006955 (immune response) as of summer 2021²⁴⁴.
- the proportion of housekeeping genes defined as genes with stable expression across many human tissues, listed in Eisenberg & Levanon²⁴⁵.
- the overall dN/dS ratio estimated by Busted for the orthologous CDS alignments.

We match the overall dN/dS between VIPs and control non-VIPs to account for an important issue of dN/dS tests: dN/dS-based tests tend to lose statistical power to detect positive selection in CDS alignments with higher selective constraint²⁴⁶. The amount of positively selected sites being equal, positive selection tests based on dN/dS tend to have lower statistical power and tend to generate more false negative results when the rest of the coding sequence is more highly constrained. VIPs tend to be much more strongly constrained than non-VIPs¹⁰⁹, which gives a

strong, unfair statistical disadvantage to VIPs when testing positive selection with BUSTED or other HYPHY tests. We limit this issue by matching VIPs and control non-VIPs for dN/dS. Thus, VIPs have an excess of adaptation compared to non-VIPs when they have a balance of the same total amount of non-synonymous changes more tilted towards advantageous rather than neutral amino acid changes. In this case non-VIPs still have less constraint (more neutral changes) than VIPs, and thus still more power to detect positive selection, but not to an extent as severe and unfair as if we did not match the overall dN/dS¹⁰⁹. In the case where VIPs do not have an excess of adaptation, then they have the same balance of advantageous and neutral amino acid changes resulting in the same overall dN/dS. This is the case of RNA VIPs in bats in this study; this internal negative control shows that the matching of dN/dS works as intended.

Gene Duplications

To quantify patterns of gene duplication and loss, we quantified the copy number of genes with human orthologs from our gene annotations for each nearctic *Myotis* genome. To calculate per-gene expansion and loss rates and their statistical significance, we ran CAFE¹³⁷ v5 on the previously described set of copy number counts using our time-calibrated species tree pruned to include only the nine nearctic *Myotis* species. *M. myotis* was excluded because of its lower quality assembly. We ran CAFE on the subset of genes with two or more copies in at least one species using a Poisson distribution for the root frequency ($-p$), first generating an error model to correct for genome assembly and annotation error ($-e$). We compared the base model (each gene family belongs to the same evolutionary rate category) to gamma models (each gene family can belong to one of k evolutionary rate categories) with different values of k . A final gamma model with $k=9$ was chosen to balance model log likelihood with the number of gene families for which the optimizer failed. The model was run three separate times to ensure convergence.

To understand if genes in these pathways have higher birth rates or are more likely to have significant changes in gene copy number than expected relative to other genes, we compared the gene copy birth rate λ and number of genes that have significantly expanded or contracted in copy number on at least one branch within our nearctic *Myotis* phylogeny. Following Huang et al.⁴⁸, we tested if VIP genes in particular underwent significant copy number changes or had significantly different birth/death rates than non-VIP genes. For each category of VIP genes (all VIPs, DNA VIPs, DNA only VIPs, RNA VIPs, and RNA only VIPs), we generated 100 bootstrap sets of control non-VIP genes with the same number of genes as the corresponding VIP gene set. We ran CAFE on each set of VIP genes and the corresponding control non-VIP genes to infer per-gene birth-death rates and per-gene, per-branch expansion/loss events.

Assessment of DNA Double-Strand Break Tolerance

We assessed each species' tolerance to DNA double strand breaks using a by measuring viability, cytotoxicity, and apoptosis across a range of doses of Neocarzinostatin, a radiomimetic drug. We measured dose response curves in wing-derived primary dermal fibroblasts across 5 bat species (*Myotis lucifugus*, $n=8$; *Myotis evotis*, $n=3$; *Rousettus langosus*, $n=2$; *Eidolon helvum*, $n=2$; *Pteropus rodrigensis*, $n=2$) using the multiplexed ApoTox-Glo assay (Promega). Using 96-well plates, two individuals and 11

doses were assessed simultaneously with four technical replicates. Results were normalized to treatment controls for each individual in R as previously described^{42,43,46,50}.

Mapping PKR exons

We further validated the annotations for the PKR locus by re-aligning the primary *M. velifer* coding sequence back to the nine nearctic *Myotis* reference genomes, as well as a non-*Myotis* outgroup, *Pipistrellus pygmaeus*, and the genome haplotypes for each of these species. Because the presence of two copies makes this task challenging for most aligners, we independently aligned the *M. velifer* reference PKR sequence to sequential sections of each genome in 50kb search regions surrounding the known loci in each genome. This alignment search was conducted for 5 regions upstream (250 kb) and 5 regions downstream (250 kb) of the known loci. In species with two known copies, the location of each copy was included in a separate search region. This was to prevent erroneous merging or loss of exons. These regions were retrieved using bedtools getfasta²⁴⁷ and alignment was performed using miniprot⁶². Miniprot settings were optimized to retain secondary alignments (-p 0 -n 1 -outsc=0.0 -outc=0.0) and find known exons with accurate boundaries (-J 18 -F 21 -O 15 -L 10). The resulting gff file was converted to bed format using AGAT²⁴⁸, sequences retrieved with bedtools getfasta, and a custom script used to remove identical duplicates. Finally, all sequences were aligned with MACSE v2.07²²². We used BISER^{125,208} to confirm the presence of segmental duplications in these regions.

PKR cell lines and vectors

All PKR experiments were performed using HeLa PKR-KO cells (kindly provided by A. Geballe, Fred Hutchinson Cancer Center, Seattle WA) that were plated either at densities of 5x10⁴ cells/mL in 24-well plates or at 1x10⁵ cells/mL in 12-well plates. The cells were maintained at 37°C with 5% CO₂ and cultured in DMEM supplemented with 5% fetal bovine serum (FBS), 1% penicillin/ streptomycin mix and 1 µg/mL puromycin (Sigma-Aldrich). All transfections were performed 24 hours after seeding, using 3 µL of TransIT-LT1 Transfection Reagent (Mirus Bio) per 1 µg of DNA and Opti-MEM media. We used previously-generated pSG5-FLAGx2 vectors encoding either *M. myotis* PKR-1 (GenBank OP006550), *M. myotis* PKR-2 (GenBank OP006559), *M. velifer* PKR-1 (GenBank OP006558), or *M. velifer* PKR-2 (GenBank OP006557)²⁸. Plasmids encoding the interferon-stimulated gene ISG20²⁴¹ and a constitutively active variant of the sterile alpha motif domain-containing protein 9-like SAMD9L-F886Lfs*11 (here, SAMD9L²⁴⁹) were used as controls in viral infections and cell translation experiments, respectively.

