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Building a Portuguese Coalition for Biodiversity Genomics

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90 Abstract

The diverse physiography of the Portuguese land and marine territory, spanning from continental Europe to the Atlantic archipelagos, has made it an important repository of biodiversity throughout the Pleistocene glacial cycles, leading to a remarkable diversity of species and ecosystems. This rich biodiversity is under threat from anthropogenic drivers,

95 such as climate change, invasive species, land use changes, overexploitation or pathogen 96 (re)emergence. The inventory, characterization and study of biodiversity at inter- and intra-97 specific levels using genomics is crucial to promote its preservation and recovery by informing 98 biodiversity conservation policies, management measures and research. The participation of 99 researchers from Portuguese institutions in the European Reference Genome Atlas (ERGA) 100 initiative, and its pilot effort to generate reference genomes for European biodiversity, has 101 reinforced the establishment of Biogenome Portugal. This nascent institutional network will 102 connect the national community of researchers in genomics. Here, we describe the 103 Portuguese contribution to ERGA's pilot effort, which will generate high-quality reference genomes of six species from Portugal that are endemic, iconic and/or endangered, and include 104 plants, insects and vertebrates (fish, birds and mammals) from mainland Portugal or the 105 106 Azores islands. In addition, we outline the objectives of Biogenome Portugal, which aims to (i) 107 promote scientific collaboration, (ii) contribute to advanced training, (iii) stimulate the participation of institutions and researchers based in Portugal in international biodiversity 108 genomics initiatives, and (iv) contribute to the transfer of knowledge to stakeholders and 109 engaging the public to preserve biodiversity. This initiative will strengthen biodiversity 110 111 genomics research in Portugal and fuel the genomic inventory of Portuguese eukaryotic species. Such efforts will be critical to the conservation of the country's rich biodiversity and 112 113 will contribute to ERGA's goal of generating reference genomes for European species.

114

115 Keywords

116 Biodiversity, Genomics, Conservation, Portugal

117

118 Sumário

A diversa fisiografia dos territórios terrestre e marinho portugueses, que se estende desde a Europa continental até aos arquipélagos do Atlântico, promoveu a sua importância como repositório de biodiversidade ao longo dos ciclos glaciares do Pleistoceno, resultando numa notável diversidade de espécies e ecossistemas. No entanto, esta biodiversidade enfrenta 123 ameaças decorrentes de atividades humanas, tais como as alterações climáticas, invasão de 124 espécies, mudanças no uso do solo, exploração excessiva e (re)emergência de agentes 125 patogénicos. A fim de promover a preservação e recuperação da biodiversidade, é crucial 126 inventariar, caracterizar e estudar a biodiversidade a níveis inter e intraespecíficos, utilizando 127 técnicas genómicas. Essa informação é essencial para informar políticas de conservação da 128 biodiversidade, medidas de gestão e pesquisas futuras. A participação de investigadores de 129 instituições portuguesas na iniciativa European Reference Genome Atlas (ERGA), e no 130 projeto-piloto para gerar genomas de referência para a biodiversidade europeia, promoveu a 131 criação do Biogenoma Portugal. Esta rede institucional em desenvolvimento visa conectar a comunidade nacional de investigadores em genómica. Neste contexto, descrevemos a 132 contribuição portuguesa para o projeto-piloto do ERGA, que tem como objetivo gerar 133 genomas de referência de alta qualidade de seis espécies de Portugal. Estas espécies são 134 135 endémicas, icónicas e/ou ameacadas e incluem plantas, insetos e vertebrados (peixes, aves e mamíferos) do território continental de Portugal e das ilhas dos Açores. Adicionalmente, 136 delineamos os objetivos do Biogenoma Portugal que visa: (i) promover a colaboração 137 científica, (ii) contribuir para a formação avançada, (iii) estimular a participação de instituições 138 139 e investigadores baseados em Portugal em iniciativas internacionais de genómica da biodiversidade e (iv) contribuir para a transferência de conhecimento e envolvimento do 140 público em geral na preservação da biodiversidade. Esta iniciativa fortalecerá a investigação 141 em genómica da biodiversidade em Portugal e contribuirá para o objetivo do ERGA de gerar 142 143 genomas de referência para espécies europeias. Estes esforços terão um papel fundamental na conservação da biodiversidade do país e na promoção da conscientização sobre a 144 importância de preservar a diversidade biológica. 145

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147 Introduction

The remarkable diversity of life on Earth is essential for maintaining ecosystem stability, fostering ecological interactions among species, regulating the climate, and providing fundamental resources that sustain human well-being and promote the development of 151 society. Today, Earth's biodiversity, ecological and evolutionary heritage, and the ecosystem 152 services they provide are under intense and extensive strain as a result of many direct and 153 indirect anthropogenic activities. These major activities include habitat destruction and 154 fragmentation caused by agricultural expansion and other forms of land conversion, 155 overexploitation of natural resources, overharvesting, illegal killing and trading of wild species, 156 climate changes, environmental pollution, and global spread of exotic species (Ceballos et al., 2017; Cowie et al., 2022; Habel et al., 2019; Karger et al., 2021; O'Hara et al., 2021). These 157 158 pressures not only result in the decline and extinction of populations, species and ecosystems, 159 but also have cascading effects on various dimensions of biodiversity, including genetic, functional or phenological diversity and the interactions within biodiversity networks. These 160 impacts harm the health of natural populations and their survival, and reduce their potential to 161 adapt to environmental challenges (Dauphin et al., 2021). Characterising and understanding 162 163 in detail the genetic variation underlying biodiversity at all levels provides essential modern tools to delineate sound and efficient strategies to mitigate the impacts of human activities, 164 design management plans for conservation and restoration, and support data-driven 165 biodiversity policy development (Browne et al., 2019; Hohenlohe et al., 2021; Ralls et al., 2020; 166 Segelbacher et al., 2022). Regional biodiversity genomics initiatives that promote research 167 and applications under the umbrella of international coalitions, can play an important role in 168 169 this global challenge.

