Title
A new sexannulate species of Orobdella (Hirudinida, Arhynchobdellida, Orobdellidae) from Yakushima Island, Japan

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Kyoto University
A new sexannulate species of *Orobdella* (Hirudinida, Arhynchobdellida, Orobdellidae) from Yakushima Island, Japan

Takafumi Nakano

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**Abstract**

A new sexannulate species of the genus *Orobdella* Oka, 1895, *Orobdella mononoke* sp. n., is described on the basis of five specimens collected from Yakushima Island, Japan. *Orobdella mononoke* sp. n. differs from other sexannulate *Orobdella* species in its possessing the following combination of characters: dorsal surface bicolor in life, I–XIII, XXVII and caudal sucker grayish purple, XIV–XXVI amber, male gonopore at XI c11/c12, female gonopore at XIII b2, 8 + 1/2 between gonopores, tubular but bulbous at junction with crop gastroporal duct, epididymides in XV–XIX, and atrial cornua ovate. Phylogenetic analyses using nuclear 18S rDNA and histone H3, and mitochondrial COI, tRNACys, tRNAMet, 12S rDNA, tRNAVal and 16S rDNA markers show that *O. mononoke* sp. n. is closely related to *Orobdella esulcata* Nakano, 2010 from Kyushu, Japan, and two species, *Orobdella dolichopharynx* Nakano, 2011 and *Orobdella shimadae* Nakano, 2011, from the Ryukyu Archipelago, Japan.

**Keywords**

Hirudinida, Hirudinea, Orobdellidae, *Orobdella*, new species, molecular phylogeny, Japan

**Introduction**

The genus *Orobdella* Oka, 1895 consists of nine terrestrial gastroporous leeches described from Japan (Nakano 2010, 2011a,b, 2012; Oka 1895; Richardson 1975). The genus *Orobdella* was formerly a member of the family Gastrostomobdellidae.
(Oceguera-Figueroa et al. 2011; Richardson 1971, 1975; Sawyer 1986), but a recent molecular phylogenetic study indicated that this genus belongs to the monotypic family Orobdellidae under Erpobdelliformes (Nakano et al. 2012).

The nine *Orobdella* species are split into three groups based on their mid-body somite annulation (Nakano 2012, Nakano et al. 2012): 1) the quadrannulate group containing five species; 2) the sexannulate containing three species; and 3) one octannulate species. Among these groups, the sexannulate *Orobdella* species consist of *O. ijimai* Oka, 1895 from Honshu, Japan, and two species, *O. dolichopharynx* Nakano, 2011 and *O. shimadae* Nakano, 2011, from the Ryukyu Archipelago, Japan. Recently, sexannulate *Orobdella* specimens were collected from Yakushima Island. These specimens are clearly distinguishable from the other three sexannulate species. *Orobdella* leeches from Yakushima Island are thus described as a new species herein. In addition, its phylogenetic position is estimated using nuclear 18S rDNA and histone H3, and mitochondrial COI and tRNA<sup>Cys</sup>, tRNA<sup>Met</sup>, 12S rDNA, tRNA<sup>Val</sup> and 16S rDNA (tRNA<sup>Cys</sup>–16S) sequence data.

**Material and methods**

Leeches were collected from Yakushima Island, Japan (Fig. 1), under rocks along mountain or forest trails. Altitude and coordinates for localities were obtained using a Garmin eTrex GPS unit.

Botryoidal tissue was taken from every specimen for DNA extraction, and the rest of the bodies were fixed in 10% formalin and preserved in 70% ethanol. Two measurements were made: body length (BL) from the anterior margin of the oral sucker to the

![Figure 1](image-url)
posterior margin of the caudal sucker, and maximum body width (BW). Examination, dissection, and drawings of the specimens were accomplished under a stereoscopic microscope with a drawing tube (Leica M125). Specimens used in this study have been deposited in the Zoological Collection of Kyoto University (KUZ).

The numbering convention is based on Moore (1927): body somites are denoted by Roman numerals, and annuli in each somite are given alphanumeric designations.

