



Data curation and modeling of compositional heterogeneity in insect phylogenomics: A case study of the phylogeny of Dytiscoidea (Coleoptera: Adephaga)



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ABSTRACT

Diving beetles and their allies are an almost ubiquitous group of freshwater predators. Knowledge of the phylogeny of the adephagan superfamily Dytiscoidea has significantly improved since the advent of molecular phylogenetics. However, despite recent comprehensive phylogenomic studies, some phylogenetic relationships among the constituent families remain elusive. In particular, the position of the family Hygrotidae remains uncertain. We address these issues by re-analyzing recently published phylogenomic datasets for Dytiscoidea, using approaches to reduce compositional heterogeneity and adopting a site-heterogeneous mixture model. We obtained a consistent, well-resolved, and strongly supported tree. Consistent with previous studies, our analyses support Aspidytidae as the monophyletic sister group of Amphizoidae, and more importantly, Hygrotidae as the sister of the diverse Dytiscidae, in agreement with morphology-based phylogenies. Our analyses provide a backbone phylogeny of Dytiscoidea, which lays the foundation for better understanding the evolution of morphological characters, life habits, and feeding behaviors of dytiscoid beetles.

1. Introduction

Just like many other fields of biology, entomology has experienced a true genomic revolution in the past few decades (Cameron, 2014; Behura, 2015; Kjer et al., 2016a; Johnson, 2019). While the first Sanger-era molecular phylogeny of insects only analyzed 18S sequences and restriction sites for 12 species (Wheeler, 1989), the most recent insect super tree included 440 transcriptomes, 1,490 mitogenomes, and DNA barcodes for some 69,000 species (Chester, 2019). However, despite the almost exponential increase in sequence data, this growth has not been matched by a growing consensus over insect phylogeny. Many old controversies in insect evolution remain unresolved, such as the relationships among the earliest-branching endognathous orders, early-branching winged insects (the ‘Palaeoptera problem’), and the phylogenetic position of highly specialized parasites such as fleas (Kjer et al., 2016b; Misof et al., 2014; Beutel et al., 2017; Johnson et al., 2018; Simon et al., 2018; Wipfler et al., 2019). Likewise, the monophyly and position of numerous traditionally well-defined taxa has been

questioned by recent phylogenomic studies (Pauli et al., 2018; Ding et al., 2019; Hamilton et al., 2019; McKenna et al., 2019; Tang et al., 2019; Wang et al., 2019; Winterton et al., 2019). It remains unclear to what extent these inconsistencies between traditional and phylogenetically-derived hypotheses represent progress in elucidating insect phylogeny and to what extent they are the result of factors confounding the reliable reconstruction of phylogenetic relationships (Borowiec et al., 2019; Cai et al., 2019).

One of the key sources of uncertainty and error in inferring phylogenies is compositional and rate heterogeneity (Bleidorn, 2017). Some of the most popular inference methods used in phylogenomics operate under the assumption that the rate of evolutionary change is equal for every position of a sequence alignment (Sheffield et al., 2009). However, this assumption is unrealistic and does not reflect the high compositional and rate heterogeneity observed in metazoan genomes (Lartillot and Philippe, 2008); not only does mutation rate vary among bases (Hodgkinson and Eyre-Walker, 2011), but different parts of the genome are under selection pressures of different intensities (Xing and

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Lee, 2006), typically resulting in a highly unequal evolutionary rate across any given sequence. Models which assume compositional and rate homogeneity can consistently recover incorrect topologies, albeit often with high statistical support (Ho and Jermiin, 2004; Jermiin et al., 2004; Cox et al., 2008; Sheffield et al., 2009). To combat these problems, an arsenal of methods has been developed to reduce site compositional heterogeneity in datasets, such as various data filtering and data recoding approaches (Bleidorn, 2017). Moreover, some recent complex site-heterogeneous models can account for both compositional and rate heterogeneity across sites. These models, such as CAT-GTR + G, have been shown to fit real data better than conventional site-homogeneous models and suppress common sources of phylogenetic error such as long branch attraction (Lartillot et al., 2007; Blanquart and Lartillot, 2008; Wang et al., 2008; Foster et al., 2009). When reanalyzed using these approaches, some of the most controversial debates in evolutionary biology today such as the origin of eukaryotes and metazoans seem to boil down to problems caused by compositional and/or rate heterogeneity (Cox et al., 2008; Feuda et al., 2017; Pisani et al., 2015; Puttik et al., 2018; Tarver et al., 2016; Williams et al., 2020).