Western blot

We assessed for the steady state protein expression of *M. myotis* Flag-PKR after transfection of 350 ng and 700 ng of DNA plasmids encoding either PKR1 alone, PKR2 alone, or both PKR1 and PKR2 (175 ng of each and 350 ng of each, respectively). Briefly, cells were re-suspended and lysed in ice-cold RIPA buffer (50 mM Tris pH8, 150 mM NaCl, 2 mM EDTA, 0.5% NP40) with protease inhibitor cocktail (Roche) and sonicated. 20 µL of the clarified fraction was denatured with 5 µL of 6x Laemmli buffer at 95°C for 5 min and loaded into 4-20% BioRad Criterion TGX Stain-Free precast gel. The wet transfer into a PVDF membrane was executed overnight at 4°C. The membranes were blocked in a 1xTBS-T buffer (Tris HCl 50 mM pH8, NaCl 30 mM, 0.05% of Tween 20) containing 10 % powder milk,

and were incubated for 1h. The membranes were incubated with primary mouse anti-Flag (Sigma F3165) and anti-Tubulin (Sigma T5168) antibodies and secondary anti-Mouse IgG-Peroxidase conjugated (Sigma A9044). Detection was made using the Chemidoc Imagina System (BioRad) with SuperSignal West Pico Chemiluminescent Substrate (ThermoFisher Scientific).

PKR co-immunoprecipitation

Hela Δ PKR cells were transfected with 1.25 μ g plenti6 HA-tagged *M. myotis* PKR1 plasmid per million cells and 1.25 μ g of either plenti6 myc empty vector, myc-tagged *M. myotis* PKR1 or myc-tagged *M. myotis* PKR2 plasmid using TransIT-LTI transfection reagent (Mirus Bio). The next day, some wells were infected with Sindbis virus expressing GFP (SINV-GFP) at MOI 2 for 24 hours. Cells were then scraped with cold PBS and pelleted. For the IP, cells were lysed in 500 μ l IP buffer (50mM Tris HCl pH7.5, 140mM NaCl, 6 mM MgCl₂, 0.1% NP40) supplemented with RNase (RiboLock, Fisher Scientific) and protease (cOmplete EDTA-free protease inhibitor cocktail, Sigma) inhibitors for 10 minutes on ice, then centrifuged at 12,000 $\times$ g for 10 min at 4°C. 5% of the volume was kept for input, while the rest was incubated with 40 μ l μ MACS anti-c-myc MicroBeads (Miltenyi Biotec) for 1h at 4°C with constant rotation. Samples were then loaded onto μ MACS columns placed in the magnetic field of a μ MACS Separator (Miltenyi Biotec), washed 4 times with cold IP buffer, and eluted with the μ MACS denaturing elution buffer. Proteins were denatured in elution buffer for 5 min at 95°C, then loaded onto a 4-20% BioRad Criterion TGX Stain-Free precast gel and transferred onto an Amersham Protran nitrocellulose membrane (Sigma) for 1h. Membranes were blocked for 1h in 5% milk in PBS (Euromedex) supplemented with 0.2% tween (Fisher) and incubated with mouse anti-myc monoclonal antibodies (Abcam 9E10, cat# ab32) then secondary anti-mouse IgG antibodies conjugated with HRP (Sigma, cat# A4416), or with rat anti-HA antibodies conjugated with HRP (Roche, Sigma, cat# 12013819001). Images were taken on a Fusion FX imager (Vilber) with SuperSignal West Femto Chemiluminescent Substrate (ThermoFisher Scientific).

Cell viability assay

Hela PKR-KO cells were transfected 24h after plating in 96 well plates, with 100 or 200 ng of pSG5 plasmid: empty or coding for *M. myotis* or *M. velifer* PKR1, PKR2 or PKR1+2 equal mix (50%-50%). 24 hours post-transfection, positive control cells were treated with an apoptosis-inducing drug, Etoposide, at different doses (250, 200 or 100 μ M). 48 hours post transfection, cells were harvested and lysed to quantify luminescent signal according to CellTiter-Glo® Luminescent Cell Viability Assay (Promega) kit protocol.

VSV and SINV infections

VSV infections. Cells were transfected 24 h after plating with 350 ng of pSG5 plasmid: empty, or encoding *M. myotis* or *M. velifer* PKR1, PKR2, or equal input of PKR1 and PKR2 (175 ng per plasmid), or a plasmid encoding interferon-stimulated exonuclease gene 20 (ISG20), due to its known antiviral activity against VSV as positive control²⁴¹. Cells were infected 24 h post transfection with replicative VSV-GFP virus¹⁴² at a MOI of 3. Cells were fixed with 4% paraformaldehyde 16-18 hours post infection. VSV infection was quantified by measuring the percentage of GFP positive cell populations with BD

FACSCanto II Flow Cytometer (SFR BioSciences). Fold change results were normalized to the empty pSG5 condition across at least three independent experiments.

SINV infections. Hela Δ PKR cells were transfected with 5 μ g pSG5 empty vector, *M. myotis* PKR1, *M. myotis* PKR2 or 2.5 μ g *M. myotis* PKR1 + 2.5 μ g *M. myotis* PKR2 per million cells using TransIT-LTI transfection reagent (Mirus Bio). The next day, some wells were infected with SINV-GFP at MOI 0.2. Cells were then placed into a CellCyte X live cell imaging system (Cytex) and pictures of every well were taken every 2h for 48h. The fraction of GFP+ cells over the total cell area was measured and averaged from six photos of 2 individual wells per condition, and repeated for a total of three independent experiments.

Luciferase reporter assays

Luciferase reporter assays were carried out to investigate whether the two PKR paralogs have synergistic, additive or dominant negative effect in translation shutdown. Transfection was performed as previously described with additional 50 ng of FFLuc firefly luciferase reporter plasmid per well. Sterile alpha motif domain-containing proteins 9L (SAMMD9L gain-of-function mutant) was used as a positive control of translational repression²⁴⁹. 24 h post transfection, cells were briefly washed with PBS, lysed by a 5 $\times$ reporter lysis buffer (Promega) and incubated overnight at -20°C. Cells were then collected and 100 μ l of the luciferase substrate (Promega) was added to 20 μ l of the lysis supernatant. Alternatively, cells were lysed using BrightGlow Lysis Reagent (Promega E2620). The relative luminescence units (RLUs) were immediately quantified with LUMIstar Omega microplate reader optima (BMG Labtech). All luciferase assays were conducted in technical duplicates in at least five independent experiments. Fold change results were normalized to the empty pSG5 condition within each independent experiment.

