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1 **Title:** Genome assembly of the dyeing poison frog provides insights into
2 the dynamics of transposable element and genome-size evolution

3
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1 **Abstract**

2 Genome size varies greatly across the tree of life and transposable elements are an important
3 contributor to this variation. Among vertebrates, amphibians display the greatest variation in
4 genome size, making them ideal models to explore the causes and consequences of genome
5 size variation. However, high-quality genome assemblies for amphibians have, until recently,
6 been rare. Here, we generate a high-quality genome assembly for the dyeing poison frog,
7 *Dendrobates tinctorius*. We compare this assembly to publicly available frog genomes and find
8 evidence for both large-scale conserved synteny and widespread rearrangements between frog
9 lineages. Comparing conserved orthologs annotated in these genomes revealed a strong
10 correlation between genome size and gene size. To explore the cause of gene-size variation,
11 we quantified the location of transposable elements relative to gene features and find that the
12 accumulation of transposable elements in introns has played an important role in the evolution
13 of gene size in *D. tinctorius*, while estimates of insertion times suggest that many insertion
14 events are recent and species-specific. Finally, we carry out population-scale mobile-element
15 sequencing and show that the diversity and abundance of transposable elements in poison frog
16 genomes can complicate genotyping from repetitive element sequence anchors. Our results
17 show that transposable elements have clearly played an important role in the evolution of large
18 genome size in *D. tinctorius*. Future studies are needed to fully understand the dynamics of
19 transposable element evolution and to optimise primer or bait design for cost-effective
20 population-level genotyping in species with large, repetitive genomes.

21

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23

24 **Keywords** Amphibians, genome evolution, transposable elements, long-reads, MobiSeq

1 **Significance**

2 Amphibians display more variation in genome size than any other vertebrate lineage.
3 Complexities associated with large genomes frequently hamper genome assembly and
4 population genetic studies. Here we use long-read HiFi sequences to generate a high-quality
5 6.3 Gb genome assembly of the poison frog *Dendrobates tinctorius*. We use this genome and
6 leverage comparative genomics and *de novo* annotations to quantify aspects of genome
7 evolution driven by repetitive transposable genetic elements. Our results provide support for the
8 dynamic role that transposable elements play in driving the evolution of “genomic gigantism” in
9 amphibians. We also show how transposable elements can be leveraged for cost-efficient
10 population genetic studies using limited input material.

11

1 Introduction

2 Interspecific variation in genome size is a fundamental feature of biodiversity, and
3 transposable elements play an important role in contributing to variation in genome size and
4 structure (Kidwell 2002; Hawkins et al. 2006; Lee and Kim 2014). While historically referred to
5 as “junk” DNA, it has been long known that the evolution of transposable elements can have
6 profound effects on an organism's phenotype. For example, the ability of transposable elements
7 to drive gene expression and mosaic coloration in maize accompanied their discovery in the late
8 1940s by Barbara McClintock (McClintock 1950). With the advancement of computational
9 capabilities and the availability of cost-effective genetic methods, evidence of the phenotypic
10 effects of transposable elements has increased. It is now widely appreciated that transposable
11 elements can have a profound effect on evolution by influencing exon structure, telomeres,
12 gene expression, and ultimately, adaptation or speciation (Almojil et al. 2021; Casacuberta and
13 González 2013; Feschotte 2008; Serrato-Capuchina and Matute 2018). It has also been
14 suggested that the mobilisation and insertion of transposable elements can drive adaptation to
15 global change when organisms are subject to environmental stress (Pimpinelli and Piacentini
16 2020; Rey et al. 2016). For a comprehensive classification and in-depth review of the impact
17 transposable elements can have on the genome, we recommend the recent works of (Almojil et
18 al. 2021; Bourge et al. 2018; Kidwell and Lisch 1997; Sotero-Caio et al. 2017; Wicker et al.
19 2007).

20 Characterising the genomic landscape of transposable elements—such as the
21 abundance of different types of elements and where they are located in the genome relative to
22 genes, exons, and introns—is one approach that can be used to illuminate aspects of their
23 evolution. With the increasing availability of whole-genome sequence data collected from many
24 species, studies are increasingly quantifying the genomic landscape and “ecology” of
25 transposable elements (Lamichhaney et al. 2021; Stitzer et al. 2021; Gozashti et al. 2023). A

1 general pattern emerging from these studies is that different types of transposable elements
2 contribute to genome evolution in different species. For example, long terminal repeat (LTR)
3 retrotransposons play an important role in genome size variation among plants (Lee and Kim
4 2014); while long and short interspersed nuclear elements (LINEs and SINEs, respectively) are
5 more abundant than LTRs in mammals (Chalopin et al. 2015; Platt et al. 2018), LINEs are more
6 abundant than SINEs and LTRs in bird (Zhang et al. 2014) and squamate genomes (Pasquesi
7 et al. 2018), and SINEs are nearly absent in amphibian genomes (Zuo et al. 2023). In addition
8 to the types of repetitive elements present within genomes, estimates of the timing of when
9 different repetitive elements insert themselves vary both across species (Sun et al. 2015) and
10 among the types of repetitive elements present within single genomes (Sun et al. 2015; Stitzer
11 et al. 2021). Finally, different types of repetitive elements can display different insertion-site
12 preferences or biases, with some inserting themselves into non-random locations in the
13 genome, such as in intergenic regions, promoters, or introns (Bourque et al. 2018; Stitzer et al.
14 2021). Gaining a better understanding of the insertion-site preferences or biases displayed by
15 different repetitive elements across different species is important because it will help facilitate a
16 predictive understanding of how repetitive elements contribute to the evolution of genomic and
17 genetic variation.

18 Among vertebrates, amphibians exhibit remarkable variation in genome size, surpassing
19 that of any other group (Liedtke et al. 2018). The variation in amphibian genome size is
20 influenced by the activity and accumulation of transposable elements (Sotero-Caio et al. 2017).
21 However, the limited availability of high-quality amphibian genome assemblies—in large part
22 due to the challenges of assembling large repetitive genomes with short-read sequencing
23 technologies—makes the evolutionary dynamics and genomic ecology of TEs, alongside their
24 impact on phenotypes and adaptation, difficult to test (however, see Zuo et al. 2023).

1 As complete genomes are often lacking for amphibians, the research community uses
2 reduced representation library sequencing for studies of species delimitation and population
3 genetics (Dufresnes et al. 2018; Funk et al. 2018; Homola et al. 2019; Nunziata and Weisrock
4 2018). Given their transposable element-rich genomes, it is surprising that methods that
5 leverage repetitive elements, such as MobiSeq (Rey-Iglesia et al. 2019), have not yet been
6 applied to amphibians. MobiSeq is a method for constructing a reduced representation library by
7 targeting the flanking regions of transposable elements to identify single nucleotide
8 polymorphisms (SNPs) and genotypes (Rey-Iglesia et al. 2019). This approach requires minimal
9 DNA input and does not necessarily require a reference genome, making it useful for
10 investigating a wide array of research questions with regards to population genetics,
11 evolutionary dynamics and ecological interactions.

12 Poison frogs (Family Dendrobatidae) are a group of Central- and South American forest-
13 dwelling amphibians with complex social behaviour and elaborate parental care (Stynoski et al.
14 2015). A thriving research community has focused on this model clade for studying the effects
15 of natural selection on phenotype, particularly the bright colouration coupled with chemical
16 defences that protects some species from predation (Noonan and Comeault 2009; Chouteau et
17 al. 2011; Maan and Cummings 2012; Lawrence and Rojas et al. 2019). A solid natural history
18 background stemming from field observations done in the late 1900s (e.g. Sexton 1960,
19 Silverstone 1973; Wells 1980; Myers and Daly 1983; Donnelly 1989; Summers 1989) had laid a
20 firm foundation to ask both ultimate and proximate research questions. For example, research
21 on the ultimate factors shaping their communication and territorial behaviour, parental care, and
22 space use, as well as the behaviour of their larvae, has been steadily gaining traction (e.g.,
23 Pröhl 2005; Summers et al. 2006; Amézquita et al. 2011; Ringler et al. 2013; Schulte et al.
24 2013; Tumulty et al. 2014; Rojas 2014; Stynoski et al. 2014; Carvajal-Castro et al. 2021;
25 Fouilloux et al. 2021). More recently, studies on the proximate mechanisms of such behaviours,

1 e.g., neurobiology of egg provisioning, tadpole transport and tadpole aggression (Fischer and
2 O'Connell 2020; Fischer et al. 2019, 2020), and hormonal correlates of care, territoriality and
3 space use (Fischer and O'Connell 2020; Pašukonis et al. 2022; Rodríguez et al. 2022), now
4 provide a more holistic understanding of what makes these frogs unique. However, the lack of
5 genomic resources has precluded some of these topics from being addressed in depth. The one
6 genome available for dendrobatid frogs (*Oophaga pumilio*) is highly fragmented, but still shows
7 that transposable elements comprise a large portion of it (Rogers et al. 2018). Having access to
8 reference genomes from more poison frogs would allow for more comprehensive approaches to
9 questions related to demography, conservation, behavioural ecology, disease dynamics and
10 adaptation (Brandies et al. 2019).

