**The genomic landscape, causes, and consequences of extensive phylogenomic discordance in murine mice and rats**

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# Abstract

A species tree is a central concept in evolutionary biology whereby a single branching phylogeny reflects relationships among species. However, the phylogenies of different genomic regions often differ from the species tree. Although tree discordance is widespread in phylogenomic studies, we still lack a clear understanding of how variation in phylogenetic patterns is shaped by genome biology or the extent to which discordance may compromise comparative studies. We characterized patterns of phylogenomic discordance across the murine mice and rats – a large and ecologically diverse group that gave rise to the laboratory mouse and rat model systems. Combining recently published linked-read genome assemblies for seven murine species with other available rodent genomes, we first used ultra-conserved elements (UCEs) to infer a robust time-calibrated species tree. We then used whole genomes to examine finer-scale patterns of discordance across 12 million years of divergence. We found that proximate chromosomal regions tended to have more similar phylogenetic histories, but no clear relationship between local tree similarity and recombination rates in house mice. However, we did observe a correlation between recombination rates and average similarity to the species tree. We also detected a strong influence of linked selection whereby purifying selection at UCEs led to appreciably less discordance. Finally, we show that assuming a single species tree can result in high error rates when testing for positive selection under different models. Collectively, our results highlight the complex relationship between phylogenetic inference and genome biology and underscore how failure to account for this complexity can mislead comparative genomic studies.

# Introduction

Phylogenies are the unifying concept in understanding the evolution of species, traits, and genes. However, extensive high-throughput sequencing data has now revealed that evolutionary relationships between species may not be well represented by a single representative phylogeny (Edwards 2009; Hahn and Nakhleh 2016). While a dominant signal of bifurcating speciation usually exists (*i.e.*, a species tree), phylogenetic signal that may disagree with species relationships can arise from ancestral polymorphisms (incomplete lineage sorting; ILS), gene flow (introgression), and gene duplication and loss (Maddison 1997). The theoretical prediction of phylogenetic discordance has long been appreciated (Hudson 1983; Pamilo and Nei 1988; Maddison 1997; Rosenberg 2002), but empirical evidence now emphasizes just how extensive discordance can be among a set of species (Feng et al. 2022; Gable et al. 2022; Smith et al. 2023). For example, studies of birds (Jarvis et al. 2014), mammals (Ferreira et al. 2021; Lopes et al. 2021; Foley et al. 2024), plants (Pease et al. 2016), and insects (Sun et al. 2021; He et al. 2023) have found that with extensive taxon sampling and genomic data, highly supported species tree topologies are rarely or never recovered in the underlying locus trees. Whereas these examples highlight the prevalence of phylogenetic discordance across the tree of life, we still lack a clear understanding of how phylogenetic patterns are shaped by the details of genome biology or the extent to which discordance may compromise inferences from comparative studies that assume a singular species history.

In practice, failure to acknowledge and account for phylogenetic discordance could severely affect biological inference. Analyses of molecular evolution are usually performed on a gene-by-gene basis (Pond et al. 2005; Yang 2007; Hu et al. 2019; Kowalczyk et al. 2019), but it is still common practice to assume a single genome-wide species tree for each locus. For gene-based analyses, using the wrong tree may cause erroneous inferences of positive directional selection, convergent evolution, and genome-wide inferences of correlated rate variation (Mendes et al. 2016). Phylogenetic discordance can also affect how continuous traits are reconstructed across phylogenies, as the genes that underly these traits may not follow the species history (Avise and Robinson 2008; Hahn and Nakhleh 2016; Mendes et al. 2018; Hibbins et al. 2023). In these instances, phylogenetic discordance may need to be characterized and incorporated into the experimental and analytical design. Alternatively, if a researcher’s primary questions are focused on reconstructing the evolutionary history of speciation (*i.e.*, the species tree), then phylogenetic discordance may obscure the true signal of speciation (Fontaine et al. 2015; Foley et al. 2024). In this case, knowledge about patterns of discordance across genomes could inform decisions about locus selection, data filtering, and model parameters during species tree reconstruction.

Given these considerations, a better understanding of the genomic context of phylogenetic discordance is warranted. Although often conceptualized primarily as a stochastic consequence of population history (Maddison 1997), patterns of phylogenetic discordance are likely to be non-random and dependent on localized patterns of genetic drift, natural selection, recombination, and mutation. Discordance due to ILS ultimately depends on effective population sizes across the branches of the phylogeny (Pamilo and Nei 1988; Degnan and Rosenberg 2006) and, therefore, should covary with any process that influences local patterns of genetic diversity (*e.g*., linked negative or positive selection). Likewise, discordance due to introgression may be influenced by selection against incompatible alleles or positive selection for beneficial variants (Lewontin and Birch 1966; Jones et al. 2018). Selection, ILS, and introgression, are expected to leave different genomic signals that should allow us to test hypotheses about both the cause and the scale of phylogenetic discordance (Huson et al. 2005; Kulathinal et al. 2009; Green et al. 2010; Vanderpool et al. 2020). Yet the genomic context of phylogenetic discordance has remained elusive. For example, localized patterns of phylogenetic discordance should be influenced by patterns of recombination (Hudson and Kaplan 1988) and simulation studies confirm that the closer two regions are in the genome, the more history they share (Slatkin and Pollack 2006; McKenzie and Eaton 2020). However, empirical studies have been inconclusive regarding the relationship between discordance and recombination rates, ranging from no relationship in great apes (Hobolth et al. 2007), a weak positive correlation in house mice (White et al. 2009), a strong positive correlation broadly across primates (Rivas-Gonzalez et al. (2023), or increased discordance in regions of lower recombination (Scally et al. 2012; Pease and Hahn 2013). Thus, it remains unclear how phylogenetic discordance scales locally across the genome as a function of recombination and the strength of linked selection, pointing to the need for empirical studies in systems with sufficient egenomic resources to explore the causes of discordance.

To investigate the causes and consequences of phylogenetic discordance, we took advantage of genomic resources available for house mice (*Mus musculus*). This rodent species is one of the most important mammalian model systems for biological and biomedical research and is embedded within a massive radiation of murine rats and mice (Murinae). This ecologically diverse and species-rich group is comprised of over 600 species and makes up >10% of all mammalian species, and yet is only about 15 million years old, making this system an excellent choice for phylogenetic studies over both short and long timescales. Despite the power of evolution-guided functional and biomedical analysis (Christmas et al. 2023), relatively few murine genomes have been sequenced outside of *Mus* and *Rattus*.

We analyze recently sequenced genomes for seven murine species (*Mastomys natalensis*, *Hylomyscus alleni*, *Praomys delectorum*, *Rhabdomys dilectus*, *Grammoyms dolichurus*, *Otomoys typus*, and *Rhynchomys soricoides*) sampled from across this radiation (Kumon et al. 2021). We combine these new genomes with previously sequenced genomes and genomic resources from the *M. musculus* model system to study phylogenetic relationships within Murinae as well as the landscape of discordance along rodent chromosomes. The aims of our study are two-fold. We first inferred a species tree for these and other sequenced rodent genomes, focusing on signals derived from commonly used ultra-conserved elements (UCEs). We used these UCE data to infer a robust, time-calibrated phylogeny of sequenced murine rodents, providing a useful resource for future comparative studies within this important group. Using this species tree, we then used a subset of whole genomes to study how phylogenetic discordance is related to species-level inferences of relatedness, recombination rate, and patterns of molecular evolution. Using genetic maps and functional annotation from the powerful house mouse system, we test several hypotheses linking spatial patterns of discordance to genetic drift, natural selection, and recombination. Finally, we show how the use of a single species-tree impacts gene-level inferences from common molecular evolution tests for natural selection in these species. Collectively, our results advance understanding of how core features of genome biology influence underlying phylogenetic patterns, the extent to which established model system resources can be leveraged for broader phylogenetic studies, and the consequences of ignoring phylogenetic uncertainty.