**PCR and DNA sequencing**

The extraction of genomic DNA followed Nakano (2012). Primer sets used in this study are listed in Table 1: for 18S, A and L, C and Y, and O and B (Apakupakul et al. 1999); for histone H3, H3aF and H3bR (Colgan et al. 1998); for COI, LCO1490 and HCO2198 (Folmer et al. 1994), and LCO-in (Nakano 2012) and HCO-out (Nakano 2012) and tRNA^Cys-, tRNA^Met-, 12S, tRNA^Val- and 16S (tRNA^Cys–16S), 12SA-out and 12SB-in, and 12SA-in and 12SB-out (Nakano 2012). All amplification reactions were performed using a GeneAmp PCR System 2700 (Applied Biosystems) or a MyCycler (Bi-Rad Laboratories) using an Ex *Tag* Polymerase Kit (Takara Bio Inc.). Only for primer set O and B of 18S, 10% DMSO was included in mixtures. Reaction mixtures were heated to 94°C for 5 min, followed by 35 cycles of 94°C (10 s), 42.5°C for 18S, COI and tRNA^Cys–16S or 53°C for histone H3 (20 s), and 72°C (42 s for 18S, 21 s for histone H3, 1 min 13 s for COI, and 1 min for tRNA^Cys–16S) and a final extension at 72°C for 6 min. The amplified DNA fragments were purified using polyethylene glycol (20% PEG 6000) precipitation.