11 In this study, we generate a high-quality reference genome for the dyeing poison frog,
12 *Dendrobates tinctorius*. This species is aposematic—with drastic differences in coloration and
13 toxicity across populations (Lawrence and Rojas et al. 2019)—and displays complex social
14 behaviours typical of many poison frog species including male parental care and territoriality
15 (Rojas and Pašukonis 2019; Fouilloux et al. 2021; Rojas 2014; 2015). We first leverage our
16 genome assembly and three publicly available chromosome-scale assemblies of species
17 belonging to Hyloidea to provide evidence for both large regions of synteny alongside significant
18 structural evolution. This analysis also revealed that the evolution of genome size across these
19 species is correlated with gene size, with *D. tinctorius* having both the largest genome and the
20 longest genes. We then annotate transposable elements in the *D. tinctorius* genome to explore
21 the genomic landscape of their evolution and find that transposable elements are more
22 abundant in introns than in exons, likely contributing to the evolution of large genes in *D.*
23 *tinctorius*. Finally, we explored the usefulness of the *D. tinctorius* reference genome in
24 leveraging population genetic information from cost-effective mobile element sequencing
25 (MobiSeq, Rey-Iglesia et al. 2019). Applying this method to tadpoles collected from a wild

1 population revealed that the highly repetitive nature of the *D. tinctorius* genome can generate
2 challenges to MobiSeq genotyping, with a high degree of missing data across sites and
3 individuals. We hope that the ability to use the *D. tinctorius* genome as a resource will refine the
4 development of markers—such as additional mobile-element tags or alternate approaches such
5 as baited capture—for genotyping individuals, thereby broadening the scope of behavioural
6 ecology and population genomic research in amphibians.

7

8 **Results**

9 *Genome assembly, quality control*

10 Our final *Dendrobates tinctorius* assembly consists of 6.356 Gb assembled into 830
11 contigs. This assembly has a contig N50 and L50 of 32.539 Mb and 56 contigs, respectively,
12 with a maximum contig size of 131.4 Mb. Our assembly is also highly accurate and complete,
13 achieving an error rate of less than 0.0001—as indicated by a Quality Value score (Chen et al.
14 2021) of 41—and containing 4,796 of 5,310 curated tetrapod BUSCO genes (BUSCO summary:
15 C:90.3% [S:88.4%, D:1.9%], F:2.8%, M:6.9%, n:5310). While our assembly does not contain
16 complete chromosomes, the contiguity and BUSCO scores were comparable to chromosome-
17 level assemblies of other anurans available on NCBI (fig. 1).

18

19 *Synteny between D. tinctorius and other Hyloidea genomes*

20 To explore patterns of conserved synteny between our assembly and closely related
21 Anuran species, we compared the *D. tinctorius* assembly to chromosome-scale assemblies of
22 *Bufo gargarizans* (Asiatic toad; Bufonidae), *Eleutherodactylus coqui* (Common coquí;
23 Eleutherodactylidae), and *Engystomops pustulosus* (Túngara frog; Leptodactylidae). *Bufo*
24 *gargarizans*, *El. coqui*, and *En. pustulosus* shared a common ancestor with *D. tinctorius* ~65
25 Mya (Feng et al. 2017), and, while these four species have significantly diverged from one

1 another, the former three were used for comparisons because they represent the three most
2 closely related genera to *Dendrobates* that currently have publicly available chromosome-scale
3 genome assemblies. Orthology-guided synteny analyses based on the tetrapoda BUSCO
4 geneset (Simão et al. 2015; v5.2.2) in GENESPACE (Lovell et al. 2022; v1.3.1) provided
5 evidence of both broad-scale synteny and genome evolution between *D. tinctorius* and the three
6 other species we analysed (fig. 2). The number of synteny blocks identified between *D.*
7 *tinctorius* and the chromosome-scale assemblies ranged from 116 (*En. pustulosus*) to 164 (*B.*
8 *gargarizans*), with an average size of each block of 27.36 Mb (*En. pustulosus*) to 17.86 Mb (*B.*
9 *gargarizans*). We focused this analysis on scaffolds of the *D. tinctorius* assembly that were
10 greater than 20 Mb in length (see Methods), and these *D. tinctorius* scaffolds have median and
11 mean lengths of 32.64 Mb and 41.81 Mb, respectively. Synteny between *D. tinctorius* scaffolds
12 and the chromosome-scale assemblies therefore tends to span roughly half of entire *D.*
13 *tinctorius* scaffolds, on average, while in many cases entire scaffolds showed collinearity with
14 chromosomal regions of the chromosome-scale assemblies (fig. 2B and C). In addition to
15 collinear regions, we observed numerous rearrangements between *D. tinctorius* scaffolds and
16 chromosomes of the other Anuran genomes we analysed: for example, Figure 2B highlights
17 rearrangements between *D. tinctorius* scaffolds and *B. gargarizans*'s chromosome 1 (inversions
18 highlighted in salmon pink colour). Finally, we observed a single instance where a *D. tinctorius*
19 scaffold contained synteny blocks that mapped to different chromosomes of a chromosome-
20 scale assembly (*El. coqui* chromosomes 1 and 4; fig. 2C). These results suggest that—at least
21 at the course scale of ~65 My of evolution—rearrangements within chromosomes are more
22 numerous than changes in the overall evolution of chromosome number via fission and/or fusion
23 events.

24 Our *D. tinctorius* assembly is larger than any of the genomes we used in the analyses
25 presented above (*O. pumilio*, *B. gargarizans*, *El. coqui*, *En. pustulosus*, *R. temporaria* and
26 *Xenopus tropicalis*), with genome size ranging from 1.451 Gb (*X. tropicalis*) to 6.356 Gb (*D.*

1 *tinctorius*). We therefore explored how genome size evolution affects the size of gene-regions
2 by comparing the size of BUSCO annotations across five chromosome-scale genomes and the
3 *D. tinctorius* assembly. For this analysis, we excluded the *O. pumilio* assembly due to high
4 levels of fragmentation and included an assembly of *R. temporaria* to increase the phylogenetic
5 diversity of genomes being analysed — *R. temporaria* is outside of Hyloidea yet more closely
6 related to Hyloidea than *X. tropicalis* (fig. 3A). Across 1,949 single copy orthologs that were
7 present in all genomes, *D. tinctorius* orthologs ranged from 39.8 to 207.8% longer, on average,
8 than the same orthologs in the *R. temporaria* and *X. tropicalis* assemblies, respectively. This
9 percent difference in gene size was negatively correlated with the percent difference in genome
10 size (Spearman's $\rho = 0.9$; $P = 0.083$), with larger genomes (i.e., *B. gargarizans* and *R.*
11 *temporaria* versus *D. tinctorius*) showing less difference in genes size than when genomes
12 differed in size (i.e., *X. tropicalis* versus *D. tinctorius*). In general, we found a positive correlation
13 between genome size and the average size of BUSCO gene regions ($\rho = 0.94$; $P = 0.017$, fig.
14 3B), which was independent of phylogeny (Spearman's correlation on phylogenetically
15 independent contrasts: $\rho = 0.9$; $P = 0.083$; fig. 3C), indicating that the evolution of larger
16 genomes in *D. tinctorius*, *B. gargarizans*, and *R. temporaria* has resulted in the evolution of
17 larger gene regions. Because we focused on conserved orthologous gene regions that include
18 introns, this pattern is likely driven by larger introns in larger genomes, and genome size is not
19 correlated with predicted BUSCO amino acid lengths ($\rho = -0.14$; $P = 0.80$).

21 *Genome annotation and transposable element diversity*

22 The majority of the *D. tinctorius* genome consists of repetitive elements: annotation with
23 RepeatModeler (Flynn et al. 2020; v2.0.3) + RepeatMasker (Smith et al. 2021; v4.1.2-p1)
24 identified 74.72% of our assembly as repetitive. Many of the repeats classified by
25 RepeatMasker could not be assigned to specific types of repetitive elements (39.15%). Among

1 assigned elements, 23.13% were retroelements and 12.45% were DNA transposons. The most
2 common retroelements classified by RepeatMasker were LTR retrotransposons (15.69% of the
3 assembly) and LINEs (7.4%). LTR retrotransposons Ty3/*DIRS1* were the most abundant
4 superfamily (13.79% of the assembly), while DNA transposons made up 12.45% of the
5 assembly. Of DNA transposons, *Tc1-IS630-Pogo* (*Tc1*) and *hobo-Activator* (*hAT*) elements
6 were the most abundant (6.57% and 3.91% of the assembly, respectively).

7 Independent analyses with LtrDetector (Valencia and Girgis 2018) annotated 262,486
8 LTRs spanning 2,343,246,106 bp, or 36.9% of our assembly. While this represents a higher
9 percentage of LTRs in our assembly than when annotated with RepeatMasker, consistent with
10 RepeatMasker, the most abundant LTR elements were found to belong to the Ty3/*DIRS1*
11 superfamily (99,884 elements, 819.4 Mb total), followed by *BEL/Pao* elements (1,762 elements,
12 13.9 Mb total). Also consistent with results from RepeatMasker, 66,046 of the LTR elements
13 annotated by LtrDetector (spanning 588 Mb) had BLAST matches to the “Unknown” category of
14 elements identified by RepeatModeler.

15 To better understand the role that repetitive elements have played in the evolution of *D.*
16 *tinctorius* genome structure, we compared the location of repetitive elements relative to *de novo*
17 gene annotations we generated using the BRAKER2 pipeline (Brůna et al. 2021; v2.1.6). Of the
18 11,331,718 repeat annotations generated by RepeatMasker, 115,330 (1.02%) overlapped with
19 coding DNA sequence (CDS) and 4,872 of these had a reciprocal overlap of at least 75% (table
20 1 and table 2, respectively). More repetitive elements overlapped with introns compared to CDS
21 (525,441 elements; $\chi^2 = 270,118$; $P < 2.2 \times 10^{-16}$); however, proportionally fewer TEs that
22 overlapped introns showed reciprocal overlaps of greater than 75% compared to those
23 overlapping CDS (1.05 versus 4.22% respectively; $\chi^2 = 5,956.1$; $P < 2.2 \times 10^{-16}$). These
24 patterns are likely due to the fact that introns span nearly 4 times the number of bases and have

1 a median size nearly twice that of CDS features (266 Mb versus 69 Mb and 113 versus 66 bp,
2 respectively).