# Results

## Estimation of a murine species tree

Using a concatenated dataset of 2,632 aligned ultra conserved elements (UCEs), we inferred a species tree of 18 murine rodent species (Fig. 1; Table S1) that recovered the same relationships as previous reconstructions of Murinae using a small number of loci (Lecompte et al. 2008; Steppan and Schenk 2017). The species tree inferred from a quartet-based summary of the gene tree topologies was identical to the concatenated tree (Fig. S1). While bootstrap and SH-aLRT values provided high support to our inferred species trees (Fig. 1), we found evidence for considerable discordance across individual UCE phylogenies. The five shortest branches in the concatenated tree had a site concordance factor (sCF) of less than 50%, suggesting that alternate resolutions of the quartet had equivocal support (Fig. S2). Gene concordance factors (gCF) for each branch in the species tree were on aggregate much higher, with all but four branches supported by almost every gene tree in the analysis and with the lowest values likely being driven by a several short internal branches (Fig. S2). This pattern was recapitulated using a quartet-based summary method (Figs. S1 and S3). At the two most discordant nodes (E and J in Fig. 1), the recovered topology was supported by approximately one third of all gene trees.

We estimated divergence times for the inferred concatenated phylogeny (Fig. 1; Table S2) using four fossil calibration points (Table S3). The murid and cricetid groups had an estimated divergence time of 22.62 Ma (node A in Fig. 1) followed by the Murinae and the Gerbillinae at 21.30 Ma (B), albeit with wide confidence intervals in both cases. With the ancestral Murinae node (C) fixed for calibration, Hydromyini arose at 12.12 Ma (D) and was followed by Otomyini and Arvicanthini at 11.67 Ma (E). The remaining Murine tribes evolved in rapid succession, with Apodemini diverging at 10.82 Ma (F) and Murini and Praomyini splitting at 10.08 Ma (H). The *Rattus* node, which was fossil calibrated, was recovered at the very youngest end of the calibration range. Although congruent with previous works (Lecompte et al. 2008; Steppan and Schenk 2017), this dated UCE phylogeny provides context on the evolutionary timescale upon which we next describe the genomic landscape of phylogenetic discordance across a collection of murine genomes.

## The landscape of phylogenetic discordance along murine genomes

We analyzed genome-wide phylogenetic histories of six recently sequenced murine rodent genomes and the *M. musculus* reference genome spanning approximately 12 million years of divergence (see Fig. 1). Using the *M. musculus* coordinate system, we partitioned and aligned 263,389 non-overlapping 10 kb windows from these seven species (Table S1). After filtering windows in repetitive regions or with low phylogenetic signal, we recovered 163,765 trees with an average of 616 informative sites per window (Fig. S4).

Phylogenetic discordance was pervasive within and between chromosomes. We inferred 597 of the 945 possible unique rooted topologies among six species (when specifying *R. soricoides* as the outgroup) across all chromosomes. The number of unique topologies per chromosome ranged from 75 to 218 (mean = 141). However, just four different topologies were ranked in the top three per chromosome. (Fig 2A; File S1) and only nine trees were present at a frequency above 1%. Among these, the top three topologies only differed in the ordering of the clade containing *Hylomyscus alleni, Mastomys natalensis,* and *Praomys delectorum* (HMP clade). This clade also showed the second lowest concordance in the species tree inferred from UCEs (Fig. 1, node J). These three topologies comprise between 13-15% of all recovered topologies (Fig. 2). Interestingly, the least common of these three trees (13.1%) matched the topology recovered via concatenation of all coding regions and the species tree recovered from UCEs (Fig. 1). That is, the robustly inferred species tree did not match the evolutionary relationships inferred for over 85% of the genome.

While visual inspection revealed no clear partitioning of topological structures along chromosomes (*e.g.*, Fig. 2C), we found that phylogenies were not randomly distributed across mouse chromosomes. Using the weighted Robinson-Foulds metric, we found that tree similarity between windows decayed logarithmically along chromosomes (Fig. 3A and B) and the distance at which tree similarity appeared random varied considerably among chromosomes ranging from 0.15 Megabases (Mb) on chromosome 17 to 141.29 Mb on the chromosome 2 (Fig. 3C, Fig. S5). While chromosomes 2, 7, 9, and 11 were autosomal outliers with distances between windows to random-like trees exceeding 25 Mb, the average distance among all other autosomes was only 2.1 Mb. The rates at which phylogenetic similarity decayed tended to be inversely proportional to the distance at which two randomly drawn phylogenies lost similarity (Fig. 3D).

Next, we performed a pairwise alignment of the reference mouse and rat genomes to assess how large structural variation, such as inversions and translocations, may influence our inferences of phylogenetic relatedness along the genome. These species span the deepest divergence of the sample for which we assessed genome-wide discordance, so the level of large structural variation present among them should give us an idea of the amount of ancestral variation in our sample. The mouse and rat genomes were mostly co-linear for large, aligned chunks, with large translocations and inversions on mouse chromosomes 5, 8, 10, 13, and 16 (Fig. S6). We also observe large-scale inversions on chromosome 16. We found that, while most chromosomes were co-linear between mouse and rat, the average size of the 307,275 contiguously aligned chunks averages under 10 kb, with the average distance between aligned segments being between 2,380 bp on the mouse genome and 4,927 bp on the rat chromosome (Fig. S7). This pattern presents two major implications for our analyses. First, we could not transpose the coordinate system from mouse to rat with enough resolution to use genetic maps from rat. Second, most other structural variation in our sample appears likely to be small insertions of transposable elements (e.g., SINEs ~150-500 bp, LINEs ~4-7kb; (Platt et al. 2018)) that should have a negligible effect on discordance analyses since our window size is much larger and we excluded windows that were made up of mostly repeats.

## Discordance with recombination rate and other genomic features

Using markers from genetic crosses within *M. musculus* (Shifman et al. 2006; Cox et al. 2009) we examined whether regions with high recombination also showed more phylogenetic discordance over short genetic distances when compared to regions with low recombination. Specifically, we calculated recombination rates within 5 Mb windows (Fig. S8) and then measured tree similarity between the first and last 10 kb window (R2 = 3.0e-9; *p* = 0.99; Fig. 4A) and the rate at which tree similarity changes between the first 10 kb window and every other 10 kb window (R2 = 0.003; *p* = 0.11; Fig. 4B). Surprisingly, we found no relationship between tree similarity and recombination rates measured at this scale. However, we did observe a slight positive correlation between recombination rate and dissimilarity to the species tree when averaging wRF over all 10 kb window trees within a 5 Mb recombination window (R2 = 0.05; *p* = 7.6e-8; Fig. 4C). We also examined regions of the genome centered on recombination hotspots identified in *M. musculus* (Smagulova et al. 2011) and found that these regions had significantly slower rates of decay in similarity over genomic distance compared to windows that were not centered on hotspots (*p* = 0.019; Fig. 5A), and that they were also significantly more phylogenetically similar over short distances (*p* = 0.015; Fig. 5B). Thus, when taken as a whole, we found that regions of higher recombination rates in house mice did not show more local phylogenetic discordance per se but did tend to show more discordance relative to the genome-wide species tree.

