

# 1 Isolation and characterisation of eight polymorphic 2 microsatellite markers from South American limpets 3 of the species complex *Nacella*

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10 Abstract

11 In this study we provide eight polymorphic microsatellite markers for the two South  
12 American patellogastropods *Nacella magellanica* and *N. deaurata*. Microsatellite  
13 amplification was carried out in multiplex PCRs, a new feature of the program pipeline  
14 STAMP. Allelic diversity ranged from 5 to 57 alleles per locus. Observed heterozygosities  
15 varied between 0.1 and 0.98. Three of the four loci designed for *N. magellanica* cross  
16 amplified also with *N. deaurata*, and two loci vice versa. Six of the microsatellites  
17 successfully cross amplified with the two sister taxa *N. mytilina* and *N. delicatissima*. This set  
18 of microsatellites provides a suitable tool for population genetic purposes and can be of  
19 important help in identifying morphologically ambiguous *Nacella* individuals.

20 *Keywords: Nacellidae, Population genetics, multiplex PCR, Patagonia, coastal organism*

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23 Species of the family *Nacella* mainly inhabit coastal areas of Chilean and Argentinian coasts  
24 in Patagonia and Tierra del Fuego as well as on the Falkland Island, UK. These South  
25 American members of the family comprise the following nine species: *Nacella clypeater*  
26 (Lesson 1831), *Nacella deaurata* (Gmelin 1791), *Nacella delicatissima* (Strebel 1907),  
27 *Nacella fuegiensis* (Reeve 1855), *Nacella flammea* (Gmelin 1791), *Nacella magellanica*  
28 (Gmelin 1791), *Nacella magellanica chilensis* (Reeve 1855), *Nacella magellanica venosa*  
29 (Reeve 1854) and *Nacella mytilina* (Helbling 1779). Criteria for classification into these  
30 species are all based on morphology, like shell shape and colour, radula morphology and  
31 tentacle pigmentation (see: Valdovinos and R uth 2005). It is yet not clear whether this  
32 classification of the different morphotypes into species is supported on a genetic level or

33 whether it is a cause of phenotypic plasticity. Several physiological studies were carried out  
34 and revealed differentiation between the two most conspicuous members of the genus, the  
35 intertidal *Nacella magellanica* and the subtidal *Nacella deaurata* (Malanga et al. 2004,  
36 Malanga et al. 2005, Gonzalez et al. 2008). The first attempt to unravel genetic differences  
37 between members of *Nacella* was undertaken by de Aranzamendi et al. (2009) using the ISSR  
38 technique (Inter simple sequence repeats). The authors could successfully demonstrate  
39 significant amounts of genetic differentiation between the species *Nacella magellanica*,  
40 *Nacella deaurata* and *Nacella mytilina*. All other species mentioned above were identical  
41 with *N. magellanica* or *N. deaurata* and therefore considered as morphotypes instead of being  
42 true species. However, multiallelic approaches like ISSR suffer from their inapplicability in  
43 many population genetic statistical tests that require biallelic markers. Therefore we  
44 developed eight polymorphic microsatellite markers to provide a tool for comprehensive  
45 studies on population structure, gene flow and demographic and evolutionary history of  
46 patagonian limpets from the genus *Nacella*.

47

48 Microsatellite isolation was carried out with individuals of *Nacella magellanica* and *Nacella*  
49 *deaurata* from Bahia Laredo and Bahia Gregorio, both sites located in the Central Magellan  
50 Strait, Chile. Genomic DNA was isolated from muscle tissue preserved in Ethanol using spin  
51 columns (QIAGEN DNeasy Mini Kit), applying the standard tissue protocol. For each species  
52 enriched microsatellites genomic libraries were produced using the reporter genome protocol  
53 by Nolte et al. (2005), modified by Held and Leese (2007). Single stranded DNA from *Mus*  
54 *musculus*, bound to Hybridization chips (Hybond N+, Healthcare), served as reporter genome  
55 probes. Enriched fragments were PCR-amplified, purified using the QIAGEN Qiaquick Kit  
56 and finally cloned into pCR2.1-TOPO vector and transformed into competent TOP10F'  
57 *Escherichia coli* (Invitrogen). After overnight growth in LB media positive clones were  
58 transferred to 96-well sequencing plates provided by GATC-Biotech (Konstanz, Germany)  
59 who also performed plasmid preparation of colonies and shotgun sequencing using a standard  
60 M13-forward primer.

