



Taxonomy of Three Oxytrichids (Protozoa, Ciliophora, Hypotrichia), With Establishment of the New Species Rubrioxytricha guangzhouensis spec. nov.

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Luo X, Huang J, Bourland WA, El-Serehy HA, Al-Farraj SA, Chen X and Song W (2021) Taxonomy of Three Oxytrichids (Protozoa, Ciliophora, Hypotrichia), With Establishment of the New Species Rubrioxytricha guangzhouensis spec. nov. Front. Mar. Sci. 7:623436. doi: 10.3389/fmars.2020.623436 Two oxytrichid ciliates collected from China, Oxytricha lithofera Foissner, 2016 and Rubrioxytricha haematoplasma (Blatterer and Foissner, 1990) Berger, 1999 were investigated based on living observations, protargol preparations, and molecular analyses. The Chinese population of O. lithofera shares highly distinctive features with the type population, e.g., lithosomes, abutting macronuclear nodules, straight undulating membranes, and long dorsal bristles. The morphology and morphogenesis of our new isolate of *R. haematoplasma* corresponds well with the type population. The 18S rRNA gene sequences of both species were obtained, and the molecular phylogeny of the genera Oxytricha Bory de Saint-Vincent in Lamouroux et al., 1824 and Rubrioxytricha Berger, 1999 was analyzed. In addition, we found that the Guangzhou population of R. haematoplasma described by Chen et al. (2015) can be distinguished from the type population and our isolate by the reddish-brown cortical granules (vs. lemon yellowish to greenish) and mitochondria-like granules present (vs. absent), as well as the slightly red cytoplasm (vs. slightly orange) and the marine or brackish water habitat (vs. freshwater). In addition, a 20-bp divergence in their 18S rRNA gene sequences indicates that they are not conspecific. Thus, we establish a new species, Rubrioxytricha guangzhouensis, for R. haematoplasma sensu Chen et al. (2015).

Keywords: hypotrich, morphology, morphogenesis, Oxytricha, phylogeny

INTRODUCTION

The family Oxytrichidae Ehrenberg, 1830, including more than 200 valid species, is one of the most morphologically and morphogenetically diverse hypotrichid ciliate groups (Berger, 1999, 2008; Foissner et al., 2002; Shao et al., 2011; Foissner, 2016; Hu et al., 2019; Kim et al., 2019; Dong et al., 2020; Wang et al., 2020a). The genus *Oxytricha* Bory de Saint-Vincent in Lamouroux et al., 1824, the type and time-honored genus of the family Oxytrichidae, comprises almost 40 nominal species isolated from various

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habitats worldwide and has been a melting pot for flexible 18cirri hypotrichs with caudal cirri for over one century (Berger, 1999; Shao et al., 2011, 2014; Weisse et al., 2013; Foissner, 2016; Kim and Min, 2019). Even though many studies have been carried out to delimit the genus more clearly (Berger, 1999; Shao et al., 2011, 2015; Foissner, 2016), it is still a polyphyletic genus with species irregularly distributed across the molecular tree (Schmidt et al., 2007; Paiva et al., 2009; Shao et al., 2014). The incongruence between morphology and molecular information suggests that this taxon is a genus with high genetic divergence (Berger, 1999; Foissner, 2016). Oxytricha lithofera, a highly distinctive species with a combination of features absent in any other described congeners (s. l.), was originally described by Foissner (2016) based on a population collected from moss on stones and soil from an old coral cave in the Golfete de Cuare del Indio, Venezuela. Here, we provide not only a description of the first Chinese population of O. lithofera but also the first analysis of the 18S rRNA gene sequence for this species.

In a revision of oxytrichids, Berger (1999) transferred two Oxytricha species, viz., Oxytricha haematoplasma Blatterer and Foissner, 1990 and Oxytricha ferruginea Stein, 1859, to the newly established genus Rubrioxytricha Berger, 1999, with the former as the type species. Rubrioxytricha was mainly characterized by the homogeneously colored cytoplasm (slightly orange to reddish in R. haematoplasma and rusty brown in R. ferruginea), the presence of one or two caudal cirri, and four to five dorsal kineties. Three more Rubrioxytricha species have since been reported, namely R. indica Naqvi et al., 2006, R. tsinlingensis Chen et al., 2017, and R. guamensis Kumar et al., 2018 (Naqvi et al., 2006; Chen et al., 2017; Kumar et al., 2018). Recently, Chen et al. (2015) reported a brackish/marine population collected from Guangzhou, China, under the name R. haematoplasma. Herein, we describe a freshwater population of R. haematoplasma based on a detailed investigation of morphology and morphogenesis. A comprehensive comparison between R. haematoplasma populations as well as congeners is provided, and a new species is established. Updated phylogenetic analyses based on 18S rRNA gene sequences were also performed, and the non-monophyly of the genera Oxytricha and Rubrioxytricha are discussed.

MATERIALS AND METHODS

Sample Collection, Observation, and Terminology

A brackish water sample containing *Oxytricha lithofera* was collected from a sandy beach near Zhuhai Fishing Girl Statue $(22^{\circ}15'41''N; 113^{\circ}35'16''E;$ **Figures 1A,B**) in Zhuhai, southern China, on 26 May 2014, when the water temperature was about 21°C and the salinity was about 3‰. The sample was maintained at about 22°C in Petri dishes with rice grains to enrich bacteria as a food source for the ciliates. *Oxytricha lithofera* was identified in this sample on 6 June 2014, when the salinity of the sample reached 6‰, and the population collapsed over the next 3 days. Investigations for *O. lithofera* were made with specimens from raw cultures rather than from cloned individuals. However, *O. lithofera* is easily distinguished by its long dorsal bristles

and no other *Oxytricha*-like morphotypes were present in the protargol preparations, indicating that our morphological and molecular studies deal with the same species.

Rubrioxytricha haematoplasma was collected on 8 and 11 April 2016 from a freshwater lake in Wuhan Botanical Garden, Chinese Academy of Sciences (30°32′57″N; 114°25′51″E; **Figures 1A,C**), Wuhan, China, when the water temperature was about 21°C. Sponge cubes, with the length about 10 cm, submerged at a depth of about 1 m for about 2 weeks, were used as artificial substrates to allow colonization by ciliates. Samples were collected according to Luo et al. (2018). Cells were selected and cultivated in Petri dishes at about 22°C with Volvic mineral water and rice grains. Although attempts to establish clonal cultures were unsuccessful, no other hypotrichid morphotypes were present in the protargol preparations, indicating that our morphological, morphogenetic, and molecular studies deal with the same species.