1882 Figures

- 1883 **Figure 1: 8 near-complete reference assemblies for North American (Nearctic) *Myotis*.**
- 1884 **Figure 2: Evolution of body size, lifespan, and cancer risk in bats and mammals.**
- 1885 **Figure 3: Selection in Nearctic *Myotis* is enriched for pleiotropic cancer resistance pathways.**
- 1886 **Figure 4: Adaptation to DNA viruses, but not RNA viruses, is enriched in *Myotis* and other bats.**
- 1887 **Figure 5: A varied structural variation landscape across 9 nearctic *Myotis* species.**
- 1888 **Figure 6: Evolutionary history and function of an actively segregating copy number polymorphism**
- 1889 **of *PKR* in *Myotis*.**

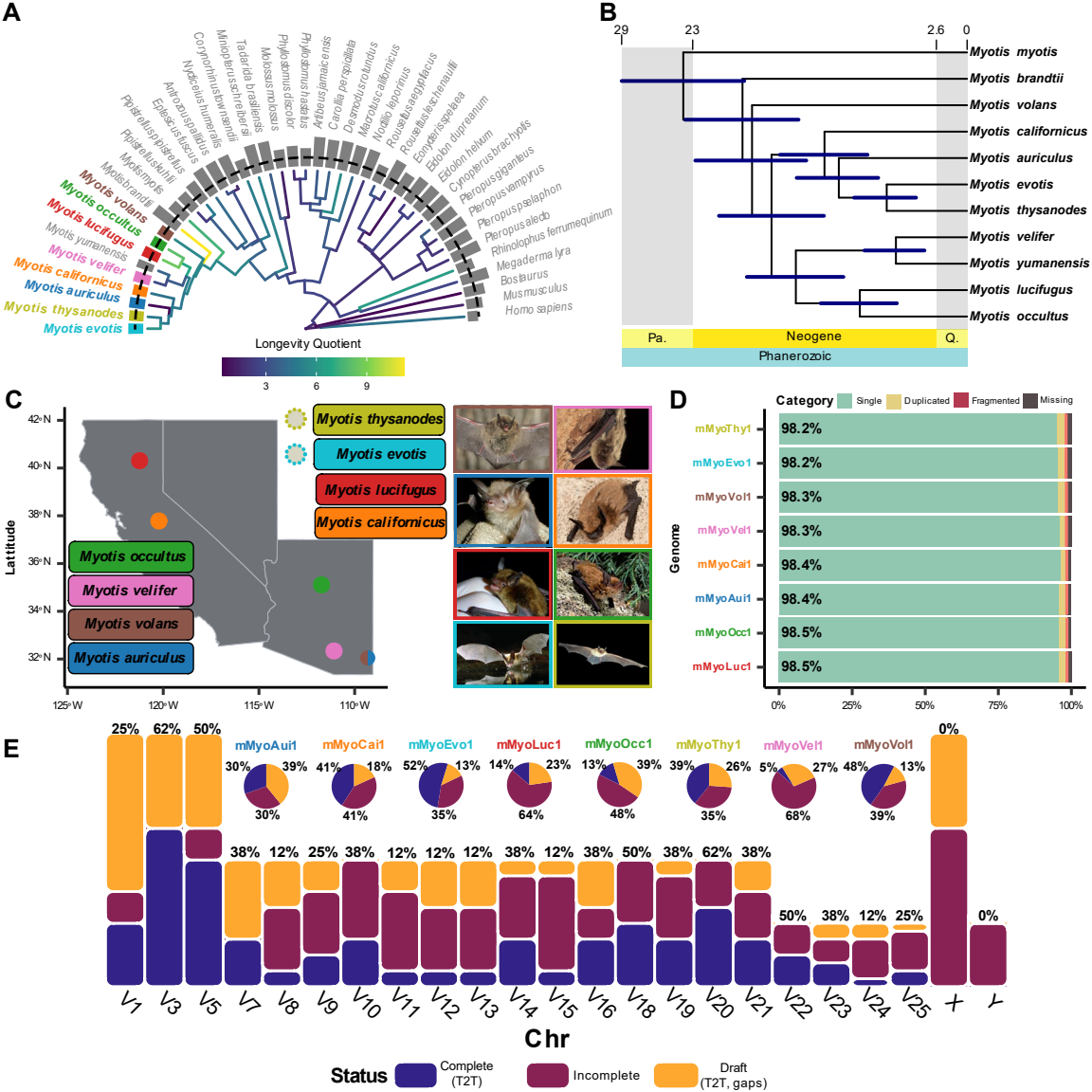
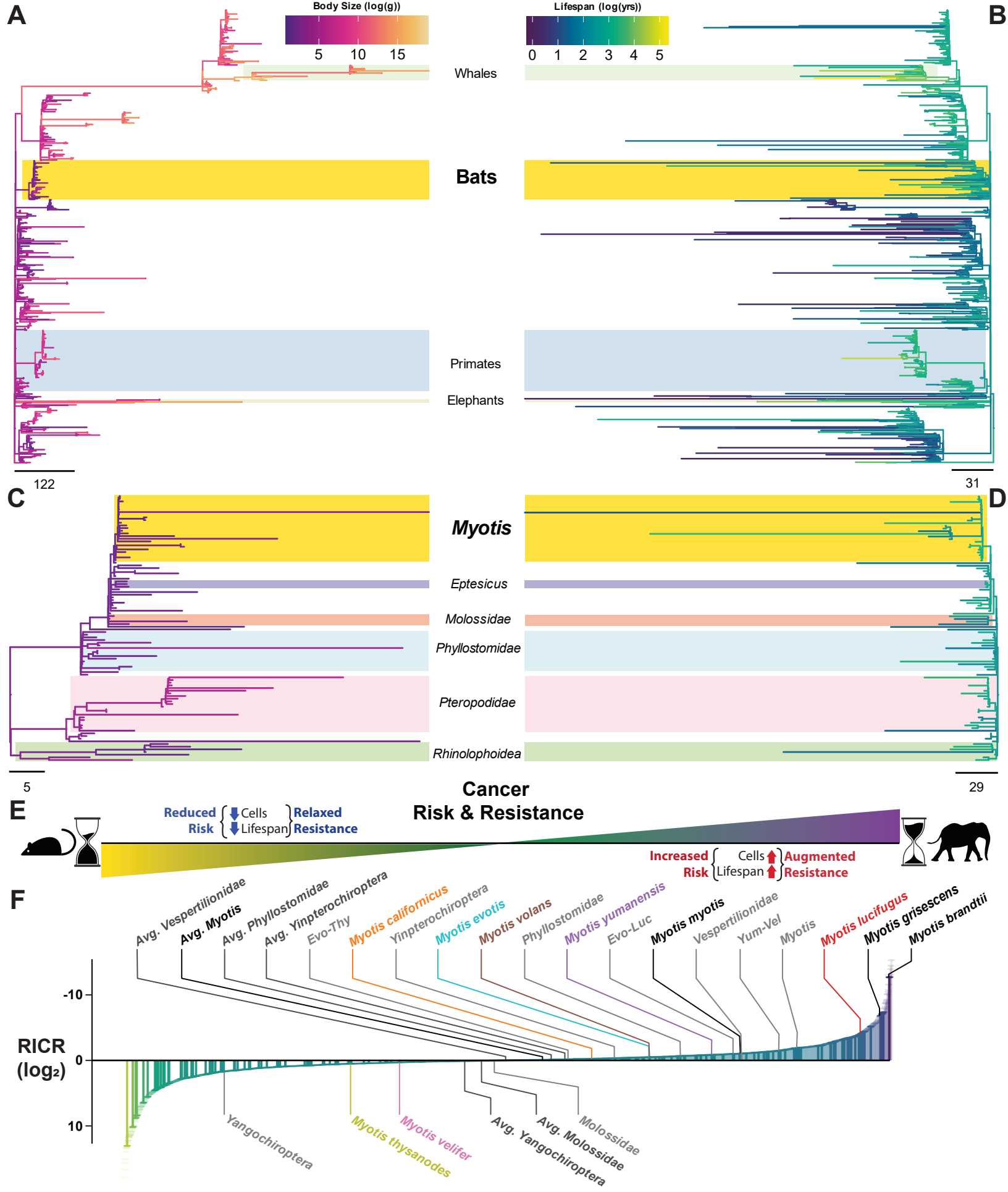
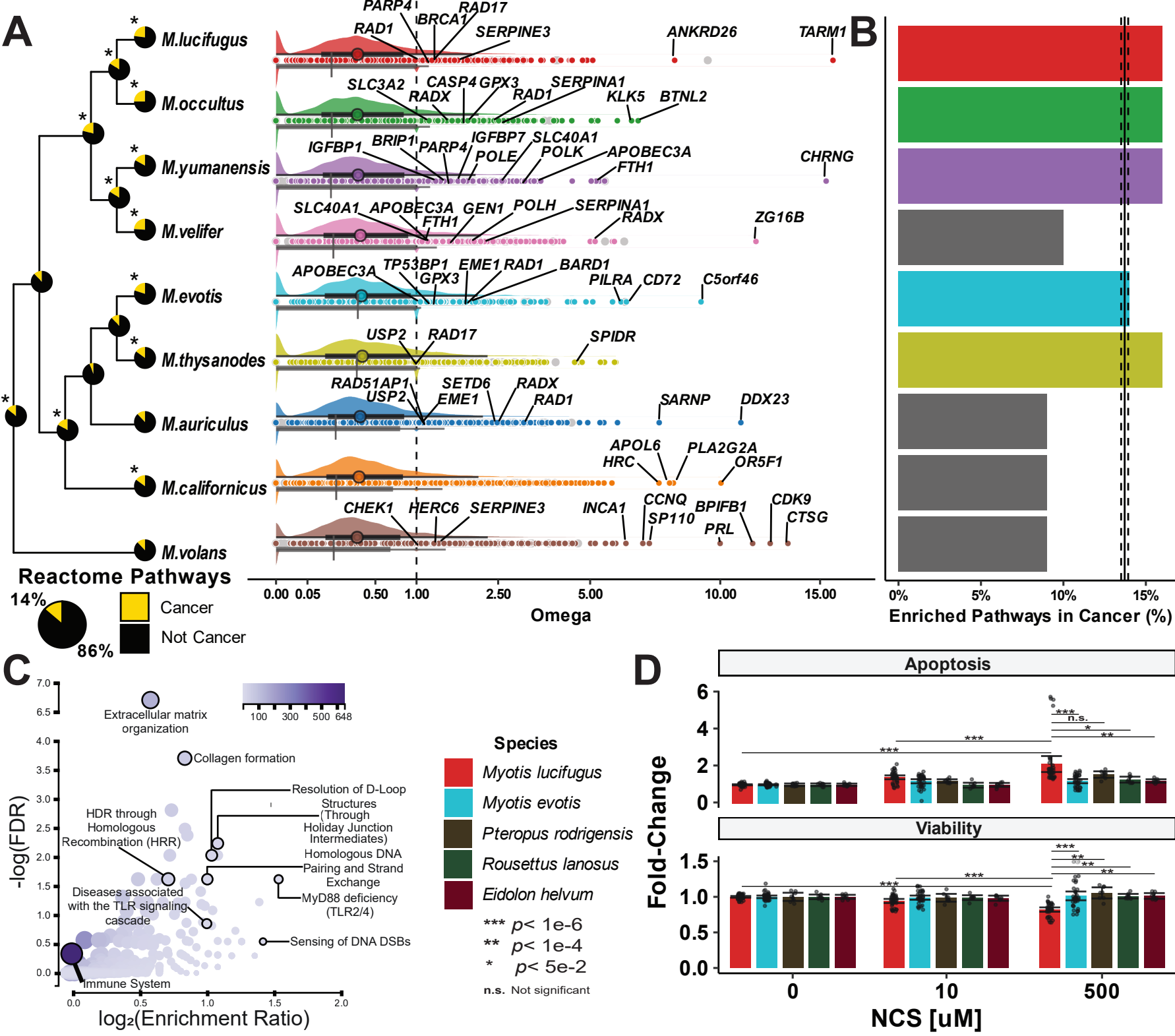


