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### Genetic Diversity of Gorontalo Local Cattle Based on Microsatellite DNA

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Journal	Advances in Animal and Veterinary Sciences					
Manuscript ID	MH20221207091228					
Manuscript Type	Research Article					
Area of Interest	Husbandry					
Date Submitted by the Author	Wed, 07 Dec 2022, 10:05 AM					
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#### ABSTRACT

19 Microsatellites are one of an option for characterizing cattle populations, those information can be used as a basis for the development and maintenance of local cattle. This 20 study aims to evaluate genetic diversity and relationships using microsatellite markers 21 between the populations of Gorontalo cattle in Gorontalo Province, Indonesia using 3 22 microsatellite markers. A total of 126 samples of local bovine blood were collected, which 23 consisted of North Gorontalo (n=28, Gorontalo (n=52), Bonebolango (n=37), PO (n=3), and 24 Bali (n=6), were used in this study. The Genomic DNA Mini Kits was used for DNA 25 extraction for analysis fragment in microsatellite DNA region using ILSTS017, HEL13 and 26 BM1818 primer. A total of 74 alleles were identified through entire populations. The 27 expected heterozygosity ranged from 0.407±0.216 (Bali) to 0.716±0.050 (Gorontalo) and the 28 observed heterozygosity ranged from 0.471±0.084 (Bonebolango) to 0.778±0.222 (PO). F 29 30 statistical analysis includes F<sub>IS</sub> 0.038, F<sub>IT</sub> 0.248, and F<sub>ST</sub> 0.231. The 3 microsatellite markers were moderate (0.25-0.5) to highly informative (PIC>0.5) The research showed that Bali 31 cattle were distinct from all other cattle populations, while Gorontalo-Bonebolango 32 admixture and North Gorontalo cattle were mixed with PO cattle population. In conclusion, 33 markers used were highly informative and polymorphic in investigating genetic diversity in 34 Bonebolango, Gorontalo and North Gorontalo Populations, while the genetic relationship 35 among cattle populations was divided into two main clusters i.e cluster Bali and PO 36 populations, closely reflects the breeding process state on research area. These information 37 will be useful for future development and maintenance of local cattle in Gorontalo Province, 38 Indonesia 39

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Key words: Microsatellite, Genetic Diversity, Genetic relationships, Local Cattle, Gorontalo

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### INTRODUCTION

The increases demand for livestock products as human food illustrates an increase human quality life, but challenges the genetic availability of livestock. The development of local livestock as a provider of genetic resources needs to be preserved, because local livestock have ability to adapt to their's own environment. This genetic ability is expected to produce more productive livestock (Gobel et al, 2021; Pribadi L W, 2015; Ilham et al., 2016)

Gorontalo Province is located in Indonesia, specifically between Central Sulawesi and 48 North Sulawesi. The potential for cattle in Gorontalo Province is 192,229 heads (BPS 2020). 49 50 There are several types of cattle that are well developed in the Gorontalo region, such as: Bali cattle, PO, Brahman cattle, and other local cattle that have not been identified. Local cattle 51 that have not been identified are often called Diiti/ordinary/Local cattle. The existence of 52 local cattle has a different phenotype appearance, compared to Bali and PO cattle. Naturally, 53 livestock that adapt to the environment will bring up different phenotype variations in the 54 population. Muladno (1994) state that genetic change can be used for the process of evolution 55 of livestock populations/breeds by looking at the genetic distance of a population. 56 Adaptability of local livestock is the ability of gene expression resulting in gene expression 57 so that these livestock can adapt well to that environment and are able to produce fertile 58 offspring. Efforts to maintain the genetic abilities that are formed in local cattle require 59 adequate information, from qualitative characteristics, morphometric, productivity and 60 61 genetic information, so that these local livestock can be developed in the future.

Microsatellites are an option for characterizing cattle breeds or populations (Metta et al., 2004; Rincon et al., 2007), also used to answer questions related to genetic diversity and genetic relationships among cattle populations (Rincon et al., 2007; Sun et al. al., 2008; Chaudhari et al., 2009). Microsatellite loci are widely used because of high polymorphism, co-dominance, and relatively abundant in the genome (Rincon et al., 2007; Karthickeyan et

al., 2009). The study of microsatellite DNA diversity in local Indonesian cattle has been 67 reported previously (Satriani et al., 2002; Sarbaini, 2004; Winaya et al., 2007; Abdullah, 68 2008; Sutarno et al., 2015; Septian and Sumantri, 2015; Agung et al., 2016; Agung et al., 69 70 2019), but has never been reported in local cattle in Gorontalo. It is necessary to have an overview of the genetic diversity analysis of local Gorontalo cattle based on microsatellite 71 DNA analysis. The importance of this information can be used as a basis for the development 72 and maintenance of local cattle. This study aims to evaluate genetic diversity and 73 relationships using microsatellite markers between the populations of Gorontalo cattle in 74 75 Gorontalo Province, Indonesia.