3 We next considered positions of the four most abundant classes/families of repetitive
4 elements annotated by RepeatMasker in the *D. tinctorius* genome—LTRs, LINEs, *Tc1* DNA
5 transposons, and *hAT* DNA transposons—in relation to CDS and introns. Of these four
6 classes/families of repeats, LTRs and *Tc1* transposons showed the highest overlap with CDS
7 and introns, with 0.33% (3,500) of LTR elements and 1.14% (10,227) of *Tc1* transposons
8 showing at least partial overlap with CDS, and 4.09% (42,844) and 5.11% (45,806) overlapping
9 with introns, respectively. Only *Tc1* transposons were enriched in the proportion overlapping
10 with CDS and introns compared to all non-*Tc1* elements (CDS: 1.14% versus 1.01%,
11 respectively; introns: 5.27% versus 4.65%, respectively).

12 Given that a large portion of the *D. tinctorius* assembly is comprised of LTR elements—
13 15.69% to 36.9% of the assembly annotated by RepeatModeler or LtrDetector, respectively—
14 we next explored length distributions of these elements and estimated the timing of their
15 insertion by estimating divergence between the left and right LTR of each element, assuming a
16 substitution rate of 2.5×10^{-9} substitutions per site per year (Lau et al. 2020). Average estimates
17 of insertion times for different types of LTR retroelements ranged from 10 to 28 Mya (fig. 4).
18 Retroelements with BLAST matches to the *DIRS* order of elements had the oldest average
19 estimated insertion time (mean = 23.35 Mya, 95% empirical range = 1 - 60.4 Mya) while *ERV1*
20 elements had the youngest insertion times (mean = 10.4 Mya, 95% empirical range = 0 - 45.1
21 Mya). The broad range of insertion times we estimated indicate that some LTR retroelements
22 are old and may be shared with other species of poison frog (the MRCA of Dendrobatidae is
23 estimated to have occurred ~36 Mya; Hime et al. 2018), while others are young, potentially
24 active, and species-specific (e.g. *Dendrobates* split from its sister genus *Oophaga* ~16 Mya;
25 Guillory et al. 2019).

1 *A test of the utility of MobiSeq for population genetics*

2 We next used the *D. tinctorius* reference genome to compare the genotyping success of
3 three genotyping approaches using a MobiSeq dataset generated from 87 tadpoles collected
4 from 17 phytotelmata at the Nouragues Research Station, French Guiana. Specifically, we
5 compared the number of usable SNPs generated by either (1) de novo assembly of MobiSeq
6 reads, (2) mapping the reads to the other poison frog genome currently available on NCBI (*O.*
7 *pumilio*), or (3) mapping the reads to the *D. tinctorius* genome we generated as part of this
8 study. We also called SNPs using two approaches: either the stacks pipeline (Catchen et al.
9 2011, 2013; v2.64) designed to assemble RADseq data de novo or using a reference genome;
10 or the original MobiSeq pipeline using the program "analysis in next generation sequencing
11 data" (ANGSD; Korneliussen et al. 2014; v0.940) mapping sequence reads to the *D. tinctorius*
12 assembly (Rey-Iglesia et al. 2019) (details in Methods).

13 The number of SNPs differed considerably between the two primers and different
14 mapping approaches (de novo vs. reference genome) and showed high degrees of missing
15 genotypes when genotyped using the stacks pipeline (LINE109: $93.21 \pm 3.06\%$, TE644: $90.46 \pm$
16 1.46% ; mean percent missing data across sites \pm sd) and low coverage (supplementary table
17 S1, Supplementary Material online). Regions amplified with the LINE109 primer resulted in
18 considerably fewer SNPs than the TE644 primer. When we mapped sequences to the *O.*
19 *pumilio* genome, gstacks only incorporated 10-12% of the reads and called considerably fewer
20 SNPs than either the de novo approach or mapping to the *D. tinctorius* genome. The de novo
21 approach with 50bp fragments called the highest numbers of SNPs (n SNPs TE644 = 92,433);
22 in contrast, the 100bp fragments called the lowest numbers (n SNPs TE644 = 15,987). These
23 differences were expected due to the large loss of SNPs due to truncation. Mapping to the *D.*
24 *tinctorius* reference genome for both primers called the second most SNPs (n SNPs; LINE109 =
25 2,476, TE644 = 82,863), but missingness was still high and coverage low (supplementary table

1 S1 and S2, Supplementary Material online). After applying a strong filter, where SNP had to be
2 present in at least 50% of individuals, the numbers of SNP dropped dramatically (table S2 and
3 fig. 5B).

4 The SNP calling with the ANGSD pipeline gave $17.67 \pm 0.17\%$ missing genotypes and
5 345 SNPs were called when the output was restricted to be present in at least 70 individuals. If
6 SNPs were restricted to be present in at least 80 individuals, 291 SNPs could be called with
7 $11.07 \pm 0.08\%$ genotype missingness (fig. 5A and B).

8 We used our called SNP dataset to test their usefulness in parentage analysis.
9 Specifically, we ran a relatedness analysis with COLONY (Jones and Wang 2010; v2.0) on our
10 dataset of 87 tadpoles. We included 5 sets of duplicate samples to test whether these samples
11 will be reliably grouped together in our analysis. In general, we found significant differences in
12 COLONY results depending on which SNP dataset we used for the analysis. Due to the low
13 number of SNPs called with the LINE109 primers, we focused our COLONY analyses to data
14 obtained using the TE644 primers. We considered the successful resolution of our duplicate
15 samples, grouped together with a probability of 0.9 or greater, as evidence that a SNP data set
16 was reliable for parentage analysis. Generally, reliability was low when using SNPs that were
17 genotyped using Stacks: neither the de novo nor the reference genome mapping approaches
18 reliably identified the 5 duplicates in the dataset with high probability (fig. 5C). When genotyped
19 using the ANGSD pipeline, COLONY identified 4/5 duplicates when SNPs from at least 80
20 individuals were retained, and 5/5 duplicates when 70 individuals were the minimum threshold
21 for calling SNPs, suggesting that the more complete datasets generated with ANGSD—with
22 only 11-17% missing genotypes—provided sufficient information for identifying full-sibs (fig. 5A
23 and C). This method also gave more realistic estimated population sizes than the stacks method
24 (supplementary table S2, Supplementary Material online).

1 Discussion

2 Access to genomic resources and tools holds the potential to transform our
3 understanding of the ecology, evolution, life-history, and conservation of amphibians (e.g.
4 Liedtke et al. 2018; Womack et al. 2019; Schloissnig et al. 2021; Kosch et al. 2022); yet
5 amphibians have lagged behind other groups of animals in available genomic resources
6 (Hotaling et al. 2021). The lack of genomic resources for frogs and toads (Anurans), in
7 particular, is at least in part due to some species possessing large and complex genomes
8 (Rogers et al. 2018; Nowoshilow et al. 2018; Sun et al. 2015); among frogs and toads genome
9 size is highly variable ranging from 0.99 Gb in the plains spadefoot toad, *Spea bombifrons*
10 (GenBank accession GCA_027358695.2) to 10.2 Gb in the mountain yellow-legged frog, *Rana*
11 *muscosa* (GenBank accession GCA_029206835.1). Here we report a 6.8 Gb assembly of the
12 dyeing poison frog, *D. tinctorius*, with a contig N50 of 32.5 Mb and 50% of the assembly being
13 represented by only 56 contigs (fig. 1). The lack of fully assembled chromosomes in our
14 assembly limits our ability to test for chromosomal evolution in *D. tinctorius* compared to other
15 species; however, we found only a single contig that mapped to multiple contigs of other
16 available Hyloidea genomes (fig. 2). Our analysis of synteny between these genomes suggests
17 that the evolution of intrachromosomal rearrangements occurs much more rapidly than the
18 evolution of chromosome structure, despite two-fold differences in genome size. Recent work
19 comparing 8 distantly related anurans supports the idea that the evolution of genome size
20 greatly outpaces chromosomal changes (Bredeson et al. 2024).

21 We used multiple annotation approaches to show that over three quarters of the *D.*
22 *tinctorius* genome consists of repetitive elements. These repetitive elements are more abundant
23 in introns than exons, contributing to the evolution of gene sizes. Below, we discuss how our
24 results advance the understanding of the genome structure evolution in frogs. Additionally, we

1 highlight an application of the *D. tinctorius* genome to facilitate marker development for cost-
2 effective, population-scale multilocus genotyping using MobiSeq (Rey-Iglesia et al. 2019).

