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To cite this article: Diana Aviles, Vincenzo Landi, Juan Vicente Delgado, José Luis Vega-Pla & Amparo Martinez (2015) Isolation and Characterisation of a Dinucleotide Microsatellite Set for a Parentage and Biodiversity Study in Domestic Guinea Pig (*Cavia Porcellus*), Italian Journal of Animal Science, 14:4, 3960, DOI: [10.4081/ijas.2015.3960](https://doi.org/10.4081/ijas.2015.3960)

To link to this article: <https://doi.org/10.4081/ijas.2015.3960>



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Published online: 14 Mar 2016.



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## PAPER

## Isolation and characterisation of a dinucleotide microsatellite set for a parentage and biodiversity study in domestic guinea pig (*Cavia porcellus*)

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### Abstract

The domestic guinea pig is a valuable genetic resource because it is part of local folklore and food tradition in many South American countries. The economic importance of the guinea pig is due to its high feed efficiency and the quality of animal protein produced. For these reasons, our study is aimed to design a complete dinucleotide microsatellite marker set following international recommendation to assess the genetic diversity and genealogy management of guinea pigs. We selected a total of 20 microsatellites, looking for laboratory efficiency and good statistical parameters. The set was tested in 100 unrelated individuals of guinea pigs from Ecuador, Peru, Colombia, Bolivia and Spain. Our results show a high degree of polymorphisms with a total of 216 alleles and a mean number of  $10.80 \pm 3.49$  for markers with a combined exclusion probability of 0.99.

### Introduction

The guinea pig (*Cavia porcellus*), also called *cavy*, is originally from the Andean regions of southern Colombia, Ecuador, Peru, and Bolivia, where the species was domesticated between 7000 and 5000 BC (Morales, 1995). Today, a stable population of 35 million animals is reared in this area (DAD-IS, 2014). The guinea pig has several uses and is a valuable economic resource for indigenous populations in the South American marginal areas where

they originate. The guinea pig is a unique source of food due to their ability to convert poor vegetable resources to protein. Additionally, the guinea pig has a strong presence in local folklore and in popular medicine and is an important resource in the cultural patrimony of local nations, especially the Quechuas and Aymaras. The guinea pig has been introduced to other countries since the Spanish colonisation of the American continent, and today, they are used as exotic pets or for scientific experimentation (Guerrini, 2003). Owing to its great capacity of growing and the poor feeding needs, many efforts have also been made to promote guinea pig husbandry in developing countries. The guinea pig was introduced in several West African countries. Even if no official statistics are available (Manjeli *et al.*, 1998), there are some stable reared populations in Cameroon, Democratic Republic of Congo and Tanzania (Maass *et al.*, 2005, 2010; Matthiesen *et al.*, 2011). To date, no complete genetic study has been carried out on the domestic guinea pig although great advances have been reached with the completion of the genomic sequence ([http://www.ensembl.org/Cavia\\_porcellus/Info/Index](http://www.ensembl.org/Cavia_porcellus/Info/Index); Broad Institute, 2015). Only a few studies have been conducted looking at microsatellites in guinea pigs, and they have centred on wild subspecies of the *Cavia* genus such as *Cavia aperea* and *Cavia magna* (Kanitz *et al.*, 2009) or have been limited to a small marker panel (Burgos-Paz *et al.*, 2011). The large number of guinea pig animals and breeds reared in South America necessitated the development of molecular tools to perform genetic characterizations and population structure studies as well as a parentage testing strategy for modern breeding approaches. To respond to this demand, the aims of our study were to design a polymorphic set of dinucleotide microsatellites useful both for analysing the genetic diversity of the domestic *Cavia* and as for parentage control, following the Food and Agriculture Organization (FAO) and International Society for Animal Genetics (ISAG) recommendations on this type of research in domestic animals.

