# A new quadrannulate species of Orobdella (Hirudinida,Arhynchobdellida, Orobdellidae) from central Honshu, Japan 

Takafumi Nakano'<br>I Department of Zoology, Graduate School of Science, Kyoto University, Kyoto 606-8502, Japan<br>Corresponding author: Takafumi Nakano (nakano@zoo.zool.kyoto-u.ac.jp)

Academic editor: Fredric Govedich | Received 31 May 2014 | Accepted 05 September 2014 | Published 13 October 2014
http://zoobank.org/E3A379D5-EF2B-4378-BA88-6C662EF1 BEA7
Citation: Nakano T (2014) A new quadrannulate species of Orobdella (Hirudinida, Arhynchobdellida, Orobdellidae) from central Honshu, Japan. ZooKeys 445: 57-76. doi: 10.3897/zookeys.445.7999


#### Abstract

A new quadrannulate species of Orobdella, Orobdella masaakikuroiwai sp. n., from the mountainous region of central Honshu, Japan is described. This is only the second small species known within this genus, with a body length of less than 4 cm for mature individuals. Phylogenetic analyses using nuclear 18S rDNA and histone H 3 as well as mitochondrial COI, tRNA ${ }^{\text {Cys }}$, tRNA ${ }^{\text {Met }}, 12 \mathrm{~S}$, tRNA ${ }^{\text {Val }}, 16 \mathrm{~S}$, and ND1 markers showed that $O$. masaakikuroiwai $\mathbf{s p}$. $\mathbf{n}$. is the sister species of the quadrannulate $O$. whitmani Oka, 1895. Phylogenetic relationships within $O$. masaakikuroiwai sp. n. conducted using mitochondrial markers reveled a distinction between eastern and western phylogroups.


## Keywords

Hirudinea, Hirudinida, Orobdella, new species, gastroporous, molecular phylogeny, Japan

## Introduction

The genus Orobdella Oka, 1895 is an East Asian terrestrial macrophagous leech taxon assigned to the family Gastrostomobdellidae Richardson, 1971, along with the Southeast Asian terrestrial macrophagous genus Gastrostomobdella Moore, 1929 (Richardson 1971). Gastrostomobdellidae was once classified within the suborder Hirudiniformes, which includes jawed blood-feeding taxa (Sawyer 1986). Recent molecular phyloge-
netic studies revealed that Orobdella is part of the suborder Erpobdelliformes, which contains only predaceous leech taxa (Nakano et al. 2012, Oceguera-Figueroa et al. 2011). The monotypic family Orobdellidae Nakano, Ramlah \& Hikida, 2012 was established for Orobdella based on both morphological differences between Orobdella and Gastrostomobdella (gastroporal duct of Orobdella is tubular and positioned on top of the female organ while in Gastrostomobdella, this duct is columnar and vertical in position to the gastropore) and the results of an analysis by Nakano et al. (2012) which rejected the monophyly of these two taxa. Despite their failure to reconstruct the precise phylogenetic relationships of Orobdella and Gastrostomobdella within Erpobdelliformes, the classification by Nakano et al. (2012) is followed here.

Orobdella now consists of 11 nominal leech species from East Asia: nine species known from the Japanese Archipelago (Nakano 2010, 2011b, 2012a, b, c); one present in the Korean Peninsula and adjacent islands (Nakano 2011a, Nakano and Seo 2012, 2014); and the remainder in Taiwan (Nakano and Lai 2012). Species of Orobdella are usually large in size, with the body length of mature individuals reaching to around 10 cm (e.g. Nakano (2011b)). The largest species in this genus is the octannulate O. octonaria Oka, 1895 recorded from Honshu, Japan, with a body length often greater than 20 cm (Nakano 2012c, Oka 1895). In contrast, the smallest species is the quadrannulate $O$. koikei Nakano, 2012b found in Hokkaido, Japan with a body length of less than 4 cm , but which were considered to be mature due to the presence of developed testisacs. It is also noteworthy that the distribution of $O$. koikei in Hokkaido overlaps with that of the quadrannulate $O$. kawakatsuorum Richardson, 1975, which is present in Hokkaido as well as its adjacent islands and attains a body length of ca. 10 cm (Nakano 2012b, Nakano and Gongalsky 2014).

Several small Orobdella leeches were recently collected from east-central Honshu, Japan. Although the bodies of the specimens are up to 3.5 cm in length, some of them already possess an obvious clitellum and they are thus considered to be mature individuals. These leeches are described herein as a new species. The phylogenetic position of this new species was reconstructed using nuclear 18S and histone H3 (H3), and mitochondrial COI, $\mathrm{tRNA}^{\mathrm{Cys}}, \mathrm{tRNA}^{\text {Met }}, 12 \mathrm{~S}$, $\mathrm{tRNA}^{\mathrm{Val}}$ and 16 S rDNA , and ND1 sequence data.

## Materials and methods

## Sampling and morphological examination

Leeches were collected from seven localities in east-central Honshu, Japan (Fig. 1). These seven collection localities are numbered referring to the locality name listed in Table 1. When possible, altitudes above sea level and geographical coordinates for localities were obtained using a Garmin eTrex ${ }^{\oplus}$ GPS unit.

The specimens were relaxed by the gradual addition of absolute ethanol to fresh water. For DNA extraction, botryoidal tissue was taken from the posterior part of the body around the caudal sucker of every specimen, and then preserved in abso-


Figure I. Map showing the collection localities of the specimens examined in this study. Open circle (4) indicates the Orobdella masaakikuroiwai sp. n. type locality, and closed circles (1-3,5-7) indicate additional localities.

Table I. Collection localities in this study with the information on locality names.

| Locality number | Locality name |
| :--- | :--- |
| 1 | Akiruno, Tokyo Metropolis, Japan |
| 2 | Namesawakeikoku Valley, Izu Shizuoka Prefecture, Japan |
| 3 | Shibunoyu, Kitayama, Chino, Nagano Prefecture, Japan |
| 4 | Mr. Mitsugaisan, Ina, Nagano Prefecture, Japan |
| 5 | Shirabisotoge Pass, Ida, Nagano Prefecture, Japan |
| 6 | Ikuta, Matsukawa, Nagano Prefecture, Japan |
| 7 | Shiojidaira Nature Park, Iizuna, Nagano Prefecture, Japan |

lute ethanol. The rest of the body was fixed in $10 \%$ formalin and then preserved in $70 \%$ ethanol. Four measurements were taken: body length (BL) from the anterior margin of the oral sucker to the posterior margin of the caudal sucker, maximum body width (BW), caudal sucker length (CL) from the anterior to the posterior margin of the sucker, and caudal sucker width (CW) from the right margin to the left margin of the sucker. Examination, dissection, and drawing of the specimens were accomplished using 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 the annuli in each somite are given alphanumeric designations.

