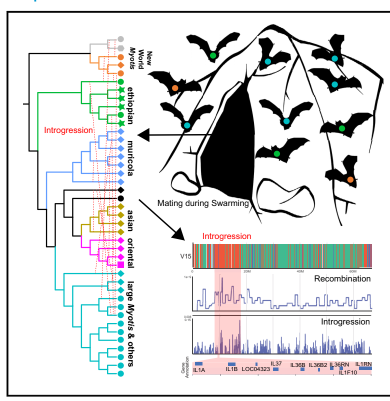
# Karyotypic stasis and swarming influenced the evolution of viral tolerance in a species-rich bat radiation

## **Graphical abstract**



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## In brief

In their recombination-aware phylogenomic analyses, Foley et al. demonstrate pervasive introgression among *Myotis* bats, likely promoted by their use of cryptic swarming behavior during mating. Introgression has acted to restrict the species tree to low recombining regions of the genome. In contrast, immune loci are enriched in introgressed regions.

## **Highlights**

- Phylogenomics reveals pervasive introgression among Old World Myotis bats
- A cryptic bat mating behavior called swarming promotes interspecific introgression
- Genomic regions of low recombination repel introgression and retain the species tree
- Immune loci are among those most frequently introgressed







## **Article**

# Karyotypic stasis and swarming influenced the evolution of viral tolerance in a species-rich bat radiation

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#### **SUMMARY**

The emergence of COVID-19 and severe acute respiratory syndrome (SARS) has prioritized understanding bats' viral tolerance. *Myotis* bats are exceptionally species rich and have evolved viral tolerance. They also exhibit swarming, a cryptic behavior where large, multi-species assemblages gather for mating, which has been hypothesized to promote interspecific hybridization. To resolve the coevolution of genome architecture and their unusual antiviral tolerance, we undertook a phylogenomic analysis of 60 Old World *Myotis* genomes. We demonstrate an extensive history of introgressive hybridization that has replaced the species phylogeny across 17%–93% of the genome except for pericentromeric regions of macrochromosomes. Introgression tracts were enriched on microchromosome regions containing key antiviral pathway genes overexpressed during viral challenge experiments. Together, these results suggest that the unusual *Myotis* karyotype may have evolved to selectively position immune-related genes in high recombining genomic regions prone to introgression of divergent alleles, including a diversity of interleukin loci responsible for the release of pro-inflammatory cytokines.

## **INTRODUCTION**

The observation that several bat species are tolerant to some viruses of significant concern to human and animal health<sup>1,2</sup> has led global public health initiatives to prioritize understanding the basis of this phenotype.3 Hypotheses for this enhanced tolerant, antiviral immunity include the coordinated evolution of bat flight, unique immune system adaptations, and bat/virus coevolution.<sup>2,5-9</sup> The hypothesized coevolution of bats and viruses<sup>1,10</sup> suggests that interrogating phylogenomic discordance may be instrumental in understanding the genetic basis of viral tolerance. Resolved species-level phylogenies may also aid in monitoring future disease emergence through the identification of related species that host similar viruses. 11 Myotis bats comprise the most species-rich bat genus and have an almost global distribution. 12 However, their phylogenomic relationships lack consensus, most likely due to some combination of gene flow and incomplete lineage sorting during their evolution, the resolution of which has been hampered by limited surveys of genomic variation. 12-15

Adaptive introgression of immune loci is frequently observed between species pairs across the mammalian phylogeny. 16-20 Additionally, mating strategy has been shown to result in systematic differences in immune responses, conferring enhanced protection again sexually transmitted diseases in promiscuous primates and adenovirus infection in promiscuous bats.<sup>21,22</sup> Temperately distributed bats, including numerous Myotis species, exhibit autumnal swarming, a cryptic behavior where large, multi-species assemblages gather for mating.<sup>23</sup> During the swarming period between August and October, thousands of bats from multiple bat genera form nightly gatherings at cave entrances.<sup>24</sup> At these transient meetings, bats exhibit intense flight activity, chasing, and increased communication, including echolocation and human-audible social calls, which facilitate interspecific communication. 24-26 Additionally, these assemblages are typically 80% male biased, 27,28 suggesting intense competition for mates.<sup>29</sup> Although the phenology of swarming varies by species, elevation, and seasonal timing, <sup>26,30,31</sup> many bat species and genera still overlap at swarming sites.



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Initially, the purpose of swarming was thought to be locating suitable hibernation sites and orienting juveniles to their whereabouts.<sup>25</sup> Later, swarming was shown to facilitate gene flow between isolated populations of the same species. 32 Most recently, genetic analyses of microsatellites have suggested that swarming may promote interspecific hybridization among Myotis species, but the strength and consistency of this process during the evolution of the genus is unknown.<sup>23</sup> We hypothesized that an excess of historical gene flow, facilitated by swarming, has contributed to the acquisition and maintenance of novel and divergent genomic diversity. Some of this novel variation may include immune gene loci that enhance viral tolerance.

#### **RESULTS**

#### Whole-genome phylogenies

To test this hypothesis, we sequenced and aligned 60 new Myotis genomes representing 42 species to a chromosome-level long-read reference genome assembly for Myotis myotis (Tables S1 and S2).33 We improved this assembly by initially using the associated gene annotation and previously generated ZooFISH maps<sup>34</sup> to predict the location of unassigned scaffolds.

The original Hi-C data<sup>33</sup> were remapped to the assembly, and assignments were made if contacts in the Juicebox Analysis Toolkit<sup>35,36</sup> matched predictions made by the ZooFISH and gene annotation analysis. In this way, we placed 37 scaffolds that were previously chromosomally unassigned (Table S3). Notably, these contained large clusters of immune-related genes (e.g., major histocompatibility complex [MHC], immunoglobulin heavy-chain genes, and interleukin receptors) (Figure S1), which have previously been implicated in adaptive introgression. 37,38 Our analysis localized these scaffolds predominantly to chromosome ends, which typically possess elevated recombination rates (Figure S2).

To establish a baseline phylogeny for the Old World Myotis, we sampled phylogenomic signal across the chromosome alignments, analyzing 1,463,340 single-nucleotide variants (SNVs) with site pattern frequency-based coalescent methods in SVDquartets.<sup>39</sup> The same dataset was also concatenated to generate a maximum likelihood tree using IQ-TREE 2.40 The concatenation- and coalescent-based SNV phylogenies were highly congruent. Both trees identified six primary Old World Myotis clades (Figure 1A) and differed only in the positions of the whiskered and muricola clades and a few rogue taxa, including M. annatessae and M. welwitschii (Figures 1A and S3). By contrast, there was a considerable degree of phylogenomic conflict between the nuclear and mitogenome topologies. Widespread clade switching occurred for several focal species (Figures 1A and 1B), similar to results from small-scale analyses of nuclear and mitochondrial gene fragments (Figures 1B, 1C, and S4).12 Resolving the basis for this discordance was critical to distinguishing species relationships and associations that may arise due to gene flow and other types of phylogenetic noise.41-45

Mitonuclear discordance is generally predictive of gene flow in other mammals, although there are some exceptions. 46,47 Among species with a history of post-speciation gene flow, it has been demonstrated that the species tree is not the dominant

phylogenomic signal across the genome. 42,43 However, majority-rule phylogenomic approaches assume that the most frequent topology represents the species tree and therefore often mask underlying signatures of discordance. 41,43,44 In several hybridizing radiations, episodes of post-speciation gene flow have erased the historical branching events of the species tree from large swaths of the genome. 42,43,45 Therefore, we implemented a recombination-aware locus tree approach, which has proved crucial to distinguishing the phylogenetic signal of speciation from introgression.

All surveyed Myotis species shared an identical G-banded karyotype (2n = 44) except M. annectans, with (2n = 46),  $^{34,48}$ an interesting trait given that animal radiations with histories of post-speciation gene flow typically possess conserved karvotypes. 41-43,49 The Myotis karyotype is also unusual among mammals, comprising three large autosomes (~228-217 Mb) and 18 much smaller chromosomes (~108–14 Mb) (Figure 2A),<sup>34</sup> which superficially resemble macro- and microchromosomes commonly observed in avian and reptilian karyotypes. 50,51 We used ReLERNN, 52 a deep learning approach, to infer a sex-averaged recombination map for M. myotis (Figure S5). A comparative analysis indicated the recombination landscape is conserved between M. brandtii and M. myotis (Figure S6). Therefore, we proceeded to use the M. myotis map as a representative for all Myotis. Low recombining regions were narrowly restricted to the pericentromeric regions of the largest macrochromosomes (V1, V3, and V5) and the X chromosome (chrX) (Figure 2B). The highest recombination rates were predictably found on the smallest autosomes, which also exhibit elevated gene densities and GC content (Figure 2C), similar to the microchromosomes of birds and some reptiles. 50,51

## **Recombination-aware phylogenomics**

To investigate phylogenomic variation across the recombination landscape, we computed 36,264 maximum likelihood (ML) locus trees for 60 species from the whole-genome dataset using the GTR + GHOST model of rate heterogeneity<sup>53</sup> in IQ-TREE 2 from 50 kilobase (kb) alignment windows covering 91% of the M. myotis genome. The GHOST model was chosen given that recent studies have demonstrated that some tests of introgression are highly sensitive to rate variation across clades.<sup>54</sup> Ten subtrees (Table S4) were pruned from these datasets to determine the genomic stability of relationships among the principal clades and between individual species whose positions were difficult to resolve in the whole-genome and mitogenome analyses (Figures 1A and 1B). These analyses revealed widespread locus tree discordance otherwise masked in the whole-genome concatenation and coalescence phylogenies (Figures 3 and S7-\$15). Locus tree frequencies varied markedly between chrX and the autosomes and between high and low recombining regions (Figure S16).

Phylogenomic relationships within the whiskered bats and most Myotis clades exhibited asymmetric tree topology frequencies, considered one of the clearest indicators of the presence of gene flow (Figures 3 and S16).55 Previous studies have shown that in the presence of gene flow, the species tree is enriched in low recombining regions of the genome (particularly pericentromeric regions and the chrX) (Figure S17), while trees

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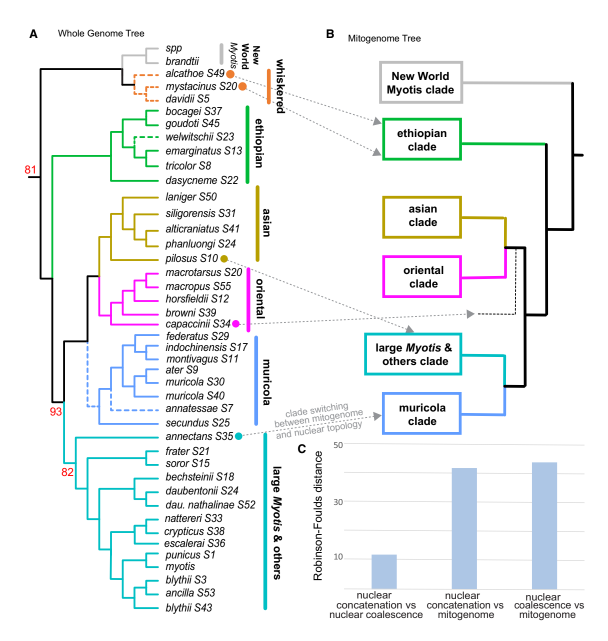


Figure 1. Clade switching between the whole-genome nuclear and mitogenome topologies

(A) Maximum likelihood cladogram from ~1.5 million genome-wide SNVs, showing only one sample per species where multiple exist. Bootstrap support is 100% for all nodes unless indicated otherwise. Dashed branches show which relationships differ between concatenation and coalescent analyses of the same dataset. Concatenation bootstrap values are shown in red.

(B) The mitogenome phylogeny is illustrated to highlight, via gray dashed arrows, clade switching relative to the nuclear phylogeny for the species indicated with colored dots.

(C) Robinson-Foulds (RF) distances between the nuclear and mitogenome topologies.

reflective of gene flow will be enriched in high recombining regions. 41,43-45 Using a combined approach to infer the species tree, our distributions of phylogenomic signal were validated by three statistical measures that distinguish introgression from incomplete lineage sorting (ILS): (1) whole-genome D-statistics<sup>56</sup> (Table S5), (2) windowed f(D) statistics (Figures 3C, S16, and S17), and (3) QuIBL (Table S6). While D-statistics can yield insights into the genome-wide presence of introgression, f(D) can provide information on more precise genomic locations of introgression. QuIBL takes a distribution of internal branch lengths from a set of locus trees and uses a coalescence-based model to infer the proportion of locus trees derived from introgression or ILS. For the whiskered clade, topology 1 (t1) is most frequent in low recombining regions and in regions with reduced introgression, while t2 becomes more frequent in introgressed, high recombining regions (Figure 3B). Similar observations across numerous eukaryotic clades have increasingly demonstrated the predictive power of using regions of low



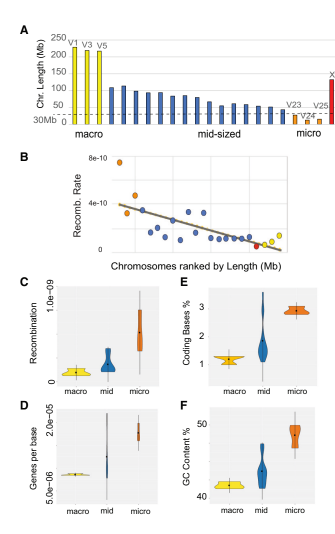


Figure 2. An unusual mammalian karyotype

(A) The dashed line indicates 30 Mb, which previous studies have used as a cutoff to define microchromosomes.50

- (B) Chromosome length is inversely related to recombination rate.
- (C-F) Comparison of recombination rate (C), genes per base (D), coding bases (E), and GC content (F) between macro-, mid-sized, and microchromosomes.

recombination to infer the species tree, as they are depleted of ILS and gene flow. 41,43-45,57

Remarkably, the inferred species tree uniting the whiskered clade with New World species was restricted to less than  $\sim$ 60% of the genome, clustering in the pericentromeric regions of the three macrochromosomes, and chrX, which possessed the lowest rates of recombination (Figures 3 and S16). This pattern, as well as patterns of alpha and internal branch length (ibl) parameter distributions along chromosomes (Figure 3C), was mirrored in a highly predictable manner across other Myotis clades (Figure S18) where the species tree was commonly observed in less than 50% of the genome. Crucially, there are numerous differences between the recombination-aware inferred species tree (Figure 4A) and the whole-genome concatenation and coalescence analyses inferred from SNVs. D-statistics show that analyses of the genome-wide SNV dataset return

the most frequent introgressed topology for the relationships between the muricola, large myotis, Asian, and Oriental clades, the respective positions of M. annatessae, M. capaccinii, and M. pilosus, and the relationship among M. pilosus, M. capaccinii, and M. laniger (Figure 4B; Table S5). D-statistics and QuIBL (Table S6) also show that clade switching between nuclear and mitogenome topologies (Figure 1B) is due to historical interclade introgression likely due to range overlap between Asian-distributed M. annectans and M. indochinensis (Figure 4).