Live cells were isolated with micropipettes and then observed and measured with bright field and differential interference contrast microscopy at 40–1,000 × magnification using a Zeiss Axioplan 2 imaging microscope equipped with a Zeiss AxioCam HRc camera. The protargol staining method of Wilbert (1975) was used to reveal the nuclear apparatus and the ciliature. Counts, measurements, and photographs for protargol-stained specimens were made at a magnification of 1,000 × using an Olympus BX53 compound microscope (Tokyo, Japan) equipped with a Pixelink M5DC-SE-CYL camera (Ottawa, Canada). Drawings were made at a magnification of 1,000 × or 1,250 × with the aid of a drawing attachment or photomicrographs as templates. In the drawings of the morphogenetic stages, parental structures are shown as outlines, whereas new ones are shaded in black. Terminology is according to Berger (1999) and Foissner (2016).

DNA Extraction and Gene Sequencing

One to four randomly isolated cells were washed five times with filtered (0.22 µm) habitat water or Volvic mineral water and then transferred to a 1.5-ml microfuge tube as described by Zhang et al. (2020). Total genomic DNA was extracted using the DNeasy Blood and Tissue Kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions with modification according to Lu et al. (2018). The 18S rRNA gene was amplified using the primers 18S-F (5'-AACCTGGTTGATCCTGCCAGT-3') and 18S-R (5'-TGATCCTTCTGCAGGTTCACCTAC-3') (Medlin et al., 1988). Cycling parameters of touchdown PCR were as follows: one cycle of initial denaturation at 94°C for 5 min, followed by 18 cycles of amplification (94°C, 30 s; 66-49°C touch down, 40 s; 72°C, 2 min) and another 18 cycles (94°C, 30 s; 48°C, 40 s; 72°C, 2 min), with a final extension of 72°C for 7 min. The purified PCR product of O. lithofera of the appropriate size was inserted into the pEASY-T1 Cloning Vector (pEASY-T1 Cloning Kit, TransGen Biotech, Beijing, China) and sequenced using primers M13F (5'-CGCCAGGGTTTTCCCAGTCACGAC-3') and M13R (5'-AGCGGATAACAATTTCACACAGGA-3') and two internal primers at Sunny Biotechnology (Shanghai, China) (Zhang et al., 2019). For R. haematoplasma, cloning and sequencing in both directions were performed at Icongene



(Wuhan, China) using primers M13F and M13R and four internal primers E528F (5'-CGGTAATTCCAGCTCC-3'), E1146F (5'-TGCATGGCCGTTCTTAG-3'), RevD (5'-GGA GCTGGAATTACCG-3'), and RevE (5'-CTAAGAACGG CCATGC-3'). Contigs were assembled using Seqman V. 7.1.0 (DNAStar).

Phylogenetic Analyses

The 18S rRNA gene sequences of 70 hypotrichids and four oligotrichous ciliates, i.e., Strombidinopsis acuminate (FJ790209), Strombidium apolatum (DQ662848), Novistrombidium orientale (FJ422988), and Parastrombidinopsis minima (DQ393786), selected as outgroup taxa, were downloaded from the National Center for Biotechnology Information (NCBI) Database¹. The following stylonychines were selected: Gastrostyla steinii (AF508758), Histriculus histrio (FM209294), Laurentiella strenua (JX893368), Onychodromus grandis (AJ310486), Paraparentocirrus sibillinensis (KF184655), Parentocirrus sp. (KR063273), Pattersoniella vitiphila (JX885704), Pleurotricha lanceolata (AF508768), Rigidohymena candens (KC414885), Steinia sphagnicola (JX946276), Sterkiella cavicola (GU942565), Sterkiella histriomuscorum (FJ545743), Sterkiella nova (AF508771), Stylonychia ammermanni (FM209295), Stylonychia lemnae (AF508773), Stylonychia mytilus (AF508774), Stylonychia

notophora (FM209297), Styxophrya quadricornuta (X53485), Tetmemena bifaria (FM209296), and Tetmemena pustulata (AF508775). The sequence alignment was performed on the GUIDANCE web server² with the MUSCLE algorithm (Sela et al., 2015). The MUSCLE alignment score is 0.997502 and unreliable columns below confidence score 0.987 (97% of columns remain) were removed. The program BioEdit 7.2.5 (Hall, 1999) was used to remove the primers from the aligned sequences. The final alignment used for phylogenetic tree construction included 76 taxa with 1,703 positions. The GTR model of nucleotide substitution was selected as the best model by the program MrModeltest v.2.2 (Nylander, 2004) and was then used for Bayesian inference (BI) analysis. BI analysis was performed with MrBayes 3.2.7 on XSEDE (Ronquist et al., 2012) on the CIPRES Science Gateway³, with 2,000,000 generations, a sampling frequency of every 100th generation, and a burn-in of 5,000 trees. The remaining trees were used to calculate the posterior probabilities with a majority rule consensus. Maximum likelihood (ML) analysis was carried out using RAxML-HPC2 on XSEDE v 8.2.12 in CIPRES Science Gateway with the GTRGAMMA model (Miller et al., 2010; Stamatakis, 2014). The reliability of the internal branches was assessed by a non-parametric bootstrap method using 1,000 resamplings.

¹https://www.ncbi.nlm.nih.gov/

²http://guidance.tau.ac.il/

³https://www.phylo.org

Tree topologies were displayed with SeaView v 4.6.1 (Gouy et al., 2010) and MEGA 7.0.26 (Tamura et al., 2013). The systematic classification follows Lynn (2008).

The statistical probability of the monophyletic hypothesis of the genus *Rubrioxytricha* was evaluated using AU tests (Shimodaira, 2002) in the CONSEL package (Shimodaira and Hasegawa, 2001).

RESULTS

ZooBank Registration

Present work: urn:lsid:zoobank.org:pub:C09DE513-8C3A-422C-8953-2AFBF5581313.

Oxytricha lithofera Foissner, 2016 Deposition of Voucher Specimens

Two voucher slides with protargol-stained specimens were deposited in the Laboratory of Protozoology, OUC, China, with registration numbers LXT2014052601-01~02.

Morphology Based on the Zhuhai Population (Figures 2A–P, 3A–J and Table 1)

Cells 80–115 \times 25–40 μ m *in vivo*, ellipsoid in outline with both ends rounded, left margin more or less convex, right margin usually more or less straight (**Figures 2A,B, 3A–E**). Dorsoventrally flattened 2–3:1. Body flexible, not contractile. One contractile vacuole slightly above mid-body at left cell margin, about 8 μ m in diameter in diastole (**Figures 2A**,



