1	How spiders make their eyes: Systemic paralogy and function of retinal determination
2	network homologs in arachnids
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## 16

## 17 Abstract

18

19 Arachnids are important components of cave ecosystems and display many examples of 20 troglomorphisms, such as blindness, depigmentation, and elongate appendages. Little is 21 known about how the eves of arachnids are specified genetically, let alone the mechanisms 22 for eye reduction and loss in troglomorphic arachnids. Additionally, paralogy of Retinal 23 Determination Gene Network (RDGN) homologs in spiders has convoluted functional 24 inferences extrapolated from single-copy homologs in pancrustacean models. Here, we 25 investigated a sister species pair of Israeli cave whip spiders (Arachnopulmonata, 26 Amblypygi, Charinus) of which one species has reduced eyes. We generated the first embryonic transcriptomes for Amblypygi, and discovered that several RDGN homologs 27 28 exhibit duplications. We show that paralogy of RDGN homologs is systemic across 29 arachnopulmonates (arachnid orders that bear book lungs), rather than being a spider-specific 30 phenomenon. A differential gene expression (DGE) analysis comparing the expression of 31 RDGN genes in field-collected embryos of both species identified candidate RDGN genes 32 involved in the formation and reduction of eyes in whip spiders. To ground bioinformatic 33 inference of expression patterns with functional experiments, we interrogated the function of 34 three candidate RDGN genes identified from DGE in a spider, using RNAi in the spider 35 Parasteatoda tepidariorum. We provide functional evidence that one of these paralogs, sine 36 oculis/Six1 A (soA), is necessary for the development of all arachnid eye types. Our results 37 support the conservation of at least one RDGN component across Arthropoda and establish a 38 framework for investigating the role of gene duplications in arachnid eye diversity. 39 40 Keywords: cave blindness | *sine oculis* | *Six1* | *Parasteatoda tepidariorum* | Amblypygi |

- 41 RNAi
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## 43 Introduction

44

45 Cave habitats offer apt systems for investigating the genetic basis of morphological 46 convergence because communities of these habitats are similarly shaped by environmental 47 pressures, such as absence of light and diminished primary productivity (Howarth, 1993; 48 Juan, Guzik, Jaume, & Cooper, 2010). Troglobites, species exclusive to cave environments 49 and adapted to life in the dark, exhibit a suite of characteristics common to cave systems 50 around the world, such as reduction or complete loss of eyes, depigmentation, elongation of 51 appendages and sensory structures, and decreased metabolic activity (Jemec, Škufca, 52 Prevorčnik, Fišer, & Zidar, 2017; Protas & Jeffery, 2012; Riddle et al., 2018). Previous work 53 has shown that troglomorphism can evolve over short time spans (<50 kyr) despite gene flow 54 (Bradic, Teotónio, & Borowsky, 2013; Coghill, Darrin Hulsey, Chaves-Campos, García de 55 Leon, & Johnson, 2014; Herman et al., 2018) and that parallel evolution of troglomorphic 56 traits (e.g., depigmentation; eye loss) in independent populations can involve the same 57 genetic locus (Protas et al., 2005; Protas, Trontelj, & Patel, 2011; Re et al., 2018). 58 Troglomorphism and troglobitic fauna have been analyzed across numerous taxonomic 59 groups with respect to systematics and population genetics. However, one component of the 60 troglobitic fauna that remains poorly understood is cave arachnids. Most orders of Arachnida 61 are prone to nocturnal life history and some orders broadly exhibit troglophily; in fact, 62 troglobitic species are known from all the extant terrestrial arachnid orders except Solifugae and Uropygi (Cruz-López, Proud, & Pérez-González, 2016; Esposito et al., 2015; Harvey, 63 64 2002; 2007; Hedin & Thomas, 2010; Mammola, Mazzuca, Pantini, Isaia, & Arnedo, 2017; 65 Miranda, Aharon, Gavish-Regev, Giupponi, & Wizen, 2016; Santibáñez López, Francke, & 66 Prendini, 2014; Smrž, Kováč, Mikeš, & Lukešová, 2013). In addition to eye and pigment 67 loss, troglomorphism in arachnids manifests in the form of compensatory elongation of 68 walking legs and palps, appendages which harbor sensory structures in this group 69 (Derkarabetian, Steinmann, & Hedin, 2010; Mammola & Isaia, 2017; Mammola et al., 70 2018a; Mammola, Cardoso, Ribera, Pavlek, & Isaia, 2018b). 71 Thorough understanding of the developmental genetic basis for the evolution of 72 troglomorphic traits has been largely spearheaded in two model systems: the Mexican cave 73 fish Astyanax mexicanus (Bradic et al., 2013; Coghill et al., 2014; Herman et al., 2018; 74 Porter, Dittmar, & Pérez-Losada, 2007; Protas et al., 2005; Protas & Jeffery, 2012) and the 75 cave isopod Asellus aquaticus (Jemec et al., 2017; Re et al., 2018; Stahl et al., 2015). Both

76 model systems have more than one hypogean population, can be maintained in laboratories,

77 and are amenable to approaches such as genetic crosses and quantitative trait locus mapping. 78 The advent of short read sequencing technology in tandem with experimental approaches has 79 transformed the potential to triangulate regulatory differences between hypogean 80 (subterranean) and epigean (surface-dwelling) lineages (Protas et al., 2005; Re et al., 2018; 81 Riddle et al., 2018; Stahl et al., 2015), and to study a broader range of cave taxa. 82 Among arthropods, work on the isopod A. aquaticus in particular has made significant 83 advances in the identification of loci regulating pigmentation and size of arthropod eves 84 (Protas et al., 2011; Re et al., 2018), complementing forward and reverse genetic screening 85 approaches in other pancrustacean models (e.g., Drosophila melanogaster, Tribolium castaneum, and Gryllus bimaculatus) (Cagan, 2009; Kumar, 2009; Takagi et al., 2012; 86 87 ZarinKamar et al., 2011). However, developmental and genetic insights into the evolution of 88 blindness illuminated by A. aquaticus and other pancrustacean models are not directly 89 transferable to Arachnida for two reasons. First, the eyes of arachnids are structurally and 90 functionally different from those of pancrustaceans. Typically, the main eyes of adult 91 Pancrustacea (e.g., A. aquaticus) are a pair of faceted (or apposition) eyes, which are 92 composed of many subunits of ommatidia. In addition, adult Pancrustacea have small median 93 ocelli (typically three in holometabolous insects), often located medially and at the top of the

94 head.

95 By contrast, extant arachnids lack ommatidia and typically have multiple pairs of eyes 96 arranged along the frontal carapace. All arachnid eyes are simple-lens eyes or ocelli; each eye 97 has a single cuticular lens, below which are a vitreous body and visual cells. The retina is 98 composed of the visual cells and pigment cells. These eyes are divided in two types, namely 99 the principal eyes and the secondary eyes (Foelix, 2011; Land, 1985). Principal and 100 secondary eyes differ in the orientation of their retina (Homann, 1971): the principal eyes are 101 of the everted type, with the visual cells lying distally, and lack a reflective layer; the 102 secondary eyes are inverted, with the light-sensitive rhabdomeres pointing away from 103 incoming light (analogous to vertebrate eyes). All secondary eyes possess a reflective layer of 104 crystalline deposits called a tapetum, which is responsible for the "eye shine" of spiders. The 105 principal eyes are the median eyes (ME, also known as anterior medium eyes). The 106 secondary eyes comprise the anterior lateral eyes (ALE), posterior lateral eyes (PLE), and medium lateral eyes (MLE; also known as posterior medium eyes) (Fig. 1A) (Foelix, 2011; 107 108 Land, 1985) (nomenclature used here follows Schomburg et al 2015). Certain orders and 109 suborders of arachnids have lost one type of eye altogether, with the homology of eyes

110 clarified by the fossil record and embryology (Foelix, 2011; Garwood, Sharma, Dunlop, &

111 Giribet, 2014; Morehouse, Buschbeck, Zurek, Steck, & Porter, 2017).

112 The second concern in extending the model derived from pancrustaceans is that a subset 113 of Arachnida exhibits an ancient shared genome duplication, resulting in numerous paralogs 114 of developmental patterning genes. Recent phylogenetic and comparative genomic works on 115 Arachnida have shown that Arachnopulmonata (Ballesteros & Sharma, 2019; Ballesteros, 116 Santibáñez López, Kováč, Gavish-Regev, & Sharma, 2019; Sharma, Kaluziak, Pérez-Porro, 117 González, Hormiga, et al., 2014a), the clade of arachnids that bear book lungs (e.g., spiders, 118 scorpions, whip spiders), retain duplicates of many key transcription factors, such as 119 homeobox genes, often in conserved syntenic blocks (Leite et al., 2018; Schwager et al., 120 2017; Sharma, Santiago, González-Santillán, Monod, & Wheeler, 2015a; Sharma, Schwager, 121 Extavour, & Wheeler, 2014b). Many of the ensuing paralogs exhibit non-overlapping 122 expression patterns and a small number have been shown to have subdivided the ancestral 123 gene function (subfunctionalization) or acquired new functions (neofunctionalization) (Leite 124 et al., 2018; Paese, Leite, Schönauer, McGregor, & Russell, 2018; Turetzek, Pechmann, 125 Schomburg, Schneider, & Prpic, 2015).