Figure 1: 8 near-complete reference assemblies for North American (Nearctic) *Myotis*. **A)** Phylogeny of 38 bat genomes with 3 outgroup species: cow (*bosTau9*), mouse (*mm39*); and human (*T2T-CHM13v2.0*). Bars at the tips of the phylogeny indicate the AuNG score of each genome (lower values equal more contiguous genomes); the dotted line represents the AuNG score for complete (T2T) genome assemblies as represented by T2T-CHM13v2.0. **B)** The time-calibrated phylogeny of 9 Nearctic and two representative Palaearctic *Myotis* species based on orthologous codon alignments. Blue bars represent age uncertainties. **C)** Map of capture sites with representative images (see “Acknowledgements” for attributions) for the individuals and species sequenced in this study; cell lines for *M. evotis* and *M. thysanodes* were provided by Richard Miller and were not collected for this study. **D)** Mammalian BUSCO scores for annotations generated for the 8 new *Myotis* genomes. **E)** Ideogram bar plot indicating completion status of each chromosome in assembly. Pie graphs indicate completion status of all chromosomes in assembly. All chromosomes were positively identified based on size, synteny, and homology to human chromosomes⁵⁷. “Complete (T2T)” status indicates that a chromosome is fully assembled telomere-to-telomere without gaps; “Draft (T2T, gaps)” status indicates that a chromosome is fully scaffolded with both telomeres, but has one or more gaps in the assembly; “Incomplete” status indicates that a chromosome was positively identified, but was not scaffolded from telomere to telomere (only contains one telomere).



1975 Supplemental Information

- 1976 **Document S1.** Figures S1-S6
- 1977 **Table S1.** Genome Statistics
- 1978 **Table S2.** Phylogeny time calibration and evolutionary modeling data
- 1979 **Table S3.** aBSREL significant gene lists and Reactome enrichments
- 1980 **Table S4.** RERConverge and RELAX results and enrichments
- 1981 **Table S5.** List of VIPs and VIP subclasses
- 1982 **Table S6.** SyRI-identified structural variants (SVs)
- 1983 **Table S7.** Experimental data for Neocarzinostatin and PKR experiments
- 1984



1906 **Figure 2: Evolution of body size, lifespan, and cancer risk in bats and mammals. A, B)** Cophylo plot
 1907 of the evolution of body size (**A**) and lifespan (**B**) across *Eutheria*. **C, D)** Cophylo plot of the evolution of
 1908 body size (**C**) and lifespan (**D**) in bats. Branch lengths in **A-D** are scaled proportional to the rate of change
 1909 of the trait over time, and tree scales are shown below their respective phylogenies. **E)** Diagram illustrating
 1910 the relationship between changes in body size and lifespan with changes in cancer risk and resistance. **F)**
 1911 Reduced Intrinsic Cancer Risk (RICR) for every node in *Eutheria*, ranked from greatest reduction in cancer
 1912 risk to greatest increase in cancer risk. RICR relative to the most recent ancestor of select nodes are
 1913 highlighted, as well as the average RICR across for all nodes within select clades.

