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Multiple nuclear and mitochondrial genes resolve the branching order of a rapid radiation of crested newts (Triturus, Salamandridae)

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1. Introduction

The task of molecular phylogenetics is to reconstruct the order by which species have originated. However, when the different lineages split over short time spans or when the studied markers contain insufficient information, phylogenies can remain unresolved. One way forward is to collect large amounts of independent data. Independence is essential to resolve the gene tree—species tree problem, as linked loci (for example, from mitochondrial DNA) will have the same history. Because of stochasticity in the lineage sorting process, incomplete lineage sorting and horizontal gene transfer, gene trees generally differ from one another and from the actual species tree (which is tried to be recovered) (Avise and Wollenberg, 1997). This also brings up analytical problems. Over the last decades, phylogenetic analyses have mostly followed three strategies (reviewed in Huelsenbeck et al., 1996): the total evidence approach (Kluge, 1989), the separate analysis for each marker followed by an analysis for congruence (Miyamoto and Fitch, 1995), and conditional data combination (de Queiroz, 1993). More recently, phylogenetic methods have been proposed that incorporate biologically more realistic models of evolution, by taking into account processes like incomplete lineage sorting or horizontal gene transfer. Two new methods are phylogenetic networks (see Huson and Bryant, 2006) and an approach that calculates species trees explicitly, (hereafter called ‘the hierarchical method’; Edwards et al., 2007).

The genus Triturus (Caudata: Salamandridae) is composed of two groups, with two and five species, respectively (the marbled newts—T. marmoratus and T. pygmaeus—from south-western Europe and the crested newts—or T. cristatus superspecies—from western, central and eastern Europe). Support for a monophyletic origin is strong for either of the groups, i.e., crested newts, marbled newts and Triturus (Arntzen et al., 2007; Steinfartz et al., 2007). The five crested newt species have mutually exclusive, parapatric ranges across Europe: the Italian crested newt (T. carnifex), the northern crested newt (T. cristatus), the Dobrogean crested newt (T. dobrogicus), the southern crested newt (T. karelinii) with two subspecies: T. k. karelinii and T. k. arntzenii and the Macedonian crested newt (T. macedonicus). All species meet in the Balkans which is considered to be the centre of origin of the group (Cnobić et al., 1997; see Fig. 1). Crested newts come in four morphologically distinct lineages that differ in the number of presacral vertebrae and relative appendage size (e.g., forelimb length divided by interlimb distance—a ratio
known as the ‘Wolterstorff-Index’ and negatively correlated with the number of presacral vertebrae). Modal presacral vertebrae counts are 14 in \( T. \) karelinii, 15 in \( T. \) carnifex and \( T. \) macedonicus, 16 in \( T. \) cristatus and 17 or 18 in \( T. \) dobrogicus, with little intraspecific variation (Arentzen, 2003). \( T. \) Triturus marmoratus and \( T. \) pygmaeus have a modal presacral vertebrae count of 13. Note however that the evolutionary tree suggested by this morphological series, as seen from the root \((T. \) karelinii, \((T. \) carnifex, \( T. \) macedonicus), \((T. \) cristatus, \( T. \) dobrogicus)), is not supported by molecular genetic analyses (Arentzen et al., 2007).

As a result of that analysis, we forwarded the hypothesis that the speciation of crested newts was near-simultaneous at around 11–10 Ma and occurred in the Balkans (Arentzen et al., 2007). The temporal calibration was obtained from a 24 Myr old fossil record ascribed to \( T. \) marmoratus (Estes, 1981; Steinfartz et al., 2007). We found external support for this hypothesis in the observations that genetic and species diversity is highest in the Balkans and that a polytomous speciation scenario matches with the paleogeography of the Balkan region (Popov et al., 2004). It is not expected that more precise paleogeographic reconstructions will become available soon, so we would suggest, on the contrary, that an established phylogeny—either resolved or polytomous—might actually assist the reconstruction of the paleogeography of south-eastern Europe.

In our previous work on the phylogeny and historical biogeography of the genus \( T. \) Triturus we failed to resolve the basal node of the crested newt radiation (Arentzen et al., 2007). Indeed, no firm phylogenetic resolution has been found among five crested newt lineages, other than the sister-species status of \( T. \) carnifex and \( T. \) macedonicus (see below). We here apply some of the above methods with a set of seven nuclear and mitochondrial genes, in an attempt to break the crested newt polytomy.