FIGURE 2 | (A–P) Morphology and ciliature of the Zhuhai population of *Oxytricha lithofera* from life (A–D) and after protargol impregnation (E–P). (A) Ventral view of a representative individual. (B) Ventral views, showing body shape and lithosomes (arrows). (C,D) Distribution of cortical granules on the ventral (C) and dorsal (D) side. (E–G) Ventral (E) and dorsal (F,G) views of representative specimens, showing ciliature and nuclear apparatus; arrowheads in (E) mark pretransverse ventral cirri; arrow in (E) indicates buccal cirrus; arrowheads in (F,G) mark caudal cirri; arrows in (F,G) indicate micronuclei; double arrowheads in (F,G) mark the indeed very indistinct gap in dorsal kinety 1. (H,I) Ventral views showing the abutting macronuclear nodules and the large micronuclei (arrowheads) attached to the macronuclear nodules. (J) Ventral view, showing a specimen with an ingested ciliate (arrowhead). (K) Ventral view of a specimen with three frontoventral cirri; arrow heads indicate the long dorsal birstles of dorsal kinety 6. (L,M) Ventral (L) and dorsal (M) views of a representative specimen, the same one as (E,F); arrowheads mark caudal cirri. (N) Ventral view, showing ciliature, the abutting macronuclear nodules, and an ingested diatom (double arrowheads); arrowheads); arrowhead indicates paroral membrane; endoral membrane. (O,P) Show the slightly conspicuous one-bristle-wide gap in dorsal kinety 1. AZM, adoral zone of membranelles; e, endoral membrane; FC, frontal cirri; FVC, frontoventral cirri; RMR, right marginal row; TC, transverse cirri; 1–6, dorsal kineties. Scale bars = 30 μm (A,E–P) and 10 μm (C,D).



FIGURE 3 | (A–J) Morphology of the Zhuhai population of *Oxytricha lithofera* from life (A–E, bright field; F–J, differential interference contrast). (A–E) Ventral views of representative individuals, showing body shape, flexibility, contractile vacuole, lithosomes (arrowheads), and the long dorsal bristles (arrows). (F,G) Anterior (F) and posterior (G) part of the same specimen, showing three lithosomes (arrowheads), two macronuclear nodules (double arrowheads), two large-sized micronuclei (arrows), and crystals. (H) Ventral view, showing macronuclear nodules (double arrowheads), micronuclei (arrows), food vacuole (arrowhead), and inclusions. (I,J) Mitochondria-like cortical granules (arrowheads) on ventral (I) and dorsal (J) side; arrows in (J) indicate the long dorsal bristles. AZM, adoral zone of membranelles; CT, crystals; CV, contractile vacuole. Scale bars = 30 μm (A–E) and 10 μm (J).

3A). Mitochondria-like cortical granules present, globular, about 1.0-1.5 µm across, irregularly distributed on the ventral and dorsal sides (Figures 2C,D, 3I,J). Cytoplasm colorless, usually packed with numerous irregularly shaped shining granules (3-5 µm-sized), minute crystals (Figure 3F), and some food vacuoles, containing ciliates (Figure 2J, arrowhead), diatoms (Figure 2N, double arrowheads), and green algae (Figure 3H, arrowhead). One, two, or three ring-shaped lithosomes, $4-7 \,\mu m$ in diameter, with wall 1.0-1.5 µm thick, located subapically and/or subterminally (Figures 2B, 3A-D,F,G). Two abutting macronuclear nodules, positioned in mid-body and left of midline, broadly to elongate ellipsoid, on average 21 \times 15 μ m in protargol preparations, with anterior one extending slightly anterior to the buccal vertex (Figures 2F-I). Consistently two globular micronuclei about 4 µm across, each one attached to one macronuclear nodule, usually on the left side (Figures 2F-I,

3F-H). Locomotion by moderately fast crawling on the substrate, occasionally swimming.

Adoral zone extending 35–51% of body length (average 42%) in protargol preparations, composed of 21–29 membranelles (**Table 1**), with cilia 13–15 μ m long *in vivo*. Paroral and endoral membrane almost equal in length, more or less straight, optically side by side, or superimposed, with paroral extending slightly more anteriorly than endoral (**Figures 2E,L,N**).

Ventral cirral pattern as shown in **Figures 2E,K,L,N**. Three frontal cirri with the right one located behind the distal end of the adoral zone, easily misinterpreted as an adoral membranelle. One buccal cirrus, located anterior to the endoral membrane and right of the anterior end of the paroral membrane. Usually four frontoventral cirri (only one out of 20 cells observed with three, **Figure 2K**) arranged in V shape, three of them (IV/3, VI/3, VI/4) arranged in a line, cirrus III/2 slightly above the level of cirrus

Character	Species	Min	Max	Mean	М	SD	CV	n
Body, length (μm)	Ol	60	105	88.6	90	11.5	12.9	19
	Rh	122	175	151.1	154	15	10	17
Body, width (µm)	Ol	29	51	41.7	42	6.2	14.9	19
	Rh	45	74	62.1	63	7.9	12.7	17
Body width: length, percentage	Ol	40	60	47.3	45.7	5.4	11.5	19
	Rh	31	47	41.2	41.4	4	9.8	17
Adoral zone length (μm)	Ol	27	46	37	37	5.7	15.3	19
	Rh	47	62	53.4	53	3.7	7	17
Adoral zone length: body length, percentage	Ol	35	51	41.8	41	4.4	10.4	19
	Rh	29	41	35.6	35.1	2.8	8	17
Adoral membranelles, number	Ol	21	29	25.8	27	2.7	10.5	19
	Rh	34	41	37.6	38	2	5.4	17
Frontal cirri, number	Ol	3	3	3	3	0	0	19
	Rh	3	3	3	3	0	0	17
Buccal cirrus, number	Ol	1	1	1	1	0	0	19
	Rh	1	1	1	1	0	0	17
Frontoventral cirri, number	Ol	3	4	4	4	0.2	5.7	20
	Rh	4	4	4	4	0	0	17
Postoral ventral cirri, number	Ol	3	3	3	3	0	0	19
	Rh	3	3	3	3	0	0	13
Pretransverse ventral cirri, number	Ol	2	2	2	2	0	0	19
	Rh	2	2	2	2	0	0	16
Transverse cirri, number	Ol	5	5	5	5	0	0	19
	Rh	4	6	5	5	0.4	7.1	17
Left marginal cirri, number	01	14	18	15.4	15	1	6.6	19
	Rh	29	40	35.2	36	2.7	7.7	17
Right marginal cirri, number	01	13	16	14.9	15	0.8	5.4	19
	Rh	29	40	34.9	35	3	8.5	17
Caudal cirri, number	01	3	3	3	3	0	0	19
	Rh	1	1	1	1	0	0	17
Dorsal kineties, number	01	6	6	6	6	0	0	18
	Rh	4	4	4	4	0	0	17
Dikinetids in dorsal kinety 1, number	01	13	26	21.7	23	3.9	18.2	18
	Rh	18	27	23.6	24	2.3	9.9	17
Dikinetids in dorsal kinety 2, number	01	11	25	18.6	20	3.7	19.7	17
	Rh	19	30	22.8	23	2.7	11.6	17
Dikinetids in dorsal kinety 3, number	OI Dh	10	21	14.4	13.5	3.2	22	16
Dilipatida in deveal kinaty 4, number	Rh	19	28	22.6	23	2.4	10.6	17
Dikinetids in dorsal kinety 4, number	OI Dh	3	8	5.4	5	1.5	27.8	18
Dikinatida in daraal kinaty E. number	Rh Ol	14 5	26 18	19.1 9.1	19	3.1 2.8	16.1 30.9	16 18
Dikinetids in dorsal kinety 5, number Dikinetids in dorsal kinety 6, number		5 4		9.1 6.4	9		30.9 22.9	10
Macronuclear nodules, number	01	4	9		6 2	1.5 0	0	
Macionucieal nodules, number	Ol Rh	2	2 2	2 2	2	0	0	19 15
Anterior macronuclear nodule, length (μ m)	01	14	26			4	19.1	18
menor macronaciear noutre, ιθηθητη (μπη)	Oi Rh	14	26 33	21.2 24.8	22.5 25	4.8	19.1	15
Anterior macronuclear nodule, width (μ m)	Ol	9	33 20	24.8 15	25 15.5	4.8 3.5	23.5	18
niterior macionaciear nodale, wiath (µm)	0i Rh	9 8	20 13	10.1	10	3.5 1.8	23.5 17.4	15
Posterior macronuclear nodule, length (μ m)	Ol	o 14	30	22.2	22.5	4.6	20.8	18
ספנפווטו ווומטוטווטטופמו ווטעעופ, ופווטנוו (μΠ)	Oi Rh	14	30	22.2	22.5	4.6 3.4	20.8 13.8	15
Posterior macronuclear nodule, width (μ m)	Ol	19	30 18	25 13.4	25 14	3.4 2.6	13.8	18
	UI	10	10	10.4	14	2.0	19.0	10