61 Subsequent analyses of inserts containing microsatellites comprising vector clipping,  
62 redundancy detection and primer design were conducted using STAMP (Kraemer et al. 2009),  
63 a program pipeline based on the sequence analysis package STADEN (Staden 1996).  
64 Extensions to the basic program were the tandem repeat detection and analysis software  
65 PHOBOS (Mayer 2008 and the primer design program Primer 3 (Rozen and Skaletsky 2000).

66 Only inserts containing microsatellites with a perfection of 95% or higher were chosen using  
67 phobos and considered for primer design.

68 For *N. magellanica* 14 redundant inserts were found in 79 sequenced clones. The remaining  
69 65 unique inserts yielded 12 suitable loci, for which primers were designed using the  
70 multiplex option in PRIMER3 with a  $T_m$  of 55°C. For *N. deaurata* 9 redundant inserts were  
71 found in 87 sequenced clones. The remaining 78 unique inserts resulted in 12 suitable loci.  
72 Also here the multiplex option was applied with a  $T_m$  of 55°C.

73 Primer pairs were tested on a gradient PCR over a variety of annealing temperatures ranging  
74 from 48°C to 65°C. PCRs were carried out in total volume of 25µl, containing ~10 ng  
75 genomic DNA, 0.2 mM dNTPs, 0.5 µM primer, 0.5 M Betaine, 2.5 mM MgCl<sub>2</sub>, 0.03 U/µL  
76 Hotmaster Taq (Eppendorf). Following PCR conditions were applied: 2min 94°C, 32 cycles  
77 of 20sec at 94°C, 10 sec at different annealing temperatures, 60 sec at 65°C and a final  
78 extension of 45 min at 65°C.

79 For *N. magellanica*, 10 of the 12 loci produced distinct PCR products, for *N. deaurata* 11 out  
80 of 12. These remaining 21 loci were chosen as candidate loci and fluorescent labelled primers  
81 were developed with the dyes HEX and FAM. PCRs were repeated as described above using  
82 the labelled primers. PCR products were purified using ExoSAPit (Fermentas). Exonuclease I  
83 (Exo) degrades remaining primers and Shrimp Alkaline Phosphatase (SAP) inactivates  
84 remaining dNTPs. 5 µl of PCR products were mixed with 0.25 µl Exo I (20 U/µl) and 1 µl  
85 SAP (1 U/µl) and incubated at 37°C for 30 min. Enzyme activities were subsequently  
86 inactivated by an incubation step of 15 min at 80°C. The purified PCR products were  
87 denatured and analysed on an ABI 3130xl sequencer using ROX GS500 size standard (ABI).  
88 Genotyping was performed using the software genemapper 4.0.

89 For each species four microsatellite loci could be reliably genotyped. The remaining seven for  
90 *N. magellanica* and six for *N. deaurata* had to be excluded due to the presence of more than  
91 two alleles per individual or inconsistency during genotyping.

92 The remaining eight microsatellite loci were validated regarding their suitability for  
93 population genetic approaches. First the data were examined for possible scoring errors  
94 during the genotyping process using the software microchecker 2.2.3 (Van Oosterhout et al.  
95 2004). Diversity measures and deviations from Hardy-Weinberg equilibrium were tested  
96 using the program Arlequin 3.11 (Excoffier et al. 2005) and tests for linkage disequilibrium  
97 were computed using genepop 4.0.6 (Rousset 2008). The unbiased probability of identity was  
98 calculated using gimlet 1.3.3 (Valiere 2002).