### Materials and methods

#### Samples used and DNA extraction

Hair samples from a total of 100 unrelated animals belonging to several domestic guinea pig populations were used in our study. Some samples were collected from several breeding lines from Ecuador (40) divided in 10 sample for type/line (Andina, Peru, Inti and commer-

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**Key words:** *Cavia porcellus*; STRs; Biodiversity; Power of exclusion.

**Acknowledgements:** the authors wish to express thanks to the different breeders and research groups who kindly provided biological samples: Angelika Stemmer (University of San Simon, Cochabamba, Bolivia), Niltón Gómez (Universidad Nacional del Altiplano, Puno, Perú), Luz Angela Franco (Universidad Nacional de Colombia, Palmira, Colombia) and D. Carlos San José Marqués (BioDonostia, Spain). The authors gratefully thank the members of the CONBIAND network for valuable cooperation over the years.

**Funding:** the authors wish to acknowledge the financial support received by FUNDACION CAROLINA and the Programme Centro De Investigaciones CENI (Universidad Técnica de Ambato) for financial support for this project.

Received for publication: 15 March 2015.

Accepted for publication: 22 August 2015.

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Italian Journal of Animal Science 2015; 14:3960

doi:10.4081/ijas.2015.3960

cial local type) and others from Colombia (15), Bolivia (13) and Perú (15); also, some samples were collected in Spain from commercial lines (20) reared as pets. DNA was obtained by incubating 3 hair roots in the presence of 100 µL of 5% Chelex® (Biorad, Göttingen, Germany) resin suspension at 95°C for 10 minutes and 99°C for 3 min.

#### *In silico* identification of microsatellites and primer design

The cavPor3 (high-coverage 6.79X assembly) genome release of the guinea pig (*Cavia porcellus*) was used to search for microsatellite sequences ([http://www.ensembl.org/Cavia\\_porcellus/Info/Index](http://www.ensembl.org/Cavia_porcellus/Info/Index)) using the NCBI finder tool (Appendix Table 1). Sequence repeat motifs of  $\geq 18$  bp including poly AG, AC, AT, TC, CA, and GT were searched. A total of 25 sequences were selected. The primer pairs used for polymerase chain reaction (PCR)

amplification were designed using Primer3 software version 0.4.0 (Rozen and Skaletsky, 2000). Our parameter sets included an optimum primer size of  $20 \pm 5$  bp, an optimum melting temperature of  $\sim 60 \pm 5^\circ\text{C}$  and a GC content between 20 and 80%. The software was allowed to design primer pairs with expected PCR product sizes of 80 to 350 bp.

### Microsatellite locus selection

Our primer pairs were synthesised by Stabvida, Costa de Caparica (Portugal) without further modifications. PCR was performed separately for each locus in a reaction volume of 25  $\mu\text{L}$  containing  $\sim 10$ -30 ng of genomic DNA, 0.2  $\mu\text{M}$  each primer pair, 1X  $\text{NH}_4\text{SO}_4$  PCR buffer, 2.5 mM  $\text{MgCl}_2$ , 200  $\mu\text{M}$  each dNTP, and 1U Taq polymerase (AIDLAB, Beijing, China). The annealing temperature was  $56^\circ\text{C}$  for 35 cycles. PCR products were visualised on a 3% agarose gel, stained with ethidium bromide, in TBE buffer at 150 V/cm, using a 100-bp ladder as a reference (Thermo Fisher Scientific Inc., Waltham, MA, USA).

Based on the amplification efficiency and the absence of a nonspecific PCR product, the samples were sequenced using the BigDye cycle sequencing kit 2.0 (Life Technologies, Carlsbad, CA, USA), and the sequences were deposited in GenBank (Table 1) after sequencing a control sample from the original clone (Appendix Table 1). Additionally, four microsatellite loci (Kanitz *et al.*, 2009) were included in our study with some modifications

and discarding tetranucleotide repeat motifs loci (Table 1).