### Table 1. PCR and cycle sequencing (CS) primers used in this study.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer name</th>
<th>Reaction</th>
<th>Primer sequence (5' → 3')</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>18S</td>
<td>1 A</td>
<td>PRC &amp; CS</td>
<td>AACCTGGTTGTATCCTGCCAGT</td>
<td>Apakupakul et al. (1999)</td>
</tr>
<tr>
<td></td>
<td>L</td>
<td>PRC &amp; CS</td>
<td>CCAACTACGAGCTTTTTTAACTG</td>
<td>Apakupakul et al. (1999)</td>
</tr>
<tr>
<td></td>
<td>2 C</td>
<td>PRC &amp; CS</td>
<td>CGGTAATTCAGCAGATCCATAG</td>
<td>Apakupakul et al. (1999)</td>
</tr>
<tr>
<td></td>
<td>Y</td>
<td>PRC &amp; CS</td>
<td>CAGACAAATCGCTCCACCAAC</td>
<td>Apakupakul et al. (1999)</td>
</tr>
<tr>
<td></td>
<td>3 O</td>
<td>PRC &amp; CS</td>
<td>AAGGGACCACCCAGAAGGGAG</td>
<td>Apakupakul et al. (1999)</td>
</tr>
<tr>
<td></td>
<td>B</td>
<td>PRC &amp; CS</td>
<td>TGATCCCTTCCGCAGGTTCACCT</td>
<td>Apakupakul et al. (1999)</td>
</tr>
<tr>
<td>Histone H3</td>
<td>H3aF</td>
<td>PRC &amp; CS</td>
<td>ATGGCTCTCGTACCAAGCAGACVGC</td>
<td>Colgan et al. (1998)</td>
</tr>
<tr>
<td></td>
<td>H3bR</td>
<td>PRC &amp; CS</td>
<td>ATATCCTTRGGCATATRGTGAC</td>
<td>Colgan et al. (1998)</td>
</tr>
<tr>
<td>COI</td>
<td>1 LCO1490</td>
<td>PRC &amp; CS</td>
<td>GGTCAACAAATCATAAAGATATG</td>
<td>Folmer et al. (1994)</td>
</tr>
<tr>
<td></td>
<td>HCO2198</td>
<td>CS</td>
<td>TAAACTTCAGGCTGCCAATAAATCA</td>
<td>Folmer et al. (1994)</td>
</tr>
<tr>
<td></td>
<td>2 LCO-in</td>
<td>CS</td>
<td>TCAGAAACGTATTCGTTATTTC</td>
<td>Nakano (2012)</td>
</tr>
<tr>
<td></td>
<td>HCO-out</td>
<td>PCR &amp; CS</td>
<td>TACACATCTGGCATGCGATTG</td>
<td>This study</td>
</tr>
<tr>
<td>tRNA^Cys–16S</td>
<td>1 12SA-out</td>
<td>PCR &amp; CS</td>
<td>TTGTATGAAACAACATTAATG</td>
<td>Nakano (2012)</td>
</tr>
<tr>
<td></td>
<td>12SB-in</td>
<td>CS</td>
<td>TAAGCTGCACTTTGACCTGA</td>
<td>Nakano (2012)</td>
</tr>
<tr>
<td></td>
<td>2 12SA-in</td>
<td>CS</td>
<td>AATTAAAACAGGTTAGATACTC</td>
<td>Nakano (2012)</td>
</tr>
<tr>
<td></td>
<td>12SB-out</td>
<td>PCR &amp; CS</td>
<td>AACCCATAATGCAAAGGTAC</td>
<td>Nakano (2012)</td>
</tr>
</tbody>
</table>
All samples were sequenced in both directions. Sequencing reactions were performed using a BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems). Each sequencing reaction mixture was incubated at 96°C for 2 min, followed by 40 cycles of 96°C (10 s), 50°C (5 s), and 60°C (42 s for 18S, 21 s for Histone H3, 45 s for COI, and 40 s for tRNA<sub>Cys</sub>–16S). The products were collected by ethanol precipitation and sequenced on an ABI 3130xl Genetic Analyzer (Applied Biosystems). The obtained sequences were edited using DNA BASER (Heracle Biosoft S.R.L.). In this study, the following DNA sequences were newly obtained and deposited in GenBank (Table 2): 1) 18S sequences from the holotype (KUZ Z224) of the new species, the holotype (KUZ Z156) of <i>O. koikei</i> Nakano, 2012 and the topotype (KUZ Z181) of <i>O. octonaria</i> Oka, 1895; 2) histone H3 sequences from ten <i>Orobdella</i> species, <i>Erpobdella japonica</i> Pawłowski, 1962 (Erpobdellidae), <i>Gastrostomobdella monticola</i> Moore, 1929 (Gastrostomobdellidae) and <i>Mimobdella japonica</i> Blanchard, 1897 (Salifidae); 3) COI and tRNA<sub>Cys</sub>–16S sequences from the holotype (KUZ Z224) and two of the paratypes (KUZ Z221, 223) of the new species. Among the new species, DNA sequences of the holotype (KUZ Z224) were analyzed in the present study. The other DNA sequences were taken from GenBank (Table 2).

### Table 2. Samples used for the phylogenetic analyses. The information on voucher, collection locality, and GenBank accession numbers is indicated. Acronym: UNIMAS, the Universiti Malaysia Sarawak. Sources: aNakano (2012), bNakano et al. (2012).

