Markers for genetic change

Giovanni Forcina¹, ², Miguel Camacho-Sanchez¹, ², Fred Y.Y. Tuh³, Sacramento Moreno⁴, Jennifer A. Leonard¹, ²

¹ Conservation and Evolutionary Genetics Group, Estación Biológica de Doñana (EBD-CSIC), 41092 Seville, Spain
² CIBIO/InBIO, Centro de Investigación em Biodiversidade e Recursos Genéticos, Universidade do Porto, Campus Agrário de Vairão, 4485-661 Vairão, Portugal
³ Sabah Parks, Lot 45 & 46 KK Times Square Coastal Highway, 88100 Kota Kinabalu, Sabah, Malaysia
⁴ Department of Ethology and Biodiversity Conservation, Estación Biológica de Doñana (EBD-CSIC), 41092 Seville, Spain

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A B S T R A C T

Background and aims: Wildlife conservation has focused primarily on species for the last decades. Recently, popular perception and laws have begun to recognize the central importance of genetic diversity in the conservation of biodiversity. How to incorporate genetic diversity in ongoing monitoring and management of wildlife is still an open question.

Methods: We tested a panel of multiplexed, high-throughput sequenced introns in the small mammal communities of two UNESCO World Heritage Sites on different continents to assess their viability for large-scale monitoring of genetic variability in a spectrum of diverse species. To enhance applicability across other systems, the bioinformatic pipeline for primer design was outlined.

Results: The number of loci amplified and amplification evenness decreased as phylogenetic distance increased from the reference taxa, yet several loci were still variable across multiple mammal orders.

Conclusions: Genetic variability found is informative for population genetic analyses and for addressing phyogeographic and phylogenetic questions, illustrated by small mammal examples here.

1. Introduction

Genetic diversity is important for the long-term survival of species, and thus is of high conservation concern (Frankham, 2002; Spielman et al., 2004; O’Grady et al., 2006). This is even more true in an epoch of unprecedented human-caused disturbance and environmental change, where the preservation of adaptive potential and functional diversity are key for populations to respond to changing environments (Hoelzel et al., 2019; Razgour et al., 2019). Nevertheless, laws and policy have long focused primarily on more visible elements of biodiversity, such as species or populations, with genetics being most often neglected (Laikre, 2010). Genetic diversity is increasingly taking an important role in the international political agenda. It has been included in well-known international mandates such as the Habitats Directive 92/43/EEC and the Birds Directive 79/409/EEC, which apply across the European Union (EU). More recently, the United Nations (UN) has declared 2020 the Post 2020 Biodiversity Framework (OECD) becoming an obligation for the EU member states as well as other signatories. The EU has the stated objective to slow the loss of biodiversity and ecosystem services by 2020, and restore, as possible, the environment in order for its member states to contribute to the maintenance of global biodiversity. One of the official criteria to fulfill this task is the maintenance of genetic diversity.

However, there is no consensus on how this diversity should be measured. In the context of research, a number of studies have looked at changes in genetic diversity through time, especially before and after certain events, often periods of overexploitation (e.g., Baker et al., 2000; Pinsky and Palumbi, 2013; Sanchez-Donoso et al., 2014) or pollution exposure (for a review, see van Straalen and Timmermans, 2002). These studies have been based primarily on population level analyses of mitochondrial DNA markers such as the control region or cytochrome b (cyt b), or autosomal markers such as microsatellite loci. Mitochondrial DNA (mtDNA) markers are generally variable within populations, easy to amplify with standard protocols, applicable across a wide taxonomic range, yield sequences comparable between studies, and there is a lot of...
comparative data available (e.g., Kohlmüller et al., 2012, 2016). They are also easier than nuclear markers to type from degraded material, such as feces and old bones, because of their higher per-cell copy number (Templeton et al., 2013). The downside of using mtDNA to track changes through time is that it is a single marker with necessarily limited power, and also it only informs about the history of the female lineage (Heled and Drummond, 2009). These constraints, among others, mean that mtDNA diversity does not necessarily correlate with nuclear diversity (Teske et al., 2018) and the historical demography of the species being studied (Bazin et al., 2006; Nabholz et al., 2008).

Microsatellites are other popular markers for population level studies. Even though these loci are more difficult to isolate and optimize, the system generally yields data from multiple, independently inherited genetic markers useful for population level metrics (Pinsky and Palumbi, 2013). Generally very abundant in the genomes of most species, now with next generation sequencing (NGS) methods they are easier to isolate than before (Yang et al., 2014). Their loci are often very variable within populations, and so offer higher power to identify changes in genetic diversity through time (Hausl and Payseur, 2011; Putman and Carbone, 2014). The primary drawback of microsatellite studies is that the results are generally not comparable between laboratories (Moran et al., 2006), or even between different projects in the same laboratory. Moreover, ascertainment bias from the marker discovery approaches (Dufresne et al., 2014; de Groot et al., 2016) and the low number of loci in a typical microsatellite dataset (Fischer et al., 2017) can limit their power for reliable estimates of genetic diversity and population structure (Camacho-Sanchez et al., 2020) as well as the many assumptions associated with their analysis and interpretation (Putman and Carbone, 2014). Overall, this makes it very difficult to build on previous batches of data, which is fundamental to ongoing genetic monitoring. Ideally there would be a panel of markers that could be applied to a variety of non-model organisms in a way such that genotypes produced at different times and/or in different laboratories, and/or with different methods would be comparable.