3 ***Transposable elements and genome evolution***

4 Amphibians are particularly useful models to provide insights into relationships between
5 environmental or ecological factors and the dynamics of genome size evolution (Liedtke et al.
6 2018). Transposable elements are an interesting component of genome architecture, as their
7 abundance and diversity has the potential to contribute to genetic diversity and subsequent
8 adaptations and divergence within and between species (Schrader and Schmitz 2018; Ding et
9 al. 2016). We find that over three quarters of the *D. tinctorius* genome consists of repetitive
10 elements and that different elements can be found within—or overlapping—both exons and
11 introns (tables 1 and 2). This finding is consistent with the few studies that have generated
12 chromosome-level genome assemblies for other amphibians. Notably, analyses of the Mexican
13 axolotl (*Ambystoma mexicanum*; 32 Gb genome) and Tibetan frog (*Nanorana parkeri*; 2 Gb
14 genome) genomes have reported LTR elements as the most abundant class of repetitive
15 elements (Nowoshilow et al. 2018; Sun et a. 2015). By contrast, in the relatively small
16 amphibian genome of the Western clawed frog (*Xenopus tropicalis*; 1.5 Gb) LTR elements are
17 less abundant, while DNA transposons are more abundant (Hellsten et al. 2010). The fact that
18 LTR retroelements are abundant in large amphibian genomes has led to the hypothesis that
19 these elements play a particularly important role in the evolution of “genomic gigantism” (Sun et
20 al. 2012). However, the mechanism underlying the expansion of LTR elements in large
21 amphibian genomes, or the phenotypic consequences of their proliferation, remain to be tested.
22 Our analyses of LTR abundances, size, and insertion times corroborate past findings
23 (Nowoshilow et al. 2018, Sun et al. 2015) and indicated that LTRs have played a significant role
24 in the genome size evolution of *D. tinctorius* (fig. 4).

1 In addition to amphibians as a whole, transposable elements have likely played an
2 important role in the evolution of genome size among poison frogs (family Dendrobatidae). For
3 example, analyses of a fragmented assembly of *O. pumilio* estimated that this genome consists
4 of over 4.5 Gb of repetitive sequence (Rogers et al. 2018). Consistent with our findings in *D.*
5 *tinctorius*, Rogers et al. (2018) reported *Ty3/DIRS1* LTRs (referred to as “Gypsy-like”) and *Tc1*
6 retroelements as among the most abundant (1.0 Gb and 250 Mb of sequence, respectively).
7 Rogers et al. (2018) also found that *Ty3* and *Tc1* elements are actively expressed in the oocytes
8 of *O. pumilio*. Taken with the evidence of recent insertion events we estimate here (fig. 4), it is
9 likely that many elements are actively evolving and contributing to ongoing genome size
10 evolution in Dendrobatid frogs. Research into the mechanisms governing TE insertion and
11 regulation in anurans will be important to understand the mechanisms underlying genome size
12 evolution in this group.

13 SINEs are a highly abundant class of transposable element in mammals and many other
14 vertebrates, although they are almost entirely absent in amphibian genomes (Chalopin et al.
15 2015; Zuo et al. 2023), including *D. tinctorius*. This difference illustrates how the abundance of
16 different transposable elements varies greatly across the tree of life. Interestingly, the majority of
17 transposable elements we annotated in the *D. tinctorius* genomes could not be classified as
18 known elements using our approach (39.15%). This is likely because reference libraries of
19 described transposable elements used in annotation pipelines lack amphibian-specific
20 elements, and this pattern of abundant “unclassified” elements is common in studies of
21 amphibian genomes (Sotero-Caio et al. 2017; Zuo et al. 2023). Future work that leverages ever-
22 increasing amphibian genomic resources to describe the diversity and structure of the
23 amphibian ‘dark matter’ elements would open doors for comparative analyses of transposable
24 element evolution across taxa and greatly increase our understanding of the evolutionary history
25 of parasitic DNA across the tree of life.

1 An ultimate consequence of TE evolution is their effect on genome size. For example,
2 genome size increases with the abundance of repetitive elements in anurans (Zuo et al. 2023).
3 Our results show that effects on genome size are not restricted to intergenic regions, with TEs
4 being found within both exons and introns (tables 1 and 2). Analyses of the Mexican axolotl,
5 *Ambystoma mexicanum*, have shown a similar pattern, with introns being ten times larger in this
6 species of salamander compared to other vertebrates (Smith et al. 2009). We also found that
7 TEs affect the size of gene regions within the *D. tinctorius* genome (fig. 3). These findings are
8 contrary to a lack of relationship between gene size and genome size reported in a recent
9 comparison of 14 anuran genomes (Zuo et al. 2023). A possible explanation for this discrepancy
10 is that the largest poison frog genome, in Zuo et al. (2023)'s analysis was found to contain
11 significant contamination and has since been retracted (Stuckert et al. 2021, retracted).

12 While it is now clear that TEs comprise a large proportion of many amphibian genomes,
13 and they are not restricted to intergenic regions (Sotero-Caio et al. 2017), there is still a
14 knowledge gap in our understanding of how TEs influence the evolution and adaptability in this
15 group. The fact that many repetitive elements are within or overlap gene features (introns and
16 exons; tables 1 and 2) suggests that their evolution may have important phenotypic
17 consequences. For example, studies in other non-amphibian species have shown that TEs can
18 have diverse effects on gene expression (Lanciano and Cristofari 2020; Rech et al. 2022);
19 however, more work is needed to understand the phenotypic consequences of TEs across
20 ecologically and behaviourally diverse amphibian species. TEs have also been shown to
21 contribute to the evolution of colour phenotypes in a wide range of animals (Galbraith and
22 Hayward 2023), yet links between TEs and colouration in amphibians are largely lacking
23 (Galbraith and Hayward 2023). A possible mechanism linking aspects of genome size
24 evolution—such as the accumulation of TEs in introns—and phenotypic evolution, is the effect
25 that intron size can have on gene expression (Castillo-Davies et al. 2002; Taft et al. 2007).

1 Future work that leverages an increasing number of amphibian genomes with transcriptomic
2 analysis—for example analysing expression of different isoforms across species and
3 developmental stages—could provide a fruitful avenue towards addressing the ‘consequences
4 of genome-size evolution’ knowledge gap.

5 ***Multilocus genotyping using MobiSeq***

6 We used our genome assembly to show that leveraging repetitive elements to design
7 and anchor primers may be a useful way to generate multilocus genotypic data at the population
8 scale. These types of datasets would open exciting possibilities for understanding the adaptive
9 processes and evolutionary dynamics of amphibian populations. The low amount of DNA that is
10 needed for sequencing over larger parts of the genome makes MobiSeq a powerful tool for non-
11 invasive sampling of target species. The target primers can be developed from the reference
12 genome (if available), or from closely related species, as we have shown in our study. However,
13 we recommend using a reference genome whenever possible to design species-specific
14 primers and to map reads and genotype SNPs.

15 Our test of the MobiSeq approach showed that primers can differ considerably in their
16 amplification and sequencing success. The element LINE109 could have been not abundant
17 enough throughout the genome to give enough sequences: the LINE109 element was found
18 109 times in the *D. tinctorius* assembly, while the unknown transposable element TE644 was
19 found 644 times. Therefore, we suggest that future marker development should design primers
20 for markers which have at least 400 to 600 copies in the genome.

21 When genotyping and calling SNPs, de novo assembly with the program stacks gives
22 better results than mapping to the *O. pumilio* genome where the primers were developed from
23 (the *D. tinctorius* assembly was not available when we started the MobiSeq test). Although this
24 is a closely related species, high variability in TE composition could contribute to poorer

1 mapping and genotyping when relying on the *O. pumilio* genome. The de novo approach also
2 gave better results than the mapping to the new *D. tinctorius* reference genome. A limitation of
3 the stacks method we used for genotyping is that this approach resulted in high genotype
4 missingness. Additionally, the stacks program was developed for restriction site digested (RAD)
5 approaches; therefore, we highly recommend using the original ANGSD pipeline, provided by
6 the team that developed MobiSeq when genotyping MobiSeq data (Rey-Iglesias et al. 2019).
7 This method gave the best results considering genotyping and further analysis with COLONY.
8 Specifically, filtering SNPs to those called within a minimum of 70 (out of 92) individuals gave
9 the most reliable relationship estimates in terms of identifying 5/5 of the duplicates in our
10 dataset, as well as giving realistic population-size estimates, given our knowledge of the sample
11 population and the biology of *D. tinctorius*.

12 MobiSeq was developed using mammalian genomes, which have lower levels of
13 transposable elements, and focused on SINEs and LINEs, which are less common in
14 amphibians (Sotero-Caio et al. 2017). Therefore, there are challenges in applying this method to
15 amphibians with large genomes and high transposable element content, such as primer
16 selection and mapping to an existing reference genome.

17 **Conclusions**

18 Transposable elements are a major component of amphibian genomes and they play an
19 important role in genome size evolution. By generating and analysing a long-read assembly of
20 the poison frog, *Dendrobates tinctorius*, we have shown that TE evolution impacts genome size,
21 not only through their insertion in intergenic regions, but also within exons and introns. This
22 “ecology” of TEs provides a possible mechanism that links genome size evolution to phenotypic
23 evolution. We also provide an example use of MobiSeq to generate low-input cost-effective
24 population genetic data. This method could be used to study the evolutionary dynamics of
25 amphibian populations, alongside aiding their conservation. Overall, our study adds to the

1 growing body of knowledge on the evolution of amphibian genomes. We hope the data and
2 analyses we report here will be a valuable resource for future studies of amphibian genetics,
3 evolution, behaviour, and conservation.

4

5 **Materials and Methods**

6 *Sample collection, DNA extraction, and sequencing for whole genome assembly*

7 We generated a reference genome for *Dendrobates tinctorius* from a single male of the
8 “azureus” morph / population. This individual was captive bred by hobbyists in the United
9 Kingdom, kept under licence of the Home Office at Bangor University, and sacrificed by
10 overdose of Tricaine Methanesulfonate followed by immediate pithing and decapitation. DNA
11 was extracted in four parallel extractions performed at the NERC Environmental Omics Facility
12 (NEOF) at the University of Sheffield using the Macherey-Nagel NucleoBond High Molecular
13 Weight DNA kit (see SI for details). DNA was then cleaned and sheared to an average size of
14 19 kb before generating four HiFi libraries using SMRTbell template express kit 2.0 (PacBio).
15 Final libraries were size-selected in the size range of 7-50kb and sequenced across 18 SMRT
16 cells (see SI for details).