126 While comparatively little is known about the genetics of arachnid eye development, gene 127 expression surveys of insect retinal determination gene network (RDGN) homologs of two 128 spiders (Cupiennius salei and Parasteatoda tepidariorum) have shown that this phenomenon 129 extends to the formation of spider eyes as well (Samadi, Schmid, & Eriksson, 2015; Schomburg et al., 2015). Different paralog pairs (orthologs of *Pax6*, *Six1*, *Six3*, *eves absent*, 130 131 atonal, dachshund and orthodenticle) exhibit non-overlapping expression boundaries in the 132 developing eye fields, resulting in different combinations of transcription factor expression in 133 the eye pairs (Samadi et al., 2015; Schomburg et al., 2015). While these expression patterns 134 offer a potentially elegant solution to the differentiation of spider eye pairs, only a few 135 studies with the spider *P. tepidariorum* have attempted to experimentally test the role of these 136 genes in the formation of arachnid eyes. Ptep-orthodenticle-1 maternal RNA interference (RNAi) knockdown results in a range of anterior defects, including complete loss of the head, 137 138 which precluded assessment of a role in the formation of the eyes (Pechmann, McGregor, 139 Schwager, Feitosa, & Damen, 2009). Ptep-dac2 RNAi knockdown results in appendage 140 segment defects, but no eye patterning defects were reported by the authors (Turetzek et al., 2015). More recently, a functional interrogation of *Ptep-Six3* paralogs, focused on labrum 141 142 development, reported no discernible morphological phenotype, despite a lower hatching rate 143 than controls and disruption of a downstream target with a labral expression domain

144 (Schacht, Schomburg, & Bucher, 2020). Thus, gene expression patterns of duplicated RDGN 145 paralogs have never been linked to eye-related phenotypic outcomes in any 146 arachnopulmonate model. Similarly, the functions of the single-copy orthologs of RDGN 147 genes in groups like mites (Grbić et al., 2007; Telford & Thomas, 1998), ticks (Santos et al., 148 2013), and harvestmen (Garwood et al., 2014; Sharma, Schwager, Giribet, Jockusch, & 149 Extavour, 2013; Sharma, Tarazona, Lopez, Schwager, Cohn, Wheeler, et al., 2015b) are 150 entirely unexplored, in one case because an otherwise tractable arachnid species lacks eves 151 altogether (the mite Archegozetes longisetosus (Barnett & Thomas, 2012; 2013a; 2013b; 152 Telford & Thomas, 1998). Investigating the evolution of eye loss in arachnids thus has the potential to elucidate 153 154 simultaneously (1) the morphogenesis of a poorly understood subset of metazoan eyes 155 (Foelix, 2011; Morehouse et al., 2017), (2) developmental mechanisms underlying a 156 convergent trait (i.e., eye loss in caves) in phylogenetically distant arthropod groups (Protas 157 & Jeffery, 2012; Re et al., 2018), (3) shared programs in eye development common to Arthropoda (through comparisons with pancrustacean datasets) (Cagan, 2009; Stahl et al., 158 159 2015; Takagi et al., 2012; ZarinKamar et al., 2011), and (4) the role of ancient gene 160 duplicates in establishing the diversity of eyes in arachnopulmonates (Leite et al., 2018; 161 Samadi et al., 2015; Schomburg et al., 2015). 162 As first steps toward these goals, we first developed transcriptomic resources for a sister 163 species pair of cave-dwelling *Charinus* whip spiders, wherein one species exhibits typical eye morphology and the other highly reduced eyes (a troglobitic condition). We applied a 164 165 differential gene expression (DGE) analysis to these datasets to investigate whether candidate 166 RDGN genes with known expression patterns in model spider species (C. salei, P. 167 *tepidariorum*) exhibit differential expression in non-spider arachnopulmonates, as a function 168 of both eye condition and developmental stage. To link bioinformatic inference of expression 169 patterns with functional outcomes, we interrogated the function of three candidate RDGN 170 genes identified from DGE in a model arachnopulmonate, using RNAi in the spider P. tepidariorum, which exhibits the same number and types of eyes as whip spiders. We provide 171 172 functional evidence that one of these candidates, *sine oculis/Six1*, is necessary for the 173 development of all spider eye types. 174 175 **Results** 

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177 Charinus ioanniticus and Charinus israelensis embryonic transcriptomes

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179 As an empirical case of closely related, non-spider arachnopulmonate sister species pair 180 that constitutes one epigean and one troglobitic species, we selected the whip spider species 181 Charinus ioanniticus and C. israelensis (Fig. 1 B-C). Whip spiders, arachnopulmonates of 182 the order Amblypygi, are commonly found in cave habitats ranging from rain forests, 183 savannahs and deserts (Weygoldt, 2000). The recently described troglobitic species Charinus 184 israelensis (reduced-eyes) occurs in close proximity to its congener Charinus ioanniticus 185 (normal-eyes) in caves in the Galilee, northern Israel (Miranda et al. 2016). Given that the formation of Levantine cave refuges is considerably recent, C. israelensis and C. ioanniticus 186 are likely sister species with a small time of divergence, an inference supported by their 187 188 similar morphology (Miranda et al 2016). We collected ovigerous females from both species 189 in caves in Israel and extracted RNA from embryos (SI Appendix, Table S1). Embryos of 190 whip-spiders (*Phrynus marginemaculatus*) achieve a deutembryo stage around 20-25 days 191 after egg laying (dAEL), a stage where most external features of the embryo, such as 192 tagmosis and appendages are fully formed, but not the eyes (Weygoldt, 1975). The 193 deutembryo hatches from the egg membrane inside the broodsac carried by the mother, but 194 remains in this stage relatively unchanged for around 70 days. The eyes begin to form around 195 50 dAEL, but the eye spots become externally visible and pigmented only close to hatching 196 (90 dAEL) (Weygoldt, 1975). 197 For *de novo* assembly of the embryonic transcriptomes of *C. ioanniticus* and *C.* 198 israelensis, we extracted RNA from all embryonic deutembryo stages collected in the field 199 (see Supplementary Information; table 1 for localities and sample explanations). Assemblies 200 include two deutembryo stages before eyespot formation and one deutembryo stage bearing

201 eyespots for *C. ioanniticus*; and two early deutembryo stages for *C. israelensis* (SI Appendix,
202 Fig. S1).

The assembly of *C. ioanniticus* reads resulted in 219,797 transcripts composed of 143,282,365 bp with and N50 of 1122 bp (more than 50% of transcripts are 1122 bp or longer) (SI Appendix, Table S2). Universal single copy ortholog benchmarking with BUSCO v3.0 (Waterhouse et al., 2017) indicated 93.8% completeness, with 5.7% of BUSCO genes exhibiting duplication.

The assembly *C. israelensis* resulted in a higher number of transcripts: 663,281 transcripts composed of 230,044,656 bp and with N50 of 1045 bp. The BUSCO analysis shows 95.2% completeness, which is similar to the value for *C. ioanniticus* assembly.

