-					
ĸ	ACA6	arch	Λr	tic	Δ
1/	COU	иu	Δ	uu	ľ

- 2 Mitochondrial genomes infer phylogenetic relationships among the
- 3 oldest extant winged insects (Palaeoptera)
- 4 Sereina Rutschmann^{1,2,3}, Ping Chen⁴, Changfa Zhou⁴, Michael T. Monaghan^{1,2*}
- 5 Addresses:
- 6 Leibniz-Institute of Freshwater Ecology and Inland Fisheries (IGB), Müggelseedamm 301, 12587
- 7 Berlin, Germany
- 8 ²Berlin Center for Genomics in Biodiversity Research, Königin-Luise-Straße 6-8, 14195 Berlin,
- 9 Germany

15

18

19

- ³Department of Biochemistry, Genetics and Immunology, University of Vigo, 36310 Vigo, Spain
- ⁴The Key Laboratory of Jiangsu Biodiversity and Biotechnology, College of Life Sciences, Nanjing
- Normal University, Nanjing 210046, China
- sereina.rutschmann@gmail.com; pingc918@hotmail.com; zhouchangfa@njnu.edu.cn;
- 14 <u>monaghan@igb-berlin.de</u>
- *Correspondence: sereina.rutschmann@gmail.com
- 17 Department of Biochemistry, Genetics, and Immunology, University of Vigo, 36310 Vigo, Spain

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

Abstract

Background: The relationships among the oldest winged insects (Palaeoptera), including the Ephemeroptera (mayflies) and Odonata (dragonflies and damselflies), remain unclear. The Palaeoptera together with the Neoptera have evolved as result of a rapid divergence from a common ancestor in the distant past. Thus they are thought to be more susceptible to systematic inadequacies, including taxon sampling, choice of outgroup, marker selection, and phylogenetic methods. Here we reconstruct their phylogenetic relationship using newly sequenced mitochondrial genomes in combination with 90 additional insect mitochondrial genomes. In particular, we investigate the impact of the increased mayfly taxon sampling, the effect of rogue taxa, and the used phylogenetic framework (Bayesian inference (BI) vs. maximum likelihood (ML)) approach. Results: We found support for the clustering of the Odonata as most ancient, extant winged insects, using BI based on an optimized data matrix. Overall, we found no support for the basal Ephemeroptera clustering and the sister relationship between the Ephemeroptera and Odonata. Our newly sequenced mitochondrial genomes of Baetis rutilocylindratus, Cloeon dipterum, and Habrophlebiodes zijinensis showed the complete set of 13 protein coding genes and a conserved gene orientation with the exception of two inverted tRNAs for *H. zijinensis*. Conclusions: The increase of palaeopteran taxon sampling in combination with a Bayesian phylogenetic framework was crucial to infer phylogenetic relationships within the three ancient insect lineages of Odonata, Ephemeroptera, and Neoptera. Pruning of rogue taxa improved the number of supported nodes in all phylogenetic trees. It remains to be tested weather an increased taxon sampling might also reveal the elusive phylogenetic positions of other insect orders. Keywords: Baetidae, Bayesian Inference, Ephemeroptera, Long-branch attraction, Mayfly, Mitochondrial genomics, Odonata, Palaeoptera problem, Polyneoptera, Pterygota, Rogue taxa

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

Background Insects are the most diverse branch of metazoan life, yet the relationships among the basal branches of the winged insects (Pterygota) remains one of the major open questions in the evolution of life. The unresolved relationship of the basal orders Odonata (dragon- and damselflies) and Ephemeroptera (mayflies) to the Neoptera (i.e., to the rest of the Pterygota) has been termed the "Palaeoptera problem" [1,2]. The name Paleoptera ("old wings") reflects the inability of the Ephemeroptera and Odonata to fold their wings flat over the abdomen. This has long been considered to be the ancestral condition, in contrast to the more derived Neoptera ("new wings") (reviewed by Trautwein et al. [3]). The monophyly of the Neoptera, including the three major lineages Polyneoptera, Paraneoptera, and Holometabola, is widely accepted, although relationships among polyneopteran lineages [4] and the monophyly of the Paraneoptera [5] remain unresolved. There are three competing hypotheses relating to the Palaeoptera problem: the Palaeoptera hypothesis ((Ephemeroptera + Odonata) + Neoptera), the basal Ephemeroptera hypothesis (Ephemeroptera + (Odonata + Neoptera)), and the basal Odonata hypothesis (Odonata + (Ephemeroptera + Neoptera)) [3,6,7]. All hypotheses are supported by morphological as well as molecular data to varying degrees (reviewed by Trautwein et al. [3]). Different authors, using the same set of genes and taxa set but different phylogenetic approaches, namely Bayesian inference (BI) vs. maximum likelihood (ML) [2,8,9], and nucleotide vs. amino acid (aa) sequences [10] obtained results supporting distinct hypotheses. The Palaeoptera hypothesis has received support from several molecular studies based on rRNA genes and nuclear DNA [5,11-13], and from nDNA phylogenomic analyses [9,14,15]. Thomas et al. [12] and Kjer et al. [11] included the most comprehensive taxon sampling with 35, respectively seven species from each of the three lineages (i.e., Ephemeroptera, Odonata, Neoptera) analyzing seven, respectively eight genes (rRNAs, nuclear DNA (nDNA), mitochondrial DNA (mtDNA)). The basal Ephemeroptera hypothesis was supported by previous studies using mitochondrial genome data [10,16,17], and a combined analysis of rRNA

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

and one nuclear gene [2]. Other studies have concluded the relationships are ambiguous [2]. Notably, studies supporting the basal Ephemeroptera hypothesis have included either a limited number of mayfly species (e.g., one [17,18], two [16,19], four [20], or five [10] species) or a limited number of unlinked loci (e.g., one [10,16,17,20], two [18,19,21], or three [2]). The basal Odonata hypothesis has received support from previous studies based on mitochondrial genome data [22,23], rRNA genes [8,24-27], combining rRNA and nDNA [28], and phylogenomic nDNA [9,29]. These studies, with the exceptions of few studies based on 18S rRNA [24,25,27], included in total between two [8.29] and 13 species [26] from the orders Ephemeroptera and Odonata. The conflicting phylogenetic signals may result from the ancient radiation of the lineages Ephemeroptera, Odonata, and Neoptera from a common ancestor in the distant past [4]. These three ancient lineages appear to have diverged rapidly, leaving few characteristics to determine their phylogenetic relationships [4]. Evolutionary rate heterogeneity across clades and the representation of old clades by recent extant taxa make ancient relationships such as those of early winged insects the most difficult challenges for phylogenetics [4,7]. Given the weak phylogenetic signal, reconstructions are more susceptible to systematic errors [4,12,30-32]. Other sources of conflicting signal in studies of the relationships among these three lineages may result from differences in taxon sampling, sequence data, alignment methods, and phylogenetic methods used including models of evolution [3,12,33]. The use of mitochondrial protein-coding genes (PCGs) derived from mitochondrial genomes can circumvent some of these difficulties, since they are easier to align and also appropriate models of molecular evolution are well established [34-36]. On the other hand, mitochondrial genes include several drawbacks, most importantly the possible presence of pseudogenes [37-39]. However, data derived from high-throughput sequencing, depending on the coverage bears the potential to distinguish between functional genes and pseudogenes based on higher mutation rates of the later [40].

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

116

117

Overall, mitochondrial markers are well studied and therefore the most widely employed genetic markers in insects, being considered a promising "instrument" for insect systematics [41]. Insect mitochondrial genomes are highly conserved, ranging from 15 to 18 kb in length, containing 37 genes: 13 PCGs, two ribosomal RNAs (rRNAs, rrnL and rrnS), and 22 transfer RNAs (tRNAs, trn*) [42]. A non-coding region of variable length, thought to be the origin of initiation of transcription and replication, is typically present [43,44] and referred to as the AT-rich region. The typical ancestral insect mitochondrial genome differs from the ancestral arthropod mitochondrial genome only by the location of trnL [45]. Significant differences in structure, gene content, and gene arrangement have been found to be the exception for highly derived taxa. Importantly there is still an unbalance in the numbers of available molecular markers between the different insect lineages. In particular the early winged insects (i.e., Ephemeroptera and Odonata) are relatively under-represented. This is (partly) because so far most genomic data were obtained for insect orders that are of economical interest (e.g., as pollinators, model species, agricultural pests, and vectors of human diseases, [46]). As of January 2017, there were 39 complete or nearly complete palaeopteran mitochondrial genomes (Ephemeroptera: 18, Odonata: 21) from 22 families (eleven of each) available on GenBank; 22 of which were included in publications (Ephemeroptera: ten, Odonata: twelve). Compared to the number of families that have been described (Ephemeroptera: 42 [47], Odonata: 30 [48]) this is still a relatively low number. The insect order Ephemeroptera comprises over 3,000 species, comprising as the two most species rich families the Baetidae (833 species) and the Leptophlebiidae (608 species) [47]. Their biogeographic origin is probably Pangean, including a greater diversity and higher endemism rate in the Neotropics and Australasia for the Baetidae, and the Neotropical and Afrotropical regions for the Leptophlebiidae, respectively [47]. We investigated the relationships of the oldest extant winged insects and newly sequenced three mayfly mitochondrial genomes to improve sampling of this under-represented order. New taxa

included one representative of the family Leptophlebiidae, for which no mitochondrial genome data were available, and two representatives of the Baetidae, one from each subfamily. We used BI and ML approaches and created matrices that eliminated taxa that reduced overall tree support (rogue taxa) and that were affected by long-branch attraction (LBA). Our analysis included 29 palaeopteran mitochondrial genomes from 20 families and 64 other insect mitochondrial genomes. We expect the increased taxon sampling to improve the phylogenetic signal of the basal nodes and add support for one of the three competing hypotheses.

Methods

Taxon sampling

We newly sequenced three mitochondrial genomes of the two most diverse Ephemeroptera families. *Habrophlebiodes zijinensis* GuI, ZHANG & Wu, 1996 is the first representative of the family Leptophlebiidae to have its complete mitochondrial genome sequenced. *Baetis rutilocylindratus* WANG, QIN, CHEN & ZHOU, 2011 and *Cloeon dipterum* L. 1761 are both members of the Baetidae and representative each subfamily: *B. rutilocylindratus* from Baetinae, and *C. dipterum* from Cloeninae. There was a mitochondrial genome from one other member of the Baetinae available on GenBank prior to our study (*Alainites yixiani*, GQ502451, Jia & Zhou). Our analysis also included four species from Archaeognatha, three from Zygentoma (formerly Thysanura), ten from Odonata, 16 additional species from Ephemeroptera, four from Plecoptera, four from Ensifera of Orthoptera, nine from Caelifera of Orthoptera, 13 from Phasmatodea, two from Mantodea, six from Blattodea, 13 from Isoptera, and one each from Collembola, Dermaptera, Grylloblattodea, and Mantophasmatodea (Table 1). Amino acid sequences were obtained from GenBank using a custom Python script (mitogenome ncbi.py, https://github.com/srutschmann/python scripts).