Europe hosts a remarkable biodiversity richness despite its temperate climate and historically 170 171 anthropogenic landscapes (Ette & Geburek, 2021), and has an important biodiversity hotspot 172 along the Mediterranean basin (Habel et al., 2019; Myers et al., 2000). Most of the continental territory of Portugal, located in south-western Europe in the Iberian Peninsula, is part of the 173 174 Mediterranean biodiversity hotspot (Myers et al., 2000). The country also includes the volcanic 175 Atlantic archipelagos of Azores, Madeira and Selvagens (Figure 1). Even though Portugal is the 19th largest European country (land area ~92000 km²), it holds the 5th largest marine territory 176 (~1.7 million km²) (Pauly et al., 2020). As a result of its unique geographic location, climate 177 and geophysical features, Portugal comprises a variety of landscapes and ecosystems and 178

179 harbours a large diversity of species. The Iberian Peninsula provided refugia for species throughout the glacial periods and remains one of the most important repositories of 180 181 biodiversity in Europe, both at inter- and intraspecific level (Gómez & Lunt, 2007; Hewitt, 182 2004). In addition, the geographic isolation of the archipelagos of Azores, Madeira and 183 Selvagens, which together with the Canary and Cabo Verde archipelagos form the 184 biogeographical unit of Macaronesia, has led to the divergence and speciation of many 185 lineages that resulted in many endemisms (Florencio et al., 2021). According to the 186 Portuguese National Strategy for the Conservation of Nature and Biodiversity 2030 (ENCNB 187 2030; https://dre.pt/dre/detalhe/resolucao-conselho-ministros/55-2018-115226936), Portugal holds ~35,000 animal and plant species, representing a relevant proportion of these species 188 groups present in Europe. The IUCN Red List database (https://www.iucnredlist.org; 27 April 189 190 2023) lists ~800 of the assessed species as endemic to the continental mainland or 191 archipelagos of Portugal. There are 414 protected areas in Portugal encompassing 22.28% of the land and 2.46% of marine water, and 404 species and 102 habitats are protected under 192 EU (Biodiversity Information 193 law System for Europe: https://biodiversity.europa.eu/countries/portugal). The ENCNB 2030 recognizes the 194 195 importance of a systematic inventory and characterization of biodiversity at all levels ecosystems, species and genetic diversity - to properly anchor conservation strategies. 196 Genomics provides unprecedented power to understand biodiversity at the inter- and 197 intraspecific levels (Formenti et al., 2022; Theissinger et al., 2023). Thus, gathering scientific 198 expertise in the area, coordinating research efforts and promoting genomic studies of native 199 200 Portuguese species, can give an invaluable contribution to preserving the rich biodiversity of Portugal and its ecosystem services. 201

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203 ERGA-Portugal community and the Pilot effort

The European Reference Genome Atlas (ERGA; <u>https://www.erga-biodiversity.eu/</u>) is an ambitious and ground-breaking initiative that aims to revolutionise our understanding of biodiversity by creating a comprehensive genomic resource for European species. ERGA is 207 implemented under the umbrella of the Earth Biogenome Project (EBP; 208 https://www.earthbiogenome.org/), the global network of networks that aims to catalogue the 209 eukaryotic life diversity of Earth through genome sequencing. ERGA gathers researchers, 210 scientists, and biodiversity enthusiasts to collect and sequence the genomes of species across 211 Europe, providing a valuable tool for studying and preserving biodiversity, understanding 212 evolutionary processes, and addressing pressing environmental challenges. The network leverages the power of genomics to unlock the secrets of European biodiversity and foster 213 214 collaborative efforts towards its conservation and sustainable management.

215 ERGA includes more than 50 members in Portugal, distributed across seven research units, five universities and two natural history museums (May 31st, 2023; Figure 1). This community 216 217 of ERGA members based in Portugal is called ERGA-Portugal, and covers a wide range of 218 expertise that is relevant for biodiversity genomics research, from taxonomy and systematics across diverse biological groups (e.g., microorganisms, plants, insects, fish, amphibians, birds 219 220 or mammals), to comparative and population genomics and bioinformatics. This community 221 communicates through open meetings, mailing lists and message sharing channels to 222 facilitate the exchange of biodiversity genomic expertise across Portugal. Many Portuguese community members are also active participants in ERGA's scientific committees. 223