211

## 212 RDGN gene duplication in *Charinus* whip spiders

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214 Amblypygi is inferred to be nested stably in Arachnopulmonata, the clade of arachnids 215 that bear book lungs (Ballesteros & Sharma, 2019; Giribet, 2018; Lozano-Fernandez et al., 216 2019; Rota-Stabelli et al., 2010; Sharma, Kaluziak, Pérez-Porro, González, Hormiga, et al., 217 2014a). Recent evidence suggests that the common ancestor of arachnopulmonates has 218 undergone a whole- or partial-genome duplication affecting large gene families, such as 219 homeobox genes (Leite et al., 2018; Schwager et al., 2017; Sharma, Santiago, González-220 Santillán, Monod, & Wheeler, 2015a). The stable phylogenetic position of Amblypygi in 221 Arachnopulmonata predicts that genes in RDGN that are duplicated in spiders, should also be 222 duplicated in *Charinus* whip spiders. To test this hypothesis, we performed phylogenetically-223 informed orthology searches on the newly assembled embryonic transcriptomes of both 224 *Charinus* species, and conducted phylogenetic analysis with orthologs across selected 225 arthropod species. We discovered that homologs of atonal, Pax6, dachshund, sine oculis 226 (Six1), Optix (Six3), and orthodenticle are duplicated in Charinus, whereas eyegone and eyes 227 *absent* occur as single-copy orthologs (these latter two also occurring single-copy in spiders) 228 (Fig. 2).

229 atonal: The atonal gene tree showed poor resolution (SI Appendix, Fig. S2), hampering 230 unambiguous assignment of the whip spider genes to *atonal* copies previously annotated in 231 spiders (Samadi et al., 2015; Schwager et al., 2017). D. melanogaster copies of atonal and 232 *amos* clustered together forming a clade with other pancrustacean and myriapod sequences, 233 suggesting these paralogs are restricted to Mandibulata. The fruit fly *cousin of atonal (cato)* 234 formed a clade including the *Cupiennius salei* sequence of *atonalB* whereas the second copy 235 of C. salei, atonalA, is found in an independent clade with only arachnid sequences. It is in 236 this later clade that the only sequences of *Charinus* related to *atonal* are found, in turn 237 forming two separate clades with clear amino acid differences between these copies (SI 238 Appendix, Dataset S1; atonal alignment). Herein, these copies are labeled atonalA (atoA) and 239 atonalB (atoB). Note that the reference genomic sequences, annotated as "atonal like 240 homolog 8 like" (Ptep XP 0159181091), is found orthologous to the gene net in D. 241 melanogaster.

*Pax6*: In *D. melanogaster*, there are two paralogous copies of the vertebrate *Pax6*, *eyeless*and *twin of eyeless*. This duplication seems to be shared across all arthropods and both *Pax6*copies have been characterized in spiders (Samadi et al., 2015; Schomburg et al., 2015). The
gene tree of *Pax6* homologues clearly identified a clade for *toy* including chelicerate and

246 mandibulate copies, but no *Charinus* sequences are found in this clade (SI Appendix, Fig. 247 S3). The sister clade (eyeless) consists only of pancrustacean sequences whereas the 248 chelicerate copies, previously annotated as *eveless* orthologs, are found in a separate clade. 249 Among these, two distinct genes, herein dubbed *Pax6A* and *Pax6B*, are present in both 250 Charinus species. Sequence similarity searches (blastp) of both Pax6A and Pax6B against the 251 genome of *Drosophila melanogaster* points to *Dmel-toy* as the best hit, followed by *Dmel-ey*. 252 Therefore, although the homology of these copies with *Dmel-ey/toy* is evident, it is not trivial 253 to assign these to either of these genes or if these represent taxon-restricted duplicates of 254 eyeless. eyegone/twin of eyegone: These members of the Pax gene family are paralogous in D. 255 256 melanogaster but occur as single copy in arachnids. Single copy orthologs of eyg/toe are 257 present in the two target Amblypygi species (SI Appendix, Fig. S3). 258 dachshund: Spiders and scorpions have two paralogous copies of dachshund (Nolan, 259 Santibáñez López, & Sharma, 2020; Turetzek et al., 2015). Two copies are present in the 260 transcriptomes of both Charinus species and are here termed dacA and dacB (SI Appendix, 261 Fig. S4). The *C. israelensis dacB* is assembled in two different gene fragments that overlap 262 by three amino acids (SI Appendix, Fig. S4; see *dachshund* alignment in SI Appendix 263 Dataset S1). The C. ioanniticus dacA copy is also assembled as two different gene fragments 264 with little sequence overlap but being part of the *dacA* clade (SI Appendix, Fig. S4). 265 eyes absent: This single-copy orthologs are found in arthropods and arachnids alike and is represented in both *Charinus* species. The association of transcript to this gene is 266 267 unambiguous for both amblypigid species (SI Appendix, Fig. S5). 268 orthodenticle: As with spiders, there are two copies homologous to Dmel-otd in 269 *Charinus.* The resolution of the gene tree is poor and does not allow uncontroversial 270 association to spider orthologs (SI Appendix, Fig. S6). Charinus copies are termed otdA and 271 otdB.

272 *Optix*: There are two very similar copies of *Optix* in *C. israelensis* and one in *C.* 

273 *ioanniticus* (SI Appendix, Fig. S7). The *C. ioanniticus* copy is termed *OptixA*. The two copies

of *C. israelensis* show very conserved amino acid sequences but clear nucleotide differences.

Although the gene tree with the reference genome shows them more closely allied to one of

the spider paralogous copies of *Optix* (Ptep NP 00130752.1), a reduced analysis including

- 277 Cupiennius salei and P. tepidariorum copies, suggests that the Charinus copies are
- 278 independent duplications. Here the two whip spider copies are dubbed *OptixA* and *OptixB* but
- they should not be considered orthologous to the spider *OptixA/B*.

280 sine oculis: Two copies of sine oculis are found in C. israelensis and one in C. 281 ioanniticus. Both copies are nested in a clade with Ptep-soA (SI Appendix, Fig. S8). These 282 are herein dubbed as *soA* and *soB* given that orthology with either spider copy is unclear. 283 284 RDGN genes in whip spider eye formation: comparing early and late stages of C. ioanniticus 285 286 The expression of paralog pairs of *Pax6*, sine oculis, Optix, eves absent, atonal, 287 dachshund and orthodenticle in the developing eyes of the spiders (Samadi et al., 2015; Schomburg et al., 2015), and the occurrence of the same paralogs in *Charinus* whip spiders, 288 289 suggest that these genes may also be involved in the formation of eyes in whip spiders. We 290 investigated this idea by comparing the expression levels of these RDGN genes in the stages 291 before eye-spot formation versus a stage after eye-spot formation in the eye-bearing whip 292 spider Charinus ioanniticus (henceforth "Comparison 1"; Fig. 3A). 293 We mapped reads of both treatments to the reference transcriptome of C. ioanniticus 294 using the quasi-alignment software Salmon v. 1.1.0 (Patro, Duggal, Love, Irizarry, & 295 Kingsford, 2017) and conducted a differential gene expression analysis of Comparison 1 296 using DESeq2 v 1.24.0 (Love, Huber, & Anders, 2014) (SI Appendix, Fig. S9). These 297 comparisons showed that Cioa-dacA, Cioa-otdA, Ciao-eya and Cioa-soA are significantly 298 over-expressed ( $p_{adj} < 0.05$ ) in the eyespot stage in comparison with the stage before eyespot 299 formation (Fig. 3A). While we cannot rule out that the differences in gene expression are due 300 to other developmental differences between the two stages sequenced, these results 301 highlighted these four RDGN genes as promising candidates involved in the formation of 302 eyes in whip spiders. 303 304 RDGN genes in whip spider eye reduction: comparing C. ioanniticus and C. israelensis 305 306 Blindness in adults of the model cave fish Astyanax mexicanus is a result of an embryonic 307 process in which the rudimentary eye of the embryo is induced to degenerate by signals

308 emitted from the lens tissue (Jeffery, 2009). Both early and late expression of RDGN genes,

309 such as Pax6, are responsible for the reduction of eyes in fish from cave populations

310 (Jeffery, 2009; Strickler, Yamamoto, & Jeffery, 2001). Likewise, in the isopod crustacean

311 Asellus aquaticus cave blindness has a strong genetic component and mechanisms of eye

312 reduction also act at embryonic stages (Mojaddidi, Fernandez, Erickson, & Protas, 2018;

313 Protas et al., 2011). The embryonic development of the reduced-eyes whip spider *C*.

314 *israelensis* has not been explored to date, but we expect that reduction of eyes results from 315 changes in embryonic gene expression during the deutembryo stage (Weygoldt, 1975). We 316 investigated this possibility by quantifying the relative gene expression of RDGN genes in 317 comparable embryonic stages of C. israelensis (reduced eyes) and C. ioanniticus (normal 318 eves) embryos before eye-spot formation (SI Appendix Table S1; Figure S1). Using the DGE 319 approach from Comparison 1, we conducted a heterospecific analysis using as the reference 320 either the C. israelensis transcriptome (henceforth "Comparison 2.1") or the C. ioanniticus 321 transcriptome (henceforth "Comparison 2.2"). 322 Both analyses are anchored on the premise that a hybrid mapping between the sister 323 species is possible given the recent divergence between them. The mapping rate of the C.