The adephagan superfamily Dytiscoidea (Amphizoidae, Aspidytidae, Dytiscidae, Hygrobiidae, Meruidae, and Noteridae) is a well-established group of beetles (e.g. Baca et al., 2017; Beutel et al., 2013; Dressler et al., 2011; but see López-López and Vogler, 2017). Dytiscoid species occur in diverse freshwater habitats, including springs, rivers, acidic swamps, lakes, and even in hypersaline and hygropetric habitats. Bell (1966) suggested a clade, Dytiscoidea, comprising aquatic (or semi-aquatic) families such as Noteridae, Amphizoidae, Hygrobiidae, and Dytiscidae. The monophyly of Dytiscoidea has been confirmed in many phylogenetic analyses of morphological characters (Beutel and Haas, 1996; Beutel, 1998; Beutel and Haas, 2000) as well as analyses of molecular data (Ribera et al., 2002a,b; McKenna et al., 2015).

Although the phylogenetic relationships of dytiscoids have been extensively investigated based on morphology, gland chemical compounds, fossils, and molecular data (e.g. Alarie et al., 2011; Alarie and Bilton, 2005; Baca et al., 2017; Balke et al., 2008; Beutel et al., 2006, 2008, 2013; Beutel and Haas, 1996; Burmeister, 1976; Dettner, 1985; Kavanaugh, 1986; López-López and Vogler, 2017; McKenna et al., 2015; Ribera et al., 2002b; Toussaint et al., 2015), these different datasets do not yield a congruent topology (Vasilikopoulos et al., 2019). Both morphology and molecular based phylogenies have indicated that Meruidae + Noteridae represent the sister clade of the remaining four dytiscoid families (summarized in Vasilikopoulos et al., 2019). However, the phylogenetic relationships among Amphizoidae, Aspidytidae, Dytiscidae and Hygrobiidae, remain unresolved. A recent phylogenomic study based on transcriptomes provided new insights into the backbone phylogeny of Dytiscoidea (Vasilikopoulos et al., 2019): Aspidytidae (cliff water beetles) was recovered as a monophyletic group, which is sister to the relictual family Amphizoidae. However, this phylogenomic study could not present conclusive evidence for some of the interfamilial relationships. After accounting for potential tree confounding factors, Vasilikopoulos et al. (2019) concluded that Hygrobiidae (squeak beetles) is most likely a sister group to a clade comprising Amphizoidae, Aspidytidae, and Dytiscidae. While this result has also been supported by previously published Sanger sequence data and a combination of molecular and morphological data (Balke et al., 2005, 2008), other recent phylogenomic-scale studies have arrived at yet different results. The largest phylogeny of beetles published to date, based on 4818 genes (McKenna et al., 2019), and an analysis of Adephaga based on ultraconserved elements (Gustafson et al., 2019) have both recovered Hygrobiidae as a sister to Amphizoidae + Aspidytidae. In addition, the results of Vasilikopoulos et al. (2019) contradict the conventional hypothesis inferred from comparative morphological studies. For example, a clade consisting of Dytiscidae and Hygrobiidae is strongly supported by some critical morphological features (Beutel et al., 2006; Dressler and Beutel, 2010) such as the presence of

prothoracic glands (Beutel, 1986, 1988).

To understand the systematic position of Hygrobiidae and the backbone phylogeny of Dytiscoidea, we re-analyzed the recently published phylogenomic data for Dytiscoidea of Vasilikopoulos et al. (2019) using a site-heterogeneous mixture model (CAT-GTR + G in PhyloBayes). We also investigated the effects of different approaches of reducing compositional heterogeneity of the datasets by the data block mapping and gathering using entropy (BMGE) method.