(Continued)

TABLE 1 | Continued

Character	Species	Min	Max	Mean	М	SD	CV	n
Micronuclei, number	OI	2	2	2	2	0	0	9
	Rh	2	3	2.1	2	0.3	12.9	14
Micronucleus, length (μ m)	OI	3	5	4.1	4	0.5	13.2	11
	Rh	5	8	6.5	6.5	0.7	11.5	28
Micronucleus, width (μ m)	OI	3	5	3.8	4	0.6	15.1	12
	Rh	3	5	4.2	4	0.5	11.8	28

Abbreviations: CV, coefficient of variation in%; M, median; Max, maximum; Mean, arithmetic mean; Min, minimum; n, number of cells measured; SD, standard deviation.

VI/3. Three postoral ventral cirri arranged in a triangular pattern. Two pretransverse ventral cirri, with the posterior one almost at the same level as transverse cirrus III/1. Five enlarged transverse cirri, arranged in a check-mark shape, with cilia 18–20 μ m long *in vivo*, conspicuously protruding beyond the posterior end of the cell (**Figures 2A, 3A–E**). Single marginal row on either body side, separated by a distinct gap, left one with 14–18 cirri, right one with 13–16 cirri, with cilia about 13 μ m in length.

Invariably six dorsal kineties (DK): DK1, the leftmost kinety, strongly shortened anteriorly, reaches the posterior end, with an indeed very indistinct gap in most specimens (Figures 2F,G, double arrowheads), only six out of 25 cells observed with a slightly conspicuous gap (Figures 20,P, arrowhead); DK2 almost extending the entire body length; DK3 extends to the anterior body end and splits to form a short row of DK4 (on an average of five dikinetids) at the subterminal end, usually one or two dikinetids between DK3 and DK4; DK5 and DK6, the dorsomarginal kineties, shortened anteriorly, with DK5 reaching the mid-body, DK6 extends slightly above the midbody (Figures 2F,G,M). Bristles conspicuous in vivo, even at low magnification (Figures 3A-C,J, arrows), anterior bristles of DK1–3 about 5 μ m long, gradually increase to 9 μ m posteriorly; bristles of DK4 usually 8-9 µm long, bristles of DK5 and DK6 usually 5-7 µm long. Three long caudal cirri, with cilia 18-20 µm long, one each at end of DK1, 2, and 4, optically positioned in gap between ends of marginal rows (Figures 2F,G,M).

Rubrioxytricha haematoplasma (Blatterer and Foissner, 1990) Berger, 1999

Deposition of Voucher Specimens

Voucher slides with protargol-stained specimens were deposited in the Laboratory of Protozoology, OUC, China, with registration numbers LXT2016040802-01~03 and LXT2016041102-01~02.

Morphology Based on the Wuhan Population (Figures 4A–N and Table 1)

Cells 125–160 × 40–60 μ m *in vivo*. Body soft, flexible and not contractile. Body long ellipsoid in shape, both anterior and posterior ends rounded, some with broad ends, some with narrower ends, right and left margins more or less convex (**Figures 4A,D,G-I**). Dorsoventrally flattened about 2:1. Single contractile vacuole slightly behind the buccal vertex and ahead of the mid-body, about 45% of body length, at the left cell margin, approximately 12 μ m in diameter in diastole, two longitudinally oriented collecting canals present (Figures 4A,D,G-I). Cortical granules of single type, spherical, 0.8-1.0 µm across, lemon yellowish to greenish, arranged in lines or small groups then in long lines on both body sides (Figures 4B,C,J-L, slightly squeezed cells). Cytoplasm slightly orange (Figures 4J,L), usually packed with numerous irregularly shaped granules (2-6 µm-sized) and food vacuoles, containing diatoms, sometimes rendering cells yellow-brown to slightly darkish at low magnification. Consistently two macronuclear nodules, positioned in the mid-body and more or less left of midline, with the anterior one almost at the level of the buccal vertex, each nodule elongate ellipsoidal in shape, on average $25 \times 10 \ \mu m$ after protargol preparation. Usually two ellipsoidal micronuclei, 5-8 \times 3-5 μ m in size, with one attached to or near each macronuclear nodule (Figures 4F,M,N). Locomotion by slowly crawling on the substrate, occasionally swimming with cells rotating around the longitudinal axis.

Adoral zone 29–41% of body length (average 36%) in protargol preparations, composed of 34–41 membranelles (**Table 1**). Cilia of membranelles 18–20 μ m long *in vivo*, widest base about 10 μ m. Undulating membranes in *Australocirrus* pattern (Kumar and Foissner, 2015), that is, paroral moderately to distinctly curved but not recurved distally, paroral extending slightly anteriorly to endoral and almost equal in length, both membranes optically intersecting at middle of paroral (**Figures 4E,M**).

Ventral cirral pattern as shown in **Figures 4E,M**. Three slightly enlarged frontal cirri continuous with frontoventral cirri arranged in a V-shaped pattern, rightmost three (IV/3, VI/3, VI/4) almost arranged in a line, cirrus III/2 slightly ahead level of the posteriormost one (cirrus IV/3) and distant from cirrus VI/3. Single buccal cirrus, almost at the same level with the right frontal cirrus. Three postoral ventral cirri and two pretransverse ventral cirri. On average five slightly enlarged transverse cirri, with cilia $20-22 \,\mu$ m long, extending slightly behind the rear end of the cell, arranged in a check-mark shape, just behind the pretransverse ventral cirri. Single left and right marginal row, more or less confluent at the posterior end.