*ioanniticus* reads was similar regardless of the reference species, (96.74% and 96.59%

325 respectively for *C. ioanniticus* and *C. israelensis*). In the case of the reads from *C. israelensis* 

embryos, mapping rate to the conspecific (96.8%) transcriptome was higher than when

327 mapping against *C. ioanniticus* (82.45%). The similar mapping rate of *C. ioanniticus* reads

328 suggests that the two whip spiders are sufficiently closely related to generate interspecific

329 comparisons of gene expression. Comparisons 2.1 and 2.2 yielded similar results with respect

to the direction of differentially expressed RDGN genes (Fig. 3B–C). In comparison 2.1,

331 Pax6A, OptixA and OptixB are significantly over-expressed in the normal-eyes species, with

332 expression levels at least 4 times higher than in the reduced-eyes species ( $log_2FC > 2$ ;  $p_{adj} <$ 

333 0.05) (Fig. 3B; SI Appendix, Fig. S10). In comparison 2.2, Pax6A and OptixA are also over-

expressed in *C. ioanniticus* ( $p_{adj} < 0.05$ ), and so is *eyes absent* ( $p_{adj} < 0.05$ ; Fig. 3C). In

335 comparison 2.2, *orthodenticle-B* appears under-expressed in the normal-eyes species ( $p_{adj} <$ 

336 0.05 (Fig. 3C; SI Appendix, Fig. S11). We note that the magnitude of log<sub>2</sub>FC and

337 significance values differed considerably between analysis. Nonetheless, *Pax6A* and *OptixA* 

338 were consistently over expressed in the normal-eyes species, highlighting these two genes as

339 promising candidates involved in the reduction of eyes in *Charinus israelesis*.

340

*sine oculis* is necessary for principal and secondary eye development in a model

342 arachnopulmonate

343

344 Our bioinformatic analysis in the whip spider system suggested that *eyes absent* and

345 paralogs of *sine oculis*, *orthodenticle*, and *dachshund* may be involved in the normal

346 formation of eyes in *C. ioanniticus* (Comparison 1). We also found evidence that *Pax6* and a

347 paralog of *Optix* may be involved in the reduction of eyes in the cave whip spider *C*.

348 *israelensis*. To link bioinformatic reconstructions of gene expression with functional

349 outcomes, we interrogated the function of RDGN genes using parental RNA interference

350 (RNAi) in the spider Parasteatoda tepidariorum. We selected Ptep-soA (Ptep-sol sensu

351 Schomburg et al. 2015), *Ptep-otdB* (*Ptep-otd2 sensu* Schomburg et al. 2015) and *Ptep-OptixB* 

352 (*Ptep Six3.2 sensu* Schomburg et al. 2015). In *P. tepidariorum*, these genes are known to be

353 expressed in all eye types, in the median eyes only, and in the lateral eyes, respectively (Fig.

354 1D) (Schomburg et al., 2015).

Early expression of *Ptep-soA* is detected in lateral domains of the head lobes (stage 10)

356 corresponding to the principal and secondary eyes, and continues until the pre-hatching stage

357 14 (Schomburg et al., 2015). Expression of *Ptep-soA* on wild type stage 14.1 embryos is

bilaterally symmetrical on all eyes and uniformly strong (Fig. 4A–B). By stage 14.2, it

remains strong on the principal eyes but it is stronger at the periphery of the secondary eye spots (Fig. 4A, C).

*P. tepidariorum* hatchlings, or postembryos, initially have no externally visible lenses and pigment. The red pigment and lenses of all eyes, and the reflective tapetum of the lateral eyes, become progressively recognizable in the 48 hours (at 26°C) until the animal molts into the first instar with fully formed eyes (SI Appendix, Video S1) (see also Mittmann & Wolff, 2012). We fixed embryos from *Ptep-soA* dsRNA-injected and dH2O-injected treatments between 24h-48h, which encompasses stages where the eyes of postembryos are already recognizable until the first instar.

Negative control experiments (dH<sub>2</sub>O-injected females) yielded postembryos with eve 368 369 morphology indistinguishable from wild type animals: the median eyes (ME; principal eyes) 370 have an inferior semi-lunar ring of red pigment and lack the tapetum; and all pairs of lateral 371 eyes (secondary eyes) have the canoe-shaped tapetum type (Foelix, 2011; Land, 1985), which 372 is split in the middle and surrounded by red pigment (Fig. 5A; panel 1). We observed 373 misshaped tapeta on the lateral eyes of some postembryos on the earlier side of the 374 developmental spectrum of fixed animals, but that was never observed on postembryos close 375 to molting or first instars (SI Appendix, Fig. S12). It is unclear if this reflects a natural 376 variation of early developing tapetum or an artifact of sample preparation.

Embryos from *Ptep-soA* dsRNA-injected females are also able to hatch into postembryos and continue molting to adulthood (SI Appendix, Video S2). However, a subset of the embryos of dsRNA-injected treatment (9.5%; n=195/2049) exhibits a spectrum of eye defects that was not observed on the controls (Fig. 5A–B; SI Appendix, Fig. S13). The defects occurred on all eyes, namely medium eyes (ME), anterior lateral eyes (ALE), posterior lateral 382 eves (PLE), and medium lateral eves (MLE) (Fig. 5A). Affected medium eves have reduced 383 pigmentation or complete absence (Fig. 5A, panels 2-6), while lateral eyes also exhibited 384 defects of the tapetum or complete absence of the eye (Fig. 5A, panels 4–6). 385 We selected a subset of the knockdown postembryos initially scored as having any eye 386 defect (n=48) for quantifying the degree of effect per eye type, and the proportion of 387 symmetrical and mosaic eve phenotypes in our sample. Medium eves are affected in almost 388 all cases (97%), whereas the three lateral eye types were similarly lowly affected (MLE: 389 14%; PLE: 8%; ALE: 10%) (Fig. 5C; SI Appendix, Fig. S12; detailed scoring criteria in 390 Material and Methods). The majority of defective eyes are mosaics, meaning that a given eye 391 pair is affected only on one side of the animal (Fig. 5C; SI Appendix, Fig. S12). 392 Parental RNAi against *Ptep-soA* did not completely abolish its expression, as detected by 393 in situ hybridization (Fig. 4D; see Material and Methods). Nevertheless, we detected 394 asymmetrical reduction of *Ptep-soA* expression on single eyes of a subset of stage 14 395 embryos (n=6/16; Fig. 4D), which closely correlates with the predominance of mosaic 396 phenotypes observed in late postembryos (Fig. 5C). 397 Parental RNAi experiments using the same protocol targeting *Ptep-otdB* and *Ptep-OptixB* 398 did not result in any detectable phenotypic effects on the eyes of embryos from dsRNA-399 injected treatment (two and six females injected, respectively; counts not shown). These 400 results accord with a recent study that knocked down both Optix paralogs P. tepidariorum 401 and did not recover eye defects (Schacht et al., 2020) 402 403 Discussion 404 405 Paralogy of RDGN members in arachnopulmonates 406

407 Amblypygi have a critical placement within arachnid phylogeny, as they are part of a trio 408 of arachnid orders (collectively, the Pedipalpi, comprised of Amblypygi, Thelyphonida, and 409 Schizomida), which in turn is the sister group to spiders. Whereas the eyes of spiders have 410 greatly diversified in structure, function, and degree of visual acuity (particularly the eyes of 411 hunting and jumping spiders), the arrangement and number of eyes in Amblypygi likely 412 reflects the ancestral condition across Tetrapulmonata (= spiders + Pedipalpi), consisting of 413 three pairs of simple lateral ocelli and a pair of median ocelli; a similar condition is observed in basally branching spider groups like Mesothelae and Mygalomorphae, as well as 414 415 Thelyphonida (vinegaroons). However, while developmental genetic datasets and diverse

416 genomic resources are available for spiders and scorpions (Oda & Akiyama-Oda, 2020; 417 Posnien et al., 2014; Schwager et al., 2017; Sharma, Schwager, Extavour, & Wheeler, 418 2014b), the developmental biology of the other three arachnopulmonate orders has been 419 virtually unexplored in the past four decades beyond a single work describing the 420 embryology of one North American amblypygid species (Weygoldt, 1975). To address this 421 gap, we focused our investigation on a sister species pair of cave whip spiders and generated 422 the first embryonic transcriptomes for this order. These datasets are immediately amenable to 423 testing the incidence of RDGN duplicates previously known only from two spiders (Samadi 424 et al., 2015; Schomburg et al., 2015) and their putative effects in patterning eyes across 425 Arachnopulmonata broadly.

