Applications in Plant Sciences 2015 3(11): 1500071

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...markers to compare the genetic variation in the seed bank of viable seeds (Leck, 1989). We developed 21 microsatellite...1

Cyperus fuscus L. (Cyperaceae) is an annual herb that is native in the Mediterranean region and temperate Eurasia and introduced in North America. It grows on muddy, sandy, or gravelly substrata, on shores of rivers or lakes, and is also found in anthropogenic habitats like gravel pits, wet fields, and traditionally used fish ponds. It has a short life cycle, taking just two to three months from seedling to ripe fruits (von Lampe, 1996). Cyperus fuscus is anemophilous and self-compatible. With 0.24 pg/1C (or 234.72 Mbp; Doležel et al., 2003), the genome size of...2

Premise of the study: Microsatellite markers were characterized in the extremely specialized ephemeral wetland plant species Cyperus fuscus (Cyperaceae). The markers will be used for studying population genetics in natural vs. anthropogenic habitats, on a European scale, and the role of the soil seed bank in the life cycle of this ephemeral species.

Methods and Results: Twenty-one microsatellite loci were established and scored in two populations, with mean number of alleles of 2.6 and 2.9 and mean expected heterozygosity of 0.405 and 0.470, respectively. Forty-four additional loci with the number of alleles ranging from one to four (mean = 2.1) were successfully amplified in seven individuals.

Conclusions: The novel microsatellite markers will be useful for studying the genetic structure of populations of this ephemeral plant as well as their seed bank.

Key words: 454 sequencing; Cyperaceae; Cyperus fuscus; Isoëto-Nanojuncetea; microsatellites.

METHODS AND RESULTS

Plants were grown in the greenhouse from ripe seeds collected in the field (Appendix 1). Genomic DNA of fresh leaves from one plant was extracted with the DNeasy Plant Mini Kit (QIAGEN, Hilden, Germany) following the manufacturer’s instructions and sent to LGC Genomics (Berlin, Germany) for next-generation sequencing (NGS) on a Genome Sequencer FLX Titanium Instrument (454 Life Sciences, a Roche Company, Branford, Connecticut, USA). In this first run, 143,027 sequence reads with an average length of 238 bp were obtained (Table 1). NGS data are deposited in the GenBank Sequence Read Archive (BioProject no. PRJNA275048). MSATCOMMANDER version 0.8.2 (Faircloth, 2008) was used to detect 520 sequences with simple sequence repeat (SSR) motifs (options: dinucleotide repeats ≥10 repeat units, tri- and tetranucleotide repeats ≥6 repeat units, combine multiple arrays within a sequence if within 50 bp distance). Primers for microsatellite-containing sequences were also designed in MSATCOMMANDER using Primer3 (Rozen and Skaletsky, 1999), with a GTTT PIG-tail (Brownstein et al., 1996) added to the 5’ end of one primer and a CAG or M13R tail (CAG: 5′-CAGTCGCGGCTATCA-3′; M13R: 5′-GGAACAGCTATGACCAT-3′) added to the 5’ end of the other primer (Schuelke, 2000). Due to the shortness of the sequences (range = 7–762 bp, mean = 238 bp), only 101 out of the 520 SSR-containing sequences were suitable for primer design. PCR amplifications were performed in a 25-µL final volume of REDTaq ReadyMix PCR Reaction Mix (Sigma-Aldrich, St. Louis, Missouri, USA) with 0.40 µM 5’ FAM-labeled universal CAG or M13R primer, 0.40 µM GTTT-tailed primer, 0.04 µM CAG- or M13R-tailed primer, and 1 µL diluted DNA extract (2–20 ng DNA). Reactions were performed using a touchdown PCR protocol in an Eppendorf Mastercycler gradient (Eppendorf, Hamburg, Germany), with an initial 5 min of denaturation at 95°C, 24 cycles with denaturation at 95°C for 45 s, annealing at 63–48.6°C (0.6°C decrease per cycle) for 90 s, and extension at 72°C for 60 s; 19 cycles with denaturation at 95°C for 45 s, annealing at 50°C for 90 s, and extension at 72°C for 60 s; and a final extension at 72°C for 5 min and 60°C for 30 min. Amplified fragments were analyzed on a 3500 Genetic Analyzer (Applied Biosystems, Foster City, California, USA) and sized using GeneMarker 2.4 (SoftGenetics, State College, Pennsylvania, USA). The markers were tested on seven individuals from different localities (Appendix 1). Seven loci could be unambiguously scored in all seven test individuals. Four of these were applied to a larger number of individuals (primers with the prefix Cf in Table 2; remaining loci are shown in Appendix 2). A second NGS run of an SSR-enriched library was performed at ecogenics (Balgach, Switzerland), starting from a mix of genomic DNA of two individuals (Appendix 1). Size-selected fragments from genomic DNA were enriched for SSR content by using magnetic streptavidin beads and biotin-labeled CT, GT, AAG, and ATGT repeat oligonucleotides. The SSR-enriched library...
was analyzed on a Roche 454 platform using the GS FLX Titanium reagents (454 Life Sciences, a Roche Company). In total, 4877 reads with a mean length of 415 bp were obtained and deposited in the GenBank Sequence Read Archive (BioProject no. PRJNA275048), of which 967 contained SSR motifs (MSATCOMMANDER search and primer design settings same as above; Table 1). Four hundred ninety-four reads were suitable for primer design. Ego-
genics sent 80 primer pairs also designed with Primer3, containing an M13 tail at the 5′ end of the forward primer (5′-GTGAAAACGACGGCCAGT-3′; Schuelke, 2000) and no PIG-tail. For primer testing, the concentrations and volumes for PCR were the same as above, but we used JumpStart REDTaq ReadyMix Reaction Mix (Sigma-Aldrich) and a regular PCR protocol, with an initial 5 min of denaturation at 95°C; 38 cycles of denaturation at 95°C for 45 s, annealing at 56°C for 60 s, and extension at 72°C for 1 min; and a final extension at 72°C for 5 min and 60°C for 30 min. Of these 80 markers, 22 showed no PCR product or had a weak signal, failures, or were unspecific. The remaining 58 mark-
ers showed clear peaks. Ten of these were monomorphic and 48 polymorphic. Seventeen polymorphic markers were selected for further analysis and com-
bined into four multiplex PCRs with Multiplex Manager version 1.0 (Holleley and Geerts, 2009; PCR multiplex sets 1–4 in Table 2). The remaining loci are