Another molecular tool used in some studies and partly meeting these criteria is intron sequences. These have some of the beneficial characteristics of mtDNA, such as being sequence-based markers that can be easily shared and compared between projects, and for the applicability of methods across different species as exemplified by Transcriptome Ortholog Alignment Sequence Tools (TOASTs: Jiang et al., 1998; Weisel et al., 2020) and Comparative Anchor Tagged Sequences (CATs: Lyons et al., 1997). Likewise, introns share some of the advantages of microsatellites in that they are biparentally inherited, numerous, and evenly distributed throughout the genome. Similar to other neutral nuclear markers, introns may represent a good proxy to assess functional adaptive potential or functional diversity (Vilas et al., 2015). This combination of features offers great potential for monitoring genetic diversity in wildlife. Although generally much less variable than nuclear microsatellites or mitochondrial control region sequences (but more than respective exons: Igea et al., 2010), introns are abundant, so their power to evaluate intra-specific genetic diversity can be increased by genotyping more loci. Multilocus PCR panels can be easily established profiting from the generally conserved flanking exonic regions, which enables the use of the same primers across different taxa. However, amplification and sequencing of multiple loci can quickly become expensive and logistically complicated. Perhaps for this reason, panels of introns have mostly been used in systematic studies that look at the relationships between different and sampling individuals (e.g., Hafner et al., 2012; Igea et al., 2013). With recent methodological advances such as molecular indexing and next generation sequencing, some of these problems can be overcome (Meyer and Kircher, 2010). The large number of publicly available genomes has enabled the development of panels of possibly suitable loci (Igea et al., 2010; Rodriguez-Prieto et al., 2014). Recently, panels of introns have been used in population level studies (Pons et al., 2010; Tollis et al., 2012; Cordero et al., 2014; Kuchta et al., 2016; Camacho-Sanchez et al., 2018).

Small mammals (defined as terrestrial mammals weighing 5 kg or less: Merritt, 2010) play a pivotal role in their ecosystems as they account for a considerable biomass proportion and include both primary and secondary consumers, seed dispersers, and predators (Ostfeld et al., 1996; Ashwanden et al., 2007). By being easy to collect and handle, and mostly habitat specialists, small mammals represent useful bioindicators and are good candidates for comparing geographically distant study areas (Talmage and Walton, 1991; Smith et al., 2002). Moreover, with their short lifespan and rapid life-history responses to environmental changes, taxa falling into this category are particularly suitable for understanding environmental effects on animal population dynamics (Rowe and Terry, 2014; Hope et al., 2017) and for singling out specific anthropogenic drivers (Byrom et al., 2015). Since in situ species conservation is primarily achieved by protected areas, the preservation of genetic diversity is also likely to be dependent on them. Documenting the genetic diversity in these sites will therefore be of key concern in implementing the new mandate to reserve genetic diversity.

In this study we assess a panel of nuclear markers for utility in characterizing genetic diversity in the small mammal communities (Table 1) of two UNESCO World Heritage Site national parks located at opposite ends of Eurasia (Figure 1). This panel of markers (Table 2, Table S1) is amplified in a single multiplexed PCR reaction, decreasing lab time and reagent cost. The indexed PCR products are pooled and sequenced using high-throughput sequencing platforms. We test the applicability of this panel in different taxa with increasing phylogenetic distance from the species for which it was developed, the brown rat (Rattus norvegicus), and evaluate the power for population genetics purposes. We also test the comparability of data generated through different library preparation protocols and on different NGS platforms. In order to facilitate the applicability of this method in other systems, the bioinformatics pipeline for primer development is described in detail. Finally, the phylogenetic utility of the panel is evaluated by a comparison with phylogenies based on mtDNA.

2. Results

2.1. Amplification and polymorphism across taxa

The number of loci yielding reads varied from 40 (the entire panel: Table 1, Table 3, Table S2, Appendix S1) in Rattus (and other representatives of the subfamily Murinae) to roughly half among Eulipotyphla (here represented by shrews and gymnures) and even less in other distantly related taxa from other orders such as Scandentia (tree-shrews: only 17). Two loci (Dhc2r4, Smo) amplified across all orders tested. Polymorphisms were detected in all the assembled loci, with the highest values in the subfamily Murinae. In terms of species, the maximum value was recorded in Leopoldamys sabanus (34 loci, 85%), while the number of alleles per species and locus ranged from one to eleven (Table 3). The lowest number of segregating sites (S) were recorded in non-rodent taxa, although the minimum value (0) was observed in a squirrel, Sundasciurus jentinki, and a murid, Chiropodomys pusillus, other than in a shrew, Suncus etruscus (Table 1, Table 3, Table S2 Appendix S1). Five loci (Dhc2r4, P2rx1, Smo, Usp20, and Wls) were the most variable, with Dhc2r4 and Smo yielding in total more than 100 and 80 alleles across 16 and 13 species, respectively. Genetic diversity indexes revealed a higher variability among murids, with values generally exceeding those inferred at the reference nuclear loci, the growth hormone receptor exon 10 (Ghr) and the retinol binding protein 3 exon 1 (Rbp3) (Appendix S1). Some of the loci that did not work in the amplicon libraries yielded sequences from shotgun libraries (i.e., Wls, Rtras). Variation in coverage among loci in the same PCR and sequencing platforms did not seem to be driven by taxonomic affiliation (Figure 2). We discarded 85 out of a total of 97 (87.6%) alleles obtained exclusively with GS Junior 454 due to sequence variation in homopolymeric regions which might be attributable to sequencing errors. Sequences at issue were not employed for tree
reconstruction or computation of standard genetic diversity indexes, but were scored as successful amplifications.