Evolutionary relationships around certain conserved genomic features may also be shaped by locally reduced effective population sizes due to a history of pervasive linked negative or positive selection. To test for this, we measured tree similarity in 10 kb windows around all annotated protein-coding genes, ultra-conserved elements (UCEs), and protein-coding genes identified as evolving rapidly (*i.e.*, significantly elevated *dN*/*dS*) due to positive directional selection and compared these patterns relative to chromosome-wide trends (i.e., windows without annotated features). In general, UCEs showed more local phylogenetic similarity among adjacent windows (i.e., less discordance) than regions surrounding recombination hotspots (*p* = 2.42e-12), coding genes (*p* = 4.65e-14), rapidly evolving coding genes (*p* = 1.56e-6), and windows that did not include any of these features (*p* = 5.02e-14; Fig. 5A). In contrast, protein-coding genes (including rapidly evolving genes) were indistinguishable from background rates of discordance observed in windows without annotated genomic features (Fig. 5A). Likewise, UCEs were also much more similar to the overall species tree when compared to any other feature (Fig. 5B). Unlike our test of local discordance, protein-coding genes also showed less species tree discordance than windows containing no features or recombination hotspots, but the effect was much less pronounced than observed at UCEs.

## Consequences of tree misspecification on analyses of molecular evolution

Next, we examined how phylogenetic discordance influenced inferences on the evolution of protein-coding sequences. Among a set of 22,261 *M. musculus* protein-coding transcripts, the average distance between the start and end of the coding sequence was 37.02 kb, or roughly 4 non-overlapping 10 kb windows. At this distance, tree similarity is predicted to diminish considerably (*e.g.*, by 0.10 wRF units), such that the phylogenetic history of individual genes may often contain some phylogenetic discordance (Mendes and Hahn 2016; Mendes et al. 2019). We also found that out of the 67,566 times the coding sequence in a gene overlapped with a 10 kb window, the inferred topology of the gene tree exactly matched the topology of the corresponding window tree only 11% of the time. Thus, the common practice of inferring gene trees on concatenated coding exons from a single transcript is still likely averaging over multiple possible albeit correlated histories.

Finally, we tested how tree misspecification might impact standard *dN*/*dS* based phylogenetic analyses for positive directional selection. Specifically, we used the still common practice of assuming a single species tree for all genes and compared that to using individually inferred gene trees in three common statistical tests for positive selection: PAML’s M1a vs. M2a test (Yang 2007), HyPhy’s BUSTED test (Murrell et al. 2015), and HyPhy’s aBSREL test (Smith et al. 2015a). We found evidence that tree misspecification likely induces both false positive (type I) and false negative (type II) errors. For example, many genes were inferred as having experienced positive directional selection when using a single species tree, but not when using local gene trees and vice versa (Fig. 6). Assuming the locally inferred gene tree is more accurate than the single tree inferred from concatenation of all gene sequences, this resulted in varying rates and types of error (Table 1). For BUSTED, we observe that 28% of genes inferred as having evolved under positive directional selection when using the gene tree were not inferred when using the concatenated species tree (likely false negatives). The opposite was true for M1a vs. M2a, where, among genes showing inconsistent evidence for positive selection across the two scenarios, 76% do so when using the concatenated species tree but not individual gene trees (likely false positives). In general, genes found to be evolving under positive selection using both tree types tended to be more concordant with the species tree than those that had evidence for positive selection either using only the concatenated tree or the gene tree (Fig. 6).

# Discussion

Phylogenies provide insight into the relationships of species and serve as a framework for asking questions about molecular and trait evolution. However, phylogenetic histories can vary extensively across regions of a genome, and evolutionary relationships between species may not often be well represented by a single representative species-level phylogeny. Here, we combine the resources of the house mouse (*Mus musculus*) with new and recently published (Kumon et al. 2021) genomes from seven species to understand the systematics of murine rodents and causes and consequences of phylogenetic discordance along murine genomes. These new analyses help to place this important model system in a stronger evolutionary context and begin to fill the gap in genome sampling of murine rodents which, despite their exceptional morphological and ecological diversity and species richness, have had relatively few whole genomes sequenced. They further provide us with the resources to study the landscape of phylogenetic discordance across the genome, understand how recombination and natural selection shape phylogenetic histories, and evaluate how assuming a single evolutionary history can compromise the study of molecular evolution in an important biomedical model system.

## Phylogenomic relationships of murine rodent lineages from conserved genomic regions

The extraordinary species richness of murine rodents complicates phylogenetic analyses because of the resources required to sample, sequence, and analyze such widely distributed taxa. Earlier work either attempted to resolve specific groups such as *Mus* (Lundrigan et al. 2002; Suzuki et al. 2004) and *Apodemus* (Serizawa et al. 2000; Liu et al. 2004), or to uncover broader relationships across the subfamily (Martin et al. 2000; Steppan et al. 2005) based on a few genetic markers. Evidence from these datasets suggested appreciable phylogenetic discordance across Murinae that lead to different genomic regions supporting incompatible phylogenetic reconstructions (Suzuki et al. 2004; Steppan et al. 2005; White et al. 2009). Lecompte et al. (2008) provided one of the earliest well-supported phylogenetic reconstructions from across Murinae and the tribal classifications they proposed remain generally supported. More recent work has increased the number of taxa sampled, both for analyses of Murinae specifically (Pagès et al. 2016) and for their placement within Muridae and Muroidea (Schenk et al. 2013; Steppan and Schenk 2017; Rowe et al. 2019), but the number of loci used for phylogenetic inference remained limited. Other recent studies have greatly expanded the number of loci used for phylogenetic inference, including the use of 1,245 exons (Roycroft et al. 2020) and 1,360 exons (Roycroft et al. 2021), but have focused mainly on the Hydromyini group.

Our inferred species tree based on 2,632 UCEs from 18 species across the radiation (Fig. 1) is consistent with previous studies (Lecompte et al. 2008; Steppan and Schenk 2017; Aghova et al. 2018)*.* Branch support was uniformly high, and gene trees unambiguously support the tribal classification of Lecompte et al. (2008). However, four shorter branches show more substantial gene tree discordance (Fig. 1, branches D, E, H, and J), with two recovered clades (E and J) being supported by less than half of all gene trees. We also estimated divergence times on our inferred species tree using four fossil calibration points (Table S3), recovering times that are roughly consistent with the relatively young estimates found by (Steppan and Schenk 2017) (see Supplement). This dated species tree provides the evolutionary timescale upon which to evaluate the genomic landscape of phylogenetic discordance across ~12 my of murine evolution.

## The genomic landscape of phylogenetic discordance

Limiting the number and nature of the loci used to resolve species relationships is often useful to get an initial picture of the history of speciation across many taxa. However, such targeted approaches may fail to capture the degree of discordance driven by incomplete lineage sorting and introgression (Alexander et al. 2017; Chan et al. 2020; Vanderpool et al. 2020; Alda et al. 2021). Our results highlight the limitations of a reduced marker-based approach and the general relationships between phylogenetic patterns and functional attributes of the genome in several interesting ways. Using the house mouse genome annotation, we found that the species tree inferred from only genes or UCEs did not match evolutionary relationships inferred for over 85% of the genome. Although similar frequencies were observed among these three most common trees (Fig. 2), the topology robustly inferred from genes or UCEs was not that common overall and only the third most frequent topology among 10 kb windows genome-wide. This result was driven mainly by discordance among three more closely related (Praomyini) species sampled for this study, which had nodes with low concordance in the UCE species tree (Fig 1, node J). In the window analysis, each alternate topology of this clade occurred at a frequency of ~14% while the rest of the topology remained consistent with the species tree (Fig. 2), indicating that the alternate topologies are caused by high levels of ILS at these nodes. Increased discordance at unresolved nodes is a common feature of phylogenomic datasets. These patterns illustrate how extensive discordance can arise even in a small sample of species and underscores that a single inferred species tree often may not capture the history of individual regions of the genome.