Invariably four dorsal kineties (DK): three bipolar ones (DK1–3) with about 23 dikinetids each, one slightly shortened dorsomarginal kinety (DK4) comprising about 19 dikinetids. Always one parental dorsal kinety fragment located between DK3 and DK4, with sparsely arranged monokinetids (**Figures 4F,N**, arrowheads). Dorsal bristles 3–4 μ m long *in vivo* (**Figure 4L**, arrowheads). Single caudal cirrus located at the end of DK3 (**Figures 4F,M**, arrow).





Morphogenesis (Figures 5A-H, 6A-H and Table 1)

In a very early divider (**Figures 5A**, **6A**), stomatogenesis commences with appearance of the oral primordium as groups of basal bodies between the buccal vortex and transverse cirri, including a large group just left to the postoral ventral cirri. At this stage, all parental cirri remain intact and do not contribute to the formation of the oral primordium. Somewhat later (**Figures 5B, 6B**), the anterior two postoral ventral cirri (IV/2, V/4) dedifferentiate and contribute to the oral primordium. In an early divider (**Figures 5C, 6C**), the posterior postoral ventral cirrus (V/3) is no longer recognizable and the oral primordium

gives rise to new adoral membranelles. Simultaneously, the parental undulating membranes dedifferentiate anteriorly to form the undulating membranes anlage (UMA) (Figure 5C, arrow) and the frontal-ventral-transverse cirral anlagen (FVTA) begin to form. In a middle divider (Figures 5E, 6D), two sets of FVTA are formed. Subsequently (Figures 5G, 6F), the UMA forms the new undulating membranes and gives rise to the left frontal cirrus. At the same time, the other streaks differentiate into new cirri in the pattern of 3:3:3:4:4 from left to right. The parental adoral zone, which is inherited by the proter, remains unchanged throughout the whole process (Figures 5C,E,G).



FIGURE 5 | (A–H) Ciliature of the Wuhan population of *Rubrioxytricha haematoplasma* during the ontogenetic process after protargol impregnation. (A,B) Ventral views of very early dividers, showing the formation of the oral primordium. (C,D) Ventral (C) and dorsal (D) views of an early divider, showing the formation of frontal–ventral–transverse cirral anlagen; arrow in (C) indicates undulating membranes anlage of proter (= anlage I) dedifferentiated from old structure; arrowhead in (C) marks right marginal anlage of proter; arrows in (D) show newly formed dorsal kinety anlagen; arrowheads in (D) indicate grandparental dorsal bristles. (E,F) Ventral (E) and dorsal (F) views of a middle divider, showing two sets of frontal–ventral–transverse cirral anlagen and dorsal kinety anlagen (arrows); arrowheads in (E) indicate left marginal anlage of proter and opisthe; arrowheads in (F) indicate grandparental dorsal bistles. (G,H) Ventral (G) and dorsal (H) views of a late divider, showing frontal–ventral–transverse cirral anlagen of proter and opisthe; arrowheads in (F) indicate grandparental dorsal bristles. (G,H) Ventral (G) and dorsal (H) views of a late divider, showing newly formed frontal–ventral–transverse cirral anlagen (G) mark organization (G) mark organization (G) mark dorsol (H) views of a late divider, showing newly formed frontal–ventral–transverse cirral mass and micronucleus; arrowheads in (G) mark dorsol with anlagen formed near the anterior ends of right marginal anlagen; arrowheads in (H) indicate grandparental dorsal bistles; arrow marks new caudal cirrus formed at the end of dorsal kinety 3. LMA, left marginal anlagen; Ma, macronuclear nodules; Mi, micronuclei; OP, oral primordium; RMA, right marginal anlagen; I–VI, frontal–ventral–transverse cirral anlagen; 1–4, parental dorsal kineties. Scale bars = 50 μm.

The marginal anlagen develop by dedifferentiation of the parental structures, and the right marginal anlage of the proter appears earlier than others (Figures 5C,E, 6C,D). Then, all the anlagen extend longitudinally and generate new cirri in both proter and opisthe (Figures 5G, 6F).

Dorsal morphogenesis occurs by two groups of primordia: one group develops by dedifferentiation of the parental structures

at the middle stage, that is, each of dorsal kineties 1–3 forms an anlage in both proter and opisthe (**Figures 5F**, **6E**, arrows). Later, all the anlagen lengthen at both ends, and one caudal cirrus is generated at the end of dorsal kinety 3 (**Figures 5H**, **6H**). The second group, that is, the dorsomarginal kinety primordia, develops *de novo* to the anterior right of the right marginal row anlage during the late stage (**Figures 5G**, **6F**, arrowheads).





During the late stage of the ontogenetic process, the macronuclear nodules fuse with one another and the micronuclei fuse into a single mass as well (**Figures 5H**, **6G**).

18S rRNA Gene Sequences and Phylogenetic Analyses

The 18S rRNA gene sequence of *Oxytricha lithofera* (excluding both primers) has been deposited in GenBank with accession number MT364897. It has a length of 1,726 bp and a G + C content of 45.48%. The closest sequence to *O. lithofera* is *Urosoma salmastra* (KF951419), with the sequence similarity of 98.3% (28 nucleotide difference). The 18S rRNA gene sequence of *Rubrioxytricha haematoplasma* (excluding both

primers) has been deposited in GenBank with accession number MT364898. It has a length of 1,724 bp and a G + C content of 45.01%. The 18S rRNA gene sequences of all six *Rubrioxytricha* isolates differ from each other by 0 to 26 nucleotides, with sequence identities from 98.4 to 100% (**Table 2**).

In the phylogenetic trees, *Oxytricha* species are separated into several clades. *Oxytricha lithofera* is sister to *U. salmastra* (KF951419) and distant from the type species of the genus, *O. granulifera* (AF508762), and *O. granulifera chiapasensis* (KX889988) although this relationship is not robust as indicated by the low support in the ML tree (ML/BI, 54%/1.00). All species of *Rubrioxytricha* fall within a wellsupported assemblage (ML/BI, 91%/1.00) although this genus

Species (accession number)	1	2	3	4	5	6
1. R. haematoplasma (MT364898)	-	1	1	6	9	20
2. R. tsinlingensis (KR817675)	0.999	-	0	5	10	21
3. Rubrioxytricha sp. (MG603612)	0.999	1.000	-	5	10	21
4. R. ferruginea (AF370027)	0.996	0.996	0.996	_	15	26
5. R. guamensis (KY947508)	0.994	0.993	0.993	0.990	-	17
6. R. guangzhouensis spec. nov. (KJ645977)	0.987	0.987	0.987	0.984	0.989	-

TABLE 2 | Numbers of unmatched nucleotides (upper right) and distribution of percentages of sequence identity (lower left) to all available 18S rRNA gene sequences of Rubrioxytricha species.

is not monophyletic as two sequences under the names of *Polystichothrix monilata* (KT192639) and *Pseudogastrostyla flava* (KP266627) nest within it. *Rubrioxytricha guamensis* (KY947508) and *R. guangzhouensis* spec. nov. (KJ645977) branched off first in the large assemblage. The new sequence *R. haematoplasma* (MT364898) is most closely related with a cluster including three congeners, *R. tsinlingensis* (KR817675), *Rubrioxytricha* sp. (MG603612), and *R. ferruginea* (AF370027). However, the sister relationship between the four abovementioned species of *Rubrioxytricha* and the group of *Polystichothrix–Pseudogastrostyla* does not receive high support in the ML tree (ML/BI, 59%/1.00). The morphological hypothesis that *Rubrioxytricha* is a monophyletic group was rejected by the AU test (*p* value = 4×10^{-17}).

