New species of the genus *Inversidens* Haas, 1911 (Unionoida, Unionidae, Gonideinae) from Jiangxi Province, China

Ruiwen Wu¹, Xiongjun Liu², Takaki Kondo³, Shan Ouyang⁴, Xiaoping Wu⁴

1 School of Life Science, Shanxi Normal University, Linfen 041000, China 2 School of Life Science, Jiaying University, Meizhou 514015, China 3 Division of Natural Science, Osaka Kyoiku University, Osaka 582-8582, Japan 4 School of Life Sciences, Nanchang University, Nanchang 330031, China

Corresponding authors: Xiongjun Liu (609449126@qq.com); Xiaoping Wu (xpwu@ncu.edu.cn)

Abstract
We diagnose and describe a new freshwater mussel species of the genus *Inversidens*, *I. rentianensis* sp. nov. from Jiangxi Province, China based on morphological characters and molecular data. This paper includes a morphological description and photograph of the holotype, and partial sequences of mitochondrial COI as DNA barcode data.

Keywords
COI, freshwater mussel, genetic distances, morphology, taxonomy

Introduction

The genus *Inversidens* Haas, 1911 belongs to the subfamily Gonideinae in the family Unionidae. The genus was first depicted by Haas (1911) as a subgenus of *Nodularia* with two species, i.e., *Unio brandtii* Kobelt, 1879 and *Nodularia parcedentata* Haas, 1911, both restricted to Japan. Later, Haas (1969) further classified *Unio reinianus* Kobelt, 1879, *Unio haconensis* Ihering, 1893, *Unio japonensis* Lea, 1859, *Unio pantoensis* Neumayr, 1899 within *Inversidens*. All species were restricted to Japan, except for...
*U. pantoensis*, which was distributed in China. By comparing the conchological characters, Kondo (1982) believed that *U. brandtii* was not morphologically distinct from *N. parcedentata*, and regarded *N. parcedentata* as a variety of *U. brandtii*. Based on the morphology of the glochidium, Habe (1991) removed *U. reinianus* from *Inversidens*, and established a new genus *Inversiunio*. Based on morphological characteristics of the shell, Kondo (1998) moved *Unio haconensis*, which was regarded as a synonym of *Unio jokohamensis* (Ihering, 1893), into *Inversiunio*. Furthermore, Starobogatov (1970) used *Unio japanensis* as the type species for his newly established genus *Pronodularia*.

Currently, only two species are recognized within *Inversidens*, the Japanese endemic *I. brandtii* and *I. pantoensis* in China (Fig. 1A, B; Kondo 2008; He and Zhuang 2013; Lopes-Lima et al. 2020; Graf and Cummings 2021a, b; MolluscaBase eds. 2021).

In this study, we diagnose and describe a new *Inversidens* species from Jiangxi Province, China. In addition, we provide estimations of the intraspecific and interspecific genetic distances within *Inversidens* based on the mitochondrial COI barcode to examine species validity.

**Materials and methods**

**Specimen collection and identification**

In March 2018, four samples were collected from the Mianshui River, Rentian Town, Ganzhou City, Jiangxi Province, China (25.989557°N, 116.131333°E). All type and voucher specimens are deposited at the Biological Museum of Nanchang University, China (NCFM180325-NCFM180328).

**DNA extraction and COI amplification**

Of the four individuals, only two samples had tissues. Total genomic DNA was extracted from dissected somatic tissue using TIANamp Marine Animals DNA Kit (Tiangen Biotech, Beijing, China) according to the manufacturer’s instructions.

Mitochondrial cytochrome oxidase subunit I (COI) gene sequences have been widely used for species delimitation of freshwater mussels based on genetic distance and the criteria of monophyly (Elderkin et al. 2016; Lopes-Lima et al. 2019; Smith et al. 2019). Polymerase chain reaction (PCR) amplification of the COI gene with a 680-base pair fragment was performed using a primer pair consisting of LCO1490 and HCO2198 (Folmer et al. 1994). Thermal cycling conditions were 98 °C for 10 s, followed by 35 cycles of 94 °C for 1 min, 50 °C for 1 min, 72 °C for 1–2 min, and a final extension of 72 °C for 7 min, following the TaKaRa Ex manufacturer’s protocol. The amplified PCR products were purified and sequenced by Sangon Biotech (Shanghai). The PCR product size for the COI amplicon was 680 bp. The sequences obtained in this study have been uploaded to GenBank.
DNA barcode dataset construction

We constructed a mitochondrial COI dataset with the newly obtained sequences from this study and the available *Inversidens brandtii* sequences from GenBank.