Given the fundamental role that recombination should play in shaping patterns of genetic variation within genomes, it is reasonable to assume that patterns of ILS should be broadly shaped by local recombination rate. We did not observe a clear relationship between local recombination rates in mice (*M. musculus*) and the degree of local phylogenetic discordance (i.e., phylogenetic similarity over 5 Mb intervals). However, we did find that regions of high recombination rate tended to me more discordant with the inferred species tree, in line with findings in mammals (Pease and Hahn 2013; Foley et al. 2023; Rivas-Gonzalez et al. 2023) and *Drosophila* (Pease and Hahn 2013). Recombination rates evolve fairly rapidly both within (Kong et al. 2002; Cox et al. 2009; Stapley et al. 2017) and between mammalian species (Jensen-Seaman et al. 2004; Ptak et al. 2005; Stevison et al. 2016; Stapley et al. 2017) due, in part, to the high turnover of hotspots due to the changing landscape of binding sites for PRDM9 (Baudat et al. 2010; Singhal et al. 2015). Similar to findings in great apes (Hobolth et al. 2007), our results suggest that high-resolution genetic maps from a single species provide some weak predictive value for understanding broader patterns of species tree discordance. However, these limited estimates may not be predictive of finer-scale patterns in a sample spanning over 12 million years of mammalian evolution (but see (Foley et al. 2023). Overall, the evolution of recombination landscapes across closely related species remains an important empirical question in evolutionary genetics (Dapper and Payseur 2017), especially as the generation of chromosome-scale genome assemblies continues to greatly outpace estimates of patterns of recombination within those genomes.

One source of evolution in the recombination map may be changes in synteny. Our reference-guided analyses assume collinearity between *Mus* and the other lineages we are comparing (*i.e.*, no karyotype variation), at least at the window-based scale we are comparing variation. Structural variation, including substantial variation in chromosome numbers, is likely to be widespread in rodents (Stanyon et al. 1999; Yalcin et al. 2011; Romanenko et al. 2012; Keane et al. 2014) and has the potential to skew our results when comparing tree similarity between regions of the genome using multiple species. Generating chromosome-scale assemblies for many non-*Mus* and *Rattus* species may prove limiting given that most tissue resources for this group are derived from natural history collections that often lack high molecular weight DNA. Nonetheless, whole genome alignments between mouse and rat indicate high degrees of chromosomal synteny and co-linearity (Fig S6), suggesting that many regions will be colinear in our sample.

Natural selection reduces the effective population size (*Ne*) of genomic regions through the processes of genetic hitchhiking of variation linked to the fixation of positively selected mutations (*i.e.*, selective sweeps; Smith and Haigh 1974; Kaplan et al. 1989) and the purging of deleterious mutations (*i.e.*, background selection; Charlesworth et al. 1993; Hudson and Kaplan 1995). Thus, variation in parameters dependent on *Ne* – such as standing levels of nucleotide variation and patterns of incomplete lineage sorting – should be reduced by linkage to functional elements subject to selection. Consistent with this, we observed the lowest rates of local discordance (Fig. 5A) and overall gene tree/species tree discordance (Fig. 5B) near UCEs when compared to all other genomic features we studied. These results suggest that a history of recurrent purifying selection on UCEs (Katzman et al. 2007) strongly reduces patterns of discordance through a persistent local reduction in *Ne*. In contrast, protein coding genes showed rates of local discordance that were similar to background levels, even when considering genes rapidly evolving due to positive directional selection (Fig. 5A). However, both classes of genes did show less species tree discordance than background consistent with previous results (Scally et al. 2012; Rivas-Gonzalez et al. 2023), but this effect was much weaker than as observed at UCEs (Fig. 5B). Collectively, these data suggest that the frequency and strength of selection plays an important role in structuring patterns of incomplete lineage sorting across the genome over deeper evolutionary timescales.

One practical consequence of this is that phylogenetic inferences based on UCE markers would seem less prone to discordance and may provide cleaner estimates of species tree history than randomly chosen or protein-coding regions. Indeed, windows centered on UCEs have a higher degree of similarity to the species tree than other genomic features (i.e., 17% concordance with the species tree, versus 13% genome-wide or 15% for protein-coding genes). However, it is worth noting that UCEs are also more likely to provide a potentially misleading underestimate of genome-wide levels of discordance. Given this relationship, species tree inferences based on UCEs should likely not, for example, be extended to related population genetic parameters of interest (*e.g.*, ancestral population sizes, estimates population genetic diversity), and could mislead the reconstruction of trait evolution across phylogenies (Avise and Robinson 2008; Hahn and Nakhleh 2016; Mendes et al. 2018; Hibbins et al. 2023). Finally, despite the relative ease of generating UCE data, such markers are likely not suitable for genetic inferences within populations given the pervasive effects of linked selection.

## Discordance and Molecular Evolution

We also found that the choice of tree topology drastically affects the results from various common tests for positive selection. Previous studies have used simulations to show that tree misspecification can lead to incorrect placement of substitutions on branches, possibly leading to spurious results for tests of positive directional selection within empirical datasets (Mendes and Hahn 2016). Here, we use empirical data in mice to show that these errors result in false positive (detected signal for selection only when using the gene tree) and false negative results (detected signal for selection only when using the species tree).

For each of three selection tests run, HyPhy’s BUSTED and aBSREL and PAML’s M1a vs. M2a some genes showed evidence of positive selection whether the species tree or gene tree was used while many other genes had signatures of positive selection restricted only to a single tree. The genes unique to the type of tree used were often discordant with the species tree. In contrast, the genes that showed evidence of positive selection regardless of tree had levels of discordance comparable to all genes (85%, Fig. 6, numbers in parentheses). This suggests that mis-mapping substitutions by supplying these tests with the wrong tree (*i.e.*, the species tree when gene trees are discordant) can lead to inflated false positive and false negative rates when inferring genes under positive selection. The magnitude and direction of these biases were dependent on the underlying model. So-called branch-site models that allow substitution rates to vary among both branches and codon sites, such as HyPhy’s BUSTED and aBSREL models, resulted in more genes inferred with evidence for positive selection when using the inferred gene tree (*i.e.*, the correct tree, assuming no errors in gene tree reconstruction). This means that using a single species tree for these tests reduces the power to detect positive selection. On the other hand, models that only allow rates to vary among sites, such as PAML’s M1a vs. M2a test, showed an increase in the number of putative false positives inferred when using the wrong tree. That is, tree mis-specification results in spurious increases in *dN/dS* that mimics positive directional selection.

