- 1 Sequencing, de novo assembly of *Ludwigia* plastomes, and comparative analysis within
- 2 the Onagraceae family
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#### 11 Abstract

12 The Onagraceae family, which belongs to the order Myrtales, consists of approximately 657 13 species and 17 genera. This family includes the genus *Ludwigia* L., which is comprised of 82 14 species. In this study, we focused on the two aquatic invasive species Ludwigia grandiflora 15 subsp. hexapetala (Lgh) and Ludwigia peploides subsp montevidensis (Lpm) largely distributed 16 in aquatic environments in North America and in Europe. Both species have been found to 17 degrade major watersheds leading ecological and economical damages. Genomic resources for 18 Onagraceae are limited, with only Ludwigia octovalvis (Lo) plastid genome available for the 19 genus Ludwigia L. at the time of our study. This scarcity constrains phylogenetic, population 20 genetics, and genomic studies. To brush up genomic ressources, new complete plastid genomes 21 of Ludwigia grandiflora subps. hexapetala (Lgh) and Ludwigia peploides subsp. montevidensis 22 (Lpm) were generated using a combination of MiSeq (Illumina) and GridION (Oxford 23 Nanopore) sequencing technologies. These plastomes were then compared to the published 24 Ludwigia octovalvis (Lo) plastid genome, which was re-annotated by the authors. We initially 25 sequenced and assembled the chloroplast (cp) genomes of *Lpm* and *Lgh* using a hybrid strategy 26 combining short and long reads sequences. We observed the existence of two Lgh haplotypes 27 and two potential Lpm haplotypes. Lgh, Lpm, and Lo plastomes were similar in terms of genome 28 size (around 159 Kb), gene number, structure, and inverted repeat (IR) boundaries, comparable 29 to other species in the Myrtales order. A total of 45 to 65 SSRs (simple sequence repeats), were 30 detected, depending on the species, with the majority consisting solely of A and T, which is 31 common among angiosperms. Four chloroplast genes (matK, accD, ycf2 and ccsA) were found 32 under positive selection pressure, which is commonly associated with plant development, and 33 especially in aquatic plants such as Lgh, and Lpm. Our hybrid sequencing approach revealed the presence of two Lgh plastome haplotypes which will help to advance phylogenetic and 34 35 evolutionary studies, not only specifically for Ludwigia, but also the Onagraceae family and Myrtales order. To enhance the robustness of our findings, a larger dataset of chloroplast 36 37 genomes would be beneficial.

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#### 39 Keywords

Water primrose, *Ludwigia* sp., Onagraceae, chloroplast genome, long and short reads, hybrid
assembly, haplotype

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#### 45 Introduction

46 The Onagraceae family belongs to the order Myrtales which includes approximately 657

47 species of herbs, shrubs, and trees across 17 genera grouped into two subfamilies: subfam.

48 Ludwigioideae W. L. Wagner and Hoch, which only has one genus (*Ludwigia* L.), and subfam.

49 Onagroideae which contains six tribes and 21 genera [1]. Ludwigia L. is composed of 83

50 species[2][3]. The current classification for *Ludwigia* L., which are composed of several hybrid

51 and/or polyploid species, lists 23 sections. A recent molecular analysis is clarified and

52 supported several major relationships in the genus but has challenged the complex sectional

53 classification of *Ludwigia* L.[4].

54 The diploid species Ludwigia peploides (Kunth) Raven subsp. montevidensis (Spreng.) [5] 55 (named here Lpm) (2n=16), and the decaploid species, Ludwigia grandiflora (Michx) Greuter 56 & Burdet subsp. hexapetala (Hook. & Arn) Nesom & Kartesz (named here Lgh) (2n=80), 57 reproduce essentially by clonal propagation, which suggests that there is a low genetic diversity 58 within the species [6]. Lgh and Lpm are native to South America and are considered as one of 59 the most aggressive aquatic invasive plants [7]. Largely distributed in aquatic environments in 60 North America and in Europe [8], both species have been found to degrade major watersheds 61 as well as aquatic and riparian ecosystems [9] leading ecological and economical damages. In 62 France, both species occupied aquatic habitats, such as static or slow-flowing waters, riversides, 63 and have recently been observed in wet meadows [10]. The transition from an aquatic to a 64 terrestrial habitat has led to the emergence of two Lgh morphotypes [11]. The appearance of 65 metabolic and morphological adaptations could explain the ability to acclimatize to terrestrial conditions, and this phenotypic plasticity involves various genomic and epigenetic 66 67 modifications [12].

68 Adequate genomic resources are necessary in order to be identify the genes and metabolic 69 pathways involved in the adaptation process leading to plant invasion [13] with genomic 70 information making it possible to predict and control invasiveness [14]. However, even though 71 the number of terrestrial plant genomes has increased considerably over the last 20 years, only 72 a small fraction (~ 0.16%) have been sequenced, with some clades being significantly more 73 represented than others [15]. Thus, for the Onagraceae family (which includes *Ludwigia* sp.), 74 only a handful of chloroplast sequences (plastomes) are available, and the complete genome 75 has not yet been sequenced. If Lpm is a diploid species (2n=2x=16) with a relatively small 76 genome size (262 Mb), Lgh is a decaploid species (2n=10x=80) with a large size genome of 77 1419 Mb [16]. Obtaining a reference genome for these two non-model species without having 78 a genome close to the Ludwigia species is challenging and development of plastome and/or

79 mitogenome will be a first step to generate genomic resource. As of April 2023, there are 10,712 80 reference plastomes listed on GenBank (Release 255: April 15 2023), with the vast majority 81 (10,392 genomes) belonging to Viridiplantae (green plants). However, in release 255, the 82 number of plastomes available for the Onagraceae family is limited, with only 36 plastomes 83 currently listed. Among these, 15 plastomes are from the tribe Epilobieae, with 11 in the 84 Epilobium genus and 4 in the Chamaenerion genus. Additionally, there are 23 plastomes from 85 the tribe Onagreae, with 17 in the Oenothera genus, 5 in the Circaea genus, and only one in the 86 Ludwigia genus. The Ludwigia octovalvis chloroplast genome was released in 2016 as a unique 87 haplotype of approximately 159 kb [17]. L. octovalvis belongs to sect. Macrocarpon (Micheli) 88 H.Hara while Lpm and Lgh belong to Jussieae section [18][19]. Generally, the inheritance of 89 chloroplast genomes is considered to be maternal in angiosperms. However, biparentally 90 inherited chloroplast genomes could potentially exist in approximately 20% of angiosperm 91 species [20][21]. Both maternal and biparental inheritance are described in the Onagraceae 92 family. In tribe Onagreae, Oenothera subsect. Oenothera are known to have biparental plastid 93 inheritance [22][23]. In tribe Epilobieae, biparental plastid inheritance was also reported in 94 Epilobium L. with mainly maternal transmission, and very low proportions of paternally 95 transmitted chloroplasts [24].

96 The chloroplast is the symbolic organelle of plants and plays a fundamental role in 97 photosynthesis. Chloroplasts evolved from cyanobacteria through endosymbiosis and thereby 98 inherited components of photosynthesis reactions (photosystems, electron transfer and ATP 99 synthase) and gene expression systems (transcription and translation)[25]. In general, 100 chloroplast genomes (plastomes) are highly conserved in size, structure, and genetic content. 101 They are rather small (120-170 kb,[26]), with a quadripartite structure comprising two long 102 identical inverted repeats (IR, 10-30 kb) separated by large and a small single copy regions 103 (LSC and SSC, respectively). They are also rich in genes, with around 100 unique genes 104 encoding key proteins involved in photosynthesis, and a comprehensive set of ribosomal RNAs 105 (rRNAs) and transfer RNAs (tRNAs)[27]. Plastomes are generally circular but linear shapes 106 also exist [28]. Chloroplast DNA usually represents 5-20% of total DNA extracted from young 107 leaves and therefore low-coverage whole genome sequencing can generate enough data to 108 assemble an entire chloroplast genome [29].

109 If we refer to their GenBank records, more than 95% of these plastomes were sequenced by 110 so-called short read techniques (mostly Illumina). However, in most seed plants, the plastid 111 genome exhibits two large inverted repeat regions (60 to 335 kb,[29]), which are longer than 112 the short read lengths (< 300 bp). This leads to incomplete or approximate assemblies [30]. 113 Recent long-read sequencing (> 1000 bp) provides compelling evidence that terrestrial plant 114 plastomes exhibit two structural haplotypes. These haplotypes are present in equal proportions 115 and differ in their inverted repeat (IR) orientation [31]. This shows the importance of using the 116 so-called third generation sequence (TGS, PacBio or Nanopore) to correctly assemble the IRs 117 of chloroplasts and to identify any different structural haplotypes. The current problem with 118 PacBio or Nanopore long read sequencing is the higher error rate compared to short read 119 technology [32][33][34]. Thus, a hybrid strategy which combines long reads (to access the 120 genomic structure) and short reads (to correct sequencing errors) could be effective [30][35].