2.2. Phylogenetic reconstructions

The phylogenies based on nuclear markers (Figure 3) were consistent with recent work combining multiple exonic and mitochondrial markers (Rowe et al., 2019) for the subset of taxa included in both studies. However, the mtDNA phylogeny (Figure 4A and B) showed internal discordances in the clade from the Dacnomys division (Rowe et al., 2019) for the subset of taxa included in both studies. With recent work combining multiple exonic and mitochondrial markers, potential for widespread application. There is a growing body of literature showing efforts to take stock of the plethora of molecular approaches that are being applied for biodiversity monitoring, especially among mammals (e.g., Larsen and Matoq, 2019; Forcina and Leonard, 2020). In this study, we tested a multiplexed panel of intron loci in spatially and taxonomically different small mammal communities of major ecological relevance to develop a tool for genetic monitoring which could be applied to many different taxa with limited effort.

3. Discussion

Anthropogenic activities are wiping out biological diversity at an alarming rate, raising serious concerns for ecosystem functioning and the delivery of associated services in an epoch of increasing environmental stochasticity (e.g., Kremer et al., 2012; Valiente-Banuet et al., 2015; Seddon et al., 2016). Now that the broader society has started recognizing the importance of preserving genetic diversity to counteract this trend, a vital question being tackled is how to measure and compare it. Major advances in genomic technologies have heralded a new era, enabling researchers to produce massive amounts of genome-wide data on multiple individuals. Although budget, computational, and other logistical constraints still impairs the large-scale adoption of whole genome approaches (Fuentes-Pardo and Ruzzante, 2017), using genomic tools to genotype targeted loci with fast, cheap and easy protocols has the potential for widespread application. There is a growing body of literature showing efforts to take stock of the plethora of molecular approaches that are being applied for biodiversity monitoring, especially among mammals (e.g., Larsen and Matoq, 2019; Forcina and Leonard, 2020). In this study, we tested a multiplexed panel of intron loci in spatially and taxonomically different small mammal communities of major ecological relevance to develop a tool for genetic monitoring which could be applied to many different taxa with limited effort.

Primers designed with the pipeline used in this study (Figure S1) and the laboratory rat genome largely worked as a single multiplex. None of the primer pairs appeared to amplify other loci or duplicated regions, suggesting that these loci will be of value across mammals, and could be worth testing in other vertebrate groups as initially suggested by Igea et al. (2010). We successfully amplified all the loci tested in at least some species, and obtained sequences across all the surveyed taxa from at least a few loci. Amplification success decreased with phylogenetic distance from the model species used for designing primers, R. norvegicus. The panel worked well across the entire family Muridae, whose most recent common ancestor is estimated at 95 MYA (Dos Reis et al., 2015). Two loci (Dhcr24, Smo) even worked across all the tested orders, whose most recent common ancestor is estimated at 80–95 MYA (Dos Reis et al., 2012; Foley et al., 2016; Upham et al., 2019). Given the higher rate of mutation in rodents than other groups of mammals (e.g., Cooper et al., 2004), primers may be better conserved in other orders. These results show that the genome of the target species is not necessary for primer development, as primers in evolutionary related species worked consistently well. Hence, when setting up a new project, it is recommendable to...
check the fast-increasing list of newly available genomes and select the closest one to the study species for primer design. Alternatively, the primer design step could be partially avoided and the loci genotyped through sequencing of enriched shotgun libraries. This, however, will require more complex laboratory protocols, deeper sequencing (which is relatively cheap), and more importantly, will be bioinformatically more complicated to analyze.

The intron sequences contained a good amount of information as compared to those of commonly targeted nuclear loci such as Rbp3 or Ghr (Appendix S1), with fairly high genetic diversity indices in Muridae and, to a lesser extent, across other families of the order Rodentia (Table 3, Appendix S1). These results indicate that this panel may prove useful for addressing population level genetic questions (Camacho-Sánchez et al., 2018). It will still be necessary to establish appropriate species or population specific baselines from which power estimates can be made. In many cases populations of wildlife have been subject to historic changes in population size and connectivity, so it may be most appropriate to establish these baselines with historic data.

This panel of markers was also sufficiently informative to construct a high confidence evolutionary phylogeny of the Rattini in our dataset. In comparison to the whole mitochondrial genome phylogeny with the same set of taxa, we show that the nuclear loci were better able to resolve the difficult nodes in the clade containing Leopoldamys, Lenothrix, and Niviventer.

Overall, this protocol should work in any animal taxa. Additionally, these particular loci should work at least across most mammals, and these primers in most Rattini and many rodents. Nevertheless, it is best to design taxon-specific primers for the target species for each particular study to ensure the highest amplification success, and this will not impair the comparability across studies. Indeed, the primers tested here worked best in the target group, and success dropped off substantially in other orders. To apply this panel in other groups of mammals, it would be advisable to design new, taxon specific primers, if possible. The other way to get around this- targeting the loci trough enrichment instead of PCR - required more sequencing and is bioinformatically more complex. Also, power calculations should be made for each species and question in order to ensure that the number of loci sequenced is sufficient to achieve the level of taxonomic resolution needed. Further, in this test we have used high quality samples. If low quality samples such as feces are used, PCR replicates may be necessary.