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### **MATERIALS AND METHODS**

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#### **Blood Samples and DNA Isolation**

All procedures related to animal use in this study were approved by the Animal Care 78 79 and Use Committee of Brawijaya University under regulation number 145-KEP-UB-2022 (Ethical Clearance). A total of 126 samples of local bovine blood were collected from 3 80 different regions, in Gorontalo (n=52), Bonebolango (n=37) and North Gorontalo districts 81 (n=28), plus the PO (n=3) and Bali (n=6) population samples in the Gorontalo province, were 82 used in this study. Meanwhile, blood samples were analyzed at the Biotechnology 83 Laboratory, Faculty of Animal Science, Universitas Brawijaya, Malang. Blood was collected 84 from the vena coxygealis of the cattle. The DNA was isolated using Genomic DNA Mini 85 Kits. The DNA isolation procedures followed the protocol instructions. 86

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### Primer and DNA Amplification

A total of 3 bovine microsatellite markers were used in the PCR process. The markers consisted of ILSTS017, HEL13 and BM1818 (Table 1). The PCR reaction contained the mixture of 1  $\mu$ L DNA template (50-100 ng/ $\mu$ L) and 30  $\mu$ L PCR premix. The PCR premix consisted of 0.4  $\mu$ L primer (10 pmol/ $\mu$ L), 15  $\mu$ L Go Taq Green Master Mix (Promega,

USA), and 14.2 µL Nucleus Free Water (NFW). The PCR thermal cycler conditions were
shown in Table 2. The PCR product visualization methods were based on Susilorini *et al.*,
(2022). Microsatellite fragment analysis was conducted at First Base Laboratory, Selangor,
Malaysia.

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### Microsatellite Data Analysis

The fragment analysis result was processed to get allele size data. Allele data then 97 converted using the Convert version 1.31 for further analysis. The converted data were 98 analyzed using Cervus version 3.0.7 to generate observed heterozygosity (HO), expected 99 100 heterozygosity (HE), frequency/number of alleles, polymorphism information content (PIC) values and Hardy-Weinberg (HW) equilibrium. GenAlEx 6.51 b2 version was used to 101 generate genetic differentiation (F<sub>ST</sub>), the rate of inbreeding between populations (F<sub>IS</sub>), the 102 103 rate of inbreeding in populations (F<sub>IT</sub>) and Principal coordinate analysis (PCoA) to determine the genetic relationship between livestock breeds. Genetic structures were analyzed using 104 POPTREEW (POPTREEW website version) (Takezaki et al., 2014) to generate the 105 reconstruction of phylogeny trees between population and genetic distance. 106

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#### **RESULT AND DISCUSSION**

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#### **Microsatellite Polymorphisms and Genetic Diversity**

109 Studies on determination of genetic diversity on Gorontalo cattle based on 110 microsatellite markers aren't available, so this study was conducted to find information about 111 its diversity. Polymorphism of molecular genetics and F-statistic belonging to each locus are 112 summarizes in Table 3 and Table 4. A total of 74 alleles were detected through the 3 markers 113 analyzed in entire populations; 8 alleles detected in Bali cattle, 9 alleles in PO cattle, 20 114 alleles in Bonebolango cattle, 21 alleles in Gorontalo cattle and 16 alleles in North Gorontalo 115 cattle. The observed heterozygosity (OH) was 0.474, 0.407, 0.737 and the expected

heterozygosity (EH) was 0.712, 0.658, and 0.829 for ILSTS017, HEL013 and BM1818, respectively. Genetic diversity belonging to each population was shown in Table 5. EH ranged from  $0.407\pm0.216$  (Bali) to  $0.716\pm0.050$  (Gorontalo), whereas OH ranged from  $0.471\pm0.084$  (Bonebolango) to  $0.778\pm0.222$  (PO). The EH value was higher than OH in all populations, except in PO and Bali populations.

Expected heterozygosity in all loci showed a higher value than observed 121 heterozygosity, indicated that all loci were deviated from HWE. The deviation of loci from 122 HWE can be caused by several factors such as selection, genetic drift, inbreeding, the 123 presence of null alleles, selection in favor of homozygote and Wahlund effect (Rahal et al., 124 2020). The EH value was lower compare to Algerian cattle (Rahal et al., 2020), but higher 125 than previous study in Indonesian cattle (Sutarno et al., 2015). The highest level of 126 heterozygosity was in locus BM1818 was higher than previous research in Red Steppe cattle 127 using BM1818 loci which obtained heterozygosity value of 0.692 (OH) and 0.701 (EH) 128 (Kramarenko et al., 2018). The highest heterozygosity in Gorontalo population (0.716±0.050) 129 could be explained due to the lack of breeding program. The EH > OH value indicates low 130 heterozygosity. Kriswiyanti (2014) states that heterozygosity influenced by the variety of 131 alleles and the frequency of each allele at each locus. Heterozygosity values range from 0 to 132 133 1. If heterozygosity equals 0 (zero), then the population being measured has a very close genetic relationship, and if the value is equal to 1 (one), then the population being measured 134 has no genetic relationship. 135