[Table 1 should be added here]

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

159

160

161

162

163

164

165

166

167

Sequencing and assembly The three mayfly species were sequenced with either 454/FLX pyrosequencing (C. dipterum) or Sanger sequencing (B. rutilocylindratus and H. zijinensis. We extracted DNA from C. dipterum from twelve to twenty pooled reared subimago specimens, prepared a shotgun library, and sequenced four lanes on a Roche (454) GS FLX machine at the Berlin Center for Genomics in Biodiversity Research (BeGenDiv, Berlin, Germany) (see also [49]). The obtained sequence reads were trimmed and de novo assembled using the software Newbler v.2.5.3 (454 Life Science Cooperation) under default settings for large data sets. In order to extract the mitochondrial genome of C. dipterum, we performed BLASTN searches [50] using as query all assembled contigs against the NCBI database. We mapped all matching reads back to the mitochondrial genome with BWA [51], using the BWA-SW algorithm [52] with settings suggested for 454 data by CORAL (match score = 2, mismatch penalty = 2, and gap open penalty = 3, [53]). Specimens of B. rutilocylindratus and H. zijinensis were collected in Zijin Hill, Nanjing, China. The DNA was extracted from between two and four larvae using the DNeasy® Blood & Tissue (Qiagen, Leipzig, Germany) kit. Four DNA fragments of each species were amplified with universal primers (B. rutilocylindratus: cob, cox1, cox3, rrnL; H. zijinensis: cob, cox1, nad4, rrnL, [54]). Subsequently, based on the previously obtained sequence information we designed six (B. rutilocylindratus) respective four (H. zijinensis) specific primer pairs (Additional file 1: Table S1; see also Li et al. [55]). Standard and long polymerase chain reactions (PCRs) were performed on a DNA Engine Peltier Thermal Cycler (Bio-Rad, Shanghai, China). Therefore, we used the rTagTM DNA polymerase (TaKaRa Bio, Dalian, China) for fragments smaller than two kb and the LA TaqTM polymerase (TaKaRa Bio, Dalian, China) for fragments larger than two kb. All PCR products were purified with the Axygen agarose-out kit. When the PCR amplification signal was too weak to sequence or sequencing resulted in overlap peaks, the products were ligated to pGEM®-T Easy Vector (Promega, Southampton, UK) by Escherichia coli, and each resulting clone was sequenced.

169

170

171

172

173

174

175

176

177

178

179

180

181

182

183

184

185

186

187

188

189

190

191

192

All purified amplification products were sequenced successively in both directions on an ABI3130xl capillary sequencer. Forward and reverse sequences were assembled and edited using CodonCode Aligner v.3.5.6 (CodonCode Corporation). Sequence alignments and data matrices We prepared two data matrices of an sequences based on the annotated mitochondrial genomes (see below), including a set with all 93 taxa (all taxa matrix), and a matrix for which we removed taxa that were either identified as rogue taxa using the RogueNaRok algorithm [56,57] and showing LBA [58,59] (optimized taxa matrix; 88 taxa, see Table 1). We used GUIDANCE2 [60,61] to align the sequences, and identify and remove positions detected as ambiguously aligned regions. To run GUIDANCE we applied the default settings, using as multiple sequences alignment program MAFFT v.7.050b (L-INS-I algorithm with default settings, [62]). The best-fitting partitioning schemes and corresponding as substitution models were estimated with PartitionFinder v.2 (https://github.com/brettc/partitionfinder, [63]). We used the Bayesian Information Criterion (BIC) to choose the best model, linked branch lengths, and a greedy search [63] with RAxML v.8.2.8 [64]. Thereby each gene was defined as one data block (i.e., possible partition). To identify rogue taxa, we applied the RogueNaRok algorithm [56,57]. We used the bootstrap replicates and the best supported maximum likelihood tree based on the all taxa matrix (see below). Identified rogue taxa were removed from the all taxa matrix, resulting in the optimized taxa matrix (Table 1). Taxa showing evidence of LBA sensu Bergsten [65] were identified using heterogeneities in sequence divergence based on a model of evolution derived from the entire data matrix. For this, the Relative Composition Frequency Variability (RCFV) values of each species were calculated with BaCoCa [66] using the partitioned concatenated data matrix (i.e., all taxa matrix). Taxa with the highest values were excluded in the optimized taxa matrix, apart from the Ephemeroptera, which were retained because they are the focus of our study and their monophyly is well-established [1,2].

194

195

196

197

198

199

200

201

202

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

Phylogenetic reconstruction Phylogenetic reconstructions were carried out using MrBayes v.3.2.4 [67], and RAxML v.8.1.7 [64]. We analyzed both matrices (all taxa, optimized taxa) using the best-fitting partitioning scheme and model for each (see Table 2). For BI, we unlinked the frequencies, gamma distributions, substitution rates and the proportion of invariant sites across partitions. For each matrix, we run two independent analyses of four MCMC chains with 10⁷ generations, sampling every 10³ generations and discarding a burn-in of 25%. Maximum likelihood inferences were performed with 200 bootstrap replicates. Trees reconstructed from the all taxa matrix were rooted with Tetrodontophora bielanensis (Collembola). Trees reconstructed from the optimized taxa matrix were rooted with Nesomachilis australica (Archaeognatha, according to Song et al. [10]) because Tetrodontophora bielanensis was found to be a rogue taxon (see Results). All trees were visualized with the ape v.4.1 [68], phangorn v.2.1.1 [69], and ggtree v.1.4.20 [70] packages for R v.3 [71]. Annotation and characterization All mitochondrial genomes were annotated using the MITOS webserver (http://mitos.bioinf.unileipzig.de/index.py, [72]), DOGMA [73], and MFannot (http://megasun.bch.umontreal.ca/cgibin/mfannot/mfannotInterface.pl). The predicted PCGs were checked for stop codons and manually adjusted by comparison with the longest predicted open reading frames as implemented in Geneious R7 v.7.1.3 (Biomatters Ltd.), and by comparison with the respective homologous insect sequence alignment. For the tRNA prediction, we additionally used ARWEN v.1.2.3 [74] and tRNAscan-SE v.1.21 (http://lowelab.ucsc.edu/tRNAscan-SE/, [75]). The annotated mitochondrial genomes were visualized with OrganellarGenomeDRAW (http://ogdraw.mpimp-golm.mpg.de/cgi-bin/ogdraw.pl, [76]) and manually edited. Nucleotide contents were retrieved using Geneious. To correct biases in the AT content due to incomplete mitochondrial genomes mostly missing the AT-rich region, we removed all AT-rich

regions and the two rRNAs including the five close-by tRNAs (trnL1-trnM) and recalculated the

nucleotide base pair (bp) compositions. The AT and GC composition skewness were calculated as follows: AT-skew = (A - T) / (A + T), and GC-skew = (G - C) / (G + C) [77].

Results

Data matrices

The final matrices had lengths of 3,675 aa (all_taxa matrix, including 93 taxa) and 3,673 aa (optimized_taxa matrix, including 88 taxa, see Table 1). As best-fitting partitioning scheme for the two matrices, we identified either five partitions (all_taxa matrix, Table 2) or four partitions (optimized_taxa, Table 2). As best-fitting models of aa sequence evolution, we identified MtZoa [78] and MtArt [35].

Table 2 Overview of data matrices, including the best-fitting partitioning and model schemes, and amino acid (aa) sequence length

Matrix	Model	Length [aa]
all_taxa		
partition_1: atp6, atp8, nad2, nad3, nad6	$MTZOA + \Gamma + I + F$	840
partition_2: cox1	$MTZOA + \Gamma + I$	528
partition_3: cob, cox2	$MTZOA + \Gamma + I + F$	608
partition_4: cox3	$MTART + \Gamma + I$	268
partition_5: nad1, nad4, nad4L, nad5	$MTZOA + \Gamma + I + F$	1431
optimized_taxa		
partition_1: atp6, atp8, nad2, nad3, nad6	$MTZOA+\Gamma+I+F$	843
partition_2: cox1	$MTZOA + \Gamma + I$	529
partition_3: cob, cox2, cox3	$MTZOA + \Gamma + I + F$	878
partition_4: nad1, nad4, nad4L, nad5	$MTZOA + \Gamma + I + F$	1423

The RogueNaRok analysis identified the outgroup *T. bielanensis* as well as *Gryllotalpa orientalis* (Orthoptera) and *Phraortes* sp. (Phasmatodea) as taxa with uncertain phylogenetic positions, leading to less accurate phylogenetic reconstructions. *G. orientalis* (Orthoptera) and *Phraortes* sp.

(Phasmatodea) clustered within the corresponding order using the all_taxa matrix, although the nodes containing these species were not supported by ML. In the test for LBA, *Aposthonia japonica* (Embioptera) and *Challia fletcheri* (Dermaptera) had high RCFV values (0.0237 and 0.0307, respectively) suggesting they were affected by LBA. Both species clustered within other orders: *Challia fletcheri* (Dermaptera) within the Ephemeroptera as sister taxon to a clade containing all representatives of the family Baetidae, and *A. japonica* (Embioptera) within the Phasmotodea as sister taxon to a clade containing all species of the order with the exception of *T. californicum* (Table 3, Additional file 2: Figure S1). The mayflies Teloganodidae sp. (0.0305) and the three baetids *A. yixiani* (0.0235), *B. rutilocylindratus* (0.0232), and *C. dipterum* (0.0201) showed relatively high RCFV values.

Phylogenetic reconstruction

The Bayesian reconstruction based on the optimized_taxa matrix produced the tree with most support compared to ML or to either analysis of the all_taxa matrix (Figure 1, Table 3, Additional files 2-4: Figures S1-S3). The basal Odonata hypothesis was highly supported in this tree (Bayesian posterior probability (BPP) = 1.00), whereas neither the basal Ephemeroptera hypothesis nor the Palaeoptera hypothesis received support in any analysis (Table 3). The Odonata were monophyletic in all analyses (BPP = 1.00, BS = 100%) whereas the Ephemeroptera were monophyletic only using the optimized_taxa matrix (Figure 1, Table 3, Additional file 3: Figures S2).