<table>
<thead>
<tr>
<th>Species</th>
<th>Voucher</th>
<th>18S</th>
<th>Histone H3</th>
<th>COI</th>
<th>tRNA&lt;sub&gt;Cys&lt;/sub&gt;–16S</th>
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</thead>
<tbody>
<tr>
<td>&lt;i&gt;Orobdella esulcata&lt;/i&gt;</td>
<td>KUZ Z29</td>
<td>AB663655&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698873</td>
<td>AB679664&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679665&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella dolichopharynx&lt;/i&gt;</td>
<td>KUZ Z120</td>
<td>AB663665&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698876</td>
<td>AB679680&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679681&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella ijimai&lt;/i&gt;</td>
<td>KUZ Z110 Topotype</td>
<td>AB663659&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698877</td>
<td>AB679672&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679673&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella kawakatsuwom&lt;/i&gt;</td>
<td>KUZ Z167 Topotype</td>
<td>AB663661&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698878</td>
<td>AB679704&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679705&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella koikei&lt;/i&gt;</td>
<td>KUZ Z156 Holotype</td>
<td>AB698883</td>
<td>AB698882</td>
<td>AB679688&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679689&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella mononoke sp. n.&lt;/i&gt;</td>
<td>KUZ Z221</td>
<td>AB698862</td>
<td>AB698863</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella mononoke sp. n.&lt;/i&gt;</td>
<td>KUZ Z223</td>
<td>AB698864</td>
<td>AB698865</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella mononoke sp. n.&lt;/i&gt;</td>
<td>KUZ Z224 Holotype</td>
<td>AB698868</td>
<td>AB698869</td>
<td>AB698866</td>
<td></td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella octonaria&lt;/i&gt;</td>
<td>KUZ Z181 Topotype</td>
<td>AB698870</td>
<td>AB698871</td>
<td>AB679708&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679709&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella shimadai&lt;/i&gt;</td>
<td>KUZ Z128 Holotype</td>
<td>AB663663&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698875</td>
<td>AB679676&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679677&lt;sup&gt;a&lt;/sup&gt;</td>
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<td>&lt;i&gt;Orobdella tashimensis&lt;/i&gt;</td>
<td>KUZ Z134 Holotype</td>
<td>AB663653&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698872</td>
<td>AB679662&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679663&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Orobdella whitmani&lt;/i&gt;</td>
<td>KUZ Z45 Topotype</td>
<td>AB663657&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698874</td>
<td>AB679668&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679669&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Erpobdella japonica&lt;/i&gt;</td>
<td>KUZ Z178</td>
<td>AB663648&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698879</td>
<td>AB679654&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679655&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Gastrostomobdella monticola&lt;/i&gt;</td>
<td>UNIMAS/A3/ BH01/10</td>
<td>AB663649&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698880</td>
<td>AB679656&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679657&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>&lt;i&gt;Mimobdella japonica&lt;/i&gt;</td>
<td>KUZ Z179</td>
<td>AB663650&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AB698881</td>
<td>AB679658&lt;sup&gt;a&lt;/sup&gt;</td>
<td>AB679659&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

### Phylogenetic analyses

Histone H3 and COI sequences were aligned by eye since there were no indels. Nuclear 18S and mitochondrial tRNA<sub>Cys</sub>–16S sequences were aligned using MAFFT X-INS-i (Hofacker et al. 2002; Katoh and Toh 2008; McCaskill 1990; Tabei et al. 2008) taking
into account RNA secondary structure information, and then refined with GBLOCKS (Castresana 2000). The length of aligned sequences of 18S was 1787 bp, that of histone H3 was 327 bp, that of COI was 1266 bp, and that of tRNA\textsuperscript{Cys}–16S was 787 bp. The concatenated sequences thus yielded a total of 4167 bp positions.

Phylogenetic trees were constructed using maximum likelihood (ML) and Bayesian inference (BI). ML phylogenies were calculated using TREEFINDER v October 2008 (Jobb et al. 2004) with the tool package PHYLOGEARS v 2.0 (Tanabe 2008), and then non-parametric bootstrapping (Felsenstein 1985) was conducted with 500 replicates. The best-fit models for each partition were selected using the Akaike Information Criterion (Akaike 1974) by using KAKUSAN4 (Tanabe 2011): for 18S, the Jobb 2008 model (J2) with gamma distribution (+G) and proportion of invariant sites (+I) was selected; for the 1st position of histone H3, the Tamura-Nei model (TN93); for the 2nd position of histone H3, the Jukes-Cantor model (JC69); for the 3rd position of histone H3, J2+G; for the 1st position of COI, TN93+G+I; for the 2nd position of COI, the transversion model (TVM)+I; for the 3rd position of COI, the transition model (TIM)+G; and the general time reversal model (GTR)+G was selected for tRNA\textsuperscript{Cys}–16S. BI and Bayesian posterior probabilities (BPPs) were estimated using the MPI version of MRBAYES v 3.1.2 (Altekar et al. 2004; Huelsenbeck et al. 2001; Ronquist and Huelsenbeck 2003). The best-fit models for each partition were identified using the Bayesian Information Criterion (Schwarz 1978) also by using KAKUSAN4: for 18S, the Kimura 1980 model (K80)+I; for histone H3 1st and 2nd position, JC69; for histone H3 3rd position, the Hasegawa-Kishino-Yano model (HKY85)+G; for COI 1st position, GTR+I; for COI 2nd position, the Felsenstein 1981 model (F81)+I; for COI 3rd position, HKY85+G; and for tRNA\textsuperscript{Cys}–16S, GTR+G. Two independent runs for four Markov chains were conducted for 7 million generations and the tree was sampled every 100 generations. Based on checking the parameter estimates and convergence using TRACER v 1.5 (Rambaut and Drummond 2009), the first 15,001 trees were discarded.