F-statistics consisting of FIS, FIT and FST were 0,038, 0248 and 0.231, respectively. All microsatellite loci were deviated from the Hardy-Weinberg Equilibrium (HWE), meanwhile for all loci in each population, only locus ILSTS and HEL013 in Gorontalo cattle showed the deviation from HWE. The presence of null alleles in the study was between 6.15% (BM1818) and 24.01% (HEL013). The F<sub>ST</sub> value of 23.1% indicated that 23.1% of the

total genetic variation was due to alleles differentiation between breed, while the remaining 141 76.9% due to the difference among individuals within the breed across the 3 markers. The 142 average of 0.231 F<sub>ST</sub> value indicated that the genetic difference in the population was high 143 (0.15-0.25). Hartl and Clark (2007) state that F<sub>ST</sub> of 0-0.05 (small), 0.05-0.15 (medium), 144 0.15-0.25 (high) and >0.25 (very high). Obtained  $F_{ST}$  value were higher than Egyptian cattle 145 (El Saveed et al., 2016), Algerian cattle (Rahal et al., 2020), South African Nguni cattle 146 (Sanarana et al., 2016) and cattle raised in Turkey (Demis and Balciogly, 2019), but these 147 value were lower compare than previous research in Indonesian cattle (Agung et al., 2019). 148 The F<sub>IS</sub> value was 0.412, 0.067 and -0.017 for ILSTS017, HEL013 and BM1818, 149 respectively. The obtained F<sub>IS</sub> value was used to get a deeper insight about inbreeding degree 150 and endangered potentiality. F<sub>IS</sub> value less than 0.05 means the breeds are not in danger, 0.05-151 0.15 means the breeds potentially endangered, 0.15-0.25 means the breeds minimally 152 endangered, 0.25-0.40 means the breeds endangered, and >0.40 means the breeds critically 153 endangered (Simon and Bchenauer, 1993; El Sayeed et al., 2016). 154

PIC value were found polymorphic between 0.601 (HEL013) and 0.803 (BM1818). 155 The PIC value based on each population ranged from 0.353 to 0.674 which indicated as 156 moderate to highly informative value. In the present study, PIC value of three markers were 157 highly informative (>0.5), meanwhile PIC value for ILSTS017 loci in Bali and PO 158 populations had low informative value (0.25-0.5). The PIC values in previous research on 159 Egyptian cattle were 0.45 and 0.64 (El Sayeed et al., 2016), Taro White cattle was 0.536 160 (Hervani et al., 2019), Zimbabwean Sanga cattle was 0.666 - 0.682 (Gororo et al., 2018), and 161 that of Eastern European cattle was 0.752 (Illie et al., 2015). The PIC result showed that the 162 markers are highly informative for characterization of three populations used in study, i.e. 163 Bonebolango, Gorontalo and North Gorontalo populations. Botstein et al., (1980) state that 164 PIC value should be > 0.5 for identifications of genetic diversity. 165

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### **Genetic Structure and Relationships**

The result of genetic distance was a basic data to create the visualization of phylogeny 167 tree (Table 6) revealed the closest genetic distance between Gorontalo - Bonebolango 168 (0.037), following by Gorontalo – North Gorontalo (0.043) and Bonebolango – North 169 Gorontalo (0.078), respectively, while the farthest genetic distance between Bali - PO 170 (2.109). Estimation of pair-wise FST value revealed the highest genetic differentiation 171 between PO and Bali (0.206) and the lowest genetic differentiation between Bonebolango -172 North Gorontalo (0.011), Gorontalo – Nort Gorontalo (0.012), Bonebolango – Gorontalo 173 174 (0.013) and PO – North Gorontalo (0.029). Low genetic differentiation indicated high genetic similarity between populations. 175

Genetic distance analysis shows that the genetic distance between populations ranged 176 from 0.037 to 2.109, while the pair-wise FST value ranged from 0.011 to 0.206. The 177 reconstruction of phylogeny tree based on Nei's genetic distance was shown in Figure 1. The 178 figure revealed two main clusters of cattle populations. The first cluster consisted by Bali 179 cattle, the second cluster consisted of PO cattle. Cluster of PO population were divided into 180 two subpopulations, i.e North Gorontalo population and Bonebolango - Gorontalo 181 population. The principal coordinate analysis (PCoA) result supported the visualization of 182 183 phylogeny tree. It is shown that Bali cattle clearly separate from other cattle, while PO cattle population form a cluster which is contain local cattle in three different population as 184 subpopulations. 185

The relationship among five cattle populations using reconstruction of phylogeny tree indicated a high share of gene between Gorontalo – Bonebolango. The PCoA analysis confirmed that Bali populations is separated from other studied populations, while Gorontalo – Bonebolango admixture and North Gorontalo were mixed with PO population. This is due to the distribution pattern of the arrival of cattle to Gorontalo province starting from North