## PCR and DNA sequencing

The extraction of genomic DNA from botryoidal tissues preserved in absolute ethanol followed Nakano (2012b). Primer sets for the PCR and cycle sequencing (CS) reactions used in this study were as follows: for 18S, A and L (PCR and CS), C and Y (PCR and CS), and O and B (PCR and CS) (Apakupakul et al. 1999); for histone H3 (H3), H3aF and H3bR (PCR and CS) (Colgan et al. 1998); for COI, LCO 1490 (PCR and CS) and HCO 2198 (CS) (Folmer et al. 1994), and LCO-in (CS) and HCOout (PCR and CS) (Nakano 2012b); for tRNA ${ }^{\text {Cys }}$, $\mathrm{tRNA}^{\mathrm{Met}}$, 12 S , tRNA ${ }^{\text {Val }}$, and 16 S (tRNA ${ }^{\text {Cys }}-16 \mathrm{~S}$ ), 12 SA -out (PCR and CS) and 12SB-in (CS), and 12SA-in (CS) and 12SB-out (PCR and CS) (Nakano 2012b); for tRNA ${ }^{\text {Leu }}$ and ND1 ( $t$ RNA ${ }^{\text {Leu }}-N D 1$ ), LND3000 and HND1932 (PCR and CS) (Light and Siddall 1999). The PCR reaction and DNA sequencing were performed using the modified methods outlined by Nakano (2012a). The 18S, H3 and ND1, and COI and $\mathrm{tRNA}^{\mathrm{Cys}}-16 \mathrm{~S}$ reactions were respectively performed using a GeneAmp PCR System 2700 and a GeneAmp PCR System 9700 (Applied Biosystems). The PCR reaction mixtures were heated to $94{ }^{\circ} \mathrm{C}$ for 5 min , followed by 40 cycles at $94^{\circ} \mathrm{C}(10 \mathrm{~s}$ each $), 48^{\circ} \mathrm{C}$ for $18 \mathrm{~S}, \mathrm{H} 3$, and $\mathrm{RRNA}{ }^{\text {Leu }}-$ ND1 or $45^{\circ} \mathrm{C}$ for COI and tRNA ${ }^{\text {Cys }}-16 \mathrm{~S}$ (20 s), and $72{ }^{\circ} \mathrm{C}$ ( 48 s for 18 S , H3 and tRNA ${ }^{\text {Leu }}-N D 1$ or $1 \min 12 \mathrm{~s}$ for COI and $\mathrm{tRNA}^{\text {Cys }}-16 \mathrm{~S}$ ), and a final extension at 72 ${ }^{\circ} \mathrm{C}$ for 6 min . The sequencing mixtures were heated to $96^{\circ} \mathrm{C}$ for 2 min , followed by 40 cycles at $96^{\circ} \mathrm{C}(10 \mathrm{~s}$ each $), 50^{\circ} \mathrm{C}(5 \mathrm{~s}$ each $)$, and $60{ }^{\circ} \mathrm{C}(48 \mathrm{~s}$ each $)$. The obtained sequences were edited using DNA BASER (Heracle Biosoft S.R.L.). The DNA sequences listed in Table 2 were newly obtained in this study, and were deposited with the International Nucleotide Sequence Database Collaboration (INSDC).

## Molecular phylogenetic and genetic distance analyses

Sixty-six previously published sequences (Nakano 2012a, b, Nakano and Gongalsky 2014, Nakano and Lai 2012, Nakano et al. 2012, Nakano and Seo 2014) were obtained from the INSDC and used for the molecular phylogenetic analyses (Table 2). Four erpobdelliform species, Erpobdella japonica Pawłowski, 1962, G. monticola Moore, 1929, Mimobdella japonica Blanchard, 1897, and Odontobdella blanchardi (Oka, 1910), were used as outgroup taxa.

The phylogenetic position of the new species within the genus Orobdella was estimated based on sequences of nuclear 18 S and H 3 and mitochondrial COI, $\mathrm{tRNA}^{\mathrm{Cys}}$ 16 S , and ND1. Sequences of nuclear H3 and mitochondrial COI were aligned by eye because there were no indels. Nuclear 18 S and mitochondrial $t \mathrm{RNA} \mathrm{A}^{\mathrm{Cys}}-16 \mathrm{~S}$ and tRNA ${ }^{\text {Leu }}-$ ND1 were aligned using MATTF L-INS-I (Katoh et al. 2005). Then, the $t R N A^{\text {Leu }}$ region was removed from each sequence of $t R N A^{\text {Leu }}-N D 1$. The length of the aligned 18 S sequences was 1845 bp , that of H 3 was 327 bp , that of COI was 1266 bp , that of tRNA ${ }^{\text {Cys }}-16 \mathrm{~S}$ was 1107 bp , and that of ND1 was 633 bp . The concatenated sequences thus yielded $5,124 \mathrm{bp}$ positions.
Table 2. Samples used for the phylogenetic analyses. The information on the vouchers is accompanied by the collection locality numbers for Orobdella masaakikuroiwai sp. n. (see Fig. 1 and Table 1) and the INSDC accession numbers. Acronym: KUZ, the Zoological Collection of Kyoto University; UNIMAS, the Universiti Malaysia Sarawak.

## Species

Orobdella masaakikuroiwai sp. n.
Orobdella masaakikuroiwai sp. n.
Orobdella masaakikuroiwai sp. n.
Orobdella masaakikuroiwai sp. n.
Orobdella masaakikuroiwai sp. n.
Orobdella masaakikuroiwai sp. n.

Orobdella masaakikuroiwai sp. n. Orobdella dolichopharynx Nakano, Orobdella esulcata Nakano, 2010 Orobdella ijimai Oka, 1895 Orobdella kawakatsuorum Richardson, 1975 | Orobdella ketagalan Nakano and Lai, | KUZ Z208 Holotype | AB704785d |
| :--- | :--- | :--- | 2012 Orobdella koikei Nakano, 2012b Orobdella mononoke Nakano, 2012a Orobdella octonaria Oka, 1895 Orobdella shimadae Nakano, 2011b Orobdella tsushimensis Nakano, 2011a Orobdella whitmani Oka, 1895 Erpobdella japonica Pawłowski, 1962 Gastrostomobdella monticola Moore, 1929 Mimobdella japonica Blanchard, 1897 Odontobdella blanchardi (Oka, 1910)

Sources: a Nakano (2012a); b Nakano (2012b); c Nakano and Gongalsky (2014); d Nakano and Lai (2012); e Nakano et al. (2012); f Nakano and Seo (2014).