### Table 1. Characteristics of the two 454 GS FLX Titanium sequencing runs.

| Sequencing run | Total no. of reads | Range of read lengths (bp) | Average read length (±SD; bp) | GC content (%) | SSR-containing sequences (total no. of SSRs encountered) | No. of reads useful for primer design |
|----------------|-------------------|---------------------------|-----------------------------|----------------|--------------------------------------------------------|-------------------------------------|
| First run      | 145,027           | 7–762                     | 238 (±130)                  | 40.2           | 520 (539)                                              | 101                                 |
| Second run     | 4877              | 34–801                    | 415 (±165)                  | 40.7           | 967 (990)                                              | 494                                 |

*Note: SD = standard deviation.

*In the first run, a crude extract of genomic DNA of a single Cyperus fuscus individual was used. In the second run, an enriched library, generated from genomic extracts of two C. fuscus individuals, was used. See Appendix 1 for origin of sequenced individuals.*

The 21 newly developed microsatellite markers were applied to 25 individu-
als from each of two fish pond populations in the Czech Republic (Appendix 1). Interpretation of electropherograms in all loci and all individuals is compatible with a diploid cytotype. The number of alleles, observed ($H_o$) and expected heterozygosity ($H_e$), fixation index, and exact test for Hardy–Weinberg equilibri-

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Table 2. Characteristics of 21 SSR loci developed in *Cyperus fuscus*.