Gorontalo, and then the incoming cattle will be selected. After that, the cattle will be 191 distributed to other areas such as Gorontalo and Bonebolango. Cattle that entered North 192 Sulawesi province (before Gorontalo province separated from North Sulawesi province) were 193 dominated by PO cattle. Then in 2000 (UU No. 38 of 2000) when Gorontalo officially 194 became a separate province from North Sulawesi, the province of Gorontalo was dominated 195 by Bali cattle. This distribution pattern allows for differences between the cattle population in 196 North Gorontalo and other areas. The similarity in cattle between the populations of 197 Gorontalo and Bonebolango can be due to the fact that the two regions are close together, 198 199 which allows for interbreeding between livestock and causes the appearance of the same genetics. 200

The population of Bali and PO cattle form a different cluster. This is because Bali and 201 PO cattle are different species, where Bali cattle are included in Bos Sondaicus, while PO 202 203 cattle are in Bos Indicus. The already existing of PO cattle for more than 2 decades can cause a high share of genetic pool between PO cattle and local cattle in Gorontalo province. 204 Samples of cattle in the population of North Gorontalo, Gorontalo and Bonebolango are local 205 cattle in each of these regions. The phenotypic appearance of the three cattle in the area has a 206 dominant mixture of PO phenotype, a little of Bali phenotype and both cattles phenotypes 207 (Suvadi et al., 2014; Lava et al, 2020; Domili et al, 2021; Gobel et al, 2021). However, they 208 are not said to be PO and Bali cattle (Figure 4, 5 & 6), but as local cattle for each region. The 209 study could be as guidelines for future genetic studies on the development and maintenance 210 of local cattle in Gorontalo Province, Indonesia. 211

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#### CONCLUSION

Three markers used were highly informative and polymorphic in investigating geneticdiversity on three cattle populations in Gorontalo Province, i.e Bonebolango, Gorontalo and

North Gorontalo. There are 2 main classified clusters, i.e., (1) Bali cattle were distinct from all other cattle populations, and (2) PO population as second main cluster consist of North Gorontalo, Bonebolango and Gorontalo subpopulations, closely reflects the breeding process state on research area. These information will be useful for future development and maintenance of local cattle in Gorontalo Province, Indonesia.

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### ACKNOWLEDGEMENTS

The authors thank the team of Department of Animal Husbandry, Faculty of Agriculture, Gorontalo University for supporting the data and the member of Genomic and Proteomic Research Group, Faculty of Animal Science, University of Brawijaya for data analysis.

225	AUTHORS CONTRIBUTIONS
226	SD, SS, VMAN & GC: Idea and design. SD, NKL, SIG, AA & DW: Material sample
227	collection & lab analysis. DW, SD & NKL: Write the manuscript. SD & SS: Revision
228	CONFLICT OF INTEREST
229	The authors confirm that there is no conflict of interest in the manuscript.
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**Table 1.** Bovine microsatellite markers information used in study.

Marker	Chromosome	Primer sequences (5'>3')	Primer	Length of	Genebank
		forward	attachment	DNA base	access
		reverse	temperatures	(Bp)	

			(°C)		
ILSTS017				105-125	
HEL13	11	TAAGGACTTGAGATAAGGAG	52-57	178-200	X65207
(D11S15)		CCATCTACCTCCATCTTAAC			
BM1818	23	AGCTGGGAATATAACCAAAGG	56-60	248-278	G18391
(D23S21)		AGTGCTTTCAAGGTCCATGC			

350

### **Table 2**. The PCR thermal cycler condition

	PCR Step	Temperature (°C)	Time	Number of Cycle
	Pre-denaturation	95	5 minutes	Once
	Denaturation	95	30 s	
	Annealing	58	45 s	35 times
	Extension	72	1 minute	
	Final extension	72	5 minutes	Once
352				
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### **359 Table 3.** F Statistics (F<sub>IS</sub>, F<sub>IT</sub>, F<sub>ST</sub>) between five cattle population.

Lokus	Ν	Ho	He	PIC	HW	FIS	FIT	Fst	Nm
ILSTS017	95	0.474	0.712	0.699	***	0.412	0.530	0.200	0.999
HEL013	86	0.407	0.658	0.601	**	0.067	0.214	0.157	1.342

	BM1818	95	0.737	0.803	0.803	***	-0.017	0.157	0.068	3.442
	Mean						0.154	0.265	0.142	1.928
	SD						0.131	0.140	0.039	0.764
360	Note: N=nu	umber of	sample;	Ho=obse	erved h	eterozygo	sity; He=	expected	heterozy	gosity;
361	PIC=p	olymorphis	sm informa	tion cont	ent; HW	= Hardy-	Weinberg	Equilibriu	ım; FIS=	inter-
362	popula	ation inb	reeding r	ate; FIT=	= inbree	ding rat	te in po	pulation;	FST= g	genetic
363	differe	entiation; 1	Nm=gene fl	ow.						
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272										
373	Table 4 Nue	mber of all	eles (Na)	bserved h	eterozvo	sity (Ho)	and expec	ted (He)	and	
375	Polymorphis	n Informat	tion Conten	t in five ca	attle popu	lation in e	ach loci.	, (110 <i>)</i> , t		
	P		Loku	<u>c</u> N	No	No	PIC	Но	п	