To contribute to the establishment and mission of ERGA, ERGA-Portugal participated in the 224 pilot test of ERGA (Cartney et al., 2023). This pilot aimed at testing and developing the 225 226 biodiversity genomic networks across Europe through the sequencing of several European species, with the contribution of expert ERGA members and European institutions and without 227 a centralised source of funding. In Portugal, the goal of this project was to consolidate the 228 ERGA-Portugal community, and initiate collective efforts to generate reference genomes for 229 Portuguese species. This pilot effort in Portugal focused on endemic, endangered and iconic 230 species from several taxonomic groups and different ecosystems in both the mainland and 231 the Azores islands. From an initial list of 53 species proposed by members of ERGA-Portugal, 232 233 11 were prioritised based on ERGA's feasibility criteria defined by the ERGA Sampling and Sample Processing Committee (Böhne et al., 2023). These species were then ranked based
on an anonymous online survey open to all members of ERGA-Portugal at the time. The six
species with the highest score were included in the pilot effort. This shortlist included one fish,
one mammal, one bird, one invertebrate and two plants (Figure 2).

238 Each species was championed by a genome team, led by one or two species ambassadors, 239 and included ERGA-Portugal members and other ERGA members with transdisciplinary skills, who were responsible for successfully implementing each step for generating a reference 240 241 genome. These steps included, e.g., sampling, taxonomic identification, vouchering, 242 laboratory work, sequencing, assembly, annotation and downstream analysis. These six genome teams, detailed below for each species, promoted new national and international 243 collaborations. The implementation of this project also promoted interactions with national 244 authorities, for example, to obtain permits for capture, storage and export of samples, in 245 246 coordination with the Portuguese National authority on Nature Conservation (Institute for Nature Conservation and Forests - ICNF), and the Azorean Regional Directorate for the 247 Environment and Climate Change (DRAAC). For mainland Portugal, the capture and 248 collection of samples of wildlife specimens were authorised by the ICNF, while for the 249 250 Autonomous regions of Azores permission was given by the DRAAC. For the Azores, collections complied with the Access and Benefit Sharing provisions codified in the Nagoya 251 Protocol (Convention on Biological Diversity, 2010). For all species, the sampling steps 252 followed the guidelines provided by the ERGA SSP committee (Böhne et al., 2023). Sampling 253 was optimised to ensure that High Molecular Weight (HMW) DNA and RNA could be obtained 254 from at least one individual and ideally from multiple tissues (see details for each species 255 256 below). The immediate preservation and transport of tissues in liquid nitrogen was prioritised 257 and tissues were subsequently maintained at -80°C conditions.

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259 ERGA-Portugal pilot species

260 Iberian minnowcarp - Anaecypris hispanica (Steindachner, 1866)

261 Common name in Portuguese: saramugo

262 The **Iberian minnowcarp** (Anaecypris hispanica) is a strictly freshwater fish with a short 263 lifespan that inhabits a restricted geographical area in two river basins of the Southern Iberian 264 Peninsula (Cardoso, 2023; De Miguel et al., 2010), a region which is severely affected by the 265 effects of global climate change. This endemic species is considered to be the most 266 endangered strictly freshwater fish in Iberian rivers, and is listed as Endangered on the IUCN 267 Red List (Crivelli, 2006) and as Critically Endangered on the Portuguese Red List (Cabral et al., 2005). These fish live in vulnerable freshwater ecosystems and are exposed to multiple 268 269 threats, including increased temperatures and propensity for drought, pollution, habitat 270 fragmentation, dams, intensive water use and invasive species proliferation (Cabral et al., 2005; Cardoso, 2023). The Iberian minnowcarp is considered an iconic species as it is the 271 only living member of the genus Anaecypris, representing a phylogenetically unique old 272 lineage that persisted in the Iberian Peninsula (Perea et al., 2010). 273

274 Sampling and sample processing: An adult female was collected by hand net from an openair breeding tank at Aquário Vasco da Gama (Cruz Quebrada-Dafundo, Portugal), where an 275 ex-situ conservation program is ongoing with captive-born fish derived from 37 individuals of 276 River Chança (Guadiana river basin, Portugal). The individual was sampled, euthanized and 277 278 dissected on the same day, according to permits from the ICNF (Permit P-026382/2021) and animal welfare regulations of the host institution (ORBEA-MARE 02/2021). The head of the 279 fish was flash-frozen and used for biobanking. Fin clips were used for barcoding and preserved 280 in ethanol. Fin clips were also used to derive cell lines, which were flash-frozen and can be 281 used for karyotyping and further genetic studies in the future. Tissues were collected 282 immediately and flash-frozen in liquid nitrogen. Liver, spleen, muscle, eyes and ovary tissues 283 were collected. 284

285 Sequencing and assembly: Aliquots of these tissues were shipped to the Max Planck Institute 286 of Molecular Cell Biology and Genetics (MPI-CBG, Germany), where the remaining steps for 287 DNA extraction and sequencing were performed. The HMW DNA extraction and library 288 preparation for PacBio HiFi was done using the spleen, whereas Hi-C library preparation for 289 Illumina was done using muscle. RNA-Sequencing and Iso-Seq libraries from eye and gonad tissues were produced and sequenced. Sequencing data was used to obtain a genome assembly, which was done and completed by the genome team members from MPI-CBG (Germany) using the VGP standards. A manually curated assembly is completed, and the annotation of the genome is ongoing in a collaborative effort involving researchers from ERGA-Portugal, as well as institutions and national computational resources.