### Microsatellite typing

A final set of 20 polymorphic microsatellites was selected from the microsatellites we tested. The forward primer for each locus was 5' end labelled with fluorescent dye (Figure 1). PCR was performed separately for each locus in a final reaction volume of 25  $\mu\text{L}$  containing  $\sim 10$ -30 ng of genomic DNA, 0.2  $\mu\text{M}$  each primer pair, 1X  $\text{NH}_4\text{SO}_4/\text{KCl}$  PCR buffer, 3 mM  $\text{MgCl}_2$ , 200  $\mu\text{M}$  each dNTP, and 1U Taq polymerase (AIDLAB, Beijing, China). Multiplex reactions were performed following the size range and dye availability using ABI dye set D (Figure 1). The optimal annealing temperature was established by a gradient amplification of 8 samples (annealing temperature from 50 to  $62^\circ\text{C}$ ) on a Biometra Tgradient Thermal cycler (Biorad).

The sizes of the microsatellite alleles were visualised using an ABI PRISM 3130 Genetic Analyzer (Life Technologies), using a POP7 polymer and the internal size standard GeneScan500-Rox (Life Technologies). Genotypes were read with the ABI PRISM GeneScan 3.1.2 software (Applied Biosystems, Carlsbad, CA, USA) and interpreted with the ABI PRISM Genotyper 3.7 NT software (Applied Biosystems).

### Statistical analysis

The mean number of alleles, observed and

unbiased expected estimates of gene diversity, and their standard deviations, together with the polymorphic information content (PIC) were obtained using MICROSATELLITE TOOLKIT software (Park, 2001). We estimated non-exclusion probabilities considering the first (NE-1P), second (NE-2P) or parent pairs (NE-PP) and individual (NE-I) and sib identity (NE-SI) as well as the Hardy Weinberg Equilibrium (HWE), using Cervus software version 3.0.3 (Kalinowski *et al.*, 2007). The combined posterior probability (PEC) was calculated with the algorithm of Jamieson (1994). Deviations from HWE and  $F_{is}$  based on locus by locus AMOVA calculations were assessed using ARLEQUIN 3.5.1.3 (Excoffier and Lischer, 2010).

## Results

### Fluorescent polymerase chain reaction design and microsatellite genotyping

Based on amplification efficiency, success rate, and the absence of non-specific amplification of our primer pairs, a total of 16 microsatellites were selected for the panel design. We named these microsatellites CUY1, CUY2, CUY3, CUY4, CUY5, CUY6, CUY7, CUY8, CUY9, CUY10, CUY12, CUY16, CUY17, CUY18, CUY20, and CUY22. Additionally, 4 din-