Рор	Lokus	Ν	Na	Ne	PIC	Но	Не	HW
Bali	ILSTS017	6	3	1.946	0.424	0.167	0.530	ND
	HEL013	6	5	3.600	0.680	0.833	0.788	ND

	BM1818	6	6	5.143	0.777	0.833	0.879	ND
РО	ILSTS017	3	2	1.800	0.346	0.000	0.533	ND
	HEL013	3	3	2.571	0.535	1.000	0.733	ND
	BM1818	3	4	3.600	0.671	1.000	0.867	ND
Bolongo	ILSTS017	22	8	2.907	0.627	0.364	0.671	ND
	HEL013	17	4	2.429	0.513	0.412	0.606	ND
	BM1818	22	8	4.523	0.754	0.636	0.797	ND
Gorontalo	ILSTS017	41	9	3.543	0.682	0.488	0.727	*
	HEL013	37	5	2.700	0.564	0.351	0.638	*
	BM1818	41	7	5.033	0.777	0.732	0.811	ND
Gorontalo Utara	ILSTS017	23	4	2.574	0.553	0.696	0.625	ND
	HEL013	23	3	2.266	0.489	0.304	0.571	ND
	BM1818	23	9	5.290	0.785	0.783	0.829	ND

376 Note: N=number of sample; Na=Number of alleles; Ne= Mean number of effective alleles;
377 Ho=observed heterozygosity; He=expected heterozygosity; PIC=polymorphism information
378 content; HW= Hardy-Weinberg Equilibrium.

379

381 Table 5. Number of alleles (Na), observed heterozygosity (Ho), and expected (He), and
382 Polymorphism Information Content in five cattle breeds observed

Breed	Ν	Na±SD	Ne±SD	Ho±SD	He±SD	PIC
А	6	4.67±0.88	3.563±0.923	0.611±0.222	0.671±0.096	0.627
В	3	3.00±0.58	2.657±0.521	0.667±0.333	0.593±0.081	0.518
С	22	6.68±1.33	3.286±0.634	$0.471 \pm 0.084$	$0.674 \pm 0.056$	0.631
D	41	7.00±1.16	3.759±0.682	0.524±0.111	0.716±0.050	0.674

	Е	23	5.33±1.86	3.377±0.961	0.594±0.147	$0.660 \pm 0.077$	0.609
383	Note: A=	=Bali cattle;	B=PO cattle;	C=Bolongo; D	=Gorontalo; E=N	orth Gorontalo; N	N=number of
384	sar	nple; Na=N	Number of all	eles; Ne= Mea	an number of e	ffective alleles;	Ho=observed
385	het	terozygosity	; He=expected	d heterozygosi	ty; PIC=polymo	rphism informati	on content;
386	SD	=standart de	eviation.				

Table 6. Population pair-wise Fst (bottom diagonal) and Nei's standard genetic distance (top diagonal) 

U	,				
Populasi	Bali	РО	Bolongo	Gorontalo	Gorontalo Utara
Bali	***	0.053	0.959	0.714	0.807
РО	0.015	***	1.233	0.931	1.030
Bolongo	0.134	0.174	***	0.037	0.078
Gorontalo	0.104	0.142	0.008	***	0.043
Gorontalo	0.128	0.166	0.017	0.010	***
Utara					



Figure 1. The reconstruction of the UPGMA phylogeny tree of Gorontalo cattle populations with Nei genetic distance. 

395



399 Gorontalo Utara= <sup>×</sup>



401 Figure 3. Principal coordinate analysis (PCoA) based on each population.

402



404 Figure 4. Local cattle population of Gorontalo Regency



407 Figure 5. Local cattle population of North Gorontalo

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410 Figure 6. Local cattle of the Bonebolango population



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### Genetic Diversity of Gorontalo Local Cattle Based on Microsatellite DNA

Journal	Advances in Animal and Veterinary Sciences					
Manuscript ID	MH20221207091228-R2					
Manuscript Type	Research Article					
Area of Interest	Husbandry					
Complete List of Authors:	<ul> <li>Dr Safriyanto Dako, Department of Animal Husbandry, Faculty of Agriculture, Gorontalo University, Indonesia</li> <li>Mrs Nibras Laya, Department of Animal Husbandry, Faculty of Agriculture, Gorontalo University, Indonesia</li> <li>Mr Syukri Gubali, Department of Animal Husbandry, Faculty of Agriculture, Gorontalo University, Indonesia</li> <li>Mr Ari Ardiantoro, Biotechnology Laboratory, Faculty of Animal Science, Universitas Brawijaya Malang, Indonesia</li> <li>Prof Ani Nurgiartiningsih, Faculty of Animal Science, Universitas Brawijaya Malang, Indonesia</li> <li>Prof Gatot Ciptadi, Faculty of Animal Science, Universitas Brawijaya Malang, Indonesia</li> <li>Ms Desinta Wulandari, Faculty of Animal Science, Universitas Brawijaya Malang, Indonesia</li> <li>Prof Suyadi Suyadi, Faculty of Animal Science, Universitas Brawijaya Malang, Indonesia</li> </ul>					