Phylogenetic trees were constructed using maximum likelihood (ML) and Bayesian inference (BI) models. ML phylogenies were calculated using TREEFINDER v. October 2008 (Jobb et al. 2004) with the PHYLOGEARS v. 2.0 tool package (Tanabe 2008), followed by nonparametric bootstrapping (BS) (Felsenstein 1985) conducted with 1,000 replicates. The best-fit models for each partition were selected based on the Akaike Information Criterion (Akaike 1974) using KAKUSAN4 (Tanabe 2011): for 18S, TN93 with gamma distribution $(+G)$ and proportion of invariant sites $(+\mathrm{I})$; for the first, second, and third positions of H 3 , respectively, a homogenous $(+\mathrm{H})$ TN93 model, JC69+H, and $\mathrm{J} 2+\mathrm{G}$; for the first, second, and third positions of COI, respectively, TN93+G+I, $T V M+I$, and $T I M+G$; for $t R N A{ }^{\text {Cys }}-16 S$, GTR $+G$; and for the first, second, and third positions of ND1, respectively, GTR $+\mathrm{G}+\mathrm{I}$, HYK85+G, and J2+G. BI and Bayesian posterior probabilities (BPPs) were estimated using MRBAYES v. 3.2 (Ronquist et al. 2012). The best-fit models for each partition were identified with the Bayesian information criterion (Schwarz 1978) using KAKUSAN4: for 18S, K80+G; for the first, second and third positions of H 3 , respectively, JC69+H, JC69+H, and $\mathrm{HKY}+\mathrm{G}$; for the first, second, and third positions of COI, respectively, GTR $+\mathrm{G}+\mathrm{I}, \mathrm{F} 81+\mathrm{I}$, and $\mathrm{HKY}+\mathrm{G}$; for $\mathrm{tRNA}^{\mathrm{Cys}}-16 \mathrm{~S}$, GTR+G; and for the first, second, and third positions of ND1, respectively, GTR+G, HKY85+G, and HKY85+G. Two independent runs of four Markov chains were conducted for 10 million generations, and the tree was sampled every 100 generations. The parameter estimates and convergence were checked using TRACER v. 1.5 (Rambaut and Drummond 2009), and the first 25,001 trees were discarded based on these results.

The phylogenetic relationships of the specimens of the new species were reconstructed based on sequences of mitochondrial regions. The alignment of the sequences as well as the reconstruction of the ML and BI phylogenies was accomplished followed the methods described above. The length of the aligned COI was 1266 bp , that of tRNA ${ }^{\text {Cys }}-16 \mathrm{~S}$ was 1056 bp , and that of ND1 was 579 bp . Thus, the concatenated sequences yielded $2,901 \mathrm{bp}$ positions. The best-fit models for each partition selected for the ML phylogenies were as follows: for the first, second, and third positions of COI, respectively, TN93+G, TVM +H , and TN93+G; for tRNA ${ }^{\text {Cys }}-16 \mathrm{~S}$, GTR+G; and for the first, second, and third positions of ND1, respectively, TN93+G, HKY85+H, and HYK85+G. The best-fit models identified for each partition for the BI analyses were as follows: the first, second, and third positions of COI, respectively, GTR+G, F81+H, and HKY85+G; for $t R N A^{C y s}-16 S$, GTR+G; and for the first, second, and third positions of ND1, respectively, GTR+G, F81+H, and HKY85+G. For BI and BPPs, two independent runs of four Markov chains were conducted for 6 million generations, and the tree was sampled every 100 generations. The first 15,001 trees were eliminated based on the results of the parameter estimates and convergence.

Nodes with BS values higher than 70\% were considered sufficiently resolved (Hillis and Bull 1993). Nodes with BPPs higher than $95 \%$ were considered statistically significant (Leaché and Reeder 2002).

Pairwise comparisons of the Kimura-2-parameter (K2P) distance (Kimura 1980) for the COI sequences ( 1266 bp ) obtained from the specimens of the new species were calculated using MEGA5 (Tamura et al. 2011).

## Taxonomy

Family Orobdellidae Nakano, Ramlah \& Hikida, 2012<br>http://zoobank.org/5F5BABE8-BD26-4FC7-9593-F73E62E26122

Genus Orobdella Oka, 1895
http://zoobank.org/FA8333ED-8C17-41FD-AFC1-62A4F98D4AC1

## Orobdella masaakikuroiwai sp. n.

http://zoobank.org/72F9627C-763A-49D9-9C97-E15B2FD856AA
Figs 2-5

Diagnosis. Body length of mature individual less than 4 cm . Somite IV uniannulate, somites VIII-XXV quadrannulate. Clitellum in XI b5 to XIII a2. Pharynx reaching to XIV. Gastropore conspicuous in middle of XIII a1. Gastroporal duct bulbous, winding at junction with gastropore. Male gonopore in middle of XI b6, female gonopore inconspicuous in middle of XIII a1, behind gastropore, gonopores separated by $1 / 2+4+1 / 2$ annuli. Paired epididymides in XV/XVI-XVI b5/b6 to XVII b5/b6-XVIII/XIX, occupying 7-10 annuli (i.e. one and a half to two and a half somites). Atrial cornua developed, ovate.

Type materials (see Fig. 1 for the locality number). Holotype. KUZ Z694, holotype, dissected, collected from under a rock along a forest road at Mt. Mitsugaisan, Ina, Nagano Pref., Japan ( $35^{\circ} 47.72^{\prime} \mathrm{N}, 138^{\circ} 04.70^{\prime} \mathrm{E}$; Alt. 875 m ; locality number 4), by TN on 20 July 2012.

Paratypes. Four paratypes from the type locality by TN on 20 July 2012: KUZ Z690, Z691 ( $35^{\circ} 47.72^{\prime} \mathrm{N}, 138^{\circ} 04.69^{\prime} \mathrm{E}$; Alt: 872 m ), and KUZ Z692, Z693 ( $35^{\circ} 47.74^{\prime} \mathrm{N}, 138^{\circ} 04.69^{\prime} \mathrm{E}$; Alt: 872 m ). KUZ Z693, dissected.

Additional materials (see Fig. 1 and Table 1 for the locality numbers). In total, 11 specimens examined. KUZ Z684-Z686 (three specimens), collected from under rocks in Akiruno (locality number 1), by TN: KUZ Z684, from along a mountain trail at Mt. Kariyoseyama ( $35^{\circ} 42.37^{\prime} \mathrm{N}, 139^{\circ} 12.03^{\prime} \mathrm{E}$; Alt. 341 m ) on 29 March 2010; KUZ Z685, from along Ohikagedori Road ( $35^{\circ} 43.33^{\prime} \mathrm{N}, 139^{\circ} 11.98^{\prime} \mathrm{E}$; Alt. 230 m ) on 30 March 2010; KUZ Z686, from along Bonborisen Forest Road ( $35^{\circ} 47.73$ 'N, $139^{\circ} 11.01^{\prime} \mathrm{E}$; Alt. 284 m ) on 30 March 2010. KUZ Z687, Z688 (two specimens), from under rocks along a forest road in Namesawakeikoku Valley (locality number 2), by TN on 9 July 2011: KUZ Z687 ( $34^{\circ} 50.59^{\prime} \mathrm{N}, 138^{\circ} 54.69^{\prime} \mathrm{E}$; Alt. 551 m ); KUZ Z688 ( $34^{\circ} 50.50^{\prime} \mathrm{N}, 138^{\circ} 54.59^{\prime} \mathrm{E}$; Alt. 576 m ). KUZ Z689, from under fallen leaves along a forest road at Shirabisotoge Pass $\left(35^{\circ} 26^{\prime} \mathrm{N}, 138^{\circ} 01^{\prime} \mathrm{E}\right.$; Alt. 1840 m ; locality number 5), by Yoshiko Yamane on 14 October 2011. KUZ Z695, Z696 (two specimens), from under rocks in Shiojidaira Nature Park (locality no 7), by TN on 10 August 2012: KUZ Z695 (35º40.62'N, 137º 50.48'E; Alt. 1304 m); KUZ Z696 ( $35^{\circ} 40.66^{\prime} \mathrm{N}, 137^{\circ} 50.48^{\prime} \mathrm{E}$; Alt. 1315 m ). KUZ Z697, Z698 (two specimens), from under rocks along a mountain stream in Ikuta (locality no 6), by TN on 10 August 2012: KUZ Z697 ( $35^{\circ} 33.67^{\prime} \mathrm{N}, 138^{\circ} 00.04^{\prime} \mathrm{E}$; Alt. 1098 m ); KUZ Z698 ( $35^{\circ} 33.68^{\prime} \mathrm{N}$, $138^{\circ} 00.04^{\prime} \mathrm{E}$; Alt. 1099 m ). KUZ Z699, from under fallen leaves near Shibunoyu