295 Expected results: A high-quality reference genome will open the door to reconstruct the 296 evolutionary history of this monospecific genus, elucidating its relationship with other fish 297 lineages. Given that the Iberian minnowcarp is fragmented into small populations, inhabiting 298 a region highly impacted by drought and water scarcity (Sousa-Santos et al., 2014), the availability of its reference genome will be crucial to understand adaptation to intermittent 299 environments, to inform conservation management, and predict responses to increasing water 300 temperatures. The Iberian minnowcarp is included in ongoing *ex-situ* conservation programs, 301 302 hence these genomic resources will allow quantifying the loss of genetic diversity due to consecutive generations bred from small initial stocks of wild breeders, as well as optimise 303 and monitor future ex-situ efforts. 304

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306 Iberian hare - Lepus granatensis (Rosenhauer, 1856)

307 Common name in Portuguese: lebre Ibérica

308 The **Iberian hare** (*Lepus granatensis*), also known as Granada hare, is a lagomorph species endemic to the Iberian Peninsula, and the only hare species present in Portugal (Alves et al., 309 310 2023). It is an iconic species of major ecological and economic importance in the Iberian 311 ecosystems, acting as an important prey to apex predators and a relevant small game species in Portugal and Spain. Genetic studies have promoted this species as an outstanding model 312 313 to understand diverse evolutionary questions, such as the impact of glacial cycles in repeated 314 changes in species ranges, and the influence of introgressive hybridization on their adaptive potential (Seixas et al., 2018). It is currently classified as Least Concern in the IUCN Red List 315 (Soriguer & Carro, 2018), but a marked decline over the last decade led to its classification as 316 Vulnerable in the last review of the Red Book of Mammals of Portugal (Mathias et al., 2023). 317

The rate of population decline has increased due to the emergence of a natural recombinant derived from the rabbit *Myxoma virus*, the *hare Myxoma virus* (ha-MYXV) (Águeda-Pinto et al., 2019; Dalton et al., 2019).

Sampling and Sample Processing: One male individual Iberian hare was sampled in Mértola, southern Portugal, during the regular permitted hunting season, duly authorised by ICNF (Permit 012/2022/CAPT). Tissues were collected shortly after the animal's death and flash frozen in liquid nitrogen. Tissues from several organs were collected during the necropsy and kept at -80°C for biobanking: liver, kidney, spleen, lung and testis. The whole body will be prepared for deposition at the Natural History and Science Museum of the University of Porto (MHNCUP_MAM 0812).

Sequencing and Assembly: Tissue samples were shipped to the University of Antwerp for 328 Omni-C and RNA-Sequencing library preparation, and to the University of Florence for PacBio 329 330 DNA library preparation. Final Omni-C libraries were generated using kidney tissue, while RNA-Sequencing libraries were produced for liver, kidney, spleen, lung and testis tissues. 331 These libraries along with PacBio DNA libraries were sequenced at the University of Florence. 332 Long-read DNA sequencing was performed in HiFi mode in a PacBio Sequel IIe platform using 333 334 five 8 million ZMW SMRT cells. Sequencing data was used to produce a genome assembly following the gold standard instructions implemented in the VGP Galaxy Pipeline (Batut et al., 335 2018; Lariviere et al., accessed June 2023). The genome manual curation and annotation is 336 ongoing, and involves ERGA-Portugal members, as well as institutions and national 337 338 computational resources.

Expected results: Generating a high-quality reference genome will improve genomic studies in this species, upgrading the genomic resources for this and related species. This will open the door for fundamental research in speciation and adaptation genomics. Furthermore, it will anchor research aiming to quantify the impact of emerging pathogens on the naïve host hare populations, which can be used to inform conservation measures.

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345 Azores laurel - Laurus azorica (Seub.) Franco

346 Common name in Portuguese: louro-da-terra and louro-bravo

The Azores laurel (Laurus azorica) is a micro or mesophanerophyte dioecious perennial tree 347 348 endemic to the Azores Islands. It can be commonly found in submontane Laurus forests (Elias 349 et al., 2016) but also occupies habitats such as coastal scrublands, among other native 350 Azorean vegetation (Elias et al., 2016). It is listed as Least Concern in the IUCN Red List (Silva 351 & Beech, 2016), yet it is facing habitat loss due to competition with invasive species and alteration of habitats resulting from the implementation of production forests and pastures 352 353 (Borges Silva et al., 2018; Dutra Silva et al., 2019; Hortal et al., 2010; Lourenço et al., 2011). 354 The genus Laurus L. (Lauraceae) is currently restricted to isolated refugia in the southern Black Sea area, Mediterranean Basin, Northern Africa, and the Macaronesian archipelagos. 355 Its taxonomic classification remains uncertain, with some recognizing the existence of two 356 species, Laurus azorica (Seub.) Franco, endemic from the Azores and Laurus nobilis L. 357 358 However, while some molecular studies contradict this distinction (Rodríguez-Sánchez et al., 2009), others even recognise a third species, Laurus novo-canariensis Rivas Mart., Lousã, 359 Fern.Prieto, E.Dias, J.C.Costa & C.Aguiar (Rivas-Martínez et al., 2002). 360