426 The inference of a partial or whole genome duplication (WGD) in the most recent 427 common ancestor of Arachnopulmonata is supported by the systemic duplications of 428 transcription factors and synteny detected in the genomes of the scorpion *Centruroides* 429 sculpturatus, and the spider Parasteatoda tepidariorum, as well as homeobox gene 430 duplications detected in the genome of the scorpion Mesobuthus martensii and transcriptome 431 of the spider *Pholcus phalangioides* (Leite et al., 2018; Schwager et al., 2017). Additional 432 evidence comes from shared expression patterns of leg gap gene paralogs in a spider and a 433 scorpion (Nolan et al., 2020). Embryonic transcriptomes are particularly helpful in the 434 absence of genomes, as several duplicated genes, such as some homeobox genes, are only 435 expressed during early stages of development (Leite et al., 2018; Sharma, Santiago, González-Santillán, Monod, & Wheeler, 2015a; Sharma, Schwager, Extavour, & Wheeler, 436 2014b). Our analysis of Charinus embryonic transcriptomes shows that RDGN gene 437 438 duplicates observed in spiders also occur in whip spiders, supporting the hypothesis that these 439 paralogous copies originated from a shared WGD event in the common ancestor of

440 Arachnopulmonata.

441 The conservation of some transcription factors patterning eyes is widespread in the 442 Metazoan tree of life (Vopalensky & Kozmik, 2009). In the model fruit fly D. melanogaster, 443 the homeobox Pax6 homolog, eveless, was the first transcription factor identified as a "master 444 gene", necessary for compound eye formation and capable of inducing ectopic eye formation 445 (Gehring & Ikeo, 1999; Kumar, 2009). The Pax6 protein is essential for eye formation across 446 several metazoan taxa, which has fomented ample debate about the deep homology of gene 447 regulatory networks in patterning structurally disparate eyes (Carroll, 2008; Shubin, Tabin, & 448 Carroll, 2009; Vopalensky & Kozmik, 2009). In the case of sine oculis (Six1/2), orthologs 449 are found across metazoans (Bebenek, Gates, Morris, Hartenstein, & Jacobs, 2004; Byrne et

450 al., 2017; Rivera et al., 2013). Evidence that *sine oculis* is required for the eye patterning in 451 other bilaterians comes from expression pattern in the developing eyes of the annelid 452 Platynereis dumerilii (Arendt, Tessmar, Medeiros de Campos-Baptista, Dorresteijn, & 453 Wittbrodt, 2002), and functional experiments in the planarian Girardia tigrina (Pineda et al., 454 2000). Therefore, studies interrogating the genetic bases of eye formation in chelicerate 455 models have the potential to clarify which components of the eye gene regulatory network of 456 Arthropoda evolved in the MRCA of the phylum, and which reflect deep homologies with 457 other metazoan genes. 458 459 A conserved role for a *sine oculis* homolog in patterning arachnopulmonate eyes 460 461 The eyes of arthropods are diverse in number, arrangement, structure and function (Paulus, 1979). Both types of eyes observed in Arthropoda, the faceted eyes (compound) and 462 463 single-lens eyes (ocelli), achieve complexity and visual acuity in various ways. To mention 464 two extremes, in Mandibulata the compound eyes of mantis shrimps (Stomatopoda) achieve a 465 unique type of color vision and movements by using 12 different photoreceptive types and 466 flexible eye-stalks (Daly, How, Partridge, & Roberts, 2018; Marshall, Cronin, & Kleinlogel, 467 2007; Thoen, How, Chiou, & Marshall, 2014). In Arachnida, the simple-lens median eyes of 468 some jumping spiders (Salticidae) have exceptional visual acuity in relation to their eye size, 469 achieve trichromatic vision through spectral filtering, and can move their retina using 470 specialized muscles (Harland, Li, & Jackson, 2012; Land, 1985; Zurek et al., 2015). 471 Comparative anatomy suggests that the common ancestor of Arthropoda had both lateral 472 compound eyes and median ocelli that then became independently modified in the arthropod 473 subphyla (Morehouse et al., 2017; Paulus, 1979). While in situ hybridization data for selected 474 RDGN genes across arthropods generally support the hypotheses of eye homology, 475 comparative developmental datasets remain phylogenetically sparse outside of Pancrustacea 476 (Samadi et al., 2015; Schomburg et al., 2015) 477 We therefore applied a bioinformatic approach in a study system that lacked any genomic 478 resources (Amblypygi) to assess whether RDGN homologs are transcriptionally active during 479 the formation of eyes in the eye-bearing C. ioanniticus (Comparison 1), as well as those that may be putatively involved in eye loss in its troglobitic sister species (Comparison 2). As first 480

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steps toward understanding how arachnid eyes are patterned, our experiments demonstrated

that soA, a sine oculis paralog identified as differentially expressed during the formation of

eyes in *C. ioanniticus*, is necessary for patterning all eyes of a model arachnid system with

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the same eye configuration (*Parasteatoda tepidariorum*). Thus, we provide the first
functional evidence that part of the RDGN is evolutionarily conserved in the most recent
common ancestor (MRCA) of insects and arachnids, and by extension, across Arthropoda.

487 The advantage of such a bioinformatic approach is that it can potentially narrow the range 488 of candidate genes for functional screens, due to the inherent challenges imposed by paralogy 489 when assessing gene function. Eve reduction in the cave fish Astvanax mexicanus has been 490 shown to involve differential expression of genes known to be involved in eye patterning in 491 model organisms, such as hedgehog and Pax6 (eyeless/toy) (Jeffery, 2009; Protas & Jeffery, 492 2012). In addition, other "non-traditional" candidates have been identified, such as hsp90 493 (Jeffery, 2009). Likewise, evidence from quantitative trait loci mapping in cave populations 494 of the troglobitic crustacean Asellus aquaticus shows that eye loss phenotype is correlated 495 with loci that are not part of the RDGN (Protas et al., 2011; Protas & Jeffery, 2012). The 496 results of the DGE analysis in whip spiders underscore the potential of a DGE approach to 497 triangulate targets among candidate genes in non-model species more broadly. Future efforts 498 in the Charinus system should focus on dissecting individual eye and limb primordia of 499 embryos of both species, in order to identify candidate genes putatively involved in the 500 reduction of each eye type, as well as compensatory elongation of the sensory legs of the 501 troglobitic species, toward downstream functional investigation.