Here, we report the newly sequenced complete plastid genomes of *Ludwigia grandiflora* subps. *hexapetala* (*Lgh*) and *Ludwigia peploides* subsp *montevidensis* (*Lpm*), using a combination of different sequencing technologies, as well as a re-annotated comparative genomic analysis of the published *Ludwigia octovalvis* (*Lo*) plastid. The main objectives of this study are (1) to assemble and annotate the plastomes of two new species of *Ludwigia* sp., (2) to reveal the divergent sequence hotspots of the plastomes in this genus and in the Onagraceae (3) to identify the genes under positive selection.

128 To achieve this, we utilized long read sequencing data from Oxford Nanopore and short read 129 sequencing data from Illumina to assemble the Lgh plastomes and compared these assemblies 130 with those obtained solely from long reads of Lpm. We also compared both plastomes to the 131 published plastome of Lo. Our findings demonstrated the value of de novo assembly in reducing 132 assembly errors and enabling accurate reconstruction of full heteroplasmy. We also evaluated 133 the performance of a variety of software for sequence assembly and correction in order to define 134 a workflow that will be used in the future to assemble Ludwigia sp. mitochlondrial and nuclear 135 genomes. Finally, the three new Ludwigia plastomes generated by our study make it possible 136 to extend the phylogenetic study of the Onagraceae family and to compare it with previously 137 published analyses [4][36][37].

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#### 139 Material and Methods

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## Plant sampling and experimental design

141 The original plant materials were collected in June of 2018 near to Nantes (France) and 142 formal identified by D. Barloy. *L. grandiflora* subsp. *hexapetala (Lgh)* plants were taken from 143 the Mazerolles swamps (N47 23.260, W1 28.206), and *L. peploides* subsp. *montevidensis (Lpm)* 144 plants from La Musse (N 47.240926, W -1.788688)). Plants were cultivated in a growth 145 chamber in a mixture of  $\frac{1}{3}$  soil,  $\frac{1}{3}$  sand,  $\frac{1}{3}$  loam with flush water level, at 22°C and a 16 h/8 h (light/dark) cycle. A single stem of 10 cm for each species was used for vegetative propagation in order to avoid potential genetic diversity. *De novo* shoots, taken three centimeters from the apex, were sampled for each species. Samples for gDNA extraction were pooled and immediately snap-frozen in liquid nitrogen, then lyophilized over 48 h using a Cosmos 20K freeze-dryer (Cryotec, Saint-Gély-du-Fesc, France) and stored at room temperature. All the plants were destroyed after being used as required by French authorities for invasive plants (article 3, prefectorial decree n°2018/SEE/2423).

Due to high polysaccharide content and polyphenols in *Lpm* and *Lgh* tissues and as no standard kit provided good DNA quality for sequencing, genomic DNA extraction was carried out using a modified version of the protocol proposed by Panova et al in 2016, with three purification steps [38].

157 40 mg of lyophilized buds were ground at 30 Hz for 60 s (Retsch MM200 mixer mill, 158 FISHER). The ground tissues were lysed with 1 ml CF lysis buffer (MACHEREY-NAGEL) 159 supplemented with 20 µl RNase and incubated for 1 h at 65°C under agitation. 20 µl proteinase 160 K was then added before another incubation for 1 h at 65°C under agitation. To avoid breaking 161 the DNA during pipetting, the extracted DNA was recovered using a Phase-lock gel tube as 162 described in Belser [39]. The extracts were transferred to 2 ml tubes containing phase-lock gel, 163 and an equal volume of PCIA (Phenol, Chloroform, Isoamyl Alcohol; 25:24:1) was added. 164 After shaking for 5 min, tubes were centrifuged at 11000 g for 20 min. The aqueous phase was 165 transferred into a new tube containing phase-lock gel and extraction with PCIA was repeated. 166 DNA was then precipitated after addition of an equal volume of binding buffer C4 167 (MACHEREY-NAGEL) and 99% ethanol overnight at 4°C or 1 h in ice then centrifuged at 800 168 rpm for 10 min. After removal of the supernatant, 1 ml of CQW buffer was added then the 169 pellet of DNA was re-suspended. Next, DNA purification was carried out by adding a 2 ml 170 mixture of wash buffer PW2 (MACHEREY-NAGEL), wash buffer B5 (MACHEREY-171 NAGEL), and ethanol at 99% in equal volumes, followed by centrifugation at 800 rpm for 10 172 min. This DNA purification step was carried out twice. Finally, the DNA pellet was dried in 173 the oven at 70°C for 30 min then re-suspended in 100 µl elution buffer BE (MACHEREY-174 NAGEL) (5 mM Tris solution, pH 8.5) after 10 min incubation at 65°C under agitation.

A second purification step was performed using a PCR product extraction from gel agarose kit from Macherey-Nagel (MN) NucleoSpin® Gel and PCR Clean-up kit and restarting the above protocol from the step with the addition of CQW buffer then PW2 buffer. The third purification step consisted of DNA purification using a Macherey-Nagel (MN)
NucleoMag kit for clean-up and size selection. Finally, the DNA was resuspended after a 5 min
incubation at 65°C in 5 mM TRIS at pH 8.5.

- 181 The quantity and quality of the gDNA was verified using a NanoDrop spectrometer, 182 electrophoresis on agarose gel and ethidium bromide staining under UV light and Fragment 183 Analyzer (Agilent Technologies) of the University of Rennes1.
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## Library preparation and sequencing

185 MiSeq (Illumina) and GridION (Oxford Nanopore Technologies, referred to here as 186 ONT) sequencing were performed at the PGTB (doi:10.15454/1.5572396583599417E12). Lgh 187 and Lpm genomic DNA were re-purified using homemade SPRI beads (1.8X ratio). Lgh has a 188 large genome size of 1419 Mb, 5-fold larger than Lpm genome 262 Mb [16]. SR (Illumina, one 189 run) and LR (Oxford Nanopore, three runs) sequencing were therefore carried out for Lgh and 190 only LR sequencing for Lpm (one run). For Illumina sequencing, 200 ng of Lgh DNA was used 191 according to the QIAseq FX DNA Library Kit protocol (Qiagen). The final library was checked 192 on TapeStation D5000 screentape (Agilent Technologies) and quantified using a QIAseq 193 Library Quant Assay Kit (Qiagen). The pool was sequenced on an Illumina MiSeq using V3 194 chemistry and 600 cycles (2x300bp). For ONT sequencing, around 8 µg of Lgh and Lpm DNA 195 were size selected using a Circulomics SRE kit (according to the manufacturer's instructions) 196 before library preparation using a SQK-LSK109 ligation sequencing kit following ONT 197 recommendations. Basecalling in High Accuracy - Guppy version: 4.0.11 (MinKNOW 198 GridION release 20.06.9) was performed for the 48 h of sequencing. Long reads (LR) and short 199 reads (SR) were available for Lgh and only LR for Lpm.

#### 200

## Chloroplast assemblies

201 Quality controls and preprocessing of sequences were conducted using Guppy v4.0.14 for 202 long reads (via Oxford Nanopore Technology Client access) and fastp v0.20.0 [40] for short 203 reads, using Q15, since increasing the Phred quality to 20 or higher has no effect on the number 204 of sequences retained (66%). A preliminary draft assembly was performed using Lgh short-205 reads (SR, 2\*23,067,490 reads) with GetOrganelle v1.7.0 [41] and NOVOPlasty v4.2.1 [42], 206 and chloroplastic short and long reads were extracted by mapping against this draft genome. 207 Chloroplastic short reads were then *de novo* assemble using Velvet (version 1.2.10) [43], 208 ABySS (version 2.1.5 [44][45]), MEGAHIT (1.1.2,[46]), and SPAdes (version 3.15.4,[47]), 209 without and with prior error correction. The best k-mer parameters were tested using kmergenie 210 [48] and k=99 was found to be optimal. For ONT reads, Lgh (550,516 reads) and Lpm (68,907 211 reads) reads were self-corrected using CANU 1.8 [49] or SR-corrected using Ratatosk [50] and 212 *de novo* assembly using CANU [49] and FLYE 2.8.2 [51] run with the option --meta and -

213 plasmids. For all these assemblers, unless otherwise specified, we used the default parameters.

## 214

## Post plastome assembly validation

215 As we used many assemblers and different strategies, we produced multiple contigs that 216 needed to be analyzed and filtered in order to retain only the most robust plastomes. For that, 217 all assemblies were evaluated using the QUality ASsessment Tool (QUAST) for quality 218 assessment [52] and visualized using BANDAGE [53], both using default parameters. 219 BANDAGE compatible graphs (.gfa format) were created with the megahit toolkit for 220 MEGAHIT [46] and with gfatools for ABySS [45]. Overlaps between fragments were manually checked and ambiguous "IUPAC or N" nucleotides were also biocured with Illumina reads 221 222 when available.