**Table 1. Summary of the general characteristics of the twenty selected microsatellite loci.**

Locus	GB	RP	MX	Tm	SR	Forward primer (5'-3')	Reverse primer (5'-3')	Reference
CUY1	KP115879	GT	2	55	271-285	cttccagcgaataggcatcc	gcagcttgactacagagca	This work
CUY2	KP115880	CA	2	55	250-262	caagatccatcaacttctgt	tgttgcagatgctgcttt	This work
CUY3	KP115881	GT	1	55	212-252	gcaagtcaaatcatccctga	gagtcctccaagcaaaatc	This work
CUY4	KP115882	GT	2	55	210-230	tcatctgccttcagcatttg	aatggtcagggctaggatt	This work
CUY5	KP115883	CA	2	55	141-163	ggccaaagcaggaatgtcta	tagggcaagcattgatgatg	This work
CUY6	KP115884	CA	4	55	158-168	tggcttgccttctcttgg	ctgtgctcagcattgcattt	This work
CUY7	KP115885	CA	2	55	183-197	gatgcagtcagagagtgca	tgtgtggtttgtgtgtgagg	This work
CUY8	KP115886	TC	1	55	181-217	tgatgcacctgagaagtgg	ccaagtgtcttggctgctt	This work
CUY9	KP115887	GT	2	55	116-130	gctggaatgcaagcaagc	tgagtttcagctgtgatgag	This work
CUY10	KP115888	GT	1	55	106-128	ttccaagcatttcagaaaaca	tgacttcccaaccaaggaaa	This work
CUY12	KP115889	AG	4	55	232-250	ggaatggtggcaactccta	tctcctcctcctcctc	This work
CUY16	KP115890	AT	3	60	223-247	tttgagtcgaagcctgaaca	gcctgtttgaaactgtttactg	This work
CUY17	KP115891	TC	4	55	152-170	tgatggacaataactgggaacc	tagcatgatgaagccctaa	This work
CUY18	KP115892	CA	2	55	176-214	tgtcacttctcactccaca	tcccaaacctctgttctgt	This work
CUY20	KP115893	AT	4	55	218-258	tcttggaaatggcctacattt	tggtctctagggtatccatt	This work
CUY22	KP115894	TC	4	55	206-232	cgaacatgccaagcagatta	acaccagttccttgccacat	This work
Cavy2	AJ496560	AC	2	55	124-154	ggccattatgcccccaac	agctgctcctgtgctgtag	Kanitz <i>et al.</i> (2009)
Cavy3	AJ496561	CT	1	55	195-225	acagcgatcacaatctgcac	gcagtggaaccagcaatgg	Kanitz <i>et al.</i> (2009)
Cavy11	AC192015	CT	1	55	140-180	ccgtgcttttctgctcttg	tggaccacaatctgacatag	Kanitz <i>et al.</i> (2009)
Cavy12	AC182323	AG	1	55	143-187	agaatgccttgggactgg	agatctgctctgcacttg	Kanitz <i>et al.</i> (2009)

GB, GenBank accession number; RP, microsatellite repeat motif; MX, polymerase chain reaction multiplex reaction where the locus amplified; Tm, annealing temperature of polymerase chain reaction; SR, size range in base pairs.

ucleotide markers were selected from the Kanitz *et al.* (2009) based on sequence length and marker polymorphisms (Table 1) with no modification except for Cavy11 and Cavy 12, where the primer sequence was re-designed to improve the melting temperature parameter. A 4 colour system (ABI D Dye set) and a ~20 bp minimum predicted distance between loci was used to design the electrophoresis pattern. The unusually large distance between loci was designed because of a lack of references about this species, specifically information about expected allelic range. The panel of PCR amplification resulted in four PCR multiplexes divided into three electrophoresis sets (Figure 1). The gradient amplification resulted in an optimal hybridisation temperature, based on the broadness of the band, of  $55\pm 0.5^\circ\text{C}$  for all of the multiplexes, with the exception of the CUY 16 maker ( $60\pm 0.5^\circ\text{C}$ ).

### Marker polymorphism and quality

The allelic range (a region of the electropherogram where a locus specific allele can be found) we obtained was generally high. The mean difference between two alleles in the same individual ranged from 1.5 in CUY7 to 10.91 Cavy2.

A total of 216 alleles were found with a mean value of  $10.80\pm 3.49$ . All microsatellites were highly polymorphic with a minimum of 6 alleles (CUY6) and a maximum of 19 (Cavy12). The allelic richness ranged from a minimum of

4.002 for CUY9 and a maximum of 9.969 for Cavy12. We found observed and expected heterozygosity to have an average mean value of  $0.590\pm 0.115$  and  $0.778\pm 0.080$ , respectively, which is considered high (Table 2). To evaluate the polymorphisms of each marker, the PIC value was calculated and found to range from 0.503 for CUY9 and 0.902 for Cavy12. Deviations from HWE were found in 9 of the 20 loci (Appendix Table 2); Cavy12 and CUY7 were found in disequilibrium in 6 populations, CUY2, CUY10 and CUY17 ( $P<0.05$ ). The sample from Bolivia showed the highest number markers in disequilibrium (8) while the Spanish population showed the lowest ones (2).  $F_{is}$  values with a total mean value of 0.173.