502

503 Do gene duplications play a role in the functional diversification of arachnopulmonate eyes?504

505 A challenge in studying arachnopulmonate models to understand ancestral modes of eve 506 patterning in Arthropoda is the occurrence of RDGN duplicates in this lineage. Our orthology 507 searches and phylogenetic analysis showed that the evolutionary history of genes is not 508 always resolved using standard phylogenetic methods, as short alignable regions and/or 509 uncertainty of multiple sequence alignments can result in ambiguous gene trees. One way to 510 circumvent this limitation is by analyzing expression patterns via in situ hybridization 511 between paralogs in different arachnids in order to determine which patterns are 512 plesiomorphic (Leite et al., 2018; Nolan et al., 2020; Turetzek et al., 2015). Nonetheless, the 513 possibility of subfunctionalization and neofunctionalization may also complicate such 514 inferences because discerning one process from the other is analytically challenging (Sandve, 515 Rohlfs, & Hvidsten, 2018). 516 Genetic compensation of gene paralogs is another confounding variable, which can be

517 accounted for by experimental advances in model organisms (e.g., Shull et al., 2020). We

518 note that the overall penetrance in this experiment is low (9.5%) when compared to some 519 studies in *P. tepidariorum* (e.g., Khadjeh et al 2012; >59% in *Ptep-Antp* RNAi). Wide 520 variance in penetrance has been reported by several research groups in this system, with 521 phenotypic effects varying broadly even within individual experiments (e.g., Fig. 5 of 522 Akiyama-Oda & Oda, 2006; Fig. S5 of Schwager, Pechmann, Feitosa, McGregor, & Damen, 523 2009). Furthermore, some genes have empirically proven intractable to misexpression by 524 RNAi in *P. tepidariorum*, with one case suggesting functional redundancy of posterior Hox 525 genes to be the cause (Khadjeh et al., 2012). Double knockdown experiments have been 526 shown to exhibit poor penetrance (0-1.5%) in P. tepidariorum as well (Fig. S3 of Khadjeh et 527 al. 2012; Fig. S1 of Setton et al., 2017), and to our knowledge, no triple knockdown has ever 528 been achieved. While we cannot rule out functional redundancy with other RDGN paralogs 529 in the present study, the low penetrance we observed may also be partly attributable to our 530 conservative phenotyping strategy (see Material and Methods), which did not assess a 531 possible delay in eye formation and emphasized dramatic defects in eye morphology for 532 scoring.

533 The occurrence of RDGN gene duplications in Arachnopulmonata, in tandem with 534 improving functional genetic toolkits in P. tepidariorum (e.g., Pechmann, 2016), offers a 535 unique opportunity of studying the role of sub- and neofunctionalization in the development 536 of their eyes, and a possible role of this process in the diversification of number, position and 537 structure of the eyes in an ancient group of arthropods (Harland et al., 2012; Land, 1985; 538 Morehouse et al., 2017; Paulus, 1979; Zurek et al., 2015). The genomes of mites, ticks, and 539 harvestmen (Grbić et al., 2011; Hoy et al., 2016) (Gulia-Nuss et al., 2016) reveal that 540 apulmonate arachnid orders have not undergone genome duplication events as seen in 541 Arachnopulmonata (Schwager et al., 2017), or horseshoe crabs (Kenny et al., 2015; Nossa et 542 al., 2014; Zhou et al., 2020). Future comparative studies focused on understanding the 543 ancestral role of chelicerate RDGN genes should additionally prioritize single-copy orthologs 544 in emerging model systems independent of the arachnopulmonate gene expansion, such as 545 the harvestman Phalangium opilio (Sharma et al., 2013; Sharma, Schwager, Extavour, & 546 Giribet, 2012).

- 548 **Materials and Methods**
- 549
- 550 Animal collection
- 551

552 Three ovigerous females of the normal-eyes species, *Charinus ioanniticus* (ISR021-2; 553 ISR021-3; ISR021-4), and two egg-carrying females of the reduced-eyes species, Charinus 554 israelensis (ISR051-4; ISR051-6), were hand collected in caves in Israel in August 2018 555 (Supplementary Information; Table 1). Females were sacrificed and the brood sacs 556 containing the embryos were dissected under phosphate saline buffer (PBS). For each female, 557 a subset of the embryos (5 to 13 individuals) was fixed in RNAlater solution after poking a 558 whole into the egg membrane with fine forceps, while the remaining embryos of the clutch 559 were fixed in a 4% formaldehyde/PBS solution to serve as vouchers (SI Appendix, Table S1). 560 Adult animals and embryos of *Parasteatoda tepidariorum* were obtained from the colony at 561 UW-Madison, US. 562 563 Transcriptome assembly for Charinus whip spiders 564 565 RNAlater-fixed embryos were transferred to 1.5mL tubes filled with TRIZOL 566 (Invitrogen) after two months, and subject to RNA extraction. Total RNA extracted from 567 each sample of the embryos of C. ioannicitus (three samples) and C. israelensis (two 568 samples) (SI Appendix, Table S1) was submitted for library preparation at the Biotechnology 569 Center of the University of Wisconsin-Madison. Each sample was sequenced in triplicate in 570 an Illumina High-Seq platform using paired-end 100 bp-long read strategy at the same 571 facility. Read quality was assessed with FastQC (Babraham Bioinformatics). Paired-end 572 reads for C. ioanniticus (ISR021) and C. israelensis (ISR051) were compiled and de novo 573 assembled using Trinity v.3.3 (Grabherr et al., 2011) enabling Trimmomatic v.0.36 to remove adapters and low-quality reads (Bolger, Lohse, & Usadel, 2014). Transcriptome quality was 574 575 assessed with the Trinity package script 'TrinityStats.pl' and BUSCO v.3 (Waterhouse et al., 576 2017). For BUSCO, we used the 'Arthropoda' database and analyzed the transcriptomes 577 filtered for the longest isoform per Trinity gene. 578 579 RNA sequencing for differential gene expression 580 581 The total RNA extraction of each sample of C. ioanniticus and C. israelensis embryos

582 was sequenced in triplicate in an Illumina High-Seq platform using a single-end 100 bp-long

583 read strategy in the same facility as described above. For *C. ioanniticus* (normal-eyes), we

584 sequenced two biological replicates of embryos at an early embryonic stage, before eye-spot

formation (ISR021-2, ISR021-3), and one sample of late embryos, after eye-spot formation

- 586 (ISR021-4); For *C. israelensis* (reduced-eyes), we sequenced embryos at an early embryonic
- 587 stage (ISR051-6; ISR051-4) comparable to the early stage in *C. ioanniticus* (ISR021-2,
- 588 ISR021-3), as inferred by the elongated lateral profile of the body and marked furrows on the
- 589 opisthosomal segments (SI Appendix, Fig. S1).
- 590
- 591 Differential gene expression analysis in *Charinus* and identification of eye gene orthologs592
- 593 Orthologs of Drosophila melanogaster eyeless and twin of eyeless (Pax6A, Pax6B), sine
- 594 oculis (soA, soB), orthodenticle (otdA, otdB), Optix (Six3.1, Six3.2), dachshund (dacA, dacB),
- and eyes absent (eya) had been previously isolated in Parasteatoda tepidariorum
- 596 (Schomburg et al., 2015, and references therein). We used as reference sequences the
- 597 complete predicted transcripts for these genes from *P. tepidariorum* genome (Schwager et al.,
- 598 2017), Cupiennius salei (Samadi et al., 2015) (for atonal and Pax6), and D. melanogaster,
- 599 including also *atonal* and *eyegone* from the latter species. The sequences were aligned with
- 600 MAFFT (v7.407) (Katoh & Standley, 2013) and the resulting alignment used to build hidden
- 601 Markov model profiles for each gene (hmmbuild, from the hmmer suite v.3.3) (Finn et al.,
- 602 2015). Matches to these profiles were found using hmmsearch in the reference transcriptomes
- 603 of *C. ioanniticus* and *C. israelensis* as well as in the genomes of representative arthropods
- 604 including D. melanogaster (GCA 000001215.4), Tribolium castaneum (GCA 000002335.3),
- 605 Daphnia magna (GCA 003990815.1), Strigamia maritima (GCA 000239455.1),
- 606 Dinothrombium tinctorium (GCA 003675995.1), Ixodes scapularis (GCA 002892825.2),
- 607 Tetranychus urticae (GCA 000239435.1), Limulus polyphemus (GCA 000517525.1),
- 608 Tachypleus tridentatus (GCA 004210375.1), Centruroides sculpturatus (GCA 000671375.2),
- 609 Parasteatoda tepidariorum (GCA 000365465.2) and Trichonephila clavipes (GCA
- 610 002102615.1). These species were selected from a pool relatively recent genome assembly
- 611 resources and well curated reference genomes.
- Homologous sequences (those with hmmer expectation value,  $e < 10^{10}$ ) to the genes of
- 613 interest were then compiled into individual gene FASTA files, combined with the reference
- 614 sequences used for the homology search, aligned (MAFFT), trimmed of gap rich regions
- 615 (trimAL v.1.2, -gappyout) (Capella-Gutiérrez, Silla-Martínez, & Gabaldón, 2009) and used
- 616 for maximum likelihood gene tree estimation (IQTREE v.1.6.8, -mset
- 617 LG,WAG,JTT,DCMUT -bb 1000) (Nguyen, Schmidt, Haeseler, & Minh, 2015). The
- 618 association of transcripts in the *Charinus* species with the genes of interest is based on the
- 619 gene phylogeny and was followed by inspection of the coding sequences to distinguish

