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#### Abstract

We describe the development of 13 microsatellite markers in the pine processionary moth *Thaumetopoea pityocampa*. In a single Portuguese population tested, the number of alleles per locus ranged from 2 to 12, and observed heterozygosity from 0.07 to 0.83. Cross amplifications gave satisfactory results for most loci in the *T. pityocampa/T. wilkinsoni* complex but proved to be of low interest for other species. These markers will be valuable to investigate the population genetic structure of this forest pest, particularly for the expanding populations at the northern limits of its range, and for a new phenological race discovered in Portugal.

The isolation of microsatellite markers in Lepidoptera is known to be difficult, and therefore only few loci are available for most species (Zhang 2004). The pine processionary moth Thaumetopoea pityocampa is a forest pest, responsible for heavy loss of production in pine forests in Southern Europe through the consumption of needles by its gregarious larvae. It is also known from public-health and veterinary services because of its urticating hairs that can induce allergic reactions. The distribution area of this species is clearly limited both by the availability of suitable hosts and by climatic conditions during larval development, i.e. high summer temperatures in the south and low winter temperatures in the northern areas. Its distribution range is expanding northward and in higher altitudes due to the ongoing climate warming, and in particular to the improvement of winter conditions (Battisti et al. 2005). Another matter of interest is the presence in Portugal of a peculiar, unique population that exhibits a shifted phenology, with reproduction in May and larval development in summer (summer population) while the classical life cycle implies a reproduction in autumn and larval development during winter. The summer population is supposed to have originated from the local winter population (Santos et al. 2011a) and seems to be adapted to higher temperature during larval development (Santos et al. 2011b). The shift in phenology is such that it prevents crosses between the sympatric winter and summer populations (Santos et al. 2011a).

A first set of 6 loci was previously developed for this species (Rousselet *et al.* 2004; Santos *et al.* 2007). Yet, this was far too few to develop landscape genetics approaches and investigate the genetic structure and origin of newly settled populations around the expansion fronts, either at the northern limits of the natural range or at higher elevation in mountains. It was also insufficient to test scenarios about the foundation and origin of the allochronic population in Portugal. We thus decided to develop new polymorphic microsatellite markers, and we describe here the development of 13 new loci.

An enriched library was made by Ecogenics GmbH (Zurich, Switzerland) from size selected genomic DNA ligated into SNX forward/SNX reverse linker (Hamilton *et al.* 1999) and enriched by magnetic bead selection with biotin-labelled (CT)<sub>13</sub>, (GT)<sub>13</sub>, (GTAT)<sub>7</sub> and (GATA)<sub>7</sub> oligonucleotide repeats (Gautschi *et al.* 2000a, b). Of 528 recombinant colonies screened, 242 gave a positive signal after

hybridization. Plasmids from 126 positive clones were sequenced and primers were designed for 24 microsatellite inserts, of which 21 were tested for polymorphism. 13 primer pairs were finally selected that gave satisfying experimental results, and grouped in 4 multiplex.

Amplification for each multiplex was optimized to be performed in a 10  $\mu$ l reaction volume containing <10 ng of DNA, 5  $\mu$ l 2X QIAGEN multiplex PCR master mix, distilled water, and 0.3  $\mu$ M of forward and reverse primers each, 10% to 50% of the forward primer being labeled with a fluorescent dye (either 6-Fam, Hex, Ned or Vic, Applied Biosystems). We used the following thermotreatment: 35 cycles at 94°C for 30 seconds, 56°C for 90 seconds, and 72°C for 60 seconds. Before the first cycle, a prolonged denaturation step (95°C for 15 min) was included and the last cycle was followed by a 30 min extension at 72°C. Fragments were run on an ABI 3730 automatic sequencer using the size standard GeneScan-600 LIZ, and sized with GENEMAPPER 4.0 software (Applied Biosystems).