Figure 2. Orobdella masaakikuroiwai sp. n., holotype, KUZ Z694. A Dorsal and B ventral views. Scale bar, 5 mm .
( $36^{\circ} 02.1^{\prime} \mathrm{N}, 138^{\circ} 19.5^{\prime} \mathrm{E}$; Alt. 1860 m ; locality number 3), by Yume Imada on 6 October 2012. KUZ Z684, Z687, Z689, Z696, Z697 and Z699 (six specimens), dissected.

Etymology. The specific name is a noun in the genitive case formed directly from the name of Mr Masaaki Kuroiwa, who generously accompanied the field survey in Nagano Prefecture.

Description of holotype. Body firm and muscular, elongate, with constant width in caudal direction, dorsoventrally compressed, BL 34.0 mm , BW 3.42 mm (Fig. 2). Caudal sucker ventral, elliptic, CL 1.7 mm (minor axis), CW 1.9 mm (major axis) (Figs 2B, 3D).

Somite I completely merged with prostomium (Fig. 3A). Somites II-IV uniannulate, II not separated from I (Fig. 3A). Somite V biannulate, $(a 1+a 2)=a 3$; $a 3$ forming posterior margin of oral sucker (Fig. 3A, B). Somites VI, VII triannulate, a1 =a2 = a3 (Fig. 3A, B). Somites VIII-XXV quadrannulate, $\mathrm{al}=\mathrm{a} 2=\mathrm{b} 5=\mathrm{b} 6$ (Fig. 3A-E); b5 of X and a2 of XIII respectively being first and last annuli of clitellum (Fig. 3E). Somite XXVI triannulate, with slight furrow in a 3 , $\mathrm{a} 1>\mathrm{a} 2<\mathrm{a} 3(\mathrm{~b} 5=\mathrm{b} 6)$; a 3 being ventrally last complete annuls (Fig. 3C, D). Somite XXVII biannulate, with slight dorsal furrow in last annulus; anus behind it with no post-anal annulus (Fig. 3C).

Anterior ganglionic mass in VI a2 and a3. Ganglia VII-X, of each somite, in a2 (Fig. 4A). Ganglion XI in a2 and b5 (Fig. 4A). Ganglia XII-XVIII, of each somite, in a2 (Fig. 4A). Ganglia XIX, XX, of each somite, in a1 and a2. Ganglia XXI, XXII, of each somite, in a2. Ganglion XXIII in a1 and a2. Ganglion XXIV in a1. Ganglion XXV in XXIV b6 and XXV a1. Ganglion XXVI in b5 and b6 of XXV. Posterior ganglionic mass in al-a3 of XXVI.

Eyes in three pairs, first pair dorsally on anterior margin of III, second and third pairs dorsolaterally on posterior margin of $\mathrm{V}(\mathrm{a} 1+\mathrm{a} 2)$ (Fig. 3A). Nephridiopores in 17 pairs, one each situated ventrally at posterior margin of al of each somite in VIII-XXIV (Fig. 3B, E). Papillae numerous, minute, hardly visible, one row on every annulus.

Pharynx agnathous, euthylaematous, reaching to XIV a1/a2 (Fig. 3G). Crop tubular, reaching to XIX b5/b6 (Fig. 3G). Gastropore conspicuous, ventral in middle of XIII a1 (Fig. 3E, F). Gastroporal duct bulbous, slightly winding at junction with


Figure 3. Orobdella masaakikuroiwai sp. n., holotype, KUZ Z694. A Dorsal and B ventral views of somites I-VIII. C Dorsal and D ventral views of somites XXV-XXVII and caudal sucker. E Ventral view of somites X-XIII. F Ventral view of gastropore and female gonopore. G Ventral view of gastroporal duct. Scale bars, $1 \mathrm{~mm}(\mathbf{E}), 0.5 \mathrm{~mm}(\mathbf{A}-\mathbf{D}, \mathbf{G})$ and $0.25 \mathrm{~mm}(\mathbf{F})$. Abbreviations: af, annular furrow; an, anus; cl, clitellum; cp, crop; fg, female gonopore; gd, gastroporal duct; gp, gastropore; mg, male gonopore; np, nephridiopore; and ph, pharynx.
gastropore, joining with crop in XIV b5 (Fig. 3G). Intestine tubular, acecate, reaching to XXIV a1/a2. Rectum tubular, thin-walled, descending to anus.

Male gonopore in middle of XI b6 (Fig. 3E). Female gonopore in middle of XIII a1, inconspicuous, located posterior to gastropore (Fig. 3G). Gonopores separated by $1 / 2+4+1 / 2$ annuli (Fig. 3E). Testisacs multiple, one or two on each side in each an-


Figure 4. Orobdella masaakikuroiwai sp. n., holotype, KUZ Z694. A Dorsal view of reproductive system including ventral nervous system. B Dorsal, $\mathbf{C}$ lateral, and $\mathbf{D}$ ventral views of male atrium: $\mathbf{B}$ including position of ganglion XI. E Dorsal view of female reproductive system including position of ganglion III. Scale bars, $1 \mathrm{~mm}(\mathbf{A})$ and $0.25 \mathrm{~mm}(\mathbf{B}-\mathbf{E})$. Abbreviations: ac, atrial cornua; at, atrium; cod, common oviduct; ed, ejaculatory duct; ep, epididymis; gp, gastropore; od, oviduct; ov, ovisac; and ts, testisacs.
nulus, in XVIII a2 to XXV a1 (Fig. 4A). Paired epididymides in XVI a2 to XVIII a1, occupying 8 annuli (Fig. 4A). Ejaculatory bulbs absent. Paired ejaculatory ducts in XI a2/b5 to XVI a2, coiled in position posterior to ovisacs; each duct crossing ventrally beneath each ovisac, then loosely curved in position anterior to ovisacs; each widening from respective junction with epididymis, narrowing at junction with atrial cornua, then turning sharply inward toward atrial cornua with pre-atrial loop reaching to anterior margin of XI b5 (Fig. 4A). Pair of muscular atrial cornua ovate, in XI b5 and b6 (Fig. 4A-D). Atrium short, muscular, globular in XI b6 (Fig. 4B-D). Penis sheath and penis absent. Paired ovisacs elongated globular, one each in XIII a2-b6 (Fig. 4A, E). Oviducts thin-walled, left oviduct crossing ventrally beneath nerve cord; both oviducts converging into common oviduct in XIII a1/a2 (Fig. 4A, E). Common oviduct thinwalled, short, directly descending to female gonopore (Fig. 4E).

