The Development of Three Long Universal Nuclear Protein-Coding Locus Markers and Their Application to Osteichthyan Phylogenetics with Nested PCR

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Abstract

**Background:** Universal nuclear protein-coding locus (NPCL) markers that are applicable across diverse taxa and show good phylogenetic discrimination have broad applications in molecular phylogenetic studies. For example, RAG1, a representative NPCL marker, has been successfully used to make phylogenetic inferences within all major osteichthyan groups. However, such markers with broad working range and high phylogenetic performance are still scarce. It is necessary to develop more universal NPCL markers comparable to RAG1 for osteichthyan phylogenetics.

**Methodology/Principal Findings:** We developed three long universal NPCL markers (>1.6 kb each) based on single-copy nuclear genes (KIAA1239, SACS and TTN) that possess large exons and exhibit the appropriate evolutionary rates. We then compared their phylogenetic utilities with that of the reference marker RAG1 in 47 jawed vertebrate species. In comparison with RAG1, each of the three long universal markers yielded similar topologies and branch supports, all in congruence with the currently accepted osteichthyan phylogeny. To compare their phylogenetic performance visually, we also estimated the phylogenetic informativeness (PI) profile for each of the four long universal NPCL markers. The PI curves indicated that SACS performed best over the whole timescale, while RAG1, KIAA1239 and TTN exhibited similar phylogenetic performances. In addition, we compared the success of nested PCR and standard PCR when amplifying NPCL marker fragments. The amplification success rate and efficiency of the nested PCR were overwhelmingly higher than those of standard PCR.

**Conclusions/Significance:** Our work clearly demonstrates the superiority of nested PCR over the conventional PCR in phylogenetic studies and develops three long universal NPCL markers (KIAA1239, SACS and TTN) with the nested PCR strategy. The three markers exhibit high phylogenetic utilities in osteichthyan phylogenetics and can be widely used as pilot genes for phylogenetic questions of osteichthyan at different taxonomic levels.

Introduction

Over the past two decades, nuclear protein-coding locus (NPCL) markers have become popular tools for inferring the evolutionary history between vertebrate species at different taxonomic levels [1–10]. NPCL markers are based on nuclear exons, but these exons are usually short (less than 500 bp). Therefore, most NPCL markers are short and cannot provide sufficient information to resolve relationships among broadly diverged vertebrate taxa, i.e., taxa that diverged 20–420 Ma (million years ago). For example, the popular marker c-mos is often represented by very short (375 bp) fragments. In general, long markers comprise more phylogenetic signals than short ones, the resolution of resulting trees from long markers is normally higher than those from short ones. Therefore, for practical purposes, systematists are commonly willing to use some long markers (>1,000 bp) with tested good phylogenetic performance to address their questions at hand first.

RAG1 is one commonly used long NPCL marker. It takes advantage of a long (~3 kb) and uninterrupted exon that is found across osteichthysans, has an overall evolutionary rate that is appropriate for evolutionary events from 20 to 420 Ma and furthermore, contains slightly faster- or slower-evolving regions that could resolve problems at different taxonomic levels. Due to these advantages, RAG1 has been widely used for osteichthyan phylogenetic studies, and more than 15,000 RAG1 sequence records have been deposited in the NCBI GenBank. If there are more NPCL markers like RAG1, which can be easily amplified across osteichthysans and are long enough to be phylogenetically informative at different taxonomic levels, people can use them to quickly investigate framework relationships for many taxa of interest. However, such long universal NPCL markers remain...
relatively limited. Therefore, it is necessary to develop more number of long universal markers suitable for osteichthyan phylogenetics. The first step toward developing these markers is to locate exons that are of the appropriate length (more than 3 kb), that are uninterrupted by introns in a diverse range of osteichthyan taxa, and that contain both fast- and slow-evolving regions.

In our previous study [10], we analyzed multiple genome alignments and developed 21 new NPCL markers for use in tetrapods. However, these markers are not long enough (normally <1,000 bp) and are difficult to be applied in bony fishes and amphibians. Therefore, we reinvestigated these 21 nuclear protein-coding genes based on a large set of genome data available from the ENSEMBL database. We found that three single-copy genes (KIAA1239, SACS and TTN) contain large exons (3.9 kb, 11.6 kb and 17.1 kb, respectively) and are fairly well conserved from ray-finned fishes to mammals. Further analyses of these exons indicated that they contain regions with variable evolutionary rates. These properties make these three nuclear genes potential candidates for long universal NPCL markers.