In conclusion, here we propose an efficient and effective molecular tool for the genetic screening of small mammal communities. We describe a multiplexing strategy which, in combination with the bar-coding of multiple individuals, represents a cost- and time-efficient as well as easy-to-implement procedure for use by academics, governmental agencies, and wildlife managers. This has a major advantage over mitochondrial DNA alone because it relies on many independent markers. Unlike microsatellites, the sequence-based markers proposed here generate data which can be compared between projects and/or laboratories making them particularly suited to ongoing management of genetic diversity. The panel tested shows its utility across evolutionary scales, from population genetics to intra-species phylogenies, from the same genotyped dataset. The protocol applied is easily transferable to other study systems, thus making a substantial contribution to the establishment of standardized monitoring strategies for counteracting the harmful biodiversity loss in this epoch of accelerating global change.

4. Materials and methods

4.1. Study sites and species

The study focuses on two important national parks, Doñana National Park (DNP, Spain) and Kinabalu National Park (KNP, Malaysia) (Figure 1). Located within the Mediterranean Basin and Sundaland biodiversity hotspots (Myers et al., 2000), these protected areas were awarded World Heritage Site by UNESCO in 1994 and 2000, respectively. The parks contain a diversity of small mammals with no species shared, the only exceptions being the invasive black (R. rattus) and brown (R. norvegicus)
Table 2. Summary of the nuclear loci and associated primers used in this study. For the sake of clarity, amplicon size incorporates target sequence plus both primers. int.: intron; ex.: exon.

| Rat gene | Intron | Size - bp (R. norvegicus) | Forward primer 5'-3' | Reverse primer 5'-3' | Reference |
|----------|--------|---------------------------|----------------------|----------------------|-----------|
| Apeh 14 | (int.) 447 | GACCATCAGCCTACAGACATC | CCCAGTTCTCCACACCCA | Camacho-Sanchez et al., 2018 |
| Alkbh7 3 | (int.) 429 | GCTGGAGGTGGCTCTTCTG | CTGGCCTTTCCCTGTTGTCT | Camacho-Sanchez et al., 2018 |
| Agxt 10 | (int.) 560 | GGCTACAACTGGAGGGACATC | GTGCAGGGCCTCCYTCAGGGCCT | Rodríguez-Prieto et al., 2014 |
| Abcg8 9 | (int.) 430 | TTTCCAATGACTTCCGGGAC | GGCAAAGAAATAAGGACCAGCA | Camacho-Sanchez et al., 2018 |
| Abcb9 2 | (int.) 584 | GCATYGTSATCCAGAARAGCAYGGA | GCTGTGCGRTTCTCRTCRAARAAGCT | Rodríguez-Prieto et al., 2014 |
| Camacho-Sanchez et al., 2018 | |
| Alkbh7 3 | (int.) 429 | GCTGGAGGTGGCTCTTCTG | CTGGCCTTTCCCTGTTGTCT | Camacho-Sanchez et al., 2018 |
| Agxt 10 | (int.) 560 | GGCTACAACTGGAGGGACATC | GTGCAGGGCCTCCYTCAGGGCCT | Rodríguez-Prieto et al., 2014 |
| Abcg8 9 | (int.) 430 | TTTCCAATGACTTCCGGGAC | GGCAAAGAAATAAGGACCAGCA | Camacho-Sanchez et al., 2018 |
| Abcb9 2 | (int.) 584 | GCATYGTSATCCAGAARAGCAYGGA | GCTGTGCGRTTCTCRTCRAARAAGCT | Rodríguez-Prieto et al., 2014 |
| Camacho-Sanchez et al., 2018 | |
| Alkbh7 3 | (int.) 429 | GCTGGAGGTGGCTCTTCTG | CTGGCCTTTCCCTGTTGTCT | Camacho-Sanchez et al., 2018 |
| Agxt 10 | (int.) 560 | GGCTACAACTGGAGGGACATC | GTGCAGGGCCTCCYTCAGGGCCT | Rodríguez-Prieto et al., 2014 |
| Abcg8 9 | (int.) 430 | TTTCCAATGACTTCCGGGAC | GGCAAAGAAATAAGGACCAGCA | Camacho-Sanchez et al., 2018 |
| Abcb9 2 | (int.) 584 | GCATYGTSATCCAGAARAGCAYGGA | GCTGTGCGRTTCTCRTCRAARAAGCT | Rodríguez-Prieto et al., 2014 |
| Camacho-Sanchez et al., 2018 | |
| Alkbh7 3 | (int.) 429 | GCTGGAGGTGGCTCTTCTG | CTGGCCTTTCCCTGTTGTCT | Camacho-Sanchez et al., 2018 |
| Agxt 10 | (int.) 560 | GGCTACAACTGGAGGGACATC | GTGCAGGGCCTCCYTCAGGGCCT | Rodríguez-Prieto et al., 2014 |
| Abcg8 9 | (int.) 430 | TTTCCAATGACTTCCGGGAC | GGCAAAGAAATAAGGACCAGCA | Camacho-Sanchez et al., 2018 |
| Abcb9 2 | (int.) 584 | GCATYGTSATCCAGAARAGCAYGGA | GCTGTGCGRTTCTCRTCRAARAAGCT | Rodríguez-Prieto et al., 2014 |
| Camacho-Sanchez et al., 2018 | |