### Panel set power statistics

In Table 3, the non-exclusion probability values are shown. The first two values (NE-1P and NE-2P) give the non-exclusion probability when the parents were considered one by one (the first parent and then the second parent of the opposite sex, respectively). In both cases, the higher value was for CUY9 (0.84 and 0.68), and the lower value was for Cavy12 (0.31 and 0.18). When parent pairs were considered, the results were comparable for identity and sibling identity non-exclusion probability, with a maximum value obtained for CUY9 (0.50, 0.25, and 0.54, respectively) and a lower probability for Cavy12 (0.05, 0.02 and 0.30, respectively).

Following the Jamieson (1994) algorithm

the combined posterior probability (PEC) was calculated (Table 3). The results show a high value for all types (0.99) but a smaller value for the sibling identity exclusion combined probability (0.84).

## Discussion

The aim of our study was to construct a polymorphic marker panel of microsatellites that would be useful for both genetic diversity studies and kinship and parentage analysis in *Cavia porcellus* populations. Microsatellites are very powerful genetic markers that can be used for identifying the genetic structure, pedigree analysis and genetic variation of closely related species. Until the present work, only a few studies had been carried out on wild guinea pigs using either a reduced microsatellite loci panel (Asher *et al.*, 2008; Kanitz *et al.*, 2009; Kouakou *et al.*, 2015) or AFLP loci (Burgos-Paz *et al.*, 2011). Some biodiversity studies have been carried out in Africa using the Kanitz *et al.* (2009) marker panel, such in Côte d'Ivoire (Kouakou *et al.*, 2015) although these authors did not find clear genetic differences among the three analysed populations. The most complete study on the genus *Cavia* was performed on mitochondrial DNA (Dunnum and Salazar-Bravo, 2010). Domestic

**Table 2. Descriptive statistics of the twenty designed microsatellite marker loci.**

Locus	NA	AR	Ho	He	PIC	HW	Fis
Cavy02	9	5.093	0.670	0.723	0.675	ns	-0.038
Cavy03	13	6.469	0.573	0.816	0.788	**	0.228
Cavy11	17	8.193	0.777	0.872	0.854	ns	0.068
Cavy12	18	9.969	0.500	0.913	0.902	nd	0.427
CUY01	8	5.196	0.588	0.756	0.718	*	0.076
CUY02	7	5.131	0.447	0.728	0.682	**	0.283
CUY03	11	5.997	0.650	0.790	0.756	ns	0.034
CUY04	9	5.832	0.500	0.708	0.680	**	0.186
CUY05	12	6.863	0.728	0.835	0.810	ns	0.097
CUY06	6	4.266	0.461	0.723	0.668	**	0.270
CUY07	7	4.551	0.373	0.690	0.639	**	0.457
CUY08	17	8.166	0.621	0.860	0.841	**	0.180
CUY09	7	4.002	0.398	0.547	0.503	ns	0.242
CUY10	11	6.299	0.573	0.790	0.760	ns	0.232
CUY12	9	5.278	0.703	0.754	0.716	ns	0.039
CUY16	11	7.229	0.767	0.829	0.808	ns	0.025
CUY17	10	6.857	0.713	0.840	0.816	ns	0.097
CUY18	10	6.287	0.578	0.800	0.770	**	0.212
CUY20	14	6.800	0.578	0.737	0.713	ns	0.178
CUY22	10	7.068	0.590	0.847	0.825	*	0.193
Mean±SD		6.270±1.430	0.590±0.115	0.778±0.080	0.750±0.100	**	0.173

NA, total number of alleles; AR, allelic richness; Ho, observed heterozygosity; He, expected heterozygosity; PIC, polymorphic information content; HW, deviation from Hardy-Weinberg equilibrium; Fis, fixation index within populations; SD, standard deviation. \* $P<0.05$ ; \*\* $P<0.01$ ; ns, not significant.