17

18 *Genome assembly, quality control, and synteny*

19 We assembled HiFi reads greater than 10kb in length using HiFiASM (Cheng et al.
20 2021, 2022; v0.16.1-r375) run with a bloom filter of 39 bits (option -f f39) and “aggressive”
21 purging of haplotigs (option -l 2). To identify and correct assembly errors, we ran our primary
22 assembly through the Inspector pipeline using inspector.py and inspector-correct.py scripts,
23 respectively (Chen et al. 2021). Inspector also provides an estimate of assembly accuracy,
24 reported as a Quality Value ($QV = -10\log_{10}(\text{base-level errors} / \text{total assembly length})$). After
25 error-correction we identified and removed bacterial or viral contaminants using Kraken2 (Wood
26 et al. 2019; v2.1.2) run using the standard Refseq database containing archaea, bacteria, viral,

1 plasmid, human, and UniVec_Core indexes (downloaded 27-5-2021). We estimated assembly
2 completeness using BUSCO (Simão et al. 2015; v5.2.2) run in genome mode using the
3 tetrapoda_odb10 dataset, which consists of 5,310 single-copy orthologs derived from 38
4 genomes (created 2021-02-19). All other BUSCO options were left as default. We also
5 compared assembly statistics of the *D. tinctorius* genome to the only other Dendrobatid genome
6 currently available on NCBI (*Oophaga pumilio*) and four chromosome-scale assemblies of other
7 Anurans (*Bufo bufo* [NCBI RefSeq ID: GCF_905171765.1], *Rana temporaria* [RefSeq ID:
8 GCF_905171775.1], *Engystomops pustulosus* [GenBank ID: GCA_019512145.1], and *Xenopus*
9 *tropicalis* [RefSeq ID: GCF_000004195.4]).

10

11 *Synteny between D. tinctorius and other Hyloidea genomes*

12 We compared the *D. tinctorius* assembly to chromosome-scale assemblies of *Bufo*
13 *gargarizans* (Asiatic toad; Bufonidae; GenBank accession: [GCA_014858855.1]),
14 *Eleutherodactylus coqui* (Common coquí; Eleutherodactylidae; [GCA_019857665.1]), and
15 *Engystomops pustulosus* (Túngara frog; Leptodactylidae; [GCA_019512145.1]) using orthology-
16 guided synteny map construction in GENESPACE (Lovell et al. 2022; v1.3.1) and BUSCO
17 (Simão et al. 2015; v5.2.2) annotations of the 5,310 gene tetrapoda_odb10 dataset (created
18 2021-02-19) as input. BUSCO annotations for *Xenopus tropicalis* [GCA_000004195.4] were
19 used as outgroup sequences in the initial GENESPACE analysis. Because we were interested
20 in identifying large regions of synteny, we constrained our analysis to scaffolds that have been
21 assigned to chromosomes in the chromosome-scale assemblies, and contigs at least 20 Mb
22 long in the *D. tinctorius* assembly. This approach resulted in the number of annotated single-
23 copy BUSCOs per genome being: 4,482, 3,579, 3,337, 3,440, and 4,939 for *B. gargarizans*, *En.*
24 *pustulosus*, *El. coqui*, *D. tinctorius*, and *X. tropicalis*, respectively. We present summaries of
25 synteny between *D. tinctorius* and the three chromosome-scale assemblies as the proportion of
26 the *D. tinctorius* assembly contained within a synteny block and the average size of synteny

1 blocks between genomes. Syntenic alignments were visualised using GENESPACE's
2 `plot_riparian` function (Lovell et al. 2022; v1.3.1).

3 We also tested whether the size of annotated BUSCO genes was positively correlated
4 with genome size using a Spearman's rank correlation test. We included the five genomes listed
5 above and the chromosome-scale assembly of *Rana temporaria* [GCA_905171775.1]. We
6 focused this analysis on BUSCO annotations rather than *de novo* annotations generated for
7 each genome because we annotated BUSCO genes using the same pipeline in all genomes,
8 and BUSCO genes are chosen based on orthology across different vertebrate genomes,
9 thereby reducing the likelihood that comparisons between paralogous genes affected our
10 results. We note that restricting this analysis to conserved BUSCO genes could result in biases
11 in gene size and evolutionary rate compared to all genes present in these genomes. To test
12 whether the size of BUSCO gene regions correlated with genome size we carried out
13 Spearman's rank correlations on the raw data as well as on phylogenetically independent
14 contrasts (PICs) of both genome and BUSCO gene region sizes. PICs were calculated using
15 the `pic()` function from the `ape` package in R (Paradis and Schliep 2019) and a pruned version of
16 the phylogeny provided by Hime et al. (2021). Finally, we compared the *B. gargarizans*
17 assembly—as it was the closest in size to that of *D. tinctorius*—to *D. tinctorius* using a non-
18 ortholog based nucleotide alignment with `minimap2` (Li 2018, 2021; v2.24-r1122; options:-x
19 `asm20 -l10G -B3 -O4,24 -N10`).

21 *Genome annotation and transposable element diversity*

22 We annotated repetitive elements in our *D. tinctorius* assembly by first using
23 RepeatModeler (Flynn et al. 2020; v2.0.3) with `rmbblast` (v2.11.0+) to identify and generate a
24 species-specific repeat library, followed by RepeatMasker (Smith et al. 2021; v 4.1.2-p1) run
25 using the custom repeat library generated by RepeatModeler. RepeatModeler was run with
26 default settings and RepeatMasker was run with `-xsmall -a -gff` options. Rather than conducting

1 LTR structural analysis with RepeatMasker (-LTRStruct option), we independently annotated
2 Long Terminal Repeat (LTR) retrotransposons using LtrDetector (Valencia and Girgis 2019).
3 We used BLAST and the custom repeat library generated by RepeatModeler to classify LTR
4 retroelements annotated by LtrDetector. To estimate LTR insertion times we extracted left and
5 right LTR regions from the *D. tinctorius* genome in fasta format using bedtools getfasta function
6 (Quinlan and Hall 2010) and the annotated coordinates generated by LtrDetector. We retained
7 annotations where BLAST was able to match greater than 50% of the LTR retroelement with an
8 element contained within the library generated by RepeatModeler (Flynn et al. 2020; v2.0.3).
9 We then aligned left and right elements for each LTR retroelement using MAFFT (Katoh and
10 Standley 2013; v7.490; options: --globalpair --maxiterate 1000) and estimated insertion times
11 from divergence between the left and right LTR, which we estimated from each MAFFT
12 alignment using the dist.dna function in R run assuming Kimura's 2-parameter model (Kimura
13 1980; option: model = "K80"). To convert divergence estimates to time, we assumed a
14 substitution rate of 2.5×10^{-9} substitutions per site per year (Lau et al. 2020).

15 In addition to repeat elements, we annotated protein-coding genes *de novo* using the
16 BRAKER2 pipeline (Brůna et al. 2021; v2.1.6). Before annotation we masked the genome using
17 RepeatModeler and RepeatMasker with default settings. We ran BRAKER2 using evidence
18 from 22 RNAseq libraries from brain, eggs, gut, liver, and skin. RNAseq reads were trimmed
19 using Trimmomatic (Bolger et al. 2014; v. 0.39) to remove adaptor contamination and low-
20 quality bases (options: LEADING:9 TRAILING:9 SLIDINGWINDOW:4:15 MINLEN:80). Trimmed
21 reads were then mapped to the genome using STAR (Dobin et al. 2013; v. 2.7.8a) specifying
22 the twopass Mode Basic option. Mapped RNAseq reads were then used as evidence in the
23 BRAKER2 pipeline. UTR predictions were added to these predictions using the --addUTR=on
24 option. Finally, UTR annotations were trimmed following a gap of over 1000bp. We used these
25 annotations—alongside those generated by RepeatModeler + RepeatMasker—to test whether
26 repetitive elements played a role in the evolution of intron / exon structure in the *D. tinctorius*

1 genome using bedtools intersect (Quinlan and Hall 2010) with options -wa for the full
2 comparison or -f 0.75 -r when testing for 75% reciprocal overlap between annotated gene
3 features and repetitive elements.

4

5 *Tadpole sampling and DNA extraction for relatedness analysis*

6 The aim of this part of the study was to test the MobiSeq protocol as a method for
7 generating a reduced representation library in a non-model organism, the dyeing poison frog (*D.*
8 *tinctorius*). We wanted to use single nucleotide polymorphism (SNP) data to resolve the
9 relatedness between tadpoles in different, small, confined environments such as phytotelmata
10 (small bodies of water in tree holes or palm bracts, for natural history see Rojas and Pašukonis
11 2019).

12 We sampled 87 tadpoles of *D. tinctorius* from 17 phytotelmata in the Nouragues Nature
13 Reserve, French Guiana in 2020, by clipping a small part of the tail tip. Tissue samples were
14 stored in 70% ethanol (EtOH) at -20°C until further processing. To increase reliability and to test
15 the appropriateness of the relatedness analysis, we included 5 samples as duplicates, giving a
16 total sample size of 92. A modified salting-out method was used to extract DNA from small parts
17 of the tail clips (supplementary methods, section 1, Supplementary Material online). The
18 extracted DNA was eluted in 100µl TE buffer, DNA concentration measured with a NanoDrop™
19 spectrophotometer (Thermo Fisher) and stored at -20°C until further processing.