## 223 Chloroplast genome annotation

224 Plastomes were annotated via the GeSeq [54] using ARAGORN and tRNAscan SE to 225 predict tRNAs and rRNAs and tRNAscan SE to predict tRNAs and rRNAs and via Chloe 226 prediction site [55]. The previously reported Lo chloroplast genome was also similarly re-227 annotated to facilitate genomic comparisons. Gene boundaries, alternative splice isoforms, 228 pseudogenes and gene names and functions were manually checked and biocurated using 229 Geneious (v.10). Finally, plastomes were represented using OrganellarGenomeDRAW 230 (OGDRAW)[56]. These genomes were submitted to GenBank at the National Center of 231 Biotechnology Information (NCBI) with specific accession numbers (for Lgh haplotype 1, 232 (LGH1) OR166254 and Lgh haplotype 2, (LGH2) OR166255; for Lpm haplotype, (LPM) 233 OR166256) using annotation tables generated through GB2sequin [57].

#### 234

## SSRs and Repeat Sequences Analysis

235 Simple Sequence Repeats (SSRs) were analyzed through the MISA web server [58], with 236 parameters set to 10, 5, 4, 3, 3, and 3 for mono-, di-, tri-, tetra-, penta-, and hexa-nucleotides, 237 respectively. Direct, reverse and palindromic repeats were identified using RepEx [59]. 238 Parameters used were: for inverted repeats (min size 15 nt, spacer = local, class = exact); for 239 palindromes (min size 20 nt); for direct repeats (minimum size 30 nt, minimum repeat similarity 240 97%). Tandem repeats were identified using Tandem Repeats Finder[60], with parameters set 241 to two for the alignment parameter match and seven for mismatches and indels. The IRa region 242 was removed for all these analyses to avoid over representation of the repeats.

243 <u>Comparative chloroplast genomic analyses</u>

*Lgh* and *Lpm* plastomes were compared with the reannotated and biocurated *Lo* plastome
using mVISTA program [61], with the LAGAN alignment algorithm [62] and a cut-off of 70%
identity.

247 Nucleotide diversity (Pi) was analyzed using the software DnaSP v.6.12.01 [63] [64] with 248 step size set to 200 bp and window length to 300 bp. IRscope [65] was used for the analyses of 249 inverted repeat (IR) region contraction and expansion at the junctions of chloroplast genomes. 250 To assess the impact of environmental pressures on the evolution of these three Ludwigia 251 species, we calculated the nonsynonymous (Ka) and synonymous (Ks) substitutions and their 252 ratios ( $\omega = Ks/Ks$ ) using TB tools [66] to measure the selective pressure. Genes with  $\omega < 1, \omega =$ 253 1, and  $1 < \omega$  were considered to be under purifying selection (negative selection), neutral 254 selection, and positive selection, respectively.

## 255 Phylogenetic analysis of Ludwigia based on MatK sequences

We performed a phylogenetic analysis on the Ludwigia genus using the MatK, only protein 256 257 coding barcode available for a large number of Ludwigia species. All MatK amino acid 258 sequences were aligned with the FFT-NS-2 (Fast Fourier Transform-based Narrow Search) 259 algorithm and BLOSUM62 scoring matrix using MAFFT 7 [67]. The phylogenetic tree analysis 260 was conducted using the rapid hill-climbing algorithm (command line: -f d) in RAxML 8.2.11 261 [68], with GAMMA JTT (Jones-Taylor-Thornton) protein model. Node support was assessed 262 through fast bootstrapping (-f a) with 1,000 non-parametric bootstrap pseudo-replicates. 263 Circaea MatK were selected as outgroup, and all accession numbers are indicated on the 264 phylogenetic tree labels.

## 265 Graphic representation

Statistical analyses were performed using R software in RStudio integrated development
environment (R Core Team, 2015, RStudio: Integrated Development for R. RStudio, Inc.,
Boston, MA, <u>http://www.rstudio.com/</u>). Figures were realized using ggplot2, ggpubr, tidyverse,
dplyr, gridExtra, reshape2, and viridis packages. SNPs were represented using trackViewer [69]
and genes represented using gggenes packages.

271

#### 272 Results

## 273 Plastome short read assembly

The chloroplastic fraction of *Lgh* short reads (SR) was extracted by mapping against the two draft haplotypes generated by GetOrganelle, which differ only by a "flip-flop" of the SSC region (Figure 1). Since the assembly by NOVOplasty did not provide any additional information compared to GetOrganelle, it was not retained. This subset (1,360,507 reads) was assembled 278 using ABySS, Velvet, MEGAHIT and SPAdes in order to identify the best assembler for this 279 plant model. As shown in Figure 2, both the number and size of contigs depend greatly on the 280 algorithms used and the correction step. The effect of prior read correction is notable for 281 MEGAHIT and Velvet, especially concerning the increase in the size of the large alignment 282 (Add. Figure 1A), loss of misassemblies, and reduction of the number of mismatches (Add. 283 Figure 1B). Investigating results via BANDAGE (Add. Figure 2), we observed that ABySS and 284 SPAdes suggest the tripartite structure with the long single-copy (LSC) region as the larger 285 circle in the graph (blue), joined to the small single-copy region (green) by one copy of the 286 inverted repeats (IRs, red), both IRs being collapsed in a segment of approximately twice the 287 coverage. For Velvet and MEGAHIT, graphs confirm the significant fragmentation of the 288 assemblies, which is improved by prior correction of the reads.

In conclusion, none of the short-read assemblers tested in our study produced a complete plastome. The best result was achieved by SPAdes using corrected short reads (mean coverage 1900 X) to assemble a plastome consisting of three contigs: 90,272 bp (corresponding to LSC),

292 19,788 bp (corresponding to SSC), and 24,762 bp (corresponding to one of the two copies of

293 the IR).

## 294 Plastome long read assembly

295 Chloroplast fractions of Lgh long reads (28,882 reads) were assembled using CANU or 296 FLYE. With raw data, CANU generates a unique contig (NGA50 112648) corresponding to 297 haplotype 2, whereas FLYE makes two contigs (NGA50 133687) that reconstruct haplotype 1. 298 Self-corrected LR leads to fragmentation into two (CANU) or three (FLYE) contigs which both 299 reconstruct haplotype 1, with an large gap corresponding to one of the IR copies for CANU. 300 Finally, SR-correction by RATATOSK allows CANU to assemble two redundant contigs 301 reproducing haplotype 2 while FLYE makes two contigs corresponding to haplotype 1 (Add. 302 Figure 3A). In conclusion, the two Lgh haplotypes were reconstructed (average coverage 700X) 303 and the most complete and accurate hybrid assemblies (99.94% accuracy, Additional Figure 304 3B) were submitted to GenBank.

Unfortunately, due to the absence of short read data, we could only perform self-corrected long read assembly for *Lpm* using CANU. We also compared CANU and FLYE assembler efficiency, and found that assembly using CANU produces 13 contigs whereas FLYE produces 12 contigs. In both cases, only three contigs are required to reconstitute a complete cpDNA assembly (no gap, no N), with an SSC region oriented like those of the *Lgh* haplotype 2 and the *Lo* plastome. Although it is more than likely that these two SSC region orientations also exist for *Lpm*, the low number of nanopore sequences generated (68907 reads) and absence of 312 Illumina short reads prevented us from demonstrating the existence of both haplotypes. As a

result, only the "haplotype 2" generated sequence was deposited to Genbank.

- 314 Annotation and comparison of *Ludwigia* plastomes
- 315 1.

#### 1. General Variations

316 Plastomes of the three species of Ludwigia sp., Lgh, Lpm and Lo, are circular double-317 stranded DNA molecules (Figure 3) which are all (as shown in Table 1) approximately the same 318 size: Lo is 159,396 bp long, making it the smallest, while Lgh is the largest with 159,584 bp, 319 and Lpm is intermediate at 159,537 bp. The overall GC content is almost the same for the three 320 species (37.4% for Lo, 37.3 % for Lgh and Lpm) and the GC contents of the IR regions are 321 higher than those of the LSC and SSC regions (approximately 43.5 % compared to 35% and 322 ca.32% respectively). Between the three species, the lengths of the total chloroplasts, LSC, 323 SSC, and IR are broadly similar (approximately 90.2 kb for LSC, 19.8 kb for SSC and 24.8 kb 324 for IB, see details Table 1) and the three plastomes are perfectly syntenic if we orient the SSC 325 fragments the same way.