guinea pigs were included in these studies as an out-group. Our main objective was to compare the genetic diversity of the domestic guinea pig to the overall rearing area of the species. For this reason, we designed a panel of microsatellite markers to examine recent evolutionary events to infer the population structure and the genetic differentiation among different commercial lines and locally recognised guinea pig breeds. In addition, the importance of the guinea pig for the rural economy of several Latin American countries increases the need for molecular tools to further initiatives for their genealogical management and breeding design (Mommens *et al.*, 1998; Tozaki *et al.*, 2001; Bonnet *et al.*, 2002). Despite the diffusion into local communities and the low technological level needed for guinea pig farming, there exists intense commercial activity for these animals. Dinucleotide microsatellites are being used as genetic markers for the identification of population structure, genome mapping, and pedigree analysis and to resolve taxonomic ambiguities in many other animals in addition to the guinea pig (Xu and Liu, 2011; Martinez *et al.*, 2012; Gama *et al.*, 2013; Abdul-Muneer, 2014).

We successfully isolated, by scaffold genome sequencing, 25 microsatellite sequences, of which 16 were selected for the final panel based on their technical quality. All markers proposed here can be easily amplified in multiplex PCR reactions using crude sample lysates. Generally, all of the loci had a very high number of alleles ( $10.8 \pm 3.40$ ), which was higher than the values found by Kanitz *et al.* (2009) and Kouakou *et al.* (2015), as well as a high mean allelic range (25 bp). Even if only 11 loci out of 20 were in HWE in overall sample, the F index values were very high (0.173). These findings, despite the high number of alleles, can be explained by the small sample number used in this preliminary study, possibly leading to the maximization of heterozygous excess values (Wahlund, 1928), as highlighted also by the HW disequilibrium calculated by separated populations that showed a significant value for the sixth population only in the markers Cavy12 and CUY7. These results can be due by the particular mating system based on using inbreeding animals added to the great interchange of males and females in the country markets. The total combined exclusion probability highlighted that the 20 loci are enough to obtain a good efficiency for parentage testing and traceability purposes in this species.

## Conclusions

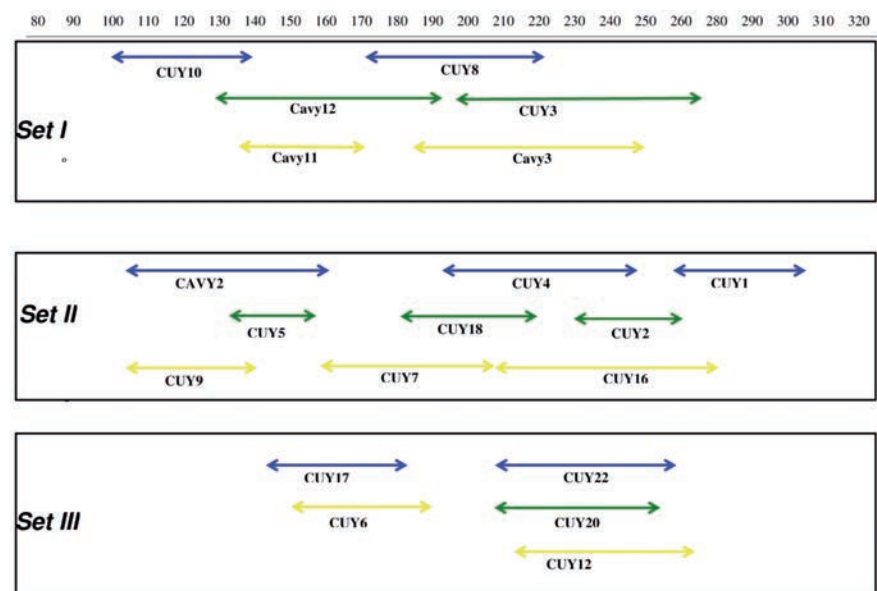
We have identified a set of 16 microsatellite loci for domestic *Cavia porcellus* genetic diver-

sity research, and we have also established their standardised genotype analysis parameters. These markers could potentially resolve parentage and individual assignment cases. The high degree of genetic diversity and poly-