DISCUSSION

Identification of the Chinese Population of *Oxytricha lithofera* and Emended Diagnosis of the Species

Based on a population collected from Golfete de Cuare del Indio (salinity 10‰), Venezuela, Foissner (2016) reported a highly distinctive species, O. lithofera, with a combination of features not present in any other Oxytricha species. Our population, collected from a sandy beach (salinity 3-6%) of Zhuhai, shares most of the following typical features with the Venezuelan population: (1) cirral pattern; (2) straight and parallel undulating membranes; (3) abutting macronuclear nodules, usually with a globular micronucleus attached to each; (4) large lithosomes present; (5) dorsal bristles increasing in length from the anterior end to the posterior end of dorsal kineties, up to 10 µm long; and (6) mitochondria-like granules present. Compared with the type population, the Zhuhai population has (1) a relatively larger body size in vivo (80-115 \times 25-40 vs. 60–85 \times 23–32 μ m) and after protargol impregnation (60–105 \times 29–51 vs. 53–75 \times 20–28 μ m) (shrinkage differed between the two methods); (2) more adoral membranelles (21-29 vs. 19-21); and (3) more marginal cirri (left, 14-18 vs. 12-15; right, 13-16 vs. 9-12). However, all of these differences, most of which overlap with each other, can be considered as population-dependent geographic variations (Berger, 1999; Chen et al., 2020). One difference that cannot be ignored is that there is a one-bristle-wide gap in dorsal kinety 1 of the type population, while the gap in most of the specimens of the Zhuhai

population is absent. This may be seen as an intraspecific or intrapopulational difference. Therefore, we consider the Zhuhai population conspecific with the Venezuelan population. Based on the information of the Zhuhai population and the type population, an emended diagnosis is supplied here.

Emended Diagnosis of Oxytricha lithofera

Size $60-115 \times 23-40 \ \mu m$ *in vivo*; body usually elongate ellipsoid. Mitochondria-like cortical granules present, globular, about 1.0–1.5 $\ \mu m$ across. One, two, or three ring-shaped lithosomes, located subapically and/or subterminally. Adoral zone about 39% of body length, with about 23 membranelles. Undulating membranes straight, optically side by side. Eighteen frontoventral, transverse cirri. Six dorsal kineties with two dorsomarginal rows included, dorsal bristles gradually increasing in length from the anterior end to the posterior end. Three conspicuous caudal cirri. Two abutting ellipsoid macronuclear nodules and two globular micronuclei. Brackish habitat.

Identification of the Populations of *Rubrioxytricha haematoplasma*

Hitherto, five *Rubrioxytricha* species have been reported, namely *R. haematoplasma*, *R. ferruginea*, *R. indica*, *R. tsinlingensis*, and *R. guamensis* (Berger, 1999; Naqvi et al., 2006; Chen et al., 2017; Kumar et al., 2018). The type species of the genus, *R. haematoplasma*, was originally reported by Blatterer and Foissner (1990) based on a German population. Subsequently, Shin and Kim (1993) described a Korean population. Recently, Chen et al. (2015) reported a brackish/marine population collected from Guangzhou, China, based on detailed morphological, morphogenetic, and phylogenetic analyses.

The main features of the Wuhan population correspond well with those of the type population of *R. haematoplasma*, e.g., body shape, cirral pattern, dorsal ciliature, especially the color of cortical granules and cytoplasm. There is a tiny difference in terms of the distribution pattern of cortical granules, that is, the cortical granules of the type population are arranged in regular lines as shown in the drawings (Figure 5d in Blatterer and Foissner, 1990), while the cortical granules in the Wuhan population seem more or less irregular, with some in small groups and then in long lines. In addition, the postoral ventral cirri in the type population are distributed in a line, while those in the Wuhan population are in a V pattern. However, we consider that these differences are not enough to separate them as different species. The Wuhan population of *R. haematoplasma* can be easily distinguished from the other four congeners by the characteristics shown in **Table 3**, while the differences of the 18S rRNA gene sequences of the five *Rubrioxytricha* members ranges from 1 to 20 nucleotides (**Table 2**). It is noteworthy that there is only one nucleotide difference of the 18S rRNA gene between *R. haematoplasma* and *R. tsinlingensis*, while morphologically, they have distinct different dorsal ciliature (four dorsal kineties with only one caudal cirrus vs. six dorsal kineties with three caudal cirri) (**Table 3**). The highly conserved nature of the 18S rRNA gene has been noted in other oxytrichid morphospecies as well (Kumar et al., 2017; Fan et al., 2021). Thus, the Wuhan population can be identified as *R. haematoplasma*.

There was no information about cortical granules or cytoplasm color of the Korean population (Shin and Kim, 1993). The differences in numbers of adoral membranelles and in the dorsal ciliature were considered as geographical variations by Berger (1999) and population differences by Chen et al. (2015). Herein, we tentatively accept these opinions until more information (cortical granules, cytoplasm, molecular data) is available for the Korean population.

Chen et al. (2015) did mention that the paroral and endoral membranes of the Guangzhou population of R. haematoplasma are more curved than those of the type population. However, this might be a preparation artifact, because photomicrographs of live specimens do not show a wide and deep buccal cavity as discussed by Kumar et al. (2018), and the undulating membranes of the early dividers (Figures 3a,b in Chen et al., 2015) do not show a strong degree of curvature either. Therefore, there is no distinct difference of the undulating membranes among the populations, all of which are arranged in an Australocirrus pattern. Compared with the Guangzhou population described by Chen et al. (2015), the Wuhan population and the type population of R. haematoplasma show significant differences in the following aspects: (1) cortical granules (lemon yellowish to greenish in color vs. reddish-brown), (2) mitochondria-like granules (absent vs. present), (3) color of the cytoplasm (slightly orange vs. slightly red), and (4) the habitat (freshwater vs. brackish/marine) (Table 3). Hence, we consider the Guangzhou population as a different species from R. haematoplasma. Furthermore, the 18S rRNA gene sequences (20 nucleotide difference between the Wuhan and the Guangzhou populations, Table 2) indicate that these two populations represent separate species.