Sampling and sampling processing: Fresh branches with healthy leaves were collected using 361 362 gardening scissors from one mature male individual in the Azores islands, specifically, São Miguel Island, in Lombadas. The branches were loosely wrapped in moist absorbent paper, 363 364 placed inside an open plastic bag and sent on the same day through express mail to the University of Lisbon, where they were kept at -20°C until DNA extraction. The corresponding 365 366 voucher duplicate was dried using standard herbarium procedures and stored in the AZB herbarium at the Biology Department of the Faculty of Sciences and Technology of the 367 University of the Azores, with voucher AZB4382, and collector code LZ001. A second 368 369 sampling was conducted during March 2023 and followed the same methodology. The Nagoya 370 certificate has CCIR number 61/2021/DRCTD.

371 *Sequencing*: The HMW DNA extraction was performed at the Department of Plant Biology of 372 the Faculty of Sciences of the University of Lisbon (FCUL), using a modified 373 cetyltrimethylammonium bromide (CTAB) protocol (Doyle & Doyle, 1987, later modified by Weising et al., 1994) adjusted based on the exchange of protocols within the ERGA community. DNA extraction was challenging, as extractions did not pass the quality control for PacBio Hi-Fi library preparation and sequencing. Leaf tissue was thus shipped in dry ice to the University of Florence, where the libraries for Pac-Bio were prepared and sequenced. The assembly of the genome is ready to start, and will involve ERGA-Portugal members, as well as institutions and national computational resources.

Expected results: A high-quality reference genome will allow performing phylogenetic and population genetic studies to reconstruct the evolutionary history of this group, contributing to clarify the diversity and taxonomy of the *Laurus* genus.

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384 Black wheatear - Oenanthe leucura (Gmelin, 1789)

385 Common name in Portuguese: chasco-preto

386 The **black wheatear** (*Oenanthe leucura*) is a passerine bird species confined to North Africa and southwest Europe, a region where global environmental changes (namely desertification 387 and agriculture abandonment) are impacting the distribution of many bird species (Fernández-388 Nogueira & Corbelle-Rico, 2018, 2020; Fusco et al., 2021). Black wheatears are no exception, 389 390 with findings supporting that some edge populations have already vanished (France) whilst others continue to decrease (Western populations in Portugal and Spain). Despite the decline 391 and fragmentation of these populations, an isolated population located in northern Portugal 392 (~100 pairs) still subsists, confined mainly to cultivated lands (primarily vineyards and olive 393 orchards) of the Portuguese section of the Douro River, internationally known for the Port 394 wine. For this reason, despite being classified globally as Least Concern by the IUCN Red 395 396 List, Portuguese populations are classified as Critically Endangered in the Portuguese Red 397 List (Cabral et al., 2005).

Sampling and sample processing: Due to the small size of the black wheatear population in Portugal, the sacrifice of a bird would be unjustified and, therefore, ICNF authorised the collection of blood samples from live birds (Permit 720/2021/REC). Three individuals (one female and two males) were caught using spring traps at the Douro Valley, near the Tua river 402 mouth. Each bird was ringed, photographed and measured, and a blood sample (~100 403 microliters) was taken by venepuncture at the ulnar vein with a microcapillary. Each blood 404 sample was immediately stored in liquid nitrogen and transferred to a -80°C freezer on the 405 same day.

406 Sequencing and assembly: Samples were sent to the ERGA partners in dry ice, to generate 407 Omni-C (University of Antwerp) and PacBio HiFi data (University of Florence) from the female 408 (C96097) and RNA-Sequencing (University of Antwerp) from one of the males (E29638). All 409 DNA libraries were sequenced at the University of Florence. Long-read DNA sequencing was 410 performed using two 8 million ZMW SMRT cells. Genome assembly was performed following the gold standard instructions implemented in the VGP Galaxy Pipeline (Batut et al., 2018; 411 Lariviere et al., accessed June 2023). The genome manual curation and annotation is ongoing, 412 involving ERGA-Portugal members, as well as institutions and national computational 413 414 resources.

Expected results: A high-quality reference genome, combined with population genomics data, 415 will allow to characterise the structure and history of this population, which is closely 416 associated with anthropogenic agricultural lands in the Douro valley, yet geographically 417 418 isolated from other Iberian populations that occur mostly in natural habitats. This isolation and fragmentation could have irreversible effects on the survival of these populations and their 419 420 conservation may require the implementation of specific conservation measurements. Moreover, it will establish the baseline for genomic analysis of the Portuguese populations, 421 422 providing information on individuals' relatedness and dispersal capacity.