These results have wide-ranging implications for phylogenetics and comparative genomic analysis. First, it is imperative that when testing a specific locus for positive selection, discordance among loci must be accounted for. This is most easily achieved by simply using the gene tree (or other locus type) as input to the test for selection (Good et al. 2013; Mendes and Hahn 2016; Roycroft et al. 2021). However, as Mendes and Hahn (2016) pointed out, this may not completely mask the effects of discordance on substitution rates, as sites within a single gene may still have evolved under different histories because of within-gene recombination. Indeed, we found that tree similarity diminished at scales that were less than the average genomic distance between the beginning and end of a coding sequence in mice (~37 kb in this data set). Nevertheless, starting with an inferred gene tree is advisable whenever possible, followed by a secondary analysis of evidence for within-gene variation in phylogenetic history. Our results also imply that studies of molecular evolution may benefit by from approach that reduces genome-wide levels of discordance, such as through *post hoc* pruning of species that disproportionately contribute to unresolved nodes.

Incorporating discordance into a comparative framework is not trivial and many comparative genomic methods assume a single species tree that test for changes in substitution rates in a phylogeny (Pollard et al. 2010; Hu et al. 2019; Partha et al. 2019). Even methods that allow the use of different trees for different loci (like PAML and HyPhy) are still commonly applied with a single species tree across loci (Carbone et al. 2014; Foote et al. 2015; van der Valk et al. 2021; Treaster et al. 2023). While we used the simplifying assumption that the results from the gene tree are more likely to be correct, this may not always be the case given that errors can also occur during gene tree inference. Still, our results confirm that the use of a single tree for such tests that rely on accurate estimation of substitution rates are likely to lead to both inaccurate inferences of positive selection and strongly encourage the use of individual gene trees for such analyses.

*Conclusions*

Murine rodents as a study system allow us to use the high-quality *M. musculus* genome to begin to examine fine-scale patterns and effects of phylogenetic discordance along chromosomes. Our analysis reveals how discordance varies with genome biology across evolutionary timescales, as well as the limits of inference inherent to extrapolating information from a single model system to a phylogenetic sample. We also demonstrate how phylogenetic discordance can mis-lead common tests for selection if only a single species tree is used. Through these results, we can better understand the complexities of phylogenomic datasets and ensure that steps are taken to accommodate these details in future comparative studies.

# Materials & Methods

## Sample collection and assembly

We collected genomes from 16 murine species and 2 other rodents from several sources, including NCBI and several recently sequenced in Kumon et al. (2021) (see Table S1 for full list of samples and sources). We also report the sampling of *Otomys typus* (FMNH 230007) from Ethiopia in 2015. While DNA extraction and sequencing on the 10x Genomics platform for *O. typus* is the same as described in (Kumon et al. 2021), the library quality for this sample was too low for chromosome level assembly. Here, we instead assembled it into scaffolds with the express purpose of obtaining UCEs for phylogenetic analysis. Adapters and low-quality bases were trimmed from the reads using illumiprocessor (Faircloth 2013), which makes use of functions from trimmomatic (Bolger et al. 2014). All cleaned reads were de novo assembled using ABySS 2.3.1 (Jackman et al. 2017) with a Bloom filter (Bloom 1970) de Bruijn graph. The final *O. typus* scaffold assembly was 2.14GB (N50=9,211; L50=64,014; E-size=12,790).

In parallel, for six of these species (see Fig. 1; Table S1), we generated reference-based pseudo-assemblies with iterative mapping using an updated version pseudo-it v3.1.1 (Sarver et al. 2017) that incorporates insertion-deletion variation to minimize reference bias in our genome-wide phylogenetic analyses and to maintain collinearity between assemblies (<https://github.com/goodest-goodlab/pseudo-it>). We used the *Mus musculus* (mm10) genome as the reference for our pseudo-assembly approach. Briefly, pseudo-it maps reads from each sample to the reference genome with BWA (Li 2013), calls variants with GATK HaplotypeCaller (Poplin et al. 2018), and filters SNPs and indels and generates a consensus assembly with bcftools (Danecek et al. 2021). The process is repeated, each time using the previous iteration’s consensus assembly as the new reference genome to which reads are mapped. In total, we did three iterations of mapping for each sample.

## Ultraconserved element (UCE) retrieval and alignment

We first set out to reconstruct a phylogeny of sequenced murine rodents to provide both a general resource for future comparative genomic studies within this important group as well as a time-calibrated phylogeny to frame an in-depth analysis of phylogenetic discordance across a subset of murine whole genomes (see below). We combined our seven recently sequenced genomes with nine publicly available genomes from other Old World mice and rats (subfamily Murinae) as well as the genomes of two non-murine rodents, the great gerbil (*Rhombomys opimus*; (Nilsson et al. 2020) and the Siberian hamster (*Phodopus sungorus*; (Moore et al. 2022) as outgroups. We extracted UCEs from each species, plus 1000 flanking bases from each side of the element using the protocols for harvesting loci from genomes and the *M. musculus* UCE probe set provided with phyluce v1.7.1 (Faircloth et al. 2012; Faircloth 2016). In total, we recovered 2,632 unique UCE loci, though not all UCE loci were found in all taxa (Table S1).

We brought the extracted UCE sequences for each species into a consistent orientation using MAFFT v7 (Katoh and Standley 2013) and then aligned them using FSA (Bradley et al. 2009) with the default settings. We trimmed UCE alignments with TrimAl (Capella-Gutierrez et al. 2009) with a gap threshold of 0.5 and otherwise default parameters. We performed alignment quality checks using AMAS (Borowiec 2016). We processed all alignments in parallel with GNU Parallel (Tange 2018).

## Species tree reconstruction from UCEs

We constructed a species-level rodent phylogeny with two approaches. First, using the alignments of all UCEs found in four or more taxa (2,632), we reconstructed a maximum-likelihood (ML) species tree with IQ-TREE v2.2.1 (Minh et al. 2020b). Each UCE alignment was concatenated and partitioned (Chernomor et al. 2016) such that optimal substitution models were inferred for individual UCE loci with ModelFinder (Kalyaanamoorthy et al. 2017). We also reconstructed individual gene trees for each UCE alignment. For all IQ-TREE runs (concatenated or individual loci), we assessed branch support with ultrafast bootstrap approximation (UFBoot) (Hoang et al. 2018) and the corrected approximate likelihood ratio test (SH-aLRT) (Guindon et al. 2010). We collapsed branches in each UCE tree exhibiting less than 10% approximated bootstrap support using the nw\_ed function from Newick Utilities (Junier and Zdobnov 2010). We used these trees as input to the quartet summary method ASTRAL-III v5.7.8 (Zhang et al. 2018) to infer a species tree. We generated visualizations of phylogenies with R v4.1.1 (R Core Team 2021) using phytools v1.9-16 (Revell 2012) and the ggtree package v3.14 (Yu et al. 2017; Yu 2020) and its imported functions from ape v5.0 (Paradis and Schliep 2019) and treeio v1.16.2 (Wang et al. 2020).

We then used two methods to assess phylogenetic discordance across the reconstructed species tree. First, we calculated site and gene concordance factors (sCF and gCF) with IQ-TREE 2 (Minh et al. 2020a; Minh et al. 2020b) to assess levels of phylogenetic discordance between the inferred UCE trees and the concatenated species tree. gCF is calculated for each branch in the species tree as the proportion of UCE trees in which that branch also appears (Baum 2007). sCF represents the proportion of alignment sites concordant with a given species tree branch in a randomized subset of quartets of taxa (Minh et al. 2020a). We visualized gCF and sCF (Lanfear 2018) for each branch in each species tree using methods in R v4.3.0 (Lanfear 2018; R Core Team 2021). Next, we used PhyParts (Smith et al. 2015b) to identify topological conflict between the UCE trees and the species tree from ASTRAL-III. For this analysis, we rooted all trees with *Phodopus sungorus* as the outgroup using the nw\_reroot function in the Newick Utilities (Junier and Zdobnov 2010) package and excluded 204 UCE trees that did not contain the outgroup.