Table 3 Support for the three hypotheses and node support for clades of interest. Node support is based on Bayesian inference (BI) and maximum likelihood (ML) analyses of two data matrices: optimized_taxa and all_taxa, whereby optimized_taxa had 5 terminal nodes removed based on the occurrence of rogue taxa and long branch attraction. Asterices indicate hypothesis/node support (** = Bayesian posterior probability (BPP) \geq 0.95 and Bootstrap support (BS) = 90%) * = BPP \geq 0.90

and $BS \ge 80\%$). Superorders are underlined, and suborders are italicised. Letters in brackets refer to monophyletic clades in the Figure 1

Hypothesis/Clade	BI		ML	ML		
Hypothesis/Clade	optimized_taxa	all_taxa	optimized_taxa	all_taxa		
Palaeoptera hypothesis						
Ephemeroptera hypothesis						
Odonata hypothesis	**					
Winged insects (A)	**					
Palaeoptera						
Odonata (B)	**	**	**	**		
Ephemeroptera (C)	**		**			
Polyneoptera			*			
Plecoptera (D)	**	**	**	**		
Orthoptera						
Ensifera (E)	**	**	**			
Caelifera (F)	**	**	**	**		
Phasmatodea ^a (G)	**	**	**	**		
Dictyoptera (H)	**	**	**	**		
Mantodea (I)	**	**	**	**		
Isoptera (J)	**	**	**	**		

^a, Includes *Aposthonia japonica* (Embioptera) in reconstructions using the all taxa matrix.

Most other nodes of interest were consistently supported across different matrices and analyses (Table 3). The Polyneoptera were moderately supported (BS \geq 80%) for the optimized_taxa matrix using the ML. Within the Polyneoptera, all orders except Orthoptera and Blattodea were monophyletic (BPP \geq 0.95, and BS \geq 90%, Table 3). The two Orthoptera suborders Ensifera and Caelifera were monophyletic (Table 3), whereby the Ensifera clustered together with the Plecoptera for the optimized_taxa matrix using both ML and BI. The Grylloblattodea, Mantophasmatodea, and Phasmatodea were all well supported using the optimized_taxa matrix. For the all_taxa matrix also the Embioptera were recovered in this clade as sister taxa to the Phasmatodea excluding *Timema*

271

272

273

274

275

276

277

278

279

280

281

282

283

284

285

286

287

288

289

290

291

292

293

294

californicum (Figure 1, Additional files 2, 4: Figures S1, S3). The superorder Dictyoptera was monophyletic clade, consisting of the monophyletic Mantodea and the Blattodea including the Isoptera. Mitochondrial genomes The mitochondrial genome sequences have been deposited at GenBank with the accession numbers XXXXXXXX, XXXXXXXX, and XXXXXXXX. The pyrosequencing run of C. dipterum using the 454 GS FLX system resulted in 651,306 reads, of which 1.14% mapped to the assembled mitochondrial genome. The depth of coverage for the C. dipterum mitochondrial genome was 249.2x (± 80.6 SD, Figure 2a). The three mayfly mitochondrial genomes were 15,407 bp (C. dipterum), 14,883 bp (B. rutilocylindratus), and 14,355 bp (H. zijinensis) long (Table 4). For B. rutilocylindratus, the complete mitochondrial genome was sequenced (Figure 2b). The AT-rich regions were incomplete for genomes of C. dipterum and H. zijinensis. The three tRNAs between the AT-rich region and *nad2* were also missing in *H. zijinensis* due to the incomplete sequencing (Figure 2c). All three sequenced mitochondrial genomes contained the entire set of 13 PCGs, two rRNAs, and either 22 tRNAs (B. rutilocylindratus, C. dipterum) or 19 for the incomplete mitochondrial genome of *H. zijinensis*, including 23 coded at the (+) strand (21 for *H. zijinensis*) and 14 at the (-) strand (13 for *H. zijinensis*) (Figure 2). The order of the genes was conserved, with the exceptions of the two inverted tRNAs Arginine and Alanine for H. zijinensis (Figure 2c). All PCGs started with the ATN codons (ATT, ATG, ATA),

The order of the genes was conserved, with the exceptions of the two inverted tRNAs Arginine and Alanine for H. zijinensis (Figure 2c). All PCGs started with the ATN codons (ATT, ATG, ATA), and mostly ended with the complete termination codon (TAA or TAG). Exceptions were an incomplete T termination codon for nad4 in B. rutilocylindratus and cox1 started with CTC in C. dipterum. The AT content of the mitochondrial genomes that were corrected for the missing regions, was lowest in B. rutilocylindratus (59.9%, Table 4, Additional file 5: Table S2). The mean AT content for all available mayfly mitochondrial genomes was 65.34%. The average AT-skew was -0.03 (\pm 0.04 SD) and the average GC-skew was -0.13 (\pm 0.14 SD).

Table 4 Overview of sequenced ephemeropteran mitochondrial genomes. Total sequence length in base pairs (bp) and individual nucleotide compositions of mayfly mitochondrial genomes calculated based on the whole available sequences and the ones corrected for the incomplete mitochondrial genomes (see Methods; for the complete list of all ephemeropteran mitochondrial genomes see Additional file 5: Table S2)

Family	Species	Length [bp]	A%	C%	G%	Т%	GC%	AT%
Baetidae	Baetis	14,883 ^a	27.3	19.8	20.1	32.8	39.9	60.1
	rutilocylindratus	12,234	26.7	19.8	20.4	33.2	40.2	59.9
Baetidae	Classes directory	14,355	30.6	14.7	16.3	38.4	30.9	69.0
	Cloeon dipterum	12,197	29.9	14.5	16.9	38.7	31.4	68.6
Leptophlebiidae	Habrophlebiodes	15,407	33.8	20.2	11.0	35.1	31.2	68.9
	zijinensis	12,417	33.6	20.6	11.5	34.3	32.1	67.9

^a, Indicates complete mitochondrial genomes; rest are incomplete mitochondrial genomes.

Discussion

Phylogenetic relationships of Palaeoptera

Our Bayesian reconstruction based on the optimized_taxa matrix provided strong support for the basal Odonata hypothesis. Previous work based on mitochondrial genome data is (partially) congruent with our findings [10,22,23]. In contrast to our results, a handful of studies based on nucleotide and aa sequences of mitochondrial PCGs and mitochondrial RNA (mtRNAs) found overall strong support for the basal Ephemeroptera hypothesis [10,16,17,20]. Interestingly, Song et al. [10], supporting overall the Ephemeroptera hypothesis, found evidence for the Odonata hypothesis when using aa sequence data. Our results confirm the trend towards a better support for the ancestral position of the Odonata when using an increased mayfly taxon sampling (more than ten species per lineage) [24,25,27]. The exceptions for this are the work by Ogden and Whiting [2] and

Thomas et al. [12], that overall support the Ephemeroptera respective Palaeoptera hypothesis but contain some data sets in favor of the Odonata hypothesis. Early inconsistencies based on different sequence alignment strategies [2] appear to become less relevant due to the advances of multiple sequence alignment programs. Overall, the vast majority of studies were based on BI and ML. Thomas et al. [12] recovered increased resolution using BI in comparison to ML. Different data partitioning (i.e., gene trees resulting in different tree topologies) by Kjer et al. [11] supported different hypotheses for different genes. Evidently, using rRNAs (nDNA and mtDNA) and the nDNA EF1-α the Odonata hypothesis was recovered whereas the analyses based on the combined and on the mtDNA data matrix resulted in a monophyletic palaeopteran clade. It is questionable whether an increase in analyzed genomic data (i.e., phylogenomic studies) will be able to resolve the Palaeoptera problem. Notably, Misof et al. [79] and Regier et al. [15], using 1,478 PCGs and seven palaeopteran species, respective 62 PCGs and four palaeoptera species, recovered the Ephemeroptera and Odonata as sister clades (i.e., the Palaeoptera hypothesis) but without high support values. Overall, more knowledge about the use and limitation of individual markers are needed. For example Simon et al. [80] found that proteins involved in cellular processes and signaling harbor the most phylogenetic signal. The increased mayfly taxon sampling highlighted the phylogenetic diversity (as indicated by the branch lengths of the tip taxa, Figure 1, Additional files 2-4: Figures S1-S3) of the Palaeoptera. The overall long-branch lengths and high rate-heterogeneity across sites might be one of the explanations for the "Palaeoptera problem". Especially the family of the Baetidae, which was long thought to be the most ancestral mayfly family [81], showed high base composition heterogeneity possibly evidencing LBA. However, in agreement with other more recent studies we also recovered S. chinensis as sister taxa to all other mayflies [55,82]. It remains to be resolved whether an increased taxon sampling may obviate LBA sensu Bergsten [65].

Phylogenetic relationships of Polyneoptera

314

315

316

317

318

319

320

321

322

323

324

325

326

327

328

329

330

331

332

333

334

335

336

337

340

341

342

343

344

345

346

347

348

349

350

351

352

353

354

355

356

357

358

359

360

361

362

Besides the Palaeoptera, a universal consensus on the placement of the Plecoptera, Dermaptera, Embioptera, and Xenoptera (i.e., Grylloblattodea and Mantophasmatodea) remains elusive [10]. The monophyly of the Polyneoptera has become more widely accepted [5,10,23,79,83,84], although several studies and data matrices do not support their monophyly [10,11,80]. This is also reflected by our results, recovering the Polyneoptera as moderately supported monophyletic clade when using the optimized taxa matrix with ML. The placement of the order Plecoptera also remains elusive. The order often clusters among the more ancestral polyneopteran orders [10,23,79] and in a close relationship to the Dermaptera, although earlier studies using few species and the aa sequences of the mitochondrial PCGs also recovered a sister relationship to the Ephemeroptera [22,85]. Notably, Song et al. [10] was the only study including four dermapteran species. The monophyly of the Orthoptera has been established by previous studies based on mitochondrial genome data and a large number of PCGs [10,79,86,87]. However, in a recent study, the Ensifera also clustered as more ancestral clade [10]. As for this study, Song et al. [10] also used BI and ML based on mitochondrial genome data. Notably, Song et al. [10] explain the more ancestral position of Ensifera by more similar sequence compositional bias in comparison to the outgroups. The relationship between Grylloblattodea and Mantophasmatodea (i.e., Xenoptera superorder) remains elusive. Mostly this is due to the limited amount of available molecular data, including ordinal-level phylogenies based on two [5,10] or three xenopteran species [79]. Using mitochondrial genome data and one species per order, also Song et al. [10] did not recover them as a monophyletic clade. Their monophyly was supported by previous studies using one Mantophasmatodea and one Grylloblattodea species [5], respective one Mantophasmatodea and two Grylloblattodea species [79]). The Phasmatodea have also been found in previous studies as paraphyletic clade due to the outside position of T. californicum [10,16]. In contrast, Misof et al. [79], using a large set of nuclear PCGs also found *Timema* as the sister species to all other representatives of the order. The monophyly of the superorder Dictyoptera is generally

accepted and the hierarchical clustering (Mantodea + (Blattodea + Isoptera)) has been well supported [5,10,11,16,23,79,88,89] using nucleotide and as sequences of both mtDNA and nDNA.