2. Materials and methods

2.1. Dataset selection

We used the amino acid transcriptome alignments from Vasilikopoulos et al. (2019). The authors produced and analyzed different variants of nucleotide and amino acid alignments of their data. Among the eleven amino-acid supermatrices they generated, their focal analyses were principally based upon the full dataset (Supermatrix A: 14 taxa, 1,661,023 amino-acid sites) and two datasets reduced to increase data coverage and phylogenetic information (Supermatrix E: 14 taxa, 948,772 amino-acid sites) and diminish the negative effects of among-species compositional heterogeneity (Supermatrix H: 14 taxa, 211,275 amino-acid sites) (Vasilikopoulos et al., 2019). Here we focused on the same three supermatrices download from MENDELEY DATA (<https://doi.org/10.17632/j8wxdtbyb.1>) to test hypotheses about the backbone phylogeny of Dytiscoidea.

To reduce among-site compositional heterogeneity and ease the convergence of runs, we used data block mapping and gathering using entropy (BMGE). BMGE identifies phylogenetically informative sites by computing entropy-like scores weighted with BLOSUM similarity matrices in order to distinguish among biologically expected and unexpected variability for each aligned character (Criscuolo and Gribaldo, 2010). BMGE can select characters associated with a score value below a fixed threshold. We prepared four stringently filtered datasets (Supermatrices A', A'', E' and H') by trimming the previously published supermatrices A, E and H using BMGE v.1.1 and a threshold value (set using the -h flag) of 0.4 (supermatrices A', E' and H') or 0.3 for a more conserved supermatrix A''. BLOSUM95 (Henikoff and Henikoff, 1992) similarity scores were used as the studied taxa are relatively closely related and similarity between their amino acid sequences is expected to be relatively high. A test was also performed, using supermatrix A (threshold value of 0.4) where BLOSUM62, that assumes sequences to be more distantly related, was used instead of BLOSUM95.

2.2. Phylogenetic analyses of amino-acid sequence

We employed both site-heterogeneous (CAT-GTR + G) and site-homogenous (LG4X + R) models to evaluate competing hypotheses on the phylogenetic relationships among the main groups of Dytiscoidea. The CAT-GTR + G analyses were performed in PhyloBayes MPI 1.7 (Lartillot et al., 2009) for all trimmed datasets. For the CAT-GTR + G analyses, two independent Markov chain Monte Carlo (MCMC) chains were run until convergence (maxdiff < 0.3). For each PhyloBayes run, we used the bpcomp program to generate output of the largest (maxdiff) and mean (meandiff) discrepancy observed across all bipartitions. In addition, all trimmed alignments (supermatrices A', A'', E' and H') were used for maximum-likelihood (ML) phylogenetic reconstruction under the LG4X + R model (Le et al., 2012) as implemented in IQ-TREE v.1.6.10 with analyses run using 1000 ultra-fast bootstraps (Nguyen et al., 2015).