The nodes with bootstrap value (BS) higher than 70% were regarded as sufficiently resolved (Hillis and Bull 1993). Nodes with BPP higher than 95% were considered statistically significant (Leaché and Reeder 2002).

**Systematics**

**Genus Orobdella** Oka, 1895
urn:lsid:zoobank.org:act:FA8333ED-8C17-41FD-AFC1-62A4F98D4AC1

**Orobdella mononoke** sp. n.
urn:lsid:zoobank.org:act:8B4ED1DA-E1B9-49A8-8B58-014A0921695C
http://species-id.net/wiki/Orobdella_mononoke
Figs 2–5

**Diagnosis.** In life, dorsal surface of somites I–XIII, XXVII and caudal sucker grayish purple and of somites XIV–XXVI amber, ventral surface grayish white. Somite VI
quadrannulate on dorsal, \( b_1 = b_2 < a_2 = a_3 \), and triannulate on venter, \( a_1 = a_2 = a_3 \). Somite VII quadrannulate, somites VIII–XXV sexannulate, somite XXVI quinquannulate. Pharynx reaching to XIV. Gastropore conspicuous at XIII \( b_2 \) (slightly anterior to middle of annulus). Gastroporal duct, winding at junction with gastropore, tubular but slightly bulbous at junction with crop. Male gonopore at XI \( c_11/c_12 \), female gonopore at XIII \( b_2 \), behind gastropore, gonopores separated by \( 8 + 1/2 \) annuli. Paired epididymides in XV–XIX (approximately four somites). Atrial cornua developed, ovate.

**Type materials.** KUZ Z224, **holotype**, dissected, collected from under a rock along a mountain trail at Shiratani–unsuikyo, Yakushima, Kagoshima Pref. (Yakushima Island), Japan (30°22.78’N, 130°34.49’E; Alt. 648 m), by Takafumi Nakano on 29 October, 2011.

Four **paratypes** collected from under rocks along mountain trails in Yakushima, Kagoshima Pref. (Yakushima Island), Japan, by Takafumi Nakano. Two specimens from the type locality: KUZ Z221 (30°22.87’N, 130°34.68’E; Alt. 649 m), dissected, on 28 October, 2011, and KUZ Z225 (30°22.75’N, 130°34.49’E; Alt. 646 m), on 29 October, 2011. Two specimens from Kusugawa on 28 October, 2011: KUZ Z222 (30°23.76’N, 130°35.25’E; Alt. 363 m), and KUZ Z223 (30°23.75’N, 130°35.25’E; Alt. 363 m), dissected.

**Etymology.** The specific name is from the Japanese animation movie title ‘Mononoke-hime (Princess Mononoke)’. The type locality of this new species is the origin of an epic forest in that movie. The specific name is a Japanese word, not a Latin or latinized word.

**Description of holotype.** Body firm, muscular, elongated, gaining regularly in width in caudal direction, dorso-ventral depressed, sides nearly parallel from mid length to point just anterior to caudal sucker, BL 139.3 mm, BW 9.2 mm (Figs 2, 3).
A new sexannulate species of Orobdella (Hirudinida, Arhynchobdellida, Orobdellidae)...