Previous studies have demonstrated that whole-genome D-statistics maintain sensitivity when there is up to 20% genetic distance among the ingroup species in the test<sup>62</sup> and are also relatively robust to the genetic distance between the outgroup and ingroup species. In our analysis, the genetic distance between component species of Old and New World clades is  $\sim$ 16%, which spans the deepest divergence in our dataset. In conclusion, our findings support a scenario for Old World Myotis evolution where a long history of interspecific hybridization involving species from multiple clades has resulted in massive depletion of the species tree across the arms of the macrochromosomes increasing toward telomere ends, with a near-complete replacement by introgression signatures on the smallest microchromosomes.

#### A link between introgression and swarming behavior?

Introgression is exceptionally pervasive among Myotis bats, even when compared to other mammalian clades with histories of ancient introgressive hybridization. 43,49,63 This may be due to the use of swarming behavior in many Myotis species. Swarming behavior is characterized by multi-species, male-biased mating assemblages often numbering from hundreds to thousands of individuals.<sup>23</sup> Many records of species present at swarming sites come from predominantly European studies and overlap our sampling (Figure 5A; Table S7). Therefore, we pruned subtrees derived from the 50 kb windows from the whole-genome 60 species sliding-windows dataset (Table S4) to test the hypothesis that hybridization is common among species within and between clades recorded at swarming sites (Figure 4B). Asymmetry in genome-wide frequency and D-statistics estimated among swarming species indicated that introgression was the dominant driver of phylogenetic discordance in these analyses (Figures 5B, S19, and S20). Additionally, we show that introgressed topologies with the highest f(D) values are the most enriched on microchromosomes (Figure S21). Introgression was observed not only among species within clades but also between species from divergent clades (Figure 5B), like the large Myotis, whiskered, and New World clades. Note, due to our sampling scheme, we could only infer the presence of historical introgression among Myotis species. Population-level data or data sampled from species found contemporaneously at swarming sites would be required to accurately determine the prevalence of ongoing hybridization at swarming sites, which has been documented for some of the species we sampled. 15,63,64

The Ethiopian clade (Figure 1A) contains important contrasts to other predominantly swarming Myotis clades. At least one species lacks swarming behavior: M. bocagei employs a harem system where a single male and 6-8 females gather in a banana leaf for mating. 65 This behavior should minimize the probability of introgression. Although not all mating strategies employed by

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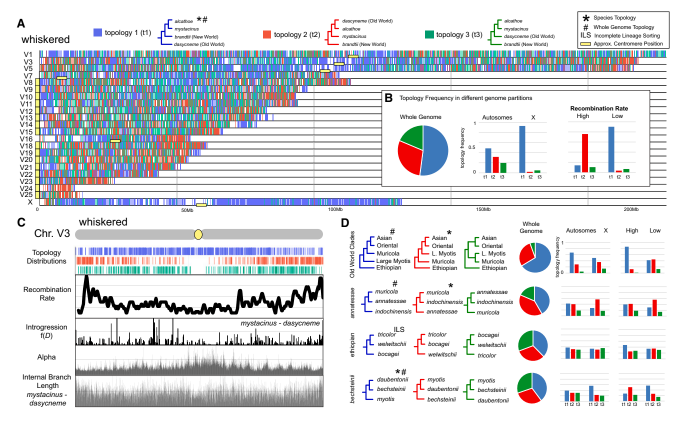


Figure 3. Genome-wide distribution of phylogenomic discordance across the Old World Myotis radiation

(A) Distribution of phylogenomic signal for the whiskered clade visualized using Tree House Explorer<sup>61</sup> in 50 kb locus alignment windows across the genome. Vertical bars along each chromosome are color coded to depict the distribution of the most frequent topologies.

(B) Comparison of the frequency of t1, t2, and t3 across the whole genome (left) and between the autosomes and the X (middle) and between high and low recombining regions of the genome (right).

(C) Whole-chromosome view for ChrV3 illustrates local relationship between recombination rate, f(D), the alpha parameter of the gamma model, and the internal branch length (ibl) for relationships between M. mystacinus and M. dasycneme. Note the depletion of topologies associated with introgression (t2 [red], t3 [green]) in the low recombining region at the center of the chromosome.

(D) Distributions of phylogenomic signal for additional Myotis clades in relation to chromosome type and recombination rate (reference the key in B).

the Ethiopian clade species are well characterized, the roughly equal frequency of locus trees for the clade are consistent with ILS, and not introgression, being the dominant source of phylogenomic discordance (Figure 3C). This is notable given that *M*. bocagei, M. tricolor, and M. welwitschii are sympatric over much of their current distributions. Conversely, two members of the Ethiopian clade, M. emarginatus and M. dasycneme, have a European distribution and are frequently reported at swarming sites (Figures 4A and 5A). Whole-genome D-statistics consistently show significant ancient introgression signatures between M. emarginatus and numerous European distributed species, yet there is minimal evidence of ancient introgression with M. bocagei (Figure 5C). In summary, these contrasting phylogenomic patterns between swarming and non-swarming species support the hypothesis that swarming sites have acted as crucibles of genetic exchange.

## Introgression at immune loci

We hypothesized that viral tolerance in Myotis bats may be enhanced by a long history of immune gene introgression involving species from multiple phylogenetic lineages interacting at swarming sites. 66 Introgression from Neanderthals into modern humans has had measurable impacts on the expression of genes associated with immunity. 16,67,68 Most notable among these is differential expression of an OAS1 isoform, which has previously been shown to contribute to differential susceptibility and severity of COVID-19 outcomes in humans.<sup>69</sup> To further examine if the apparent genome-wide intersection between introgression and immune-related processes has had any potential effect on gene expression in Myotis bats, we used a database of differentially expressed genes from a study that performed viral challenge experiments on *M. daubentonii* cell lines.<sup>70</sup> These cells were treated with a modified Rift Valley fever virus or supplemented with interferon, and gene expression was compared 6 h before and after treatment. 70 Our results demonstrate that genes that were differentially expressed in response to viral challenges in M. daubentonii were among the most frequently introgressed across microchromosomes (Figure S22) and were also significantly introgressed genome-wide across numerous Myotis clades (Figure S23). Notably, these include



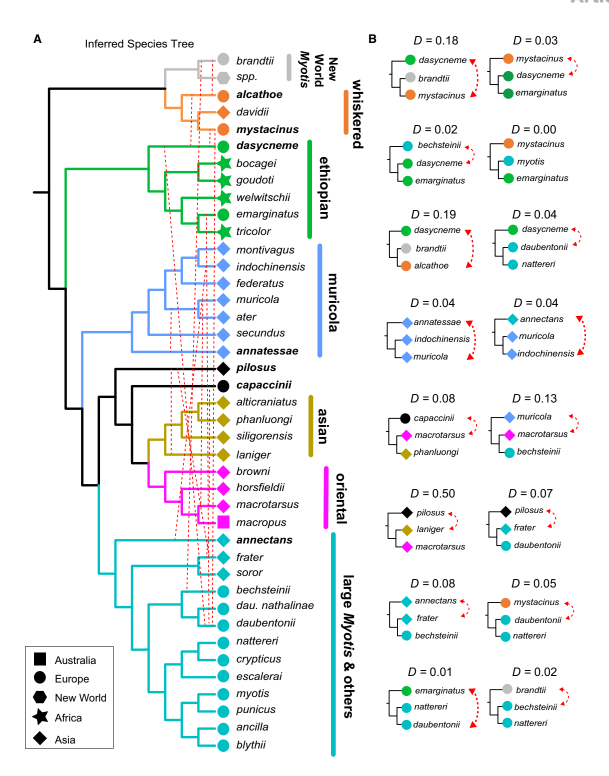


Figure 4. Discordance due to gene flow

(A) Species tree inferred from recombination-aware phylogenomic approaches. Crucially, our analyses show that standard whole-genome analyses in Figure 1 do not recover the species tree and instead depict relationships consistent with introgression for some species (e.g., M. capaccinii). Dashed lines indicate where within and between introgression is detected. Note that we illustrate introgression only for the analyses presented in (B) for the purpose of clarity. Additional instances of genome-wide introgression are depicted in Figure 5.

(B) Whole-genome-based D-statistics generated using ABBA-BABA tests in ANGSD illustrate significant evidence for introgression (Z score > 3) in (A). Trees are represented top to bottom in the format (H1, (H2, H3), O), where the outgroup O (S. latirostris) is not shown. See Table S5 for detailed results.



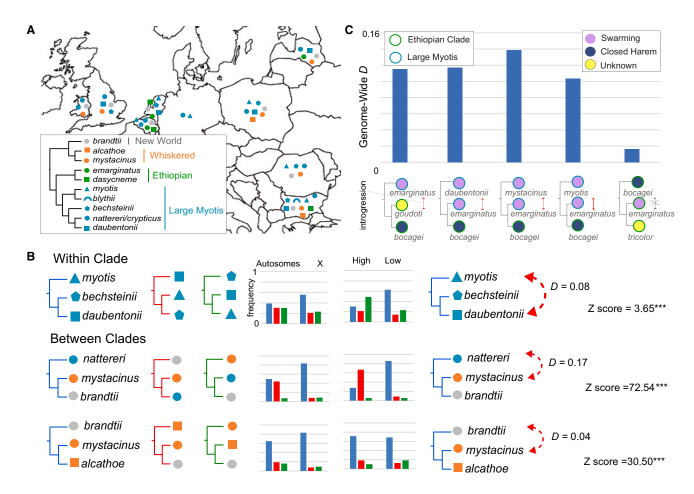


Figure 5. Introgression occurs among species that co-occur at swarming sites

(A) Species assemblages reported at swarming sites throughout Europe (Table S6). The tree inset shows the species tree and clade membership of each species. (B) Examples of within and between clade introgression. Variation in topological frequency is observed between autosomal, X, and high and low recombining genomic partitions. Whole-genome D-statistics demonstrate phylogenomic conflict is due to introgression.

(C) Comparisons of the genome-wide D-statistic for species representing the large myotis clade, with a promiscuous mating system, and M. bocagei from the Ethiopian clade, which employs a conservative (closed harem) mating system. The same analysis was repeated using M. yumanensis as the reference genome to demonstrate that our results are not confounded by reference bias (Figure S25; Table S5). Trees for introgression tests are represented top to bottom in the format (H1, (H2, H3)), O), where the outgroup O (S. latirostris) is not shown. See Table S5 for detailed results.

microchromosome-linked genes OAS1, OAS3, and OASL, which exhibit up to a 6-fold change in expression in response to viral challenge. 70 Because one of the common outcomes of interspecific hybridization is altered gene expression, 68,71-73 these results suggest that frequent hybridization-induced modulation of immune gene expression in bats might be adaptive and is worthy of further investigation in natural and experimental populations.

Recombination acts to break down introgressed haplotype blocks over time. While long blocks representing more recent introgression events are expected to be rare in our dataset due to low population genomic sampling, one of the largest haplotypes spans  $\sim$ 10 Mb on the subtelomeric region of V15 and possesses an introgressed topology uniting M. myotis and M. daubentonii (Figures 6 and S24). Measures of recombination rate, ibl, and f(D) were all elevated inside the block relative to the remainder of the chromosome. A STRING analysis of this haplotype block identified a significant enrichment (false discovery rate [FDR] < 0.05) of genes associated with cytokine receptor activity, the interleukin-36 pathway, and the interleukin-1 family signaling pathway (Table S8). IL36 is notable as a member of the IL1 superfamily of pro-inflammatory cytokines, which help regulate the adaptive and innate immune systems.<sup>74</sup> IL36 promotes antiviral immunity by increasing cellular sensitivity to interferon<sup>75</sup> in human and mouse models to herpes simplex virus<sup>75</sup> and a host of other viruses.74

## **DISCUSSION**

Collectively, our results support a scenario where swarmingmediated interspecific hybridization can impact antiviral immune responses in bats. High rates of introgression have also been reported in animal clades where multiple, closely related species are present during mating, including species of Stenella and



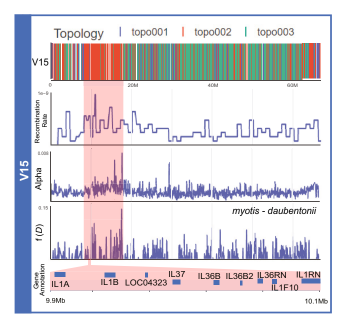


Figure 6. Evidence for a potential link between immunity and the recombination landscape in Myotis bats

Distribution of phylogenomic signal, recombination rate, ibl, and f(D) values for chromosome V15 for the bechsteinii clade. A large introgressed block for myotis and daubentonii (Figure S24) is highlighted in red and is significantly enriched for IL-36 and IL-1 signaling pathways (FDR < 0.05) (Table S8).

Tursiops dolphins<sup>76–78</sup> and Anopheles mosquitoes.<sup>42,79</sup> Swarming has specifically contributed to the introgression-mediated evolution of insecticide resistance at several loci in anopheline mosquitoes. 80,81 Promiscuous mating behaviors may frequently evolve to promote adaptive interspecific hybridization, providing access to divergent genomic variants and/or quantitative variation in gene expression as a rapid means to counter frequent and novel pathogen exposure.<sup>71</sup> The extraordinarily large number of mammalian congeners present in large colonies of bats (sometimes thousands of individuals) and mating during swarming would impose a unique burden on mucosal viral immunity. Under this model, frequent introgression at swarming sites would conceivably afford myriad opportunities for adjusting viral symbiont management and/or defense mechanisms using pooled immunogenomic resources from species across the genus. As such, swarming behavior may have provided a singular opportunity for the evolution of a novel mechanism for viral management/symbiosis through introgression biased toward chromosome termini enriched for immune loci. Such innovations are not unprecedented in nature. Radical and profound innovations in immunogenetic diversification have previously evolved from horizontal genetic transfer in other taxa (Table S9)-for example, the CRISPR-Cas9 prokaryotic adaptive immune system from IscB and IS200/IS605 transposons82 and the somatic cell recombination capability of immunoglobulin and T cell receptor genes via domestication of ProtoRAG transposase83 in an early chordate. Further investigations into understanding how karyotype evolution, the recombination rate landscape, and the introduction of divergent genomic variants via introgression and/or ILS intersect to hone bats' genomes for viral and cancer tolerance may yield novel innovations for viral immunotherapeutics and even oncology.84 Finally, our results suggest that swarming sites may merit recognition as special sites of conservation given the unique evolutionary processes to which they play host. 23,28,31,85

## Limitations of the study

All data were mapped to M. myotis, which is part of the ingroup in our analysis, and so may lead to spurious inference of introgression due to reference bias. To account for this potentially confounding factor, we also mapped the data to the long-read M, vumanensis genome assembly<sup>86</sup> from the New World clade. Although this analysis confirmed our initial results for the Old World ingroup, ideally a closely related outgroup, which is roughly equidistant to all ingroup species, should be used to mitigate any potential reference bias and data loss due to mapping to a divergent reference genome. Our analysis was also constrained to the detection of historical introgression. Although some studies indicate that contemporary introgression occurs at swarming sites,<sup>23</sup> our dataset was not selected to formally test this hypothesis. Further comparative phylogenomic analyses of species with conservative and promiscuous mating behaviors would be useful to fully characterize the contribution of swarming behavior to introgression in bats.