326 All three *Ludwigia* sp. plastomes contain the same number of functional genes (134 in total) 327 encoding 85 proteins (embracing 7 duplicated in the IR region: ndhB, rpl2, rpl23, rps7, rps12, 328 ycf2, ycf15), 37 tRNAs (including trnK-UUU which contains matK), and 8 rRNAs (16S, 23S, 329 5S, and 4.5S as duplicated sets in the IR). Among these genes, 18 contain introns, of which six 330 are tRNAs (Table 2). Only the rps12 gene is a trans-spliced gene. A total of 46 genes are 331 involved in photosynthesis, and 71 genes related to transcription and translation, including a 332 bacterial-like RNA polymerase and 70S ribosome, as well as a full set of transfer RNAs 333 (tRNAs) and ribosomal RNAs (rRNAs). Six other protein-coding genes are involved in 334 essential functions, such as *accD*, which encodes the  $\beta$ -carboxyl transferase subunit of acetyl-335 CoA carboxylase, an important enzyme for fatty acid synthesis; matK encodes for maturase K, 336 which is involved in the splicing of group II introns; *cemA*, a protein located in the membrane 337 envelope of the chloroplast is involved in the extrusion of protons and thereby indirectly allows 338 the absorption of inorganic CO2 in the plastids; *clp*P1 which is involved in proteolysis, and; 339 *ycf1*, *ycf2*, two ATPases members of the TIC translocon. Finally, a highly pseudogenized *ycf15* 340 locus was annotated in the IR even though premature stop codons indicate loss of functionality.

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#### 2. Segments Contractions/Expansion

The junctions between the different chloroplast segments were compared between three *Ludwigia* sp. (*Lpm*, *Lgh* and *Lo*), and we found that the overall resemblance of *Ludwigia* sp. plastomes was confirmed at all junctions (Figure 4A). In all three genomes, *rpl22*, *rps19*, *and rpl2* were located around the LSC/IRb border, and *rpl2*, *trn*H, and *psb*A were located at the 346 IRa/LSC edge. The JSB (junction between IRb and SSC) is either located in the *ndh*F gene or 347 the *ycf1* gene depending on the orientation of the SSC region (Figure 4B). The *ycf1* gene was 348 initially annotated as a 1139 nt pseudogene that we biocurate as a larger gene (5302 nt) with a 349 frameshift due to a base deletion, compared to Lg and Lo which both carry a complete *ycf1* 350 gene.

351 If we compare Ludwigia sp. chloroplastic LSC/SCC/IR junctions (via IRscope) with 352 representative Onagraceae plastomes of Chamaenerion sp. conspersum (MZ353638) and sp. 353 angustifolium (NC 052848), Circaea sp. cordata (NC 060876) and sp. alpina (NC 061010), 354 Epilobium amurense (NC 061015) and Oenothera villosa subsp. strigosa (NC 061365) and 355 Oenothera lindheimeri (MW538951) (Figure 5), we can observe that the gene positions at the 356 JLB (junction of LSC/IRb) and JLA (junction of IRa/LSC) boundary regions are well-preserved 357 throughout the entire family, whereas those at the JSB and JSA regions differ. Concerning JSB 358 (junction of IRb/SSC), in the five Onagraceae genera studied, ndhF is duplicated, with the 359 exception of Circaea sp. and Ludwigia sp. For Oenothera villosa, the first copy of ndhF, which 360 is located in the IRb, overlaps the JSB border, whereas for *Oenothera lindheimeri*, *Epibolium* 361 amurense and Chamaenerion sp., ndhF is only located in inverted repeats. Only Circaea sp. and 362 Ludwigia sp. have a unique copy of this locus, and it is found in the SSC segment (Figure 5). 363 At the JSA border (junction of SSC/Ira), in Circaea sp., the ycfl gene crosses the IRa/SSC 364 boundary and extends into the IRa region.

When comparing the respective sizes of chloroplast fragments (IR/SSC/LSC) in Onagraceae, it can be observed that *Ludwigia* species exhibit expansions in the SSC and LSC regions which are not compensated by significant contractions in the IR regions. This is likely due to the relocation of the *ndh*F in the SSC region and *rps19* in the LSC region. Additionally, there may be significant size variations in the intergenic region between *trnI* and *ycf2*, as well as the intergenic segment containing the *ycf15* pseudogene (Add. Figure 4).

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## 3. Repeats and SSRs analysis

372 In this study, we analyzed the nature and distribution of single sequence repeats (SSR), as 373 their polymorphism is an interesting indicator in phylogenetic analyses. A total of 65 (Lgh), 48 (Lpm) and 45 (Lo) SSRs were detected, the majority being single nucleotide repeats (38-21), 374 375 followed by tetranucleotides (12–10) and then di-, tri- and penta-nucleotides (Add. Figure 5A). 376 Mononucleotide SSRs are exclusively composed of A and T, indicating a bias towards the use 377 of the A/T bases, which is confirmed for all SSRs (Add. Figure 5B). In addition, the SSRs are 378 mainly distributed in the LSC region for the three species, which is probably biased by the fact 379 that LSC is the longest segment of the plastome (Add. Figure 5C). The analysis of SRR locations revealed that most were distributed in non-coding regions (intergenic regions andintrons, Add. Figure 5D).

The chloroplast genomes of the three *Ludwigia* species were also screened for long repeat sequences. They were counted in a non-redundant way (if smaller repetitions were included in large repeats, only the large ones were considered). Four types of repeats (tandem, palindromic inverted and direct) were surveyed in the three *Ludwigia* sp. plastomes. No inverted repeats were detected with the criteria used.

387 For the three other types of repeats, here are their distributions:

388 *Tandem repeats* (Table 3A): Perfect tandem repeats (TRs) with more than 15 bp were 389 examined. Twenty-two loci were identified in the three Ludwigia sp. plastomes (Lgh, Lpm, Lo), 390 heterogeneously distributed as shown in Table 3A: 13 loci (plus one imperfect) in Lo, nine loci 391 (plus one imperfect) in Lgh and seven loci (plus two imperfect) in Lpm. It can therefore be seen 392 that the TR distributions (occurrence and location) are specific to each plastome, since only 393 four pairs are common to the three species. Thus, nine TRs are unique to Lo, three to Lpm and 394 three to Lgh. Two pairs are common to Lgh and Lpm and one is common to Lo and Lgh. TRs 395 are mainly intergenic or intronic but are detected in two genes (accD and ycfl). These genes 396 have accelerated substitution rates, although this does not generate a large difference in their 397 lengths. This point will be developed later in this article.

*Direct repeats* (Table 3B): There are few direct (non-tandem) repeats (DRs) in the chloroplast genomes of *Ludwigia* sp. A single direct repeat of 41 nt is common to the three species, at 2 kb intervals, in *psaB* and *psaA* genes. This DR corresponds to an amino acid repeat [WLTDIAHHHLAIA] which corresponds to a region predicted as transmembrane. We then observe three direct repeats conserved in *Lpm* and *Lgh* in *ycf1*, *accD* and *clp*P1 respectively, two unique DRs in *Lo* (in the *accD* gene and *rps12-clp*P1 intergene) and one in *Lgh* (in the *clp*P1 intron 1 and *clp*P1 intron 2).

405 *Palindromes* (Table 3C): Palindromic repeats make up the majority of long repetitions, 406 with the numbers of perfect repeats varying from 19, 24 and 26 in *Lo*, *Lgh* and *Lpm*, 407 respectively, and the number of quasi-palindromes (1 mutation) varying between 8, 3 and 6. 408 They are mainly found in the intronic and intergenic regions, with the exception of six genic 409 locations in *psbD*, *ndh*K, *ccs*A and *rpl22*, and two palindromic sequences in *ycf2*. These gene 410 palindromic repeats do not seem to cause genetic polymorphism in *Ludwigia* and can be 411 considered as silent.

Thirteen palindromes are common to the three species (including 2 with co-variations in *Lo*). 13 others present in *Lpm* and *Lgh* correspond to quasi-palindromes (QPs) in *Lo* due to

414 mutated bases, and conversely, three *Lo* perfect palidromes are mutated in *Lpm* and *Lgh*. 415 Finally, only five palindromes are species specific. Two in particular are located in the 416 hypervariable intergenic spacer ndhF-rpl32, and are absent in *Lo* due to a large deletion of 160 417 nt.

418

## 4. Repeat distribution in LSC, SSC and IR segments

In the IRa/IRb regions, repeats are only identified in the first 9 kb region between rpl2 and ycf2: a tandem repeat in the Lpm rpl2 intron, and a tetranucleotide repeat, [TATC]\*3, located in the ycf2 gene in the 3 species. In ycf2 we also found 1 common palindrome (16 nt), a single palindrome in Lo (20 nt, absent following an A:G mutation in the 2 other species), as well as a shared tandem repeat (24 nt), and an additional 15 nt tandem repeat in Lo which adds 4 amino acids to protein sequence.