3. Results

3.1. Data filtering

Using the BMGE filtering method we obtained four new datasets,

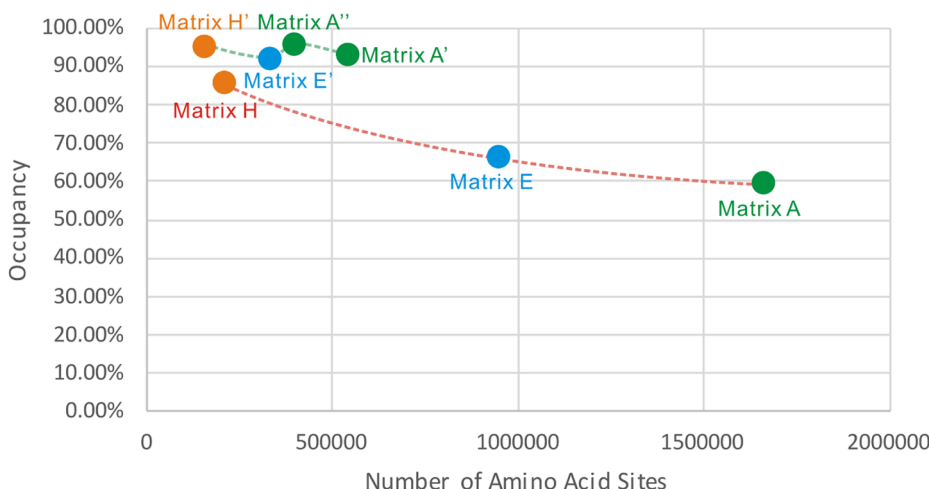


Fig. 1. Data occupancies and amino acid site numbers of original (Matrices A, E and H) and trimmed (Matrices A', A'', E' and H') supermatrices that were used in the present study.

which represent subsets of the more conserved amino acid sites of the original supermatrices A, E, and H. The amino acid occupancy of all matrices was significantly improved, especially for larger datasets such as Supermatrices A and E: the data occupancy of Supermatrix A (1,661,023 sites) increased from 59.76% to 92.98% in Supermatrix A' (542,493 sites) and to 95.48% in Supermatrix A'' (399,769 sites), Supermatrix E (948,772 sites) increased from 66.54% to 91.97% in Supermatrix E' (334,457 sites), and Supermatrix H (211,275 sites) increased from 85.92% to 95.22% in Supermatrix H' (156,395 sites) (Fig. 1).

3.2. Site-homogeneous model analyses

Our maximum likelihood (IQ-TREE) LG4X + R analyses of the amino-acid supermatrices E' and H' resulted in identical topologies (Fig. 2) to those based on the original supermatrices E and H under optimized schemes, respectively (Fig. 2a,b in Vasilikopoulos et al., 2019). Moreover, the support values are interestingly correlated to those obtained in the original analyses. For instance, for supermatrices A', A'' and H', the nodes uniting Amphizoidae + Aspidytidae and Dytiscidae were weakly supported (MLB = 73 for supermatrix H'). Similarly, within the family Dytiscidae the node between *Lioterpes haemorrhoidalis* and *Cybister lateralimarginalis* + *Thermonectus intermedius* was moderately supported (MLB = 90 for supermatrix H'). Unlike the 10-partitioned ML tree of the original supermatrix A (Supplementary Fig. 45 in Vasilikopoulos et al., 2019), our maximum likelihood analyses of the filtered supermatrices A' and A'' both yielded a topology identical to the one under supermatrix H' or supermatrix H, in which Hygrobiidae is the sister group to the weakly supported (MLB = 54 in supermatrix A' and 58 in supermatrix A'') clade (Aspidytidae + Amphizoidae) + Dytiscidae (Fig. 2). Based on the maximum likelihood analyses of supermatrices A' and A'', we found that a more conserved dataset with slower-evolving sites can produce an identical but better supported topology under the same model (Fig. 2).

3.3. Site-heterogeneous model analyses

The largest discrepancies (maxdiff) in all PhyloBayes runs equal to 0 (maxdiff < 0.1), indicating they all represent 'good' runs (Lartillot et al., 2013). Like the analyses of amino acid sequence data in Vasilikopoulos et al. (2019), all analyses in the present study supported the monophyly of Dytiscoidea and of each dytiscoid family, and indicated a sister group relationship between Noteridae and the other families of Dytiscoidea, including Amphizoidae, Aspidytidae, Dytiscidae, and Hygrobiidae. All the above relationships received maximal

statistical support (Bayesian Posterior Probabilities [BPP] = 1) in all analyses (Fig. 3). Our PhyloBayes analysis of the original amino-acid supermatrix H, which were not trimmed using BMGE to reduce the compositional heterogeneity of amino acids, suggested Hygrobiidae as the sister group to Dytiscidae + (Aspidytidae + Amphizoidae) with maximal support (BPP = 1), a topology identical to the one based on the same dataset (Supermatrix H) but under a site-homologous model (Fig. 2a in Vasilikopoulos et al., 2019). In addition to this analysis based on the original supermatrix (Supermatrix H), the PhyloBayes analyses based on our new filtered datasets (Supermatrices A', A'', E' and H') all resulted in an identical and fully supported topology: a clade composed of Amphizoidae and Aspidytidae, a clade composed of Dytiscidae and Hygrobiidae, and Noteridae as a sister group to both clades (Fig. 3). Trimming supermatrix A with BLOSUM62 -h 0.4 and subsequently analyzing this dataset with the CAT-GTR + G model yielded the same topology as the CAT-GTR + G analyses that performed BMGE trimming using BLOSUM95 -h 0.4 (Fig. S1).