In terms of cytoplasm color, cortical granules, cirral pattern, and dorsal ciliature, the Guangzhou population of *R. haematoplasma* should be compared with the other four species of the genus. The Guangzhou population of *R. haematoplasma* can be distinguished from all the others by the color of cortical granules (reddish-brown vs. *R. ferruginea*, brownish; *R. guamensis*, yellowish; *R. indica*, dark green; *R. tsinlingensis*, yellow-green) and the presence of mitochondria-like granules (vs. absent in all the congeners). For more differences among all the species, see **Table 3**. Thus, the Guangzhou population represents a separate species of the genus *Rubrioxytricha*.

Establishment of a New *Rubrioxytricha* Species

Rubrioxytricha guangzhouensis spec. nov. (Figures 7A–J)

2015 *Rubrioxytricha haematoplasma* (Blatterer and Foissner, 1990) Berger, 1999 – Chen et al., Int. J. Syst. Evol. Microbiol., 65, 309–320.

Remarks

As discussed above, a new species should be established for the Guangzhou isolate which was described under the name *R. haematoplasma* by Chen et al. (2015).

Chen et al. (2015) stated "the Chinese population came from two shrimp-farming ponds, one brackish (salinity 8.6%) and the other marine (34.4%). However, the brackish water was created artificially by mixing together marine and freshwater, and material was regularly transferred between the two ponds, so it is unclear whether the Chinese population of *R. haematoplasma* was originally from a marine or a freshwater habitat." We think it is possible that the Guangzhou isolate was originally from the marine pond and then transferred to the brackish pond. Therefore, we tentatively treat the marine pond as the type locality.

Diagnosis

Size 90–180 \times 30–70 μ m *in vivo*; body elongate oval, brownreddish in color. Cortical granules, approximately 0.8 μ m in diameter, reddish-brown, grouped in lines on dorsal; mitochondria-like granules about 3 μ m in diameter, colorless and spherical, densely packed. Adoral zone about 30% of body length, with about 37 membranelles. Eighteen frontoventral, transverse cirri. Three bipolar dorsal kineties and one slightly shorter dorsomarginal row; one caudal cirrus at the posterior end of dorsal kinety 3. Two macronuclear nodules and one to three micronuclei. Brackish/marine habitat.

Etymology

The species-group name *guangzhouensis* refers to the geographic locality where the ciliate population was discovered.

Type Locality

Marine shrimp-farming pond (salinity 34%) at the campus of the South China Normal University, Guangzhou, China ($23^{\circ}08'19''$ N; $113^{\circ}21'22''$ E).

Type Specimens

The protargol-stained slide containing the holotype specimen (**Figures 7C–E**) and 10 paratype slides were deposited in the Laboratory of Protozoology, OUC, China, with registration numbers CXM08112701-01~11.

ZooBank Registration

Rubrioxytricha guangzhouensis spec. nov. is registered in ZooBank under: urn:lsid:zooBank.org:act:036912E2-4662-40D1-B666-B4091FD50DE1.

TABLE 3 | Morphological and morphometric comparison of Rubrioxytricha species.

Character	Rubrioxytricha haematoplasma	Rubrioxytricha haematoplasma	Rubrioxytricha haematoplasma	Rubrioxytricha guangzhouensis spec. nov.	Rubrioxytricha ferruginea	Rubrioxytricha guamensis	Rubrioxytricha indica	Rubrioxytricha tsinlingensis
BL ^a (μm)	125-160	120–180	100-180	90–180	150-200	85–110	80	100–180
BW ^a (μm)	40-60	40–55	50-90	30–70	/	25–45	30	35–60
BL ^b (μm)	122-175	91–143	112-160	133–195	122-162	74–96	65-74	144–186
BW ^b (μm)	45-74	29–51	42-80	50–95	39–51	20–38	24–32	64–89
AZM, n	34–41	30–48	21-30	33–45	35–41	23–32	27–31	36–44
CC, n	1	0.8 (0-1)	0.7 (0-1)	1	1.7 (1-2)	1.1 (1–2)	1	3
DK, n	4	4	4.1 (4-5)	4	5	4	5	6
DM, n	1	1	1	1	/	1	2	/
DKF	Absent	Absent	Absent	Absent	/	Absent	Absent	/
Cortical granules	Lemon yellowish to greenish	Lemon yellowish to greenish	/	Reddish-brown	Brownish	Yellowish	Dark green	Yellow-green
Color of the cytoplasm	Slightly orange	Slightly orange to reddish	/	Slightly red	Rusty	Colorless	More or less colorless	Slightly brown
Color of the cell	Yellow–brown to slightly darkish	Darkish at low magnification	/	Dark-brownish to ashblack	Rusty brown	/	/	Yellow-brown
Habitat	Freshwater pond	Freshwater river	Freshwater river	Brackish/marine	Eutrophic pond	Freshwater pond		Freshwater pond
Data source	Present work	Blatterer and Foissner (1990); Berger (1999)	Shin and Kim (1993)	Chen et al. (2015)	Song and Wilbert (1989); Berger (1999)	Kumar et al., 2018	Naqvi et al. (2006)	Chen et al. (2017)

Characters marked with "/" were not available from the source cited. Abbreviations: AZM, adoral zone of membranelles; BL, body length; BW, body width; CC, caudal cirri; DK, dorsal kineties; DKF, dorsal kinety fragmentation; DM, dorsomarginal kineties; n, number. ^a Data based on specimens in vivo. ^b Data based on specimens after protargol impregnation.



FIGURE 7 | (A–J) Morphology and ciliature of *Rubrioxytricha guangzhouensis* spec. nov. from life (A,B,F,G, bright field; H–J, differential interference contrast) and after protargol impregnation (C–E). (A–J) From Chen et al. (2015). (A) Ventral view of a representative individual. (B) Ventral view of a bending cell to show the flexibility of the body; arrow marks the frontal scutum. (C–E) Ventral (C,E) and dorsal (D) views of the holotype specimen, showing ciliature and nuclear apparatus; arrowheads in (C) mark pretransverse ventral cirri; double arrowheads in (C,E) mark buccal cirrus; double arrowheads in (D) show micronuclei; arrow in (D,E) indicates the single caudal cirrus; arrows in (E) show frontal cirri. (F) Ventral view of the typical individual; arrowhead marks the contractile vacuole. (G) Ventral view of bending cell to show the flexibility of the body; arrow marks the transparent membrane-like structure at the anterior part of the adoral zone. (H) Ventral view of anterior cell part; arrow marks the transparent membrane-like structure at the anterior (I) and middle (J) cell parts; arrows indicate the reddish-brown, spherical, type-I cortical granules arranged in lines; arrowheads mark the colorless, densely distributed, mitochondria-like granules. AZM, adoral zone of membranelles; CV, contractile vacuole; e, endoral membrane; FC, frontal cirri; FVC, frontoventral cirri; LMR, left marginal row; Ma, macronuclear nodules; p, paroral membrane; PVC, postoral ventral cirri; RMR, right marginal row; TC, transverse cirri; UM, undulating membranes; 1–4, dorsal kineties. Scale bars = 60 μm.