Caudal sucker ventral, oval, its diameter smaller than BW (Figs 3B, 4D). In life, dorsal surface of somites I–XIII, XXVII and caudal sucker grayish purple, and of somites XIV–XXVI amber (Fig. 2), ventral surface grayish white. Color faded in preservative, without any dark lines (Fig. 3).

Figure 4. Orobdella mononoke sp. n., holotype, KUZ Z224. A Dorsal view of somites I–VIII B ventral view of somites I–VIII C dorsal view of somites XXV–XXVII and caudal sucker D ventral view of somites XXV–XXVII and caudal sucker E ventral view of somites XI–XIII F ventral view of gastroporal duct; and G ventral view of gastropore and female gonopore. Scale bars, 2 mm (A–F) and 0.5 mm (G). Abbreviations: an, anus; cp, crop; fp, female gonopore; gd, gastroporal duct; gp, gastropore; mp, male gonopore; np, nephridiopore; and ph, pharynx.
Figure 5. Orobdella mononoke sp. n., holotype, KUZ Z224. A Dorsal view of reproductive system including ventral nervous system B dorsal view of male atrium including position of ganglion XI C lateral view of male atrium D ventral view of male atrium; and E dorsal view of female reproductive system including position of ganglion XIII. Scale bars, 5 mm (A) and 1 mm (B–E). Abbreviations: ac, atrial cornu; at, atrium; cod, common oviduct; ed, ejaculatory duct; ep, epididymis; gp, gastropore; o, ovisac; od, oviduct; and ts, testisac.
Somite I completely merged with prostomium (Fig. 4A). Somites II and III unian- 
nulate (Fig. 4A). Somites IV and V biannulate, \((a_1+a_2) = a_3\) (Fig. 4A), \(V\) \(a_3\) forming posterior margin of oral sucker (Fig. 4B). Somite VI quadranullate on dorsal, \(b_1 = b_2 < a_2 = a_3\), triannulate on venter, \(a_1 = a_2 = a_3\) (Fig. 4A–B). Somite VII quadranullate, 
\(a_1 = a_2 = b_5 = b_6\) (Fig. 4A–B). Somites VIII–XX sexannulate. \(b_1 = b_2 = a_2 = b_5 = c_11 = c_12\) (Fig. 4A–E). Somite XXVI quinquanullate, \(b_1 = b_2 = a_2 < b_5 = b_6\), \(b_5\) and \(b_6\) with slight furrows on dorsal (Fig. 4C–D), XXVI \(b_5\) being last complete annulus on venter (Fig. 4D). Somite XXVII comprising a few furrows; anus behind it with no post-anal annulus (Fig. 4C).

Anterior ganglionic mass in VI \(a_2\) and \(a_3\). Ganglion VII in \(a_1\) and \(a_2\). Ganglia VIII–XV, XXII and XXIII in \(a_2\) of each somite (Fig. 5A). Ganglia XVI–XXI and XXIV in \(b_2\) and \(a_2\) of each somite (Fig. 5A). Ganglion XXV in \(b_2\). Ganglion XXVI in XXV \(c_{12}\) and XXVI \(b_1\). Posterior ganglionic mass in XXVI \(a_2\) and \(b_5\).

Eyes three pairs, first pair dorsally on posterior margin of II (Fig. 4A), second pair dorsolaterally on middle of \(V\) \((a_1 + a_2)\). Nephridiopores in 17 pairs, ventrally at posterior margin of \(a_1\) of each somite of VIII–XXIV (Fig. 4B, E). Papillae numerous, minute, hardly visible, one row on every annulus.

Pharynx agnathous, euthylaematous, reaching to XIV/XV (Fig. 4F). Crop tubular, acaecate, in XIV/XV to XXI \(b_2/a_2\). Gastropore conspicuous, ventral, located slightly anterior to middle of XIII \(b_2\) (Fig. 4E, G). Gastroporal duct, winding at junction with gastropore, tubular but slightly bulbous at junction with crop, joining with crop in XIV \(c_{11}\) (Fig. 4F). Intestine tubular, acaecate, in XXI \(b_2/a_2\) to XXIV \(b_2/a_2\). Rectum, tubular, thin-walled.