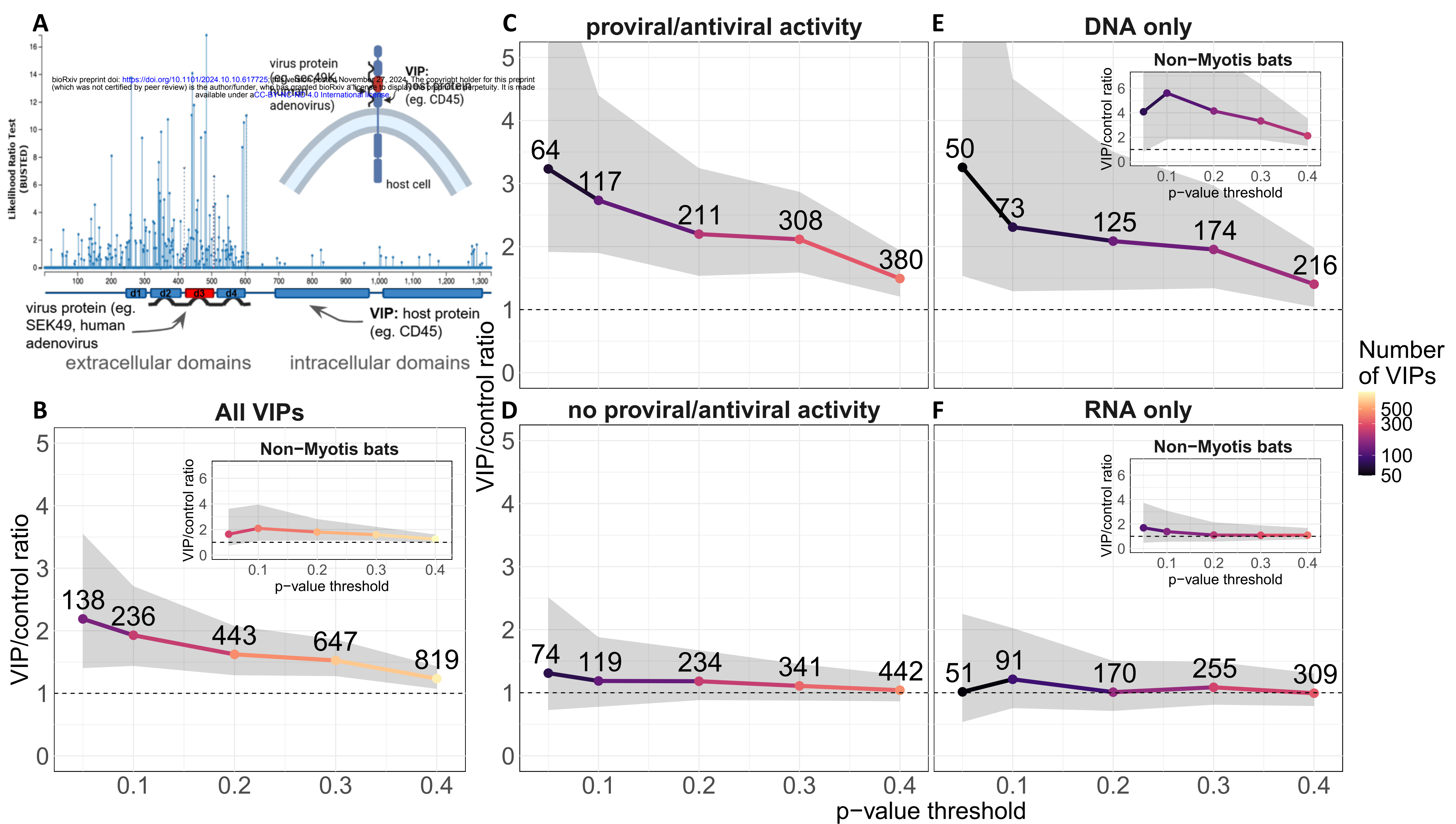


Figure 3: Selection in Nearctic *Myotis* is enriched for pleiotropic cancer resistance pathways. A) Left: phylogeny of Nearctic *Myotis*; **Right:** raincloud plot of omega values for all genes in each species since its most recent ancestor. The distribution of omega (ω) values for significant ($p \leq 0.05$ after multiple testing correction) genes and all genes is shown in color above the line. The 95% confidence interval and median for significant ω 's are represented by the black bar and circle, respectively; the overall 95% confidence interval and median are shown in grey below. Individual genes' ω 's are represented by colored points. Individual genes' omega values and grey, respectively. **Left inset:** Proportion of cancer-associated Reactome pathways among the top 100 pathways overrepresented among genes under selection at each node. Below, pie chart indicates expected proportion of pathways out of 100 that are cancer-associated after 1000 random samples. Nodes with proportions greater than the expected value with $p \leq 0.05$ using Fisher's exact test are indicated with an asterisk. **B)** Proportion of cancer-associated Reactome pathways among the top 100 pathways overrepresented among genes under selection across all nodes in a species' evolutionary history. **C)** Volcano plot of overrepresented pathways in Reactome among the union set of genes under selection across all nodes in the evolutionary history for *M. lucifugus*. **D)** Viability and Apoptosis fold-change in 5 bat species in response to different doses of neocarzinostatin, a potent inducer of DNA double-strand breaks. Points represent individual replicates normalized to each species' control, while bars represent mean $\pm$ 95% confidence intervals.