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Abstract | Microsatellites are one option for characterizing cattle populations; that information can be used as a basis for the development and maintenance of local cattle. This study aims to evaluate genetic diversity and relationships using microsatellite markers between the populations of Gorontalo cattle in Gorontalo Province, Indonesia, using 3 microsatellite markers. A total of 126 samples of local bovine blood were collected, which consisted of North Gorontalo (n=28), Gorontalo (n=52), Bonebolango (n=37), PO (n=3), and Bali (n=6), were used in this study. The Genomic DNA Mini Kits were used for DNA extraction for an analysis fragment in the microsatellite DNA region using ILSTS017, HEL13, and BM1818 primers. A total of 74 alleles were identified across entire populations. The expected heterozygosity ranged from 0.407±0.216 (Bali) to 0.716±0.050 (Gorontalo), and the observed heterozygosity ranged from 0.471±0.084 (Bonebolango) to 0.778±0.222 (PO). F statistical analysis includes  $F_{IS}$  0.038,  $F_{IT}$  0.248, and  $F_{ST}$ 0.231. The three microsatellite markers were moderate (0.25-0.5) to highly informative (PIC>0.5). The research showed that Bali cattle were distinct from all other cattle populations, while Gorontalo-Bonebolango admixture and North Gorontalo cattle were mixed with the PO cattle population. In conclusion, markers used were highly informative and polymorphic in investigating genetic diversity in Bonebolango, Gorontalo, and North Gorontalo populations, while the genetic relationship among cattle populations was divided into two main clusters, i.e., the Bali and PO populations, which closely reflect the breeding process in the research area. This information will be useful for future development and maintenance of local cattle in Gorontalo Province, Indonesia.

Keywords | Microsatellite, Genetic diversity, Genetic relationships, Local cattle, Gorontalo

Received | December 07, 2022; Accepted | February 01, 2023; Published | ---- 00, 2023 \*Correspondence | Suyadi Suyadi, Faculty of Animal Science, Universitas Brawijaya Malang, East Java 65145, Indonesia; Email: suyadi@ub.ac.id Citation | Dako S, Laya NK, Gubali SI, Ardiantoro A, Nurgiartiningsih VMA, Ciptadi G, Wulandari D, Suyadi S (2023). Genetic diversity of Gorontalo local cattle based on microsatellite DNA. Adv. Anim. Vet. Sci. 7(s1):XXX-XXX. DOI | http://dx.doi.org/10.17582/journal.aavs/2023/....... ISSN (Online) | 2307-8316



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### **INTRODUCTION**

The increased demand for livestock products as human food illustrates an increase in human quality of life but challenges the genetic availability of livestock. The development of local livestock as a provider of genetic resources needs to be preserved because local livestock have the ability to adapt to their own environment. This genetic

ability is expected to produce more productive livestock (Gobel *et al.*, 2021; Pribadi *et al.*, 2015; Ilham *et al.*, 2016).

Gorontalo province is located in Indonesia, specifically between Central Sulawesi and North Sulawesi. The potential for cattle in Gorontalo Province is 192,229 heads (BPS, 2020). There are several types of cattle that are well developed in the Gorontalo region, such as Bali cattle, PO cattle,

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Table 1: Bo	vine microsatelli	te markers information used in study.					
Marker	Chromosome	Primer sequences (5'>3')	Primer attachment temperatures (°C)	Length of DNA base (Bp)	Genebank access		
ILSTS017				105-125			
HEL13 (D11S15)	11	F: TAAGGACTTGAGATAAGGAG R: CCATCTACCTCCATCTTAAC	52-57	178-200	X65207		
BM1818	23	F: AGCTGGGAATATAACCAAAGG	56-60	248-278	G18391		

R: AGTGCTTTCAAGGTCCATGC

Brahman cattle, and other local cattle that have not been identified. Local cattle that have not been identified are often called Diiti/ordinary/Local cattle. The existence of local cattle has a different phenotypic appearance compared to Bali and PO cattle. Naturally, livestock that adapts to the environment will bring up different phenotype variations in the population. Muladno (1994) state that genetic change can be used for the process of evolution of livestock populations/breeds by looking at the genetic distance of a population. Adaptability of local livestock describes a relative ability of an individual to survive and reproduce next generation to ensure continued survival of the population. Adaptations are mutations or genetic changes that help an organism or animal survive in its environment. Genetic adaptation is a biological characteristic with a heritable basis that improves reproduction and/or survival and results from evolution by natural selection. Increased animal performance based on genetic improvement results in more product produced per animal. Genetic improvements made in one generation are passed on to the next (Naskar et al., 2012; Mueller and Eenennaam, 2022). Efforts to maintain the genetic abilities that are formed in local cattle require adequate information, including qualitative characteristics, morphometrics, productivity, and genetics information, so that these local livestock can be developed in the future.