Previously published sequences were downloaded from GenBank and added to the dataset, i.e., 17 species of the subfamily Gonideinae and four species of the subfamily Unioninae for the ingroup, and one species of the subfamily Parreysiinae as the outgroup.

As a result, a total of 29 COI sequences were used for this study. Sequence details and GenBank accession numbers are shown in Table 1.

All COI nucleotide sequences were translated to amino acid sequences using MEGA 5.0 (Tamura et al. 2011) and aligned based on the amino acid sequences using the program MUSCLE (Edgar 2004) with default settings. We calculated and compared inter-and intraspecific distances with MEGA 5.0 using the uncorrected $p$-distance. Standard error was assessed using 1000 bootstrap replicates.

| Taxa | GenBank accession number |
|------|--------------------------|
| UNIONIDAE Rafinesque, 1820 | |
| Parreysiinae Henderson, 1935 | |
| *Indonaia andersoniana* (Nevill, 1877) | KX865835 |
| Unioninae Rafinesque, 1820 | |
| *Acuticosta chinensis* (Lea, 1868) | MG462919 |
| *Inversiunio jokohamensis* (Ihering, 1893) | LC518985 |
| *Inversiunio reinianus* (Kobelt, 1879) | LC518976 |
| *Nodularia douglasiae* (Griffith & Pidgeon, 1833) | |
| Gonideinae Ortmann, 1916 | |
| *Pseudodon bogani* Bolotov, Kondakov & Konopleva in Bolotov et al. 2017 | MF352216 |
| *Pseudodon manueli* Konopleva, Kondakov & Vikrev in Bolotov et al. 2017 | MF352228 |
| *Monodontina cambodjensis* (Petit de la Saussaye, 1865) | |
| *Pilsbryoconcha exilis* (Lea, 1838) | KP795024 |
| *Chamberlainia hainesiana* (Lea, 1856) | KX822635 |
| *Sinobyropsis cuninghamii* (Lea, 1852) | NC_011763 |
| *Sinobyropsis schlegelii* (Martens, 1861) | NC_015110 |
| *Lamprotula caveata* (Heude, 1877) | KX822646 |
| *Lamprotula leati* (Griffith & Pidgeon, 1833) | NC_023346 |
| *Potomida littoralis* (Cuvier, 1798) | JN243905 |
| *Pronodularia japonensis* (Lea, 1859) | KX822659 |
| *Gonidea angulata* (Lea, 1838) | DQ272371 |
| *Leguminatia wheatleyi* (Lea, 1862) | KX822651 |
| *Microcondylata bonelli* (Féussac, 1827) | KX822652 |
| *Sinovolenatia carinata* (Heude, 1877) | KX822669 |
| *Psychrohynchus pflisteri* (Heude, 1874) | KY067440 |
| *Parasolenatia virilarii* (Heude, 1877) | KX966393 |
| *Inversidens brandtii* (Kobelt, 1879) | AB040827 |
| *Inversidens brandtii* (Kobelt, 1879) | MT020598 |
| *Inversidens brandtii* (Kobelt, 1879) | MT020597 |
| *Inversidens brandtii* (Kobelt, 1879) | LC519005 |
| *Inversidens rentianensis* sp. nov. 1* | LC519004 |
| *Inversidens rentianensis* sp. nov. 2* | MZ073336 |
| *Inversidens rentianensis* sp. nov. 1* | MT020598 |
| *Inversidens rentianensis* sp. nov. 2* | MT020597 |
Phylogenetic analysis

Bayesian inference (BI) analyses were inferred in MrBayes Version 2.01 (Ronquist et al. 2012), using GTR+GAMMA model of nucleotide substitution. Four chains were run simultaneously for 10 million generations and trees were sampled every 1000 generations. The first 25% of these trees were discarded as burn-in when computing the consensus tree (50% Majority Rule). Sufficient mixing of the chains was considered to have been reached when the average standard deviation of split frequencies was below 0.01. Additionally, IQ-TREE was run for Maximum Likelihood (ML) tree reconstruction, using partition models with 1000 ultrafast bootstraps (Minh et al. 2013).

Taxonomy

Inversidens rentianensis Wu & Wu, sp. nov.
http://zoobank.org/62424717-9514-4C7D-9C0E-240F1D95F03E

Type specimens. Holotype. China • Jiangxi Province, Ganzhou City, Rentian Town (壬田镇), Mianshui River (25.989557°N, 116.131333°E), 13 March 2018, coll. Xiongjun Liu (NCFM180325). Paratypes. Same data as holotype (NCFM180326-NCFM180328).