## **STAR**\*METHODS

Detailed methods are provided in the online version of this paper and include the following:

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## SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j. xgen.2023.100482.

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## **Article**



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#### **AUTHOR CONTRIBUTIONS**

Conceptualization, W.J.M. and N.M.F.; data curation, N.M.F. and K.R.B.; investigation, N.M.F., A.J.H., K.R.B., M.F.C., and W.J.M.; software, A.J.H.; project administration, W.J.M.; samples, M.R., S.J.P., and E.C.T.; writing original draft, W.J.M. and N.M.F.; writing - review & editing, all authors.

#### **DECLARATION OF INTERESTS**

The authors declare no competing interests.

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## Article



## **STAR**\***METHODS**

## **KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Deposited data		
HLmyoMyo6	Jebb et al. <sup>33</sup>	GCA_014108235.1
Myotis myotis Hi-C data	Jebb et al. <sup>33</sup>	SRR11732436
Myotis myotis ZooFISH map	Volleth et al. <sup>34</sup>	N/A
Myotis yumanensis reference genome	Curti et al.87	mMyoYum1.0.hap1
Myotis myotis SRA data	Zoonomia Consortium <sup>88</sup>	SRS3678524
List of differentially expressed genes from a study that treated <i>Myotis daubentonii</i> cell lines with a modified Rift Valley Fever virus or supplemented with IFN to measure response to a viral challenge	Holzer et al. <sup>69</sup>	Supplementary Material
WGS phylogenomic data generated in this study	This study	PRJNA1035163
WGS data used to generate Myotis myotis recombination map	This study	PRJNA1035163
WGS data used to generate Myotis brandtii recombination map	Laine et al. <sup>89</sup>	SRR18788643, SRR18788644, SRR18788646, SRR18788647
Myotis myotis recombination map	This study	https://doi.org/10.5281/zenodo.7044479
Myotis brandtii recombination map	This study	https://doi.org/10.5281/zenodo.7044479
Alignments	This study	https://doi.org/10.5281/zenodo.7044479
Newick Tree files	This study	https://doi.org/10.5281/zenodo.7044479
Software and algorithms		
Trim Galore! v0.6.5	Krueger et al. 90	https://github.com/FelixKrueger/TrimGalore
bwa-mem v0.7.17	Li et al. <sup>91</sup> ; Li <sup>92</sup>	https://github.com/lh3/bwa
Qualimap	Garcia-Alcalde et al. <sup>93</sup>	http://qualimap.conesalab.org/
Samtools v1.9	Li et al. <sup>94</sup>	https://github.com/samtools/samtools
GATK v4.1.2	McKenna et al. 95; DePriesto et al. 96; Poplin et al. 97; Van der Auwera et al. 98	https://github.com/broadinstitute/gatk
ANGSD v0.916	Korneliussen et al. <sup>56</sup>	http://www.popgen.dk/angsd/ index.php/Installation
SOAPdenovo2 vr242	Luo et al. <sup>98</sup>	https://github.com/aquaskyline/ SOAPdenovo2
Geneious Prime 2019.04	Dotmatics	https://www.geneious.com/
RYcode	Braun et al. <sup>99</sup>	https://github.com/ebraun68/RYcode)
IQtree2 v2.2.1	Minh et al. <sup>40</sup>	https://github.com/iqtree/iqtree2
Quast v5.2.0	Gurevich et al. 100	https://github.com/ablab/quast
Bcftools v1.14	Danecek et al. <sup>101</sup>	https://github.com/samtools/bcftools
VCFtools v1.16	Danecek et al. 102	https://vcftools.sourceforge.net/
ReLERNN	Adrion et al. <sup>52</sup>	https://github.com/kr-colab/ReLERNN
SVDquartets	Chifman and Kubatko <sup>39</sup>	https://paup.phylosolutions.com/
SNP-sites v2.5.1	Page et al. <sup>103</sup>	v2.5.1
PAUP* v4a168	Posada et al. 104	https://paup.phylosolutions.com/
trimAl v1.4.1	Capella-Gutierrez et al. 105	https://vicfero.github.io/trimal/
Tree House Explorer (THEx)	Harris et al. <sup>61</sup>	https://github.com/harris-2374/THEx
R v4.2.2	R Core Team <sup>106</sup>	https://www.r-project.org/
f(D)	Martin et al. <sup>87</sup>	https://github.com/simonhmartin/ genomics_general

(Continued on next page)



Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
QuIBL	Edelman et al.41	https://github.com/miriammiyagi/QuIBL
STRING v12.0	Szklarczyk et al. <sup>87</sup>	https://string-db.org/
Dfoil	Pease and Hahn <sup>88</sup>	https://github.com/jbpease/dfoil
Critical commercial assays		
Qiagen Puregene kit	Qiagen	Cat. No./ID: 158063
NEBNext Ultra II DNA Library Prep Kit for Illumina	New England Biolabs	E7805L
NEBNext Multiplex Oligos for Illumina	New England Biolabs	E7600S

#### RESOURCE AVAILABILITY

#### **Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, William Murphy (wmurphy@cvm.tamu.edu).

## **Materials availability**

This study did not generate new unique reagents.

## Data and code availability

All code used to generate the analysis is publicly available and are cited in the text. All short-read data generated as part of this study have been submitted to the NCBI Short Read Archive under the following BioProject PRJNA1035163. Recombination maps for M. myotis and M. brandtii, whole-genome SNP datasets used for concatenation and coalescence analyses, a FASTA-formatted version of the updated M. myotis genome assembly, and whole-genome alignments are deposited in the following Zenodo repository: https://doi.org/10.5281/zenodo.7044479. The updated M. myotis genome assembly can be found under BioProject PRJNA628559.

## **METHOD DETAILS**

## Taxonomic sampling and sequencing

We isolated genomic DNA from museum specimens, sampling 58 individual bats, putatively representing 58 individuals and 42 putative Old World Myotis species for the ingroup, selected to index the major clades and nodes in the tree. 12 An unknown New World species, Myotis spp. (see below), and Submyotodon latirostris were also sequenced as outgroup species (Table S1). BLASTn<sup>107</sup> was used to try determine a more specific nomenclature for Myotis spp. using the mitogenome and Cyt-b sequences derived from this sample. The top hit for the mitogenome was Myotis martiniquensis with 94.6% pairwise identity and 91.81% query coverage. The top hit for the Cyt-b sequence was the GenBank: JN020572, a Myotis sp. individual from Suriname which grouped in a clade including Myotis nyctor and Myotis nesopolus in the following analysis. <sup>108</sup> Pairwise identity was 94.7% and query coverage was 99.91%.

DNA was extracted using the Qiagen Puregene kit following the manufacturer's protocol, except 10μL of linear acrylamide was added to the lysis buffer to aid the recovery of DNA from small starting amounts of tissue. Standard Illumina fragment libraries (300-bp) were prepared for each DNA sample using the NEBNext Ultra II DNA Library Prep Kit for Illumina. Samples were sequenced to ~20× genome-wide depth of coverage with 2 × 150-bp reads on the Illumina HISeq X platform. We combined these data with published genomes from two other bat species: Myotis brandtii<sup>109</sup> and Myotis myotis.

## Sequence QC and read mapping

Raw reads were trimmed using Trim Galore (http://www.bioinformatics.babraham.ac.uk/projects/trim\_galore). All but one species of Myotis possess a karyotype of 2n = 44, the exception being Myotis annectans (2n = 46). G-banding similarities indicate a high degree of collinearity across the genus. 48 All trimmed and filtered reads, using default trimgalore parameters, 90 were mapped to the Myotis myotis reference genome, HLmyoMyo6. Sequence alignment was carried out using bwa-mem<sup>91,92</sup> using the -M and -R parameters. Mapping results were evaluated and summarized using the Qualimap function bamqc. 93 Samtools 4 was used to remove duplicate reads. Local realignment and variant calling was performed using GATK. 95-98 Consensus pseudohaploid variants were called for each species using ANGSD<sup>56</sup> where the minimum required mapping quality was 30 and minimum base quality of 20. Short-read IIlumina assemblies struggle to accurately resolve highly repetitive and duplicated regions within a genome. This can introduce erroneous variant calls which can confound branch length inference in downstream phylogenomic analyses. To control for this, we also used ANGSD to identify and exclude all alignment regions with read depth greater than 200% of the average coverage per genome and positions with a minimum depth of 5 reads. Average mapped read depth and the proportion of the reference covered by mapped reads was calculated to assess the degree of missing data per species across the genome (Table S2)

## **Article**



## Mitogenome assembly and analysis

To avoid incorporating nuclear copies of mitochondrial genes in our mitogenome assemblies, we aimed to sample small subsets of reads to capture only reads belonging to the mitochondria, as nuclear copies of mitochondrial genes would be rare in such small subsets. Random subsets (2, 4, and 8 million) of raw reads were extracted for each sample and trimmed as described above. SOAPdenovo2<sup>110</sup> was used to assemble the mitogenome by evaluating a range of k-mer sizes (71, 81, and 91), with a read length of 150bp and 300bp as the insert size; all other parameters were set to default. Typically, the longest scaffold represented the majority of the assembled mitogenome. The assembly of the highly repetitive D Loop region was incomplete and highly variable across species and so was excluded from further analysis. To orient, annotate and align scaffolds for downstream analyses, each scaffold was mapped to the *M. myotis* mitogenome, <sup>111</sup> with a final alignment of ~17kb. Coding regions were extracted from the mitogenome using Geneious Prime 2019.04. Coding regions were RY coded using RYcode (https://github.com/ebraun68/RYcode)<sup>99</sup> to reduce the potential impact of compositional bias. <sup>112</sup> Alignments were then concatenated to form a single matrix. We prepared a second mitochondrial dataset which included the whole mitogenome, D Loop excluded. For both datasets, a Maximum likelihood (ML) tree was generated using IQtree2<sup>40</sup> under a GTR+I+G4 model of sequence evolution. The tree was evaluated with 1000 bootstrap replicates using the ultrafast bootstrap approximation (Figure S4). <sup>113,114</sup>

## Improving the Myotis myotis reference genome

The reference genomes for bats described in<sup>33</sup> reported 96–99% of data assembled to 'chromosome-level scaffolds'. However, upon examination of the assembly for key regions like the pseudoautosomal region and the major histocompatibility complex (MHC) locus, we determined that many genic regions were not assembled on chromosomes and would limit downstream inferences. We used two complementary approaches to further improve the completeness of the chromosome-level scaffolds. First, we used the gene annotation and a previously published ZOO-FISH map of human autosomal probes painted on Myotis myotis autosomes<sup>34</sup> to predict, based on gene content, which scaffolds in the HLmyoMyo6 assembly corresponded to chromosome paints in the ZOO-FISH map. We re-mapped the original Myotis myotis Hi-C (SRR11732436) data to test these predictions and re-scaffolded HLmyoMyo6 (GCA\_014108235.1).33 The Hi-C contact map was visualized using Juicebox.36 We searched for contacts uniting scaffold associations identified using the gene annotations and ZOO-FISH maps and adjusted the assembly accordingly. This way, more data were assigned to chromosomes, most often to chromosome ends (Figure S2). Differences between the original assembly and the improved version are summarized via Quast 100 in Table S3. Assembling regions with high rates of recombination regions is key to polarizing and accurately interpreting the distribution of phylogenomic signal. Further, we noted that many previously unassigned scaffolds contained immune genes, including the Major Histocompatibility Complex (MHC) and immunoglobin clusters (Figure S1), genomic regions of particular interest given bats' exceptional immunity<sup>115</sup> and role in emerging pathogens. 116 Placing these important genomic regions in the correct genomic context furthers our understanding of the evolution of immune genes in bats.

## A recombination map for Myotis myotis

We generated a recombination map for *M. myotis* to help interpret the chromosomal distribution of phylogenomic signal in our data. Studies have shown that hybridization occurs between the closely related sister species *M. myotis* and *Myotis blythii*, where they occur in sympatry throughout much of South and Central Europe. 117,118 To avoid bias due to potential genomic introgression between the species, we selected bats from Brittany, north-west France, which occur outside the range of *M. blythii*. Procedures were approved and permits were issued by 'Arrete' by the Prefet du Morbihan. *M. myotis* were sampled in western France, 2013 from three roost locations Ferel (FER), La Roche Bernard (LRB) and Beganne (BEG). Bats were captured in custom harp traps as they left the roost and were placed in individual cloth bags until sampled. A 3mm biopsy punch was obtained from the wing membrane. Samples were flash frozen in liquid nitrogen. Before release all bats were offered food and water. As the recombination land-scape of genomes has been shown to vary between different sexes of the same species, 119 we selected samples from two males MMY78 (LRB) and MMY243 (BEG) and one female (MMY4) to construct a sex averaged map.

DNA was extracted, and Illumina libraries were prepared and sequenced as described above. Raw reads were filtered, and mapped to the M. myotis reference genome, and the resulting bam files were processed with GATK as before. Variants were called, and all samples were jointly genotyped, then filtered to remove variants in repeatmasked regions using GATK. Variants were further filtered, removing variants within 5bps of an indel and those which did not meet the following quality criteria -e'%QUAL<30 | INFO/DP < 16 | INFO/DP > 62 | QD < 2 | FS > 60 | SOR>10 | ReadPosRankSum <-8 | MQRankSum <-12.5 | MQ < 40' in bcftools (https://github.com/samtools/bcftools). VCFtools (https://vcftools.github.io/man\_latest.html) was used to remove indels, leaving 12,273,471 biallelic SNPs (referred to as Mm recombination dataset) for further analysis. ReLERNN, a deep learning approach that uses recurrent neural networks, was used to model the genome-wide recombination rate for M. myotis. The average mammalian mutation rate 2.2  $\times$  10–9120 was used for this analysis. ReLERNN was run using the simulate, train, predict and bscorrect modules with default settings. Finally, inferred recombination rates were averaged in 2Mb blocks in 50kb sliding windows for use in downstream analyses.