Optimizing data matrix

363

364

365

366

367

368

369

370

371

372

373

374

375

376

377

378

379

380

381

382

383

384

385

386

387

The optimization of the data matrix overall resulted in a better-supported tree (as measured by the total number of supported nodes and the fact that all nodes supported in the all taxa matrix were also recovered in the optimized taxa matrix). In a study on butterfly phylogenetics, the authors also removed rogue taxa and found dramatically increased bootstrap support values [90]. Comparing the number of supported nodes from the BI and ML approach, the former always resulted in more resolved nodes. This result is consistent with previous work on the "Palaeoptera problem", recovering more support for analyses using BI [12]. Generally, the choice of the outgroup species was reported as being crucial for resolving problematic splits in the tree of life such as insects origin [12]. Thus, finding the outgroup as a rogue taxon was perhaps not surprising. The close phylogenetic relationship of the Dermaptera and Ephemeroptera (Additional files 2, 4: Figures S1, S3) might be misleading due to LBA. Li et al. [55] also found the order Dermaptera as being closely related to the mayflies. On the other hand studies based on phylogenomic data reported the Dermaptera as sister taxa to the orders Plecoptera [5,16,23,80,91] Zoraptera [28,79]. Using mitochondrial genome data and an increased taxon sampling (i.e., four dermapteran species), Song et al. [10] found the Dermaptera and/or Plecoptera being sister to the remaining taxa within Polyneoptera. Notably, when only using one species of Dermaptera, C. fletcheri emerged as sister taxa to the Ephemeroptera [10]. This result in agreement with our findings is presumably due to LBA. Phylogenomic data matrices (e.g., expressed sequence tags data: [83], 1,478 PCGs: [79]) and nuclear genes [5] found the Embioptera to be sister clade to the Phasmatodea. Using mitochondrial genomes data and BI, the Embioptera were also found as sister group to the Phasmatodea (excluding *T. californicum*) [10]. However, in the same study, they also found support for the sister-relationship of Embioptera-Zoraptera using ML [10]. Overall, more phylogenetic studies will be needed to clarify this issue; mostly also because the use of few taxa from anomalous orders (i.e., Dermaptera and Embioptera) tend to evoke LBA [10].

Characterization of ephemeropteran mitochondrial genomes

388

389

390

391

392

393

394

395

396

397

398

399

400

401

402

403

404

405

406

407

408

409

410

411

412

Mayfly mitochondrial genomes are widely conserved in gene content and nucleotide composition across eleven families, with only a few differences in the content of their tRNAs. The coverage of the Cloeon-mitochondrial genome was in agreement with other studies, using the same sequencing platform (e.g., 59-281x, [92]). The missing of part of the AT-rich region is due to reduced sequencing and assembly efficiency of this low complexity region and common among insect mitochondrial genomes [55,93]. The gene order and orientation were with the exception of two shifted tRNAs for *H. zijinensis* identical to the ancestral insect mitochondrial genome [41,42,54]. Other mayflies are also reported to miss complete T termination codons in the genes cox2 and nad5 [55,93,94]. The two rRNAs were located between trnL and trnV (rrnL), and between trnV and the AT-rich region (rrnS), respectively. All ephemeropteran mitochondrial genomes contain an AT-rich region, which is placed between the rrnS (- strand) and trnI (+ strand). Li et al. [55] reported two distinct parts within the AT-rich region in Siphluriscus chinensis, which also seems to be present in C. dipterum. Therein, they described the so called CR₁ which is located close to the rrnS and has a high AT content (71.6%), including six identical 140 bp sequences, and the CR₂, which is close to the trnI and has a lower AT content (58.1%). E. orientalis contains two identical 55 bp long sequences in the AT-rich region [94]. Few mayfly mitochondrial genomes differ in their gene content from the ancestral insect mitochondrial genome, possessing one additional tRNA. The two heptageniid species Parafronurus youi and Epeorus sp. encode a second copy of the trnM (AUG, trnM2) gene located between trnI and trnO [17,93]. For S. chinensis, an additional trnK2 (AAA) gene is described [55]. Overall, Song et al. [10] found similar genetic distances for the early branching insects lineages. The AT contents for the Ephemeroptera were very similar to the ones of Odonata (62.6%-68.5% for *cox1*, [95]).

414

415

416

417

418

419

420

421

422

423

424

425

426

427

428

429

430

431

432

433

434

435

Conclusions Our analysis based on an increased mayfly taxon sampling and data matrix optimization supported the Odonata hypothesis and unravelled fine structural changes within the palaeopteran mitochondrial genomes. While both BI and ML overall resulted in highly supported trees, only the BI based on an optimized taxa matrix highly supported the sister-relationship of the Ephemeroptera and Neoptera. The optimized taxa matrix, excluding rogue taxa and taxa with LBA, resulted in an overall better supported tree (as measured by the number of supported nodes) for both BI and ML. Our findings highlight the essential need to increase the taxon sampling of under-represented lineages (such as the Palaeoptera) in order to resolve their phylogenetic position. Here we demonstrated the need for an increased taxon sampling in combination with data matrix optimization, and the use of different phylogenetic approaches in order to resolve an ancient radiation. Establishing general recommendation for data matrix optimization requires additional analyses on a broader range of lineages. **Declarations** Ethics approval and consent to participate Not applicable. **Consent for publication** Not applicable. Availability of data and material Newly generated sequences are available on GenBank (XXXXXXXX, XXXXXXXX, and XXXXXXXX).

437

438

439

440

441

442

443

444

445

446

447

448

449

450

451

452

453

454

455

456

Competing interests The authors declare that they have no competing interests. **Funding** Research was funded by the Leibniz Association PAKT für Forschung und Innovation ('FREDIE' project: SAW-2011-ZFMK-3). Authors' contributions SR and MTM conceived the study; SR sequenced and analyzed the *Cloeon dipterum* genome, and performed all combined analyses; PC and CZ obtained and analyzed the Baetis rutilocylindratus and Habrophlebiodes zijinensis genomes; SR and MTM wrote the manuscript. All authors made contributions to subsequent revisions and agreed to the final version. Acknowledgements We are grateful to D. H. Funk from the Stround Water Research Center for providing the Cloeon dipterum specimens, S. Mbedi, K. Preuß, and L. Wächter for their great help with laboratory work, G. Glöckner, and C. Mazzoni for constructive comments about the analysis of the mitochondrial genomes, and Zedat-HPC at the Freie Universität Berlin, Germany for providing access to highperformance computing clusters. SR thanks the Janggen-Pöhn-Stiftung (http://www.janggenpoehn.ch/) for a Postdoctoral stipend. We also want to thank the members of our research groups for constructive working environments.

<u>Tables</u> Table 1 Set of mitochondrial genomes with according GenBank accession numbers