One of the difficulties in developing universal NPCL markers is in ensuring a high success rate of PCR amplification across divergent taxa. In general, degenerate primers are designed based on the conserved protein sequences. However, primers with high degeneracy often lack of amplification specificity, producing many non-specific amplicons or amplification failures. In contrast, primers with low degeneracy typically only work in a subset of samples due to a lack of sensitivity across diverse taxa. For example, Fong and Fujita [11] explored 75 new protein-coding genes across vertebrates and tested degenerate primers in three species, but nearly 53% of the tested fragments were not amplified successfully. Recently, nested PCR has been popular for amplifying specific sequences [1,12–14]. Nested PCR is a modification of standard PCR that uses two sets of primers in two separate PCR rounds to amplify the target fragment, in which the product of the first round of PCR serves as the DNA template for the second round of PCR. The advantage of nested PCR is that it is extremely sensitive and specific when amplifying target sequences from complex genomic environments compared to standard PCR. Therefore, evaluating the technical differences between nested PCR and standard PCR will have practical implications on the application of NPCL markers and the development of universal markers.

With the advent of next generation sequencing (NGS) technologies, phylogenomic studies based on whole genome sequences or transcriptomes are becoming more and more common. Nevertheless, although the NGS-based approach is a promising way to reconstruct the tree of life, taxon sampling in such studies are normally restricted because of economic consideration and the difficulties on sample manipulations. In contrast, the conventional PCR-based method is still a more practical and cost-efficient way to generate sequences for many taxa. On the other hand, considering the bulk of worldwide museum-preserved specimens, the PCR-based method seems to be the only solution at present to analyze those samples. In this study, our goal is to increase the number of long universal NPCL markers comparable to RAG1 that can be used as "standard and pilot" markers for quick phylogenetic investigations among osteichthians at different taxonomic levels. We investigated the phylogenetic utility of three long NPCL markers (KIAA1239, SACS and TTN) together with the reference marker RAG1 among osteichthians. Meanwhile, we compared the ease of amplification of the three new NPCL markers in nested PCR and standard PCR. We showed that these three long NPCL markers are useful tools for phylogenetic studies of osteichthians at broad taxonomic levels and that the nested PCR strategy is much more sensitive and specific than the conventional PCR strategy.

Results

General Features of the Large Exons in the Four Marker Genes

The lengths of the large exons in RAG1, KIAA1239, SACS, and TTN that were used for the development of NPCL markers are approximately 3 kb, 3.9 kb, 11.6 kb and 17.1 kb, respectively. These exons are not interrupted by introns in any of the 16 osteichthian species observed. The conservation profile for each exon is described by the conservation diagram presented in Figure S1. Regions with a high density of long black bars indicate that the genetic divergence across the given taxa is low and that sequences are highly similar across species. For the reference gene RAG1, the conservation profile across the whole exon is not uniform; the first third of the gene (20–1,200 bp) is fairly variable (Fig. S1). In contrast, KIAA1239, SACS and TTN are more uniform (Fig. S1), indicating that these exons are more suitable for the development of universal NPCL markers. In addition, we evaluated the overall mean distances (evolutionary rates) for each of four exon alignments in MEGA 5 [15]. The overall mean distances of these three exons are similar to each other (0.295 in KIAA1239, 0.333 in SACS, 0.327 in TTN), while RAG1 has a higher value, 0.423. However, if the fast-evolving forward third region of the RAG1 exon is removed, the overall mean distances of RAG1 decrease to 0.316, similar to the values measured for the three new genes.

Summary of PCR Amplifications

A total of 14 primer pairs were used to amplify the four long NPCL markers: 10 were newly designed, and 4 were published in previous studies [10,16]. The lengths of the target PCR fragments ranged from 890 to 1,210 bp. We successfully obtained a 1,488-bp fragment for RAG1, a 1,737-bp fragment for KIAA1239, a 2,211-bp fragment for SACS and 1,680-bp fragment for TTN. Newly generated sequences were deposited in GenBank under accession numbers JN979993–JN980079, JQ929565–JQ929580 (see Table S1).