To determine if the *Myotis myotis* recombination landscape was representative of the overall *Myotis* recombination landscape we used the same methods to generate a recombination map for the New World species, *Myotis brandtii*. <sup>89</sup> Publicly available SRA data were mapped to the *Myotis brandtii* pseudohaploid assembly generated as part of this study.





## **Phylogenomics**

## Coalescent and concatenation whole-genome phylogenies

SVDquartets<sup>39</sup> was chosen as the coalescence-based species tree approach to satisfy the assumption of no intra-locus recombination under the Multispecies Coalescent model (MSC). Variant sites were used to impute quartet trees which were then aggregated to infer a species tree. Many published coalescence-based analyses have included or been comprised only of protein coding regions or ultraconserved elements, thereby violating the MSC's neutrality assumption. Variable sites were extracted from the per chromosome fasta alignments in vcf format using SNP-sites v2.5.1. 103 Variable sites were thinned to a minimum of 1000bp between sites using VCFtools v1.16<sup>102</sup> and were converted to nexus format using the script vcf2phylip.py (https://github.com/edgardomortiz/ vcf2phylip). By using widely spaced variable sites, we aimed to account for the requirement of free recombination between loci, an assumption of coalescent methods like SVDquartets. The dataset was further filtered on missingness, stripping columns from the alignment with more than 10% missingness. While data were initially thinned to a minimum distance of 1000bp apart to avoid intralocus recombination, the final distance between most variable sites is likely much larger. Data from all chromosomes were concatenated using Geneious Prime 2019.04 for a final dataset comprising 1,463,340 SNVs (referred to as the genome-wide SNV dataset). A 50% majority rule tree was estimated using SVDquartets implemented in PAUP\* v4a168<sup>104</sup> using the QFM quartet amalgamation technique, sampling 99% of quartets.

The same genome-wide SNV dataset was concatenated into a supermatrix and analyzed with IQtree2 to generate a corresponding ML tree. Given that our nuclear datasets contained variable sites only, we used the +ASC option to correct for ascertainment bias due to the exclusion of constant sites in our analyses. Trees were analyzed as a single partition using a GTR + I + G model of sequence evolution to facilitate the use of the +ASC option. The +ASC option is currently incompatible with more complex mixture models. Trees were evaluated with 1000 bootstrap replicates using the ultrafast bootstrap approximation. 113,1114

## **Sliding windows**

Many phylogenomic studies show that phylogenetic signal is non-randomly distributed throughout the genome. To sample phylogenomic signal, we performed a sliding window-based analysis signal across the 60-species whole genome alignment (referred to as the whole-genome dataset). Alignments were divided into 50-kb nonoverlapping, contiguous windows across the entire genome. Alignment columns were filtered in trimAl<sup>105</sup> using the setting -qt 0.50 to allow only 50% missing data per alignment column and an alignment filtering script (https://github.com/VCMason/Foley2021) with the following parameters: cutoff = 0.02, Z score = 0.5, windsize = 50, step = 1, Myotis myotis as the reference and PCcutoff = 50, where an alignment is excluded if one species is missing more than 50% than the total alignment length. After filtering, only windows retaining 10kb or more data were retained for further analysis. Maximum Likelihood locus trees were generated using IQtree2 under a GTR model of sequence evolution and the GHOST model of rate variation (GTR\*H4). Trees were evaluated with 1000 bootstrap replicates using the ultrafast bootstrap approximation. 113,114 Relationships for the difficult-to-resolve relationships were summarized in subtrees extracted from the whole-genome dataset (Figure 1; Table S4). Data were visualized using Tree House Explorer (THEx). 61

## **Detection of gene flow**

Introgression causes phylogenomic distortion. 121,122 In a recent review, Hibbins and Hahn stated, "asymmetry in [the frequencies of] discordant tree topologies is one of the clearest signals of introgression".55 Following this and other studies, 41,43 we estimated the relative frequency of each topology for the autosomes and X separately. We also investigated the relationship between recombination rate and phylogenomic signal by calculating the frequencies of topologies in high and low recombining regions, defined as 500 windows at either end of the distribution of recombination rates.

Models of rate heterogeneity are fitted to data as part of some tree-building procedures to account for rate variation among sites. Simulations and empirical studies have shown that genome regions subject to introgression have elevated rate heterogeneity due to the accumulation of discordant donor sites in these regions. Potential introgression regions can be identified via the alpha (or shape) parameter, which is estimated as part of fitting gamma models of rate heterogeneity. Alpha parameters are lower in regions of potential introgression as multiple changes are required to account for apparent homoplasy. 122,123 To estimate the alpha parameter per locus, alignments used to generate the 60 species locus trees pruned to represent subtrees (Table S4). As the GHOST model of rate heterogeneity was used to estimate locus trees, we re-ran IQtree2 for alignment loci to fit a gamma model of rate heterogeneity to the data. The alpha parameter, number of parsimony informative sites, and the sum of internal and external branch lengths per locus were extracted from the output using the bash command grep.

The combined effect of introgression and recombination can create local patchworks of phylogenomic histories, resulting in discordant tree topologies and branch lengths. The degree of discordance varies depending on the timing and direction of introgression and the relationship between hybridizing taxa. 55 Introgression between sister taxa will not produce discordant topologies but will result in compressed branch lengths or divergence times. Introgression among non-sister species will create discordant topologies with much older or much younger branch lengths or divergence times depending on the direction and age of the introgression events. 55 As such, we examined the genome-wide distribution of internal branch lengths extracted from quartet topologies using the ape package 124 in R. 106 Given the extreme morphological conservation observed among Myotis bats 12 we did not formally estimate divergence times given the difficulty in accurately assigning fossils branches in this phylogeny, particularly in the presence of

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gene flow. In lieu of this, we examined the distribution of internal branch lengths to determine the distorting effect of introgression. In a divergence time analysis, the rate of substitution per site is simply scaled to time, thereby reflecting similar information.

Rather than relying solely upon genome-wide tests of introgression, <sup>125</sup> which do not identify specific regions of admixture, we used f (*D*), <sup>87</sup> which provides a point estimate of the admixture proportion at a given locus. More specifically, given a tree (((p1, p2), p3), O), f (*D*) describes the quantification of excess shared variation between p2 and p3 that is also not shared with p1. This approach also allows for the possibility of bidirectional introgression on a site-by-site basis. To test for evidence of admixture, we calculated f(*D*) for trios of species and an outgroup corresponding to the two most frequent introgressed topologies from each clade (Table S5). The whole-genome dataset chromosome alignments were pruned to the trio, and the outgroup and biallelic SNPs were called using SNPsites. <sup>103</sup> f(*D*) was calculated using scripts from (https://github.com/simonhmartin/genomics\_general) following (https://github.com/simonhmartin/tutorials/blob/master/ABBA\_BABA\_windows/README.md). Introgression was estimated in 10kb windows requiring a minimum of 250 SNPs per window as a cut-off for analysis.

Given that f(D) can underestimate introgression if all three species in the ingroup taxa hybridize, we also used QuIBL<sup>41</sup> to determine to what extent discordance could be explained by ILS and introgression. This approach uses the distribution of internal branch lengths to distinguish between these two processes. Under ILS, internal branch lengths are expected to be exponentially distributed, while internal branches derived from introgressed regions will have a distribution with a non-zero mode.<sup>41</sup> The fit of models where all discordance is due to ILS or a mixture of ILS and introgression are compared via likelihood, and the proportion of trees that best fit either model are returned. Trees were pruned to a trio of focal species and an outgroup and run through QuIBL using default parameters.

To determine if there was a difference in the probability of the admixture between related species based on mating behavior, we estimated *D* the whole genome statistic used to detect introgression among species, in the form (((H1, H2), H3), O), where O is the outgroup *S. latirostris* in all cases. Within the Ethiopian Clade, *M. bocagei* uses a harem-based mating strategy where one male and several females gather in a banana leaf for mating. Conversely, *M. emarginatus* is frequently reported at swarming sites in Europe. Using subsets of the whole-genome dataset, a suite of whole genome *D* statistics<sup>125</sup> were calculated in Dfoil<sup>88</sup> using mode = dstat and all other default parameters for all trees shown in Figure 5C which are shown in the format (outgroup, (P1,(P2, P3))). We varied the individual used to represent *M. bocagei*, the position of species in the tree and varied the third species selected in addition to *M. bocagei* and *M. emarginatus* to represent a variety of species which have been found to co-occur with *M. emarginatus* at swarming sites or are within the Ethiopian clade, but the status of their mating behavior is unknown.

To investigate if our results were impacted by reference bias, we repeated the analysis described above, but instead mapped the data to both the *Myotis myotis* reference genome and the *Myotis yumanensis* reference genome (mMyoYum1.0.hap1). <sup>86</sup> As a member of the New World clade, *M. yumanensis* is roughly equally distant from all Old World species and so could be used to detect any possible bias from using a nested ingroup species as the reference. ABBA-BABA or whole genome *D*-statistics were estimated using ANGSD<sup>56</sup> for combinations of BAM files resulting in highly similar estimates of *D* and Z-scores, indicating the negligible impact of reference bias. As such, the remainder of introgression tests were conducted with data mapped to the *M. myotis* reference genome. To investigate if sampling a different individual from the same species changed our introgression results, we mapped an additional *M. myotis* individual (NCBI: SRS3678524)<sup>126</sup> and used this individual in our re-analysis. These results also showed negligible differences to our original results (Figures 5C and S25; Table S5).

## **Hybridization at swarming sites?**

Typically swarming is thought to be limited to temperate bat species. However, data on the mating behavior of bats are limited and not widely known in families with large numbers of species. To better investigate this, we performed a literature search and aggregated data from multiple studies reporting species compositions at swarming sites across Europe (Table S7). As before, subtrees corresponding to the species which swarm within and between clades (Figure 5B) were pruned from the 60 species sliding windowed phylogenies derived from analyses of subsets of the whole-genome dataset. The number and frequency of genome-wide topologies were estimated using THEx. Given the observed asymmetry in topology frequencies, introgression was determined to be the most likely source of phylogenomic discordance. To test this, we calculated whole genome *D*-statistics (Figure 5B) and windowed f(*D*) statistics (Figure S17) for trios of species and an outgroup corresponding to the two most frequent introgressed topologies using data derived from the whole-genome dataset.

## **Anti-viral immunity and introgression**

Given that immune loci are frequently introgressed among mammals, <sup>66</sup> we sought to determine if genes associated with anti-viral immunity were introgressed in our dataset. Although the interferon pathway is a conserved anti-viral innate immune pathway, most species possess a repertoire of genes adapted for anti-viral defense. Given that standard gene enrichment analyses rely on data and function information for the best-studied model organisms, a traditional enrichment analysis would likely omit *Myotis*-specific components or features of anti-viral immunity. Therefore we used a custom database of differentially expressed genes from a study that treated *Myotis daubentonii* cell lines with a modified Rift Valley Fever virus or supplemented with IFN to look at differential gene expression in response to a viral challenge. <sup>70</sup> For our database, we selected all differentially expressed genes with an adjusted p value of 0.005 or less from both treatments after the 6h time point to capture an immediate acute response to viral infection. Of these genes, 731/984 were present in the windows represented by our 50kb sliding windows dataset. The 253 missing genes were likely





removed during the filtering of regions of poor alignment quality or were missing from the gene annotation. To determine if this gene set showed significant overlap with introgressed regions of the genome, defined as windows with a non-species tree topology, we performed a chi-squared test with a Yates correction in R using the native stats package.

## Large introgressed haplotype blocks

A large introgressed haplotype block was identified through visual examination of genome-wide phylogenomic signal in THEx for the bechsteinii clade on chromosome V15. This corresponded to the topology where M. myotis and M. daubentonii were sister taxa relative to M. bechsteinii (red topology Figure 3C). Genes corresponding to this region were extracted as described previously, and an STRING enrichment analysis 127 was performed choosing human for the database, where FDR > 0.05 was considered significant, and all other parameters as default to determine if any function or pathway was enriched in this region.

Cell Genomics, Volume 4

## **Supplemental information**

Karyotypic stasis and swarming

influenced the evolution of viral tolerance

in a species-rich bat radiation

Nicole M. Foley, Andrew J. Harris, Kevin R. Bredemeyer, Manuel Ruedi, Sebastien J. Puechmaille, Emma C. Teeling, Michael F. Criscitiello, and William J. Murphy

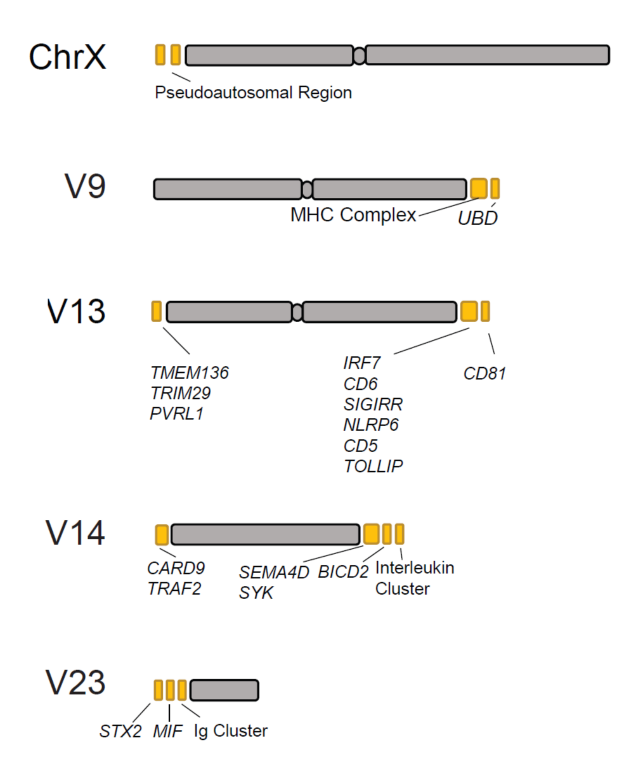


Figure S1: Previously unplaced assembly scaffolds contained many immune-related genes, related to the improved *Myotis myotis* genome assembly analyzed in Figure 2. Gray blocks depict the original chromosome-scale scaffolds from the original assembly (GCA\_014108235.1). Yellow blocks represent unplaced scaffolds currently assigned to chromosomes. The yellow blocks are not drawn to scale.