457

ArchaeognathaMachilidaePedetontus silvestriiNC_011717ArchaeognathaMachilidaePetrobius brevistylisNC_007688ArchaeognathaMachilidaeSongmachilis xinxiangensisNC_021384ArchaeognathaMachilidaeTrigoniophthalmus alternatusNC_010532ArchaeognathaMeinertellidaeNesomachilis australicaNC_006895BlattodeaBlattidaePeriplaneta americanaNC_016956BlattodeaBlattidaePeriplaneta fuliginosaNC_006076BlattodeaCorydiidaeEupolyphaga sinensisNC_014274BlattodeaCryptocercidaeCryptocercus relictusNC_018132BlattodeaEctobiidaeBlattella bisignataNC_018549BlattodeaEctobiidaeBlattella germanicaNC_012901CollembolaTetrodontophorinaeTetrodontophora bielanensis³NC_02735DermapteraPygidicranidaeChallia fletcheri³NC_018538EmbiopteraOligotomidaeAposthonia japonica³AB639034EphemeropteraBaetidaeAlainites yixianiNC_020034EphemeropteraBaetidaeBaetis rutilocylindratusXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	Order	Family	Species	Accession
Archaeognatha Machilidae Songmachilis xinxiangensis NC_021384 Archaeognatha Machilidae Trigoniophthalmus alternatus NC_010532 Archaeognatha Meinertellidae Nesomachilis australica NC_006895 Blattodea Blattidae Periplaneta americana NC_016956 Blattodea Blattidae Periplaneta fuliginosa NC_006076 Blattodea Corydiidae Eupolyphaga sinensis NC_014274 Blattodea Cryptocercidae Cryptocercus relictus NC_018132 Blattodea Ectobiidae Blattella bisignata NC_018549 Blattodea Ectobiidae Blattella germanica NC_012901 Collembola Tetrodontophorinae Tetrodontophora bielanensis ^a NC_002735 Dermaptera Pygidicranidae Challia fletcheri ^a NC_018538 Embioptera Oligotomidae Aposthonia japonica ^a AB639034 Ephemeroptera Baetidae Ameletus spl KM244682 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Archaeognatha	Machilidae	Pedetontus silvestrii	NC_011717
Archaeognatha Machilidae Trigoniophthalmus alternatus NC_010532 Archaeognatha Meinertellidae Nesomachilis australica NC_006895 Blattodea Blattidae Periplaneta americana NC_016956 Blattodea Blattidae Periplaneta fuliginosa NC_006076 Blattodea Corydiidae Eupolyphaga sinensis NC_014274 Blattodea Cryptocercidae Cryptocercus relictus NC_018132 Blattodea Ectobiidae Blattella bisignata NC_018549 Blattodea Ectobiidae Blattella germanica NC_012901 Collembola Tetrodontophorinae Tetrodontophora bielanensis NC_02735 Dermaptera Pygidicranidae Challia fletcheri NC_018538 Embioptera Oligotomidae Aposthonia japonica AB639034 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Archaeognatha	Machilidae	Petrobius brevistylis	NC_007688
ArchaeognathaMeinertellidaeNesomachilis australicaNC_006895BlattodeaBlattidaePeriplaneta americanaNC_016956BlattodeaBlattidaePeriplaneta fuliginosaNC_006076BlattodeaCorydiidaeEupolyphaga sinensisNC_014274BlattodeaCryptocercidaeCryptocercus relictusNC_018132BlattodeaEctobiidaeBlattella bisignataNC_018549BlattodeaEctobiidaeBlattella germanicaNC_012901CollembolaTetrodontophorinaeTetrodontophora bielanensis³NC_002735DermapteraPygidicranidaeChallia fletcheri³NC_018538EmbiopteraOligotomidaeAposthonia japonica³AB639034EphemeropteraBaetidaeAlainites yixianiNC_020034EphemeropteraBaetidaeBaetis rutilocylindratusXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	Archaeognatha	Machilidae	Songmachilis xinxiangensis	NC_021384
Blattidae Blattidae Periplaneta americana NC_016956 Blattodea Blattidae Periplaneta fuliginosa NC_006076 Blattodea Corydiidae Eupolyphaga sinensis NC_014274 Blattodea Cryptocercidae Cryptocercus relictus NC_018132 Blattodea Ectobiidae Blattella bisignata NC_018549 Blattodea Ectobiidae Blattella germanica NC_012901 Collembola Tetrodontophorinae Tetrodontophora bielanensisa NC_002735 Dermaptera Pygidicranidae Challia fletcheria NC_018538 Embioptera Oligotomidae Aposthonia japonica AB639034 Ephemeroptera Baetidae Ameletus spl KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Archaeognatha	Machilidae	Trigoniophthalmus alternatus	NC_010532
Blattodea Blattidae Periplaneta fuliginosa NC_006076 Blattodea Corydiidae Eupolyphaga sinensis NC_014274 Blattodea Cryptocercidae Cryptocercus relictus NC_018132 Blattodea Ectobiidae Blattella bisignata NC_018549 Blattodea Ectobiidae Blattella germanica NC_012901 Collembola Tetrodontophorinae Tetrodontophora bielanensis³ NC_002735 Dermaptera Pygidicranidae Challia fletcheri³ NC_018538 Embioptera Oligotomidae Aposthonia japonica³ AB639034 Ephemeroptera Baetidae Ameletus sp1 KM244682 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Archaeognatha	Meinertellidae	Nesomachilis australica	NC_006895
Blattodea Corydiidae Eupolyphaga sinensis NC_014274 Blattodea Cryptocercidae Cryptocercus relictus NC_018132 Blattodea Ectobiidae Blattella bisignata NC_018549 Blattodea Ectobiidae Blattella germanica NC_012901 Collembola Tetrodontophorinae Tetrodontophora bielanensisa NC_002735 Dermaptera Pygidicranidae Challia fletcheria NC_018538 Embioptera Oligotomidae Aposthonia japonicaa AB639034 Ephemeroptera Ameletidae Ameletus sp1 KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Blattodea	Blattidae	Periplaneta americana	NC_016956
Blattodea Cryptocercidae Cryptocercus relictus NC_018132 Blattodea Ectobiidae Blattella bisignata NC_018549 Blattodea Ectobiidae Blattella germanica NC_012901 Collembola Tetrodontophorinae Tetrodontophora bielanensis³ NC_002735 Dermaptera Pygidicranidae Challia fletcheri³ NC_018538 Embioptera Oligotomidae Aposthonia japonica³ AB639034 Ephemeroptera Ameletidae Ameletus sp1 KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Blattodea	Blattidae	Periplaneta fuliginosa	NC_006076
Blattodea Ectobiidae Blattella bisignata NC_018549 Blattodea Ectobiidae Blattella germanica NC_012901 Collembola Tetrodontophorinae Tetrodontophora bielanensisa NC_002735 Dermaptera Pygidicranidae Challia fletcheria NC_018538 Embioptera Oligotomidae Aposthonia japonica AB639034 Ephemeroptera Ameletidae Ameletus spl KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Blattodea	Corydiidae	Eupolyphaga sinensis	NC_014274
Blattodea Ectobiidae Blattella germanica NC_012901 Collembola Tetrodontophorinae Tetrodontophora bielanensisª NC_002735 Dermaptera Pygidicranidae Challia fletcheriª NC_018538 Embioptera Oligotomidae Aposthonia japonicaª AB639034 Ephemeroptera Ameletidae Ameletus spl KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Blattodea	Cryptocercidae	Cryptocercus relictus	NC_018132
Collembola Tetrodontophorinae Tetrodontophora bielanensis ^a NC_002735 Dermaptera Pygidicranidae Challia fletchert ^a NC_018538 Embioptera Oligotomidae Aposthonia japonica ^a AB639034 Ephemeroptera Ameletidae Ameletus sp1 KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Blattodea	Ectobiidae	Blattella bisignata	NC_018549
Dermaptera Pygidicranidae Challia fletcheri ^a NC_018538 Embioptera Oligotomidae Aposthonia japonica ^a AB639034 Ephemeroptera Ameletidae Ameletus sp1 KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Blattodea	Ectobiidae	Blattella germanica	NC_012901
Embioptera Oligotomidae Aposthonia japonica ^a AB639034 Ephemeroptera Ameletidae Ameletus sp1 KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Collembola	Tetrodontophorinae	Tetrodontophora bielanensis ^a	NC_002735
Ephemeroptera Ameletidae Ameletus sp1 KM244682 Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Dermaptera	Pygidicranidae	Challia fletcheri ^a	NC_018538
Ephemeroptera Baetidae Alainites yixiani NC_020034 Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Embioptera	Oligotomidae	Aposthonia japonica ^a	AB639034
Ephemeroptera Baetidae Baetis rutilocylindratus XXXXXXXX Ephemeroptera Baetidae Cloeon dipterum XXXXXXXXX Ephemeroptera Caenidae Caenis pycnacantha GQ502451 Ephemeroptera Ephemerellidae Ephemerella sp. KM244691 Ephemeroptera Ephemerellidae Vietnamella dabieshanensis NC_020036	Ephemeroptera	Ameletidae	Ameletus sp1	KM244682
EphemeropteraBaetidaeCloeon dipterumXXXXXXXXEphemeropteraCaenidaeCaenis pycnacanthaGQ502451EphemeropteraEphemerellidaeEphemerella sp.KM244691EphemeropteraEphemerellidaeVietnamella dabieshanensisNC_020036	Ephemeroptera	Baetidae	Alainites yixiani	NC_020034
EphemeropteraCaenidaeCaenis pycnacanthaGQ502451EphemeropteraEphemerellidaeEphemerella sp.KM244691EphemeropteraEphemerellidaeVietnamella dabieshanensisNC_020036	Ephemeroptera	Baetidae	Baetis rutilocylindratus	XXXXXXXX
EphemeropteraCaenidaeCaenis pycnacanthaGQ502451EphemeropteraEphemerellidaeEphemerella sp.KM244691EphemeropteraEphemerellidaeVietnamella dabieshanensisNC_020036	Ephemeroptera	Baetidae	Cloeon dipterum	XXXXXXXX
Ephemeroptera Ephemerellidae <i>Vietnamella dabieshanensis</i> NC_020036	Ephemeroptera	Caenidae		GQ502451
	Ephemeroptera	Ephemerellidae	Ephemerella sp.	KM244691
Ephemeroptera Ephemerellidae <i>Vietnamella</i> sp. KM244655	Ephemeroptera	Ephemerellidae	Vietnamella dabieshanensis	NC_020036
	Ephemeroptera	Ephemerellidae	Vietnamella sp.	KM244655
Ephemeroptera Ephemeridae Ephemera orientalis NC_012645	Ephemeroptera	Ephemeridae	Ephemera orientalis	NC_012645
Ephemeroptera Heptageniidae <i>Epeorus</i> sp. KM244708	Ephemeroptera	Heptageniidae	Epeorus sp.	KM244708
Ephemeroptera Heptageniidae Paegniodes cupulatus NC_020035	Ephemeroptera	Heptageniidae	Paegniodes cupulatus	NC_020035
Ephemeroptera Heptageniidae <i>Parafronurus youi</i> NC_011359	Ephemeroptera	Heptageniidae	Parafronurus youi	NC_011359
Ephemeroptera Isonychiidae Isonychia ignota NC_020037	Ephemeroptera	Isonychiidae	Isonychia ignota	NC_020037
Ephemeroptera Leptophlebiidae <i>Habrophlebiodes zijinensis</i> XXXXXXX	Ephemeroptera	Leptophlebiidae	Habrophlebiodes zijinensis	XXXXXXXX
Ephemeroptera Potamanthidae <i>Potamanthus</i> sp. KM244674	Ephemeroptera	Potamanthidae	Potamanthus sp.	KM244674
Ephemeroptera Siphlonuridae Siphlonurus immanis NC_013822	Ephemeroptera	Siphlonuridae	Siphlonurus immanis	NC_013822
Ephemeroptera Siphlonuridae Siphlonurus sp. KM244684	Ephemeroptera	Siphlonuridae	Siphlonurus sp.	KM244684
Ephemeroptera Siphluriscidae Siphluriscus chinensis HQ875717	Ephemeroptera	Siphluriscidae	Siphluriscus chinensis	HQ875717
KM244670;	Enhamantana	Talaganadidaa	-	KM244670;
Ephemeroptera Teloganodidae sp. KM244703 ^b	Epnemeropiera	reioganodidae	sp.	KM244703 ^b
Grylloblattodea Grylloblattidae Grylloblatta sculleni DQ241796	Grylloblattodea	Grylloblattidae	Grylloblatta sculleni	DQ241796
Isoptera Hodotermitidae <i>Microhodotermes viator</i> NC_018122	Isoptera	Hodotermitidae	Microhodotermes viator	NC_018122
Isoptera Kalotermitidae <i>Neotermes insularis</i> NC_018124	Isoptera	Kalotermitidae	Neotermes insularis	
Isoptera Rhinotermitidae Coptotermes lacteus NC_018125	-	Rhinotermitidae	Coptotermes lacteus	_
Isoptera Rhinotermitidae <i>Heterotermes</i> sp. NC_018127	-	Rhinotermitidae		_