620 splicing variants from other gene copies. Alignments and the list of *Charinus* sequences is

available in SI Appendix Dataset S1. These gene transcript association was then used for the

- 622 transcript to gene map required for the DGE analysis.
- 623

624 Read mapping, transcript abundance quantification

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626 For the *in-silico* analysis of gene expression, single-end raw reads were first trimmed 627 using the software Trimmomatic v. 0.35 (Bolger et al., 2014). For the intra-specific analysis 628 of early (before eyespot) and late (eyespot) embryos of *C. ioanniticus* (Comparison 1), the 629 trimmed reads were quantified in the embryonic transcriptome of C. ioanniticus. For the 630 intra-specific comparison of early embryos of C. ioanniticus and C. israelensis, two 631 reciprocal analysis were conducted: reads from both species mapped onto C. israelensis 632 transcriptome as the reference (Comparison 2.1); and reads from both species mapped onto 633 C. ioanniticus transcriptome (Comparison 2.2). 634 Transcript abundance was quantified using the software Salmon v. 1.1.0 (Patro et al., 635 2017), enabling '-validateMapping' flag. Analysis of differential gene expression was 636 conducted with the software DESeq2 v 1.24.0 (Love et al., 2014) following a pipeline with 637 the R package tximport v.1.12.3 (Soneson, Love, & Robinson, 2015). The exact procedures 638 are documented in the custom R script (SI Appendix, Dataset S2) 639 Parental RNA interference, in situ hybridization, and imaging in Parasteatoda tepidariorum 640 641 642 Total RNA from a range of embryonic stages of *P. tepidariorum* was extracted with 643 TRIZOL (Invitrogen), and cDNA was synthetized using SuperScriptIII (Invitrogen). Gene 644 fragments for *Ptep-sine oculis A* (soA), orthodenticle B (otdB) and OptixB were amplified 645 from cDNA using gene specific primers designed with Primers3Web version 4.1.0 646 (Koressaar & Remm, 2007) and appended with T7 ends. Cloning amplicons were generated 647 using the TOPO TA Cloning Kit with One Shot Top10 chemically competent Escherichia 648 coli (Invitrogen). Amplicon identities and directionality were assessed with Sanger 649 sequencing. Primer, amplicon sequences and fragment lengths are available in SI Appendix Dataset S3. Double-stranded RNA for Ptep-soA, Ptep-otdB and Ptep-OptixB was synthetized 650 651 from amplicon on plasmids using MEGAScript T7 transcription kit (Thermo Fischer) with 652 T7/T7T3 primers. Sense and antisense RNA probes for colorimetric in situ hybridization

653 were synthetized from plasmid templates with DIG RNA labeling mix (Roche) and T7/T3

654 RNA polymerase (New England Biolabs) using the manufacturer's instructions.

655 Parental RNA interference (RNAi) followed established protocols for double-stranded 656 RNA (dsRNA) injection in virgin females of *P. tepidariorum* (Oda & Akiyama-Oda, 2020). 657 Each female was injected four times with 2.5 µL of dsRNA at a concentration of 2 µg/uL, to 658 a total of 20µg. For *Ptep-soA*, seven virgin females were injected with dsRNA of a 1048bp 659 cloned fragment (SI Appendix, Fig. S13C) and 3 females were injected with the same volume 660 of dH<sub>2</sub>O as a procedural control. Two virgin females were injected with dsRNA for Ptep-661 otdB, and six females for *Ptep-OptixB*. All females were mated after the second injection, and were fed approximately every-other day after the last injection. Cocoons were collected until 662 663 the sixth clutch, approximately one per week.

664 Hatchlings for all cocoons were fixed between 24–48 hours after hatching. Freshly hatched postembryos have almost no external signs of eye lenses and pigments. The selected 665 666 fixation window encompasses a period in which postembryos have deposited eye pigments 667 until the beginning of the first instar, where eyes are completely formed (SI Appendix, Video 668 S1, S2). Hatchlings were immersed in 25% ethanol/PBST and stored at 4°C. For the Ptep-soA 669 RNAi experiment, hatchlings were scored in four classes: (1) wild type, where all eyes were 670 present and bilaterally symmetrical; (2) Eyes defective, where one or more eyes were reduced 671 in size or completely absent; (3) dead/arrested; (4) Undetermined, where embryos were 672 damaged or clearly freshly hatched. A subset of *Ptep-soA* dsRNA-injected embryos from four clutches (n=48) and of three control clutches (n=48) were further inspected in detail to assess 673 674 the effects on individual eye types. Given that there is a spectrum on the intensity of pigment 675 deposition in the medium eyes (ME), and small asymmetries on the shape of the early 676 developing tapetum of the lateral eyes (LE) in control embryos, the following conservative 677 criteria was adopted: ME were considered affected when asymmetry in pigmentation or lens 678 size was detected. Both ME were only scored as affected when they were both completely 679 missing, in order to rule out embryos were simply delayed in pigment deposition; LE were 680 considered defective only when the tapetum was completely absent (SI Appendix, Fig. S12). 681 Therefore, our coding does not allow detection of a phenotype consisting of delayed 682 pigmentation. For in situ hybridization, a subset of *Ptep-soA* dsRNA-injected embryos at stage 13/14 683

(Mittmann & Wolff, 2012) was fixed in a phase of heptane and 4% formaldehyde for 12–24
hours, washed in PBST, gradually dehydrated in methanol and stored at -20°C for at least 3

686 days before downstream procedures, after a modified protocol of Akiyama-Oda and Oda 687 (2003). In situ hybridization followed the protocol of Akiyama-Oda and Oda (2003). 688 Embryos from in situ hybridization were stained with Hoechst nuclear staining and 689 imaged in a Nikon SMZ25 fluorescence stereomicroscope mounted with a DS- Fi2 digital 690 color camera (Nikon Elements software). For postembryos, the prosoma was dissected with 691 fine forceps, gradually immersed in 70% Glycerol/PBS-T and mounted on glass slides. 692 Postembryos were imaged using an Olympus DP70 color camera mounted on an Olympus 693 BX60 epifluorescence compound microscope. 694 695 Acknowledgements 696 697 Microscopy was performed at the Newcomb Imaging Center, Department of Botany, 698 University of Wisconsin-Madison. Sequencing was performed at the UW-Madison 699 Biotechnology Center. Access to computing nodes for intensive tasks was provided by the 700 Center for High Throughput Computing (CHTC) and the Bioinformatics Resource Center 701 (BRC) of the University of Wisconsin-Madison. Specimens were collected under permit 702 2018/42037, issued by the Israel National Parks Authority to E.G.R. Fieldwork in Israel was 703 supported by a National Geographic Society Expeditions Council grant no. NGS-271R-18 to 704 J.A.B. This work was supported by National Science Foundation (grant no. IOS-1552610) to 705 P.P.S. 706 707 References 708 709 Akiyama-Oda, Y., & Oda, H. (2006). Axis specification in the spider embryo: dpp is required 710 for radial-to-axial symmetry transformation and sog for ventral patterning. Development, 711 133(12), 2347–2357. http://doi.org/10.1242/dev.02400 712 Arendt, D., Tessmar, K., Medeiros de Campos-Baptista, M. I., Dorresteijn, A., & Wittbrodt, 713 J. (2002). Development of pigment-cup eyes in the polychaete Platynereis dumerilii and 714 evolutionary conservation of larval eyes in bilateria. Development, 129(5), 1143-1154. 715 Ballesteros, J. A., & Sharma, P. P. (2019). A Critical Appraisal of the Placement of 716 Xiphosura (Chelicerata) with Account of Known Sources of Phylogenetic Error. 717 Systematic Biology. http://doi.org/10.1093/sysbio/syz011 718 Ballesteros, J. A., Santibáñez López, C. E., Kováč, L., Gavish-Regev, E., & Sharma, P. P. 719 (2019). Ordered phylogenomic subsampling enables diagnosis of systematic errors in the 720 placement of the enigmatic arachnid order Palpigradi. Proceedings. Biological Sciences, 721 286(1917), 20192426. http://doi.org/10.1098/rspb.2019.2426