## Divergence time estimation

We used IQ-TREE 2’s (Minh et al. 2020b) implementation of least square dating to estimate branch lengths of our species trees in units of absolute time (To et al. 2016). To improve divergence-time estimation, we used SortaDate (Smith et al. 2018) to identify a set of 100 UCE loci that exhibit highly clocklike behavior and minimized topological conflict with the concatenated species tree. We applied node age calibrations (Table S3) from Schenk et al. (2013) and Steppan and Schenk (2017), which in turn were sourced from fossil calibrations described on Paleobiology Database (2011). As *Rattus* is paraphyletic, the maximum age is taken from the earliest crown group fossil on Paleobiology Database (2011). In contrast, the estimated *Rattus* node age from Schenk et al (2013) was used as the minimum age. Branch lengths were resampled 100 times to produce confidence intervals. To return a single solution, least square dating typically requires that one calibration be fixed and not a range. We selected one calibration node (here, the branch leading to Murinae) and estimated dates across the tree when this node is set to its minimum, its maximum, and its midpoint ages. On the midpoint calibrated tree, we plot confidence intervals for each node representing the lowest minimum and highest maximum ages estimated across the three dating analyses.

## Genome window-based phylogenetic analysis

For the second part of our work, we wanted to quantitatively infer phylogenetic discordance across a subset of the murine genomes used to infer the species tree and relate that discordance to other features of the genome, such as recombination rate, proximity to genes, and rates of molecular evolution. To assess the distribution of phylogenetic discordance across rodent genomes, we limited subsequent analyses to *M. musculus* and the pseudo-assemblies (see above) of six of the genomes (*Mastomys natalensis*, *Hylomyscus alleni*, *Praomys delectorum*, *Rhabdomys dilectus*, *Grammomys dolichurus*, and *Rhynchomys soricoides*). *Otomys typus* was excluded from these analyses due to the inadequacy of the library outlined above.

We partitioned these genomes into 10 kilobase (kb) windows based on the coordinates in the reference *M. musculus* genome (mm10; Mouse Genome Sequencing et al. 2002) using bedtools makewindows (Quinlan and Hall 2010). These coordinates were converted between the reference and the consensus sequence for each genome using liftOver (Hinrichs et al. 2006). Note that this method assumes both collinearity of all genomes and similar karyotypes (see Discussion). We then removed windows from the subsequent analyses if (1) 50% or more of the window overlapped with repeat regions from the *M. musculus* reference RepeatMasker (Smit et al. 2013-2015) file downloaded from the UCSC Genome Browser’s table browser (Hinrichs et al. 2006) or (2) 50% or more of the window contained missing data in 3 or more samples. Overlaps with repeat regions were determined with bedtools coverage (Quinlan and Hall 2010). We then aligned the 10kb windows with MAFFT (Katoh and Standley 2013), trimmed alignments with trimAl (Capella-Gutierrez et al. 2009), and inferred phylogenies for each with IQ-TREE 2 (Minh et al. 2020b) which uses ModelFinder to determine the best substitution model for each window (Kalyaanamoorthy et al. 2017).

To assess patterns of tree similarity between windows on the same chromosome, we used the weighted Robinson-Foulds (wRF) (Robinson and Foulds 1981; Böcker et al. 2013) distance measure implemented in the phangorn library (Schliep 2011) in R (R Core Team 2021), which compares two trees by finding clades or splits present in one tree but not the other weighted by the missing branch length in each tree for each mismatch and differences in branch length between the co-occurring branches in both trees (Robinson and Foulds 1979). Consequently, the resulting measure of wRF is in units of branch length (i.e., expected number of substitutions per site for maximum likelihood trees). We compared wRF between trees from windows on the same chromosome to characterize (1) heterogeneity in patterns discordance along the chromosome and (2) whether tree similarity is correlated with distance between windows. For the second question, we sampled every window on a chromosome at increasing distance (in 10kb windows) until the distribution of wRF scores for all pairs of windows at that distance was not significantly different (Wilcox test, p > 0.01) than that of a sample of 12,000 measures of wRF between randomly selected trees on different chromosomes. We selected 12,000 as the random sample size because it roughly matched the number of windows on the largest chromosome (chromosome 1, n = 12,113). We used Snakemake 7 (Mölder et al. 2021) to compute window alignments and trees in parallel.

*Whole genome alignment between mouse and rat*

To assess how un-accounted for large-scale structural variation may impact our conclusions, we compared the reference mouse and rat genomes. We used minimap2 (Li 2018) to align the mouse (mm10) and rat (rnor6) (Gibbs et al. 2004) genomes to assess the impact of structural variation that spans the divergence of our subset of species used to in the discordance analyses. We downloaded the rat reference genome (rnor6) from the UCSC genome browser and for both genomes removed the Y chromosome and all smaller unplaced scaffolds. We then used minimap2 in whole genome alignment mode (-x asm20) to generate a pairwise alignment file from which we calculated alignment segment sizes and the distances between alignment segments. We visualized the alignment as a dot plot using the pafr package in R (<https://github.com/dwinter/pafr>).

## Recombination rate and functional annotation

We retrieved 10,205 genetic markers generated from a large heterogenous stock of outbred mice (Shifman et al. 2006; Cox et al. 2009) to assess whether phylogenetic discordance along chromosomes was correlated with mouse recombination rates. We converted the physical coordinates of these markers from build 37 (mm9) to build 38 (mm10) of the *M. musculus* genome using liftOver (Hinrichs et al. 2006). We then partitioned the markers into 5Mb windows and estimated local recombination rates defined as the slope of the correlation between the location on the *M. musculus* genetic and physical maps for all markers in the window (White et al. 2009; Kartje et al. 2020). Within each 5Mb window, we calculated wRF distances between the first 10kb window and every other 10kb window.

We also compared the chromosome-wide wRF distances to those based on phylogenies from regions around several types of adjacent to genomic features. We retrieved coordinates from 25,753 protein coding genes annotated in *M. musculus* from Ensembl (release 99; Cunningham et al. 2022), all 3,129 UCEs from the *M. musculus* UCE probe set provided with PHYLUCE (Faircloth et al. 2012; Faircloth 2016), and 9,865 recombination hotspots from Smagulova et al. (2011). The recombination hotspot coordinates were converted between build 37 and build 38 using the liftOver tool (Hinrichs et al. 2006). For each feature, the starting window was the 10kb window containing the feature’s midpoint coordinate. We then calculated wRF between this window and all windows within 5Mb in either direction and for each chromosome compared the slope and wRF distance of windows adjacent to the feature with the same metrics for the whole chromosome. We compared distributions of these measures for each genomic feature with an ANOVA (aov(feature.measure ~ feature.label)) followed by Tukey’s range test (TukeyHSD(anova.result)) to assess differences in means, as implemented in R v4.1.1 (R Core Team 2021).