Isoptera	Rhinotermitidae	Reticulitermes flavipes	NC 009498
Isoptera	Rhinotermitidae	Reticulitermes hageni	NC 009501
Isoptera	Rhinotermitidae	Schedorhinotermes breinli	NC 018126
Isoptera	Termitidae	Drepanotermes sp.	NC_018129
Isoptera	Termitidae	Macrognathotermes errator	NC 018130
Isoptera	Termitidae	Macrotermes subhyalinus	NC_018128
Isoptera	Termitidae	Nasutitermes triodiae	NC 018131
Isoptera	Termopsidae	Porotermes adamsoni	NC 018121
Isoptera	Termopsidae	Zootermopsis nevadensis	NC_024658
Mantodea	Caliridinae	Leptomantella albella	NC_024028
Mantodea	Mantidae	Tamolanica tamolana	NC_007702
Mantophasmatodea	Mantophasmatidae	Sclerophasma paresisensis	NC_007701
Odonata	Calopterygidae	Vestalis melania	NC_023233
Odonata	Coenagrionidae	Ischnura pumilio	NC_021617
Odonata	Corduliidae	Cordulia aenea	JX963627
Odonata	Epiophlebiidae	Epiophlebia superstes	NC_023232
Odonata	Euphaeidae	Euphaea formosa	NC_014493
Odonata	Gomphidae	Davidius lunatus	NC 012644
Odonata	Gomphidae	Ictinogomphus sp.	KM244673
Odonata	Libellulidae	Hydrobasileus croceus	KM244659
Odonata	Libellulidae	Orthetrum triangulare melania	AB126005
Odonata	Pseudolestidae	Pseudolestes mirabilis	NC 020636
Orthoptera	Acrididae	Acrida willemsei	NC_011303
Orthoptera	Acrididae	Calliptamus italicus	NC_011305
Orthoptera	Acrididae	Gomphocerus sibiricus	NC_021103
Orthoptera	Acrididae	Oedaleus decorus asiaticus	NC 011115
Orthoptera	Gryllotalpidae	Gryllotalpa orientalis ^a	NC 006678
Orthoptera	Pneumoridae	Tanaocerus koebelei	NC_020777
Orthoptera	Prophalangopsidae	Tarragoilus diuturnus	NC 021397
Orthoptera	Pyrgomorphidae	Atractomorpha sinensis	NC 011824
Orthoptera	Rhaphidophoridae	Troglophilus neglectus	NC 011306
Orthoptera	Romaleidae	Xyleus modestus	NC 014490
Orthoptera	Tetrigidae	Tetrix japonica	NC 018543
Orthoptera	Tettigoniidae	Gampsocleis gratiosa	NC_011200
Orthoptera	Tettigoniidae	Ruspolia dubia	NC 009876
Phasmatodea	Bacillidae	Bacillus rossius	GU001956
Phasmatodea	Diapheromeridae	Micadina phluctainoides	NC_014673
Phasmatodea	Diapheromeridae	Sipyloidea sipylus	AB477470
Phasmatodea	Heteropterygidae	Heteropteryx dilatata	NC_014680
Phasmatodea	Heteropterygidae	Orestes mouhotii	AB477462
Phasmatodea	Phasmatidae	Entoria okinawaensis	NC_014694
Phasmatodea	Phasmatidae	Extatosoma tiaratum	NC_017748
Phasmatodea	Phasmatidae	Megacrania alpheus adan	NC_014688

Phasmatodea	Phasmatidae	Neohirasea japonica	AB477469
Phasmatodea	Phasmatidae	Phobaeticus serratipes	NC_014678
Phasmatodea	Phasmatidae	Phraortes sp. ^a	NC_014705
Phasmatodea	Phasmatidae	Ramulus irregulariterdentatus	NC_014702
Phasmatodea	Timematidae	Timema californicum	DQ241799
Plecoptera	Perlidae	Dinocras cephalotes	NC_022843
Plecoptera	Perlidae	Kamimuria wangi	NC_024033
Plecoptera	Perlidae	Togoperla sp.	KM409708
Plecoptera	Pteronarcyidae	Pteronarcys princeps	NC_006133
Zygentoma	Lepidotrichidae	Tricholepidion gertschi	NC_005437
Zygentoma	Lepismatidae	Thermobia domestica	NC_006080
Zygentoma	Nicoletiidae	Atelura formicaria	NC_011197

<sup>a, Taxa were excluded in the optimized_taxa matrix (see Methods).
b, Represents sequence consisting of two genomic fragments (contigs) from the same sample.</sup>

Figures

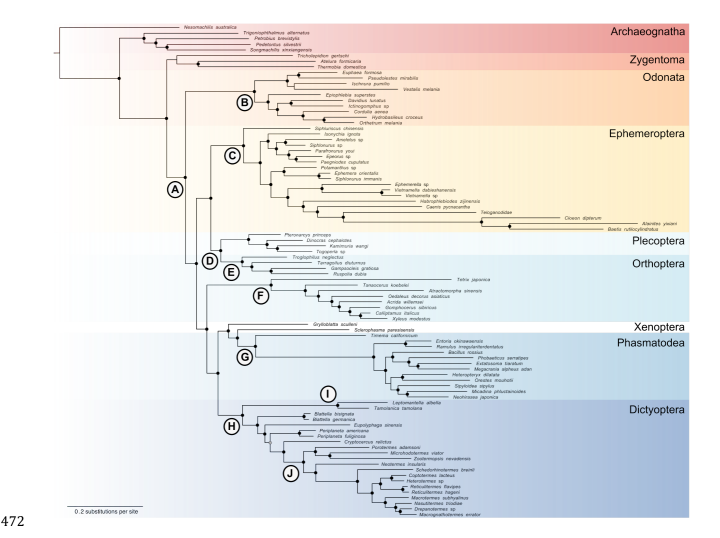
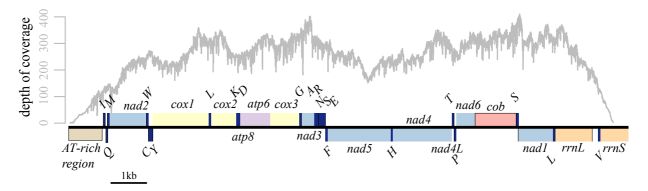
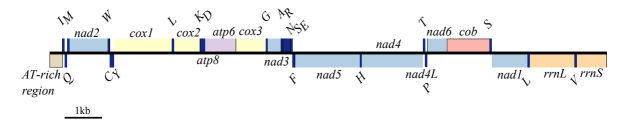


Fig. 1 Phylogenetic relationships of insect major orders using Bayesian inference reconstruction based on mitochondrial genome data using the concatenated amino acid sequences of the optimized_taxa matrix (see Table 1). Filled circles indicate supported nodes; whereby black circles represent Bayesian posterior probability (BPP) = 1, and grey circles BPP \geq 0.95. Letters indicate clades mentioned in the text (see also Table 3)

(a) Cloeon dipterum L. 1761



(b) Baetis rutilocylindratus Wang, Qin, Cheng & Zhou, 2011



(c) Habrophlebiodes zijinensis Gui, Zhang & Wu, 1996

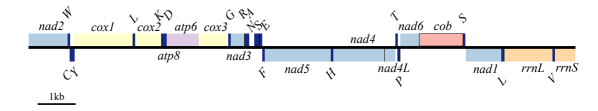


Fig. 2 Mitochondrial genome maps of three sequenced Ephemeroptera species. The nearly complete mitochondrial genome of (a) *Cloeon dipterum*, including coverage depth, the complete mitochondrial genome of (b) *Baetis rutilocylindratus*, and the incomplete mitochondrial genome of (c) *Habrophlebiodes zijinensis*. Transfer RNA genes are indicated by single-letter IUPAC-IUB abbreviations for their corresponding amino acid. Protein coding genes and ribosomal RNA genes are listed and colored in the following way: *atp6*, *atp8*, ATP synthase subunits 6 and 8 genes (violet); *cob*, cytochrome oxidase *b* gene (red); *cox1-cox3*, cytochrome oxidase *c* subunit 1-3 genes (yellow); *nad1-6*, *nad4L*, NADH dehydrogenase subunits 1-6 and 4L (light blue); *rrnS*, *rrnL*, small and large ribosomal RNA subunits (orange); AT-rich region (brown). Genes located at the (+) strand appear above the central line. Genes located on the (-) strand appear below the central line.

491

492

493

494

495

496

497

498

499

500

501

502

503

504

505

506

507

508

509

510

511

512

513

Additional files Additional file 1: Table S1. Universal primers for Sanger sequencing. List of primer pairs used for the PCR amplification and Sanger sequencing of Baetis rutilocylindratus (Baetidae) and Habrophlebiodes zijinensis (Leptophlebiidae). Additional file 2: Fig. S1. Phylogenetic relationships of insect major orders using Bayesian inference reconstruction based on mitochondrial genome data using the concatenated amino acid sequences of the all taxa matrix (see Table 1). Filled circles indicate well-supported nodes; whereby black circles represent Bayesian posterior probability (BPP) = 1, and grey circles BPP ≥ 0.95 . Scale bar indicates substitutions per site. Coloured tip labels refer to the orders, whereby light blue = Archaeognatha (AR), dark blue = Blattodea (BL), light green = Ephemeroptera (EP), dark green = Grylloblattodea (GR), red = Isoptera (IS), light orange = Mantodea (MT), dark red = Mantophasmatodea (MP), dark orange = Odonata (OD), light purple = Orthoptera (OR), dark purple = Phasmatodea, yellow = Plecoptera (PL), and brown = Zygentoma (ZY). Five taxa being excluded from the optimized taxa matrix are coloured in dark grey. Additional file 3: Fig. S2. Phylogenetic relationships of insect major orders using maximum likelihood reconstruction based on mitochondrial genome data using the concatenated amino acid sequences of the optimized taxa matrix (see Table 1). Filled circles indicate well-supported nodes; whereby black circles represent Bootstrap Support (BS) \geq 90, and grey circles BS \geq 80. Scale bar indicates substitutions per site. Coloured tip labels refer to the orders (for details see Additional file 2: Figure S1). Additional file 4: Fig. S3. Phylogenetic relationships of insect major orders using maximum likelihood reconstruction based on mitochondrial genome data using the concatenated amino acid sequences of the all taxa matrix (see Table 1). Filled circles indicate well-supported nodes; whereby black circles represent Bootstrap Support (BS) \geq 90, and grey circles BS \geq 80. Scale bar indicates

- substitutions per site. Coloured tip labels refer to the orders with the exception of taxa being
- excluded from the optimized taxa matrix coloured in dark grey (for details see Additional file 2:
- 516 Figure S1).
- Additional file 5: Table S2. Overview of ephemeropteran mitochondrial genomes. Total sequence
- length in base pairs (bp) and individual nucleotide compositions of mayfly mitochondrial genomes
- 519 calculated based on the whole available sequences and the ones corrected for the incomplete
- mitochondrial genomes (see Methods).