425 In the SSC region, the repeats are almost all located in the intergenic and/or intronic 426 regions, with a hotspot between *ndh*F and *ccs*A. There is also a shared microsatellite in *ndh*F, 427 and a palidrome (16 nt) in ccsA which is absent in Lo (due to an A:C mutation), resulting in a 428 synonymous mutation (from isoleucine to leucine). We also observed multiple and various 429 repeats in the ycfl gene: 3 common poly-A repeats (from 10 to 13 nt), 3 species-specific 430 microsatellites (ATAG)\*3 and (ACCA)\*4 in Lgh and (CAAC)\*3 in Lo, as well as two direct 431 repeats of 32 nt (37 nt spacing), which were absent from Lo due to a G:T SNP. Two tandem 432 repeats were also observed in Lo and Lgh. Neither of these repeats are at the origin of the 433 frameshift causing the pseudogenization of *ycf1* in *Lo*, this latter being due to a single deletion 434 of an A at position 3444 of the gene.

435 Finally, in the LSC region, the longest segment, which consequently contains the maximum 436 number of repeats, we still observed a preferential localization in the intergenic and intronic 437 regions since only genes atpA, rpoC2, rpoB, psbD, psbA, psbB, ndhK and clpP1 contain either 438 mononucleotic repeats (poly A and T), palindromes, or microsatellites (most often common to 439 the three species and without affecting the sequences of the proteins produced). As mentioned 440 earlier, the only exception is the accD gene, which contains several direct and tandem repeats 441 in Lgh and Lpm, corresponding to a region of 174 nt (58 amino acids) missing in Lo and, 442 conversely, a direct repeat of 40 nucleotides, in a region of 147 nt (49 aa), which is present in 443 Lo and missing in the other two species. These tandem repeats lead to the presence of four 444 copies of 9 amino acids [DESENSNEE] in Lgh and Lpm, two of which form a larger duplication 445 of 17 aa [FLSDSDIDDESENSNEE]. Similarly, the TRs present only in Lo generate two perfect 446 9 amino acid repeats [EELSEDGEE], included in two longer degenerate repeats of 27 nt (Add. 447 Figure 6). It should be noted that though these TRs do not disturb the open reading phases, it is 448 still possible for them to form an intron which is not translated. Different functional studies will 449 be necessary to clarify this point. The presence of polymorphisms of the accD gene between 450 Lo and the two species (Lpm, Lgh) is interesting because accD, that encodes a subunit of acetyl-451 CoA carboxylase (EC 6.4.1.2). This enzyme is essential in fatty acid synthesis and also 452 catalyzes the synthesis of malonyl-CoA, which is necessary for the growth of dicots, plant 453 fitness and leaf longevity, and is involved in the adaptation to specific ecological niches [70]. 454 Large accD expansions due to TRs have also been described in other plants such as Medicago 455 [71] and Cupressophytes [72]. Some authors have suggested that these inserted repeats are not 456 important for acetyl-CoA carboxylase activity as the reading frame is always preserved, and 457 they assume that these repeats must have a regulatory role [73].

458

#### 5. Sequence Divergence Analysis and Polymorphic Loci Identification

Determination of divergent regions by MVista, using *Lo* as a reference, confirmed that the three *Ludwigia* sp. plastomes are well preserved if the SSC segment is oriented in the same way (Add. Figure 7). Sliding window analysis (Figure 6) indicated variations in definite coding regions, notably *clpP*, *accD*, *ndh5*, *ycf1* with high Pi values, and to a lesser extent, *rps16*, *mat*K, *ndh*K, *pet*A, *ccs*A and four tRNAs (*trn*H, *trn*D, *trn*T and *trn*N). These polymorphic *loci* could be suitable for inferring genetic diversities in *Ludwigia* sp.

465 A comparative analysis of the sizes of protein coding genes sizes also shows that the *rps11* 466 gene initially annotated in Lo is shorter than those which have been newly annotated in Lgh and 467 Lpm (345 bp instead of 417 bp). Comparative analysis by BLAST shows that it is the long form 468 which is annotated in other Myrtales, and the observation of the locus in Lo shows a frameshift 469 mutation (deletion of a nucleotide in position 311). Functional analysis would be necessary to 470 check whether the *rps11* frameshift mutation produces shorter proteins that have lost their 471 function. And only obtaining the complete genome will verify whether copies of some of these 472 genes have been transferred to mitochondrial or nuclear genomes. Such rps11 horizontal 473 transfers have been reported for this gene in the mitochondrial genomes of various plant 474 families[74]. This also applies to *ycf1*, found as a pseudogene in *Lo* (as specified previously), 475 although it is not known if this reflects a gene transfer or a complete loss of function [75][76]. 476 Moreover, there is a deletion of nine nucleotides in the 3' region of the rpl32 gene in Lgh and 477 Lpm, leading to a premature end of the translation and the deletion of the last 4 amino acids 478 [QRLD], which are replaced by a K. However, if we look carefully at the preserved region as 479 defined by the RPL32 domain (CHL00152, member of the superfamily CL09115), we see that 480 the later amino acids are not important for rpl32 function since they are not found in the 481 orthologs.

- 482Our results show that the Ka/Ks ratio is less than 1 for most genes (Figure 7). This indicates483adaptive pressures to maintain the protein sequence except for *mat*K (1.17 between *Lgh* and484*Lpm*), *acc*D (2.48 between *Lgh* and *Lo* and 2.16 between *Lpm* and *Lo*), *ycf2* (4.3 between both485*Lgh-Lp* and *Lo*) and *ccs*A (1.4 between both *Lgh-Lpm* and *Lo*), showing a positive selection for
- these genes, and a possible key role in the processes of the species' ecological adaptations. As
  we have already described the variability in the *acc*D sequence, we will focus on *ycf2*, *mat*K,
  and *ccs*A variations.
- 489 Concerning *ccs*A, the variations observed, although significant, concern only five amino 490 acids, and modifications do not seem to affect the C-type cytochrome synthase gene function.
- 491 Concerning *ycf2*, our analysis shows that this gene is highly polymorphic with 256 SNPs
  492 that provoke 10 deletions, 7 insertions, 21 conservative and 49 non-conservative substitutions
  493 in *Lo* (Add. Figure 8), compared to *Lgh* and *Lpm* (100 % identical). This gene has been shown
  494 as "variant" in other plant species such as *Helianthus tuberosus* [77].
- 495 The *mat*K gene has been used as a universal barcoding locus to enable species discrimination 496 of terrestrial plants [78], and is often, together with the *rbc*L gene, the only known genetic 497 resource for many plants. Thus, we propose a phylogenetic tree from *Ludwigia mat*K sequences 498 (Figure 8). It should however be noted that this tree contains only 149 amino acids common to 499 all the sequences (out of the 499 in the complete protein). As only three complete Ludwigia 500 plastomes are available at the time of our study, we cannot specify whether these barcodes are 501 faithful to the phylogenomic history of Ludwigia in the same way as the complete plastome. In 502 we can that *Lo* stands case, for this tree. see apart from the any 503 other Ludwigia sp., Lpm and Lgh, and that the L. grandiflora subsp. hexapetala belongs to the 504 same branch as the species L. ovalis (aquatic taxon used in aquariums [79]), L. stolonifera 505 (native to the Nile, found in a variety of habitats, from freshwater wetlands to brackish and 506 marine waters) [80] and L. adscendens (common weed of rice fields in Asia) [81]. Lpm is in a 507 sister branch, close to the L. grandiflora subsp. hexapetala, forming a phylogenetic group 508 corresponding to subsect Jussiaea (in green, Figure 8).
- 509

#### 510 **Discussion**

- 511 In the present study, we first sequenced and *de novo* assembled the chloroplast (cp) genomes
- 512 of Ludwigia peploides (Lpm) and Ludwigia grandiflora (Lgh), two species belonging to the
- 513 Onagraceae family. We employed a hybrid strategy and demonstrated the presence of two cp
- 514 haplotypes in *Lgh* and one haplotype in *Lpm*, although the presence of both haplotypes in *Lpm*
- 515 is likely. Furthermore, we compared these genomes with those of other species in the

516 Onagraceae family to expand our knowledge of genome organization and molecular evolution517 in these species.

518 Our findings demonstrate that the utilization of solely short reads has failed to produce complete 519 Ludwigia plastomes, likely due to challenges posed by long repeats and rearrangements. On the 520 other hand, relying solely on long reads resulted in a lower quality sequence due to insufficient 521 coverage and sequencing errors. After conducting our research, we discovered that, for Lgh 522 plastomes, hybrid assembly, which incorporates both long and short read sequences, resulted 523 in the most superior complete assemblies. This innovative approach capitalizes on the 524 advantages of both sequencing technologies, harnessing the accuracy of short read sequences 525 and the length of long read sequences. In the case of our study on Lgh plastome reconstruction, 526 hybrid assembly was the most complete and effective, similarly to studies on other chloroplasts, 527 such as those in *Eucalyptus* [82], *Falcataria* [83], *Carex* [84] or *Cypripedium* [85].

In our study, we were able to identify the presence of two haplotypes in *Lgh*, which is a first for *Ludwigia* (and more broadly within Onagraceae), as the plastome of *L. octovalvis* was only delivered in one haplotype [86].