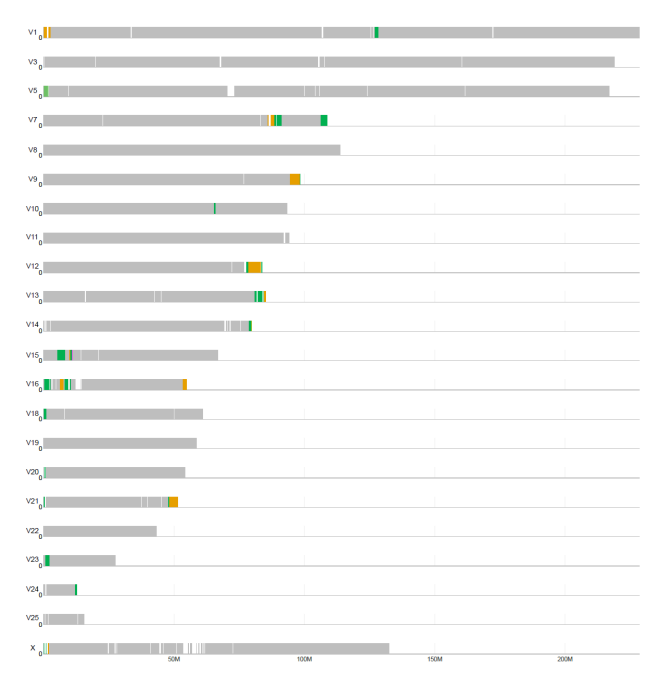


Figure S2: Most unplaced scaffolds were added to the terminal ends of chromosomes, related to the improved *Myotis myotis* genome analyzed in Figure 2. Portions of chromosomes colored gray represents the chromosome level scaffolds from the original assembly [S1]. Green and orange blocks, alternated to highlight where multiple scaffolds were added to the same region, illustrate where previously unplaced scaffolds were positioned in the new assembly based on Hi-C analysis and visualization in JBAT. All blocks and chromosomes are shown to scale. White spaces in blocks correspond to regions that were unassembled or removed due to missing data or poor alignment quality in the sliding window analysis.

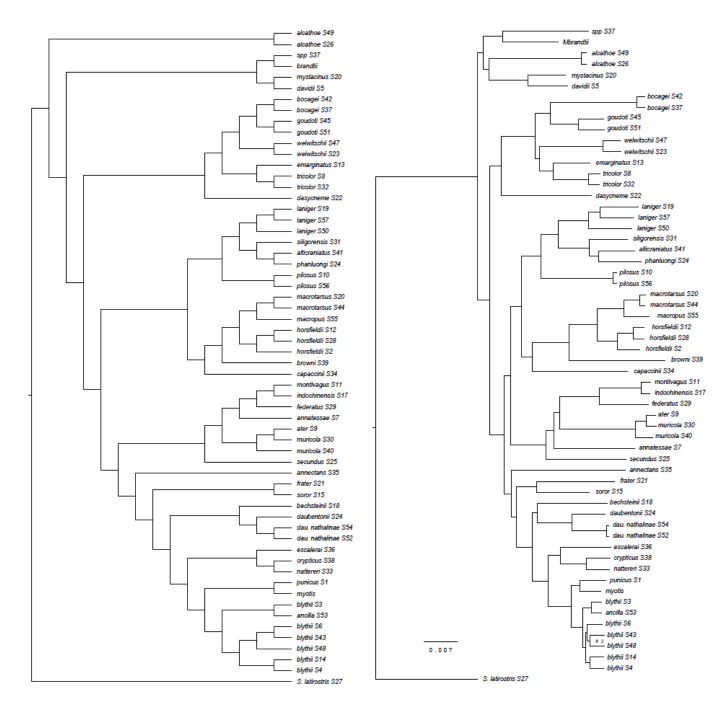
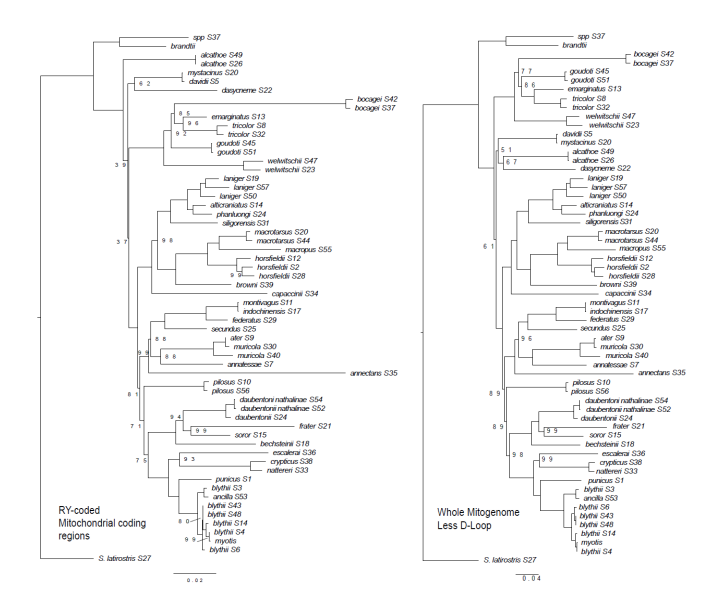
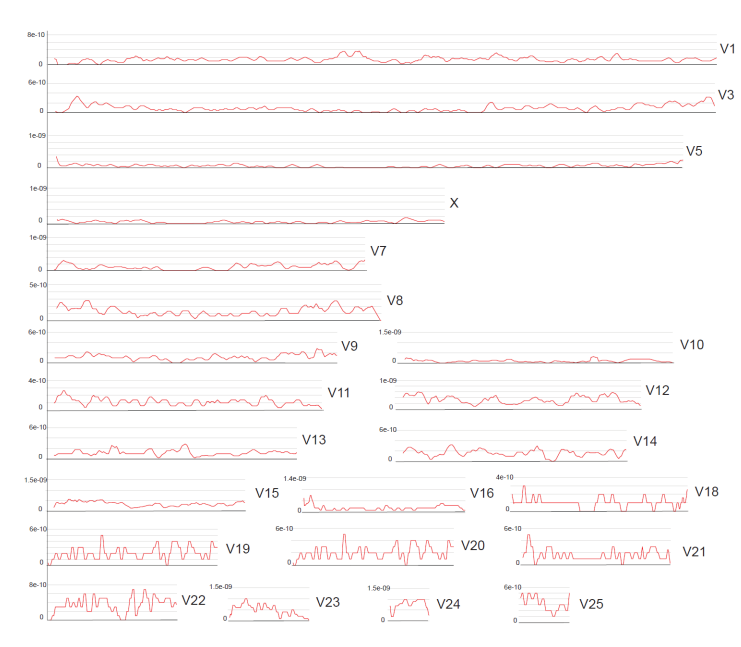


Figure S3: Whole genome nuclear topologies with all 60 individuals, related to Figure 1. The tree on the left is the 50% Majority Rule tree derived from the SVDquartets analysis. The rightmost tree is the IQtree2 Maximum Likelihood with branch lengths. All bootstrap supports are 100 unless otherwise indicated.



**Figure S4: Mitogenome topologies with all 60 individuals, related to Figure 1.** The tree on the left is RY-coded dataset. The tree on the right is the whole mitogenome with the D-Loop removed. All bootstrap supports are 100 unless otherwise indicated.



**Figure S5. Myotis myotis recombination map, related to Figure 2.** Genome-wide distribution of recombination rates inferred using ReLERNN. Rates are averaged every 2Mb in 50kb sliding windows. For illustrative purposes V1-V15 and chrX are displayed using a sliding average with a step of 50. Chromosomes V16-V25 are displayed using a sliding average with a step of 10.

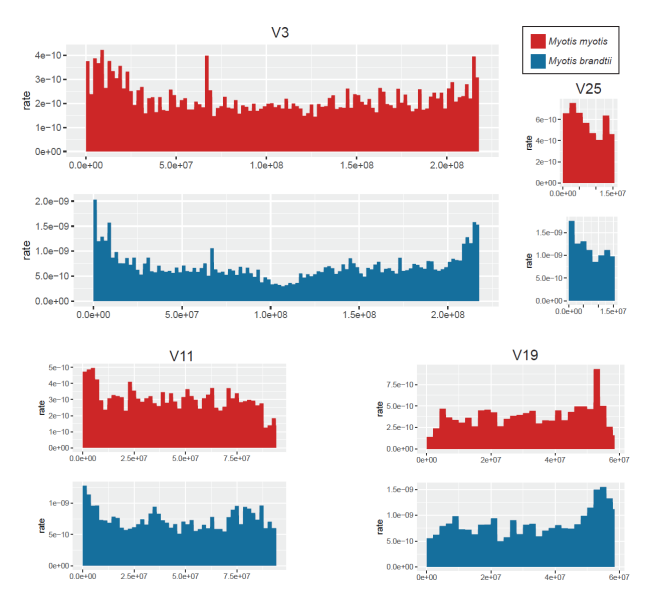


Figure S6: Comparison of select chromosomes from *M. myotis* and *M. brandtii* recombination maps, related to Figure 2.

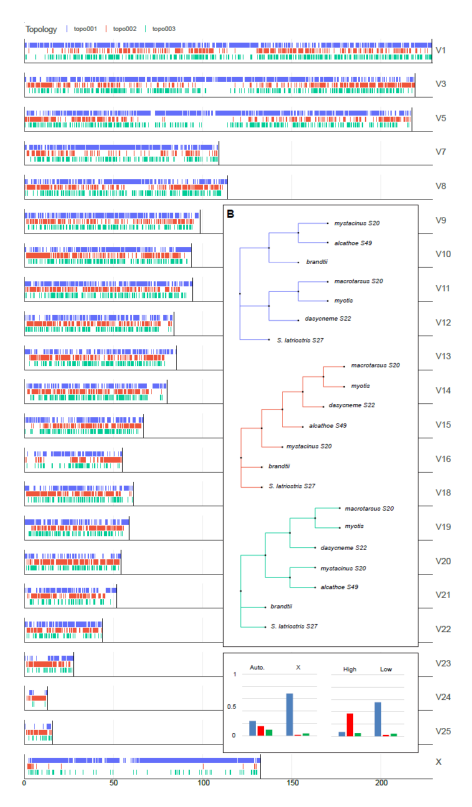


Figure S7: Genome-wide phylogenomic signal to determine the position of the whiskered bat clade, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.

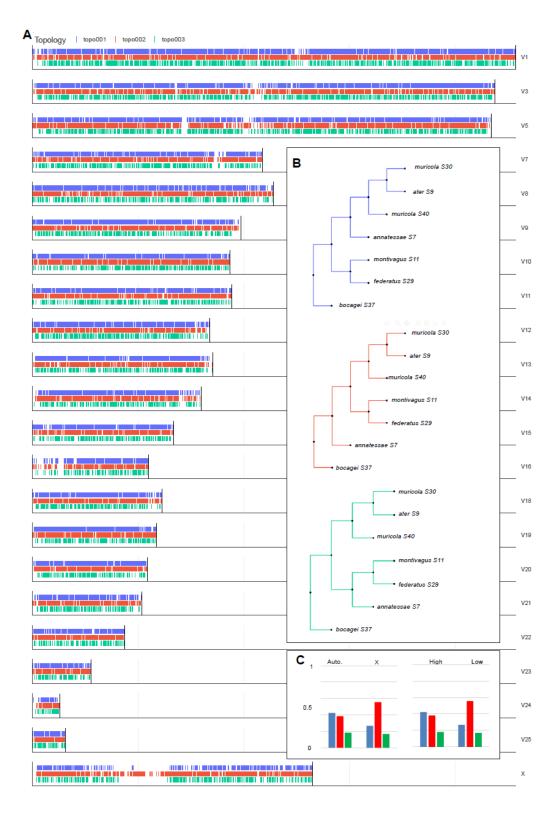


Figure S8: Genome-wide phylogenomic signal to determine the position of *M. annatessae*, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.

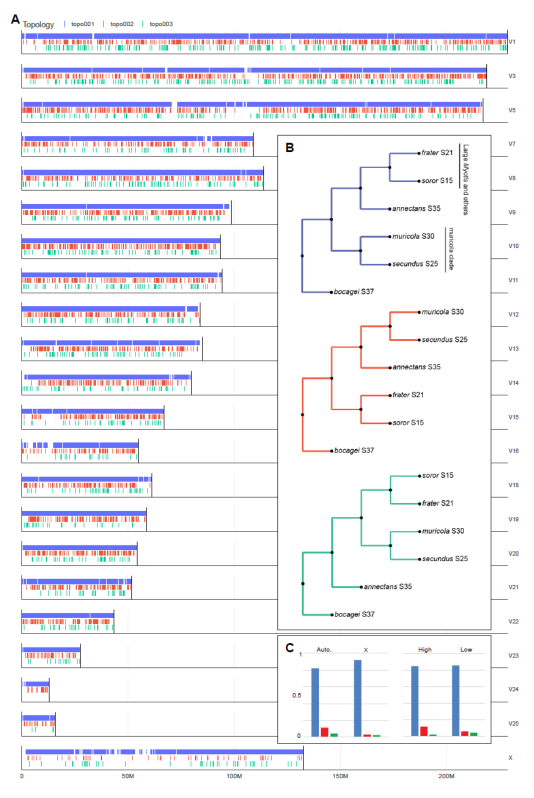


Figure S9: Genome-wide phylogenomic signal to determine the position of *M. annectans*, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.

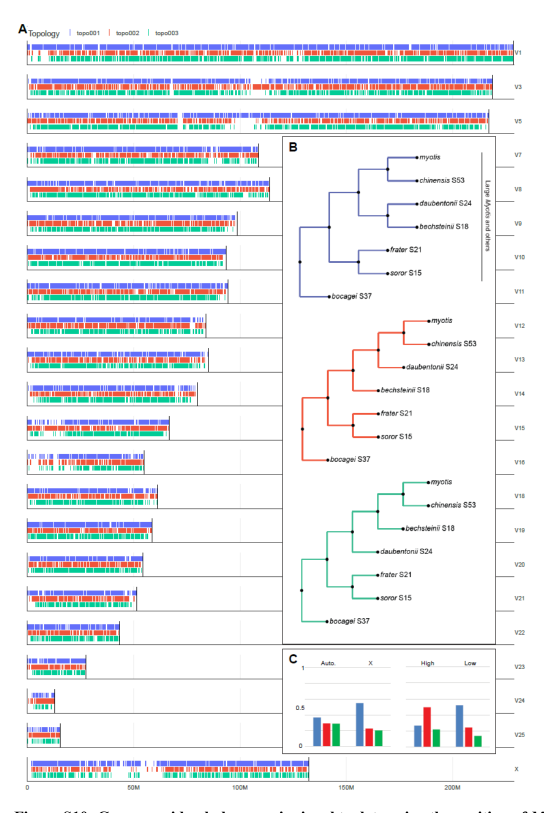


Figure S10: Genome-wide phylogenomic signal to determine the position of *M. bechsteinii*, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.