423

424 Portuguese crowberry - Corema album (L.) D.Don

425 Common name in Portuguese: Camarina and Camarinha

The **Portuguese crowberry** (*Corema album*) is a dioecious perennial woody shrub endemic to the Atlantic coast of the Iberian Peninsula (ssp. *album*), and to the Azores Islands (ssp. *azoricum*). In the Iberian Peninsula it inhabits coastal areas from Galicia to Gibraltar, and is an important species in sand dune habitats which are highly valuable for conservation 430 purposes (Council Directive 92/43/EEC of 21 May 1992 on the Conservation of Natural 431 Habitats and of Wild Fauna and Flora, 1992). The dynamic nature of these coastal ecosystems 432 provides a vast variety of habitats with unique floristic and animal richness. However, these 433 dune systems face increasing disturbances as they support various economic and leisure 434 activities, associated with the growth of the coastal population (Antunes et al., 2018). Because 435 of habitat loss, C. album has been classified as Vulnerable on the Red List of Andalusia, Spain 436 (Cabezudo et al., 2005). In the Azores Islands, it inhabits volcanic lava and ash fields (de 437 Oliveira & Dale, 2012). The fruits of the Portuguese crowberry are edible, producing bioactive 438 compounds that have been associated with chemoprotective activity and potential healthbenefits (de Oliveira & Dale, 2012; Jacinto et al., 2021; Zunzunegui et al., 2006). 439

Sampling and sample processing: Two male and two female adult plants from the same population were sampled during two field trips to Azeitão, near Arrábida Natural Park. Young expanding leaves and fruits were collected after 48 hours of dark treatment (plant covered with light-opaque paper sheets with a few holes that allowed air flow), according to permits from ICNF (21PTLX00657D). Samples from leaves and fruits for DNA and RNA extraction were flash frozen at -20°C and -80°C, respectively. Voucher specimens were deposited at the Herbarium (LISU 270092) of the MUHNAC, Universidade de Lisboa.

Sequencing: The HMW DNA extraction was very challenging. First, it was performed at the 447 Department of Plant Biology of FCUL using a CTAB protocol (Doyle & Doyle, 1987 later 448 modified by Weising et al., 1994), which resulted in low DNA quality even after purification. 449 450 Taking advantage of the ERGA network, we used a nuclei isolation protocol adapted from the ARIMA-HiC kit, shared by Narjes Yousefi (pers. comm.), a member of ERGA. This resulted in 451 higher quality and quantity of extracted DNA, but it still did not pass the quality control for 452 453 PacBio Hi-Fi library preparation and sequencing at the University of Florence. After several 454 attempts, to prevent DNA degradation during shipping, leaf tissue was directly shipped in dry 455 ice so that DNA extraction could be performed at the University of Florence, where the libraries for Pac-Bio were prepared and sequenced. The assembly of the genome is ongoing, involving 456 ERGA-Portugal members and Biogenome Portugal institutions. 457

Expected results: Building a high-quality annotated reference genome will support ongoing projects on Portuguese crowberry and related plant species from coastal areas, and it will provide the opportunity to explore evolutionary biology questions associated with adaptations to coastal environments under a scenario of global changes. Moreover, the annotated reference genome will allow to uncover the genetic basis of the production of Portuguese crowberry's bioactive compounds associated with health benefits.

464

465 <u>Cave ground beetle - Trechus terceiranus (Machado, 1988)</u>

466 Common name in Portuguese: escaravelho-cavernícola-da-Ilha-Terceira

The cave ground beetle Trechus terceiranus (Coleoptera) is endemic to Terceira Island in 467 the Azores and is restricted to subterranean habitats, including lava tubes. These tiny beetles 468 have up to 4.3 mm in length, do not fly and have several morphological traits associated with 469 470 their life in subterranean habitats (Machado, 1988). This species is listed as Vulnerable in the IUCN Red List (Borges & Amorim, 2018). Interestingly, several species of the genus Trechus 471 occur in the Azores islands, most of which inhabit subterranean habitats (seven species), but 472 two occur in surface habitats. One epigean species, T. terrabravensis Borges, Serrano & 473 474 Amorim, 2004, co-exists on Terceira Island with *T. terceiranus*.

Sampling and sample processing: Six adult individuals were sampled in a subterranean habitat (volcanic pit, lava tube) at 45-70m deep, using pitfall traps. Sampling was done according to permits issued by the Regional Government of the Azores (IRCC 23/2021/DRCTD). Adult individuals were flash frozen with liquid nitrogen. Dried vouchers were stored at the Dalberto Teixeira Pombo entomological collection at the University of the Azores, Terceira (DTP-MF1091).

Sequencing: The frozen whole organisms were shipped in dry ice to the University of Lisbon,
from where they were subsequently sent to Centro Nacional de Análisis Genómico (CNAG) –
Barcelona for RNA extraction and sequencing; and to the University of Florence where DNA
extraction was attempted. Due to constraints to obtain high quality HMW DNA for PacBio,

additional samples were sent in order to repeat the extraction to obtain enough DNA amountand quality for sequencing, which is ongoing.

Expected results: A high-quality reference genome will be crucial to pursue comparative genomics, phylogenetic and population genetic studies to reconstruct the evolutionary history of this group and to implement adequate conservation strategies. Furthermore, it will open the door to detect genes involved in adaptation to subterranean habitats and to determine the genetic architecture of traits related with such habitats (e.g., eye development, pigmentation and biological clock).