Morphogenetic Comparison of *Rubrioxytricha* Species

Blatterer and Foissner (1990) gave an incomplete morphogenetic description for the type population of *R. haematoplasma*. In the present study, we supply somewhat more detailed morphogenetic information for *R. haematoplasma* based on the Wuhan population. Chen et al. (2015) provided a detailed morphogenetic description for *R. guangzhouensis* spec. nov. Naqvi et al. (2006) reported the morphogenetic process of *R. indica* and Kumar et al. (2018) provided brief notes on the ontogenesis of *R. guamensis*. All the abovementioned populations share the following morphogenetic features: (1) the parental

adoral zone of membranelles is completely inherited, unchanged, by the proter; (2) the undulating membranes anlage of the proter originates by dedifferentiation of the parental structures; (3) parental buccal cirrus (II/2), two frontoventral cirri (III/2, IV/3), and three postoral ventral cirri (IV/2, V/3, V/4) contribute to the formation of the anlagen; (4) the frontal–ventral–transverse (FVT) cirral anlagen are probably formed *via* a primary pattern and the FVT cirri are formed in a typical *Oxytricha* pattern, i.e., in the pattern of 1:3:3:3:4:4 from left to right; (5) marginal anlagen develop intrakinetally within the parental structures; (6) dorsal kineties are formed in the *Urosomoida* pattern, i.e., dorsomarginal row present and without fragmentation of kinety 3; (7) a single caudal cirrus (seldom two caudal cirri



corresponds to 0.01 expected substitutions per site. New sequences provided in the present work are indicated in bold.

in *R. guamensis*) originates at the end of dorsal kinety 3; and (8) two macronuclear nodules fuse into a single mass during the morphogenetic process. This combination of features reveals a highly conservative morphogenetic pattern of the above *Rubrioxytricha* species (Naqvi et al., 2006; Chen et al., 2015; Song and Shao, 2017; Kumar et al., 2018).

Phylogenetic Analyses

As described in previous studies, *Oxytricha* and *Urosoma* species are irregularly distributed over the phylogenetic trees, revealing that both the genus *Oxytricha* and *Urosoma* are non-monophyletic and lack obvious synapomorphies (Schmidt et al., 2007; Paiva et al., 2009; Shao et al., 2014, 2019; Gao et al., 2016; Jung and Berger, 2019; Kaur et al., 2019; Kim and Min, 2019; Dong et al., 2020; Wang et al., 2020b; Xu et al., 2020). In the present phylogenetic trees (**Figure 8**), *O. lithofera* is placed sister to *U. salmastra* (KF951419). Morphologically, they do have the same features as follows: flexible body, single marginal row on each body side, three caudal cirri, two macronuclear nodules, and two micronuclei, which, however, are shared by most of the oxytrichids (Berger, 1999; Wang et al., 2017). Moreover, *O. lithofera* can be easily distinguished from *U. salmastra* by an adoral zone of membranelles more or less in *Stylonychia* pattern

(vs. typical *Gonostomum* pattern), having more transverse cirri (five vs. four), and more dorsal kineties (six, kinety fragmentation probably present vs. four, kinety fragmentation absent), which all have been considered as genus-level diagnostic features for the oxytrichids (Berger, 1999; Shao et al., 2015; Wang et al., 2020b). In addition, the grouping of *O. lithofera* and *U. salmastra* is far from stable as this relationship does not receive high support in the ML analysis. The topology may very likely be changed when additional molecular data from more taxa are available.

All species of *Rubrioxytricha* are intermingled with *P. monilata* (KT192639) and *P. flava* (KP266627) in a highly supported assemblage, indicating that *Rubrioxytricha* is paraphyletic. The hypothesis that *Rubrioxytricha* species form a monophyletic group was rejected by the AU test, although the internal relationships are not yet robustly resolved. The close relationship between *Rubrioxytricha* and *Polystichothrix–Pseudogastrostyla* is morphologically supported as they all have a flexible body, one marginal row on each body side, and brightly colored cortical granules. However, *Rubrioxytricha* differs significantly from *Polystichothrix* and *Pseudogastrostyla* in having a typical oxytrichid pattern of 18 FVT cirri (vs. more than 18 FVT cirri in short lines) (Fan et al., 2015; Luo et al., 2017). The phylogenetic placement

of the genera Polystichothrix and Pseudogastrostyla might be explained by the assumption that those taxa with more than 18 FVT cirri evolved by a secondary increase in the numbers of FVT anlagen and/or additional cirri originated from some of the anlagen (Berger, 1999; Luo et al., 2017). Moreover, the dorsal ciliature pattern, which also carries an informative phylogenetic signal, like the ventral cirral pattern (Berger, 1999; Shao et al., 2011, 2015; Foissner, 2016), shows great variety in the Rubrioxytricha-Polystichothrix-Pseudogastrostyla clade: (1) four dorsal kineties in R. guamensis, R. haematoplasma, R. indica, R. guangzhouensis spec. nov., and Pseudogastrostyla, with one dorsomarginal kinety present, dorsal kinety fragmentation absent, only one caudal cirrus (seldom two caudal cirri in *R. guamensis*) present at the end of dorsal kinety 3; (2) five dorsal kineties in R. ferruginea and Polystichothrix, very likely including one dorsomarginal kinety and dorsal kinety 4 originated by dorsal kinety 3 fragmentation, one or two caudal cirri present in R. ferruginea and caudal cirri lacking in Polystichothrix; (3) six dorsal kineties in R. tsinlingensis, probably with two dorsomarginal rows included and dorsal kinety 4 originated by dorsal kinety 3 fragmentation, three caudal cirri in total, one each at the posterior end of dorsal kineties 1, 2, and 4 (Berger, 1999; Naqvi et al., 2006; Chen et al., 2015, 2017; Fan et al., 2015; Luo et al., 2017; Kumar et al., 2018).

The incongruence between morphology and molecular data suggests that *Oxytricha* and *Rubrioxytricha* are genera with high genetic divergence, which need to be broken up into multiple new genera. At present, there are over 1,000 nominal hypotrich ciliates (Chen et al., 2017); however, the 18S rRNA gene sequences are only available for about 30% of this group (NCBI database). Thus, molecular data from additional taxa and even

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phylogenomic data are needed to reveal the natural phylogeny of this group of ciliates.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are publicly available. This data can be found here: GenBank (https://www.ncbi.nlm.nih.gov/genbank/) with accession numbers MT364897-MT364898.

AUTHOR CONTRIBUTIONS

XL and JH collected the samples. XL carried out the experiments (live observation, protargol impregnation, and DNA amplification) and analyses (morphometrics and phylogenetic analyses). XC and WS supervised the work. All authors wrote the manuscript and approved the final version.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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