References

521

- 1. Hovmöller R. The Palaeoptera Problem: Basal Pterygote Phylogeny Inferred from 18S and 28S
- 524 rDNA Sequences. Cladistics. 2002;18:313–23.
- 2. Ogden HT, Whiting MF. The problem with "the Paleoptera Problem:" sense and sensitivity.
- 526 Cladistics. 2003;19:432–42.
- 3. Trautwein MD, Wiegmann BM, Beutel R, Kjer KM, Yeates DK. Advances in Insect Phylogeny at
- the Dawn of the Postgenomic Era. Annu. Rev. Entomol. 2012;57:449–68.
- 4. Whitfield JB, Kjer KM. Ancient Rapid Radiations of Insects: Challenges for Phylogenetic
- 530 Analysis. Annu. Rev. Entomol. 2008;53:449–72.
- 5. Ishiwata K, Sasaki G, Ogawa J, Miyata T, Su Z-H. Phylogenetic relationships among insect orders
- based on three nuclear protein-coding gene sequences. Mol. Phylogenet. Evol. 2011;58:169–80.
- 6. Yeates DK, Cameron SL, Trautwein M. A view from the edge of the forest: recent progress in
- understanding the relationships of the insect orders. Australian Journal of Entomology. 2012;51:79–
- 535 87.
- 7. Kjer KM, Simon C, Yavorskaya M, Beutel RG. Progress, pitfalls and parallel universes: a history
- of insect phylogenetics. J R Soc Interface. 2016;13:20160363.
- 8. Mallatt J, Giribet G. Further use of nearly complete 28S and 18S rRNA genes to classify
- Ecdysozoa: 37 more arthropods and a kinorhynch. Mol. Phylogenet. Evol. 2006;40:772–94.
- 9. Meusemann K, Reumont von BM, Simon S, Roeding F, Strauss S, Kuck P, et al. A Phylogenomic
- Approach to Resolve the Arthropod Tree of Life. Mol. Biol. Evol. 2010;27:2451–64.
- 10. Song N, Li H, Song F, Cai W. Molecular phylogeny of Polyneoptera (Insecta) inferred from
- expanded mitogenomic data. Nature Publishing Group. 2016;6:36175.
- 11. Kjer KM, Carie FL, Litman J, Ware J. A Molecular Phylogeny of Hexapoda. Arthropod

- 545 Systematics & Phylogeny. 2006;64:35–44.
- 12. Thomas JA, Trueman JWH, Rambaut A, Welch JJ. Relaxed Phylogenetics and the Palaeoptera
- Problem: Resolving Deep Ancestral Splits in the Insect Phylogeny. Syst. Biol. 2013;62:285–97.
- 13. Sasaki G, Ishiwata K, Machida R, Miyata T, Su Z-H. Molecular phylogenetic analyses support
- the monophyly of Hexapoda and suggest the paraphyly of Entognatha. BMC Evol. Biol. BioMed
- 550 Central; 2013;13:236.
- 14. Reumont von BM, Jenner RA, Wills MA, Dell'Ampio E, Pass G, Ebersberger I, et al.
- Pancrustacean Phylogeny in the Light of New Phylogenomic Data: Support for Remipedia as the
- Possible Sister Group of Hexapoda. Mol. Biol. Evol. 2012;29:1031–45.
- 15. Regier JC, Shultz JW, Zwick A, Hussey A, Ball B, Wetzer R, et al. Arthropod relationships
- revealed by phylogenomic analysis of nuclear protein-coding sequences. Nature. 2010;463:1079–83.
- 16. Ma C, Wang Y, Wu C, Kang L, Liu C. The compact mitochondrial genome of *Zorotypus*
- *medoensis* provides insights into phylogenetic position of Zoraptera. BMC Genomics. 2014;15:1156.
- 17. Zhang J, Zhou C, Gai Y, Song D, Zhou K. The complete mitochondrial genome of Parafronurus
- youi (Insecta: Ephemeroptera) and phylogenetic position of the Ephemeroptera. Gene. 2008;424:18–
- 560 24.
- 18. Whiting MF, Carpenter JC, Wheeler QD, Wheeler WC. The Strepsiptera problem: phylogeny of
- the holometabolous insect orders inferred from 18S and 28S ribosomal DNA sequences and
- 563 morphology. Syst. Biol. 1997;46:1–68.
- 19. Giribet G, Ribera C. A Review of Arthropod Phylogeny: New Data Based on Ribosomal DNA
- Sequences and Direct Character Optimization. Cladistics. 2000;16:204–31.
- 20. Ma Y, He K, Yu P, Yu D, Cheng X, Zhang J. The complete mitochondrial genomes of three
- bristletails (Insecta: Archaeognatha): the paraphyly of Machilidae and insights into archaeognathan
- 568 phylogeny. Sun G, editor. PLoS One. 2015;10:e0117669.
- 21. Wheeler WC, Whiting M, Wheeler QD, Carpenter JM, Carpenter. The Phylogeny of the Extant
- 570 Hexapod Orders. Cladistics. 2001;17:113–69.
- 571 22. Lin C-P, Chen M-Y, Huang J-P. The complete mitochondrial genome and phylogenomics of a
- damselfly, *Euphaea formosa* support a basal Odonata within the Pterygota. Gene. 2010;468:20–9.
- 573 23. Wan X, Kim MI, Kim MJ, Kim I. Complete Mitochondrial Genome of the Free-Living Earwig,
- 574 Challia fletcheri (Dermaptera: Pygidicranidae) and Phylogeny of Polyneoptera. PLoS One.
- 575 2012;7:e42056.
- 576 24. Kjer K. Aligned 18S and Insect Phylogeny. Syst. Biol. 2004;53:506–14.
- 25. Misof B, Niehuis O, Bischoff I, Rickert A, Erpenbeck D, Staniczek A. Towards an 18S
- 578 phylogeny of hexapods: Accounting for group-specific character covariance in optimized mixed
- nucleotide/doublet models. Zoology. 2007;110:409–29.
- 26. Reumont von BM, Meusemann K, Szucsich NU, Dell'Ampio E, Gowri-Shankar V, Bartel D, et
- al. Can comprehensive background knowledge be incorporated into substitution models to improve

- 582 phylogenetic analyses? A case study on major arthropod relationships. BMC Evol. Biol. 2009;9:119.
- 583 27. Yoshizawa K, Johnson KP. Aligned 18S for Zoraptera (Insecta): phylogenetic position and
- molecular evolution. Mol. Phylogenet. Evol. 2005;37:572–80.
- 28. Blanke A, Greve C, Wipfler B, Beutel RG, Holland BR, Misof B. The Identification of
- Concerted Convergence in Insect Heads Corroborates Palaeoptera. Syst. Biol. 2013;62:250–63.
- 587 29. Simon S, Strauss S, Haeseler von A, Hadrys H. A Phylogenomic Approach to Resolve the Basal
- Pterygote Divergence. Mol. Biol. Evol. 2009;26:2719–30.
- 30. Baurain D, Brinkmann H, Philippe H. Lack of resolution in the animal phylogeny: closely spaced
- cladogeneses or undetected systematic errors? Mol. Biol. Evol. 2007;24:6–9.
- 31. Rokas A, Carroll SB. Bushes in the tree of life. PLoS Biol. Public Library of Science;
- 592 2006;4:e352.
- 593 32. Whitfield JB, Lockhart PJ. Deciphering ancient rapid radiations. Trends Ecol. Evol.
- 594 2007;22:258–65.
- 33. Talavera G, Vila R. What is the phylogenetic signal limit from mitogenomes? The reconciliation
- between mitochondrial and nuclear data in the Insecta class phylogeny. BMC Evol. Biol.
- 597 2011;11:315.
- 34. Abascal F, Zardoya R, Posada D. ProtTest: selection of best-fit models of protein evolution.
- 599 Bioinformatics. 2005;21:2104–5.
- 35. Abascal F, Posada D, Zardoya R. MtArt: A New Model of Amino Acid Replacement for
- 601 Arthropoda. Mol. Biol. Evol. 2006;24:1–5.
- 36. Le VS, Dang CC, Le QS. Improved mitochondrial amino acid substitution models for metazoan
- evolutionary studies. BMC Evol. Biol. BioMed Central; 2017;17:136.
- 37. Bensasson D. Mitochondrial pseudogenes: evolution's misplaced witnesses. Trends Ecol. Evol.
- 605 2001;16:314–21.
- 38. Rogers HH, Griffiths-Jones S. Mitochondrial Pseudogenes in the Nuclear Genomes of
- 607 Drosophila. PLoS One. 2012;7:e32593.
- 39. Song H, Buhay JE, Whiting MF, Crandall KA. Many species in one: DNA barcoding
- overestimates the number of species when nuclear mitochondrial pseudogenes are coamplified. Proc.
- 610 Natl. Acad. Sci. U.S.A. 2008;105:13486–91.
- 40. Claes KBM, De Leeneer K. Dealing with pseudogenes in molecular diagnostics in the next-
- generation sequencing era. Methods Mol. Biol. New York, NY: Springer New York;
- 613 2014;1167:303–15.
- 41. Cameron SL. Insect Mitochondrial Genomics: Implications for Evolution and Phylogeny. Annu.
- 615 Rev. Entomol. 2014;59:95–117.
- 42. Boore JL. Animal mitochondrial genomes. Nucleic Acids Res. 1999;27:1767–80.