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495 Building a national network for biodiversity genomics: Biogenome Portugal

496 Objectives and structure of Biogenome Portugal

497 While ERGA and ERGA-Portugal are a community of researchers, their collaboration in ERGA 498 activities and in the pilot test has played a significant role in forming a national institutional 499 network for biodiversity genomics, called Biogenome Portugal (BGP). This network is being 500 formally established with the objective of fostering scientific collaboration, facilitating the 501 exchange of expertise and infrastructure, organising advanced training programs, promoting Portugal's active participation in international biodiversity genomics initiatives, and 502 503 contributing to knowledge transfer and outreach efforts. The network will comprise members 504 with a diverse range of expertise, encompassing fields such as genomics, ecology, taxonomy, evolution, and more, spanning across various taxonomic groups. 505

506 By bringing together experts from different fields, the BGP network aims to facilitate the 507 exchange of ideas, data, and resources that will lead to the development of new research 508 projects centred on generating and using high quality reference genomes for Portuguese 509 species, and address a plethora of biodiversity applications. Such collaboration also implies 510 sharing institutional genomic and computational infrastructures and articulating with the 511 national network of infrastructures related to BGP's areas of activity. Cooperation among 512 members is crucial for promoting the exchange of expertise, a key objective of the network. 513 By sharing knowledge and skills, the network members will strengthen their collective 514 expertise and coordinate efforts. Activities towards the transfer of skills and expertise include 515 organising advanced training in the field of biodiversity genomics. Through these initiatives, 516 BGP aims to foster the training of specialised human resources able to accompany the fast 517 development of genomic data analysis tools. To broaden its scope, the network intends to facilitate the involvement of individual researchers, even if their institution is not formally 518 519 affiliated with the network.

520 BGP also aims to disseminate the results of its activities to stakeholders and the general 521 public, thus promoting knowledge transfer and outreach. These activities are important not 522 only to enrich scientific literacy about the importance of genomics for biodiversity conservation, 523 but also to promote and encourage the use of the genomic research findings across 524 stakeholders.

525 Finally, BGP also aims to coordinate and synchronise Portuguese participation in global 526 biodiversity genomics initiatives. By reinforcing and sharing the opportunities for institutional 527 representation in internationally funded projects, BGP can promote the contribution of 528 Portuguese expertise to important regional, continental or worldwide initiatives.

529

Articulation with the Portuguese National Strategy for the Conservation of Nature and Biodiversity 2030

532 The Portuguese National Strategy for the Conservation of Nature and Biodiversity (ENCNB 533 2030; https://dre.pt/dre/detalhe/resolucao-conselho-ministros/55-2018-115226936) is the main legal Portuguese document that is used to frame all national policies until 2030. It 534 535 recognizes the importance of Portugal's national biodiversity at both the European and global 536 scale and acknowledges how the nation's biogeographic specificities have led to high levels of endemic and relict species, each with a unique evolutionary history and genetic 537 composition, which is important to preserve. The national strategy has several main 538 objectives, including planning and executing action promoting the conservation and recovery 539

of species and habitats at the national level and promoting the conservation of plant and animals' genetic diversity. These objectives align with the aims of the nascent BGP. The network can provide the knowledge-based capacities necessary for the conservation and sustainable use of the national marine, terrestrial and freshwater genetic resources. BGP can reinforce a science-based approach for species conservation action plans and guide complex conservation strategies, including *in situ* and *ex situ* conservation.

546

547 <u>Fostering national research infrastructures</u>

548 In 2014, the national public agency for science, technology and innovation (Fundação para a Ciência e a Tecnologia, FCT) created a National Roadmap for Research Infrastructures of 549 Strategic Interest. This roadmap aimed at promoting cooperation, excellence and 550 internationalisation to strengthen national scientific infrastructures. A total of 56 Research 551 552 Infrastructures (RIs) have been supported during the first funding cycle in key areas, such as the Environment, Health and Food, Social and Cultural Innovation, among others. BGP 553 identified several RIs of interest, intersecting its area of intervention. Among these are 554 GenomePT (the National Laboratory for Genome Sequencing and Analysis), Biodata.pt 555 556 (ELIXIR PT - Portuguese Distributed Infrastructure for Biological Data), PORBIOTA (Portuguese E-Infrastructure for Information and Research on Biodiversity), EMBRC.PT 557 (European Marine Biological Resource Centre – Portugal), and PRISC (Portuguese Research 558 Infrastructure for Scientific Collections). Future action implies fostering the establishment of 559 strategic cooperation with the active infrastructures, which can be facilitated by the current 560 presence of several institutions of BGP in the RIs, pending the continuation of the current 561 roadmap. 562

563

Engaging the community: establishing training programs, supporting the generation of additional reference genomes and expanding funding opportunities

566 The analysis of reference genomes at scale requires standardised procedures for the 567 sampling and storing of the biological material, as well as sequencing and analysis of the 568 genomic data. Researchers working to analyse and utilise these cutting-edge genomic resources need specific training, to maintain quality standards and streamline procedures. 569 570 The development of bioinformatic pipelines by the larger ERGA community using open-source 571 platforms such as Galaxy (The Galaxy Community, 2022) allows quick implementation of 572 analyses across research centres. Within the BGP network, an online training programme is 573 being implemented to cover different topics across pipelines of genomic analyses, such as genome and transcriptome assembly, annotation, comparative and population genomics. This 574 575 programme aims to expand and facilitate training to enhance expertise in genomic analyses 576 across the Portuguese research community. The first introductory course to Galaxy and VGP 577 assembly pipeline took place in the first semester of 2023.