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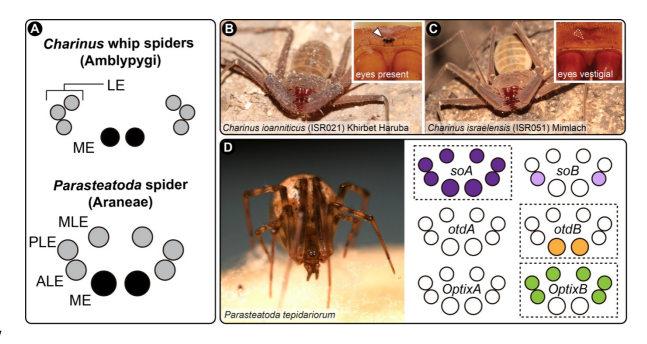
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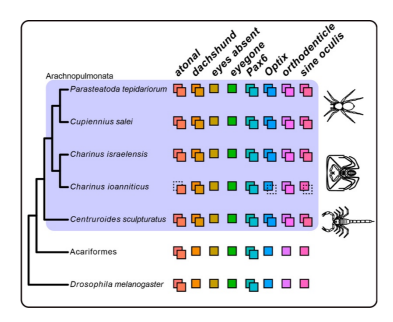
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- 1066

1068 Figure 1: Species used in this study and their eye arrangement. A: Schematic representation 1069 of the eyes of Charinus whip spiders (Amblypygi) (upper), and the spider Parasteatoda 1070 tepidariorum (Araneae; lower). B: Live specimen of C. ioanniticus from Khirbet Haruba 1071 cave (Haruva cave). Inset: detail of the median eyes. C: Live specimen of C. israelensis 1072 from Mimlach cave. Inset: detail of the reduced median eyes. D: Live specimen of 1073 Parasteatoda tepidariorum, and schematic representation of the expression patterns of 1074 paralog pairs of *Ptep-sine oculis* (soA/soB), *Ptep-orthodenticle* (otdA/otdB), and *Ptep-*1075 *Optix (OptixA/OptixB)* in the eyes. ME: median eyes; ALE: anterior lateral eyes; PLE: 1076 posterior lateral eyes; MLE: median lateral eyes; LE: lateral eyes.



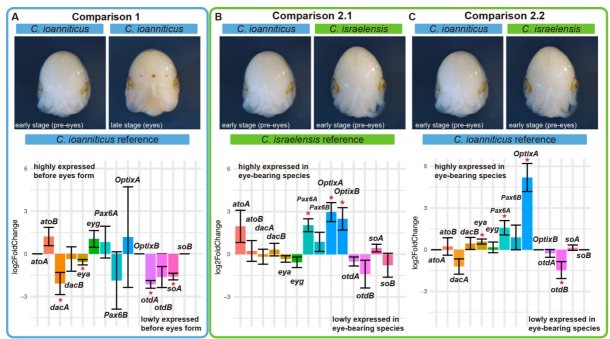
- 1078 Figure 2: Phylogenetic distribution of Retinal Determination Gene Network (RDGN) genes
- 1079 in an insect (Drosophila melanogaster), a non-arachnopulmonate arachnid group
- 1080 (Acariformes: *Dinothrombium tinctorium*; *Tetranychus urticae*) and Arachnopulmonata
- 1081 (spider: Parasteatoda tepidariorum; scorpion: Centruroides sculpturatus), including
- 1082 newly discovered orthologs in *Charinus* whip spiders (Amblypygi). Colored squares
- 1083 indicate number of gene copies for each RDGN gene. Dotted squares indicate missing
- 1084 data, not gene loss. For comprehensive list of duplicated genes in Arachnopulmonata see
- Schwager et al. (2017) and Leite et al. 2018. Gene trees and alignments for each gene areavailable in SI Appendix Dataset S1.



1087

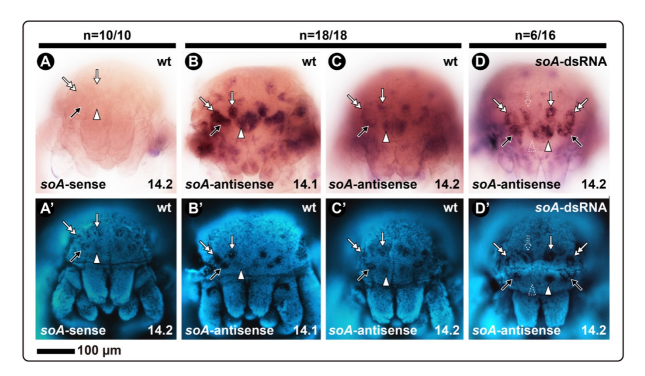


- 1090 (RDGN) genes in *Charinus* whip spider deutembryos. Bar graphs display log<sub>2</sub> fold
- 1091 change of selected RDGN genes. Direction of differential gene expression always
- 1092 follows sample to the left. A: Comparison 1; Comparison between reads of early (pre-
- 1093 eyespot) and late deutembryos (eyespot) of the eye-bearing species *C. ioanniticus*
- 1094 mapped onto *C. ioanniticus* transcriptome. B: Comparison 2.1; Comparison between
- 1095 reads of early deutembryo of *C. ioanniticus* and early deutembryo of *C. israelensis*
- 1096 mapped onto *C. israelensis* transcriptome. C: Comparison 2.2; Comparison between
- 1097 reads of early deutembryo of *C. ioanniticus* and early deutembryo of *C. israelensis*
- 1098 mapped onto *C. ioanniticus* transcriptome. *atoA/B*: *atonalA/atonalB*; *dacA/B*:
- 1099 *dachshundA/B*; eya: eyes absent; eyg: eyegone; otdA/B: orthodenticleA/B; soA/B: sine
- 1100 *oculisA/B*. Asterisks denote genes that were differentially expressed with a  $p_{adj} > 0.05$ .
- 1101  $Log_2FC = 0$  for *atoA*, *OptixB*, and *soB* for Comparison 1 and Comparison 2.2 are due to
- the absence of those paralogs in *C. ioanniticus* reference transcriptome.



1103

1105 Figure 4: In situ hybridization using DIG-labeled riboprobes for *Ptep-soA* in late embryos of 1106 the spider Parasteatoda tepidariorum. All embryos in frontal view. A-D: bright field 1107 images. A'-D': Same embryos, in Hoechst staining. A: Sense probe of a stage 14.2 1108 embryo (no signal). B: Antisense probe on a wild type stage 14.1 embryo. C: Antisense 1109 probe on a wild type stage 14.2 embryo. D: Antisense probe on a stage 14.2 embryo from 1110 the *Ptep-soA* dsRNA-injected treatment. *soA*: *sine oculis A*. White arrowhead: median 1111 eye; Black arrow: anterior lateral eye; White arrow: median lateral eye; Double white 1112 arrow: Posterior lateral eye. Dotted arrowhead/arrow indicate asymmetrical expression 1113 and eye defect. Sample sizes are indicated above each treatment.



1114

1116 Figure 5: RNA interference against *Ptep-sine oculis A*. A: Bright field images of the spider 1117 Parasteatoda tepidariorum postembryos resulting from control treatment (dH<sub>2</sub>O-1118 injected, panel 1) and double stranded RNA (dsRNA) injected treatment (panels 2–6), in 1119 frontal view. B: Frequencies of each phenotypic class per treatment from the combined 1120 clutches of all females. See SI Appendix Fig. S13 for counts per clutch. C: Frequencies 1121 of symmetrical, asymmetrical, and wild type eyes quantified from a subset of 48 1122 individuals with eye reduction phenotype. See SI Appendix Fig. S12 for figures of all specimens and coding, and Material and Methods for the scoring criteria. ME: median 1123 eyes; ALE: anterior lateral eyes; PLE: posterior lateral eyes; MLE: median lateral eyes. 1124 Schematics for the different eye types follows the nomenclature in Figure 1. 1125