To compare the nested PCR and standard PCR strategies, we separately amplified two overlapping fragments for each of three long NPCL markers using both PCR methods. The results of these PCR experiments are summarized in Figure 1. We categorized the agarose electrophoretic images of the PCR products into three groups: no target band or smear, weak target band with non-specific amplification or amplification failures. Overall, the amplification success rate and efficiency of the nested PCR were overwhelmingly higher than that of the standard PCR. The PCR success rate was 100% for the nested PCR method but only 70.3% for the standard PCR method. Furthermore, the proportion of reactions yielding strong target bands with non-specific amplification in the nested PCR was notably higher than that in the standard PCR (96.4% versus 28.6%). Finally, some species (e.g., Batrachus formosanus, Protopterus annectens) are somewhat refractory to target band amplification using standard PCR, but nested PCR was able to produce strong or weak target bands for these difficult samples. The experimental differences between the two PCR methods are demonstrated visually in Figure 2, which shows an agarose gel used to separate the products of amplification of the first fragments of SACS.
**Phylogenetic Analyses**

The refined alignments of RAG1, KIAA1239, SACS, TTN are 1,488 nt, 1,737 nt, 2,211 nt, 1,698 nt in length, respectively. The BI and ML analyses on the concatenated dataset (7,134 nt) under the three different partitioning strategies (3-partition, 4 partition, and 12-partition) produce the same topology and similar support values. Most nodes (85%) are strongly supported with $95\%$ bootstrap values (BS) and 1.0 Bayesian posterior probabilities (PP) (Fig. 3). The well-resolved tree inferred from the concatenated dataset (Fig. 3) generally agrees with the currently accepted osteichthyan phylogeny.

For each of the four NPCL markers (RAG1, KIAA1239, SACS and TTN), both partitioned BI and ML yielded almost identical trees with similar branch support values (see Figs. S2, S3, S4, S5). In all analyses, the monophyly of six animal groups (Actinopterygii, Squamata, Testudines, Aves, Crocodylia and Mammalia) are strongly supported (ML bootstrap $\geq 95\%$ and Bayesian PP $= 1.0$; Figs. S2, S3, S4, S5). The monophyly of Lissamphibia was not recovered by RAG1 (Fig. S2) but was well supported by KIAA1239 (ML bootstrap $= 95\%$; Fig. S3) and SACS (ML bootstrap $= 89\%$; Fig. S4) and weakly supported in TTN (ML bootstrap $= 50\%$; Fig. S5).

Phylogenetic relationships estimated from single long NPCL markers alone are generally similar to those estimated from the concatenated dataset. However, several relationships with weak support are incongruent among RAG1, KIAA1239, SACS and TTN (Figs. S2, S3, S4, S5). These conflicts are found mainly in currently uncertain relationships, such as the interrelationships within Neoaves, the early splitting Squamata, the placement of Testudines and the relationship between Acipenseriformes, Lepisosteiformes and Teleostei.

**Characteristics of the Four Long Universal NPCL Markers**

Figure 4 shows the phylogenetic informativeness (PI) profile curves for the four long NPCL markers tested in this study. According to these curves, the phylogenetic performance of the three developed NPCL markers are generally comparable (or even higher; SACS) to that of RAG1. Klopfstein et al. [17] argued that estimating phylogenetic informativeness profile (PI) of the marker is needed to take cautions when including more than 4 taxa. Therefore, it is necessary to compare the information content of each marker with more indexes. We thus estimated a series of indicating parameters for each marker such as GC content, gamma shape parameter ($\alpha$), proportion of invariable sites ($\Pi_{\text{var}}$), relative substitution rate, proportion of internal branch length (Treeness), and relative composition variability (RCV) (see Table 1). For each marker, none of the parameters have significant correlation with others. For example, KIAA1239 has the lowest alpha value, but shows high value of proportion of invariable sites. For relative substitution rate, four NPCL markers show slight variations, suggesting that they have similar evolutionary rates. Treeness is an indicator used by Phillips and Penny [18] to evaluate phylogenetic signal strength. The four NPCL markers also have similar Treeness values. In general, lower RCV value of a marker means a lower chance interfered by compositional bias. The RCV values of the three NPCL markers are lower than that of the reference marker RAG1. In brief, besides the PI profiles, all estimated characteristical parameters suggested that the three
Figure 2. Agarose gel electrophoretic analysis of the PCR products. The first fragments of the long NPCL marker SACS (SACS-F1) were amplified in 32 taxa using nested PCR and standard PCR, respectively. The upper image shows the results of nested PCR amplifications, and the lower image shows the results of standard PCR amplifications. Lanes 1–32 show identical PCR amplifications performed in different species. "-": negative controls, "M": DNA ladder.