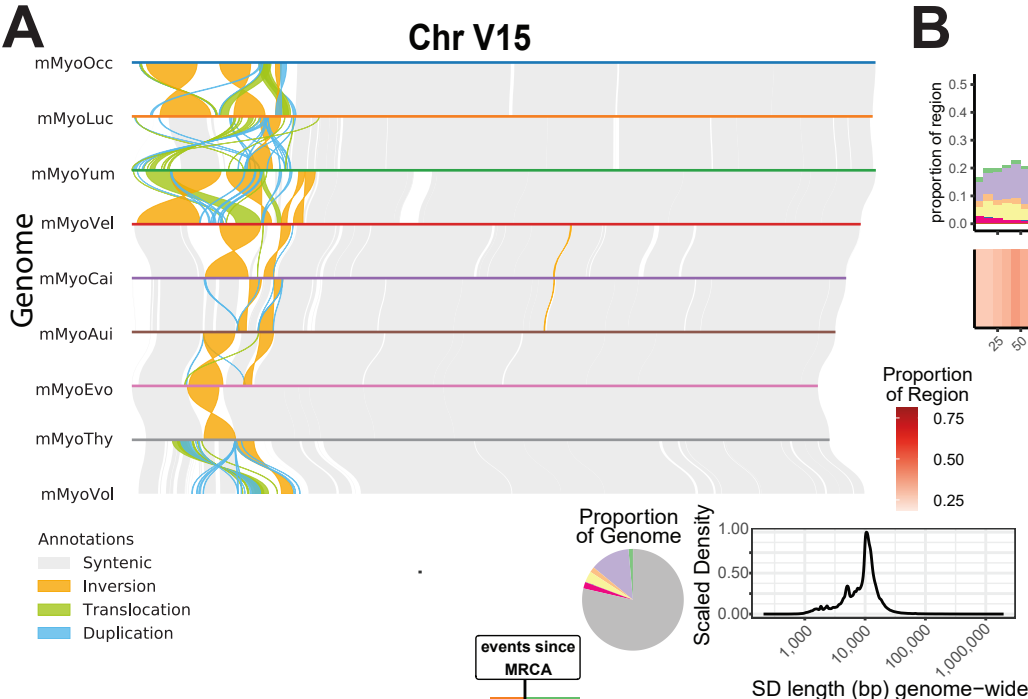
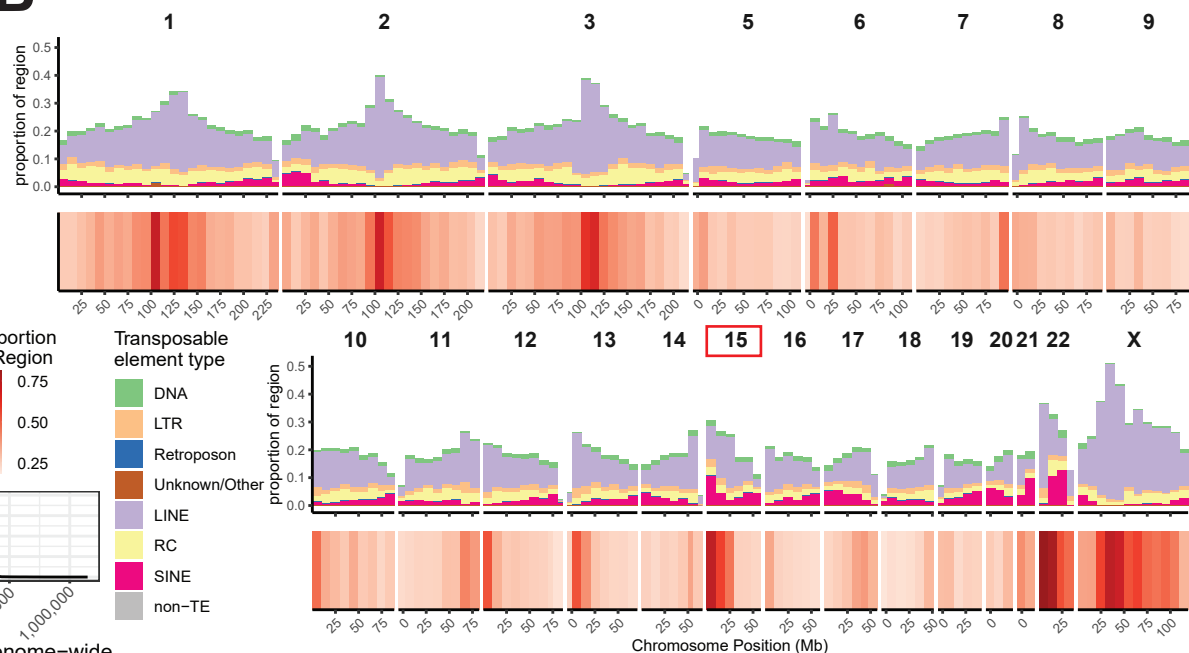
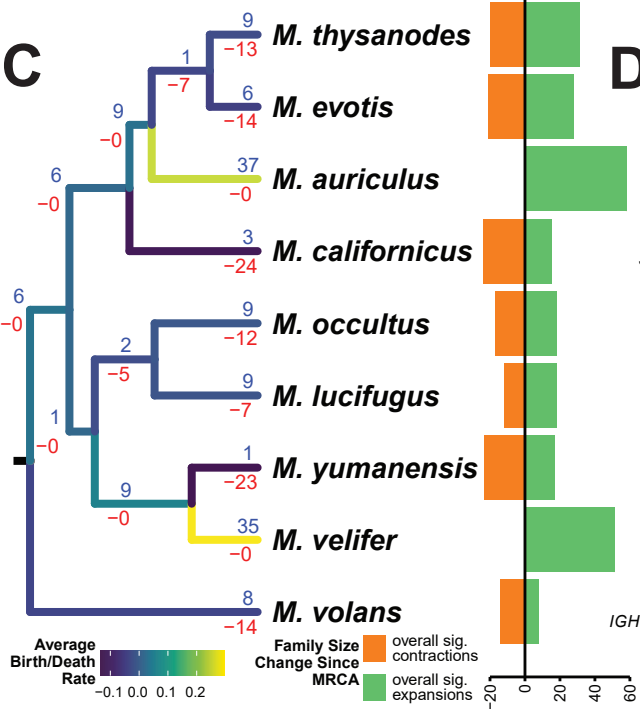
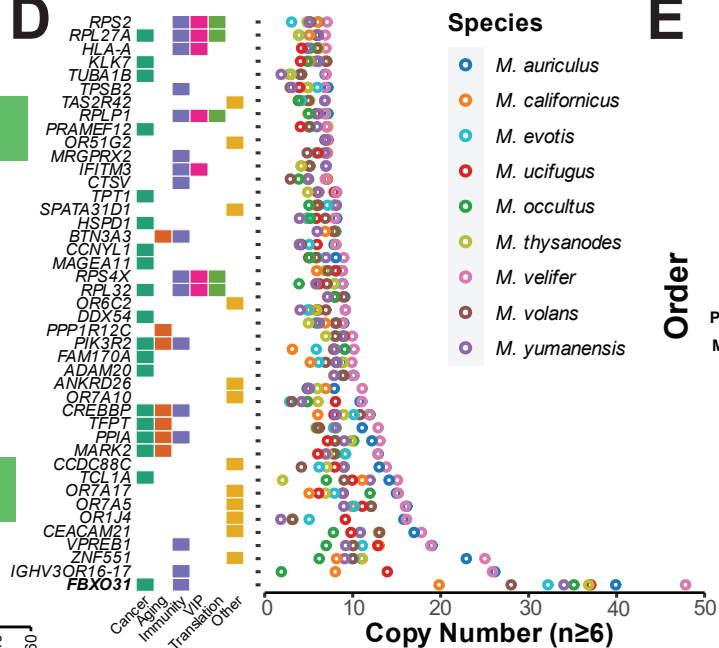
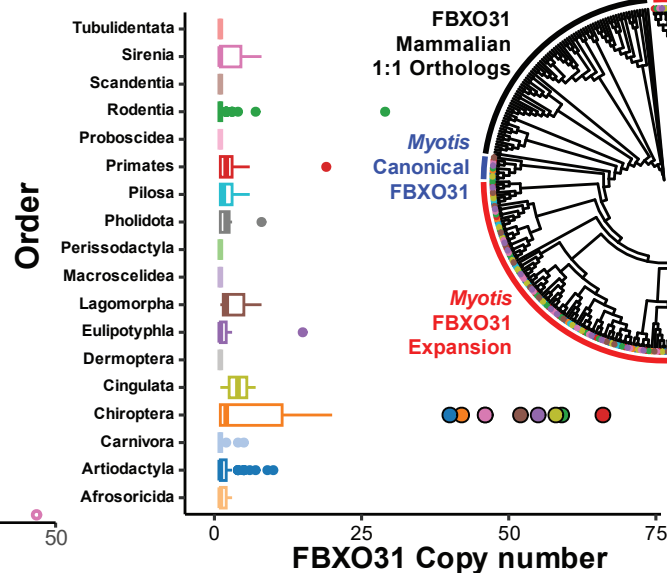
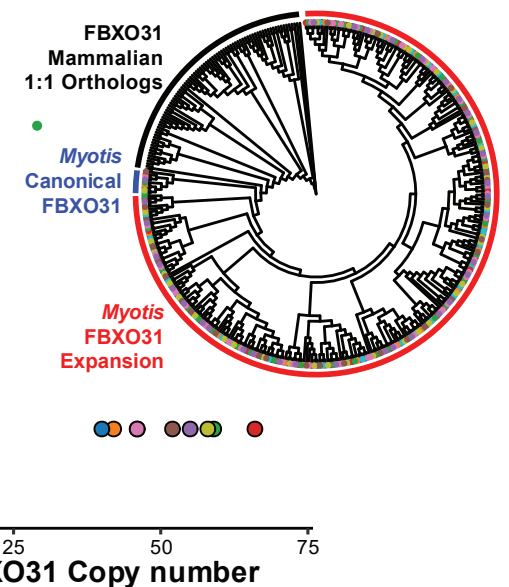
A**B****C****D****E****F**