531 Due to the unavailability of sequence data for Ludwigia octovalvis and the fact that we only 532 have long reads for Ludwigia peploides, none of which large enough to cover the SSC/IR 533 junctions, we are unable to conclusively identify the presence of these two forms in the 534 Ludwigia genus. However, we believe that they are likely to be present. Unfortunately, the 535 current representation of plastomes in GenBank primarily consists of short-read data, which 536 may result in an underrepresentation of this polymorphism. It is unfortunate that structural 537 heteroplasmy, which is expected to be widespread in angiosperms, has been overlooked. 538 Existence of two plastome haplotypes has been identified in the related order of Myrtales 539 (Eucalyptus sp.), in 58 species of Angiosperms, [87], Asparagales (Ophrys apifera orchid [88]), 540 Brassicales (Carica papava, Vasconcellea pubescens [89]), Solanales (Solanum tuberosum 541 [90]), Laurales (Avocado Persea americana [91]) and Rhamnaceae (Rhamnus crenata [92]). 542 However, the majority of reference plastomes in the current GenBank database (Release 260: 543 April 15, 2024) are described as a single haplotype, indicating an underrepresentation of 544 structural heteroplasmy in angiosperm chloroplasts. This underscores the importance of 545 sequencing techniques, as the database is predominantly composed of short-read data (98%),

which are less effective than long reads or hybrid assemblies at detecting flip-flop phenomenain the LSC region.

548 The chloroplast genome sizes for the three genera of Onagraceae subfam. Onagroideae varied 549 as follows: *Circaea* sp. ranged from 155,817 bp to 156,024 bp, *Chamaenerion* sp. ranged from 550 159,496 bp to 160,416 bp, and Epilobium sp. ranged from 160,748 bp to 161,144 bp [93]. Our 551 study revealed that the size of the complete chloroplast of Ludwigia (Onagraceae subfamily 552 Ludwigioideae) ranged from 159,369 bp to 159,584 bp, which is remarkably similar to other 553 Onagraceae plants (average length of 162,030 bp). Furthermore, Ludwigia plastome sizes are 554 consistent with the range observed in Myrtales (between 152,214 to 171,315 bp [94]). In the 555 same way, similar overall GC content was found in Ludwigia sp. (from 37.3 to 37.4%), Circaea 556 sp. (37.7 to 37.8%), Chamaenerion sp. and Epilobium sp. (38.1 to 38.2%, [93]) and more 557 generally for the order Myrtales (36.9–38.9%, with the average GC content being 37%,[94]). 558 Higher GC content of the IR regions (43.5%) found in *Ludwigia* sp. has already been shown in 559 the Myrtales order (39.7-43.5%) and in other families/orders such as Amaranthaceae (order 560 Caryophyllales [95]) or Lamiaceae (order Lamiales [96]), and is mainly due to the presence of

the four GC rich rRNA genes.

562 The complete chloroplast genomes of the three Ludwigia species encoded an identical set of 563 134 genes including 85 protein-coding genes, 37 tRNA genes and eight ribosomal RNAs, 564 consistent with gene content found in the Myrtales order, with a gene number varying from 123 565 to 133 genes with 77-81 protein-coding genes, 29-31 tRNA gene and four rRNA genes [94]. 566 Chloroplast genes have been selected during evolution due to their functional importance[97]. 567 In our current study, we made the noteworthy discovery that matK, accD, ycf2, and ccsA genes 568 were subjected to positive selection pressure. These genes have frequently been reported in 569 literature as being associated with positive selection, and are known to play crucial roles in 570 plant development conditions. Lgh and Lpm are known to thrive in aquatic environments, where 571 they grow alongside rooted emergent aquatic plants, with their leaves and stems partially 572 submerged during growth, as reported by Wagner et al. in 2007 [1]. Both species possess the 573 unique ability of vegetative reproduction, enabling them to establish themselves rapidly in 574 diverse habitats, including terrestrial habitats, as noted by Haury et al [98]. Additionally, Lo is 575 a wetland plant that typically grows in gullies and at the edges of ponds, as documented by 576 Wagner et al. in 2007 [1]. Given their ability to adapt to different habitats, these species may 577 have evolved specialized mechanisms to cope with various abiotic stresses, such as reduced 578 carbon and oxygen availability or limited access to light in submerged or emergent conditions. 579 Concerning matK, Barthet et al [99] demonstrated the relationship between light and 580 developmental stages, and MatK maturase activity, suggesting important functions in plant 581 physiology. This gene has recently been largely reported to be under positive selection in an 582 aquatic plant (Anubias sp.,[100]), and more generally in terrestrial plants (Pinus sp [101]) or 583 Chrysosplenium sp. [102]). The accD gene has been described as an essential gene required for

584 leaf development [103] and longevity in tobacco (Nicotiana tabacum)[104]. Under drought 585 stress, plant resistance can be increased by inhibiting accD [105], and conversely, enhanced in 586 response to flooding stress by upregulating accD accumulation [106]. Hence, we can 587 hypothesize that the positive selection observed on the accD gene can be explained by the 588 submerged and emerged constraints undergone by Ludwigia species. The ycf2 gene seems to 589 be subject to adaptive evolution in Ludwigia species. Its function, although still vague, would 590 be to contribute to a protein complex generating ATP for the TIC machinery (proteins importing 591 into the chloroplasts [107][108]), as well as plant cell survival [109][110]. The ccsA gene 592 positive selection is found in some aquatic plants such as Anubia sp.[100], marine flowering 593 plants as Zostera species [111], and some species of Lythraceae [105]. The ccsA gene is 594 required for cytochrome c biogenesis [112] and this hemoprotein plays a key role in aerobic 595 and anaerobic respiration, as well as photosynthesis [113]. Furthermore, we showed that Lgh 596 colonization is supported by metabolic adjustments mobilizing glycolysis and fermentation 597 pathways in terrestrial habitats, and the aminoacyl-tRNA biosynthesis pathway, which are key 598 components of protein synthesis in aquatic habitats [114]. It can be assumed that the ability of 599 Ludwigia to invade aquatic and wet environments, where the amount of oxygen and light can 600 be variable, leads to a high selective pressure on genes involved in respiration and 601 photosynthesis.

602 Molecular markers are often used to establish population genetic relationships through 603 phylogenetic studies. Five chloroplasts (rps16, rpl16, trnL-trnF, trnL-CD, trnG) and two 604 nuclear markers (ITS, waxy) were used in previous phylogeny studies of Ludwigia sp.[115]. 605 However, no SSR markers had previously been made available for the *Ludwigia* genus, or more 606 broadly, the Onagraceae. In this study, we identified 45 to 65 SSR markers depending on the 607 Ludwigia species. Most of them were AT mononucleotides, as already recorded for other 608 angiosperms [116][117]. In addition, we identified various genes with highly mutated regions 609 that can also be used as SNP markers. Chloroplast SSRs (cpSSRs) represent potentially useful 610 markers showing high levels of intraspecific variability due to the non-recombinant and 611 uniparental inheritance of the plastomes [118][119]. Chloroplast SSR characteristics for 612 Ludwigia sp. (location, type of SSR) were similar to those described in most plants. While the 613 usual molecular markers used for phylogenetic analysis are nuclear DNA markers, cpSSRs have 614 also been used to explore cytoplasmic diversity in many studies [120][121][122]. To conclude, 615 the 13 highly variable loci and cpSSRs identified in this study are potential markers for 616 population genetics or phylogenetic studies of Ludwigia species, and more generally, 617 Onagraceae.

618 Concerning the MatK-based phylogenetic tree, its topology is generally congruent with the 619 first molecular classification of Liu et al. [115] as all Ludwigia from sect Jussiaea (clade B1) 620 and sect. Ludwigia (clade A1) and sect. Isnardia (clade A2) branched together. In this MatK-621 based tree, Ludwigia prostrata, a species absent from previously published phylogenetic 622 studies, positions itself alone at the root of the Ludwigia tree. This species, sole member of 623 section *Nematopyxis*, is related as having no close relatives [123], finding supported by our 624 work. We also observed that Ludwigia ovalis branches within sect. Jussiaea, as its 258 amino 625 acids partial MatK sequence (ca. half of the complete sequence) is identical to the MatK 626 proteins of L. grandiflora, L. stolonifera and L. adscendens. Its phylogenetic placement remains 627 unresolved: classified alone by Raven (1963) [5] and Wagner (2017) [22] in sect. Miquelia, 628 later positioned by Liu et al. (2017)[4] within the Isnardia-Microcarpium section (using nuclear 629 DNA) or as sister to it (using plastid DNA). For this reason, conducting a whole plastome 630 analysis would be valuable to provide insights into L. ovalis phylogenetic positioning. Another 631 species positioned on the margins of sect. Isnardia (clade A2) is Ludwigia suffruticosa 632 (previously classified in sect. *Microcarpium*), which branches within sect. *Ludwigia* (clade A1). 633 This positioning raises questions about the current grouping of sections Isnardia, Michelia, and 634 Microcarpium into a single section Isnardia as proposed by Liu et al. (2023) [124] and 635 highlights that plastid protein coding markers can provide differing phylogenetic insights. 636 Finally, the last species positioned differently of this clade (clade B4) is Ludwigia decurrens 637 (sect. Pterocaulon) which clusters with L. leptocarpa (clade B3) and L. bonariensis (clade B4a). 638 However, it is important to note that in their study, Liu et al. (2017) indicate that clade B4 is 639 moderately supported and that the two members of sect. Pterocaulon, L. decurrens and L. 640 nervosa, diverge in all trees [4]. In summary, acquiring complete plastomes for Ludwigia sp. 641 could significantly enhance our understanding of the phylogeny of this complex genus. 642 Furthermore, comparing nuclear and plastid phylogenies would help determine if they reflect 643 the same evolutionary history and whether plastid phylogeny alone can accurately reconstruct 644 the phylogeny of Ludwigia genus.