| Locus  | Primer sequences (5′–3′)b | PCR multiplex set | Fluorescent dyec | Repeat motif | A | Allele size range (bp)i | EMBL accession no. |
|--------|----------------------------|-------------------|-----------------|--------------|---|------------------------|-------------------|
| Cf_008 | F: GGAACACGCTATGACCATAGTGATCAAAAATTACACGGATCAGGGACG<br>R: GTTTACGATGACATGACAGATAGATTAACGGATCAGGGACG | NA | ATTO 565 | (AG) 11 | 4 | 312–344 | LN848930 |
| Cf_017 | F: GGAACACGCTATGACCATGAGGCAATAGAAATTGTTGGAG<br>R: GTTTGAGACAGATTACTCACCTCTCAAG | NA | ATTO 550 | (CTT) 13 | 3 | 218–242 | LN848931 |
| Cf_019 | F: GTTTAATTGTCAGGCCACATGCC<br>R: GGAACACGCTATGACCATACAGGGAGCAACCTGAGC | NA | FAM | (CTT) 7 + (CTT) 6 | 2 | 184–205 | LN848932 |
| Cf_104 | F: GGAACACGCTATGACCATGACAGAAGATGAATTAAGGCCAC<br>R: GTTTTCGATGACAGTTTAAAGGTCCAG | NA | Yakima Yellow | (GT) 14 | 2 | 180–184 | LN848934 |
| Cypfus_0173 | F: CGCCAAAGGAGAATGAGGTG<br>R: GTTTATCGAACAATCCGATCTCGC | 1 | ATTO 532 | (GAA) 9 | 3 | 189–201* | LN848937 |
| Cypfus_0551 | F: TTTCCAATTGACGGACCAAC<br>R: GTTTAGCGTGCTATTTACAACCTTGG | 4 | FAM | (CTT) 13 | 3 | 230–236* | LN848952 |
| Cypfus_1207 | F: ATCCTTCACCTCCTCCCCCCATC<br>R: GTTTGGAGTAAACCACGGACTCG | 1 | ATTO 565 | (CTT) 12 | 4 | 218–245* | LN848954 |
| Cypfus_2257 | F: TGTGATGACGAGGAGGTTGG<br>R: GTTTGTACAGGTAAGCGCAAGCM13: TGTAAAACGACGGCCAGT | 3 | ATTO 550 | (CTT) 13 | 4 | 249–264* | LN848964 |
| Cypfus_2663 | F: TGTGATGACGAGGAGGTTGG<br>R: GTTTGTACAGGTAAGCGCAAGCM13: TGTAAAACGACGGCCAGT | 3 | ATTO 565 | (CTT) 13 | 4 | 141–162* | LN848965 |
| Cypfus_2987 | F: GGAACACGCTATGACCATGACAGAAGATGAATTAAGGCCAC<br>R: GTTTTCGATGACAGTTTAAAGGTCCAG | 1 | ATTO 550 | (CTT) 9 | 3 | 209–227* | LN848966 |
| Cypfus_3218 | F: GGAACACGCTATGACCATGACAGAAGATGAATTAAGGCCAC<br>R: GTTTTCGATGACAGTTTAAAGGTCCAG | 2 | FAM | (GAA) 8 | 3 | 163–193* | LN848967 |
| Cypfus_3300 | F: GGAACACGCTATGACCATGACAGAAGATGAATTAAGGCCAC<br>R: GTTTTCGATGACAGTTTAAAGGTCCAG | 3 | ATTO 550 | (CTT) 12 | 4 | 189–221* | LN848968 |
| Cypfus_3921 | F: TGTGATGACGAGGAGGTTGG<br>R: GTTTGTACAGGTAAGCGCAAGCM13: TGTAAAACGACGGCCAGT | 2 | ATTO 550 | (GAA) 8 | 3 | 261–270* | LN848970 |
| Cypfus_4093 | F: TGTGATGACGAGGAGGTTGG<br>R: GTTTGTACAGGTAAGCGCAAGCM13: TGTAAAACGACGGCCAGT | 2 | ATTO 550 | (GAA) 8 | 3 | 261–270* | LN848982 |
| Cypfus_4163 | F: GGAACACGCTATGACCATGACAGAAGATGAATTAAGGCCAC<br>R: GTTTTCGATGACAGTTTAAAGGTCCAG | 2 | FAM | (GAA) 8 | 3 | 163–193* | LN848983 |
| Cypfus_4236 | F: GGAACACGCTATGACCATGACAGAAGATGAATTAAGGCCAC<br>R: GTTTTCGATGACAGTTTAAAGGTCCAG | 4 | TATG | (AG) 12 | 3 | 176–184* | LN848990 |
| Cypfus_4666 | F: GGAACACGCTATGACCATGACAGAAGATGAATTAAGGCCAC<br>R: GTTTTCGATGACAGTTTAAAGGTCCAG | 3 | Yakima Yellow | (TATG) 7 | 3 | 189–221* | LN848995 |

Note: A = number of alleles sampled; EMBL = European Molecular Biology Laboratory.