In all tree reconstructions based on filtered datasets under a site-heterogeneous model, Noteridae was supported as the sister group to all remaining Dytiscoidea. Both clades of Aspidytidae + Amphizoidae and Dytiscidae + Hygrobiidae were strongly supported by all analyses based on the amino-acid datasets. We observed a confounding signal in the original amino-acid dataset (Supermatrix H), which is probably caused by compositional heterogeneity. The position of Hygrobiidae within Dytiscoidea (as a sister group to Dytiscidae) was stable and consistent in all analyses of filtered amino acid datasets.

4. Discussion

Despite extensive analyses of both morphological and molecular data, it has proven challenging to achieve a congruent reconstruction of dytiscoid phylogeny (e.g. Baca et al., 2017; Balke et al., 2005, 2008; Beutel et al., 2008, 2013; Toussaint et al., 2015; Vasilikopoulos et al., 2019). To tackle this problem, we used a large published phylogenomic dataset representing all dytiscoid families except Meruidae. Unlike the inconsistent and equivocal results under various datasets in Vasilikopoulos et al. (2019), our analyses based on a complex and better-fitting model and multiple datasets with reduced compositional heterogeneity yielded a consistent and fully supported tree of Dytiscoidea. We suggest that Noteridae (plus most likely Meruidae, Vasilikopoulos et al., 2019) is the earliest-branching lineage within Dytiscoidea, sister to a clade composed of Amphizoidae, Aspidytidae, Dytiscidae, and Hygrobiidae (McKenna et al., 2015; Vasilikopoulos et al., 2019). As confirmed in the recent phylogenomic study of Vasilikopoulos et al. (2019) and other morphological and/or molecular