## Molecular evolution

To test how tree misspecification affects common model-based analyses of molecular evolution, we retrieved 22,261 coding sequences from *M. musculus* using the longest coding transcript of each gene. Coding coordinates from the *M. musculus* coding sequences were transposed to the new assemblies via liftOver (Hinrichs et al. 2006) and sequences retrieved with bedtools getfasta (Quinlan and Hall 2010). We recovered 17,216 genes that were present in all 7 species. Using MACSE (Ranwez et al. 2018), we trimmed non-homologous regions from each ortholog using trimNonHomologousFragments, aligned the orthologs using alignSequences, and trimmed the aligned sequences with trimAlignment to remove unaligned flanking regions. Finally, we manually filtered the alignments using the following (non-mutually exclusive) criteria: 3,368 alignments were removed during filtering for gapped sites, 3,132 alignments had a premature stop codon in at least one species, 1,571 alignments had only 3 or fewer unique sequences among the 7 species, and 78 alignments were shorter than 100 bp. After filtering, 12,559 total alignments for tree reconstruction and inference of selection.

We then used IQ-TREE 2 (Minh et al. 2020b) to reconstruct a single species tree from concatenation of all gene alignments, as well as gene-trees for each individual alignment. This species tree from coding regions matches the topologies of these species inferred by concatenation of UCEs in the previous section. Next we ran several tests that use both coding alignments and a tree to infer positive selection: PAML’s M1a vs. M2a test (Yang 2007), HyPhy’s aBSREL model (Smith et al. 2015a), and HyPhy’s BUSTED model (Murrell et al. 2015). We ran each test twice on each gene, once using the species tree derived from concatenated data, and once using the tree estimated for that gene. For the HyPhy models, no target branch was selected, meaning all branches in the input phylogeny were tested.

The end point of each of these three tests is a p-value, which lets us assess whether a model that allows for positively selected sites fits better than a model that does not. For M1a vs. M2a, we obtained the p-value manually by first performing a likelihood ratio test to determine genes under selection by calculating . The p-value of this likelihood ratio is then retrieved from a one-tailed chi-square distribution with 2 degrees of freedom (Yang 2007). For BUSTED and aBSREL, p-values are computed automatically during the test using similar likelihood ratios. For the M1a vs. M2a and BUSTED tests, a single p-value is computed for each gene. P-values were adjusted by correcting for false discovery rates (Benjamini and Hochberg 1995; Yekutieli and Benjamini 1999) using the “fdr” method in the p.adjust() function in R (R Core Team 2021) and we categorized a gene as being positively selected if its adjusted p-value was < 0.01. For the aBSREL test, a p-value is generated for each branch in the input gene tree. aBSREL corrects for multiple testing internally across branches using the Holm-Bonferroni procedure (Holm 1979; Pond et al. 2005). We further correct the p-values across genes with the Bonferroni method and classify a gene as having experienced positive selection if one or more branches has a p-value < 0.01 after all corrections. We used Snakemake 7 (Mölder et al. 2021) to compute coding alignments, trees, and selection tests in parallel.

# Data availability

For the six previously assembled genomes (see Table S1), all raw reads and assemblies are available as an NCBI BioProject (Accession Number PRJNA669840). The reads and assembly for *Otomys* typus, pseudo-assemblies for the six other new samples, and locus alignments (UCEs, genes, and genomic windows) are available on Dryad (<https://doi.org/10.5061/dryad.866t1g1wq>). All code and summary data for this project are deposited on github (<https://github.com/gwct/murine-discordance>).

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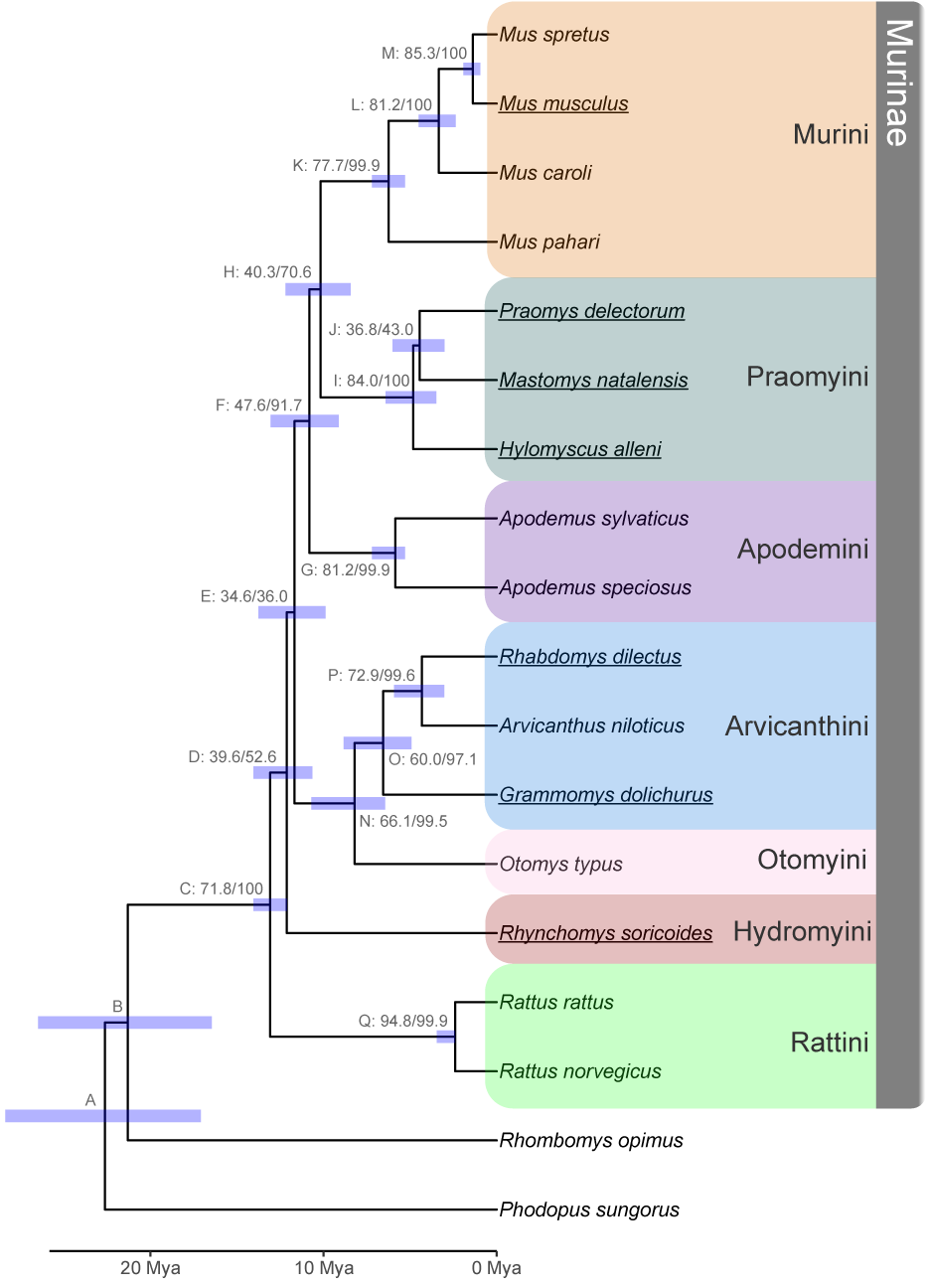
# Tables

### Table 1. Rates and types of error when using concatenated trees for gene-based selection tests. Assuming the gene tree topology is the correct topology.

|  |  |  |
| --- | --- | --- |
| Test | False positive rate | False negative rate |
| BUSTED | 0.45% | 28.10% |
| aBSREL | 0.41% | 10.60% |
| M1a vs. M2a | 2.66% | 3.20% |

# Figures

### Figure 1



**Figure 1.** Species trees inferred from concatenation of ultra-conserved elements (UCEs) from 18 rodent species. Internal nodes are labeled by a letter identifier referenced in the text and site and gene concordance factors (*i.e.*, Label: sCF/gCF) as well as a bar indicating the confidence interval for divergence time estimation. Ultrafast bootstrap/SH-aLRT values were all 100. Bottom scale represents time in millions of years before present. Fossil calibrations are described in Tables 2 and S2, with node C used as a fixed calibration point. Tribes within sub-family Murinae are highlighted on the right following the classifications used by Lecompte et al. (2008). Genomes used for the genome-wide phylogenetic discordance analyses are underlined.