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Figure 3. Phylogram derived from analysis of the concatenated four long NPCL markers. Phylogenetic relationships among osteichthyans were based on maximum likelihood and Bayesian inference analyses of the combined data set (7,134 bp) under 3-partition and 4-partition strategies (by codon and by gene). The two quarter circles above the branches represent the bootstrap proportions for partitioned ML analyses and the two quarter circles below branches represent the Bayesian posterior probabilities for partitioned BI analyses. Branch lengths were estimated in the 4-partition ML (by gene) analysis on a concatenated dataset (4 GTR +\Gamma +I models for 4 gene partitions).

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Figure 4. Phylogenetic informativeness (PI) profile of the reference NPCL marker RAG1 and three long NPCL markers. The timetree was newly estimated based on the concatenated dataset (7,134 bp). The PI profile was generated using the online program PhyDesign [52]. For more detail, please see the text.
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Nested PCR markers have potential phylogenetic performance comparable to that of RAG1.

Discussion

Three Considerations for the Development of Universal NPCL Markers

Before developing NPCL markers, researchers should first consider whether the proposed marker is a member of a gene family. When targeting genes with a large number of family members, there is a high risk of amplifying paralogous genes, which have different evolutionary histories than orthologous genes and may thus hinder correct phylogenetic inference [19]. Thus, nuclear protein-coding genes with few family members, ideally single-copy genes, are the best choices for NPCL marker candidates. In our study, the recombination-activating gene (RAG) family contains two genes (RAG1 and RAG2), but RAG1 is often recognized as a “single-copy” gene because its sequence is very different from that of RAG2. In comparison with the “single-copy” gene RAG1, none of the three nuclear genes used here have paralogs, according to a search of ENSEMBL. Therefore, these genes are suitable for the development of NPCL markers.

Recent studies [20–23] have suggested that gene size (e.g., alignment length) is positively correlated with the phylogenetic performance. In other words, appropriately chosen long NPCL markers are sufficient to build a reliable phylogeny [2,6]. Moreover, if researchers develop relatively long NPCL markers, the accuracy of phylogenetic inference, particularly those based on supertree strategies will be improved. Therefore, locating large exons that are uninterrupted across diverged taxa should also be considered when developing NPCL markers.

Finally, two favorable properties (single or low-copy gene family size and large exons) alone cannot guarantee that the developed long universal NPCL markers will be useful for phylogenetic reconstruction. Researchers must also consider the phylogenetic informativeness of their markers at different taxonomic levels. Ideally, a good candidate exon for the universal NPCL marker development should contain both slowly evolving regions and fast evolving regions to provide enough information for both deep and young nodes.

The Advantage of Nested PCR

One important criterion for NPCL markers is that they should be easily amplifiable among the taxa of interest and in other groups. The low success rate of PCR amplification, however, usually limits the range of applicability of NPCL markers. Thus, identifying a PCR strategy with high amplification efficiency is an important step toward improving the applicability of NPCL markers. Nested PCR, a modification of standard PCR, has shown to be an extremely sensitive and specific method for amplifying target sequences [1,24]. In this study, we compared the amplification efficiency of our three long NPCL markers using nested PCR and standard PCR. Our results (Fig. 1 and Fig. 2) show that the amplification efficiency of nested PCR is apparently higher than that of standard PCR.

The nested PCR strategy has been used in previous phylogenetic studies [1,13,14,24], but only for fragments that could not be amplified successfully by standard PCR. In this study, we took the nested PCR strategy as standard procedure to amplify NPCL fragments across diverse taxa from chondrichthyines to mammals. As a result, we successfully obtained all target fragments with ease, including some refractory ones in our previous study [10]. Therefore, we strongly recommend that researchers choose a nested PCR strategy rather than a standard PCR strategy when developing new phylogenetic markers or working with difficult samples.

