Coelacanth genomes reveal signatures for evolutionary transition from water to land

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Coelacanths are known as “living fossils,” as they show remarkable morphological resemblance to the fossil record and belong to the most primitive lineage of living Sarcopterygii (lobe-finned fishes and tetrapods). Coelacanths may be key to elucidating the tempo and mode of evolution from fish to tetrapods. Here, we report the genome sequences of five coelacanths, including four Latimeria chalumnae individuals (three specimens from Tanzania and one from Comoros) and one L. menadoensis individual from Indonesia. These sequences cover two African breeding populations and two known extant coelacanth species. The genome is ~2.74 Gbp and contains a high proportion (~60%) of repetitive elements. The genetic diversity among the individuals was extremely low, suggesting a small population size and/or a slow rate of evolution. We found a substantial number of genes that encode olfactory and pheromone receptors with features characteristic of tetrapod receptors for the detection of airborne ligands. We also found that limb enhancers of bmp7 and gli3, both of which are essential for limb formation, are conserved between coelacanth and tetrapods, but not ray-finned fishes. We expect that some tetrapod-like genes may have existed early in the evolution of primitive Sarcopterygii and were later co-opted to adapt to terrestrial environments. These coelacanth genomes will provide a cornerstone for studies to elucidate how ancestral aquatic vertebrates evolved into terrestrial animals.

Since Agassiz (1844) first described the coelacanths, their fossils have been found frequently in sediments from the Early Devonian to the Late Cretaceous periods, implying that they were successfully diversified in the past. However, the disappearance of coelacanths in the fossil record after the Late Cretaceous period led biologists to believe that coelacanths had died out during the mass extinction event around that period (65 million years ago [Ma]). Therefore, the discovery of the first living coelacanth, Latimeria chalumnae, off the coast of South Africa in 1938, created a sensation not only within the scientific community but also within the general public (Smith 1939). At present, coelacanths are called an “evolutionary relic” or a “living fossil” because their morphology is basically unchanged from that of the fossil record (Smith 1939). After the discovery of a second living coelacanth in the Comoros archipelagos (Smith 1953), the existence of a viable coelacanth population was confirmed in this area. In addition to the Comoros archipelagos, several coelacanths have been captured off the coast of Mozambique (Schilwen et al. 1993), Madagascar (Heemstra et al. 1996), Kenya (De Vos and Oyugi 2002), and Tanzania (Sasaki et al. 1999). Nakaido et al. (2011) recently demonstrated that a coelacanth population off the northern coastal region of Tanzania is genetically distinct from that of Comoros, indicating that
northern coastal region of Tanzania is the second habitat of coelacanths in the western Indian Ocean. Furthermore, two coelacanth individuals (L. menadoensis) were also captured off the coast of Manado, Sulawesi, Indonesia (Erdmann et al. 1998), on the opposite side of the Indian Ocean (the locations of the captured and observed coelacanths are summarized in Fig. 1; see also Supplemental Fig. 1).

The availability of coelacanth specimens enabled us to carry out genomic studies on these exceptional species (Supplemental Table 1). Coelacanths belong to the Sarcopterygii (lobe-finned fishes and tetrapods) (Forey 1988; Supplemental Fig. 2), in which lungfishes are also included. Indeed, molecular phylogenetic studies clearly indicate that coelacanths and lungfishes are more closely related to tetrapods than to teleost fishes (e.g., Zardoya and Meyer 1996), whereas the branching order of coelacanths, lungfishes, and tetrapods is still controversial (Takezaki et al. 2004). The karyotype of L. chalumnae, which was reported to be 48 chromosomes including microchromosomes, is similar to that of frogs and other species such as turtles or birds, further supporting the phylogenetic closeness of the coelacanth with tetrapods (Bogart et al. 1994). Accordingly, the coelacanths may fill an evolutionary gap between fish and tetrapods. Considering that the lungfishes are not suitable for comparative genomic analysis because of their extremely large genomes (Gregory et al. 2007), the coelacanth is a practical choice for genome-wide analysis.