99 Table 1 and 2 summarize the features of all eight polymorphic loci. Allelic diversity ranged  
100 from 5 (Nde3) to 57 (Nma6) alleles per locus. Observed heterozygosities varied between 0.1

101 (Nde3) and 0.98 (Nde2). After Bonferroni correction (Rice 1989) Locus Nde8 deviated from  
102 HWE ( $p < 0.05$ ) in the Falkland Population of *Nacella deaurata* and the Loci Nma6 and  
103 Nma12 In the Central Magellan Population of *Nacella magellanica*. These deviations could  
104 be caused by the presence of null alleles as suggested by microchecker. However, several  
105 other explanations have to be taken into consideration such as inbreeding and population  
106 expansion which also reduce heterozygosities in natural populations. No evidences for  
107 scoring errors caused by large allelic dropout or stuttering could be found. Global linkage  
108 disequilibrium analyses revealed no linkage between investigated Loci.

109

110 In addition, several cross amplification tests were carried out (Tab. 3). The four loci  
111 developed for *N. magellanica* were cross tested with individuals of *N. deaurata* and vice  
112 versa. Furthermore all eight loci were tested on the two nominal species *N. mytilina* and *N.*  
113 *delicatissima*. Two of the loci developed for *N. magellanica* also work with the other three  
114 species. Of the four loci developed for *N. deaurata* all work for *N. mytilina* and *N.*  
115 *delicatissima* and three work with *N. magellanica*. Furthermore, these amplification patterns  
116 enable us to correctly assign morphologically ambiguous specimens to the species they  
117 belong to.

118 In total we developed and provide here eight polymorphic loci of which seven are appropriate  
119 for population genetic studies with the South American limpets species *N. magellanica* and  
120 six for the species *N. deaurata*, *N. mytilina* and *N. delicatissima*. These markers enable us to  
121 develop a more profound classification of the genus *Nacella* and to study their evolutionary  
122 and demographic history.

Table 1: Genetic characterization of four polymorphic microsatellite loci isolated from individuals of *Nacella deaurata* collected on the Falkland Islands and the Central Magellan Strait region.  $N_a$  Number of alleles,  $T_a$  annealing temperature,  $H_o / H_e$  observed and expected Heterozygosity, PI probability of identity. Violations of Hardy Weinberg Equilibrium are shown in bold numbers (Bonferroni corrected 5% significant level = 0.00833)

Locus	Primer sequence (5'-3')	Dye	repeat motif	$N_a$	Size range (bp)	$T_a$ (°C)	n	Falklands		Central Magellan		
								$H_o / H_e$	PI (unbiased)	n	$H_o / H_e$	PI (unbiased)
<b>Nde1</b>	F: TAT CAA CGC ATC TTT CAT CA	Hex	(GA) <sub>18</sub>	22	213 – 234	57	52	0.92 / 0.89	1.960 x 10 <sup>-2</sup>	39	0.97 / 0.95	2.973 x 10 <sup>-3</sup>
	R: CAC GAT GTG TTG AGG TGT AG											
<b>Nde2</b>	F: TAG GTG TTA CGA GGA CGT TT	Fam	(CT) <sub>18</sub> (TC) <sub>7</sub>	25	154 - 218	57	52	0.98 / 0.92	7.703 x 10 <sup>-3</sup>	40	0.85 / 0.92	6.515 x 10 <sup>-3</sup>
	R: GAT CAA GAT TCA TCA GTG GC											
<b>Nde8</b>	F: TGT TGA TGA TGA AGG TGA TG	Hex	(GAG) <sub>6</sub> (GAA) <sub>2</sub> (GAG) <sub>3</sub>	19	108 - 151	57	46	<b>0.48 / 0.81</b>	4.133 x 10 <sup>-2</sup>	41	0.78 / 0.82	4.471 x 10 <sup>-2</sup>
	R: AGA GAG GAG CTA AAC CCA AT											
<b>Nde3</b>	F: TGA TTT AGA TAG GAG AGC GG	Hex	(AGAC) <sub>5</sub>	5	260 - 276	57	52	0.1 / 0.13	7.583 x 10 <sup>-1</sup>	42	0.12 / 0.11	7.788 x 10 <sup>-1</sup>
	R: AGG CTA AAT AAG CAT TGT CG											