Microsatellite loci were amplified in multiplex from 30 larvae belonging to a Portuguese population exhibiting the classical winter phenology, in the forest of Leiria (39°47'N, 8°58'W). We also tested cross amplification, with the same PCR conditions, for individuals from different clades of the *T. pityocampa/T. wilkinsoni* complex, namely from the *pityocampa*, *wilkinsoni* and ENA clades (Kerdelhué *et al.* 2009) and for 5 others species of the same genus.

Null allele frequencies were estimated using FreeNA (Chapuis & Estoup 2007) following the Expectation Maximization algorithm (Dempster *et al.* 1977). Genetic diversity indices were calculated, and used to test for Hardy-Weinberg equilibrium and linkage disequilibrium with ARLEQUIN 3.11 software (Excoffier *et al.* 2005).

Polymorphism characteristics in the Leiria population are provided for each locus in Table 1. Number of alleles per locus ranged from 2 (MS-*Thpit*17 and 18) to 12 (MS-*Thpit*13). The estimated null allele frequencies were below 5% for all loci except MS-*Thpit*11. This locus was also the only one exhibiting deviation from Hardy-Weinberg equilibrium. This deficit of heterozygotes is probably linked to the 13% of null alleles estimated for this locus. No significant linkage disequilibrium was detected between any pair of loci (p < 0.01).

Cross amplifications proved the usefulness of these microsatellites in the *T. pityocampa/T. wilkinsoni* complex (Table 2). Most loci can be successfully amplified in all localities and exhibit polymorphism in each taxon despite the low sampling effort (but a large geographical scale). For *T. wilkinsoni*, amplification failed for all individuals when using MS-*Thpit*7 and 9, and for some individuals with MS-*Thpit*10, 14 and 19. Such partial success was also obtained for individuals from the ENA clade with MS-*Thpit*9 and 19. Considering the others species on gymnosperm, we obtained positive and clear amplification only for 2 to 5 loci, the individual tested for each species being homozygous. Finally, no amplification product was obtained for *T. solitaria*, the only species hosted by an Angiosperm.

These results mean that we now benefit from a total of 19 microsatellite loci for *T. pityocampa* population genetic studies. This will allow to more precisely investigate the origin of the allochronic population in Portugal, and to look for the existence of natural hybrids with the sympatric winter population. It should be also possible to characterize the process of latitudinal range expansion in regard to climate warming and landscape characteristics. It opens further perspectives to link the evolution of adaptive traits and neutral genetic patterns underlying the establishment of new populations.