Implications for Osteichthyan Systematics

In general, the four independent long NPCL markers and the 4-gene concatenated datasets produced similar phylogenies for 46 tested osteichthyan species (Figs. 3, S2, S3, S4, S5). In the concatenated tree (Fig. 3), nearly all nodes are strongly supported and the interrelationships between 46 tested osteichthyans are consistent with currently accepted hypotheses. For example, our concatenated data reveals two major monophyletic clades within Teleostei: the first clade (Ostariophysi) includes members of Siluriformes and Cypriniformes, the second clade (Percomorph) comprises members of Beloniformes, Synbranchiformes, Gasterosteiformes and Tetraodontiformes. The two major clades were also found in recent multigene studies [7,25,26]. Within Lissamphibia, the combined data robustly recovers a sister-group relationship between frogs and salamanders, and is in agreement with most recent studies [8,27–31]. Within Mammalia, our concatenated data firmly shows that Monotremata (monotremes) is the sister group of other mammals and Proboscidea branched first within placentalts. Recently, Prasad et al. [32] employed 60 megabase pairs (Mb) of genomic sequences to investigate relationships for 41 mammal species. Their results also placed Monotremata (monotremes) as sister group to other mammals and Proboscidea (Xenarthra) as basal branch of placental mammals. For the avian phylogenetic tree, recent molecular studies have consistently pointed out that Aves are divided into three major Superorders: Palaeognathae, Galloanserae and Neoaves [2,33,34] and Palaeognathae as basal group in avian phylogeny. Our results also confirm this relationship. Besides the interrelationships found among turtle species are also consistent with the well-resolved phylogeny of extant turtles based on analyses of single mitochondrial or nuclear gene, mitochondrial genomes and concatenated datasets [35–38].

The concatenated tree also have several nodes without strong support (Fig. 3) that reflect currently uncertain relationships as discussed below.

Within Neoaves, the concatenated data is unable to give a decisive relationship among Columbiformes, Psittaciiformes and Passeriformes (Fig. 3). Indeed, the relationships between these avian lineages were also found to be controversial in recent nuclear and mitochondrial phylogenomic studies [34,39,40]. Hackett et al. [34] and Wang et al. [40] reported that Passeriformes and Psittaciiformes were sister groups with respect to Columbiformes based on 19 and 30 nuclear loci, respectively. This relationship was also found in our analyses of the concatenated dataset.

| Table 1. Characteristic information of the 4 NPCL markers. |
|----------------------------------------------------------|
| **Gene** | **Length (bp)** | **GC%** | **Alpha** | **Pinvar** | **Sub. rate** | **Treeness** | **RCV** |
|----------|----------------|--------|-----------|-----------|--------------|-------------|--------|
| RAG1     | 1,488          | 32.3   | 1.17      | 0.35      | 1.07         | 0.324       | 0.111  |
| KIAA1239 | 1,737          | 29.6   | 0.94      | 0.30      | 0.95         | 0.359       | 0.096  |
| SAC5     | 2,111          | 25.9   | 1.02      | 0.29      | 1.06         | 0.320       | 0.092  |
| TTN      | 1,698          | 26.4   | 1.01      | 0.25      | 0.91         | 0.356       | 0.070  |

Length, length of refined alignment; Alpha, shape parameter of the gamma distribution; Pinvar, proportion of invariant sites; Sub. Rate, relative substitution rate estimated by Mbayes; Treeness, proportion of tree distance on internal branches; RCV, relative composition variability.

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However, based on an analysis of 70 mitochondrial genomes to reinvestigate the interrelationships among major Neoaves, Pa-
checo et al. [39] suggested that Passeriformes was not a sister
group of Psittaciformes. The cause of this inconsistency is not yet
clear and deserves further exploration.

Within Squamata, the resulting tree indicates that Gekkota is the most basal lineage of living squamates but with only weak
support (Fig. 3). However, Vidal and Hedges [5] used 9 NPCL
markers to infer the relationships among the major Squamata, and
they argued that Dibamidae branched first within Squamata.

The uncertain placement of the Testudines has been debated in
various molecular studies [4,10,41–44]. To date, the most
powerful dataset is from our previous study [10], which analyzed
25 genes (21,137 bp) and produced a robust relationship as
(turtles, (birds, crocodilians)). Our concatenated analyses also
recovered the same relationship but without strong bootstrapping
support.