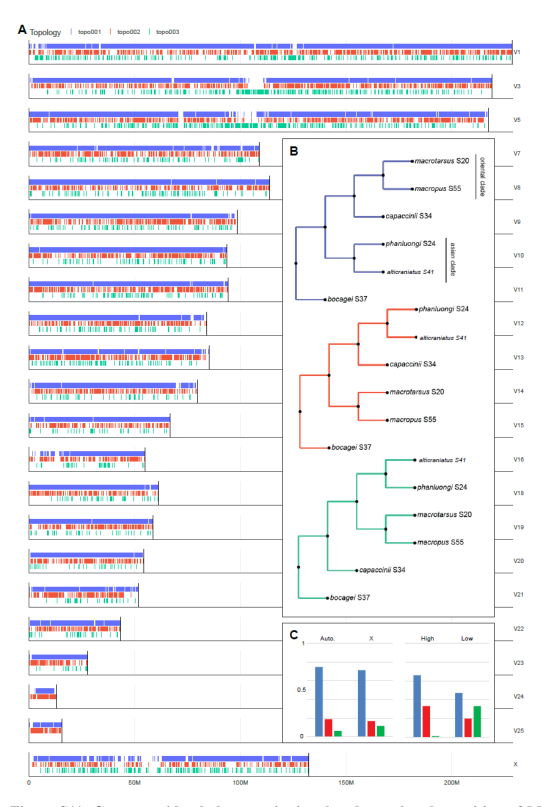


Figure S11: Genome-wide phylogenomic signal to determine the position of *M. capaccinii*, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.

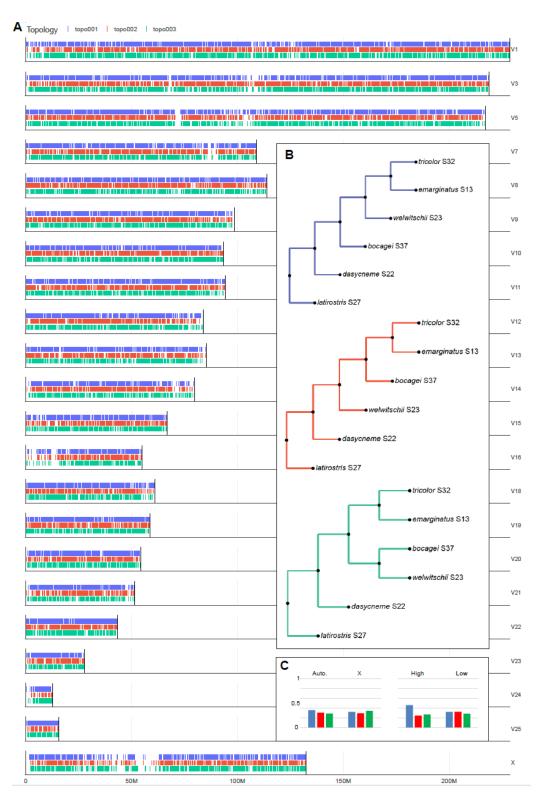


Figure S12: Genome-wide phylogenomic signal to determine the relationships among species in the Ethiopian clade, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.

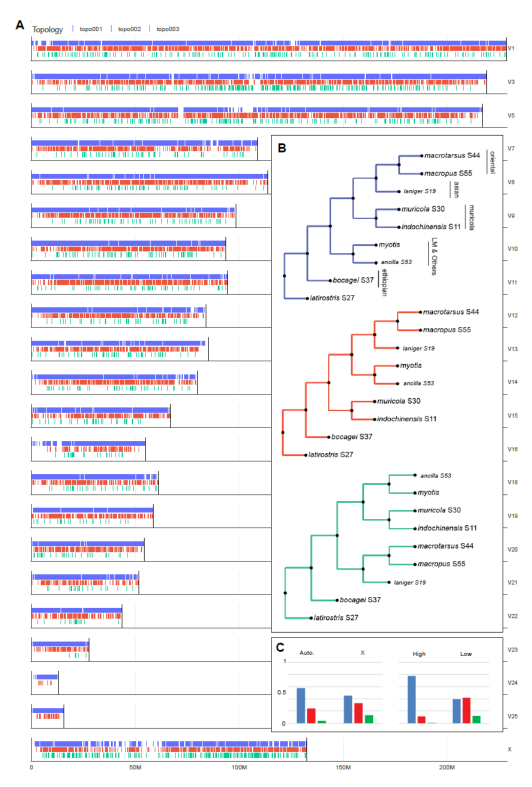


Figure S13: Genome-wide phylogenomic signal to determine the relationships among the major Old World Clades, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.

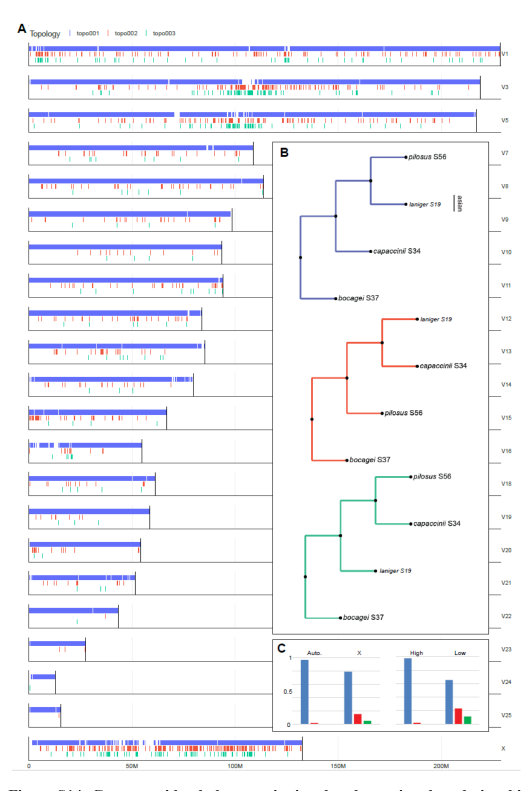


Figure S14: Genome-wide phylogenomic signal to determine the relationship between *M. pilosus*, *M. capaccinii* and *M. laniger*, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.

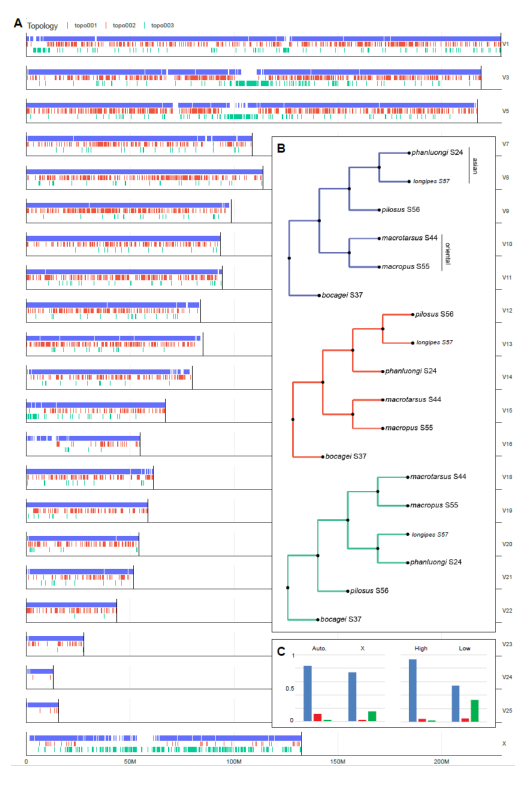


Figure S15: Genome-wide phylogenomic signal to determine the position of *M. pilosus*, related to Figure 3. A) Genome-wide distribution of topologies shown in B). C) shows the frequency of each topology on the autosomes and X and in high and low recombining regions.



Figure S16: Phylogenomic discordance, related to Figure 3. Topological frequency is calculated comparing the Autosomes and X and comparing low and high recombining regions. Average values of f(D), alpha and ibl per clade.

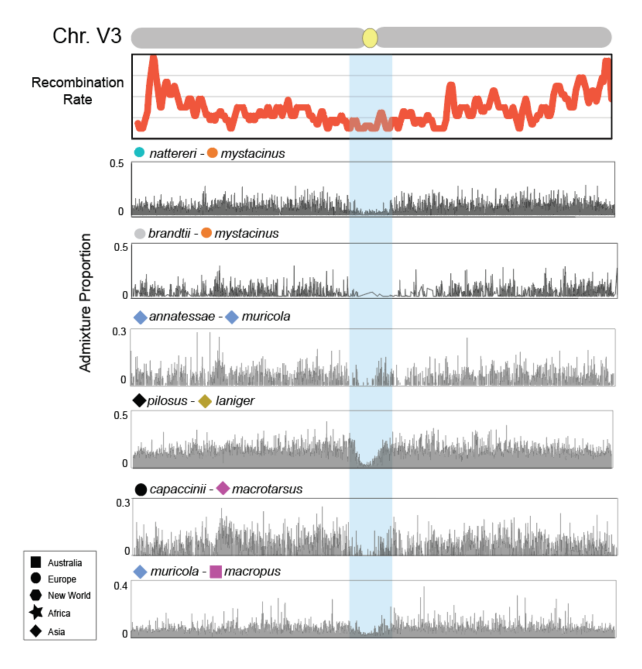


Figure S17: The f(D) statistic calculated across chromosome V3 for various species pairs, related to Figure 4. Introgression is evident along chromosome arms while low recombining pericentromeric regions are relatively devoid of signal consistent with introgression.

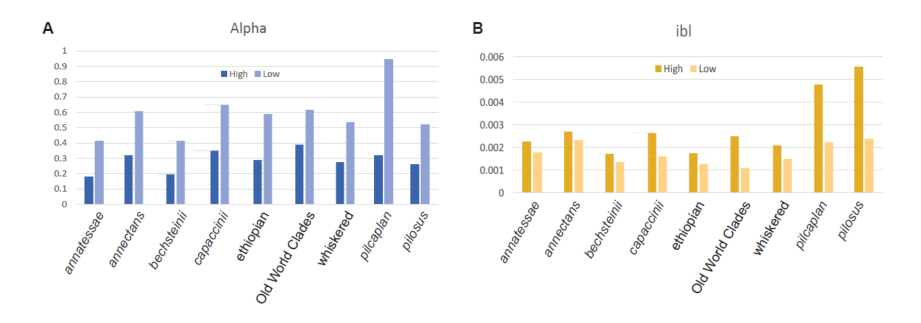


Figure S18: Distribution of alpha and internal branch length (ibl) in regions of high and low recombination, related to Figure 4.

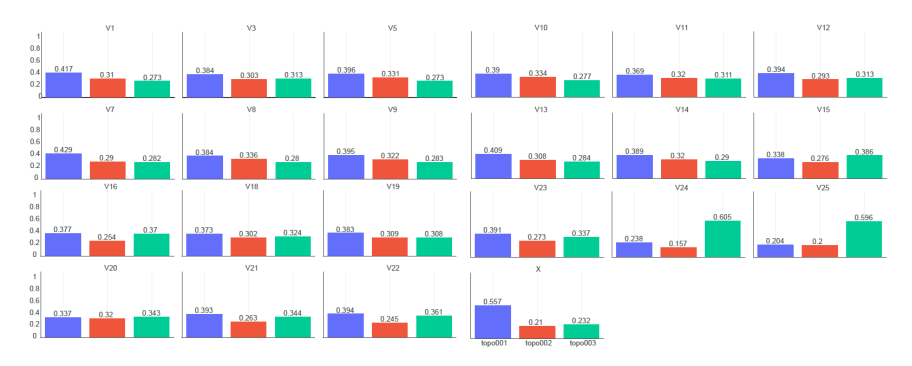


Figure S19: Topology frequency on each chromosome representing within clade relationships among species present at swarming sites, related to Figure 5.

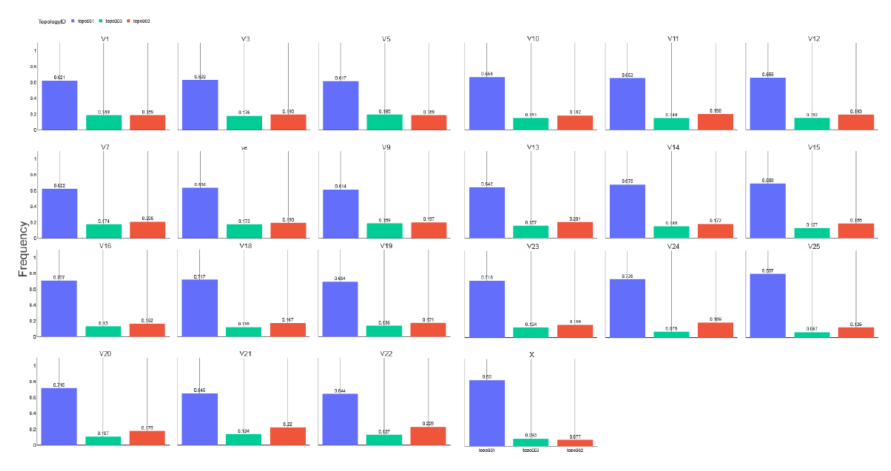


Figure S20: Topology frequency on each chromosome representing between clade relationships for *M. brandtii*, *M. mystacinus* and *M. alcathoe* at swarming sites, related to Figure 5.

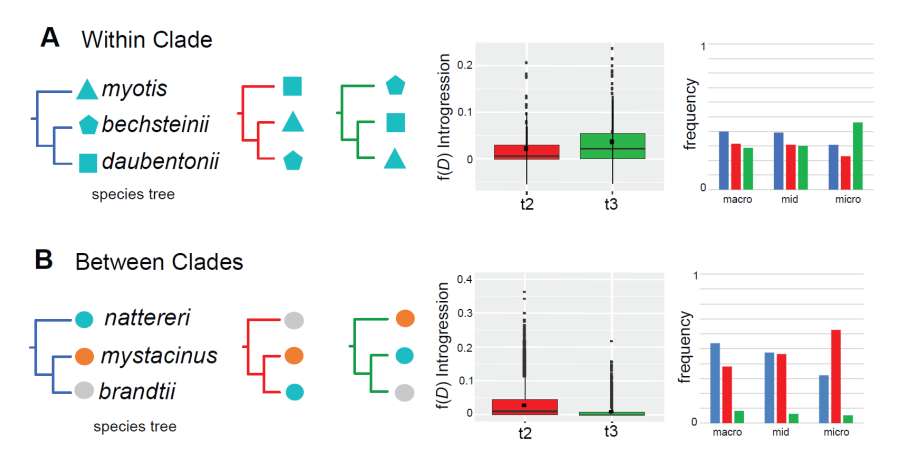


Figure S21: Introgressed topologies with the highest f(D) values are the most enriched on microchromosomes, related to Figure 6.