2. Material and methods

2.1. Sampling design

We sampled the five species of crested newts (including both subspecies of \( T. \) karelinii; \( T. \) k. karelinii and \( T. \) k. arntzenii) and one species of marbled newt as outgroup, with altogether 15 individuals from as many populations (Appendix 1). Because each species of crested newt is in spatial contact with two or more other species of the group (Fig. 1) and because hybridization and introgression is known to occur in limited areas (Wallis and Arentzen, 1989), we sampled localities at least 100 km away from documented contact zones (Arentzen, 2003). This was done to minimize confounding effects of interspecific gene flow on the phylogenetic reconstruction. Sampling effort reflected the level of intraspecific genetic variation known for the species, which is low in \( T. \) cristatus (one locality sampled) and high in \( T. \) karelinii (six localities sampled).

2.2. DNA extraction and sequencing

Tail-tip tissue samples were stored in −80°C and later transferred to 95% ethanol. Total genomic DNA was extracted with the Qiagen DNeasy tissue kit. PCR conditions consisted of an initial denaturation step at 94°C for 4 min, followed by 35 cycles of 30 s denaturation at 94°C, 45 s annealing at 57–68°C (depending on the fragment) and 90 s extension at 72°C and a final extension step of 4 min at 72°C. The molecular markers employed are two protein-coding mitochondrial genes (subunits 2 and 4 of the NADH dehydrogenase gene complex—ND2 and ND4) and five nuclear genes, namely three introns (β-Fibrinogen intron 7—fibbint7, Calreticulin intron C—CalintC, Platelet-derived growth factor receptor α—Pdgfra) and two anonymous markers (Cri1 and Cri4) (Espregueira Themudo et al., 2009). The primers used are described in Table 1. To increase accuracy of the results, both DNA strands were sequenced. Sequences were obtained commercially through Macrogen Inc. BLAST searches at Genbank confirmed the homology of the nuclear introns and the mtDNA genes with published data, while the anonymous nuclear markers remained unidentified.

2.3. Data analysis

Chromatograms were checked for basecalling errors with Chromas software (Technelysium Pty Ltd.). Sequences were aligned in BioEdit (Hall, 1999). Insertion/deletion polymorphisms (indels)
reflecting heterozygosity were reconstructed by comparing chromatograms for the forward and reverse primers (following Flot et al., 2006). Indels were excluded from the phylogenetic analysis. Because of difficulties in analyzing individuals which are heterozygous for more than one nuclear gene, phased haplotypes were combined into a consensus haplotype by coding polymorphic sites with the IUPAC ambiguity codes (IUPAC, 1974). The partition homogeneity test (or Incongruence length difference test; Farris et al., 1994) as implemented in PAUP 4.0b10 (Swofford, 2003) was used to check for congruence in phylogenetics signal among genes. Phylogenetic congruence was rejected for all pairs of markers, except the two mitochondrial genes. Nucleotide composition, transition/transversion ratio and sequence divergence under the Kimura 2-parameter model were calculated with MEGA4 (Tamura et al., 1994) as implemented in PAUP 4.0b10 (Swofford, 2003). The first quarter of the trees was discarded as burn-in and 25 million generations, with a sample taken every 1000 generations, the chain length used was 300 million generations, with one sample taken every 1000 generations. The last 10,000 trees sampled from the stationary phase were summarized under the majority-rule consensus criterion.

2.4. Phylogeny reconstruction

Phylogeny reconstruction was carried out with phylogenetic network, non-hierarchical and hierarchical Bayesian approaches. Under the non-hierarchical Bayesian approach, phylogenetic trees were estimated with MrBayes v.3.1. (Ronquist and Huelsenbeck, 2003). Following the conditional combination method described in the introduction, only the two mitochondrial genes should be concatenated. However, in order to be able to compare the results over the three methods, we still estimated trees from all five nuclear genes concatenated, and from the combined nuclear and mitochondrial genes. For each analysis, the software ran for 25 million generations, with a sample taken every 1000 generation, in two parallel runs of four Metropolis Coupled Monte Carlo Markov Chains (one cold and three incrementally heated). Stabilization of the likelihood occurred well before the runs were terminated. Convergence between runs was good, with an average standard deviation of split frequencies <0.01 (Ronquist and Huelsenbeck, 2003). The first quarter of the trees was discarded as burn-in and the remaining trees were summarized under the majority-rule consensus criterion.