a Primers with the prefix Cf are from an NGS run from raw genomic DNA libraries; primers with the prefix Cypfus are from an NGS run from an enriched library.
b GTTT PIG-tails (Brownstein et al., 1996), M13R tails (5′-GGAAACAGCTATGACCATCAGCTTCAG-3′; Cf-primers), and M13 tails (5′-TGTAAAACGACGGCCAGCTTCAGATTGAGTCGACAGCTTCAG-3′; Cypfus_4093) added to the 5′ ends of primers are underlined.
c Fluorescent dye at the 5′ ends of M13R and M13 primers (Cf-primers and Cypfus_4093) and forward primers (remaining loci).
d The allele range is based on seven test individuals (Appendix 1).
* Length of PCR products is without PIG-tail, but with M13 tail (as for other loci resulting from the second NGS run in Appendix 2).
### Table 3. Genetic diversity of 21 newly developed SSR markers in two fish pond populations of *Cyperus fuscus*.a

| Locus            | Population | A  | H₀  | Hₑ  | Fₛᵇ  | A  | H₀  | Hₑ  | Fₛᵇ  |
|------------------|------------|----|-----|-----|------|----|-----|-----|------|
| Cf_008           | Zahrádky   | 4  | 0.240 | 0.577 | 0.589*** | 4  | 0.160 | 0.565 | 0.721*** |
| Cf_017           | Zahrádky   | 2  | 0.160 | 0.470 | 0.664*** | 3  | 0.240 | 0.528 | 0.551*** |
| Cf_019           | Zahrádky   | 3  | 0.040 | 0.365 | 0.892*** | 2  | 0.120 | 0.497 | 0.762*** |
| Cf_104           | Zahrádky   | 2  | 0.040 | 0.301 | 0.870*** | 2  | 0.200 | 0.301 | 0.341  |
| Cypfus_0173      | Libohošť  | 3  | 0.120 | 0.541 | 0.782*** | 2  | 0.200 | 0.510 | 0.613*** |
| Cypfus_0551      | Libohošť  | 2  | 0.080 | 0.509 | 0.846*** | 2  | 0.080 | 0.509 | 0.846*** |
| Cypfus_1207      | Libohošť  | 3  | 0.080 | 0.223 | 0.646***  | 3  | 0.160 | 0.496 | 0.682*** |
| Cypfus_2257      | Libohošť  | 2  | 0.200 | 0.301 | 0.341  | 3  | 0.160 | 0.545 | 0.711*** |
| Cypfus_2506      | Libohošť  | 2  | 0.160 | 0.509 | 0.690*** | 3  | 0.120 | 0.667 | 0.823*** |
| Cypfus_2663      | Libohošť  | 2  | 0.080 | 0.444 | 0.823*** | 3  | 0.160 | 0.562 | 0.710*** |
| Cypfus_2987      | Libohošť  | 4  | 0.120 | 0.381 | 0.690*** | 3  | 0.120 | 0.548 | 0.784*** |
| Cypfus_2993      | Libohošť  | 3  | 0.080 | 0.401 | 0.804*** | 2  | 0.040 | 0.184 | 0.786*** |
| Cypfus_3114      | Libohošť  | 3  | 0.120 | 0.541 | 0.782*** | 3  | 0.200 | 0.601 | 0.672*** |
| Cypfus_3212      | Libohošť  | 2  | 0.040 | 0.510 | 0.923*** | 2  | 0.120 | 0.507 | 0.767*** |
| Cypfus_3218      | Libohošť  | 2  | 0.120 | 0.510 | 0.768*** | 4  | 0.240 | 0.584 | 0.594*** |
| Cypfus_3300      | Libohošť  | 4  | 0.320 | 0.706 | 0.552*** | 5  | 0.200 | 0.579 | 0.569*** |
| Cypfus_3921      | Libohošť  | 2  | 0.000 | 0.078 | 1.000*  | 2  | 0.160 | 0.490 | 0.675*** |
| Cypfus_4093      | Libohošť  | 2  | 0.120 | 0.301 | 0.607*  | 3  | 0.000 | 0.153 | 1.000*** |
| Cypfus_4216      | Libohošť  | 3  | 0.120 | 0.411 | 0.712*** | 4  | 0.000 | 0.584 | 1.000*** |
| Cypfus_4236      | Libohošť  | 2  | 0.040 | 0.350 | 0.888*** | 3  | 0.120 | 0.411 | 0.712*** |
| Cypfus_4666      | Libohošť  | 2  | 0.000 | 0.078 | 1.000*  | 2  | 0.040 | 0.040 | 0.000  |

**Note:**
- **A:** number of alleles sampled;
- **Fₛᵇ:** fixation index;
- **Hₑ:** expected heterozygosity;
- **H₀:** observed heterozygosity;
- **N:** number of individuals sampled;
- **SD:** standard deviation.