Variation. BL 22.4 (KUZ Z686) -35.2 (KUZ Z684) mm, BW 2.3 (KUZ Z691) -3.5 (KUZ Z684) mm, CL 1.1 (KUZ Z686)-1.7 (KUZ Z693) mm, CW 1.1 (KUZ Z686)-2.1 (KUZ Z689) mm. Somites III, IV uniannulate, each with slight dorsal furrow (KUZ Z695). Somite XXVI variable; often dorsally quadrannulate, ventrally triannulate, rarely with slight ventral furrow in a3; KUZ Z699 with quadrannulate; KUZ Z698, Z691 with triannulate with slight dorsal furrow in a3; KUZ Z689 with triannulate. Somite XXVII biannulate, or uniannulate with slight dorsal furrow. Eyes in three pairs; KUZ Z699 with one eye dorsoleft on posterior margin of III. Pharynx reaching to XIII/XIV-XIV $\mathrm{a} 2 / \mathrm{b} 5$. Crop reaching to XIX $\mathrm{b} 5 / \mathrm{b} 6-\mathrm{XX}$ a1. Gastropore occasionally slightly posterior to middle of XIII a1. Gastroporal duct joining with crop in XIV a1/a2-XIV b6; KUZ Z687 with thick, tubular duct. Intestine reaching to XXIII/XXIV-XXV a2. Male gonopore rarely slightly anterior to middle of XI b6, or slightly posterior to middle of XI b6. Female gonopore occasionally slightly posterior to middle of XIII a1. Testisacs in XVII b6-XIX al to XXIV b5-XXV a2. Epididymides in XV/XVI-XVI b5/b6 to XVII b5/b6-XVIII/XIX; occupying 7-10 annuli. Atrial cornua generally ovate; KUZ Z696 ellipsoid; KUZ Z687 fusiform. Pre-atrial loop absent, or reaching to middle of XI b5 (KUZ Z693, Z697). Ovisacs often in XIII a2-b6; KUZ Z687, Z699 in XIII a2, b5; KUZ Z696 right one in XIII a2-XIV a1/a2, left one in XIII a2-XIV a1. Right or left oviduct crossing ventrally beneath nerve cord; KUZ Z684, Z693 both oviducts converging into common oviduct in XIII a2.

Coloration. In life, dorsal surface ochre (Fig. 5), whitish brown, or brown, ventral surface grayish white or yellowish white; individuals from Shizuoka Pref. (KUZ Z687, Z688), dorsal surface whitish yellow. Colour faded in preservative, rarely with one dorsal black line from VII a3-IX a2 to XIX b5-XXVI b6 (KUZ Z691, Z693, Z694, Z698).


Figure 5. Orobdella masaakikuroiwai sp. n., paratype, KUZ Z690. A Dorsal view of live animal. B Live animal found curled up under a stone at the type locality: scale bar, 2 mm .

Distribution (see Fig. 1 for the locality numbers). This species was primarily collected from localities in Nagano Prefecture: the east-central part (locality number 3), and the southeastern part along the Inadani Basin (locality numbers 4-7). This species was also found in the western mountainous part of the Metropolitan Tokyo area (locality number 1), as well as in the Amagi Mountain Range in the central part of the Izu Peninsula, Shizuoka Prefecture (locality number 2). The locality data for this species suggested that $O$. masaakikuroiwai sp. n. would be widely distributed in mountainous regions such as the southwestern part of the Kanto Region and the southeastern part of the Chubu Region, Honshu, Japan. The lowest elevation among the localities was 230 m above sea level (a.s.l.) (locality number 1), and the highest was ca. 1860 m a.s.l. (locality number 3).

Natural history. This species was generally found curled up under rocks or fallen leaves in moist mountainous habitats (Fig. 5B). Soil was sometimes observed in the digestive tract during specimen dissection. This species is therefore considered an earthworm-feeder as are the other known Orobdella leeches.

Mature leeches with an obvious clitellum were collected on 20 July (KUZ Z690, Z691, Z693, Z694) and 10 August (KUZ Z697) at two sites in Nagano Prefecture (locality numbers 4 and 7 , elevation ca. 875 m and 1098 m , respectively). These findings indicate that the reproductive season of this species may begin in mid-to-late July.

Remarks. Although the leech specimens examined in this study were small (up to 35 mm ), several individuals, including the holotype, were determined to be mature due to the possession of an obvious clitellum and developed testisacs. Specimen KUZ Z687 possessed a tubular gastroporal duct and fusiform atrial cornua. Immature leeches may have these characteristics, because the sperm ducts and testisacs of specimen KUZ Z687 are undeveloped and barely detectable.

The new species unambiguously belongs to the genus Orobdella as it has all the generic diagnostic characteristics (see Nakano et al. (2012) for the generic diagnosis): post-anal annulus absent; pharynx agnathous, euthylaematous; gastropore in XIII; gastroporal duct lying on female organ; gonopores separated by more than one full somite; testisacs multiple; male atrium in XI without penis sheath and penis; ovisacs globular in XIII; female median reproductive system essentially lacking.

According to previous taxonomic studies (Nakano 2010, 2011a, 2012b, Nakano and Gongalsky 2014, Nakano and Lai 2012, Nakano and Seo 2012, 2014), O. masaakikuroiwai sp. n. differs from the six other quadrannulate species (i.e., O. esulcata Nakano, 2010, O. kawakatsuorum, O. ketagalan Nakano \& Lai, 2012, O. koikei, O. tsushimensis Nakano, 2011a, and O. whitmani Oka, 1895) by the following combination of characteristics (Table 3): body length less than 4 cm , IV uniannulate, gonopores separated by $1 / 2+4+1 / 2$, XXV quadrannulate, gastroporal duct bulbous, epididymides in XVI to XVIII, atrial cornua ovate. Among the six above-listed quadrannulate species, only $O$. whitmani is present in Honshu. Both $O$. masaakikuroiwai sp. n. and O. whitmani possess $1 / 2+4+1 / 2$ annuli between the gonopores, a bulbous gastroporal duct, and epididymides in XVI-XVIII. Thus, it is difficult to distinguish these two species using these diagnostic features. However, $O$. whitmani is a large species and
Table 3. Comparison of morphological characters between Orobdella masaakikuroiwai sp. n. and six quadrannulate congeneric species.