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Locus	Primer sequences (5'-3')	Repeat motif (cloned allele)	Multiplex	Genbank	Allele range	NA	Null allele	H <sub>0</sub>	$H_{\rm E}$
				Accession no	. (bp)		frequency		
MS-Thpit 7	F GGCTCGTGACCTAAAACTGC	(TG) <sub>13</sub>	M1	JN400258	237-247	6	0.01480	0.633	0.693
	R ACACGCCCACATACATGC								
MS-Thpit 8	F GAGTGGTCGCCCTGTCTG	$(CA)_{15}(CG)(CA)_{10}$	M1	JN400259	182-228	6	0.00000	0.600	0.543
	R CTCCAATGCCAAGGTAGACAC								
MS-Thpit 9	F TTGCACAAGACAATAAAAAGAACAG	(AG) <sub>26</sub>	M1	JN400260	85-107	6	0.02109	0.367	0.404
	R TCTCACTCTCTCACACACACTATTTC								
MS-Thpit 10	F TGAATTTGGAGGAAACGATG	(GT) <sub>10</sub>	M1	JN400261	148-164	4	0.02924	0.633	0.683
	R CCACTGCATGTTCGCATAAC								
MS-Thpit 11	F GCAGGTGGAAAGAAGTGGTC	(GT) <sub>15</sub>	M2	JN400262	198-226	6	0.12928	0.167	0.275
	R TCTCCGGAAGAGAATCCAAG								
MS-Thpit 12	F GTTCTACGTCGGGAACAAGC	(CA)19	M2	JN400263	130-176	8	0.00001	0.567	0.625
	R ACGGTTCCTGTGCGATATAC								
MS-Thpit 13	F TTTTGATTGTCAGCACTCACG	(CA)10(CG)(CA)5(CG)(CA)6	M2	JN400264	99-137	12	0.00000	0.833	0.815
	R AGAGCGTGGTTCGTTCTTTG								
MS-Thpit 14	F TCACTACTGGACTTCAACGACTG	(AC)11(GCAA)(AC)13	M3	JN400265	216-372	10	0.03082	0.667	0.701
	R CGAGAAGTAAAAGTGGTATGTGT								
MS-Thpit 15	F ACACGCGCACGCTTACTC	(AC) <sub>25</sub>	M3	JN400266	150-198	11	0.00464	0.733	0.798
_	R TCAGCAAAGTTGGAGCATAAAC								
MS-Thpit 16	F ACACGACAGGCTTGCACA	(AG) <sub>22</sub>	M3	JN400267	51-65	3	0.02592	0.400	0.468
-	R TTTTCCTTTACTAACTCTGTCCAT								
MS-Thpit 17	F AAAGGCCAAGTTCGTAGCAC	(TATG) <sub>7</sub>	M4	JN400268	180-184	2	0.00003	0.067	0.065
-	R CCACATACAAAAGTTCAAAAATACG								
MS-Thpit 18	F TTCAAGAACAATGTGGCTTTC	(TCA) <sub>9</sub>	M4	JN400269	144-147	2	0.00000	0.533	0.427
Ĩ	R GTCTACCAACCCGCACTAGG								
MS-Thpit 19	F AAGCACTCGTCCGCAGTC	(TG) <sub>11</sub> (TG) <sub>12</sub> (GT) <sub>14</sub> (GT) <sub>14</sub>	M4	JN400270	113-311	6	0.00001	0.633	0.662
	R GGGGTTCTAAAATGCCAGTTG								

Table 1 Microsatellite data and polymorphism characterization of the T. pityocampa population from Leiria

 $N_{\rm A}$ : number of alleles;  $H_{\rm O}$ : observed heterozygosity;  $H_{\rm E}$ : expected heterozygosity; values in bold indicate significant deviation from Hardy-Weinberg equilibrium (P < 0.05)

Species	host plant	Country	Ν	MS-Thpit 7	MS-Thpit 8	MS-Thpit 9	MS-Thpit 10	MS-Thpit 1	1 MS-Thpit 12	2 MS-Thpit 13	MS-Thpit 1	4 MS-Thpit 1	5 MS-Thpit 16	6 MS-Thpit 1	7 MS-Thpit 18	3 MS-Thpit 19
T. pityocampa	Pinus	Morocco, W.Turkey	2	2/3	2/4	2/4	2/4	2/4	2/3	2/3	2/3	2/3	2/3	2/1	2/3	2/4
T. p. ena clade	Pinus	Algeria, Libya, Tunisia	3	3/6	3/4	1/1	3/4	3/4	3/2	3/6	3/5	3/4	3/5	3/2	3/3	2/2
T. wilkinsoni	Pinus	Crete, Cyprus, S. & N Turkey	4	0	4/5	0	1/2	4/1	4/2	4/3	3/6	4/7	4/6	4/1	4/1	3/5
T. pinivora	Pinus	France	1	1/1	0	0	0	1/1	1/1	1/1	0	0	0	0	0	0
T. libanotica	Cedrus	Lebanon	1	1/1	0	0	0	1/1	1/1	1/1	0	0	0	0	1/1	0
T. bonjeani	Cedrus	Algeria	1	1/1	0	0	0	1/1	1/1	0	0	0	0	0	0	0
T. ispartensis	Cedrus	Turkey	1	1/1	0	0	0	1/1	0	0	0	0	0	0	0	0
T. solitaria	Pistacia	Greece	2	0	0	0	0	0	0	0	0	0	0	0	0	0

Table 2 Cross-species amplification results for *Thaumetopea* spp.: number of successful amplifications / number of alleles in the taxon

N: number of individuals tested