Table 2: Genetic characterization of four polymorphic microsatellite loci isolated from individuals of *Nacella magellanica* collected from Seno Otway and the Central Magellan Strait region. N<sub>a</sub> Number of alleles, T<sub>a</sub> annealing temperature, H<sub>o</sub> / H<sub>e</sub> observed and expected Heterozygosity, PI probability of identity. Violations of Hardy Weinberg Equilibrium are shown in bold numbers (Bonferroni corrected 5% significant level = 0.00714)

Locus	Primer sequence (5'-3')	Dye	repeat motif	N <sub>a</sub>	Size range (bp)	T <sub>a</sub> (°C)	n	Seno Otway		Central Magellan		
								H <sub>o</sub> / H <sub>e</sub>	PI (unbiased)	n	H <sub>o</sub> / H <sub>e</sub>	PI (unbiased)
<b>Nma3</b>	F: ATG AAT CAA AAC TGT TGG CT R: TGC GCT ATG ACA TAC ACA TT	Hex	(C) <sub>14</sub> (CA) <sub>14</sub>	25	189 - 220	57	31	0.81 / 0.88	1.854 x 10 <sup>-2</sup>	46	0,83 / 0.87	2,348 x 10-2
<b>Nma4</b>	F: ATC TCC GCA GAT ACA AAC AA R: GGG TAT TGG TGA GAT GTG TT	Fam	(CA) <sub>7</sub> CG(CA) <sub>3</sub>	17	184 - 202	57	31	0.77 / 0.89	1.827 x 10 <sup>-2</sup>	46	0,83 / 0.89	1,682 x 10-2
<b>Nma6</b>	F: CTT TAG CAA AAT TGG TTT CG R: GGC AGG TTT GAC AGC TAA T	Hex	(CT) <sub>5</sub> /(CT) <sub>2</sub> GT(CT) <sub>3</sub> GT(CT) <sub>2</sub> / (CT) <sub>6</sub> TG(CT) <sub>21</sub> /(CT) <sub>5</sub>	57	192 - 324	57	31	0.84 / 0.97	3.036 x 10 <sup>-4</sup>	45	<b>0,73 / 0.95</b>	2,975 x 10-3
<b>Nma12</b>	F: TGT CAT CCG TCA AAA TGT TA R: TCT TCA ATG AGA CAA AAC CC	Fam	(GA) <sub>31</sub>	28	177 - 235	57	29	0.83 / 0.95	2.948 x 10 <sup>-3</sup>	44	<b>0,75 / 0.94</b>	4,400 x 10-3

Table 3: Cross amplification tests of the eight isolated microsatellite loci. The loci isolated from *Nacella magellanica* were cross tested with individuals of *Nacella deaurata* and vice versa. All eight loci were tested for cross amplification with individuals of *Nacella delicatissima* and *Nacella mytilina*

<b>Locus</b>	<b><i>Nacella magellanica</i></b>	<b><i>Nacella deaurata</i></b>	<b><i>Nacella mytilina</i></b>
<b>Nma3</b>	189 - 220	not amplified	not amplified
<b>Nma4</b>	184 - 202	187 - 205	191 - 203
<b>Nma6</b>	192 - 324	186 - 318	190 - 198
<b>Nma12</b>	177 - 235	not amplified	not amplified
<b>Nde1</b>	210 - 246	213 - 234	210 - 235
<b>Nde2</b>	not amplified	154 - 218	168 - 194
<b>Nde3</b>	260 - 280	260 - 276	260
<b>Nde8</b>	108 - 147	108 - 151	125 - 150

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