| Character | O. masaakikuroiwai sp. n. | O. esulcata <br> Nakano 2010 | O. kawakatsuorum Richardson, 1975 | O. ketagalan Nakano \& Lai, 2012 | O. koikei <br> Nakano, 2012b | O. tsushimensis <br> Nakano, 2011a | O. whitmani Oka, 1895 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Body length of mature individual | less than 4 cm | up to ca 10 cm | up to ca 10 cm | up to ca 10 cm | less than 4 cm | up to ca 10 cm | up to ca 10 cm |
| Annulation of IV | uniannulate | uniannulate | biannulate | uniannulate | uniannulate | uniannulate | uni- or <br> biannulate |
| Number of annuli between gonopores | $1 / 2+4+1 / 2$ | $2 / 3+4+1 / 3$ | 6 | $1 / 2+4+1 / 2$ | $1 / 2+4+1 / 2$ | $1 / 2+5$ | $1 / 2+4+1 / 2$ |
| Annulation of XXV | quadrannulate | quadrannulate | quadrannulate | quadrannulate | triannulate | quadrannulate | quadrannulate |
| Gastroporal duct | bulbous | tubular, but bulbous at junction with gastropore | simple tubular | simple tubular | bulbous | bulbous | bulbous |
| Epididymides | XVI to XVIII | XVI to XX | XVI to XVII | absent | XV to XX | XVII to XIX | XVI to XVIII |
| Atrial cornua | ovate | ovate | undeveloped | undeveloped | ovate | ovate | ovate |

grows up to ca. 10 cm (Nakano 2010, Oka 1895). Therefore, O. masaakikuroiwai sp. n. clearly differs from mature individuals of $O$. whitmani in body length. However, distinguishing the new species from a small juvenile of $O$. whitmani can be complex. Because immature individuals of $O$. whitmani possess a tubular gastroporal duct ( Na kano, unpublished observation) and mature individuals of $O$. masaakikuroiwai sp. n. possess a bulbous gastroporal duct, the characteristics of the duct could be used to distinguish between the two. However, insofar as immature leeches of both species have a tubular gastroporal duct, this characteristic is not useful for discriminating between immature individuals of $O$. masaakikuroiwai sp. n. and $O$. whitmani. DNA data might be useful for identification, similar to the DNA barcoding of freshwater leeches (e.g. Oceguera-Figueroa et al. (2010)). In addition to DNA data, interbreeding experiments or karyological studies may be crucial for definitive clarification between $O$. masaakikuroiwai sp. n. and O. whitmani as is the case with the species of Hirudo Linnaeus, 1758 in Europe (Petrauskiene et al. 2009, Utevsky et al. 2009).

The quadrannulate $O$. masaakikuroiwaisp. n. is unequivocally distinguishable from the four species $O$. dolichopharynx Nakano, 2011b, O. ijimai Oka, 1895, O. mononoke Nakano, 2012a and O. shimadae Nakano, 2011b, due to their sexannulate mid-body somites, as well as $O$. octonaria, which possesses octannulate mid-body somites.

## Molecular phylogenies and genetic distances

The ML tree (ln $L=-23350.60$ ) (Fig. 6) for estimating the phylogenetic position of the new species had an identical topology to the BI tree (not shown). The monophyly of the genus Orobdella was confirmed ( $\mathrm{BS}=99 \%, \mathrm{BPP}=100 \%$ ) The genus was divided into two lineages (hereafter lineages A and B ). Lineage A consisted of $O$. kawakatsuorum and O. koikei $(\mathrm{BS}=99 \%, \mathrm{BPP}=100 \%)$. Monophyletic lineage $\mathrm{B}(\mathrm{BS}=97 \%$, $\mathrm{BPP}=100 \%$ ) included the remaining 10 species (including the new species), and was divided into two sub-lineages (hereafter lineages B1 and B2). The monophyly of lineage B1, which consisted of six species, was not well supported by the ML analysis (BS $=50 \%, \mathrm{BPP}=99 \%)$. Lineage B2 included four species, but the monophyly of this lineage was also not well supported by the ML analysis (BS $=57 \%$, BPP $=99 \%$ ). The new species, $O$. masaakikuroiwai sp. n., was part of lineage B2, and was a sister taxon of $O$. whitmani within this lineage. However, this relationship was not fully supported by the ML analysis ( $\mathrm{BS}=57 \%$, BPP $=99 \%$ ).

The ML tree ( $\ln L=-8756.30$ ) (Fig. 7) for reconstructing the phylogenetic relationships of the new species had an identical topology to the BI tree (not shown). The monophyly of the specimens identified as Orobdella masaakikuroiwai sp. n. was well supported $(B S=99 \%, ~ B P P=100 \%)$. This clade was divided into two subclades (hereafter lineages 1 and 2). Monophyletic lineage 1 ( $\mathrm{BS}=99 \%$, BPP $=100 \%$ ) consisted of two specimens, KUZ Z684 (locality number 1; Tokyo Metropolis), and Z687 (locality number 2; Shizuoka Prefecture). The monophyly of lineage 2 was well supported (BS $=99 \%$, BPP $=100 \%$ ). Lineage 2 contained five specimens from Nagano Prefecture including the holotype, and consisted of two subclades (hereafter lineages 2' and 2"). The


Figure 6. The ML tree ( $\ln L=-23350.60$ ) for $5,124 \mathrm{bp}$ of nuclear 18 S rDNA and histone H 3 , and mitochondrial COI, tRNACys, tRNAMet, 12 S rDNA, tRNAVal, 16 S rDNA, and ND1 markers. A species name of Orobdella in red indicates a quadrannulate species; in green, sexannulate; and in blue, octannulate. The numbers associated with the nodes represent the bootstrap values for ML (BS)/and Bayesian posterior probabilities (BPPs).


Figure 7. The ML tree ( $\ln L=-8756.30$ ) for 2,901 bp of mitochondrial COI, tRNACys, tRNAMet, 12 S rDNA, tRNAVal, 16 S rDNA, and ND1 markers. Voucher numbers of the specimens of Orobdella masaakikuroiwai sp. n. are accompanied by the collection locality numbers (see Fig. 1). The numbers associated with the nodes represent the bootstrap values for ML (BS)/and Bayesian posterior probabilities (BPPs).
monophyly of lineage $2^{\prime}$ was well supported ( $\mathrm{BS}=99 \%$, $\mathrm{BPP}=100 \%$ ). This lineage included two specimens, KUZ Z689 (locality number 5) and Z697 (locality number 6 ), observed in the southern part of the prefecture. The monophyly of lineage 2 " was not well supported in the ML analysis ( $\mathrm{BS}=49 \%$, $\mathrm{BPP}=94 \%$ ). Lineage 2 " contained three specimens, KUZ Z694 (holotype; locality number 4), Z696 (locality number 7), and Z699 (locality number 3) collected from the east-central and mid-southern parts of Nagano Prefecture. KUZ Z694 and Z696 formed a monophyletic lineage (BS = $99 \%, \mathrm{BPP}=100 \%)$ within lineage 2 ".

The COI K2P distance within O. masaakikuroiwai sp. n. was $0.5-6.7 \%$ (mean = 4.4\%) (Table 4). The genetic divergence between lineages 1 and 2 was 5.8-6.7\% (mean $=6.3 \%)$, and that between lineages $2^{\prime}$ and $2 "$ was $2.7-3.5 \%$ (mean $\left.=3.2 \%\right)$. The COI K2P distance between O. masaakikuroiwai sp. n. and O. whitmani (KUZ Z45, topotype) was $10.4-11.7 \%($ mean $=11.0 \%)$.