20 *MobiSeq primer design*

21 We designed specific primers for repeated elements as MobiSeq uses transposable
22 elements (TEs) in the DNA for target enrichment PCRs. No published reference genome for *D.*
23 *tinctorius* was available when we started to work on the MobiSeq approach. To find highly
24 repetitive TEs, we used the genome of a closely related species, the strawberry poison frog

1 (*Oophaga pumilio*), which was published in 2018 (Rogers et al.; NCBI GenBank:
2 [GCA_009801035.1]).

3 We used the free software RepeatMasker (Smith et al. 2021; v 4.1.2) to mask repeated
4 elements and extracted those regions with Samtools (Danecek et al. 2021) to get the 22 last bp
5 of transposable elements as a reversed complement and the number of occurrences. The list
6 was cross checked and annotated with existing libraries of transposable element families
7 provided by Dfam (Storer et al. 2021; v3.2). Based on this list we choose elements with an
8 average abundance of 400-600 times in the genome, as elements with higher abundances
9 might be clustered and not evenly spread over the genome. LINES and SINEs were generally
10 less abundant so that we included only one primer for a lower abundance LINE. We initially
11 chose six possible reverse transposable element primers and tested amplification and
12 multiplexing with *D. tinctorius* DNA. We decided on the following two primers after running a
13 target enrichment PCR, based on amplification success and minimisation of primer dimers
14 (AdapterTEsequence):

15 D_tinct_Line_109

16 (5'**GTGACTGGAGTTCAGACGTGTGCTCTTCCGATCTCCTATGTTACTATGTTACTATGT**'3)

17 D_tinct_TE_644

18 (5'**GTGACTGGAGTTCAGACGTGTGCTCTTCCGATCTTACTTTTGCCACCACTGTA**'3).

19
20 In the first step of the MobiSeq protocol, the *D. tinctorius* DNA was digested using
21 fragmentase. To ensure an even digestion across samples, the DNA was diluted to 10ng/μl. We
22 tested the correct incubation time for the fragmentase enzyme beforehand and decided on 20
23 min at 37°C for each sample based on gel images (supplementary methods, section 4.1,
24 Supplementary Material online). After fragmentase treatment, a Sera-Mag Speed beads clean-

1 up was used to remove the enzyme mix and buffers (supplementary methods, section 4.2,
2 Supplementary Material online). Clean-up was followed by end-repair of DNA fragments to
3 create blunt ended fragments using T4 polymerase. After incubation, samples were cooled
4 down to 10°C and immediately used for the next step (supplementary methods, section 4.3,
5 Supplementary Material online). For adapter ligation, double stranded modified P5 adapters
6 (Meyer and Kirchner 2010) were added to the end-repaired DNA fragments via a T4 DNA
7 ligase. The modification of the adapter results in single-stranded adapters, which allow the use
8 of universal adapter primers in the next target enrichment PCR step. The adapter ligation
9 master mix (4µl per sample) is added to the end-repaired samples. The samples were cooled
10 down to 10°C after the incubation (supplementary methods, section 4.4, Supplementary
11 Material online). The resulting product was cleaned again, using the same Sera-Mag Speed
12 beads as after DNA fragmentation (supplementary methods, section 4.2, Supplementary
13 Material online) and eluted in 25µl AE buffer and stored at 4°C. We used the primers TE644 and
14 Line109 as described above to enrich the fragments containing transposable elements in a
15 multiplex approach (supplementary methods, section 4.5, Supplementary Material online). The
16 remaining 15µl of PCR were cleaned using the Sera-Mag Speed beads and eluted in 20µl AE
17 buffer. In the final step of the sequencing library preparation, a second PCR using Illumina
18 indexed forward and reverse primers was used followed by agarose gel electrophoresis
19 (supplementary methods, section 4.6, Supplementary Material online). The uniquely tagged
20 samples were mixed, based on the brightness of the smear on the gel. The whole pool of
21 samples was loaded on another gel to cut out fragments of the lengths between 200 and 500bp.
22 The innuPREP DOUBLEpure kit (Analytik Jena) was used to extract the DNA from the gel
23 fragment, following the kit's protocol. The final library pool was sent to Novogene UK for
24 sequencing in 150bp paired end mode on a Novaseq instrument (Illumina).

25

1 *Bioinformatics for variant calling*

2 After downloading the demultiplexed fastq files from Novogene we visually checked the
3 reads for quality with qiime2 (Bolyen et al. 2019; v2021.4) using the demux plugin (function
4 summarize). As each fastq file contains Line109 and TE644 reads, we used cutadapt (Martin
5 2011; v4.1) to separate each file into two, based on the primer sequence. After separating the
6 files, we used cutadapt to remove all versions of adapters and primers from the forward and
7 reverse reads. Reverse-complement versions were used to remove primer fragments that can
8 be present on the 5' end of reads from fragments that are shorter than 150bp. After trimming,
9 the sequences were filtered for optical duplicates and remains of bacteriophage PhiX
10 contamination using the shell scripts clumpify.sh and bbduke.sh from BBTools (Bushnell 2014) .

11 For reduced representation sequencing approaches such as MobiSeq to be broadly
12 applicable to non-model species that lack genomic resources, it is important to know how reliant
13 markers are on the availability of genomic resources of closely related taxa. As such, we took
14 three approaches to calling SNPs in our dataset: (1) de novo assembly of MobiSeq fragments,
15 (2) mapping reads to the *O. pumilio* reference, and (3) mapping to the *D. tinctorius* genome we
16 generated as part of this study. The de novo assembly was conducted twice using
17 `denovo_map.pl` from the stacks pipeline (Catchen et al. 2011, 2013; v2.64), using the default
18 parameters. Therefore, all sequences were truncated to 50bp and 100bp, respectively for both
19 primers using `process_radtags`. The reason for the double truncation is that the truncation
20 process discards sequences shorter than the chosen length (50 or 100bp). The 50bp approach
21 therefore retains more but shorter sequences than the 100bp approach. We wanted to compare
22 different truncation lengths to validate the robustness of de novo assembly. The reference
23 genome mapping was done with `ref_map.pl` from the stacks pipeline (Catchen et al. 2011, 2013;
24 v2.64), using the default parameters. Before mapping, the reference genomes were indexed
25 with the BWA-mem2 index function (Vasimuddin et al. 2019; v2.2.1) and subsequently

1 sequences aligned with the mem function. The output was saved as a vcf file, with the following
2 parameters: populations: -p 1 -r 0 --write-random-snp --max-obs-het 0.5 --ordered-export --vcf.

3 The soft filtering of SNP data was done in R (R Core Team 2022; v4.2.2) with
4 Bioconductor (Morgan 2022; v3.16; BiocManager 1.30.19) and the filter_rad function from the
5 radiator package (Gosselin 2020; v1.2.8). To implement the radiator package from github we
6 used the package devtools (Wickham et al. 2022). All thresholds that we used for filtering can
7 be found in supplementary methods (section 5, Supplementary Material online). We expected a
8 high degree of missing data due to the Mobiseq approach, therefore the filter for maximum
9 missingness was quite high (0.9).

10 The strong filtering of SNP data was done with vcftools (Danecek et al. 2011; v0.1.16)
11 using the following parameters: --max-missing 0.5 --mac 2 --minDP 3.

12 Additionally, we used the ANGSD pipeline from Rey-Iglesia et al. 2019
13 (<https://github.com/shyamsg/MobiSeq/blob/master/code/pipeline.sh>), accounting for the fact that
14 stacks was developed for restriction enzyme based RAD sequencing and might bias our output.
15 We used the BWA-mem2 indexed reference genome of *D. tinctorius* to map our sequences,
16 bedtools (Quinlan and Hall 2010; v2.26.0) to merge reads and kept only sites that are present in
17 90% of the cases. We called variants using ANGSD (Korneliussen et al. 2014; v0.940) with
18 minimum quality of 30, min mapping quality of 30, filter for SNPs with a p-value of 1e-6, major
19 and minor allele were inferred directly from likelihoods, minor allele frequency was estimated
20 (fixed major unknown minor), genotypes and SNPs called. SNPs needed to be present in either
21 a minimum of 80 or 70 individuals. We randomly called one SNP per contig to avoid linkage and
22 created 5 vcf files for the 80 and 70 individuals dataset respectively.

23

24

1 *Relatedness analysis*

2 The relatedness analysis was conducted with COLONY (Jones and Wang 2010; v2.0).
3 We used the filtered SNPs of the TE644 primer only, derived from our 4 approaches (de novo
4 stacks, *O. pumilio* reference genome or our new *D. tinctorius* genome) and two methods of
5 calling SNPs (stacks vs. ANGSD). The write_colony function in the radiator package was used
6 to write a colony input function. The input parameters were as follows; the mating system of
7 males and females was set to polygamous, no inbreeding and no update of allele frequencies,
8 the length of the run was set to 2 with full likelihood analysis and high precision. We ran an
9 analysis with different random seeds to increase reliability of the clustering (1234 and 1789).

10

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18

19 **Data availability**

20 The *D. tinctorius* genome assembly and BRAKER2 annotations are available on the Open
21 Science Framework project page:
22 https://osf.io/tg24j/?view_only=13be51ec97184d888a12d4ba26dfc90f and the genome is being
23 processed to be made available on NCBI: BioProject: PRJNA977854; BioSample:

1 SAMN38117364. Illumina reads generated from MobiSeq libraries are publicly available on
2 NCBI: BioProject: PRJNA1035148.