**Table 3. Summary statistics for the non-exclusion probability values.**

Locus	NE-1P	NE-2P	NE-PP	NE-I	NE-SI
Cavy02	0.683	0.510	0.321	0.123	0.421
Cavy03	0.538	0.363	0.181	0.059	0.359
Cavy11	0.417	0.262	0.102	0.031	0.324
Cavy12	0.309	0.183	0.053	0.015	0.299
CUY01	0.640	0.460	0.269	0.096	0.398
CUY02	0.677	0.502	0.313	0.119	0.417
CUY03	0.586	0.409	0.222	0.076	0.376
CUY04	0.678	0.489	0.279	0.112	0.426
CUY05	0.502	0.332	0.156	0.049	0.347
CUY06	0.702	0.532	0.356	0.130	0.423
CUY07	0.723	0.552	0.365	0.146	0.443
CUY08	0.440	0.281	0.113	0.035	0.331
CUY09	0.836	0.676	0.499	0.249	0.540
CUY10	0.578	0.399	0.208	0.072	0.375
CUY12	0.643	0.462	0.271	0.097	0.399
CUY16	0.502	0.330	0.147	0.048	0.350
CUY17	0.493	0.323	0.149	0.047	0.344
CUY18	0.565	0.389	0.202	0.068	0.369
CUY20	0.632	0.446	0.234	0.092	0.406
CUY22	0.478	0.311	0.139	0.043	0.339
PEC	0.99	0.99	0.99	0.84	0.99

NE-1P, non-exclusion of one candidate parent; NE-2P, candidate parent given the genotype of a known parent of the opposite sex; NE-PP, candidate parent pair; NE-I, identity of two unrelated individuals; NE-SI, identity of two siblings; PEC, combined exclusion probability calculated using the Jamieson formula (Jamieson, 1994).



**Figure 1. Electrophoresis set up of the twenty *Cavia porcellus* microsatellites based on allelic range and fluorescent dye for ABI D set. Blue=6FAM; Green=HEX; Yellow=ATTO550 (ROX as internal ladder).**

morphisms indicate the potential of this microsatellite panel to be employed in future extended studies on the biodiversity of the cavy population. Therefore, genotype analyses with these standardised microsatellite panels will enhance cavy genetic selection by providing individual identification to increase the precision of measured phenotypes and for the construction of pedigrees to support the measurement of genetic estimates of phenotypic variation across generations.