Male gonopore in the furrow of XI \(c_{11}/c_{12}\) (Fig. 4E). Female gonopore located slightly anterior to middle of XIII \(b_2\), inconspicuous, located behind gastropore (Fig. 4E, G). Gonopores separated by \(8 + 1/2\) annuli (Fig. 4E). Testisacs multiple, one or two testisacs on each side in each annulus, in XIX \(c_{11}\) to XXV \(b_5\) (Fig. 5A). Paired epididymides in XVI \(b_2\) to XIX \(b_5\) (Fig. 5A). Ejaculatory bulbs absent. Ejaculatory ducts in XI \(b_5\) to XVI \(b_2\), loosely coiled, each winding from each junction with epididymis, narrowing at junction with atrial cornu, then turning sharply inward toward atrial cornu without pre-atrial loop (Fig. 5A–D). Pair of atrial cornua in XI \(b_5\) and \(c_{11}\), muscular, ovate (Fig. 5A–B, D). Atrium short, muscular, globular in XI \(c_{11}\) and \(c_{12}\) (Fig. 5B–D). Penis sheath and penis absent. Ovisacs one pair, thin-walled, globular, in XIII \(a_2\) and \(b_5\) (Fig. 5A, E). Oviducts thin-walled, right oviduct crossing ventrally beneath nerve cord, both oviducts converging into common oviduct in XIII \(b_2\) (Fig. 5A, E). Common oviduct thin-walled, short, directly ascending to female gonopore (Fig. 5E).

**Variation.** In life, color generally same as holotype (Fig. 2). Somites III and IV unianullate. Pharynx reaching to XIV \(b_5/c_{11}\)–XIV \(c_{11}/c_{12}\). Crop reaching to XXI \(b_2/a_2\)–XXI \(a_2\). Gastroporal duct joining with crop in XIV \(b_5\); immature specimen (KUZ Z223), simple tubular. Intestine reaching to XXIV \(b_1\)–XXIV \(b_5\). Testisacs in XIX \(b_1\) to XXIV \(c_{11}\). Epididymides in XV \(a_2\) to XVIII \(c_{11}\). Immature specimen (KUZ Z223), pair of atrial cornua in XI \(c_{11}\), fusiform. Left oviduct crossing ventrally beneath nerve cord.
**Distribution.** Known from mountainous regions of Yakushima Island, Japan (Fig. 1).

**Phylogenetic position.** The ML tree with \( \ln L = -14306.80 \) (Fig. 6) was nearly identical to the obtained BI tree (not shown). Monophyly of the genus *Orobdella* was confirmed (BS = 99 %, BPP = 100 %). The genus *Orobdella* then divided into two clades: clade A (BS = 99 %, BPP = 100 %) consisted of two species from Hokkaido, Japan, *O. kawakatsuorum* Richardson, 1975 and *O. koikei*; and clade B (BS = 98 %, BPP = 100 %) included all the other *Orobdella* species. Clade B comprised two subclades: subclade B1 was *Orobdella tsushimensis* Nakano, 2011 from Tsushima Island, Japan; and subclade B2 (BS = 70 %, BPP = 100 %) was further divided into two subclades. Subclade B2a (BS = 92 %, BPP = 100 %) included *Orobdella mononoke* sp. n., *Orobdella esulcata* Nakano, 2010 from Kyushu, and two *Orobdella* species from the Ryukyu Archipelago, *O. dolichopharynx* and *O. shimadae*. Subclade B2b (BS = 73 %, BPP = 100 %) consisted of three species from Honshu, Japan, *O. whitmani* Oka, 1895, *O. iijimai* and *O. octonaria*.

In subclade B2a, monophyly of *Orobdella dolichopharynx* and *O. shimadae* was well supported (BS = 93 %, BPP = 100 %). However, the precise phylogenetic position of *O. mononoke* sp. n. in the sublcade could not be determined. In the ML analysis, *Orobdella mononoke* sp. n. and *O. esulcata* formed a monophyletic clade, but this clade was not supported well (BS = 30 %). In the BI analysis, *Orobdella mononoke* sp. n. and two Ryukyu Archipelago species formed a monophyletic clade, but this relationship was not also supported (BPP = 77 %).