Diagnosis. Inversidens rentianensis sp. nov. is morphologically distinct from the other two recognized species within the genus by shell shape, beak position and nacre colour (Table 2). Diagnostic characteristics: shell irregularly subtriangular; curvature of the ventral margin slight, nearly straight; umbo situated 1/2 of shell length; nacre reddish.

Description. Shell irregularly subtriangular, medium thickness, and quite inflated. Anterior margin regularly rounded; ventral margin nearly straight; posterior margin obliquely arc-shaped. Umbo prominent and slightly eroded. Umbo sculptured with feebly wavy wrinkles. Posterior slope formed by the ventral margin and posterior margin low, triangular. Epidermis shining black or with brownish-yellow hue. Only one cardinal tooth in each valve, shape triangular. Laterals thick, a little curved, 2 in each valve. Nacre reddish-bronze in colour.

Length 43–52 mm, height 29–36 mm.

Etymology. The specific epithet is derived from the type locality, Rentian Town.

Distribution. The species is known only from Mianshui River, Rentian Town, Ganzhou City, Jiangxi Province, China (present study) (Fig. 2).

GenBank accession number. Holotype, NCFM180325: MZ073336; paratypes, NCFM180326: MZ073337.

Molecular analyses. Pairwise COI sequence divergences from Inversidens brandtii and Inversidens rentianensis sp. nov. were conducted using MEGA 5.0. Based on the uncorrected p-distance model, the intraspecific divergences of I. brandtii and I. rentianensis sp. nov. were both 0.00%. The interspecific divergence of I.
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**Figure 1.** Photographs of *Inversidens* taxa **A** *I. brandtii* **B** *I. pantoensis* **C** *I. rentianensis* sp. nov. Photos: [A, B] from the MUSSEL Project, [C] from this study, NCFM180325 (holotype), scale is 2 cm.

*brandtii* and *I. rentianensis* sp. nov. was 10.1%. Both BI and ML trees obtained a completely consistent topology. Consistent topology relationships are shown in Figure 3. In the phylogenetic trees, *I. rentianensis* sp. nov. formed a well-supported
Table 2. Conchological characters of *Inversidens rentianensis* sp. nov., *Inversidens brandtii*, *Inversidens pantoensis*. Characteristic descriptions of *I. brandtii* and *I. pantoensis* are referenced from Kondo (1982, 2008) and He and Zhuang (2013).

|                  | *I. rentianensis* sp. nov. | *I. brandtii* | *I. pantoensis*       |
|------------------|---------------------------|---------------|-----------------------|
| Shell shape      | Irregularly subtriangular | Ovate         | Inequilateral, quadrate|
| Umbo position    | 1/2 of shell length       | 1/4 of shell length | 1/3 of shell length   |
| Umbo sculpture   | Feebly wavy wrinkles      | Rippled       | Angularly wrinkled    |
| Surface sculpture| Concentric ridges         | Concentric ridges | Irregular growth lines|
| Nacre colour     | Reddish                   | Milk-white    | Bluish                |
| Posterior slope  | Sharp                     | Blunt         | Blunt                 |
| Ventral margin   | Nearly straight           | Arc-shaped    | Long and straight      |

Figure 2. Photograph of sampling site of *Inversidens rentianensis* sp. nov. in China.

sister-group relationship with *Inversidens brandtii* (PP = 1.00, BS = 100; Fig. 3). The genera *Pronodularia* and *Inversiunio* belong to different clades well-separated from *Inversidens* (Fig. 3).

Remarks. Species delineation can be problematic in the presence of morphological ambiguities due to phenotypic plasticity and convergence (e.g., cryptic species), especially in mollusks (Zieritz et al. 2010; Inoue et al. 2013). The use of molecular genetics can aid species delineation in the case of phenotypic plasticity and/or convergence (Pieri et al. 2018; Wu et al. 2018). *Inversidens rentianensis* sp. nov. can be
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Figure 3. Phylogenetic tree of freshwater mussels inferred from Bayesian Inference (BI) and Maximum Likelihood (ML) analyses of COI barcode. Support values above the branches are posterior probabilities (PP)/bootstrap support (BS). Red font indicates the new species from this study.

distinguished from congeneric species based on diagnostic characteristics of the shell. In this study, we also analyzed the interspecific divergence between *Inversidens brandtii* and *Inversidens rentianensis* sp. nov. based on the COI barcode. The results showed that the average interspecific divergence between the two species was 10.1%, which was much higher than intraspecific divergences. Genetic analysis conducted in this study supports *I. rentianensis* sp. nov. as a valid species, which can be easily distinguished by the COI barcode.

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