*a* See Appendix 1 for locality information for each population.

*b* Significant departures from Hardy–Weinberg equilibrium: *P < 0.05, **P < 0.01, ***P < 0.001.

### Appendix 1. Voucher information for *Cyperus fuscus* populations used in this study. All vouchers are deposited at the Institute of Botany, University of Natural Resources and Life Sciences, Vienna (WHB). Individuals were grown from seeds in the greenhouse.

| Voucher no. | Collection locality | Geographic coordinates | N  |
|-------------|---------------------|------------------------|----|
| 62957a      | Czech Republic, Záryby | 50°13.424’N, 14°37.717’E | 1  |
| 62959b      | Czech Republic, Semnice | 49°45.067’N, 14°39.635’E | 1  |
| 62987c      | Czech Republic, Tchořopice | 49°26.115’N, 13°48.442’E | 1  |
| 62962c      | Czech Republic, Mšec | 50°11.815’N, 13°54.651’E | 1  |
| 62960c      | Czech Republic, Hluboká nad Vltavou | 49°02.624’N, 14°25.952’E | 1  |
| 62964d      | Czech Republic, Zahrádky | 49°26.078’N, 13°54.699’E | 1  |
| 62982d      | Czech Republic, Břeclav | 48°42.710’N, 16°54.169’E | 1  |
| 62979d      | Czech Republic, Velké Němčice | 48°59.056’N, 16°39.894’E | 1  |
| 62973c      | Poland, Borków | 51°40.477’N, 16°12.239’E | 1  |
| 62955c      | Poland, Cigacice | 48°18.739’N, 16°54.224’E | 1  |
| 62968d      | Czech Republic, Zahrádky | 50°37.687’N, 14°32.595’E | 25 |
| 62964d      | Czech Republic, Libohošť | 49°42.057’N, 14°35.398’E | 25 |

**Note:** N = number of individuals sampled.

*a* Used for first NGS run at LGC Genomics (Berlin, Germany).

*b* Used for second NGS run at ecogenics (Balagch, Switzerland).

*c* Test individuals for screening of primer pairs.

*d* Test populations for assessment of genetic diversity.
## Appendix 2. Characteristics of 44 additional SSR loci with flanking regions useful for primer design in *Cyperus fuscus*

| Locus     | Primer sequences (5′–3′) | Repeat motif | A | Allele size range (bp) | EMBL accession no. |
|-----------|--------------------------|--------------|---|------------------------|--------------------|
| **First NGS run** |                          |              |   |                        |                    |
| Ct_007    | F: CAGTCGGGCGTCATCACGAGTATTTGAGATGATGAGGACC <br>R: GTTTAAGGTGCAAGATGAGTCGCGG   | (AT)₁₁       | 3 | 274–286                | LN848929           |
| Cf_020    | F: GGAACACAGCATGACCCCTCTGAGGCCACATTCACTGAGG <br>R: GTTTAGGCCATGACCCTCTCCCACTGACC   | (GGT)₃ + (GGT)₅ | 1 | 273                    | LN848933           |
| Cf_112    | F: GATTCGGGCGTCATCACGAGTATTTGAGATGAGGACC <br>R: GTTTAAGGTGCAAGATGAGTCGCGG   | (AATG)₁       | 1 | 203                    | LN848935           |
| **Second NGS run** |                          |              |   |                        |                    |
| Cypfus_0023 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₈       | 4 | 160–170                | LN848941           |
| Cypfus_0563 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (AT)₁₂       | 2 | 139–143                | LN848939           |
| Cypfus_0568 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 153–175                | LN848940           |
| Cypfus_0785 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 152–155                | LN848943           |
| Cypfus_1174 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 162–166                | LN848947           |
| Cypfus_1319 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 164–202                | LN848948           |
| Cypfus_1398 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 207–210                | LN848950           |
| Cypfus_2381 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 213–222                | LN848951           |
| Cypfus_2517 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 215–226                | LN848952           |
| Cypfus_2640 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 224–269                | LN848953           |
| Cypfus_2806 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 230–270                | LN848954           |
| Cypfus_2855 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 242–328                | LN848955           |
| Cypfus_2891 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 255–315                | LN848956           |
| Cypfus_2898 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 320–360                | LN848957           |
| Cypfus_3033 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 361–466                | LN848958           |
| Cypfus_3195 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 473–521                | LN848959           |
| Cypfus_3323 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 531–616                | LN848960           |
| Cypfus_3597 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 638–723                | LN848961           |
| Cypfus_3776 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 732–794                | LN848962           |
| Cypfus_3864 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 806–896                | LN848963           |
| Cypfus_3873 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 907–1000               | LN848964           |
| Cypfus_3898 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 1021–1110              | LN848965           |
| Cypfus_4041 | F: GTTTAACGAGCCATGAGTAAGGAAATTTGGTCAGG <br>R: TGGTTAAGGTGCAAGATGAGTCGCGG   | (GA)₁₂       | 2 | 1121–1200              | LN848966           |