For the major actinopterygian relationships, mitogenomic data
and nuclear genes produced two different relationships between
Acienceriformes, Lepisosteiformes and Teleostei. Inoue et al.
[45] employed 28 mitogenomic sequences to investigate major
relationships among actinopterygians, and pointed out that
Acienceriformes and Lepisosteiformes group as "ancient fishes"
clade closely related to Teleostei. However, actinopterygians
phylogeny based on seven nuclear genes recognized Lepisoste-
iformes as close relatives of Teleostei but not sister group to
Acienceriformes [24]. Our combined analyses favor the latter
hypothesis, this result raises confidence in the use of the four
NPCL markers among actinopterygians.

Summaries and Recommendations

In this study, we presented three long universal NPCL markers
(KIAA1239, SACS and TTN; >1,600 bp each) with comparable
or better phylogenetic performance among osteichthyans to that
of the widely used RAG1. In addition, we evaluated the differences
between nested PCR and standard PCR when amplifying NPCL
marker fragments. The amplification success rate and efficiency of
the nested PCR are overwhelmingly higher than those of standard
PCR. By using the nested PCR strategy, the three long NPCL
universal markers can be easily amplified in osteichthyans with a
success rate of over 95%. Considering their good phylogenetic
performance and high usability, these markers can be widely used
as pilot genes for phylogenetic questions of osteichthyans at
different taxonomic levels. For example, when handling large-scale
studies with many taxa, people may quickly generate data for these
pilot genes with ease to identify which nodes are more difficult to
resolved, thus directing further actions.

Materials and Methods

Taxon Sampling and DNA Preparation

The classification and source or collection locality of the 46
osteichthyan species and one chondrichthyan outgroup species
used in this study are shown in Table S1. These taxa represent eight
major osteichthyan lineages (Actinopterygii, Dipnoi, Lissam-
phibia, Squamata, Testudines, Aves, Crocodylia and Mammalia).
To repress long-branch attraction (LBA) artifacts [46], we tried to
include more than one species for each major lineage Among the
47 selected taxa, public genome data were available for 16 taxa,
while sequences for the remaining 31 taxa needed to be generated
de novo. Total genomic DNA was extracted from ethanol-
preserved tissues (liver or muscle) using the standard salt extraction
protocol. All extracted genomic DNA was stored at -20°C prior to
PCR amplification. This study was performed in strict accordance
with the guidelines developed by the China Council on Animal
Care and Use. All animal processing procedures were approved by
the Institutional Animal Care and Use Committee of Sun Yat-Sen
University (permit number: 2011–023).

The Development of Three Long NPCL Markers

The nucleotide sequences of the largest exons in RAG1,
KIAA1239, SACS and TTN were retrieved from Ensembl for 16
osteichthyan species with available genome data: Danio rerio,
Orzyxias Latipes, Gasterosteus aculeatus, Tetradon nigroviridis, Takifugu
rubripes, Silurana tropicalis, Anolis carolinensis, Gallus gallus, Tannypoggiya
gutta, Ornithorhynchus anatinus, Monodelphis domestica, Loxodonta
africana, Equus caballus, Bos taurus, Mus musculus and Homo sapiens.
Each exon was aligned based on its translated amino acid
sequence, and the subsequent alignment was used for marker
development. Our goal was to develop long NPCL markers of over
1,500 bp. However, the size is too large to be sequenced from both
ends. Therefore, we divided a long target region into two
overlapping fragments of less than 1.3 kb each to facilitate
sequencing from both ends. In addition, the strategy of using
two overlapping fragments to cover a longer marker can also be used
to check for possible cross-contamination and to ensure data
quality. This design is because although the nested PCR used in
this study (see discussion above) is extremely sensitive for
amplifying target sequences from small amounts of samples, it
may increase the risk of cross-contamination during laboratory
analyses.

We manually selected a region with an appropriate evolutionary
rate for each of the three NPCL genes (KIAA1239, SACS and
TTN). Each selected region was divided into two overlapping
fragments (less than 1.3 kb) that have two conserved blocks
flanking less conserved regions. For each target fragment, we
designed primers for a nested PCR strategy. The first round PCR
primers were used to amplify a longer region containing the target
fragment. Because the first round of PCR is only used to increase
the concentration of effective DNA templates, we designed first
round primers with high degeneracy to match as many amino acid
sequences as much as possible, i.e., to increase primer sensitivity.
In contrast, the second round of PCR is used to amplify the target
fragment, and thus, we avoided designing primers in amino acid
residues with high degeneracy (e.g., L, R and S) to increase primer
specificity. All of the primers used in this study are listed in Table 2.