Further expertise sharing will come from the expansion of reference genomes projects under 578 the BGP umbrella, which meets the technical and quality standards set by ERGA. For 579 580 example, two reference genomes of Anthozoan species belonging to octocorals are being generated under the BGP initiative: the pink sea fan (Eunicella verrucosa) and the dead man's 581 fingers (Alcyonium digitatum). Octocoral genomes remain poorly studied, with only ~10 582 reference genomes available from the more than 3,000 described species. These two species 583 584 belong to a pool of octocorals for which the sequencing of reference genomes was supported by EASI-Genomics (H2020 824110; Project ID 10240, CoGeCo). The two species are widely 585 586 distributed along the Portuguese coast in sublittoral rocky habitats, and sequencing, genome 587 assembly and annotations are underway.

Attracting national and international funding will be crucial to support the implementation of BGP's activities. Working as a network will lead to stronger research proposals, more successful grant applications, and increase collective participation in international consortia to foster biodiversity genomics cutting-edge research and knowledge transfer. This strategy will allow consolidating, strengthening and expanding the BGP network and its impact on fundamental and applied science and innovation.

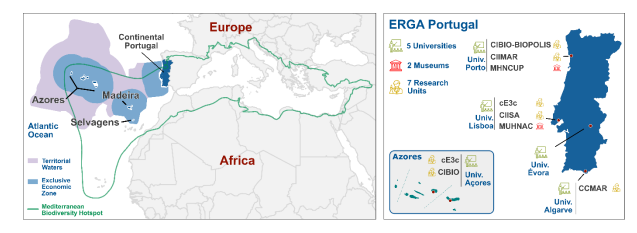
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595 <u>Producing high quality chromosome-level assemblies of Portuguese species</u>

596 The alignment of BGP with ERGA can foster the multiplication of opportunities for sequencing projects for Portuguese biodiversity. This aligns with the aim of ERGA of "propagating 597 guidelines for state-of-the-art genome establishment through training and knowledge transfer". 598 Produced genomes can therefore take advantage of the standard sampling and molecular 599 protocols and bioinformatics pipelines for sequencing, genome assembly and annotation 600 established by ERGA, and be included under the ERGA umbrella. While promoting the 601 602 visibility and accessibility of the genomes to the ERGA network and beyond, those genomes 603 will contribute to the Reference Genome Atlas of European biodiversity. Such parallel initiatives under the BGP and ERGA umbrella, coordinated with ERGA projects, will expedite 604 605 the production of a genomic inventory for all Portuguese eukaryotic species. Those efforts are invaluable to enable the use of genomic applications to preserve and protect the country's rich 606 607 biodiversity.

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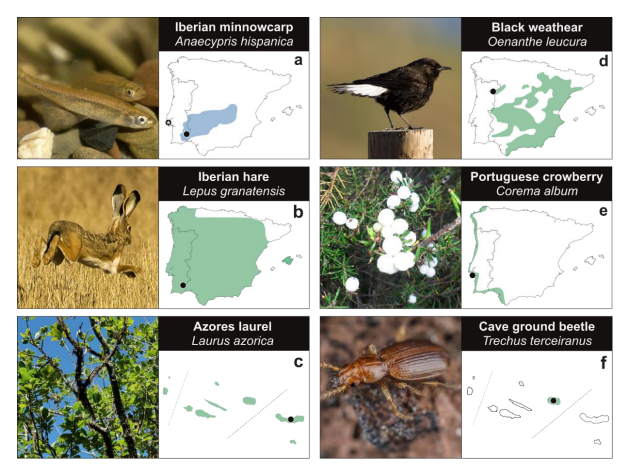
610 Figures



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<sup>Fig.1 - a) Spatial context of the Portuguese territory, including the two Autonomous Regions (Azores
and Madeira) b) Distribution of the Portugal-based members of ERGA (ERGA-Portugal), including their
host institutions (May 31, 2023).</sup>



616

617 Fig.2 - Approximate distributions and sampling localities of ERGA-Portugal species sequenced in ERGA's pilot effort: a) Iberian minnowcarp - Anaecypris hispanica; b) Iberian hare - Lepus granatensis; 618 c) Azores laurel - Laurus azorica; d) Black wheatear - Oenanthe leucura; e) Portuguese crowberry -619 620 Corema album; f) Cave ground beetle - Trechus terceiranus. Picture credits: a) Carlos Carrapato; b) 621 Pedro Moreira; c) Mónica Moura; d) Ricardo Jorge Lopes; e) Manuela Sim-Sim; f) Javier Torrent 622 (Azorean Biodiversity Group). Species distributions coloured according to terrestrial (green) or aquatic 623 (blue) territories. Sampling localities marked with a black dot • except for the Iberian minnocarp (a) that is double marked, as it was sampled in the Aquário Vasco da Gama – Lisbon ©, but it is originally 624 625 from a population of River Chança . Distributions adjusted from the IUCN Red List of Threatened 626 Species Website (https://www.iucnredlist.org), except for the Portuguese crowberry, which was based 627 on information from the Anthos geographical information system for Spanish Flora (www.anthos.es). 628

629

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