Figure 4: Adaptation to DNA viruses, but not RNA viruses, is enriched in *Myotis* and other bats. A) Diagram of an example VIP, CD45: a host cell transmembrane receptor that interacts with the human adenovirus protein sec49K. Previous work has shown that the amino acids of CD45 that participate in this direct interaction are under strong positive selection, as indicated in the graph above the cartoon. **B-F)** Enrichment plots showing the ratio of positive selection in VIPs versus matched sets of control genes at different p-value thresholds. The solid line shows the median ratio; the color of the line, and the number above each point, represents the number of VIPs with significant BUSTED-MH p-values at the given threshold; the grey band represents the 95% confidence interval generated by bootstrapping sets of matched control genes. Inset plots show the same for all bats in this study excluding *Myotis*.

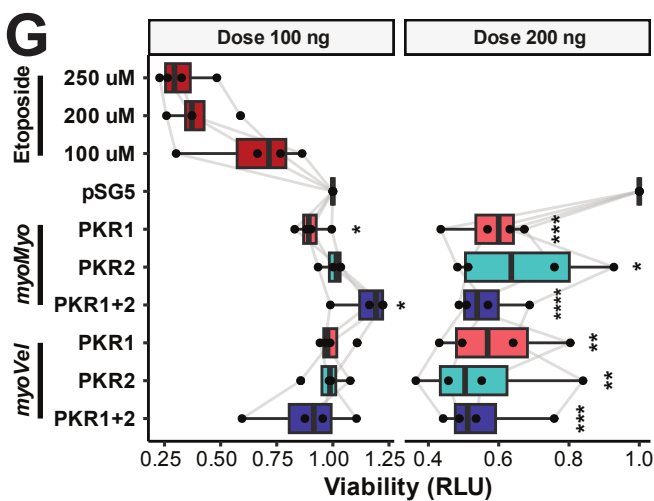
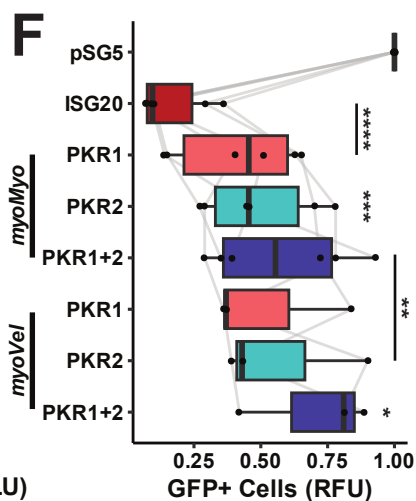
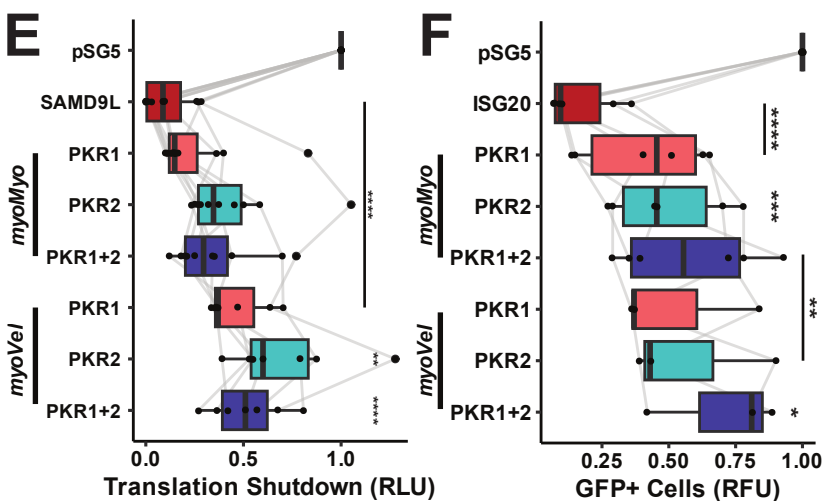
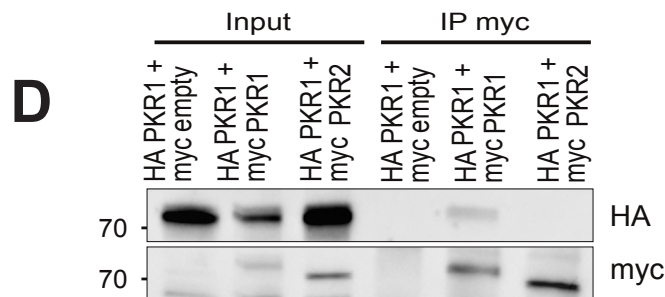
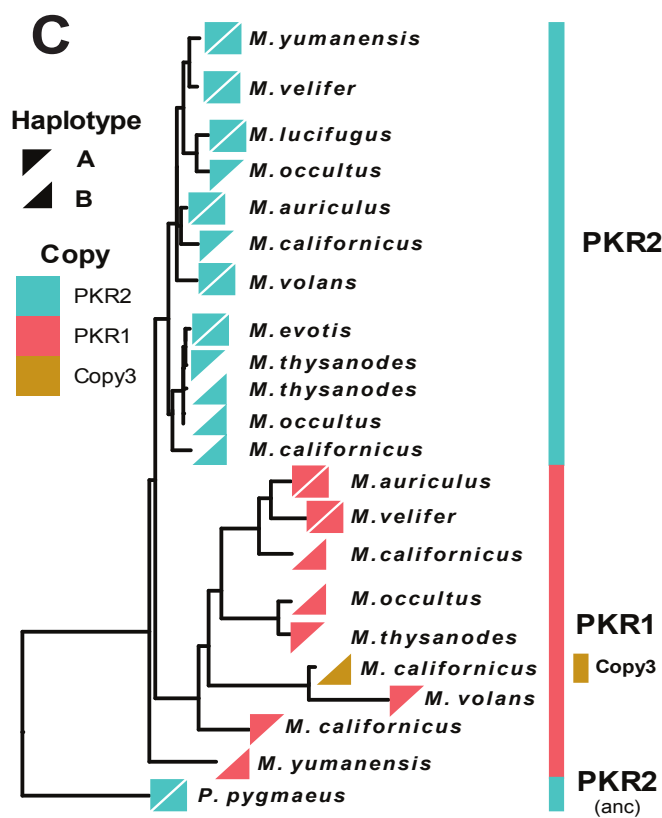
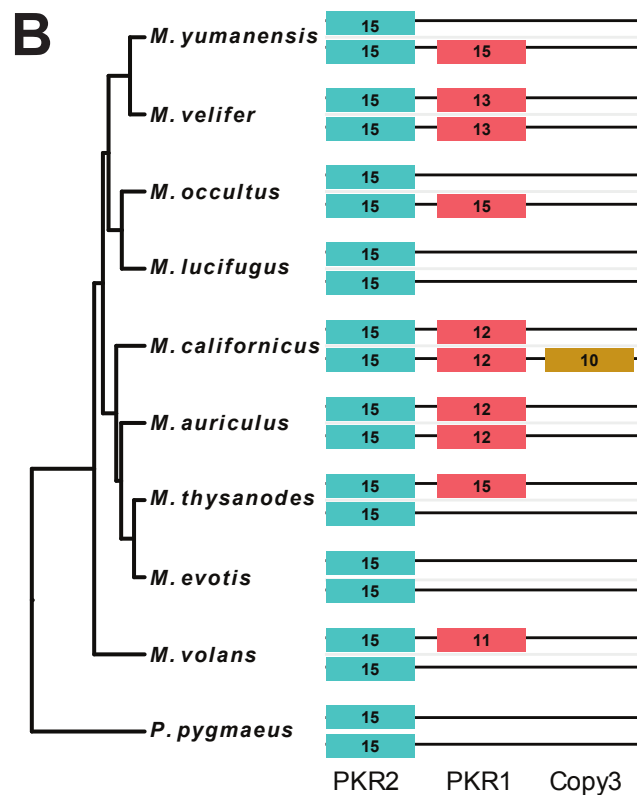
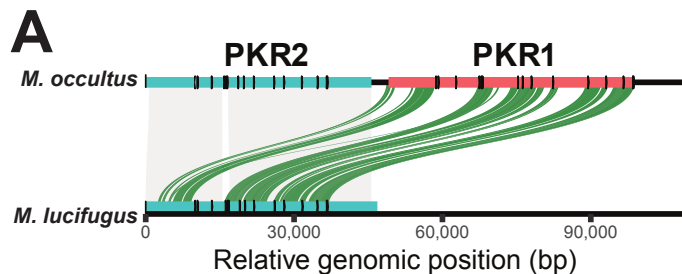