645

#### 646 Conclusion

In this study, we conducted the first-time sequencing and assembly of the complete plastomes of *Lpm* and *Lgh*, which are the only available genomic resources for functional analysis in both species. We were able to identify the existence of two haplotypes in *Lgh*, but further investigations will be necessary to confirm their presence in *Lo* and *Lpm*, and more broadly, within the *Ludwigia* genus. Comparison of all 10 Onagraceae plastomes revealed a high degree of conservation in genome size, gene number, structure, and IR boundaries. However, to further elucidate the phylogenetic analysis and evolution in *Ludwigia* and Onagraceae, additional chloroplast genomes will be necessary, as highlighted in recent studies of Iris and Aristidoideae species [125].

656

#### 657 **Declarations**

• Availability of data and materials

The datasets generated and/or analysed during the current study were available in GenBank (for *Lgh* haplotype 1, (LGH1) OR166254 and *Lgh* haplotype 2, (LGH2) OR166255; for *Lpm* haplotype, (LPM) OR166256). Chloroplastic short and long reads are available at EBI-ENA database (https://www.ebi.ac.uk/ena/browser/home) under these accession numbers for LGH plastomes (Long reads : Experiment : ERX13439011 ; Run : ERR14035997 and short reads : Experiment : ERX13439002 ; Run : ERR14035988) and for LPM plastomes ( Long reads : Experiment : ERX13439014 ; Run : ERR14036000).

666

## • Conflict of interest disclosure

668 The authors declare that they comply with the PCI rule of having no financial conflicts of669 interest in relation to the content of the article

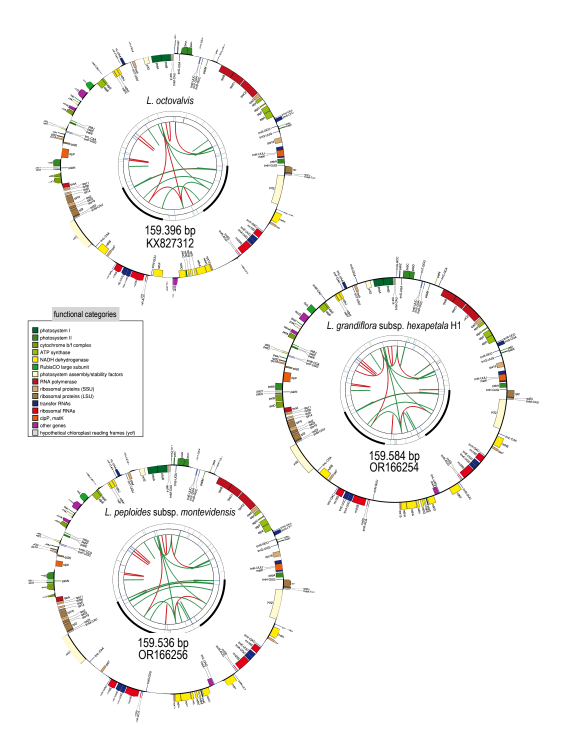
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687 Figure 3: Circular representation of annotations plastomes in Ludwigia octovalis, Ludwigia 688 grandiflora and Ludwigia peploides using ogdraw. Each card contains four circles. From the 689 center outwards, the first circle shows forward and reverse repeats (red and green arcs, 690 respectively). The next circle shows tandem repeats as bars. The third circle shows the 691 microsatellite sequences. Finally, the fourth and fifth circles show the genes colored according 692 to their functional categories (see colored legend). Only the haplotype 1 of L. grandiflora is 693 represented as haplotype 2 only diverge by the orientation of the SSC segment. Accession 694 numbers are indicated for each plastome.

# 695 Table 2. Genes present in the plastomes of *Ludwigia*

FUNCTION	Nаме							
	Photosynthesis							
Rubisco	rbcL							
Photosystem I (PSI)	psaA, psaB, psaC, psaI, psaJ							
PSI assembly factors	ycf3# (pafl), ycf4 (pafl)							
Photosystem II	psbA, psbB, psbC, psbD, psbE, psbF, psbH, psbI, psbJ, psbK, psbL, psbM, pbf1 (psbN) psbT, psbZ							
ATP synthase	atpA, atpB, atpE, atpF#, atpH, atpI							
Cytochrome b6f	petA, petB#, petD#, petG, petL, petN							
Cytochrome biogenesis	ccsA							
NADPH dehydrogenase	dehydrogenase ndhA#, ndhB**#, ndhC, ndhD, ndhE, ndhF, ndhG, ndhH, ndhI, ndhJ							
	Transcription and translation							
Transcription	rpoA, rpoB, rpoC1#, rpoC2							
Small ribosomal proteins	rps2, rps3, rps4, rps7**, rps8, rps11, rps12**#, rps14, rps15, rps16#, rps18, rps19							
Large ribosomal proteins	rpl2**#, rpl14, rpl16#, rpl20, rpl22, rpl23**, rpl32, rpl33, rpl36							
Translation initiation	infA							
Ribosomal RNA	rm5**, rm4,5**, rm16**, rm23**							
Transfer RNA	trnA-UGC**#,trnC-GCA,trnD-GUC,trnE-UUC,trnF-GAA,trnfM-CAU,trnG-GCC,trnG-UCC#,trnH-GUG,,trnI-CAU**,trnI-GAU**#,trnK-UUU#,trnL-CAA**,trnL-UAA#,trnL-UAG,trnM-CAU,trnN-GUU**,trnP-UGG,trnQ-UUG,trnR-ACG**,trnR-UCU,trnS-GCU,trnS-GGA,trnS-UGA,trnT-GGU,trnT-UGU,trnV-GAC**,trnV-UAC#,trnW-CCA,trnY-GUA							
	Other functions							
Group II intron splicing	matK							
Inorganic carbon uptake	cemA							
Protease	c/pP1#							
Fatty acid synthesis/Heat tolerance	accD							
TIC machinery (protein import)	ycf1 (Tic214), ycf2**							
Unknown function pseudogene	In function pseudogene ycf15**							
	** duplicated in IR region, # spliced genes							

## **Table 3A : Tandem repeats**

			Lgh	Lpm	Length	Region	Locus	Comments
T	IGTAGTCAGGGGTGTAGTACTAT				24	IRs	ycf2	
	AGAAGAGAGTGCAG		х	х	15	IRs	ycf2	15 nt deletion in Lgh and Lpm
A	IGAAATATCGTATAATGAAGTACCACACGAGTGGATAT	x	x		39	IRs	rpl 2 intron	39 nt deletion in Lgh and Lo
A	AAAATAGGATAGGAT		x	х	16	LSC	ycf1-trn H-GUG	56 nt deletion in Lgh and Lpm
T.	AATTAATATCTATATA		x	х	18	LSC	psb Z-trn G-GCC	18 nt deletion in Lgh and Lpm
Т	TTCTATCTATCTTATATCAA		x	x	22	LSC	tm K-UUU-rps 16	22 nt deletion in Lgh and Lpm
A	GATCCATAACATCATCAAA		x	х	20	LSC	rps 16 intron	22 nt deletion in Lgh and Lpm
T	ATTAGTTATTAATATTATTAGA		х	х	23	LSC	tm P-UGG-psa J	23 nt deletion in Lgh and Lpm
A	АТААТАТАТААТААСТТАААТА		x	х	23	LSC	rp/33-rps18	33 et 44 nt nt deletion in in Lgh et Lpm, respectively
Т	TTTTATTTAACATGCTATCAAATCAACAATGCCATACCGTAGGGCATCTGTT		x	х	53	LSC	rpl 20-clp P1	107 nt deletion in Lgh and Lpm
A	TATATTTCGATTCAATTC	х		x	19	LSC	tm H-GUG-psb A	3 copies in a 57 nt deletion in Lo and Lpm
A	TAGAAATATCAGTATTTGAGTG	х		x	23	LSC	atp H-atp I	23 nt deletion in Lo and Lpm
Т	TAATTTTAATTGAAGAA	x		x	18	LSC	psb J-psb L	17 and 24 nt deletion in Lo and Lpm, respectively
Т	TAAAGAATATTAATATTC	imperfect TR			19	LSC	trn R-UCU-atp A	A -> C mutation in second copy in Lo
T	ATTATTATTATTAAT	х	х		16	LSC	atp H-atp I	16 nt deletion in Lgh and Lo
$\mathbf{T}$	CTAAGGCTGAAATAAGG	х	х		18	LSC	pafl intron	18 nt deletion in Lgh and Lo
T	GTGAATCTATCTAT			х	15	LSC	trn S-UGA-psb Z	8 nt deletion in Lpm
Т	TTTTCTAGTA				12	LSC	pafl intron	
C	TAGTTATTGACATGG		imperfect TR	imperfect TR	16	LSC	psa J-rpl 33	G -> A mutation in second in Lpm et Lgh
A	TTTTTATTAACTCT	х		imperfect TR	15	SSC	ycf1	T->A mutation in first copy in Lpm, other sequence in first copy in Lo
A	ATCAAATAGTTGAT		х	x	15	SSC	ycf1	other sequence in first copy of Lpm and Lgh
A	ΓΑΑΤΑΑΤΑΤΑΤΤΤΑΤΤΑΑΤΤΑΑΤΑ	х			28	SSC	ndh F-rpl 32	160 nt deletion in Lo