(D23S21)

Microsatellites are an option for characterizing cattle breeds or populations (Metta et al., 2004; Rincon et al., 2007); they are also used to answer questions related to genetic diversity and genetic relationships among cattle populations (Rincon et al., 2007; Sun et al., 2008; Chaudhari et al., 2009). Microsatellite loci are widely used because of their high polymorphism, co-dominance, and relative abundance in the genome (Rincon et al., 2007; Karthickeyan et al., 2009). The study of microsatellite DNA diversity in local Indonesian cattle has been reported previously (Satriani et al., 2002; Sarbaini, 2004; Winaya et al., 2007; Abdullah, 2008; Sutarno et al., 2015; Septian and Sumantri, 2015; Agung et al., 2015, 2019), but has never been reported in local cattle in Gorontalo. It is necessary to have an overview of the genetic diversity analysis of local Gorontalo cattle based on microsatellite DNA analysis. The importance of this information can be used as a basis for the development and maintenance of local cattle. This study aims to evaluate genetic diversity and relationships using microsatellite markers between the populations of Gorontalo cattle in Gorontalo Province, Indonesia.

### MATERIALS AND METHODS

### B LOOD samples and DNA isolation

All procedures related to animal use in this study were approved by the Animal Care and Use Committee of Brawijaya University under regulation number 145-KEP-UB-2022 (Ethical Clearance). A total of 126 samples of local bovine blood were collected from 3 different regions: Gorontalo (n = 52), Bonebolango (n = 37), and North Gorontalo (n = 28), plus the PO (n= 3) and Bali (n= 6) population samples in the Gorontalo province, were used in this study. Meanwhile, blood samples were analyzed at the Biotechnology Laboratory, Faculty of Animal Science, Universitas Brawijaya, Malang. Blood was collected from the vena coxygealis of the cattle. The DNA was isolated using Genomic DNA Mini Kits. The DNA isolation procedures followed the protocol instructions.

### PRIMER AND DNA AMPLIFICATION

A total of 3 bovine microsatellite markers were used in the PCR process. The markers consisted of ILSTS017, HEL13, and BM1818 (Table 1). The PCR reaction contained a mixture of 1  $\mu$ L DNA template (50–100 ng/ $\mu$ L) and 30  $\mu$ L PCR premix. The PCR premix consisted of 0.4  $\mu$ L primer (10 pmol/ $\mu$ L), 15  $\mu$ L Go Taq Green Master Mix (Promega, USA), and 14.2  $\mu$ L Nucleus Free Water (NFW). The PCR thermal cycler conditions are shown in Table 2. The PCR product visualization methods were based on Susilorini *et al.* (2022). Microsatellite fragment analysis was conducted at First Base Laboratory, Selangor, Malaysia.

### **Table 2:** The PCR thermal cycler condition.

PCR Step	Temperature (°C)	Time	Number of cycle
Pre-denaturation	95	5 minutes	Once
Denaturation	95	30 s	35 times
Annealing	58	45 s	
Extension	72	1 minute	
Final extension	72	5 minutes	Once

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### MICROSATELLITE DATA ANALYSIS

The fragment analysis result was processed to get allele size data. Allele data was then converted using Convert version 1.31 for further analysis. The converted data were analyzed using Cervus version 3.0.7 to generate observed heterozygosity (HO), expected heterozygosity (HE), frequency/number of alleles, polymorphism information content (PIC) values, and Hardy-Weinberg (HW) equilibrium. GenAlEx 6.51 b2 was used to generate genetic differentiation (FST), the rate of inbreeding between populations (FIS), the rate of inbreeding in populations (FIT), and principal coordinate analysis (PCoA) to determine the genetic relationship between livestock breeds. Genetic structures were analyzed using POPTREEW (POPTREEW website version) (Takezaki et al., 2014) to generate the reconstruction of phylogeny trees between populations and genetic distance.

# MICROSATELLITE POLYMORPHISMS AND GENETIC DIVERSITY

**RESULT AND DISCUSSION** 

Studies on the determination of genetic diversity in Gorontalo cattle based on microsatellite markers aren't available, so this study was conducted to find information about its diversity. The polymorphism of molecular genetics and the F-statistic belonging to each locus are summarized in Tables 3 and 4. A total of 74 alleles were detected through the 3 markers analyzed in entire populations: 8 alleles were detected in Bali cattle, 9 alleles in PO cattle, 20 alleles in Bonebolango cattle, 21 alleles in Gorontalo cattle, and 16 alleles in North Gorontalo cattle. The observed heterozygosity (OH) was 0.474, 0.407, 0.737, and the expected heterozygosity (EH) was 0.712, 0.658, and 0.829 for ILSTS017, HEL013, and BM1818, respectively.