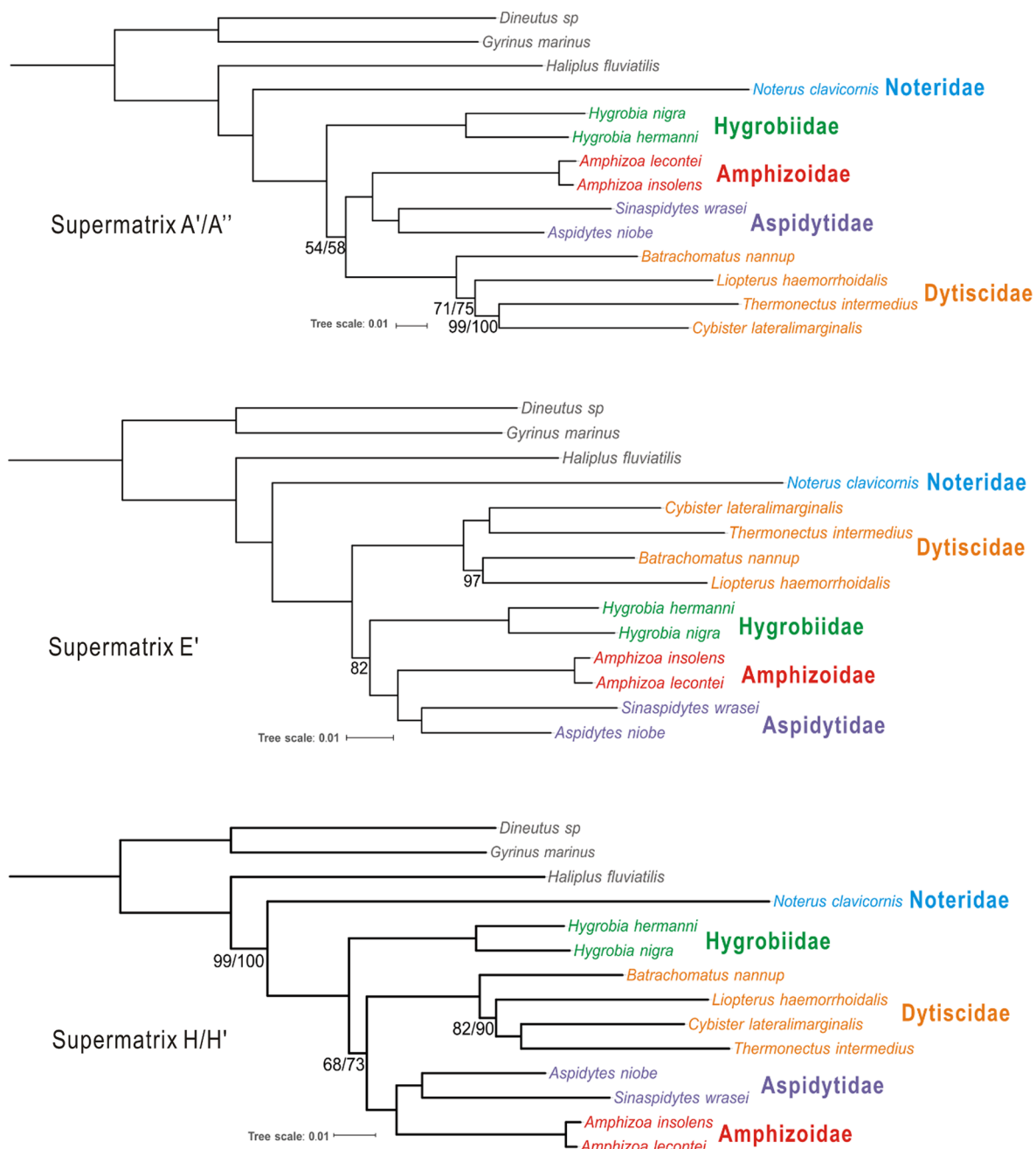


Fig. 2. Different phylogenetic hypotheses deduced from the analysis of amino-acid sequence data (Supermatrices A', E', H' and H) under the simplistic LG4X + R model. Branch support (MLB) is denoted based on 1000 ultrafast bootstrap replicates; MLB values equal to 100 are not shown.

phylogenies (e.g. Balke et al., 2005, 2008), Aspidytidae is monophyletic and sister to Amphizoidae with strong support in all Bayesian analyses of the amino-acid sequence data.

The phylogenetic position of Hygobiidae is well resolved by our re-analyses, unlike the results in Vasilikopoulos et al. (2019), in which the phylogenetic position is affected by a highly conflicting phylogenetic signal. A clade encompassing Hygobiidae and Dytiscidae, as suggested by some studies based on the analysis of morphological characters (e.g. Beutel et al., 2013; Beutel and Roughley, 1988; Dressler et al., 2011), is strongly supported in all analyses of filtered datasets. Despite several obvious anatomical differences between Hygobiidae and Dytiscidae (Alarie et al., 2004; Dettner, 2016), many studies including an analysis of molecular data (Shull et al., 2001) suggest that these families are sister groups. A close relationship between Hygobiidae and Dytiscidae is also supported by a combined phylogenetic analysis (Ribera et al.,

2002a), larval morphology (Alarie and Bilton, 2005), and traces of antimicrobial pygidial gland compounds such as benzoic acid and *p*-hydroxybenzaldehyde (Dettner, 1987). More importantly, they share a similar prothoracic defensive gland (Forsyth, 1970), which is another potential synapomorphy of the two families (Dettner, 2016).