Table 4. Kimura-2-parameter distances for the 1266 bp for the COI sequences of Orobdella masaakikuroiwai sp. n. specimens, with associated collection locality numbers (see Fig. 1 and Table 1).

| Specimen <br> (locality number) | $\mathbf{1}$ | $\mathbf{2}$ | $\mathbf{3}$ | $\mathbf{4}$ | $\mathbf{5}$ | $\mathbf{6}$ | $\mathbf{7}$ |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| 1: KUZ Z684 (1) |  |  |  |  |  |  |  |
| 2: KUZ Z687 (2) | 0.046 |  |  |  |  |  |  |
| 3: KUZ Z689 (5) | 0.065 | 0.061 |  |  |  |  |  |
| 4: KUZ Z694 (4) | 0.065 | 0.058 | 0.033 |  |  |  |  |
| 5: KUZ Z696 (7) | 0.063 | 0.059 | 0.035 | 0.005 |  |  |  |
| 6: KUZ Z697 (6) | 0.066 | 0.067 | 0.006 | 0.034 | 0.034 |  |  |
| 7: KUZ Z699 (3) | 0.063 | 0.059 | 0.027 | 0.023 | 0.023 | 0.027 |  |

## Discussion

The current molecular phylogenies showed that the specimens morphologically identified as the new species form a monophyletic group with strong support values. In addition, the K2P genetic distance of the COI sequences detected within the specimens was $0.5-6.7 \%($ mean $=4.4 \%)$. Nakano $(2012 \mathrm{~b})$ stated that the COI K2P distance between the sister species of Orobdella, O. kawakatsuorum and O. koikei, was 8.1-9.9\% (mean $=9.0 \%$ ). Therefore, the present genetic analyses support the taxonomic designation of the specimens examined in this study as belonging to the new species, $O$. masaakikuroiwai sp. n.

Orobdella masaakikuroiwai sp. n. was divided into two lineages (lineages 1 and 2) according to the molecular phylogenetic analyses. Lineage 1 consists of the individuals inhabiting the Kanto Region (KUZ Z684, locality number 1) and the Izu Peninsula (KUZ Z687, locality number 2). The Izu Peninsula is located on the Philippine Sea Plate and collided with Honshu island around 1 million years ago (Kitazato 1997).

Therefore, $O$. masaakikuroiwai sp. n. likely migrated into the peninsula after this collision event. In addition to lineages 1 and 2 composed of specimens from the mountainous region of Nagano Prefecture, the individuals of $O$. masaakikuroiwai sp. n. were sub-divided into central (lineage 2 "; locality numbers $3,4,7$ ) and southern (lineage 2'; locality numbers 5, 6) phylogroups. The Ina Basin is located in the southern part of Nagano Prefecture along the Tenryu River. Mountain districts are present to the east (including locality numbers 4-6) and west (containing locality number 7) along this basin. The specimen from Shiojidaira is the closest to the holotype from Mt. Mitsugaisan even though the Ina Basin separates the mountainous regions. In addition, the COI divergence between the two specimens from Shiojidaira (KUZ Z696) and Mt. Mitsugaisan (KUZ Z694) was low (0.5\%). This may indicate that O. masaakikuroiwai $\mathrm{sp} . \mathrm{n}$. leeches in this area have recently dispersed. The same low genetic distance ( $0.6 \%$ ) was detected between the specimens collected from the southern part of Nagano Prefecture (KUZ Z689, locality number 5, and KUZ Z697, locality number 6).

Orobdella masaakikuroiwai sp. n. is the second known species in which the body length of a mature individual is less than 4 cm . Orobdella masaakikuroiwai sp. n. is syntopic with $O$. octonaria in the Izu Peninsula (locality number 2), and the distribution of this new species partly overlaps with that of the latter species (Nakano, unpublished data). In addition, both $O$. koikei and $O$. kawakatsuorum are present in Hokkaido (Nakano 2012b). Therefore, a difference in the body size of mature individuals may allow different species of Orobdella to coexist in the same region. The phylogeny indicates that the small size of mature leeches likely evolved in parallel within Orobdella. Orobdella whitmani is the sister species of $O$. masaakikuroiwai sp. n. and grows to ca. 10 cm . In addition, $O$. ijimai and $O$. octonaria are close congeners of $O$. masaakikuroiwai sp. n. and $O$. whitmani, and they grow to ca. 10 cm and ca. 20 cm , respectively. Therefore, the intermediate size of mature individuals may be a plesiomorphic characteristic of the clade consisting of these four species. However, several undescribed species of Orobdella are known including small-sized species (Nakano, unpublished observation). Further faunal and systematic studies will help to elucidate the evolutionary and biogeographical history of the predaceous genus Orobdella.

## Acknowledgments

The author is grateful to Masaaki Kuroiwa for his generous assistance with my field survey in Nagano Prefecture. I am also grateful to Yume Imada (Kyoto University; KU ) and Yoshiko Yamane (KU) for providing specimens of the new species, and to two anonymous reviewers and Dr Fredric R. Govedich (Southern Utah University) for their constructive comments on this manuscript. A part of this study was financially supported by Grants for Biodiversity and Evolutionary Research of Global COE (A06) and for Excellent Graduate Schools, both from MEXT, Japan, to Kyoto University, and JSPS Grant-in-Aid for JSPS Fellows and Young Scientists (B) (\#26840127) to the author.

## References

Akaike H (1974) A new look at the statistical model identification. IEEE Transactions on Automatic Control 19: 716-723.
Apakupakul K, Siddall ME, Burreson EM (1999) Higher level relationships of leeches (Annelida: Clitellata: Euhirudinea) based on morphology and gene sequences. Molecular Phylogenetics and Evolution 12: 350-359. doi: 10.1006/mpev.1999.0639
Blanchard R (1897) Hirudinées du Musée de Leyde. Notes from the Leyden Museum 19: 73-113.
Colgan DJ, McLauchlan A, Wilson GDF, Livingston SP, Edgecombe GD, Macaranas J, Cassis G, Gray MR (1998) Histone H3 and U2 snRNA DNA sequences and arthropod molecular evolution. Australian Journal of Zoology 46: 419-437. doi: 10.1071/ZO98048
Felsenstein J (1985) Confidence limits on phylogenies: an approach using the bootstrap. Evolution 39: 783-791. doi: 10.2307/2408678
Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R (1994) DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. Molecular Marine Biology and Biotechnology 3: 294-299.
Hillis DM, Bull JJ (1993) An empirical test of bootstrapping as a method for assessing confidence in phylogenetic analysis. Systematic Biology 42: 182-192. doi: 10.1093/sysbio/42.2.182
Jobb G, von Haeseler A, Strimmer K (2004) TREEFINDER: a powerful graphical analysis environment for molecular phylogenetics. BMC Evolutionary Biology 4: 18. doi: 10.1186/1471-2148-4-18

Katoh K, Kuma K-i, Toh H, Miyata T (2005) MAFFT version 5: improvement in accuracy of multiple sequence alignment. Nucleic Acids Research 33: 511-518. doi: 10.1093/nar/ gki198
Kimura M (1980) A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide sequences. Journal of Molecular Evolution 16: 111-120. doi: 10.1007/bf01731581
Kitazato H (1997) Paleogeographic changes in central Honshu, Japan, during the late Cenozoic in relation to the collision of the Izu-Ogasawara Arc with the Honshu Arc. Island Arc 6: 144-157. doi: 10.1111/j.1440-1738.1997.tb00166.x
Leaché AD, Reeder TW (2002) Molecular systematics of the eastern fence lizard (Sceloporus undulatus): a comparison of parsimony, likelihood, and Bayesian approaches. Systematic Biology 51: 44-68. doi: 10.1080/106351502753475871
Light JE, Siddall ME (1999) Phylogeny of the leech family Glossiphoniidae based on mitochondrial gene sequences and morphological data. The Journal of Parasitology 85: 815-823.
Linnaeus C (1758) Systema Naturae per Regna Tria Naturae, Secundum Classes, Ordines, Genera, Species, cum Characteribus, Diferentiis, Stinontmis, Locis. Salvius, Stockholm, 824 pp.
Moore JP (1927) The segmentation (metamerism and annulation) of the Hirudinea. In: Harding WA, Moore JP (Eds) The Fauna of British India, including Ceylon and Burma Hirudinea. Taylor \& Francis, London, 1-12.