One of the most conspicuous evolutionary events of Sarcopterygii is the transition from water to land, during which a variety of organs were subject to change due to adaptation to a novel environment. For example, the olfactory organ of the extant land vertebrates detects airborne chemicals, whereas that of fish primarily detects water-soluble chemicals. Thus, an innovative change occurred in the olfactory organ of vertebrates during the habitat transition from water to land. Similarly, the robust endoskeletal structures observed in land vertebrates are believed to be a result of adaptation to terrestrial life (Coates et al. 2002). Investigation of such phenotypic alterations is quite important to elucidate how adaptation to terrestrial life was accomplished during evolution. However, the molecular mechanisms underlying such transitions are unknown. To elucidate at the molecular level the evolutionary trajectories of vertebrates from water to land, we determined the whole-genome sequences of five coelacanths and performed an extensive comparative genomic analysis from various perspectives.

Results

Assembling the coelacanth genome

First, we constructed the reference coelacanth draft genome from one of the Tanzanian specimens (TCC041-004, gender unknown) (Nikaido et al. 2011), which was recovered from the body cavity of its mother (coelacanths give birth to fully formed offspring) (Fig. 2). A micro-computed tomography (micro-CT) scanning image was taken before sampling (Fig. 2; Supplemental Fig. 3). In total, we generated 884.8 Gbp of raw sequence data, from which 780 Gbp (300× coverage) was used for the assembly using the newly developed assembler PLATANUS (Supplemental Information 2.3, 2.4, 2.5). The genome size was estimated to be 2.74 Gbp from the k-mer analysis (Supplemental Fig. 5; Supplemental Table 3).

Unique features of the coelacanth genome

Compared with the typical teleost fish, which has a genome size of ~1 Gbp (Hinegardner and Rosen 1972) (1 pg in C-values), the coelacanth genome is large (2.74 Gbp). We found that ~60% of the coelacanth genome consists of repetitive elements—including simple repeats, low-complexity regions, and small RNAs—which is higher than the corresponding percentage in frog (35%), chicken (9%), and mammalian (40%–50%) genomes (Supplemental Fig. 17). Thus, the abundance of repetitive elements may explain the relatively large genome in coelacanths. Transposable elements (TEs) also have a considerable impact on the nucleotide composition of the coelacanth genome. The GC content of the entire coelacanth genome is 42%, which is comparable to that of other terrestrial vertebrate genomes (41% in human, 42% in chicken, and 40% in frog). However, it is the TE regions in coelacanths that have a high

Figure 1. Captured or observed coelacanth individuals. The location numbers indicate the order of the captures. The location names and the dates are summarized at the right of the map. Although most of the coelacanths were recorded in the western Indian Ocean, some coelacanths were also captured and observed off the coast of Manado, Sulawesi. The names of the key African and Indonesian countries are indicated as follows: Kenya (Ken.), Tanzania (Tan.), Mozambique (Moz.), Madagascar (Mad.), South Africa (SAf.), Indonesia (Ind.), and the Philippines (Phi.).
TEs in the coelacanth genome

Most of the repetitive sequences that account for ~60% of the coelacanth genome were characterized as TEs and classified according to their types. Within coelacanths, 23% of the genome is made up of DNA transposons, and 26% is made up of retrotransposons comprising SINEs (13.6%), LINEs (10.6%), and LTR retrotransposons (2.2%) (Table 1; Supplemental Fig. 17). The age distribution of TEs reveals that all the TE classes contain both highly divergent copies (>35% sequence divergence from the consensus) and copies that have low divergence (considered young; <5% divergence), suggesting that they have transposed or retrotransposed during both early and recent evolution (Fig. 3). In mammalian genomes, TEs that transposed or retrotransposed >100–150 Ma have been detected (International Human Genome Sequencing Consortium 2001). Because the substitution rate in coelacanths is considerably lower than the rates in other vertebrates (see below) (Amemiya et al. 2010; Higasa et al. 2012), it is likely that we could find TEs that were inserted even earlier, possibly >250 Ma.

The coelacanth LINEs comprise at least 10% of the genome. Some copies of CR1, as well as those of L2, diverge from their respective consensus sequences by as little as 1% to >35%, which suggests that both families of LINEs have been retrotranspositionally active for hundreds of millions of years. In addition to CR1 and L2, seven LINE clades (Penelope, L1/Tx1, RTE, Vingi, Dong [R4], and R2) are present in the coelacanth genome (Fig. 4). The number of LINE clades in coelacanths is higher than that in tetrapods such as mammals (at most five clades), birds (one clade), anole lizard (seven clades), and frogs (five clades).