3

4 **Authors' contributions**

5 AAC, SS and BR conceived and designed the project. RM, CAF and AAC collected samples.
6 CD, FH, LSK, MH extracted DNA and prepared the libraries. SS, LAO, AF, AAC provided
7 resources (data, computational). DP, FH, CD, AAC performed bioinformatic analysis of the data.
8 CD, AAC, FH, BR and SS discussed and interpreted the results, AAC, CD, FH and LAO wrote
9 the first draft. AAC and BR provided funding. All authors read, reviewed and approved the final
10 manuscript.

11

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- 18

1 **Figure and Table Legends**

2 **Figure 1.** *Dendrobates tinctorius* assembly completeness. **(A)** Cumulative assembly size in
3 relation to size-ordered scaffolds for *D. tinctorius* and other published anuran genomes. **(B)**
4 BUSCO “tetrapoda” gene set completeness for the same genomes summarised in **(A)**.

5 **Figure 2. A)** Riparian synteny plots generated using BUSCO gene regions as anchors showing
6 broad patterns of synteny and rearrangements between *D. tinctorius* scaffolds and three publicly
7 available chromosome-scale frog assemblies. **B)** Example *B. gargarizans* chromosome showing
8 both synteny and structural evolution with *D. tinctorius* scaffolds. Inversions in panel B are
9 shown in salmon pink, collinear regions in orange. **C)** Insertion of regions on *D. tinctorius*
10 scaffold 127 (highlighted in red) into chromosomes 1 and 4 of the *E. coqui* assembly. Phylogeny
11 shown in panel A) is pruned from the RAxML “L1-L379.concatenated” tree reported in Hime et
12 al. (2021).

13 **Figure 3.** The relationship between genome size and the average size of annotated BUSCO
14 gene regions in *D. tinctorius* compared with five publicly available chromosome-scale frog
15 assemblies. A) Phylogenetic relationships between the species being compared (pruned
16 RAxML “L1-L379.concatenated” tree from Hime et al. 2021). B) The relationship between
17 genome size (in bp) and the mean size of BUSCO gene regions. C) The relationship between
18 genome size and mean size of BUSCO gene regions, corrected for phylogeny using
19 phylogenetically independent contrasts (PICs) with the phylogeny depicted in A.

20 **Figure 4.** Long terminal repeat (LTR) retroelement lengths **(A)** and insertion times **(B)**. LTRs
21 were annotated with LtrDetector and insertion times were estimated from divergence between
22 the left and right LTR regions of each element independently assuming a substitution rate of 2.5
23 $\times 10^{-9}$ substitutions per site per year (Lau et al. 2020). In **(A)** the total number of elements
24 classified to each type is given above each boxplot.

25

1 **Figure 5.** Summary of **(A)** genotype missingness in %, **(B)** number of SNPs (log scale to
 2 account for the large differences) and **(C)** percentage of duplicates found in the dataset using
 3 sequences derived from MobiSeq with the TE644 primer. Shown are the two pipeline
 4 approaches, stacks on the left (de novo assembly and mapping to reference genomes of
 5 *Oophaga pumilio* or *Dendrobates tinctorius*) with no filter SNPs, soft filter (filter_rad) and strong
 6 filter (vcftools); and ANGSD on the right (calling of SNPs present in 70 or 80 individuals). For
 7 details see Methods.

8
 9 **Table 1.** Counts of repeat elements annotated using RepeatModeler + RepeatMasker that
 10 overlap with coding DNA sequence (CDS) and introns annotated in the *D. tinctorius* assembly.

Region of overlap	All elements	LTRs	LINEs	<i>Tc1</i> transposons	<i>hAT</i> transposons
CDS	115,330	3,500	1,826	10,227	507
Introns	528,441	42,844	27,152	45,806	2,907
Total N elements	11,331,718	1,048,437	702,556	897,264	265,361

11
 12
 13
 14 **Table 2.** Counts of repetitive elements that overlap with CDS and introns (see Table 1) that
 15 show greater than 75% reciprocal overlap with those features.

Region of overlap	All elements	LTRs	LINEs	<i>Tc1</i> transposons	<i>hAT</i> transposons
CDS	4,872	13	0	11	2
Introns	5,530	914	931	1,193	23

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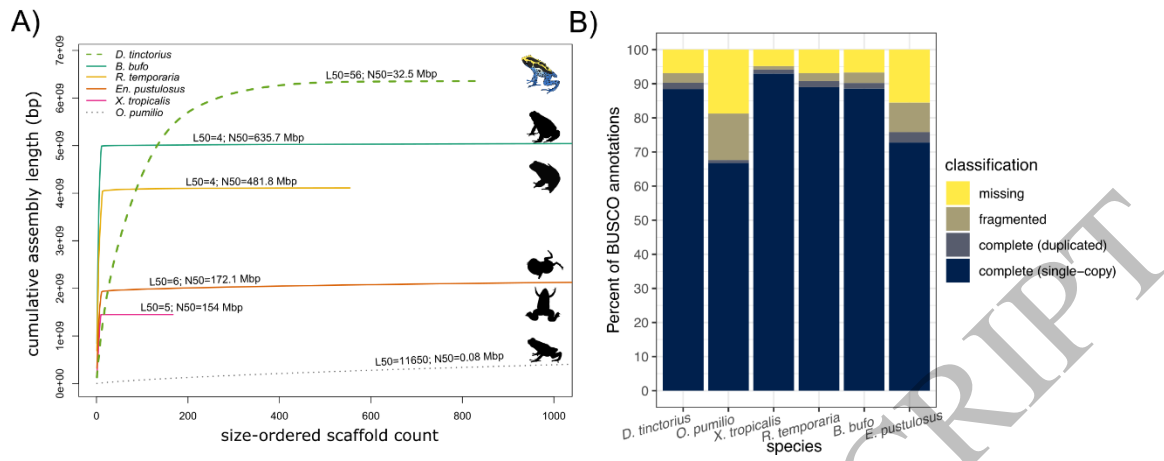


Figure 1
148x59 mm (x DPI)

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ACCEPTED MANUSCRIPT

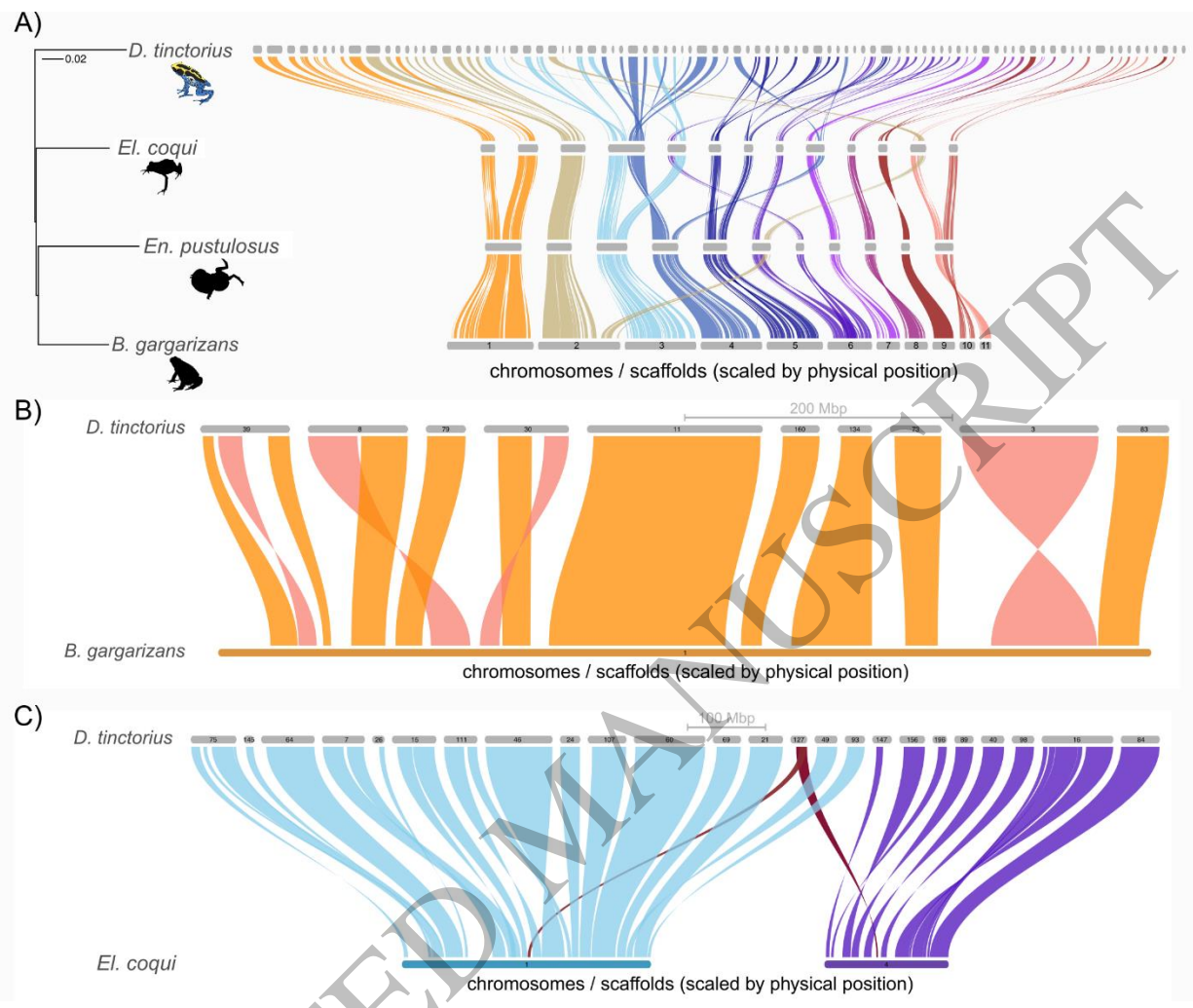


Figure 2
165x134 mm (x DPI)