Phylogenetic networks were constructed using the NeighborNet algorithm as implemented in Splitstree v.4.10 (Huson and Bryant, 2006) for (1) each of the five nuclear genes, (2) the five nuclear genes combined, (3) the two mtDNA genes combined, and (4) all seven genes combined. The statistical confidence in inferred nodes was assessed through 1000 bootstrap pseudo-replicates. A hierarchical Bayesian method was used to reconcile phylogenetic signal across gene trees into a species tree. Individuals were assigned to a particular species or subspecies, a requirement of the software, based on morphology and geographical origin. Analyses were conducted for only the nuclear genes, and for the complete dataset. BEST was run with topology, branch lengths, gene mutation rates, gamma shape parameter, character state frequencies, transition/transversion ratio, substitution rates under the GTR model, and the proportion of invariant sites set to be unlinked across genes. The chain length used was 300 million generations, with one sample taken every 1000 generations. The last 10,000 trees sampled from the stationary phase were summarized under the 50% majority-rule consensus criterion.

In this paper, we will use the term ‘species trees’ to define a phylogenetic tree calculated from a method that acknowledges heterogeneity in gene trees (Edwards, 2009), whereas ‘concatenation’ is used to define the construction of a supermatrix containing all available data.

2.5. Molecular dating

Molecular clock estimates for salamandrids are available for the ND2 gene, but vary considerably. They are contained between the extremes 0.37 (Mueller, 2006) and 0.64 (Weisrock et al., 2005) substitutions/site/100 Ma. Several other papers have used fossil calibrations and/or biogeographical data to estimate divergence times in the family Salamandridae. Steinfartz et al. (2007) used a fossil calibration of 24 million years before present (Ma) at the root of Triturus (their calibration point C4). Zhang et al. (2008) estimated the divergence of Salamandridae based on two sets of calibration: one with similar fossil calibrations as Steinfartz and another with an additional biogeographical calibration point for the split between Corsico-Sardinian Euproctus and continental Triturus. These two sets of calibrations gave an estimate for the root of Triturus of 26.6 and 11.3 (Zhang et al., 2008). We primarily followed the calibration point of 24 Ma of Steinfartz to date divergences on the species tree, and then compared the results to the

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**Table 1**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Symbol</th>
<th>Primer</th>
<th>Sequence (5’–3’)</th>
<th>Annealing (°C)</th>
<th>Reference</th>
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<td>Espregueira Themudo et al. (2009)</td>
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<td>BF CRI 1R</td>
<td>GCA CAC TGT GCT AAT CCT C</td>
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<td>PDGFRα</td>
<td>PDGFRa Fa</td>
<td>GTC ATT GAC TCC ATG CCT C</td>
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<td>Espregueira Themudo et al. (2009)</td>
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<td>CalC 3F</td>
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<td>CRI1 1Fa</td>
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<td>ND4</td>
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<td>TCT GGG TGC CAT TCA GAA GA</td>
<td></td>
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</tr>
</tbody>
</table>

**Reference**

(1) Input data from the following studies: Platelet-derived growth factor receptor-α: Babik et al. (2005); Calreticulin intron C: Arntzen et al. (2007). The partition homogeneity test (Farris et al., 1994) as implemented in PAUP 4.0b10 was used to check for congruence in phylogenetics signal among genes. Phylogenetic congruence was rejected for all pairs of markers, except the two mitochondrial genes. Nucleotide composition, transition/transversion ratio and sequence divergence under the Kimura 2-parameter model were calculated with MEGA4 (Tamura et al., 1994) as implemented in PAUP 4.0b10 (Swofford, 2003). The first quarter of the trees was discarded as burn-in and 25 million generations, with a sample taken every 1000 generations, the chain length used was 300 million generations, with one sample taken every 1000 generations. The last 10,000 trees sampled from the stationary phase were summarized under the majority-rule consensus criterion.

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other possible calibrations. The divergence times of the various lineages were determined from the relative branch lengths in the consensus tree by linear interpolation.