- 43. Saito S. Replication Origin of Mitochondrial DNA in Insects. Genetics. 2005;171:1695–705.
- 44. Zhang D-X, Hewitt GM. Insect mitochondrial control region: A review of its structure, evolution
- and usefulness in evolutionary studies. Biochem. Sys. Ecol. 1997;25:99–120.
- 45. Boore JL, Collins TM, Stanton D, Daehler LL, Brown WM. Deducing the pattern of arthropod
- 621 phylogeny from mitochondrial DNA rearrangements. Nature. 1995;376:163–5.
- 46. Robinson GE, Hackett KJ, Purcell-Miramontes M, Brown SJ, Evans JD, Goldsmith MR, et al.
- 623 Creating a buzz about insect genomes. Sills J, editor. Science. American Association for the
- 624 Advancement of Science; 2011;331:1386–6.
- 47. Barber-James HM, Gattolliat J-L, Sartori M, Hubbard MD. Global diversity of mayflies
- 626 (Ephemeroptera, Insecta) in freshwater. Hydrobiologia. Springer Netherlands; 2008;595:339–50.
- 48. Dijkstra K-DB, Bechly G, Bybee SM, Dow RA, Dumont HJ, Fleck G, et al. The classification
- and diversity of dragonflies and damselflies (Odonata). In: Zhang Z-Q, editor. 2013. pp. 36–45.
- 49. Rutschmann S, Detering H, Simon S, Fredslund J, Monaghan MT. discomark: nuclear marker
- discovery from orthologous sequences using draft genome data. Mol. Ecol. Resour. 2017;17:257–66.
- 631 50. Altschul S. Gapped BLAST and PSI-BLAST: a new generation of protein database search
- 632 programs. Nucleic Acids Res. 1997;25:3389–402.
- 51. Li H, Durbin R. Fast and accurate short read alignment with Burrows-Wheeler transform.
- 634 Bioinformatics. 2009;25:1754–60.
- 52. Li H, Durbin R. Fast and accurate long-read alignment with Burrows-Wheeler transform.
- 636 Bioinformatics. 2010;26:589–95.
- 53. Salmela L, Schroder J. Correcting errors in short reads by multiple alignments. Bioinformatics.
- 638 2011;27:1455–61.
- 54. Simon C, Frati F, Beckenbach A, Crespi B, Liu H, Flook P. Evolution, Weighting, and
- Phylogenetic Utility of Mitochondrial Gene Sequences and a Compilation of Conserved Polymerase
- Chain Reaction Primers. Annals of the Entomological Society of America. Oxford University Press;
- 642 1994;87:651–701.
- 55. Li D, Qin J-C, Zhou C-F. The phylogeny of Ephemeroptera in Pterygota revealed by the
- mitochondrial genome of *Siphluriscus chinensis* (Hexapoda: Insecta). Gene. 2014;545:132–40.
- 56. Aberer AJ, Krompass D, Stamatakis A. Pruning Rogue Taxa Improves Phylogenetic Accuracy:
- An Efficient Algorithm and Webservice. Syst. Biol. 2012;62:162–6.
- 57. Wilkinson M. Majority-rule reduced consensus trees and their use in bootstrapping. Mol. Biol.
- 648 Evol. 1996;13:437–44.
- 58. Felsenstein J. Cases in which Parsimony or Compatibility Methods Will be Positively
- Misleading. Systematic Zoology. 1978;27:401.
- 651 59. Hedtke SM, Townsend TM, Hillis DM. Resolution of phylogenetic conflict in large data sets by
- increased taxon sampling. Syst. Biol. 2006;55:522–9.

- 653 60. Landan G, Graur D. Local reliability measures from sets of co-optimal multiple sequence
- alignments. Pac Symp Biocomput. 2008;:15–24.
- 655 61. Sela I, Ashkenazy H, Katoh K, Pupko T. GUIDANCE2: accurate detection of unreliable
- alignment regions accounting for the uncertainty of multiple parameters. Nucleic Acids Res.
- 657 2015;43:W7–14.
- 658 62. Katoh K, Standley DM. MAFFT Multiple Sequence Alignment Software Version 7:
- Improvements in Performance and Usability. Mol. Biol. Evol. 2013;30:772–80.
- 660 63. Lanfear R, Calcott B, Ho SYW, Guindon S. PartitionFinder: Combined Selection of Partitioning
- Schemes and Substitution Models for Phylogenetic Analyses. Mol. Biol. Evol. 2012;29:1695–701.
- 662 64. Stamatakis A. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large
- phylogenies. Bioinformatics. 2014;30:1312–3.
- 65. Bergsten J. A review of long-branch attraction. Cladistics. 2005;21:163–93.
- 665 66. Kück P, Struck TH. BaCoCa A heuristic software tool for the parallel assessment of sequence
- biases in hundreds of gene and taxon partitions. Mol. Phylogenet. Evol. 2014;70:94–8.
- 667 67. Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Hohna S, et al. MrBayes 3.2:
- Efficient Bayesian Phylogenetic Inference and Model Choice Across a Large Model Space. Syst.
- 669 Biol. 2012;61:539–42.
- 670 68. Paradis E, Claude J, Strimmer K. APE: Analyses of Phylogenetics and Evolution in R language.
- 671 Bioinformatics. 2004;20:289–90.
- 672 69. Schliep KP. phangorn: phylogenetic analysis in R. Bioinformatics. 2011;27:592–3.
- 70. Yu G, Smith DK, Zhu H, Guan Y, Lam TT-Y. ggtree: an r package for visualization and
- annotation of phylogenetic trees with their covariates and other associated data. Methods Ecol Evol.
- 675 2017;8:28–36.
- 71. R Development Core Team. R: A language and environment for statistical computing. R
- Foundation for Statistical Computing, Vienna, Austria. International Statistical Review. 2012. pp.
- 678 357–70.
- 72. Bernt M, Donath A, Jühling F, Externbrink F, Florentz C, Fritzsch G, et al. MITOS: Improved de
- 680 novo metazoan mitochondrial genome annotation. Mol. Phylogenet. Evol. 2013;69:313–9.
- 73. Wyman SK, Jansen RK, Boore JL. Automatic annotation of organellar genomes with DOGMA.
- 682 Bioinformatics. 2004;20:3252-5.
- 74. Laslett D, Canback B. ARWEN: a program to detect tRNA genes in metazoan mitochondrial
- nucleotide sequences. Bioinformatics. 2008;24:172–5.
- 75. Lowe TM, Eddy SR. tRNAscan-SE: a program for improved detection of transfer RNA genes in
- genomic sequence. Nucleic Acids Res. Oxford University Press; 1997;25:955–64.
- 76. Lohse M, Drechsel O, Kahlau S, Bock R. OrganellarGenomeDRAW--a suite of tools for
- generating physical maps of plastid and mitochondrial genomes and visualizing expression data sets.

- 689 Nucleic Acids Res. 2013;41:W575–81.
- 77. Perna N, Kocher T. Patterns of nucleotide composition at fourfold degenerate sites of animal
- mitochondrial genomes. J Mol Evol. 1995;41.
- 78. Rota-Stabelli O, Yang Z, Telford MJ. MtZoa: a general mitochondrial amino acid substitutions
- model for animal evolutionary studies. Mol. Phylogenet. Evol. 2009;52:268–72.
- 79. Misof B, Liu S, Meusemann K, Peters RS, Donath A, Mayer C, et al. Phylogenomics resolves the
- timing and pattern of insect evolution. Science. 2014;346:763–7.
- 80. Simon S, Narechania A, DeSalle R, Hadrys H. Insect Phylogenomics: Exploring the Source of
- Incongruence Using New Transcriptomic Data. Genome Biol. Evol. 2012;4:1295–309.
- 81. Ogden TH, Whiting MF. Phylogeny of Ephemeroptera (mayflies) based on molecular evidence.
- 699 Mol. Phylogenet. Evol. 2005;37:625–43.
- 700 82. Ogden TH, Gattolliat JL, Sartori M, Staniczek AH, Soldán T, Whiting MF. Towards a new
- paradigm in mayfly phylogeny (Ephemeroptera): combined analysis of morphological and molecular
- 702 data. Sys. Entomol. 2009;34:616–34.
- 83. Letsch HO, Meusemann K, Wipfler B, Schutte K, Beutel R, Misof B. Insect phylogenomics:
- results, problems and the impact of matrix composition. Proc. R. Soc. B. 2012;279:3282–90.
- 705 84. Letsch H, Simon S. Insect phylogenomics: new insights on the relationships of lower neopteran
- orders (Polyneoptera). Sys. Entomol. 2013;38:783–93.
- 85. Zhang Y-Y, Xuan W-J, Zhao J-L, Zhu C-D, Jiang G-F. The complete mitochondrial genome of
- 708 the cockroach *Eupolyphaga sinensis* (Blattaria: Polyphagidae) and the phylogenetic relationships
- 709 within the Dictyoptera. Mol. Biol. Rep. 2009;37:3509–16.
- 710 86. Fenn JD, Song H, Cameron SL, Whiting MF. A preliminary mitochondrial genome phylogeny of
- 711 Orthoptera (Insecta) and approaches to maximizing phylogenetic signal found within mitochondrial
- 712 genome data. Mol. Phylogenet. Evol. 2008;49:59–68.
- 713 87. Ma C, Liu C, Yang P, Kang L. The complete mitochondrial genomes of two band-winged
- grasshoppers, Gastrimargus marmoratus and Oedaleus asiaticus. BMC Genomics. 2009;10:156.
- 88. Inward D, Beccaloni G, Eggleton P. Death of an order: a comprehensive molecular phylogenetic
- study confirms that termites are eusocial cockroaches. Biology Letters. 2007;3:331–5.
- 89. Legendre F, Nel A, Svenson GJ, Robillard T, Pellens R, Grandcolas P. Phylogeny of
- 718 Dictyoptera: Dating the Origin of Cockroaches, Praying Mantises and Termites with Molecular Data
- and Controlled Fossil Evidence. PLoS One. 2015;10:e0130127.
- 90. Regier JC, Mitter C, Zwick A, Bazinet AL, Cummings MP, Kawahara AY, et al. A large-scale,
- higher-level, molecular phylogenetic study of the insect order Lepidoptera (moths and butterflies).
- 722 Moreau CS, editor. PLoS One. 2013;8:e58568.
- 723 91. Wu H-Y, Ji X-Y, Yu W-W, Du Y-Z. Complete mitochondrial genome of the stonefly
- 724 *Cryptoperla stilifera* Sivec (Plecoptera: Peltoperlidae) and the phylogeny of Polyneopteran insects.
- 725 Gene. 2014;537:177–83.

- 92. Pons J, Bauzà-Ribot MM, Jaume D, Juan C. Next-generation sequencing, phylogenetic signal
- and comparative mitogenomic analyses in Metacrangonyctidae (Amphipoda: Crustacea). BMC
- 728 Genomics. 2014;15:566.

- 729 93. Tang M, Tan M, Meng G, Yang S, Su X, Liu S, et al. Multiplex sequencing of pooled
- 730 mitochondrial genomes--a crucial step toward biodiversity analysis using mito-metagenomics.
- 731 Nucleic Acids Res. 2014;42:e166–6.
- 732 94. Lee EM, Hong MY, Kim MI, Kim MJ, Park HC, Kim KY, et al. The complete mitogenome
- 733 sequences of the palaeopteran insects *Ephemera orientalis* (Ephemeroptera: Ephemeridae) and
- 734 Davidius lunatus (Odonata: Gomphidae). Genome. 2009;52:810–7.
- 95. Kim MJ, Jung KS, Park NS, Wan X, Kim K-G, Jun J, et al. Molecular phylogeny of the higher
- taxa of Odonata (Insecta) inferred from COI, 16S rRNA, 28S rRNA, and EF1-α sequences.
- 737 Entomological Research. 2014;44:65–79.