Table 1. Composition of TEs in the coelacanth genome

| Class/clade       | Copies (×10^6) | Length (kb) | [Percentage OR fraction (%)] |
|-------------------|----------------|-------------|------------------------------|
| SINEs             | 1171.4         | 359,325     | 13.56                        |
| LINES             | 490.2          | 280,452     | 10.58                        |
| L1/Tx1            | 62.9           | 59,508      | 2.25                         |
| L2                | 122.9          | 84,581      | 3.19                         |
| CR1               | 193.7          | 102,870     | 3.88                         |
| RTE               | 67.1           | 23,156      | 0.87                         |
| Vingi             | 5.0            | 3,500       | 0.13                         |
| Dong(R4)          | 0.2            | 217         | <0.01                        |
| R2                | <0.1           | 57          | <0.01                        |
| Penelope          | 38.4           | 6,618       | 0.25                         |
| LTR retrotransposons | 56.0          | 58,466      | 2.21                         |
| Gypsy             | 7.3            | 5167        | 0.20                         |
| ERV1              | 5.9            | 1513        | 0.06                         |
| Unclassified      | 9.4            | 1753        | 0.07                         |
| DIRS              | 33.4           | 50,032      | 1.89                         |
| DNA transposons   | 1,834.9        | 608,983     | 22.98                        |
| Harbinger         | 344.9          | 246,900     | 9.32                         |
| hAT/Charlie       | 90.5           | 18,045      | 0.68                         |
| Mariner/Tc1       | 12.0           | 5264        | 0.20                         |
| Kolobok           | 15.5           | 3116        | 0.12                         |
| piggyBac          | 0.2            | 41          | <0.01                        |
| Polinton          | 0.4            | 204         | <0.01                        |
| Helitron          | 7.8            | 3103        | 0.12                         |
| Nonautonomous     | 1280.7         | 294,713     | 11.12                        |
| elements          | Unclassified   | 83.0        | 37,597                       |
| Unclassified      | 130.3          | 24,838      | 0.94                         |
| Unclassified TEs  | Total          | 3682.9      | 1,332,065                    | 50.27

Estimation of the substitution rate

The evolutionary rate of the coelacanth genome has been debated for many years (Noonan et al. 2004; Amemiya et al. 2010; Higasa et al. 2012). With the entire genome available, we first compared a set of 5247 orthologous genes for coelacanth and four other vertebrates—human, chicken, frog, and zebrafish—and then constructed a maximum-likelihood tree to evaluate the difference in evolutionary rates among vertebrates (Fig. 5A). We found that the branch length of the coelacanth lineage is significantly shorter than the other branches in all substitution models applied (Supplemental Table 15). In addition, the likelihood-ratio test confirmed that the branch length for coelacanths is significantly shorter than those of the three other vertebrates ($P < 0.01$; Supplemental Information 6.1). Our data provide genome-wide confirmation of previous studies that used a small number of selected gene sets (Noonan et al. 2004; Amemiya et al. 2010; Higasa et al. 2012).

We further determined the entire genome sequence of *L. menadoensis* and mapped the sequence reads to the reference genome to compare the whole-genome sequences between the two coelacanth species (*L. chalumnae* and *L. menadoensis*). Surprisingly, the genetic divergence of the nuclear genome between them was estimated to be only 0.18% (Fig. 5B), which is at the subpopulation level. Although the slow rate of substitution was also recently reported elsewhere (Amemiya et al. 2013), the previous investigators compared the transcriptomes from only two tissues and
partial genome sequences of *L. menadoensis* to the *L. chalumnae* genome. In the present study, we report the significantly slower rate of nucleotide substitution in the nuclear genome, which was determined by a whole-genome comparison of the two coelacanth species. In addition, we calculated the $K_s/K_a$ ratio between the two coelacanth species using a total of 4531 coding DNA sequences. The resulting $K_s/K_a$ ratio was estimated to be 0.38 (Supplemental Information 6.2), which is even higher than that for the other vertebrate species. Therefore, the low rate of amino acid substitution in coelacanths cannot be explained by purifying selection (smaller $K_a$ as compared with $K_s$). Thus, the present result suggests that both the $K_s$ and $K_a$ were small, indicating a slow rate of nucleotide substitution in the coelacanth genome.

It is worth noting that the nuclear genetic divergence of 0.0018 shown here is 23-fold smaller than that of the *L. menadoensis* and *L. chalumnae* mitochondrial genomes (0.0428) (Inoue et al. 2005; Saitoh et al. 2011). The difference between the nuclear and mitochondrial genomes could be of primary importance in discussing the evolution of the coelacanth genome. If we simply integrate the divergence time of 20–30 Ma, which was estimated by the mitochondrial genome analyses (Inoue et al. 2005; Saitoh et al. 2011), the nuclear substitution rate was calculated to be 0.03–0.045 per year in the human–chimp pair, as calculated by their genetic distance and divergence time of 0.189% (Scally et al. 2012), are also consistent with the idea of a lower nucleotide substitution rate in coelacanths.