3. Results

3.1. Polymorphism and level of variation

The aligned sequence matrix (with indels removed) has a length of 4336 bp, with 694 variable positions of which 373 are parsimony-informative (Table 2). With 288 (77%) versus 85 (23%) variable positions, phylogenetic information content is higher in the two mitochondrial genes than in the five nuclear genes. The G-content was higher in nuclear genes (17.4–26.3%) than in mitochondrial genes (13.3–16.4%), as expected from the anti-G bias in mtDNA (Zhang and Hewitt, 1996). Cri4 showed the lowest level of variation and ND2 and ND4 the highest. The level of nucleotide substitution saturation was classified as ‘little’ for each gene. No significant signal for recombination was detected. The selected substitution models were F81 + I for int7, HKY + G for CalintC, PDGFRα, Cri1 and Cri4, and a mixed model for the combined mtDNA genes: GTR + I + G for ND2 and GTR + I for ND4. Across nuclear genes heterozygosity ranged from 13% to 47%.

3.2. Phylogenetic reconstruction

Non-hierarchical Bayesian phylogeny—In the mtDNA gene tree all individuals are grouped into their nominal species and subspecies, with a Bayesian posterior probability (pp) of 1.0 (Fig. 2A). The mtDNA data gives strong support to the sister-group relationship of T. carnifex and T. macedonicus (pp = 1.0) and weak to the two subspecies of T. karelinii (pp = 0.60). Triturus karelinii is the sister species to all other crested newts, but the support for this configuration is weak (pp = 0.78).

Across the nuclear genes, the results are not easy to reconcile, neither at the level of recognized species and subspecies, nor in terms of phylogenetic structure (Fig. 2B–F). Nodes with high posterior probability (pp > 0.95) that also support nominal (sub)species are found for T. carnifex (PDGFRα and Cri4), T. macedonicus (CalintC, PDGFRα and Cri4), T. k. arntzenii (βfib7int7, CalintC and Cri1) and T. dobrogicus and T. k. karelinii (all genes except βfib7int7).

Inter(sub)specific phylogenetic relationships supported with pp > 0.95 are T. dobrogicus and T. k. arntzenii by Cri1 (with exclusion of T. k. karelinii) and T. carnifex, T. cristatus, T. macedonicus and T. k. arntzenii by CalintC (with exclusion of T. k. karelinii).

The combined nuclear tree (Fig. 3B) shows all nodes resolved with a pp of 1.0. Triturus carnifex and T. macedonicus are sister species. These two species are sister to T. cristatus, and Triturus dobrogicus is sister to this group. Triturus. k. karelinii and T. k. arntzenii are sister taxa. The tree resulting from the complete dataset differs from the one in Fig. 3B only by a lower support (pp = 0.95) for the ((T. carnifex + T. macedonicus) + T. cristatus) clade (Fig. 3A).

Phylogenetic networks—Networks for the five nuclear genes separately show mixed results, with the occasional high support for the predefined taxa, such as T. carnifex, T. dobrogicus and T. macedonicus by PDGFRα (results not shown). Conversely, individuals representing a nominal species might also be found scattered over the network, such as T. carnifex for Cri4, T. dobrogicus for βfib7int7 and T. k. karelinii for CalintC, Cri1 and Cri4. The networks show generally low phylogenetic structure at the (sub)species level, with the exception of the gene PDGFRα, which supports a clade consisting of T. carnifex, T. cristatus, T. dobrogicus and T. macedonicus with a low bootstrap replication scores (brs) = 0.65 (results not shown). The phylogenetic network derived from the combined dataset resolves five species of crested newts in line with current taxonomy (Fig. 4C). Substructuring is found for T. karelinii with two groups that concur with current subspecies designations (T. k. karelinii and T. k. arntzenii). Bootstrap replication scores that would support phylogenetic structure among these taxa vary. Triturus karelinii is either monophyletic and differentiated from the other taxa (brs = 1.0), or paraphyletic (brs = 0.97) with just T. k. karelinii differentiated from all the other taxa. Moderate support is found for a clade consisting of T. carnifex, T. cristatus, T. macedonicus and T. dobrogicus (brs = 0.90). Within this clade, only the sister-group relationship of T. carnifex and T. macedonicus is strongly supported (brs = 1.0). A network based on the two mtDNA genes has the same topology but differs slightly in bootstrap replication scores (Fig. 4B). Based on the five nuclear genes, the network has the same branching order, with high support for the (sub)species, but low for interspecific relationships (Fig. 4A). The overall distances among taxa are shorter, due to the lower number of informative characters.