Previous simulation studies showed that site trimming using BMGE produces datasets with reduced heterogeneity and so the method has been widely applied to infer deep phylogenies (e.g. Zaremba-Niedzwiedzka et al., 2017; Martijn et al., 2018; Lahr et al., 2019; Philippe et al., 2019; Strassert et al., 2019). Our filtered datasets, with a significantly improved signal-to-noise ratio, are suitable for phylogenetic analyses, and the phylogenetic trees are less affected by phylogeny reconstruction artefacts due to compositional heterogeneity (e.g. Feuda et al., 2017; Lozano-Fernandez et al., 2019a). Regardless of the BLOSUM method used for trimming, the topologies were identical

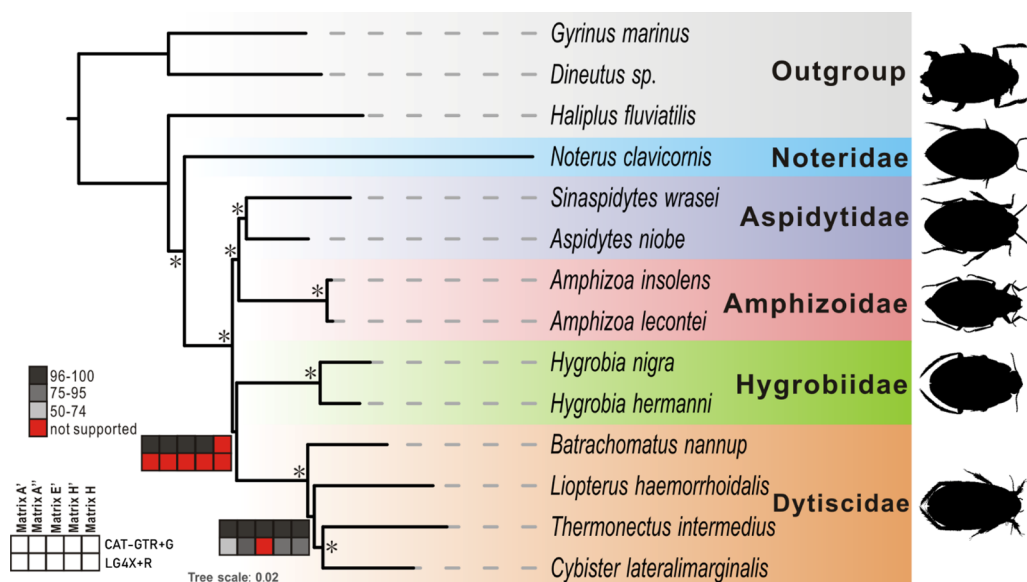


Fig. 3. Phylogenetic tree based on the PhyloBayes analysis of supermatrix A' with the site-heterogeneous CAT-GTR + G model. Supermatrix A' comprises 14 taxa (11 in-group taxa) and 542,493 amino acid positions. Support values for all analyses are plotted below respective branches as specified in the legend at the bottom-left corner. * denotes strongly supported clades in all analyses ($BPP > 0.98$ or $MLB > 95$).

further demonstrating the robustness of our analyses. Unlike the tree reconstructing methods used in Vasilikopoulos et al. (2019), we employed the more complex site-heterogeneous CAT-GTR + G model implemented in PhyloBayes, which can account for potential site-specific amino acid preferences (or compositional heterogeneity) (e.g. Lozano-Fernandez et al., 2019a; Schwentner et al., 2017; Wolfe et al., 2019). The CAT-GTR + G model has been demonstrated to be effective in suppressing artefacts such as long-branch and compositional attraction, especially for large-scale analyses (Feuda et al., 2017; Lartillot et al., 2007; Lozano-Fernandez et al., 2019). In addition, based on the comparative analyses of both amino acid and nucleotide sequence data by Vasilikopoulos et al. (2019), amino acids should be preferred to nucleotides in phylogenomic analyses of ancient relationships (e.g. Inagaki and Roger, 2006; Rota-Stabelli et al., 2013; Schwentner et al., 2017).