## **Table 3B : Direct repeats**

	Sequence	Lo	Lgh	Lpm	Size (nt)	Spacers (nt)	Region	Locus	Comments
	TTCAATTGGAACGGACGATTCGTCAATCATCT				32	37	SSC	yc/1	2 copies. In Lo, one mutation (G->A) in the second copie
	CATCGATGATGAAAGTGAAAACAGTAATGAAGAGG	х			35	28 - 22 - 11	LSC	accD	3 perfects copies and 1 mutated (G->A) copie in Lgh and Lpm . Region of 174 nt deleted in Lo
	AGATGGTGAAGAACCTTATGAAGATGGTGAAGAACCTTATG		х	х	41	22	LSC	accD	Region of 147 nt deleted in Lgh and Lpm
	TATCAAATCAACAATGCCATACCGTAGGGCAT		х	х	32	22 - 21	LSC	rps12-clpP1	3 copies
	TTAAGAGCCGTACAGGCACCTTTTGATGCATACGG	х				408 in Lpm , 406 in	LSC	clp P1	2 copies. In Lgh, one mutation (C->T) in the second copie
	TTAAGAGCCGTACAGGCACTTTTTGATGCATACGG	х		х	35	811	LSC	clpP1 intron 1- intron 2	
~ ~	TGCAATAGCCAAATGATGATGAGCAATATCAGTCAGCCATA				41	2178	LSC	psaB-psaA	

## **Table 3C : Palindromic repeats**

Common perfect palidromic repeats	Logus Comment
AGACTCTCATGAGAGTCTCATTAAAT	ImC-GCA - perN ImE-UUC-ImT-GGU
TTGGTAAATTTACCAA	<i>osb</i> 0
TTCATTTCAATTTCAATTGAAATTGAAATGAA	tml-CAU-vc/2 2 cooles in IR
GAAAAGGCCTTTTC TCTCAAATGATTAATCATTTGAGA	vc/2 2 cooles in IR tmL-UM4 intron
GGATIACTATIACTATICA	tmD-GUC-tmY-GUA
TITGAATGCATICAAA	#PG-UCC inton
ATATATTCGAATATAT	ImG-UCC -ImR-UCU
TAGTAATTAATTACTA	tr/G-GCC-tr/tM-CAU
CCAGTATGCATACTGG	ndhK
Common palidromic repeats with covariation	Locus
in L. octovalvis in L. grandiflora et L. peploio	des
ATAGAATCTATATTCTATTAGAATATAGATTCTAT ATOGAATCTATATTCTATTAGAATAT	rAGATTCGAT nahC-ImV-UAC
ATGTATATATATGGAT ATCTATATATAGAT	tmE-UUC-tmT-GGU
Common palindromic and guasi-palidromic repeats	Locus Comment
in L. catovalvis in L. granditon and L. papioi	ides
TTTAACGAATATTAATATT LGTTAAA TTAA c GAATATTAATATTCTTTAA TTTAACGAATATTAATATTCGTTAAA TTAAAGA	
AATTGTA C TTACAATT AATTGTAATTACAATT	
AGGAAGATTGATCAATCTT CT AGGAAGATTGATCAATCTT	TOCT
TTA CTAATATTACTAA TTAGTAATATTACTAA TTAGTAATATTACTAA	
ATATAGAATAT CTATAT ATATAGAATATTCTATAT	T Contraction of the second
ACATATCATGATA g GT ACATATCATGATATGT	p/22
ANTTACTANTTTCTATTACTATGTTCANTTGANCATAGTANTAGAANTTAGTANTT ANTTACTANTTCTATTACTATTTCTATTACT LTGTTCANTTGANCATA	AGTAATAGAAATTAGTAATT apH-apI
TAGTTAGAATTCTAACTA TAGTT c GAATTCTAACT.	A tmT-UGU-tmL-UAA
TATTTTTTCTAGAAAAATA TATTTTTCTAGAA.gAAA	
· · · · · · · · · · · · · · · · · · ·	
in L granditiona in L granditiona	
CCCATCAATCATGATTG : TGGG CCCATCAATCATGATTG : TGGG	GGG pstN-ImD-GUC
in L. octovalvis and L. grandiflora in L. pepiloides	
ATGAAAAAATCGATTTTTTCAT ATGATAAAAATAGATTTT a	TCAT Int-UUU-ps16
ATGAAAAAAATCCATTTTTTCAT- ATGATAAAAATCGATTTTATCAT ATGATAAAAATCGATTTTATCAT ATGATAAAAAATCGATTATA	
Unique palidromic repeats	
L. paploides	
ТТАТАТАТАТАТАТАТАТА	gr/32-ndhF Full deletion in L. actionalvis , 6 bases deletion in L. grandh
L. octovativis	
ATTGAAATTCGAATTTCAAT	psbZ-tmG-GCC Full deletion in L. grandiflora and L. pepiloides
L. peptoides and L. grandillora	
AAAAAATGGATCCATTTTT	/mL-UAG-rp/32 3 bases deleted and 3 bases mutated in <i>L. octovalvis</i>
AAAAATGGATCCATTITT AATATATTATAATAATAATATT	pr/32-ndhF Full deletion in <i>L. actionalivis</i>
AAAAAATGGATCCATTTTT	
AAAAATGGATCCATTITIT AATATATTATTATTAATAATATT	pr/32-ndhF Full deletion in <i>L. actionalivis</i>
AAAAAT GGATCOLITTIT AAITATATTATATATATATAT TATATTATTATATATA	pr/32-ndhF Full deletion in <i>L. actionalivis</i>
АААААТСБААТССАТТТІТ АКТАТАТТАТАТАТАТАТАТАТ ТАТАТТАТТАТАТАТА	대 전국 소리뷰 Full deletion in L. actowarkin
AAAAATGGATCCATTTIT AATATTATTATAATAATATAT TATATTATTATAATAA	pr32-obF     Full deletion In L activativis     pr32-oubF     Full deletion In L activativis      pab2-dmG-GCC     Full deletion In L grandition and L peptioldes
AAAAAATGGAATCAITTIT           AATATATATATATATATATATATAT           I.a.tatatatatatatatatatata           I.a.tatatatatatatatata           I.a.tatatatatatatatata           I.a.tatatatatatatata           I.a.tatatatatatatatata           I.a.tatatatatatatatata           I.a.tatatatatatatatata           I.a.tatatatatatatatata           I.a.tatatatatatatatata           I.a.tatatatatatatatatata           I.a.tatatatatatatatatatata           I.a.tatatatatatatatatatatata           I.a.tatatatatatatatatatatatata           I.a.tatatatatatatatatatatatatatatatatata	pr22-coFF Full detelon IL cobravio     pr22-coFF Full detelon In L activative     pub2-terG-GOC Full detelon In L activative     term tubic pr22     term tubic pr22     3 bases detelod and 3 bases mutated In L activative
AAAAATGGATCCATTTIT           AATATATTATTATATATATT           TATATTATTATAATAATATAT           L. Cobrevini           ATTGAATTGGATTCATT           L. Cobrevini           ATTGAATTGGATTTCAT           L. L. papioleis and L. gandifor	pr32-obF     Full deletion In L activativis     pr32-oubF     Full deletion In L activativis      pab2-dmG-GCC     Full deletion In L grandition and L peptioldes

*Lo* = *Ludwigia* octovalvis; *Lgh* = *L*. *grandiflora* subsp. *hexapetala*; *Lpm* = *L*. *peploides* subsp. *montevidensis*. *100 100 110 111 112 113*

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