rats (Wells et al., 2006). However, KNP species richness (ca. 60 species encompassing the Orders Eulipotyphla, Rodentia, and Scandentia: Nor, 2001; Phillips and Phillips, 2016) is much higher than in DNP (ca. 10 species encompassing the Orders Eulipotyphla, Rodentia, and Lagomorpha: Palomo et al., 2007). Hence, to get more comparable panels of taxa and avoid biases associated with taxonomic composition of either one or the other site, we employed only a subset of the species in KNP while complementing the DNP set with other Iberian species (Table 1). Field samples were collected according to the guidelines of the American Society of Mammalogists (Sikes and Gannon, 2016), as approved by institutional animal care and use committees (Estación Biológica de Doñana Proposal Numbers CGL 2010-21524 and 433/2016). Field work in KNP is detailed in Camacho-Sanchez et al. (2019) and was undertaken with permission from Sabah Parks (TS/PTD/5/4Jld. 47 (25)) the Economic Planning Unit (100-24/1/299), and the Sabah Biodiversity Council (JKM/MBS.1000-2/2 (104)). Biological samples were exported with permissions from the Sabah Wildlife Department (JHL.600-3/7 Jld.7/19 and JHL.600-3/7 Jld.8/7) and Sabah Biodiversity Council (Ref: TK/PP:8/8Jld.2). Field work in DNP was undertaken with permission from the Junta de Andalucía (SGYB/AFR/DBP).
Samples were also obtained from the scientific collection at Estación Biológica de Doñana.

4.2. Loci selection and primer testing

A panel of 40 intronic loci was selected including 30 from among those identified as single-copy and potentially informative in Igea et al. (2010) for addressing the phylogeny of closely related mammals and developed in Camacho-Sánchez et al. (2018), plus another 10 developed by Rodríguez-Prieto et al. (2014) specifically for rodent phylogeny (Table 2, Table S1). The 30 loci were successfully used to study intraspecific phylogeographic patterns in the summit rat (Rattus norvegicus; Camacho-Sánchez et al., 2018). Here we describe in detail the process of primer design so it can be applied in other non-model organisms. The brown rat was used for primer design due to the large amount of genomic resources for this model organism as well as its occurrence among the focal taxa of the present study. After inspecting the species tree available at http://www.ensembl.org/info/about/speciestree.html, the house mouse (Mus musculus) was found to be the closest species to our target with a fully annotated genome. In previous studies (Igea et al., 2010; Rodríguez-Prieto et al., 2014; see Table S1 for details), the target introns had been selected via multiple filtering steps including long flanking exonic regions for primer design and appropriate levels of size conservation and variation within mammals. Loci selection for the present study relied on a size criterion, namely that introns of choice were ideally between 300-400 bp long as to be completely sequenced in Roche 454 by pyrosequencing chemistry (upper read length limit of about 500 bp) and Illumina MiSeq 300PE. Lower length limit was established to ensure a satisfactory information content and because fragment size homogeneity should increase sequencing efficacy. Human gene IDs corresponding to the introns selected were obtained and the BioMart tool (Smedley et al., 2009) in Ensembl (Herrero et al., 2016) was used to find the respective orthologues in R. norvegicus (Rnor 5.0) and M. musculus (GRCm38). These sets of genes were downloaded into Geneious 8.1.5 (http://www.geneious.com; Kearse et al., 2012) using the NCBI (National Center for Biotechnology Information) plugin and aligned using the MAFFT plugin 7.453 (Katoh and Standley, 2013) with default parameters. The annotations of aligned orthologues were used to manually check for intron number and size as well as the homology of flanking regions when compared to those of the sequences from Igea et al. (2010) and Rodríguez-Prieto et al. (2014). Exonic regions complying with length and conservation criteria were targeted for primer design in Primer3 (http://primer3.ut.ee/; Koressaar and Remm, 2007; Untergasser et al., 2012). Since the panel was intended to be suitable for multiplexing in a single PCR reaction, primer pairs for the loci selected by Igea et al. (2010) were designed to have a narrow delta Tm (25 bp: PCR product size 400 to 500 bp). The primers for the loci from Rodríguez-Prieto et al. (2014) were used without modification. A 2 bp GC clamp was added to enhance primer stability, while for all the other parameters default values were applied. When Primer3 failed to retrieve solutions complying with desired features, either stringency criteria were relaxed and GC clamps removed or new exonic regions were selected for primer design. Candidate primer pairs were mapped against their homologous regions in Geneious alignments and those mapping in the most conserved regions were finally selected. If mismatches at the 3' end occurred, either the respective region was excluded or ambiguities were introduced manually to a maximum of two per primer (Table 2, Table S1). The whole workflow is illustrated in Figure S1. The primers for