When all datasets (even those filtered using BMGE) are analyzed using maximum likelihood (ML) under the less fitting LG4X + R model, a tree is supported where Amphizoidae is the sister group to Aspidytidae, but the systematic position of Hygrotiidae is, as observed in the previous study (Vasilikopoulos et al., 2019), not stable. It is noteworthy that in all ML trees of the filtered amino acid datasets the support values of the nodes between Hygrotiidae and other dytiscoid families are never well supported ($MLB = 54$ in Supermatrix A', $MLB = 58$ in Supermatrix A'', $MLB = 82$ in Supermatrix E', and $MLB = 73$ in Supermatrix H'). Similar weakly supported results, also obtained in Vasilikopoulos et al. (2019) under the site-homogeneous model, are probably artefactual. As indicated in Vasilikopoulos et al. (2019), the systematic position of Hygrotiidae cannot be resolved unambiguously under ML with the model they adopted. This difficulty is probably, in part, due to a lack of sufficient phylogenetic signal for the Hygrotiidae and Dytiscidae clade, since the internode between these two families is very short under the CAT-GTR + G model, perhaps reflecting early rapid diversification of these beetles. Such a problem is also found in other phylogenomic studies of other pancrustacean animals (e.g. Lozano-Fernandez et al., 2019; Schwentner et al., 2017), where the sister group of Hexapoda, Remipedia, can only be recovered under a site-heterogeneous model (CAT-GTR + G) but not a homogeneous model. Recent studies that have recovered Hygrotiidae as a sister to a clade containing Amphizoidae and Aspidytidae (Gustafson et al., 2019; McKenna et al., 2019) have likewise both relied on time-saving site-homogeneous models or their ML extensions which do not account for compositional heterogeneity and can lead to the recovery of misleading topologies, as demonstrated in our analyses.

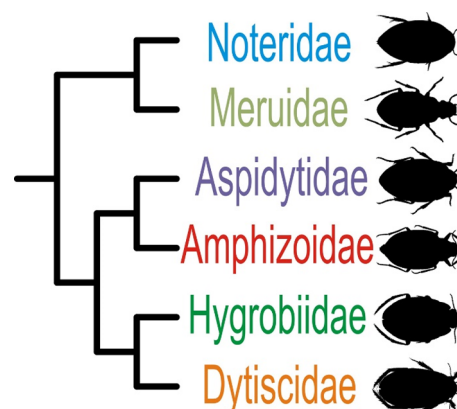


Fig. 4. Phylogenetic hypothesis on family phylogenetic relationships among Dytiscoidea based on the present study and previously published data.

Overall, our results are consistent with morphology-based views of dytiscoid relationships. The sister-group relationship between Hygrotiidae and Dytiscidae was proposed by Burmeister (1976) based on morphology of the ovipositor and by Ruhnau (1986) based on larval morphology. Both adult Dytiscidae and Hygrotiidae also share the presence of prothoracic glands, among other characters (Forsyth, 1970; Beutel, 1986; Beutel, 1988). A clade comprising the two families was recently recovered by a maximum parsimony analysis of morphological data (Beutel et al., 2020). This same analysis also recovered Aspidytidae as a sister to Amphizoidae, in congruence with our CAT-GTR + G trees. It should be noted, however, that some deeper nodes in Beutel et al., 2020 did not receive high bootstrap support values, which is a common problem in morphological phylogenies (Fig. S2). With the relationships among Dytiscoidea strongly supported in our analyses (Fig. 3), our results confirm Beutel and colleague's morphology-based phylogeny of Dytiscoidea.

5. Concluding remarks

The phylogenetic relationships presented here provide an updated hypothesis about the evolution of Dytiscoidea and the systematic position of the relictual family Hygrotiidae. By careful filtering of the original supermatrices and employing a site-heterogeneous mixture model (CAT-GTR + G), the interrelationships of the five dytiscoid families can be resolved with confidence. Our phylogenomic result is

congruent with the conventional morphology-based phylogenetic tree of Dytiscoidea. Tackling potential sources of systematic error strengthens support for a relationship between Hygrobiidae and Dytiscidae. Integrating various previous studies of the systematic position of the small family Meruidae (Baca et al., 2017; Balke et al., 2008; Beutel et al., 2013, 2020; McKenna et al., 2015; Toussaint et al., 2015), we propose a unified phylogenetic framework for the six extant families of Dytiscoidea: (Meruidae + Noteridae) + ((Aspidytidae + Amphizoidae) + (Dytiscidae + Hygrobiidae)) (Fig. 4). Based on this tree of Dytiscoidea, it will now be possible to address and test a series of hypotheses regarding the evolution of many critical morphological innovations in Dytiscoidea.

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