http://www.bioone.org/loi/apps
APPENDIX 2. Continued.

| Locus     | Primer sequences (5′–3′)a | Repeat motif | A | Allele size range (bp)b | EMBL accession no. |
|-----------|---------------------------|--------------|---|------------------------|--------------------|
| Cypfus_4074 | F: TGTAACGAGCGCGCATTTGGCCATGGGACAGCAAGAG | (TC)₃₃ | 4 | 184–198 | LN848985 |
|           | R: UCTTAAGGTAGGACACAGGGG |              |   |                        |                    |
| Cypfus_4102 | F: TGTAACGAGCGCGCATTTGGGCGTTTCAATCAAGAGAG    | (GA)₃₃ | 1 | 260      | LN848987 |
|           | R: GGGGCCACACTGAAGAAAGAA     |              |   |                        |                    |
| Cypfus_4240 | F: TGTAACGAGCGCGCATTTGGGCGTTTCAATCAAGAGAG    | (TACA)₇    | 2 | 251–255 | LN848991 |
|           | R: GGGGCCACACTGAAGAAAGAGAG   |              |   |                        |                    |
| Cypfus_4347 | F: TGTAACGAGCGCGCATTTGGGCGTTTCAATCAAGAGAG    | (TGTA)₇    | 2 | 252–256 | LN848992 |
|           | R: CAATACACTCGCACTCACTCA     |              |   |                        |                    |
| Cypfus_4468 | F: TGTAACGAGCGCGCATTTGGGCGTTTCAATCAAGAGAG    | (CT)₃₂    | 3 | 259–275 | LN848993 |
|           | R: AGATATCAAAAGCAGCACGCCACC |              |   |                        |                    |
| Cypfus_4479 | F: TGTAACGAGCGCGCATTTGGGCGTTTCAATCAAGAGAG    | (AAG)₉     | 2 | 158–233 | LN848994 |
|           | R: AGATATACAAAGCAGCACGCCACC  |              |   |                        |                    |
| Cypfus_4799 | F: TGTAACGAGCGCGCATTTGGGCGTTTCAATCAAGAGAG    | (AAG)₉     | 2 | 248–251 | LN848996 |
|           | R: AGATATACAAAGCAGCACGCCACC  |              |   |                        |                    |
| Cypfus_4849 | F: TGTAACGAGCGCGCATTTGGGCGTTTCAATCAAGAGAG    | (GA)₃₂    | 1 | 158      | LN848997 |
|           | R: AAAAAAACACCACTTCGCGTTAAGCAAGAG |              |   |                        |                    |

Note: A = number of alleles sampled; EMBL = European Molecular Biology Laboratory.

aGT TT PIG-tails (Brownstein et al., 1996), CAG and M13R tails (CAG: 5′-CAGTCGGGCGTCATCA-3′; M13R: 5′-GGAACAGCTATGACCAT-3′; only in Cf_007, Cf_020, and Cf_112), and M13 tails (5′-TGTAACGAGCGGACAGCT-3′) added to the 5′ ends of primers are underlined.
bThe allele range is based on seven test individuals (Appendix 1).