Figure 5: A varied structural variation landscape across 9 nearctic *Myotis* species. A) Synteny between *Myotis* species on chromosome V15, showing syntenic regions (grey), inversions (orange), translocations (green), and duplications (blue). Regions with high proportions of telomeric repeats were masked prior to alignment. **B)** Distribution of transposable elements and segmental duplications (red heatmap) in mMyoVel1. Pie chart indicates overall genome proportions of TEs; histogram represents the size distribution of segmental duplications genome-wide. **C)** CAFE results among our Nearctic *Myotis* relative to single-copy human orthologs. Phylogeny is colored by the estimated birth/death rate (λ) for all genes examined. Bar plot indicates the cumulative number of significant gain and loss events for each species. **D)** Per-genome copy numbers of all genes with over 6 copies in any Nearctic *Myotis* genome. Genes are classified into 5 categories (cancer, aging, immunity, VIP, translation, and “Other”) based on literature reviews on PubMed. **E)** Copy number estimates of *FBXO31* across 536 mammalian genomes identified using Reciprocal Best-Hit BLAT. **F)** Gene-tree reconciliation of *FBXO31* across mammals generated using GeneRax.

Figure 6: Evolutionary history and function of an actively segregating copy number polymorphism of *PKR* in *Myotis*. **A)** Structural comparison of the main *PKR* haplotypes in two species. Orthologous regions between the two haplotypes are indicated by grey bands, while syntenic duplications are indicated in green. Exons are annotated with black marks. **B)** Cartoon of the *PKR* locus in the two phased haplotype assemblies of each species. While *PKR2* is present across all haplotypes, *PKR1* and *PKR copy 3* are polymorphic within and across species. Each number indicates the number of exons per gene. **C)** Reconciled gene tree for *PKRs* across all haplotypes and species shown in **B**. Haplotype corresponding to the reference (A) and alternate (B) haplotype for each species are represented by upper- and lower-diagonal triangles, respectively. **D)** Co-immunoprecipitation (IP) of *PKR*-KO HeLa cells transfected with *M. myotis* HA-*PKR1* and either *M. myotis* myc-*PKR1*, *M. myotis* myc-*PKR2* or a myc-empty vector control. Proteins were pulled down with anti-myc beads and lysates from 5% input or IP samples were run on a western blot and stained for HA and myc. Representative of 3 independent experiments. **E-G)** Effect of *Myotis* *PKR* copy numbers: **E)** On luciferase translation, measured in Relative Light Units (RLU) and normalized to the empty pSG5 control; co-expression of *PKR1* and *PKR2* has an additive effect on cell translation shutdown (no synergy or dominant negative effects). Human SAMD9L-GoF is a positive control of translation inhibition²⁴⁹. **F)** On viral VSV infectivity, measured via flow cytometry as VSV-GFP-positive cells normalized to the control. Although all conditions restrict VSV, the expression of both *PKR1* and *PKR2* is not beneficial against VSV. ISG20 is a positive control of VSV-GFP restriction²⁴¹. **G)** On cell viability, normalized to the control. While no effect was observed at a low total dose of *PKRs*, at higher doses *PKRs* significantly reduce cell viability. Etoposide treatments are positive controls of cell death. For **E-G**, error bars indicate the means $\pm$ SEM for at least three independent experiments. Statistics, unpaired t-test of each condition versus control.