### Genes for limb development

Because terrestrialization was an important event during the evolution of vertebrates, and because coelacanths have been historically regarded as a missing link to that event, we looked for genes that are expected to be associated with terrestrialization. First, we looked at genes related to lobed fins. The lobed fins of the coelacanth exhibit structures that are intermediate between fish and tetrapods, as represented by the presence of ray-like dermal bones (lepidotrichia), as well as a tetrapod-like robust endochondral internal skeleton, which are the ancestral characteristics of primitive sarcopterygians (Figs. 2, 6A; Coates et al. 2002; Friedman et al. 2007). The *and* genes encode actinoidin proteins, which are essential for the formation of lepidotrichia in teleost fishes and are absent in tetrapods (Zhang et al. 2010). Gene-knockdown experiments in zebrafish suggest that the loss of *and* genes in the tetrapod lineage led to the fin-to-limb transition (Zhang et al. 2010). We found two intact putative *and* genes (*and_a* and *and_b*) in the coelacanth genome. Both genes possess the conserved domain at the N-terminal region (Fig. 6B) and the repeat regions (Supplemental Fig. 28), both of which are characteristic of *and* gene family members. This discovery is consistent with the presence of actinotrichia (fiber-like proteins that are observed when the lepidotrichia first form) in coelacanths according to the anatomical description (Gerardie and Meunier 1980), and it demonstrates at the DNA level the retention of plesiomorphic fish-like characteristics (see Supplemental Information 7.2 for details).

We also explored conserved noncoding elements (CNEs) that act as enhancers of key genes for limb development such as *bmp7*, *grem1*, *shh*, and *gli3*. These CNEs participate in gene regulatory networks for axial formation, outgrowth, and chondrogenic differentiation in limb development (Zeller et al. 2009). We found apparent sequence similarity in CNEs for the limb enhancer of *bmp7* (Adams et al. 2007) and the limb enhancer (CNE11) of *gli3* (Abbsi et al. 2010) between tetrapods and coelacanths, whereas such similarities were not observed in the corresponding genomic regions of ray-finned fishes (Fig. 6C, D). The *grem1* limb enhancer is consistent with the presence of actinotrichia (fiber-like proteins that are observed when the lepidotrichia first form) in coelacanths according to the anatomical description (Gerardie and Meunier 1980), and it demonstrates at the DNA level the retention of plesiomorphic fish-like characteristics (see Supplemental Information 7.2 for details).

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### Heterozygosity rate

To investigate the genetic diversity of coelacanths, we additionally determined the entire genome sequences of three individuals (two from Tanzania and one from Comoro). We then estimated the rate of heterozygosity for each individual. The rates of heterozygosity of the coelacanth individuals from Tanzania, Comoro, and Indonesia were estimated to be 0.0023%–0.0024%, 0.0019%, and 0.0061%, respectively (Supplemental Table 20). Thus, the heterozygosity rates in the coelacanth individuals from the western Indian Ocean were significantly lower than that in the Indonesian individual. Furthermore, the heterozygosity rate was the lowest in the Comoro individual (Supplemental Tables 20, 21). The lower heterozygosity rates in coelacanth individuals, as compared with those in human (0.069%) (Wang et al. 2008) and gorilla (0.076 to 0.189%) (Scally et al. 2012), are also consistent with the idea of a lower nucleotide substitution rate in coelacanths.

**Figure 3.** Age distribution of TEs in the coelacanth genome. The total length of each TE class is shown against the sequence divergence (%) from the consensus sequence.

**Figure 4.** Distribution of LINE clades among vertebrates. The presence and absence of the representative LINE clades are shown with a plus and minus sign, respectively. (a) These mammalian LINEs are not active at present, and only fossil sequences are found. (b) RTE and Vingi are distributed only in restricted euthenian groups, possibly because of horizontal transfer events. (c) R2 elements are not reported in anole lizard but are known in turtle and bird genomes.
The present finding of the slow rate of nucleotide substitution in the coelacanth genome may hint at a potential mechanism for the modification of phenotypic traits. Although it is widely believed that the potential to alter the phenotypic traits is limited, there is evidence suggesting that coelacanths may have evolved in a similar fashion to other vertebrates. This observation provides important insights into the evolutionary history of the coelacanth lineage and its potential role in the evolution of vertebrates.
accepted that phenotypic evolution and neutral DNA evolution are decoupled (Hay et al. 2008), the slow rate of evolution in morphology and DNA appears to be coupled in coelacanths. However, because the inbreeding or the bottleneck in each of the coelacanth populations of Tanzania, Comoro, and Indonesia could possibly lead to an inaccurate estimation of the genetic distances, we should still be cautious about conclusions in this regard.