PCR Amplification, Cloning, and Sequencing

We amplified two overlapping fragments for each of three long
NPCL markers using both nested PCR and standard PCR to
evaluate the experimental differences between the two PCR
methods. Each pair of PCR primers was tested in 25-µL reaction
volumes with ExTaq DNA polymerase (Takara, Dalian). Negative
controls were also included in every PCR to monitor possible
cross-contamination. For the nested PCR, two separate runs (first
PCR and second PCR) were conducted. The first round of PCR
settings were as follows: an initial denaturation step of 4 min at
94°C; followed by 45 cycles of a 45 sec denaturation at 94°C, a
40 sec annealing at 45°C, and a 2 min elongation at 72°C;
followed by a final 10 min elongation at 72°C. The second round
of PCR used products of the first round PCR (without dilution) as
dNA templates and the following cycling conditions: an initial
denaturation step of 4 min at 94°C; 55 cycles of a 45 sec
denaturation at 94°C, a 40 sec annealing at 45°C, and a 2 min
elongation at 72°C; followed by a final 10 min elongation at 72°C.
For the standard PCR, primers from the second round of PCR
were used alone to amplify target fragment from genomic DNA,
and the reaction cycling settings were the same as those used for the second round of PCR.

The target PCR bands were purified by gel extraction and subsequently cloned into a PMD19-T vector (Takara, Dalian). Recombinant clones were identified by colony PCR. The resulting PCR products (at least two) were purified with ExoSap and sequenced in both forward and reverse directions with an ABI3730 DNA sequencer. All sequences were confirmed as the correct target fragments by BLAST search against the human genome. Finally, the two overlapping fragments for each NPCL marker were assembled into a contiguous fragment. No conflicts were observed within overlapping regions, indicating that the two fragments were correctly generated from the same species in all cases.

### Phylogenetic Analyses

All four NPCL markers (RAG1, KIAA1239, SACS, and TTN) were aligned using the G-INS-i method from MAFFT [47,48] for phylogenetic analysis. The analysis was performed to understand the evolutionary relationships among different species. Each long NPCL marker is amplified in two contiguous and overlapping fragments (F1 and F2). * indicates NPCL marker that is amplified using both nested PCR and standard PCR in order to compare amplification difficulties between two PCR methods. For nested PCR, 1st PCR and 2nd PCR represent two separate runs, products of 1st PCR (no dilution) are used as amplification templates in 2nd PCR; For standard PCR, primers in 2nd PCR are used alone to amplify target fragments.
under the default settings according to their translated amino acid sequences. Because these genes were well aligned, alignment refinements were done manually with MEGA 5 [15]. Finally, five DNA datasets (four independent alignments and one concatenated alignment) were prepared for phylogenetic analyses. The five datasets were separately analyzed with both maximum likelihood (ML) and Bayesian inference (BI) methods under partitioned strategies. For each of the four NPCL markers, we partitioned the dataset by codon (3 partitions). For the concatenated alignment (7,134 bp), we utilized three partitioning strategies. The first strategy used 3 partitions (one partition for each codon position); the second used 4 partitions (one partition for each gene); and the third used 12 partitions (codon position partitioning across four genes). The partitioned maximum likelihood analyses (-q option) were conducted using RAxML version 7.2.6 [49]. We used the GTR+Γ+I model for each partition. A search that combined 100 separate maximum likelihood searches was applied to find the optimal tree (-f d option), and branch support for each node was evaluated with 500 rapid bootstrapping replicates (-f a option) implemented in RAxML. The partitioned Bayesian inference was conducted in MrBayes 3.2 [50]. The best-fitting model for each partitioned dataset was estimated with MrModelTest2.3 [51] using the Akaike information criterion (AIC). Nearly all partitions partitioned dataset was estimated with MrModelTest2.3 [51] using the GTR+Γ+I model. Two MCMC runs (Unlink Revmat = [all] Statefreq = [all] Shape = [all] Pinvar = [all]) were performed with one cold chain and three heated chains (temperature set to 0.2) for 3 million generations and sampled every 100 generations. The chain stationarity was visualized by plotting -lnL against the generation number using Tracer version 1.4 [http:// evolve.zoo.ox.ac.uk/beast/help/Tracer], and the first 15–50% of generations were discarded. Topologies and posterior probabilities were estimated from the remaining generations. Two runs for each analysis were compared for congruence.