**Remarks.** *Orobdella mononoke* sp. n. differs from the three other sexannulate congeneric species, *O. iijimai*, *O. dolichopharynx*, and *O. shimadae*, in the following characteristics (Table 3): 1) dorsal surface bicolor, I–XIII, XXVII and caudal sucker grayish purple, XIV–XXVI amber; 2) VI quadrannulate on dorsal; 3) VII quadrannulate; 4) VIII sexannulate; 5) gonopores separated by 8 + 1/2 annuli; 6) pharynx reaching to XIV; 7) gastroporal duct tubular but bulbous at junction with crop; 8) epididymides in XV–XIX (approximately four somites); and 9) atrial cornua ovate. *Orobdella mononoke* sp. n. is clearly distinguished from *O. esulcata*, *O. kawakatsuorum*, *O. koikei*, *O. tsushimensis*, *O. octonaria* and *O. whitmani*, in having mid-body somites that are sexannulate; they are quadrannulate in *O. esulcata*, *O. kawakatsuorum*, *O. koikei*, *O. tsushimensis* and *O. whitmani*, and octannulate in *O. octonaria*.

The trees obtained in this study are nearly identical to those obtained in other phylogenetic analyses of the genus *Orobdella* (Nakano 2012; Nakano et al. 2012). However, the phylogenetic position of *O. mononoke* sp. n. still remains uncertain. Further taxon samplings will be needed to obtain robust phylogeny of the genus *Orobdella*.

*Orobdella mononoke* sp. n. inhabits Yakushima Island, which is located in the northern part of the Ryukyu Archipelago (Fig. 1). In the Ryukyu Archipelago, two sexannulate *Orobdella* species have been described: 1) *O. dolichopharynx* from Amamioshima Island; and 2) *O. shimadae* from Okinawajima Island. These two species have the following characteristics in common: 1) long pharynx, reaching to somite XVI; 2) rudimentary gastroporal duct and absence of gastropore; 3) absence of epididymides;
A new sexannulate species of Orobdella (Hirudinida, Arhynchobdellida, Orobdellidae)...

and 4) absence of male atrial cornua. Although Orobdella mononoke sp. n. is a sexannulate species, this species does not share such morphological characteristics. Orobdella mononoke sp. n. possesses 1) normal length pharynx for the genus Orobdella, 2) developed gastroporal duct and conspicuous gastropore, 3) epididymides in XV–XIX, 4) ovate atrial cornua. Molecular phylogenetic analyses in this study also could not show...
monophyly of the three species in the Ryukyu Archipelago, *O. mononoke* sp. n., *O. dolichopharynx* and *O. shimadae*. These differences of morphological characteristics and molecular phylogenetic analyses suggest that *Orobdella mononoke* sp. n. is not closely related to *O. dolichopharynx* and *O. shimadae*. In vertebrates, the fauna of the Osumi Islands, in which Yaushima Island is included, is related to that of Kyushu (Toda et al. 2003). In the case of leeches, *Haemadipsa japonica* Whitman, 1886, which inhabits Honshu, Shikoku and Kyushu, Japan, is distributed in Yakushima Island (Itoh 2003). In islands of the Ryukyu Archipelago south of Yakushima Island, however, another species, *Haemadipsa ryukjuana* Oka, 1910, is distributed (Lai et al. 2011). A recent molecular phylogenetic study revealed that *H. japonica* and *H. ryukjuana* are not closely related species (Borda and Siddall 2011). These facts are collateral evidence that *O. mononoke* sp. n. is not very closely related to *O. dolichopharynx* and *O. shimadae*. Whether or not this is true, additional inventory surveys and molecular phylogenetic studies are needed to reveal the phylogenetic relationships within and the biogeographical history of the genus *Orobdella*.

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**References**