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**APPENDIX**

**Appendix Table 1. Total primers pair designed in available scaffold sequence of *Cavia porcellus*.**

Origin sequence description	Accession number	Repeat motive <sup>o</sup>	Amplicon length	Forward oligo	Reverse oligo
<i>Cavia porcellus</i> clone CH234-497P15, complete sequence	AC171739.3	GT(18)	273	cttccaggcaataggcatcc	gcagctggactacagagca
<i>Cavia porcellus</i> clone CH234-9K24, complete sequence	AC173430.3	CA (22)	258	caagatgccatcaactttcgt	tgttgcgatgatgctctt
<i>Cavia porcellus</i> , clone <i>Cavia porcellus</i> -24932957J7, complete sequence	AC165221.3	GT (18)	228	gcaagtcaaattcatccctga	gagtcctccaagcaaatc
<i>Cavia porcellus</i> clone CH234-14J14, complete sequence	AC175208.3	GT(22)	223	tcattcgcctcagcatttg	aatggcaggggctaggatt
<i>Cavia porcellus</i> clone CH234-497P15, complete sequence	AC171739.3	CA(20)	156	ggccaaagcaggaatgcta	tagggcaagcattgatgatg
<i>Cavia porcellus</i> clone CH234-9K24, complete sequence	AC173430.3	CA(18)	162	tggcttgcttctcttggg	ctgtgctcagcattgcatt
<i>Cavia porcellus</i> clone B64 microsatellite sequence	GU045442.1	CA(18)	187	gatgcagtcagaggagtc	tgtgtggtttgtgtgagg
<i>Cavia porcellus</i> clone CH234-402D11, complete sequence	AC175781.3	TC(21)	190	tgattgcacctgagaagtg	ccaagtgttctgtgcttg
<i>Cavia porcellus</i> clone CH234-334G9, complete sequence	AC181988.3	GT(18)	120	gctggaaatgcaagacaagc	tgagtttcagctgtgatgag
<i>Cavia porcellus</i> clone CH234-9K24, complete sequence	AC173430.3	GT(21)	117	ttccaagcattcagaaaaca	tgactccaaccaaggaaa
<i>Cavia porcellus</i> clone C15 microsatellite sequence	GU045440.1	TG(20)	156	aaaatgttccatggggatg	gcatgtgttatcgcgtctg
<i>Cavia porcellus</i> clone CH234-34N9, complete sequence	AC174609.3	AG(28)	242	ggaatggtggcaactccta	tctctcctcctcctcttc
<i>Cavia porcellus</i> clone CH234-34N9, complete sequence	AC174609.3	AG(24)	273	tgccaaatgagaaatgagta	ggggtaatggcaatgtgtc
<i>Cavia porcellus</i> clone CH234-386E16, complete sequence	AC216606.3	CA(22)	250	agcaagaggcacacaagtc	ggggtaatggcaatgtgtc
<i>Cavia porcellus</i> clone CH234-14J14, complete sequence	AC175208.3	AG(25)	153	aaagcttggactgcaaga	ttcctcctcctcctctcc
<i>Cavia porcellus</i> clone CH234-14J14, complete sequence	AC175208.3	AT(25)	248	tttgagtcaagccgtgaaca	gctggtttgaaactgtttactg
<i>Cavia porcellus</i> clone CH234-33F4, complete sequence	AC174824.3	TC(19)	154	tgatggacaatatactgggaacc	tagcatgcatgaagccctaa
<i>Cavia porcellus</i> clone CH234-33F4, complete sequence	AC174824.3	CA(21)	210	tgcaacttctcactccacca	tcccaaacctctgtttgtct
<i>Cavia porcellus</i> clone CH234-261L8, complete sequence	AC181987.3	TC(22)	196	tcccaaggctgagcatac	tggtcaaatgtcttcatgtg
<i>Cavia porcellus</i> clone CH234-261L8, complete sequence	AC181987.3	AT(22)	231	tcttggaaatggcctacattt	tggtctctagggtatccatt
<i>Cavia porcellus</i> clone CH234-176E17, complete sequence	AC171368.3	TC(27)	262	atcttctgccctctctc	tggtgccacacactgtaat
<i>Cavia porcellus</i> clone CH234-487I3, complete sequence	AC171142.3	TC(21)	248	cgaacatgcccaagcagatta	acaccagtctctgccacat
<i>Cavia porcellus</i> clone CH234-487I3, complete sequence	AC171142.3	CA(21)	195	gcaaatgtgccatctgtgt	aagtgtgtttggggattt
<i>Cavia porcellus</i> clone CH234-176E17, complete sequence	AC171368.3	CA(27)	222	tgctgcagcctcttgaata	ccacagtggttaaatgatcgag
<i>Cavia porcellus</i> clone CH234-497P15, complete sequence	AC171739.3	CA(23)	108	aaatgcctacagcaacattc	tttatggccacagagagac

<sup>o</sup>Repeat sequence length in National Center for Biotechnology Information sequence.

**Appendix Table 2. Hardy Weinberg disequilibrium P value significance for each population.**

	Andean line	Inti line	Peru line	Commercial line	Peru	Bolivia	Colombia	Spain
CAVY02								
CAVY03							*	
CAVY11						**		
CAVY12	**		**	**	**	**	**	
CUY01								
CUY02				*		**	*	**
CUY03						**		
CUY04				*				
CUY05								
CUY06				**		*		
CUY07	*	*	*		**	**		
CUY08				*			**	**
CUY09		*						
CUY10				**		**	**	
CUY12								
CUY16								
CUY17	*					*	**	
CUY18								
CUY20								
CUY22								

\*P<0.05; \*\*P<0.01.