### Table 3

| Study area | Species | n | # loci | # alleles | Introns | Ghr | S | h | Θ | π | Ghr | S | h | Θ | π |
|------------|---------|---|--------|----------|---------|-----|---|---|---|---|-----|---|---|---|---|
| DNP | Rattus norvegicus | 6 | 39 | 1-6 | 2.2 | 1.7 | 0.2 | 0.002 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0.003 |
| DNP | Rattus rattus | 2 | 38 | 1-3 | 3 | 1.6 | 0.3 | 0.004 | 1 | 2 | 1 | 0 | 0 | 0 | 0 | 0 |
| KNP | Rattus albusensis | 3 | 40 | 1-4 | 2.2 | 1.7 | 0.3 | 0.01 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 |
| KNP | Sundamsy Muelleri | 6 | 39 | 1-5 | 5.5 | 2.9 | 0.5 | 0.003 | 1 | 2 | 0.7 | 0.001 |
| KNP | Niviventer cremonivent | 4 | 32 | 1-5 | 3.7 | 2.2 | 0.4 | 0.03 | 0 | 1 | 0 | 0 |
| KNP | Leopoldamy sabanus | 5 | 39 | 1-6 | 7.6 | 3.3 | 0.6 | 0.05 | 0 | 1 | 0 | 0 |
| KNP | Lenorhis canus | 1 | 18 | 1-2 | 0.9 | 1.3 | 0.4 | 0.003 | 0 | 1 | 0 | 0 |
| KNP | Maxomys ochraceivent | 1 | 30 | 1-2 | 0.7 | 1.2 | 0.2 | 0.002 | 0 | 1 | 0 | 0 |
| KNP | Maxomys whiteheadi | 5 | 34 | 1-9 | 8.1 | 3.9 | 0.6 | 0.02 | 0 | 1 | 0 | 0 |
| DNP | Mus spretus | 8 | 25 | 1-7 | 1.9 | 2 | 0.3 | 0.002 | 0 | 1 | 0 | 0 |
| DNP | Apodemus sytycticus | 7 | 14 | 1-11 | 16.9 | 5.7 | 0.7 | 0.01 | 0 | 1 | 0 | 0 |
| KNP | Chipropodymys pusillus | 1 | 28 | 1 | 0 | 1 | 0 | 0.003 | 0 | 1 | 0 | 0 |
| KNP | Arvicola sapidus | 3 | 9 | 1-4 | 7.6 | 1.6 | 0.2 | 0.002 | 0 | 1 | 0 | 0 |
| IB | Microtus cabrerae | 1 | 11 | 1-2 | 0.2 | 1.2 | 0.2 | 3.6 × 10⁻⁴ | 0 | 1 | 0 | 0 |
| IB | Chiromys nivalis | 3 | 9 | 1-4 | 2.8 | 2.1 | 0.4 | 0.001 | 0 | 1 | 0 | 0.001 |
| DNP | Eliomyx querinus | 5 | 3 | 1-3 | 4.7 | 2 | 0.5 | 4.7 | 3 | 4 | 0.9 | 0.001 |
| IB | Sciurus vulgaris | 3 | 9 | 1-2 | 0.3 | 1.2 | 0.1 | 4.44 × 10⁻⁴ | 0 | 1 | 0 | 0 |
| KNP | Callosciurus premosti | 2 | 6 | 1-3 | 3.7 | 1.8 | 0.4 | 0.01 | 0 | 1 | 0 | 0 |
| KNP | Sundascirus everetti | 2 | 9 | 1-2 | 0.9 | 1.3 | 0.2 | 0.002 | 0 | 1 | 0 | 0 |
| KNP | Sundascirus jeninki | 1 | 6 | 1 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 |
| KNP | Sundascirus lowei | 1 | 8 | 1-2 | 0.5 | 1.1 | 0.1 | 0.001 | 0 | 1 | 0 | 0 |
| DNP | Orcytogus canalicus | 2 | 13 | 1-2 | 2.6 | 1.3 | 0.2 | 0.003 | 0 | 1 | 0 | 0 |
| KNP | Tupla montana | 2 | 4 | 1-2 | 11.8 | 1.8 | 0.5 | 0.05 | 0 | 1 | 0 | 0 |
| KNP | Hylomyx saulius | 2 | 2 | 1 | 0.5 | 1.5 | 0.25 | 5 × 10⁻⁴ | 0 | 1 | 0 | 0 |
| DNP | Suncus etruscus | 2 | 8 | 1 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 |
| DNP | Crocidura russula | 2 | 8 | 1-4 | 1.1 | 1.6 | 0.3 | 0.002 | 0 | 1 | 0 | 0 |
| DNP | Crocidura suaveolens | 2 | 7 | 1-2 | 0.6 | 1.1 | 0.1 | 0.001 | 0 | 1 | 0 | 0 |

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the loci from Rodríguez-Prieto et al. (2014) were used without modification. Additionally, two commonly sequenced nuclear genes, Rbp3 and Ghr (Table S1), were also sequenced by amplified overlapping fragments for the purpose of compared phylogeny and variability (e.g., Pages et al. 2010; Fabre et al., 2013).

4.3. DNA extraction

DNA was isolated with phenol-chloroform and ethanol precipitation (Maniatis et al., 1985) or with SeraMag™ SpeedBeads (Thermo-scientific). DNA samples were quantified by Nanodrop spectrophotometry relying on absorbance readings at 260 nm wavelength and diluted to working concentrations of 15 ng/μl.

4.4. 454 amplicon library preparation (PCR) and sequencing

We prepared amplicon libraries following a two-step PCR. The 40 selected primer pairs were equimolar in a single PCR. Reactions included 1x Multiplex PCR Master Mix (Qiagen), 3.34 μM of primer mix and 20–50 ng of template DNA (final volume 25 μl). Primers had an M13 tail on their 5’ end (fw: 5’-GTCTTTCCAGTGCGAC; rev: 5’-AACAGCTATGACCATTG). The thermal profile was 95 °C for 15 min followed by 15 cycles of touchdown: 95 °C for 30 s, 65–60 °C for 30 s, 72 °C for 30 s, and then 15 more cycles with the annealing temperature at 60 °C with a final extension at 72 °C for 10 min. PCR products were then cleaned with SPRI beads (Roehland and Reich, 2012) and diluted 1:10 before the indexing PCR. Reaction mix consisted of 0.1 μM of each Multiplexing Index sequence plus the sequencing primers, 1x PCR Gold buffer, 1.5 mM MgCl₂, 200 μM of each dNTP, 0.2 U AmpliTaq Gold DNA Polymerase (Applied Biosystems) and 2 μl of diluted PCR product (final volume 12 μl). PCR conditions were: 98 °C for 30 s and then 25 cycles of 98 °C for 10 s, 56 °C for 20 s, 72 °C for 45 s. PCR products were checked on a 2% agarose gel and viewed with Quantity-One software (Bio-Rad Laboratories) for relative quantification of 300–650 bp-long fragments prior to equimolar pooling. Products were cleaned with 1x SPRI beads and quantified with a fluorometer. Finally, an emulsion PCR was carried out with the Roche emPCR-a kit prior to sequencing on a GS Junior 454 sequencer following manufacturer’s instructions.