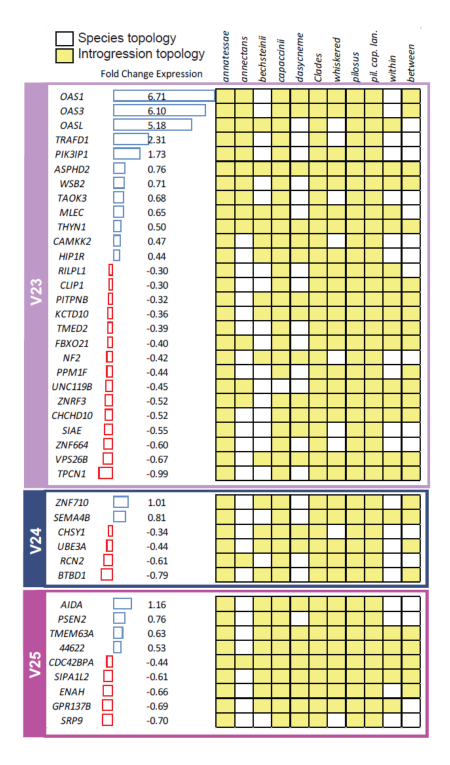


Figure S22: Genes that are differentially expressed in response to viral challenges in *M. daubentonii* were among the most frequently introgressed across microchromosomes, related to Figure 6.

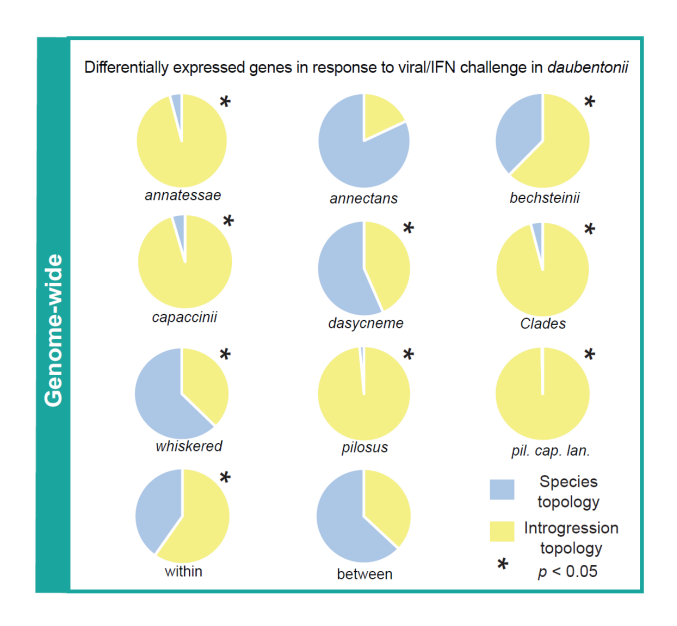


Figure S23: Significant overlap between introgressed genomic windows and genes that are differentially expressed in response to viral challenges in *M. daubentonii*, related to Figure 6.

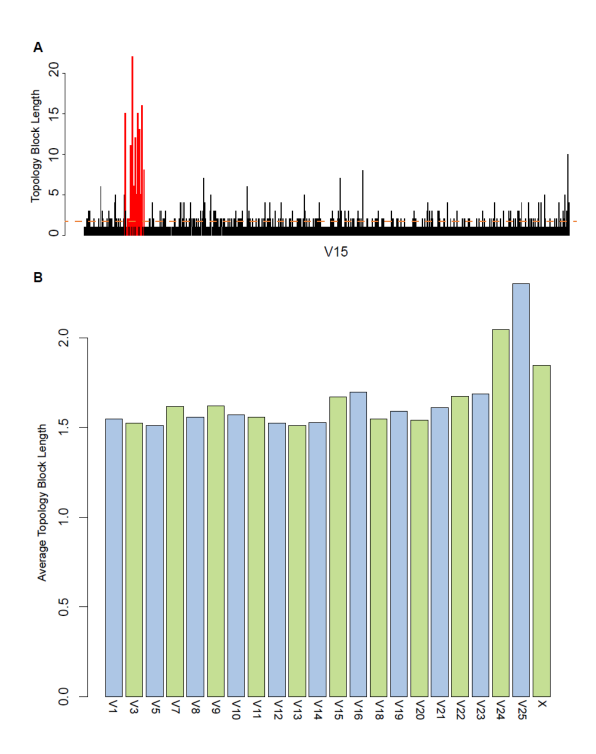


Figure S24: Plots of topology block length (=number of consecutive identical topologies) along chromosomes for the red topology uniting *M. myotis* and *M. daubentonii*, related to Figure 6. A) shows topology block length along chromosome V15. B) Microchromosomes V24 and V25 also exhibit elevated topology block lengths.

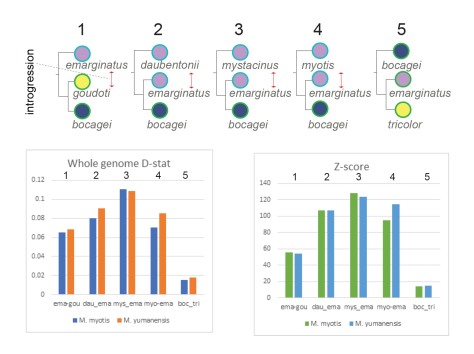


Figure S25: No impact of reference bias on whole genome tests of introgression, related to Figures 4, 5 and 6. *D*-statistics and *Z*-scores derived from data mapped to the *M. myotis* reference genome or the *M. yumanensis* genome are highly similar indicating the absence of reference bias

**Table S1: Sample Information, related to Figure 1.** Voucher or sample name, source and sampling locality for each sample used in this study. Categorization of broad scale species distributions as either tropical or temperate or spanning both. Abbreviations in sample source correspond to MHNG – Natural History Museum of Geneva, FMNH – Field Museum of Natural History Chicago, ZMMU – Zoological Museum Moscow Lomonosov State University, ROM – Royal Ontario Museum, EBD –Estacion Biologica de Doñana, AGS – Antonio Guillen-Servant, CMF – Charles M. Francis, SJP – Sebastien J. Puechmaille, MR – Manuel Ruedi

Voucher or Sample Name	Species	Sequencing Sample Number	Source	Locality	Clade
M2148	latirostris	S27	MHNG 1998.057	Taïwan.	
unknown	spp.	S37	MR		New World
M941	alcathoe	S26	MHNG 1828.073	Switzerland.	whiskered
M1575	alcathoe	S49	MHNG 1970.099	France.	whiskered
M803	davidii	S5	MHNG 1807.035	Greece	whiskered
M1993	mystacinus	S20	MHNG 1991.045	Switzerland.	whiskered
FMNH151200	bocagei	S42	FMNH	Tanzania	ethiopian
MZ004	bocagei	S37	MHNG 1971.027	Mozambique.	ethiopian
FMNH172095	goudoti	S45	MR/ FMNH	Madagascar.	ethiopian
FMNH173041	goudoti	S51	MR/FMNH	Madagascar.	ethiopian
TM39421	welwitschii	S23	MR/ Transvaal Museum	South Africa.	ethiopian
FMNH144313	welwitschii	S47	MR/ FMNH	Uganda.	ethiopian
M733	emarginatus	S13	MHNG 1807.040	Greece.	ethiopian
MZ425	tricolor	S8	MHNG 1971.030	Mozambique.	ethiopian
TM40300	tricolor	S32	MR/ Transvaal Museum	South Africa.	ethiopian
Mdas1	dasycneme	S22	MHNG 1805.052	The Netherlands.	ethiopian
M1606	laniger	S57	MHNG 1981.074	India.	asian
AG 950116.6	cf laniger	S19	AG/ SMF 86179	Laos.	asian
M629	laniger	S50	MR	Taiwan.	asian
M1637	siligorensis	S31	MHNG 1981.073	India.	asian
N194	alticraniatus	S41	MHNG 1956.089	Laos.	asian
S175154	phanluongi	S24	MR/ ZMMU		asian
CMF960418.17	pilosus	S10	CMF	Laos.	asian
M2065	pilosus	S56	MR	India.	asian
CMF960522.46	macrotarsus	S20	CMF	Borneo.	oriental
EAR122	macrotarsus	S44	MR	Philippines.	oriental
M	macropus	S55	MR	Australia.	oriental
CMF960523.40	horsfieldii	S12	CMF	Borneo.	oriental

Voucher or Sample Name	Species	Sequencing Sample Number	Source	Locality	Clade
M1551	horsfieldii	S28	MHNG 1970.049	Malaysia.	oriental
M1184	horsfieldii	S2	MHNG 1926.038	Laos.	oriental
LRH3168	browni	S39	MR	Philippines.	oriental
Mca1	capaccinii	S34	MHNG 1805.050	Italy.	oriental
AGS980326.44	montivagus	S11	AGS	Laos.	muricola
M1208	indochinensis	S17	MHNG 1926.043	Laos.	muricola
CMF960408.3	federatus	S29	CMF/ROM 106389	Laos.	muricola
M1200	annatessae	S7	MHNG 1926.044	Laos.	muricola
M1165	ater	S9	MHNG 1926.036	Laos.	muricola
M1569	muricola	S30	MHNG 1970.064	Malaysia.	muricola
M1600	muricola	S40	MHNG 1972.084	Indonesia.	muricola
M2152	secundus	S25	MHNG 1998.058	Taïwan.	muricola
CMF960406.5	annectans	S35	CMF/ ROM 106376	Laos	muricola
M2139	frater	S21	MHNG 1998.053	Taïwan.	Large Myotis & others
M2163	soror	S15	MHNG 1998.05	Taïwan.	Large Myotis & others
M2009	bechsteinii	S18	MHNG 1991.067	Switzerland.	Large Myotis & others
M2187	daubentonii	S24	MHNG 1999.098	Switzerland.	Large Myotis & others
M301	daubentonii nathalinae	S54	MR	Spain	Large Myotis & others
M303	daubentonii nathalinae	S52	MR	Spain.	Large Myotis & others
EBD25765	escalerai	S36	EBD	Spain	Large Myotis & others
M2096	crypticus	S38	MHNG 1996.094	Switzerland.	Large Myotis & others
M764	nattereri	S33	MHNG 1807.049	Greece.	Large Myotis & others
MM4SA	punicus	S1	MHNG 1805.092	Italy.	Large Myotis & others
Mb203	blythii	S3	MHNG 1805.026	Kyrgyzstan.	Large Myotis & others
631D	blythii	S14	MHNG 1805.037	Switzerland.	Large Myotis & others
M867	blythii	S6	MHNG 1807.099	Greece.	Large Myotis & others
Mb3	blythii	S4	MHNG 1805.041	Spain.	Large Myotis & others
SP.C.47	blythii	S43	SJP	Iran	Large Myotis & others
SP.C.48	blythii	S48	SJP	Iran	Large Myotis & others
Mchin1	ancilla	S53	MR	China	Large Myotis & others

**Table S2: Mapping Statistics, related to Figure 1.** 

Voucher or Sample Name	Species	Sequencing Sample Number	% Reference Covered	reads mapped	mean coverage
M2148	latirostris	S27	78.6	94.56%	25
unknown	spp.	S37	79.6	96.33%	12
M941	alcathoe	S26	82.1	97.68%	16
M1575	alcathoe	S49	84.4	97.44%	26
M803	davidii	S5	85	98.51%	23
M1993	mystacinus	S20	84.8	96.86%	22
FMNH144313	welwitschii	S47	83.1	95.94%	18
MZ004	bocagei	S37	84.8	97.17%	29
FMNH172095	goudoti	S45	81.5	95.22%	12
FMNH173041	goudoti	S51	84.8	96.80%	19
TM39421	welwitschii	S23	83.3	97.77%	17
FMNH151200	bocagei	S42	84	95.01%	19
M733	emarginatus	S13	86.1	98.32%	25
MZ425	tricolor	S8	85.4	97.52%	21
TM40300	tricolor	S32	86.2	98.11%	27
Mdas1	dasycneme	S22	79.5	98.70%	20
M1606	laniger	S57	85	98.78%	18
AG950116.6	laniger	S19	85.6	97.27%	23
M629	laniger	S50	85.8	98.33%	26
M1637	siligorensis	S31	84.8	97.39%	18
N194	alticraniatus	S41	86.6	98.29%	29
S175154	phanluongi	S24	81.3	97.45%	16
CMF960418.17	pilosus	S10	85.9	98.56%	24
M2065	pilosus	S56	86.1	98.68%	27
CMF960522.46	macrotarsus	S20	83	97.69%	14
EAR122	macrotarsus	S44	83.8	96.5%	16
M	macropus	S55	84.5	98.84%	20
CMF960523.40	horsfieldii	S12	86.1	97.91%	25

Voucher or Sample Name	Species	Sequencing Sample Number	% Reference Covered	reads mapped	mean coverage
M1551	horsfieldii	S28	83.5	97.01%	15
M1184	horsfieldii	S2	86.6	98.16%	28
LRH3168	browni	S39	83.9	96.62%	19
Mca1	capaccinii	S34	82.2	97.98%	13
AGS980326.44	montivagus	S11	85.2	97.63%	24
M1208	indochinensis	S17	85.4	97.06%	25
CMF960408.3	federatus	S29	85.7	97.62%	27
M1200	annatessae	S7	85.1	98.17%	20
M1165	ater	S9	85.9	98.04%	29
M1569	muricola	S30	84.9	96.57%	23
M1600	muricola	S40	84.6	94.99%	20
M2152	secundus	S25	86.7	96.75%	27
CMF960406.5	annectans	S35	82.4	97.80%	12
M2139	frater	S21	86.5	96.04%	20
M2163	soror	S15	86.4	97.90%	17
M2009	bechsteinii	S18	89.5	98.97%	27
M2187	daubentonii	S24	88.6	98.31%	20
M301	daubentonii nathalinae	S54	88.9	99.01%	23
M303	daubentonii nathalinae	S52	89	98.96%	26
EBD25765	escalerai	S36	90.5	96.94%	23
M2096	crypticus	S38	91.4	98.30%	28
M764	nattereri	S33	90.9	98.23%	24
MM4SA	punicus	S1	94.1	99.18%	20
Mb203	blythii	S3	94.1	98.86%	23
631D	blythii	S14	94.2	98.71%	24
M867	blythii	S6	93.5	99.35%	18
Mb3	blythii	S4	94.2	99.11%	22
SP.C.47	blythii	S43	93.3	98.64%	17
SP.C.48	blythii	S48	89.2	99.34%	17
Mchin1	ancilla	S53	93.1	98.44%	25

Table S3: Quast summary of difference between the original *Myotis myotis* assembly (HLmyoMyo6) and the improved assembly (Myotismyotis\_assembly\_V2), related to the improved *Myotis myotis* assembly described in the main text.