| Table 2 | Haplotype and alignment length, nucleotide composition, number and size of indels, number of variable sites (for the crested newt group, the genus Triturus, and parsimony-informative), observed heterozygosity (Ho), starting position of first codon, and level of substitution saturation as quantified by DAMBE (Xia and Xie, 2001) for each gene and the complete dataset. |
|---|---|---|---|---|---|---|---|---|---|---|---|---|
| Nomenclature | βfib7int7 | CalintC | PDGFRα | Cri1 | Cri4 | ND2 | ND4 | ND2* | Total |
| Alignment length | 514 | 533 | 663 | 511 | 512 | 615 | 4480 |
| After excluding indels | 4336 |
| Nucleotide composition (%) | | | | | | | | | |
| T | 31.5 | 32.9 | 32.3 | 22.5 | 26.5 | 27.8 | 26.3 | 28.4 |
| C | 20.4 | 17.8 | 21.7 | 22.0 | 22.1 | 26.1 | 28.1 | 23.4 |
| A | 30.0 | 25.7 | 28.6 | 29.2 | 31.4 | 29.7 | 32.3 | 29.9 |
| G | 18.1 | 23.6 | 17.4 | 26.3 | 20.1 | 16.4 | 13.3 | 18.4 |
| Number of indels | 1 | 9 | 9 | 6 | 3 | 0 | 1 | 29 |
| Min.–max. indel size | 14 | 1–24 | 1–12 | 1–12 | 1–2 | 0 | 1 | 1–24 |
| Number of variable sites | | | | | | | | | |
| Crested newt group | 24 | 23 | 43 | 19 | 21 | 137 | 228 | 495 |
| Triturus | 37 | 44 | 59 | 33 | 37 | 177 | 307 | 694 |
| Parsimony informative | 8 | 13 | 36 | 12 | 16 | 112 | 176 | 373 |
| Ho | 0.40 | 0.40 | 0.20 | 0.47 | 0.13 | N.A. | N.A. | 1.00 |
| Codon start | N.A. | N.A. | N.A. | N.A. | N.A. | 3rd position | 3rd position |
| Substitution saturation | Little | Little | Little | Little | Little | Little | N.A. | N.A. |

N.A., not applicable.  
* Includes 53 bp of tRNA-met + ND2 + 37 bp tRNA-trp.
Hierarchical Bayesian phylogeny—The species tree is fully resolved (pp > 0.99), except for the position of T. cristatus (Fig. 5A). The supported clades are (i) T. carnifex and T. macedonicus, (ii) group i and T. dobrogicus, and (iii) T. k. karelinii and T. k. arntzenii. When only the nuclear genes are used to build the species tree, the topology is identical, but pp values are lower (Fig. 5B).

3.3. Divergence times

The molecular clock test in DAMBE does not reject the null hypothesis of equidistance of all tips to the root of the tree. Linear interpolation on the species tree (Fig. 5A) over the 24 Myr window that covers the radiation of the genus Triturus, based on the fossil calibration (Fig. 5A) yields the following estimates of divergence times within the crested newt group: 11.2 Ma for the MRCA of the crested newts (and, by consequence, for the branching off of T. cristatus, which stems from the root), 11.1 Ma for the divergence between T. k. karelinii and T. k. arntzenii, 8.0 Ma for the origin of T. dobrogicus and 7.9 Ma for the split between T. carnifex and T. macedonicus. As molecular clock calibrations for ND2 vary considerably, the time to the MRCA of crested newts is estimated at 14–28 Ma under these calibrations, which would multiply each of the above temporal estimates by a factor 1.25–2.5.

4. Discussion

4.1. Rapid radiations, the coalescence, mitochondrial versus nuclear genes, and concatenation

The evolutionary history of rapid radiations is difficult to resolve. During the short timespan between branching events, there is limited opportunity for phylogenetic signal to accumulate. Fur-
Moreover, the older the radiation, the more difficult it will be to uncover the order of branching events, as the relative amount of informative phylogenetic change decreases through time (i.e., anagenesis masks cladogenesis). In order to resolve such apparent polytomies, large amounts of data will generally be required.