4.5. Illumina amplicon library preparation (PCR) and sequencing

Library was amplified in 20 μl reaction of 1x Phusion Master Mix (New England Biolabs), 0.05 μM of each primer, and 20–50 ng of template DNA. Primers had a tail (fw: 5’-TCCTTTCCCTACAGCAGGCTCTTGATCTC; rev: 5’-GAGTCCAGCGTGCGTCTTCGGCGATCT) complementary to indexing primers. PCR was run as follows: 98 °C for 1 min then 25 cycles of 98 °C for 10 s, 61 °C for 30 s, 72 °C for 45 s with a final denaturation at 95 °C for 3 min. PCR products were checked and cleaned with SPRI beads as above. The indexing PCR was performed in a final volume of 12 μl including 0.42 μM of each Multiplexing Index sequence plus the sequencing primers, 1x Phusion Master Mix, and 1 μl of undiluted PCR product. Thermal profile was 98 °C for 30 s, the 12 cycles of 98 °C for 10 s, 60 °C for 30 s, 72 °C for 45 s and a final denaturation at 95 °C for 3 min. Indexed PCR product were cleaned, quantified and pooled at equimolar ratios as above before sequencing of 300 bp paired end

Figure 2. Histogram showing the high variance associated with read depth across different loci in R. norvegicus and L. sabanus (one PCR per species).
Figure 3. ML tree of Rattini tribe based on the whole set of introns amplified in the representatives tested in this study and built using the coalescent approach implemented in ASTRAL. The plot at the bottom of the figure indicates the loci amplified in each taxon and their completeness (Prop.). Animal photos are not to scale; for credits see the Acknowledgements and Appendix S2.

Figure 4. MtDNA-based phylogenies of Rattini representatives sampled in this study. A. ML tree reconstructed on the basis of the whole mitogenome (10,852 bp). B. ML tree reconstructed on the basis of the cytochrome b gene.
reads on the Illumina MiSeq platform at the Johns Hopkins University Genetic Resources Core Facility (Baltimore, MD, USA).

4.6. Illumina shotgun library preparation and sequencing

Aliquots of DNA extracts were diluted to 20 ng/μl in sonication buffer (10 mM Tris, 1 mM EDTA pH 7.5–8) to 100 μl and sheared in an ultrasonic bath (Bioruptor UCD-20 0TM-EX Sonication System). Sonication cycle conditions were adjusted to the desired target size (around 500 bp); two rounds of 3 cycles of 30 s on High (H)/30 s off with a spin in the middle. Libraries were prepared with the Illumina Kapa Library Preparation Kit (Kapa Biosystems, Wilmington, Massachusetts, US) using ¼ reactions. Cleaning steps were done with SPRI beads as above. We used 10 μM Y-adapter (5'-5Phos-GATCGGAAGAGCACACGTCTGAACTCCAGTCAC and 5'-ACACTCTTTCCCTACGACGCTCTTCTCGATCTG*T; 5Phos, 5'phosphorylation; *phosphorothioate bond) previously annealed as in (Meyer and Kircher, 2010). The indexing PCR was done using a double indexing strategy as in (Kircher et al., 2012), with the PCR thermal conditions: 98 °C for 45 s then 8 cycles of 98 °C for 15 s, 60 °C for 30 s, 72 °C for 60 s with a final extension at 72 °C for 5 min. The libraries were cleaned, quantified, pooled and enriched for introns and mitochondrial DNA in independent hybridization reactions. We prepared the nuclear probes from PCR products using R. norvegicus as template for all the species tested. Each locus was amplified separately, pooled at equimolar ratios and cleaned. Mitogenomes were amplified by long-range PCR in two overlapping fragments of approximately 7 kb and 9 kb using primers from Sasaki et al. (2005) and sheared. Further details on primer and adapter sequences are reported in Camacho-Sanchez et al. (2017). In this case, the target regions across our sample were enriched with probes obtained from the same or related species. Specifically, we used R. norvegicus, S. vulgaris and C. rustula PCR products as template for Muridae-Cricetidae, Sciuridae-Gliridae and Soricidae-Erinaceidae representatives, respectively. For Leporidae and Tupaiidae we used O. cuniculus and T. montana templates. Then, the nuclear and mitochondrial baiting molecules were biotinylated as in Maricic et al. (2010) and the enrichment was done accordingly, with modifications as in Camacho-Sanchez et al. (2017). The enriched libraries were quantified by qPCR and were subsequently re-amplified as above except for the initial denaturation at 98 °C for 45 s and 15 or 20 PCR cycles depending on their concentration. The sequences of PCR indexing primers, indexing oligos and reamplification primers are as in Camacho-Sanchez et al. (2017). The re-amplified libraries were cleaned and quantified using qPCR. Final libraries were pooled in equimolar ratios and sequenced with 100 bp paired end reads on the Illumina HiSeq 2500 platform at the Johns Hopkins University Genetic Resources Core Facility.