	HLmyoMyo	Myotismyotis_assembly_V2
# contigs	92	55
Largest contig	223369599	228434895
Total Length	2003238046	2003245946
GC%	43.11	43.11
N50	94448911	108770286
N75	74216526	79764552
L50	7	6
L75	12	12
# N's per 100kbp	1468.17	1468.56

Table S4: Species composition for each subclade examined in the analysis of genome-wide phylogenomic signal, related to Figure 3.

annatessae	annectans	bechsteinii	capaccinii	ethiopian	Old World Clades	whiskered	pilcaplan	pilosus
bocagei S37	soror S15	bocagei S37	bocagei S37	latirostris S27	latirostris S27	latirostris S27	bocagei S37	bocagei S37
muricola S30	frater S21	bechsteinii S18	capaccinii S34	tricolor S32	laniger S19	mystacinus S20	capaccinii S34	macropus S55
ater S9	annectans S35	daubentonii S24	macropus S55	welwitschii S23	macropus S55	alcathoe S49	laniger S19	macrotarsus S44
muricola S40	secundus S25	ancilla S53	macrotarsus S20	bocagei S37	macrotarsus S44	brandtii	pilosus S56	pilosus S56
federatus S29	muricola S30	myotis	phanluongi S24	emarginatus S13	muricola S30	bocagei S37		phanluongi S24
montivagus S11	bocagei S37	soror S15	alticraniatus S41	dasycneme S22	montivagus S11	myotis		laniger S57
annatessae S7		frater S21			myotis			
					ancilla S53			
					bocagei S37			

**Table S5: Detailed breakdown of results for genome wide** *D***-statistic tests, related to Figure 4.** Trees were analyzed in the format (((H1, H2), H3,) Outgroup) where the outgroup was *S. latirostris* for all tests. A Z score above 3 is considered significant.

Reference genome	<u>н</u> 1	<u>H2</u>	<u>нз</u>	<u>nABBA</u>	<u>nBABA</u>	<u>Dstat</u>	<u>jackEst</u>	<u>SE</u>	<u>z</u>
Myotis myotis	MZ004_bocagei_S37	M733_emarginatus_S13	M2187_daubentonii_S24	1791427	1511330	0.084807	0.084807	0.00079	107.3416
Myotis yumanensis	MZ004_bocagei_S37	M733_emarginatus_S13	M2187_daubentonii_S24	1711426	1427716	0.090378	0.090378	0.000843	107.2569
Myotis myotis	MZ004_bocagei_S37	M733_emarginatus_S13	myo_BROAD	1800187	1563127	0.070484	0.070484	0.00074	95.21591
Myotis yumanensis	MZ004_bocagei_S37	M733_emarginatus_S13	myo_BROAD	1733777	1460703	0.085483	0.085483	0.000746	114.6314
Myotis myotis	MZ004_bocagei_S37	M733_emarginatus_S13	M1993_mystacinus_S20	1627722	1303891	0.110462	0.110462	0.000861	128.308
Myotis yumanensis	MZ004_bocagei_S37	M733_emarginatus_S13	M1993_mystacinus_S20	1548282	1244630	0.108722	0.108722	0.000881	123.4457
Myotis myotis	MZ004_bocagei_S37	fmnh172095_goudoti_S45	M733_emarginatus_S13	2226221	1954903	0.064891	0.064891	0.001166	55.66637
Myotis yumanensis	MZ004_bocagei_S37	fmnh172095_goudoti_S45	M733_emarginatus_S13	2135239	1861688	0.06844	0.06844	0.001266	54.08118
Myotis myotis	TM40300_tricolor_S32	M733_emarginatus_S13	MZ004_bocagei_S37	1981908	2042725	-0.01511	-0.01511	0.001075	-14.0531
Myotis yumanensis	TM40300_tricolor_S32	M733_emarginatus_S13	MZ004_bocagei_S37	1895093	1963687	-0.01778	-0.01778	0.001175	-15.128
Myotis myotis	myo_BROAD	M2187_daubentonii_S24	S175154_phanluongi_S24	2058223	1844105	0.054869	0.054869	0.001072	51.19208
Myotis myotis	M2009_bechsteinii_S18	M2187_daubentonii_S24	myo_BROAD	2580187	2536752	0.008488	0.008488	0.002327	3.648049
Myotis myotis	CMF960522_46_macrotarsus_S20	M2009_bechsteinii_S18	M1569_muricola_S30	2926327	3861014	-0.13771	-0.13771	0.001369	-100.559
Myotis myotis	M2009_bechsteinii_S18	M2096_nattereri_S38	Mbrandtii	1202345	1248347	-0.01877	-0.01877	0.000794	-23.6339
Myotis myotis	M2187_daubentonii_S24	M2096_nattereri_S38	M1993_mystacinus_S20	1394982	1545713	-0.05126	-0.05126	0.000799	-64.1663
Myotis myotis	M1993_mystacinus_S20	Mbrandtii	M2096_nattereri_S38	2184190	3099853	-0.17329	-0.17329	0.002389	-72.5429
Myotis myotis	M2139_frater_S21	M2009_bechsteinii_S18	CMF960406_5_annectans_S35	2760696	3263672	-0.08349	-0.08349	0.002849	-29.3023
Myotis myotis	M1993_mystacinus_S20	M941_alcathoe_S26	Mbrandtii	2430561	2617513	-0.03703	-0.03703	0.001214	-30.496

Defenence general	Ш	ш	112	A DD A	D A D A	Datat	ioalvEst	CE.	7
Reference genome	<u>H1</u>	<u>H2</u>	<u>H3</u>	<u>nABBA</u>	<u>nBABA</u>	<u>Dstat</u>	jackEst	<u>SE</u>	<u>z</u>
Myotis myotis	CMF960522_46_macrotarsus_S20	S175154_phanluongi_S24	Mca1_capaccinii_S34	3473250	4084785	-0.08091	-0.08091	0.00315	-25.6831
Myotis myotis	M941_alcathoe_S26	Mbrandtii	Mdas1_dasycneme_S22	2214338	3236368	-0.1875	-0.1875	0.00342	-54.82
Myotis myotis	M1993_mystacinus_S20	Mbrandtii	Mdas1_dasycneme_S22	2121574	3071673	-0.18295	-0.18295	0.002391	-76.5191
Myotis myotis	M733_emarginatus_S13	Mdas1_dasycneme_S22	M2009_bechsteinii_S18	2507335	2422351	0.017239	0.017239	0.001189	14.49878
Myotis myotis	M2187_daubentonii_S24	M2096_nattereri_S38	Mdas1_dasycneme_S22	1560422	1696782	-0.04186	-0.04186	0.000738	-56.7437
Myotis myotis	M733_emarginatus_S13	Mdas1_dasycneme_S22	M1993_mystacinus_S20	2131851	2029572	0.024578	0.024578	0.000942	26.09706
Myotis myotis	myo_BROAD	M733_emarginatus_S13	M1993_mystacinus_S20	2532201	2527665	0.000896	0.000896	0.001001	0.895467
Myotis myotis	M2009_bechsteinii_S18	M2187_daubentonii_S24	M2096_nattereri_S38	2621507	2579505	0.008076	0.008076	0.002248	3.592021
Myotis myotis	M629_laniger_S50	CMF960522_46_macrotarsus_S20	M2065_pilosus_S56	2361436	7045984	-0.49796	-0.49796	0.002534	-196.503
Myotis myotis	M1569_muricola_S30	M1208_indochinensis_S17	M1200_annatessae_S7	3455060	3730538	-0.03834	-0.03834	0.00132	-29.0523
Myotis myotis	M2187_daubentonii_S24	M2009_bechsteinii_S18	myo_BROAD	2536800	2581340	-0.0087	-0.0087	0.002334	-3.72829
Myotis myotis	M2139_frater_S21	M2187_daubentonii_S24	M2065_pilosus_S56	2650991	3062319	-0.07199	-0.07199	0.001695	-42.4712
Myotis myotis	M1569_muricola_S30	M1208_indochinensis_S17	CMF960406_5_annectans_S35	2093563	1918718	0.043577	0.043577	0.000709	61.43691
Myotis myotis	M2096_nattereri_S38	M2187_daubentonii_S24	M733_emarginatus_S13	1610340	1590837	0.006092	0.006092	0.000795	7.668013

**Table S6: Results of the QuIBL analysis, related to Figure 4.** The table shows the proportion of variation in discordant topologies explained by ILS alone or a combination of ILS and introgression. The difference in BIC estimates (delta) for each model indicates that the model with introgression and ILS (preferred at delta ≥-10) is the best fit for all relationships analyzed. Old World Clades refers to the relationship among the Muricola, Large Myotis, Asian, and Oriental clades shown in Figure 1. *pil.-cap.-lan.* refers to the relationship among *pilosus*, *capaccinii*, and *laniger*. Non-species trees are color-coded to correspond with data displayed in Figure 1.

Clade	non-species topology	tree	ILS	Introgression + ILS	delta
annatessae	topo002	(annatessae, (indochinensis, muricola))	0%	100%	-13383
	topo001	indochinensis, (muricola ,annatessae))	0%	100%	-17245
annectans	topo003	(annectans, (frater, muricola))	0%	100%	-995
	topo002	(frater, (annectans, muricola))	0%	100%	-3319
bechsteinii	topo002	(bechsteinii, (daubentonii, myotis))	0%	100%	-9038
	topo003	(daubentonii, (bechsteinii, myotis))	0%	100%	-9045
capaccinii	topo001	(phanluongi, (capaccinii, macrotarsus))	0%	100%	-30065
	topo002	(macrotarsus, (capaccinii, phanluongi))	0%	100%	-5825
dasycneme	topo004	(mystacinus, (alcathoe, (dasycneme, bocagei)))	0%	100%	-299
	topo002	(bocagei, (dasyceneme, (mystacinus, alcathoe)))	0%	100%	-44060
Clades	topo003	(muricola, (myotis, macrotarsus))	0%	100%	-9122
	topo001	(myotis, (macrotarsus, muricola))	0%	100%	-17445
whiskered	topo004	(alcathoe, (mystacinus, brandtii))	0%	100%	-4761

	topo006	(mystacinus, alcathoe, brandtii))	0%	100%	-4474
pil. cap. lan.	topo003	(laniger, (capaccinii,pilosus))	73%	27%	33
	topo001	capaccinii, ((pilosus, laniger))	0%	100%	-33039
pilosus	topo001	(macropus, (pilosus, laniger))	0%	100%	-32580
	topo005	(pilosus, (macropus, laniger))	28%	72%	-174

**Table S7: Studies reporting species presence at swarming sites across Europe [S2-16], related to Figure 5.** Although *M. myotis* and *M. blythii* are thought to use a lekking mating behavior

Paper	Schunger et al (2004)	Dundarova (2018)	Rivers et al (2006)	Parsons et al (2003)	Piksa et al (2011)	Suba et al (2008)	Schaik et al (2015)	Bogdanowicz et al 2012	Suba et al (2011)	Glover and Altringham (2008)	Schmidbauer and Denzinger (2019)	Pocora et al (2012)	Dekeukeleire et al (2016)	Thomas and Davison (2020)	Ignaczak et al (2019)
Country	Bulgaria	Bulgaria	UK	UK	Poland	Latvia	Netherlands	Poland	Latvia	NK	Germany	Romania	Belgium and Netherlands	Wales	Poland
myotis	X	Х			X		X				Х	Х	X		Х
blythii	X	X													
bechsteinii	X	X		X	X		X					X	X	X	X
nattereri/crypticus	X	X	X	X	X	X				X	X	X	X	X	X
daubentonii	X	X	X	X	X	X	X		X	X		X			X
brandtii	X	X	X	X	X	X	X	X	X	X		X	X	X	X
alcathoe	X	X			X			X							
mystacinus		X	X	X	X	X	X	X		X		X	X	X	
emarginatus	X	X					X						X		
dasycneme					X	X	X		X				X		

Table S8: STRING enrichment analysis for an introgressed block on V15 which unites M. myotis and M. daubentonii, related to Figure 6. Enrichments are considered significant with an FDR value of  $\geq 0.05$ .

Category	#term ID	term description	observed gene count	background gene count	strength	false discovery rate
Molecular Function	GO:0005149	interleukin-1 receptor binding	7	17	2.04	8.6E-09
Molecular Function	GO:0070851	Growth factor receptor binding	8	138	1.19	1.2E-04
Molecular Function	GO:0005126	Cytokine receptor binding	8	264	0.9	9.0E-03
KEGG Pathway	hsa04060	Cytokine-cytokine receptor interaction	7	282	0.82	3.6E-02
Reactome Pathway	HSA-446652	Interleukin-1 family signaling	7	136	1.13	2.4E-03
Reactome Pathway	HSA- 9014826	Interleukin-36 pathway	3	6	2.12	4.6E-03

**Table S9: Immunogenetic diversification mechanisms across the Tree of Life, related to Figure 6.** Note that most of these mechanisms operate within the listed taxa. MHC allelic polymorphism has long been of interests to evolutionary biologists due to its function at the population level. Now adaptive immunogenetic introgression in *Myotis* bats appears to elevate such defense diversification strategies to the level of the multispecies community.

Taxa	Process	Genes Diversified	Evolutionary Facilitation	Mechanism
archaea/bacteria	adaptive anti-phage immunity	clustered regularly interspaced short palindromic repeats (CRISPR)	horizontal gene transfer	CRISPR/cas9 [S17]
arthropods	repertoire of pattern recognition receptors	down syndrome cell adhesion molecule (DSCAM)	exon family expansion	mutually exclusive alternative RNA exon splicing [S18]
jawless fishes	VLR repertoire diversification	variable lymphocyte receptors	copy-choice recombination	APOBEC family deaminases [S19]
jawed vertebrates	somatic cell gene rearrangement of lymphocyte antigen receptors	immunoglobulin, T cell receptor	horizontal transposon capture	recombination activating genes (RAG) [S20]
jawed vertebrates	antibody affinity maturation	immunoglobulin heavy and light chain variable genes	selection of controlled somatic hypermutation	activation induced cytidine deaminase (AID) [S21]
Bovidae	ultralong cattlebody knob diversification	third complementarity determining region (CDR3) of immunoglobulin heavy chain	bias for cysteine mutation, truncations in CDR3	activation induced cytidine deaminase (AID) [S22]
jawed vertebrates	allelic polymorphism	classical MHC antigen presentation	balancing selection, pathogen co-evolution	extreme allelic diversity in peptide binding - Population [S23]
Myotis bats	tunable immunogenetic viral tolerance	interferons, antigen processing, anti-viral signaling	swarming, interspecific hybridization	adaptive immunogenetic introgression – Genus Community

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