### Figure 2



**Figure 2.** The landscape and profile of phylogenetic discordance across non-overlapping 10kb windows in murine genomes. A) Distribution of the 20 most frequent topologies recovered across all windows. Numbers above bars indicate proportion of each topology. B) The top three topologies recovered across all chromosomes 1. C) Distribution of the topologies recovered along chromosome 1. The x-axis is scaled to the length of the chromosome and each vertical bar represents one 10kb window. The three most frequent topologies occupy the first three rows while all other topologies are shown in the bottom row. See Supplemental File S1 for individual chromosome plots.

### Figure 3



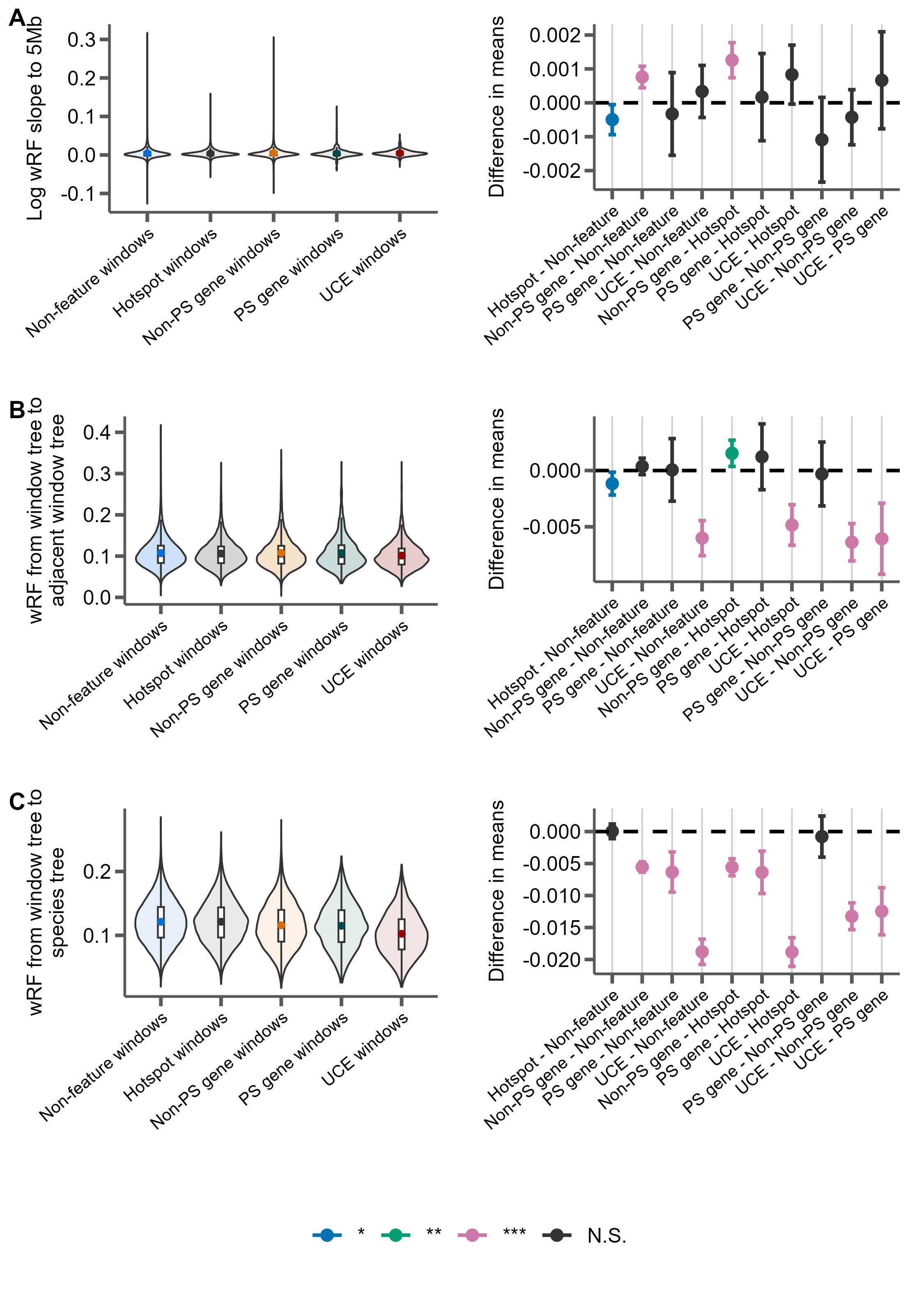
**Figure 3.** Similarity between 10kb windows decays as genomic distance between windows increases. A) The log fit to the mean of distributions of weighted Robinson-Foulds distances between trees of windows at increasing genomic distance (10kb steps). Each line represents one chromosome. B) The same, but on a log scale with a linear fit. C) For every window on each chromosome, the genomic distance between windows at which tree distance becomes random for 100 replicates of random window selection. D) Points represent the slopes of the correlation between genomic distance and tree distance from panel B represent the rate at which tree similarity decays across the genome. Dark grey dashed line is median slope and light grey dashed line is mean.

### Figure 4



**Figure 4.** Correlations between tree similarity and recombination rate in 5 Mb windows. A) Tree similarity as measured by the weighted Robinson-Foulds distance between the first and last 10 kb windows within the 5 Mb window. B) The slopes of the linear correlation between the weighted Robinson-Foulds distances between the first 10 kb window and every other 10 kb window within a 5 Mb window represent the rate at which tree similarity decays over each 5 Mb window. C) The mean wRF of all 10 kb window trees within each 5Mb window compared to the species tree.

### Figure 5



**Figure 5.** Distributions of weighted Robinson-Foulds distance from trees constructed from 10kb windows either centered on recombination hotspots (Hotspot), protein-coding genes without evidence for positive selection (Non-PS genes), protein coding genes with evidence for positive selection (PS genes), UCEs, or containing none of these features (Non-feature). For each panel, the left portion shows the distributions of the measure for each feature type and the right panel shows the differences in means for each pairwise comparison of features with significance assessed with Tukey’s range test. The labels on the x-axis indicate the feature pairs being compared, with the first feature being the reference (*i.e.* points above 0 indicate this feature has a higher mean). *P-*value thresholds: \* < 0.05, \*\* < 0.01, \*\*\* <0.001. A) The rate of decay of phylogenetic similarity is calculated as the slope of a linear regression between wRF and the log distance between each window up to 5Mb away from the feature window. B) The phylogenetic similarity of windows immediately adjacent to feature windows. C) The phylogenetic similarity between the species tree inferred from protein-coding gene trees and the feature window.

### Figure 6



**Figure 6.** Tree misspecification leads to erroneous results in tests for positive selection. The proportion of genes inferred to be under positive selection for three tests using either a single species tree (concatenated tree) or individual gene trees, as well as those found in both cases (shared). Numbers in the bars indicate raw counts, and percentages indicate the percent of genes in that category that are discordant from the species tree.

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