4.7. Intron genotyping - amplicon libraries

Size filtering, read demultiplexing, primer trimming, amplicon assignment and allele calling of FASTQ data obtained from amplicon libraries (454 and Illumina MiSeq sequencing platforms) were performed separately with AmpliSAS in AmpliSAT, a package specifically for automated amplicon assignment from NGS data (Sebastian et al., 2015). The tool AmpliCHECK was first used to perform exploratory assays and detect likely artefacts due to PCR or sequencing errors by using default analysis parameters for 454 and Illumina data, respectively. The lack of prior information on intron length in most of the surveyed taxa and the low number of individuals analyzed for each taxon did not allow adopting highly restrictive criteria for artefact detection at this step of the pipeline. However, the alleles labelled as putative errors in AmpliCHECK after read quality filtering were removed from further analyses. Hence, the tool AmpliSAS was used with default filtering and clustering parameters applying to each sequencing technology, while selecting 2 as both maximum number of alleles and minimum amplicon depth, and fixing 5000 as maximum number of reads per amplicon.

4.8. Intron genotyping - shotgun libraries

Adaptor sequences were removed with cutadapt 1.8.3 (Martin, 2011) and output files were imported into Geneious. Paired end reads were iteratively mapped (5 cycles) with medium-low sensitivity against species-specific references represented by intron sequences obtained in the same taxon by means of amplicon libraries. When a given locus was not available, homologous sequences of the closest relative available in Ensembl were used as reference (Table S3). Consensus sequence callings were performed with a 75% threshold and two read minimum. BAM files generated in Geneious were processed with SAMtools 0.1.18 (Li et al., 2009) to remove PCR duplicates. Once re-imported in Geneious, BAM files were newly inspected and consensus sequences finally obtained by applying the same parameters as above. Intron sequences retrieved from amplicon and shotgun libraries were aligned in Geneious with MAFFT plugin for each locus and species. The alignment was carefully inspected to check for mismatches between data obtained with different sequencing platforms. When more than two alleles from the same individual were retrieved across amplicon and shotgun libraries, these were visually inspected to evaluate their possibly undetected artifactual nature and purged from multiple alignments of orthologous intronic sequences. When less than five reads where available for shotgun data, polymorphisms were confirmed only if they occurred in more than one individual and/or matched amplicon data from the same individual. However, we scored a locus as having worked in a given species (e.g., being amplified) when at least one read mapped properly to the reference.

4.9. Mitogenome assembly

FASTQ files with adapters trimmed (see above) were mapped to the phylogenetically closest available mitogenome reference in GenBank (Table S4) using the mapping tool in Geneious with medium-low sensitivity and 5 iterations. We assembled one mitogenome per species. Consensus sequence callings were performed with over 75% threshold and two reads as minimum. The mapping files were exported in BAM format and the PCR duplicates removed using SAMtools, then reimported into Geneious.

4.10. Locus amplification and variability

The alleles identified as putative errors in AmpliCHECK after read quality filtering were removed from further analyses. Likewise, alleles found at a frequency below 25% in a given individual were conservatively discarded as possibly arising from undetected sequencing errors. This threshold is consistent with that proposed in recent works (e.g., O’Leary et al., 2018). The number of alleles per locus and taxon were scored to evaluate the polymorphism of loci tested.

4.11. Indices of diversity

For each species we computed number of polymorphic sites (S), number of haplotypes (h), haplotype diversity (θ), and nucleotide diversity (π) at each locus with DnaSP 5.10.1 (Librado and Rozas, 2009).

4.12. Phylogenetic reconstructions

A phylogeny of Rattini was built relying on all intron loci yielding sequences in members of this group. We retained the 9 species from Rattini to maximize the number of loci with homologous sequences. For each species we selected the most common allele. We completed the data for R. baluensis with sequences from GenBank: MG424797 (Mmp9), MG425076 (Npr2), MG425817 (Tmem87a) (Camacho-Sanchez et al.,
aligned using MAFFT. These were fragmentary data from enrichment. The sequences for each locus were RAxML using the GTR length as well as columns with gaps in more than 60% of the sites. The sequences for which 40% or more had coverage below 50% of their total 2019) to remove characters from columns with a coverage below 3 and, errez et al., 2009) to remove transex. Alignments from protein-coding genes were split into codon positions 1, 2 and 3, using AMAS (Borowiec, 2016). All alignments were concate-nated with the same software. The complete alignment contained 10,852 positions with 0.1% of ambiguous positions. The best partition scheme for the resulting 48 partitions was determined with PartitionFinder 2.1.1 (Lanfear et al., 2012). Phylogenetic reconstructions were performed in a ML framework with RAxML 8.0.0 (Stamatakis, 2014). We followed the for the same taxa.

Declarations

Author contribution statement

Giovanni Forcina: Analyzed and interpreted the data; Wrote the paper.
Miguel Camacho-Sanchez: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data.
Fred Y. Y. Tuh: Contributed reagents, materials, analysis tools or data.
Sacramento Moreno: Contributed reagents, materials, analysis tools or data.
Jennifer A. Leonard: Conceived and designed the experiments; Contributed reagents, materials, analysis tools or data.

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Data availability statement

DNA sequences from complete mitochondrial genomes and nuclear loci have been deposited in GenBank with accession numbers MW209719-MW209719 and MW394671-MW396361, respectively. Details on phylogenetic methods and phylogenetic trees are available at github.com/csmiguel/community-genetics and at https://doi.org/10.5281/zenodo.4275480.

Declaration of interests statement

The authors declare no conflict of interest.

Additional information

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