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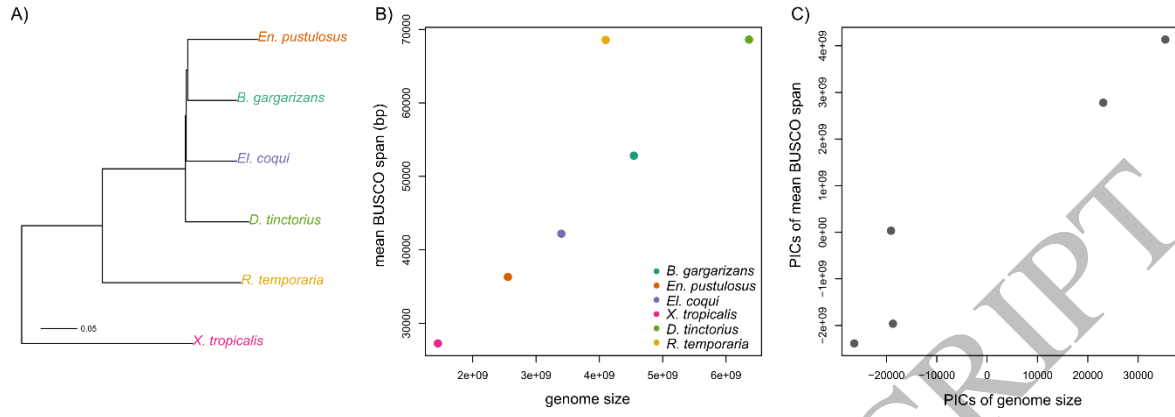
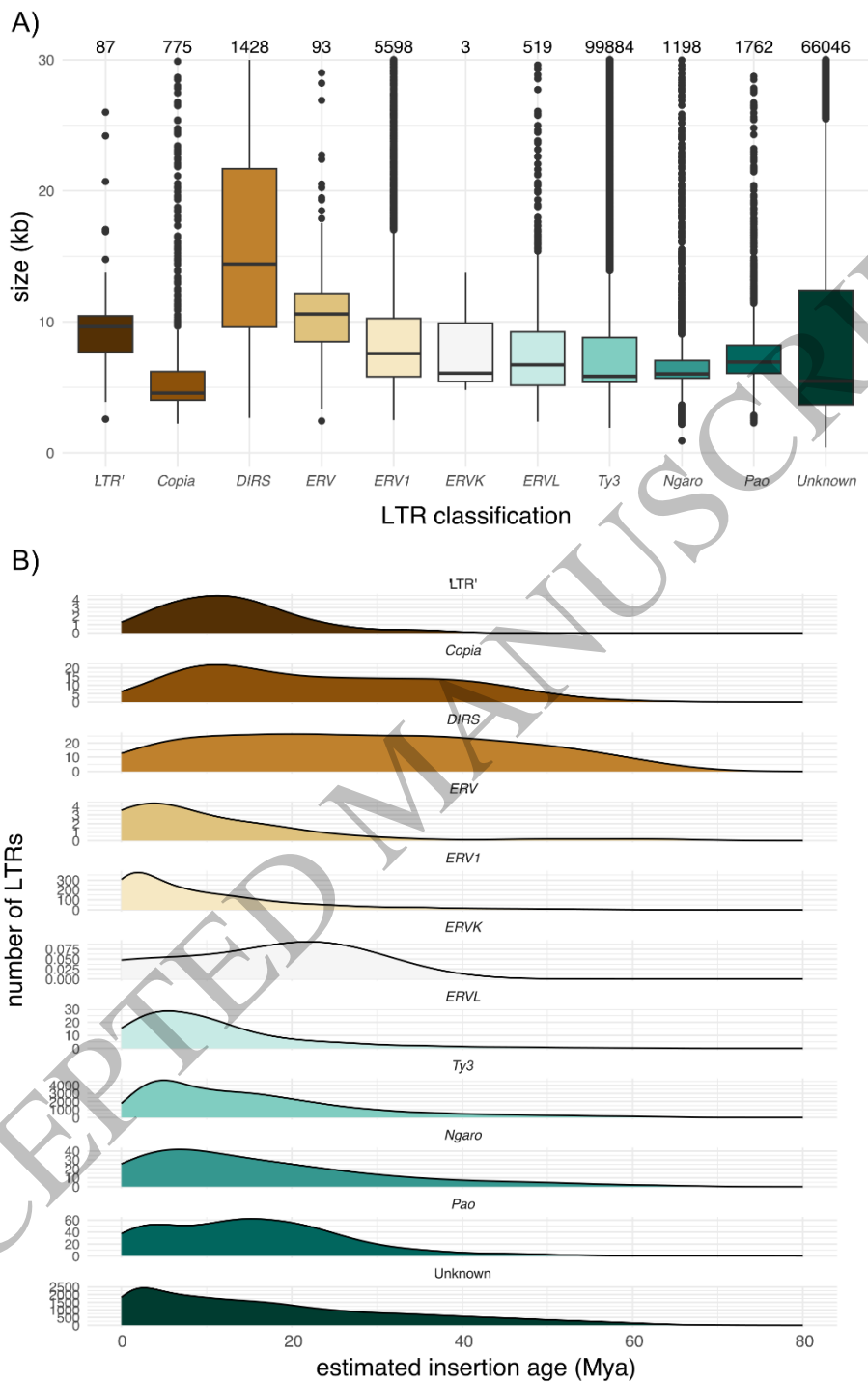


Figure 3
 165x59 mm (x DPI)

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Figure 4
144x229 mm (x DPI)

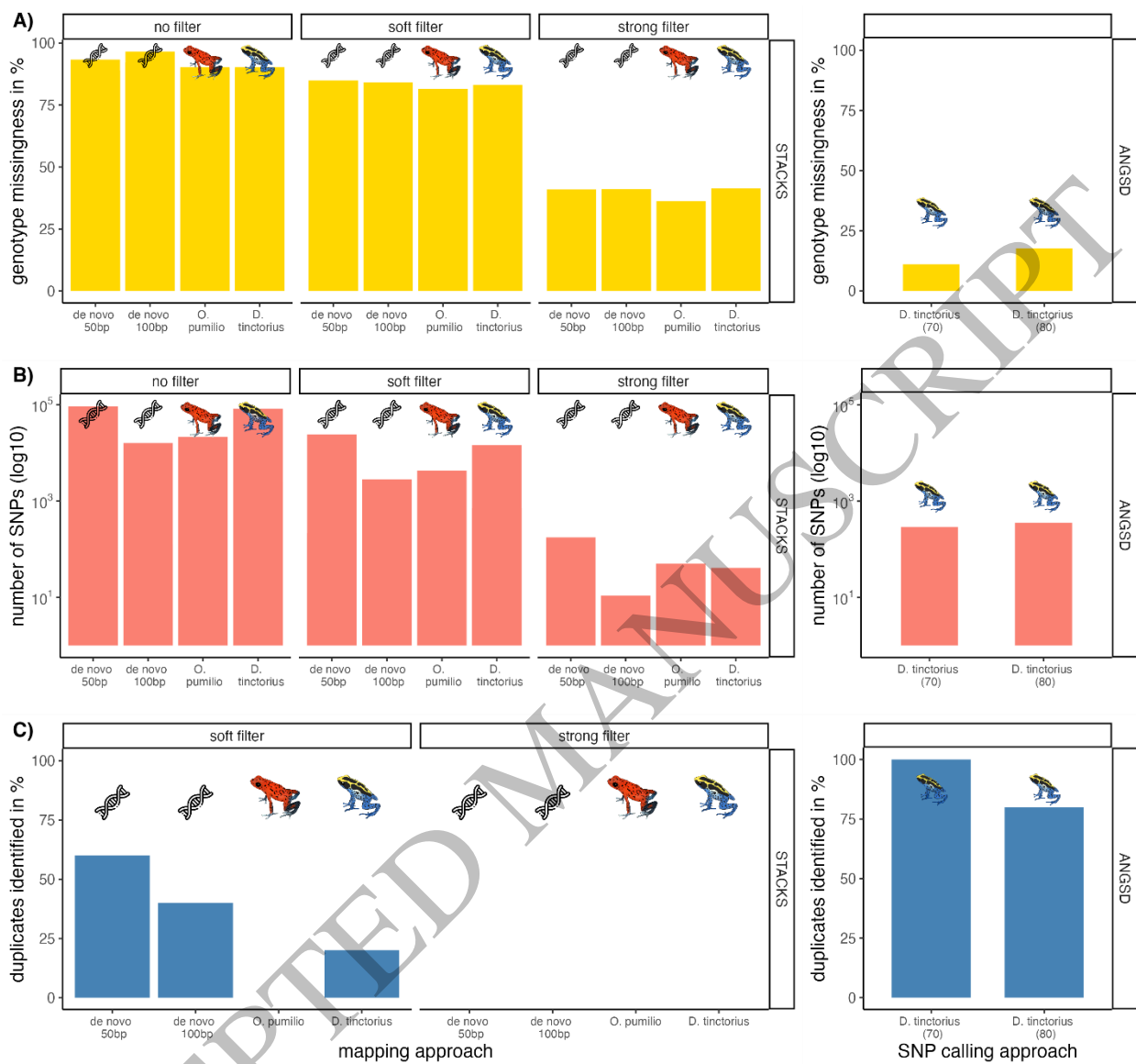


Figure 5
165x153 mm (x DPI)

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