Estimating the Phylogenetic Informativeness of the Four Long NPCL Markers

To compare phylogenetic utility more clearly, we generated phylogenetic informativeness (PI) profiles of the four long NPCL markers using the online program PhyDesign [http://phydesign.townsend.yale.edu/) [52]. An ultrametric tree file and an alignment were required for estimating phylogenetic informativeness. For the ultrametric tree, we estimated divergence times with MultiDivTime [53]. The ML tree from the concatenated DNA alignment was used as the reference tree. Chondrichthyans were used as the outgroup, and the Actinopterygii–Sarcopterygii split was regarded as the ingroup root. Here, we used twelve calibration nodes. One was Actinopterygii–Sarcopterygii split (416–422 Ma) [54]; others were the same as those used in our previous study [10]. We used the gene-partitioned concatenated DNA alignment as the input alignment. The site rate estimation was based on the time reversible model conducted in HyPhy [55], following the recommendations of Lopez-Giraldez and Townsend [52].

Supporting Information

Figure S1  Diagram of the nucleotide alignments from the largest exons in four marker genes. The nucleotide sequences of the largest exons in RAG1, KIAA1239, SACS and TTN are retrieved from Ensembl for 16 osteichthyan species with available genome. Exon location is referenced to the human genome, and the number in parentheses indicates the length of the nucleotide alignment. Arrows represent the locations and orientations of the PCR primers used in this study. In conservation profiles, nucleotide sequences that are identical in the same column are noted by a long black bar, those that are similar in the same column are given a short black bar, and those that are totally different are given a white bar. Detailed alignments of these four sequences are available upon request. (EPS)

Figure S2 Phylogram derived from analysis of the reference NPCL marker RAG1. Phylogenetic relationships among osteichthyans were inferred from codon-partitioned maximum likelihood and Bayesian inference analyses using the reference NPCL marker RAG1 (1,488 bp). The numbers closest to the nodes are ML bootstrap proportions, followed by BI posterior probabilities. Branch lengths are based on the codon-partitioned ML analysis (3 GTR +Γ+I models for codon position partitions). Hyphens indicate nodes that are not supported in the corresponding analyses. The hyphens apply to this and all subsequent tree figures. (EPS)

Figure S3 Phylogram derived from analysis of the long NPCL marker KIAA1239. Phylogenetic relationships among osteichthyans were inferred from codon-partitioned maximum likelihood and Bayesian inference analyses using the long NPCL marker KIAA1239 (1,737 bp). The numbers close to the nodes are ML bootstrap proportions, followed by BI posterior probabilities. Branch lengths are based on the codon-partitioned ML analysis (3 GTR +Γ+I models for codon position partitions). (EPS)

Figure S4 Phylogram derived from analysis of the long NPCL marker SACS. Phylogenetic relationships among osteichthyans were inferred from codon-partitioned maximum likelihood and Bayesian inference analyses using the long NPCL marker SACS (2,211 bp). The numbers close to the nodes are ML bootstrap proportions, followed by BI posterior probabilities. Branch lengths are based on the codon-partitioned ML analysis (3 GTR +Γ+I models for codon position partitions). (EPS)

Figure S5 Phylogram derived from analysis of the long NPCL marker TTN. Phylogenetic relationships among osteichthyans were inferred from codon-partitioned maximum likelihood and Bayesian inference analyses using the long NPCL marker TTN (1,698 bp). The numbers close to the nodes are ML bootstrap proportions, followed by BI posterior probabilities. Branch lengths are based on the codon-partitioned ML analysis (3 GTR +Γ+I models for codon position partitions). (EPS)

Table S1 List of all species used in this study, along with GenBank accession numbers. (DOC)

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Author Contributions

Conceived and designed the experiments: XXS DL PZ. Performed the experiments: XXS. Analyzed the data: XXS. Contributed reagents/materials/analysis tools: XXS PZ. Wrote the paper: XXS PZ.
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