Coalescence is the merging of genetic lineages backwards in time to a most recent common ancestor. Forward in time, the sorting process taking place during diversification from the ancestral lineages to the pattern observed today is a chance process. Because of this stochasticity, and because lineage sorting may be incomplete, gene trees will generally be different from one another and frequently not represent the species tree (Avise and Wollenberg, 1997). The only way out of this dilemma is to study multiple genes because the more genes involved, the more convincing the convergence towards the species tree will be. By studying many genes together, the ‘gene tree versus species tree’ problem should ultimately be eliminated.

Fig. 4. Results of a phylogenetic network analysis for the genus Triturus from DNA sequence data with T. marmoratus as outgroup. Results are shown for the concatenation of five nuclear genes (A), two mitochondrial genes (B) and all seven genes (C). Values on branches are bootstrap replication scores. Phylogenetic networks do not require congruence between fragments and conflict in the dataset is represented graphically through closed networks (or ‘rings’), indicating that interspecific relationships are not fully resolved (Huson and Bryant, 2006).

Animal molecular phylogenetics, in former times, largely relied on just the mitochondrial genome, with two main drawbacks. First, because of its small size (c. 16 kb in vertebrates) it is quickly exhausted in terms of data yield compared to the nuclear genome (for comparison Triturus newts have a nuclear genome size of c. 25 Gb; Gregory, 2008; even though a large portion is repetitive DNA and phylogenetically not-informative). Second, the mitochondrial genes are linked. The molecule is inherited as a single unit and mitochondrial gene trees are, hence, not independent. Indeed, recent studies on the phylogeny of the Salamandridae provide similar results whether based upon a small number of representative genes (Steinfartz et al., 2007) or on the complete mitochondrial genome (Zhang et al., 2008). In practice, this means that nuclear genes have to be used to get independent estimations. Unfortunately, primers for nuclear genes will often have to be newly developed (for Triturus see Espregueira Themudo et al., 2009).

4.2. Multiple independent markers solve the phylogeny of Triturus newts

The non-hierarchical Bayesian phylogenetic reconstruction yielded five nuclear gene trees and one mitochondrial tree that were all different from each other, to the extent that a consensus tree would not be informative. A striking result is that individual sequences do group according to predefined species and subspecies in the mtDNA-based tree, while for nuclear genes this is rarely the case. Possible explanations are (i) better phylogenetic resolution due to a larger number of informative characters and (ii) lineage sorting has progressed more rapidly, due to a fourfold smaller effective population size (Moore, 1995). Either way, the observed species level consistency in the individual phylogenetic placement suggests the absence or low frequency of horizontal gene exchange, which we had aimed for through a thoughtful sampling scheme, relying on the well documented species ranges (Arntzen, 2003).

The total evidence tree is completely resolved, and is identical to the combined nuclear tree, except for the lower statistical support for the position of T. cristatus (Fig. 3B). Compared to the total evidence tree, the mtDNA tree has the basal branches unresolved (Fig. 2A). This means that the nuclear data contributes significantly to the resolution of the species tree, even though this is not obvious from a gene by gene perspective.

The phylogenetic relationships supported by our 1747 bp of mtDNA data are the sister-group status of T. carnifex and T. macedonicus and T. k. karelinii and T. k. arntzenii (with pp > 0.99), while moderate support was for the clade including all crested newts that are not T. karelinii (pp = 0.78). The basal dichotomy between T. karelinii versus non-T. karelinii was not found by earlier, smaller molecular datasets (e.g. Macgregor et al., 1990; Wallis and Arntzen, 1989). Based upon 839 bp of cytochrome-b sequence data, Steinfartz et al. (2007) described the paraphyly of T. karelinii, whereas Arntzen et al. (2007) found no resolution at the basal Triturus node, from either 40 allozyme loci or 642 bp of ND4 mtDNA. The two T. karelinii subspecies (arntzenii and karelinii) are almost as old as the T. karelinii lineage itself.

In the phylogenetic network analysis, pronounced ambiguities in the total evidence approach are the unresolved branching order (T. dobrogicus, T. cristatus, T. carnifex, T. macedonicus) and the position of the T. karelinii subspecies as either monophyletic or paraphyletic (Fig. 4C). The next highest uncertainty refers to the monophyly of the group T. cristatus, T. carnifex, T. dobrogicus and T. macedonicus and, following that, the monophyly of T. carnifex and T. macedonicus.