Moore JP (1929) Leeches from Borneo with descriptions of new species. Proceedings of the Academy of Natural Sciences of Philadelphia 81: 267-295.
Nakano T (2010) A new species of the genus Orobdella (Hirudinida: Arhynchobdellida: Gastrostomobdellidae) from Kumamoto, Japan, and a redescription of $O$. whitmani with the designation of the lectotype. Zoological Science 27: 880-887. doi: 10.2108/zsj.27.880
Nakano T (2011a) A new species of Orobdella (Hirudinida: Arhynchobdellida: Gastrostomobdellidae) from Tsushima Island, Japan. Species Diversity 16: 39-47.
Nakano T (2011b) Redescription of Orobdella ijimai (Hirudinida: Arhynchobdellida: Gastrostomobdellidae), and two new species of Orobdella from the Ryukyu Archipelago, Japan. Zootaxa 2998: 1-15.
Nakano T (2012a) A new sexannulate species of Orobdella (Hirudinida, Arhynchobdellida, Orobdellidae) from Yakushima Island, Japan. ZooKeys 181: 79-93. doi: 10.3897/zookeys.181.2932
Nakano T (2012b) A new species of Orobdella (Hirudinida, Arhynchobdellida, Gastrostomobdellidae) and redescription of $O$. kawakatsuorum from Hokkaido, Japan with the phylogenetic position of the new species. ZooKeys 169: 9-30. doi: 10.3897/zookeys.169.2425
Nakano T (2012c) Redescription of Orobdella octonaria (Hirudinida: Arhynchobdellida: Orobdellidae) with designation of a lectotype. Species Diversity 17: 227-233. doi: 10.12782/ sd.17.2.227
Nakano T, Gongalsky KB (2014) First record of Orobdella kawakatsuorum (Hirudinida: Arhynchobdellida: Erpobdelliformes) from Kunashir Island, Kuril Islands. Biodiversity Data Journal 2: e1058. doi: 10.3897/BDJ.2.e1058
Nakano T, Lai Y-T (2012) A new species of Orobdella (Hirudinida, Arhynchobdellida, Orobdellidae) from Taipei, Taiwan. ZooKeys 207: 49-63. doi: 10.3897/zookeys.207.3334
Nakano T, Ramlah Z, Hikida T (2012) Phylogenetic position of gastrostomobdellid leeches (Hirudinida, Arhynchobdellida, Erpobdelliformes) and a new family for the genus Orobdella. Zoologica Scripta 41: 177-185. doi: 10.1111/j.1463-6409.2011.00506.x
Nakano T, Seo H-Y (2012) First record of Orobdella tsushimensis (Hirudinida: Arhynchobdellida: Orobdellidae) from Korea (Gageodo Island) and its molecular phylogenetic position within the genus. Species Diversity 17: 235-240. doi: 10.12782/sd.17.2.235
Nakano T, Seo H-Y (2014) First record of Orobdella tsushimensis (Hirudinida: Arhynchobdellida: Gastrostomobdellidae) from the Korean Peninsula and molecular phylogenetic relationships of the specimens. Animal Systematics, Evolution and Diversity 30: 87-94. doi: 10.5635/ASED.2014.30.2.087

Oceguera-Figueroa A, León-Régagnon V, Siddall ME (2010) DNA barcoding reveals Mexican diversity within the freshwater leech genus Helobdella (Annelida: Glossiphoniidae). Mitochondrial DNA 21: 24-29. doi: 10.3109/19401736.2010.527965
Oceguera-Figueroa A, Phillips AJ, Pacheco-Chaves B, Reeves WK, Siddall ME (2011) Phylogeny of macrophagous leeches (Hirudinea, Clitellata) based on molecular data and evaluation of the barcoding locus. Zoologica Scripta 40: 194-203. doi: 10.1111/j.14636409.2010.00465.x

Oka A (1895) On some new Japanese land leeches. (Orobdella nov. gen.). The Journal of the College of Science, Imperial University, Japan 8: 275-306.

Oka A (1910) Key to Japanese leeches. Dobutsugaku Zasshi 22: 56-64.
Pawłowski LK (1962) O występowaniu pijawki Erpobdella octoculata (L.) w Japonii. Zeszyty Naukowe Uniwersytetu Łódzkiego Nauki Matematiczno-przyrodnicze Seria II 12: 127-136.
Petrauskiene L, Utevska O, Utevsky SY (2009) Can different species of medicinal leeches (Hirudo spp.) interbreed? Invertebrate Biology 128: 324-331. doi: 10.1111/j.17447410.2009.00180.x

Rambaut A, Drummond AJ (2009) Tracer v. 1.5. http://tree.bio.ed.ac.uk/software/tracer/
Richardson LR (1971) Gastrostomobdellidae f. nov. and a new genus for the gastroporous Orobdella octonaria Oka, 1895, of Japan (Hirudinoidea: Arhynchobdellae). Bulletin of the National Science Museum (Tokyo) 14: 585-602.
Richardson LR (1975) A new species of terricolous leeches in Japan (Gastrostomobdellidae, Orobdella). Bulletin of the National Science Museum Series A (Zoology) 1: 39-56.
Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, Larget B, Liu L, Suchard MA, Huelsenbeck JP (2012) MrBayes 3.2: Efficient Bayesian phylogenetic inference and model choice across a large model space. Systematic Biology 61: 539-542. doi: 10.1093/sysbio/sys029

Sawyer RT (1986) Leech Biology and Behaviour. Clarendon Press, Oxford, 1065 pp.
Schwarz G (1978) Estimating the dimension of a model. The Annals of Statistics 6: 461-464.
Tamura K, Peterson D, Peterson N, Stecher G, Nei M, Kumar S (2011) MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Molecular Biology and Evolution 28: 2731-2739. doi: 10.1093/molbev/msr121

Tanabe AS (2008) Phylogears v. 2.0. http://www.fifthdimension.jp
Tanabe AS (2011) Kakusan4 and Aminosan: two programs for comparing nonpartitioned, proportional and separate models for combined molecular phylogenetic analyses of multilocus sequence data. Molecular Ecology Resources 11: 914-921. doi: 10.1111/j.17550998.2011.03021.x

Utevsky SY, Kovalenko N, Doroshenko K, Petrauskiene L, Klymenko V (2009) Chromosome numbers for three species of medicinal leeches (Hirudo spp.). Systematic Parasitology 74: 95-102. doi: $10.1007 /$ s11230-009-9198-2