**Difference in heterozygosity rates**

We have shown that the rates of heterozygosity are significantly different according to the coelacanth locality. Because the mutation rates are expected to be similar among extant coelacanth individuals, the difference in heterozygosity rates could result from the demographic history of their populations (i.e., a reduction in population size or inbreeding). In particular, the lowest heterozygosity rate, which was found for the Comoro individual, implies the possibility of a population bottleneck. At present, the coelacanth populations in the Comoros archipelagos are threatened because of past overexploitation (Hissmann et al. 1998). Our previous mitochondrial analysis also showed a lower genetic diversity in the Comoros population than in the Tanzanian population (Nikaido et al. 2011). Thus, our present results prompt us to investigate the population structures of coelacanths more comprehensively by adding specimens currently available for molecular research.

**Apparent similarities observed in genes and enhancers of Sarcopterygian genomes**

In the present study, we found an apparent sequence similarity in CNEs for the limb enhancer between tetrapods and coelacanth. From an evolutionary viewpoint, it is likely that these CNEs that emerged in the primitive sarcopterygians are essential for shaping tetrapod-like robust internal skeletons (Fig. 6A). Importantly, the emergence of these novel CNEs and the resulting robust internal skeletons in the primitive sarcopterygians could have been used initially for effective underwater swimming rather than for locomotion on land. It is likely that these CNEs were co-opted later during the water-to-land transition in primitive tetrapods, which was coupled to the loss of and genes. Furthermore, we revealed the
amplification of tetrapod-type chemoreceptor genes (t-V1Rs, α and γ ORs) in the coelacanth genome (Fig. 7). Because primitive sarcopterygians inhabited an underwater environment, it is unlikely that t-V1Rs or α and γ ORs of those groups received airborne chemicals. Thus, we further speculate that the initial gene expansion of these genes was not directly related to terrestrial adaptation but was subsequently co-opted in tetrapods. It is possible that the ancestral coelacanth lineage(s) once inhabited shallow water and then returned to open marine water. In that case, the presence of the robust internal skeleton as well as t-V1Rs in coelacanths may be the signature of adaptation for spending time above the surface and crawling near the shore. However, this scenario is unlikely for three reasons. First, the bodies of both extant and extinct coelacanths are very heavy, and it would be very difficult for them to lift their body against gravity. Second, their bodies are covered by armor scales, which are expected to be unfit for dryness. Third, if the expansion of t-V1Rs occurred because of the adaptation to detect airborne chemicals in the ancestral coelacanth, these genes would have subsequently been pseudonized in extant coelacanths, which inhabited a deep marine environment. These genes are, however, still intact.

It is proposed that some genes utilized for adaptation already existed before the emergence of novel fauna such as multicellular organisms (Miyata and Suga 2001). We expect that some tetrapod-like genes already existed in the genomes of ancestral Sarcopterygii before the terrestrial adaptation in spite of the fact that these genes were not originally related to the terrestrial adaptation. Other examples of genes that are expected to be critical for the water-to-land transition, such as those for hemoglobins, urea synthesis, ovoviviparity, and the swim bladder, are discussed in Section 7 of the Supplemental Information.

**Highlights in the present study**

Recently, Amemiya et al. (2013) published the *L. chalumnae* genome sequence. Here, we highlight several novel findings that were not provided in their analyses.

(1) This is the first example of the application of PLATANUS, a newly developed assembler, to the determination of an entire large eukaryotic genome using next-generation sequencing.
data. A detailed explanation of the advantage of this application is provided in Supplemental Information 2.3.

(2) The complete picture of the TEs in the coelacanth genome is presented, which might shed light on the long-debated issue regarding the phylogenetic position of the coelacanths among vertebrates using the retroposon method.