In contrast with the above results, the phylogenetic tree obtained with the hierarchical Bayesian method is fully resolved, except for the position of T. cristatus (Fig. 5A). The newly obtained phylogenetic resolution supports the notion that the radiation of crested newts lineages occurred over a relatively short period. This window of radiation, dated at 11–8 Ma (or 14–10 Ma to 28–20 Ma using the varying ND2 molecular clocks cited above), yielded all six different lineages that we observe today.

This study provides empirical support for the claim (Edwards, 2009; Edwards et al., 2007) that the method, by accounting for the differential coalescent process across independent markers, estimates the phylogenetic relationships between recently diverged taxa with high precision. Statistical support expressed by posterior probability values is high. This is a remarkable improvement over competing methods (Fig. 2 and 4). We only found a fully bifurcating and supported tree with the non-hierarchical method by concatenating incongruent markers, thus violating the prerequisites of conditional combination. A potential drawback of the hierarchical method is that additional information should be provided in the form of species identification of the individuals included. Crested newt species show distinct morphological characteristics, making species assignment straightforward. Furthermore, our sampling scheme was designed to exclude areas if interspecific gene flow. Indeed, none was detected in the mtDNA marker (Fig. 2A). Despite this, distinguishing between interspecific gene flow and incomplete lineage sorting is a complex issue (Avise et al., 1983; see also Hey and Nielson, 2004; Sang and Zhong, 2000). We are currently exploring this area of research in Triturus newts, by sampling from areas potentially subject to introgression, i.e., within 100 km of described range borders (Espregueira Themudo et al., in preparation).

As predicted by Edwards (2009), statistical support in the species tree from the hierarchical method is less than the with concatenation. The theoretical advantages of the hierarchical method over data concatenation, however, make us believe that the outcome of the hierarchical Bayesian method is more reliable.

Concatenation is also prone to “data swamping”, where one or a few of the partitions provides most of the information retained in the phylogenetic tree. This does not seem to be problematic here, as we have shown that the nuclear genes are contributing significantly to the resolution of the total evidence tree (Fig. 3A and B). In the hierarchical method, the effect of the number of informative sites per locus is still not clear (Edwards, 2009).

4.3. Phylogenetic position and taxonomic status of T. karelinii

Marked levels of genetic differentiation within T. karelinii were first apparent from mtDNA, as either monophyletic (Wallis and Arntzen, 1989) or paraphyletic (Steinfartz et al., 2007). Interestingly, the phylogenetic network gives high bootstrap support to
both the monophyly and the paraphyly of *T. karelinii* (Fig. 4A). This type of uncertainty, expressed in so called phylogenetic ‘spiderwebs’, may be indicative of reticulate evolution, c.q. hybridization (Huson and Bryant, 2006).

Based on differences in genome size, protein variation and morphological characteristics, Litvinchuk et al. (1999) described a Balkan population as the representative of a distinct subspecies, *Triturus karelinii arntzenii*. The present study suggest that the two taxa are genetically distinct, at the same level or deeper than other *Triturus* species are to each other, with a date of origin close to the basal radiation of the crested newt group (Fig. 4). Considering their long independent evolutionary history, we suggest that the two taxa should be raised to full species status, i.e., *T. karelinii* (Strauch, 1870) and *T. arntzenii* Litvinchuk, Borkin, Đukić and Kalezic, 1999 (in Litvinchuk et al., 1999).

*Triturus arntzenii* and *T. karelinii* may have their range limits at either side of the Bosphorus (Raffaelli, 2007), but geological information does not support this view, since the current connection of the Black Sea to the Mediterranean opened a mere 5000 years ago (Ryan et al., 2003). Other data suggest that the species range border could be either in Europe or in Asia. A population from the Black Sea to the Mediterranean opened a mere 5000 years since the Miocene (Elmas, 2003). A wider analysis on the spatial distribution of morphological and genetic variation in the *T. karelinii* group is underway (Olgun et al., unpublished data; Wielstra et al., unpublished data).

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Appendix 1. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.mpev.2009.03.024.

References