(3) The whole-genome sequences of multiple coelacanth individuals from Tanzania, Comoro, and Indonesia, which were determined in the present study, enabled us to establish the extremely low value of genetic divergence among individuals and to estimate an unexpectedly slow rate of nucleotide substitution. In addition, these data enabled us to determine the significant differences in the genetic diversities among each population.

(4) The possibility of co-option in the timing of terrestrial adaptation was presented. This phenomenon could be quite important for understanding the genetic origins of key innovations in natural history, providing the seed for discussions regarding the process of macro-evolution, including the fin-to-limb transition.

(5) As an example of such discussions, the chemoreceptor genes in the coelacanth genome were completely analyzed, which provides novel insight into the evolution of olfaction during the water-to-land transition.

In summary, the coelacanth genome sequences provide a cornerstone to answer, at the DNA level, the long-debated and scientifically important questions regarding how vertebrates successfully adapted to terrestrial life. Another unexpected discovery derived from the whole-genome data is the extremely slow substitution rate found within the coelacanth genome. In addition, our finding of the significantly lower heterozygosity rates of the coelacanths in the western Indian Ocean, compared to those of the Indonesian coelacanth, should promote conservation-related genetic studies (Fricke et al. 2011) to protect this “priceless heritage from the past” (Smith 1963) from extinction.

Methods

Coelacanth specimens and tissue samples

All of the coelacanth specimens described in this study were accidently caught by local fisherman and transferred later either to the Tokyo Institute of Technology; The University of Tokyo; or Aquamarine Fukushima, Marine Science Museum, under the regulation of the Convention on International Trade in Endangered species of Wild Fauna and Flora. Detailed information about each specimen is provided in the Supplemental Material. *L. chalumnae* (TCC041-004, TCC025, and S2) were transferred from the Tanzania Fisheries Research Institute to the Tokyo Institute of Technology under the Memorandum of Understanding (M.O.U.) between the two institutions. Similarly, the Comoran specimen was obtained under the Memorandum of Understanding (M.O.U.) between the Fisheries Research Institute to the Tokyo Institute of Technology and the Center National de Documentation et de Recherche Scientifique, Musee National des Comores (CNDRS) to Aquamarine Fukushima, Marine Science Museum. Frozen muscle tissue from the Comoran *L. menadoensis* was given by Sam Ratulangi of Wild Fauna and Flora. Detailed information about each specimen is provided in the Supplemental Table 1.

Genome sequencing and assembly

Sequencing libraries were prepared using the Illumina TruSeq DNA Sample Prep kit (300 bp, 500 bp, and 1.0 kb) and the SOLiD Mate-Paired Library Construction kit (2.5 and 5.0 kb; Applied Biosystems) according to the manufacturers’ instructions. All libraries were sequenced on the Illumina HiSeq2000 sequencers. The raw sequence reads were filtered for the trimming of adapter sequences in reads and for the removal of paired reads with low-quality or extremely short insert sizes. Whole-genome assembly was performed with the newly developed assembler PLATANUS, which is optimized for short-read data from high-throughput sequencers. See Supplemental Information 2.3 for details.

Genome browser

The Coelacanth Genome Browser has been established using the assembled genome sequence and RNA-seq data, and genomic data sets used in this study are freely available online (http://coelacanth.nig.ac.jp/; Supplemental Fig 16). This browser shows the SNV density of each individual, gene models, expression, repeats, comparative analysis, fosmid clone map, and coelacanth/human alignment using the Generic Genome Browser (GBrowser) (Stein et al. 2002). In addition, the browser provides a sequence similarity search function against the coelacanth assembled genome and the gene model sequences with blast/BLAT programs, and a keyword search function against gene symbols and definitions. Users can also download the entire data set described in this study, including the genome sequence, LatCha_J1.0, as well as predicted gene sequences CDS, CDS+UTR, protein sequences, gene structures, rRNAs, and ncRNAs. See Supplemental Information 4.7 for details.

More details about the genome sequencing and assembly, RNA-seq, gene annotation, data mining, bioinformatic analyses, phylogenetic analyses, and micro-CT imaging are described in the Supplemental Methods and Supplemental Information.

Data access

All nucleotide sequence reads and the genome assembly have been deposited in the DDBJ SRA under BioProject PRJDB500 and LatCha_J1.0. (http://trace.ddbj.nig.ac.jp/dra/index.shtml). The fosmid sequences have been submitted to NCBI GenBank (http://www.ncbi.nlm.nih.gov/genbank/) under accession nos. DH994576–DH995329, GA605430-GA720357, AP012980–AP012984, AP012992–AP012996.

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