Table 3: F Statistics	s (F <sub>15</sub>	$, F_{\rm IT}, F_{\rm ST})$	between fi	ive cattle	population.
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Lokus	Ν	Ho	He	PIC	HW	F <sub>IS</sub>	F <sub>IT</sub>	F <sub>st</sub>	Nm
ILSTS017	95	0.474	0.712	0.699	***	0.412	0.530	0.200	0.999
HEL013	86	0.407	0.658	0.601	**	0.067	0.214	0.157	1.342
BM1818	95	0.737	0.803	0.803	***	-0.017	0.157	0.068	3.442
Mean						0.154	0.265	0.142	1.928
SD						0.131	0.140	0.039	0.764

Note: N=number of sample; Ho=observed heterozygosity; He=expected heterozygosity; PIC=polymorphism information content; HW= Hardy-Weinberg Equilibrium; FIS= inter-population inbreeding rate; FIT= inbreeding rate in population; FST= genetic differentiation; Nm= gene flow.

Pop	Lokus	Ν	Na	Ne	PIC	Ho	He	HW
Bali	ILSTS017	6	3	1.946	0.424	0.167	0.530	ND
	HEL013	6	5	3.600	0.680	0.833	0.788	ND
	BM1818	6	6	5.143	0.777	0.833	0.879	ND
РО	ILSTS017	3	2	1.800	0.346	0.000	0.533	ND
	HEL013	3	3	2.571	0.535	1.000	0.733	ND
	BM1818	3	4	3.600	0.671	1.000	0.867	ND
Bolongo	ILSTS017	22	8	2.907	0.627	0.364	0.671	ND
	HEL013	17	4	2.429	0.513	0.412	0.606	ND
	BM1818	22	8	4.523	0.754	0.636	0.797	ND
Gorontalo	ILSTS017	41	9	3.543	0.682	0.488	0.727	*
	HEL013	37	5	2.700	0.564	0.351	0.638	*
	BM1818	41	7	5.033	0.777	0.732	0.811	ND
Gorontalo Utara	ILSTS017	23	4	2.574	0.553	0.696	0.625	ND
	HEL013	23	3	2.266	0.489	0.304	0.571	ND
	BM1818	23	9	5.290	0.785	0.783	0.829	ND

**Table 4:** Number of alleles (Na), observed heterozygosity (Ho), and expected (He), and Polymorphism Information Content in five cattle population in each loci.

Note: N=number of sample; Na= Number of alleles; Ne= Mean number of effective alleles; Ho= observed heterozygosity; He= expected heterozygosity; PIC= polymorphism information content; HW= Hardy-Weinberg Equilibrium.

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**Table 5:** Number of alleles (Na), observed heterozygosity (Ho), and expected (He), and polymorphism information content in five cattle breeds observed.

Breed	Ν	Na±SD	Ne±SD	Ho±SD	He±SD	PIC
А	6	4.67±0.88	3.563±0.923	0.611±0.222	0.671±0.096	0.627
В	3	3.00±0.58	2.657±0.521	0.667±0.333	0.593±0.081	0.518
С	22	6.68±1.33	3.286±0.634	0.471±0.084	0.674±0.056	0.631
D	41	7.00±1.16	3.759±0.682	0.524±0.111	0.716±0.050	0.674
Е	23	5.33±1.86	3.377±0.961	0.594±0.147	0.660±0.077	0.609

Note: A= Bali cattle; B= PO cattle; C= Bolongo; D= Gorontalo; E= North Gorontalo; N= number of sample; Na= Number of alleles; Ne= Mean number of effective alleles; Ho= observed heterozygosity; He= expected heterozygosity; PIC= polymorphism information content; SD= standart deviation.

The genetic diversity belonging to each population was shown in Table 5. EH ranged from  $0.407\pm0.216$  (Bali) to  $0.716\pm0.050$  (Gorontalo), whereas OH ranged from  $0.471\pm0.084$  (Bonebolango) to  $0.778\pm0.222$  (PO). The EH value was higher than OH in all populations except in the PO and Bali populations.

Expected heterozygosity in all loci showed a higher value than observed heterozygosity, indicating that all loci deviated from HWE. The deviation of loci from HWE can be caused by several factors, such as selection, genetic drift, inbreeding, the presence of null alleles, selection in favor of homozygotes, and the Wahlund effect (Rahal et al., 2021). The EH value was lower compared to Algerian cattle (Rahal et al., 2021), but higher than the previous study in Indonesian cattle (Sutarno et al., 2015). The highest level of heterozygosity was in locus BM1818, which was higher than previous research in Red Steppe cattle using BM1818 loci, which obtained heterozygosity values of 0.692 (OH) and 0.701 (EH) (Kramarenko et al., 2018). The highest heterozygosity in the Gorontalo population (0.716±0.050) could be explained by the lack of a breeding program. The EH > OH value indicates low heterozygosity. Kriswiyanti (2014) states that heterozygosity influenced by the variety of alleles and the frequency of each allele at each locus. Heterozygosity values range from 0 to 1. If heterozygosity equals 0 (zero), then the population being measured has a very close genetic relationship, and if the value is equal to 1 (one), then the population being measured has no genetic relationship.

The F-statistics consisting of  $F_{IS}$ ,  $F_{TT}$ , and  $F_{ST}$  were 0,038, 0248, and 0.231, respectively. All microsatellite loci were deviated from the Hardy-Weinberg Equilibrium (HWE), meanwhile for all loci in each population, only locus ILSTS and HEL013 in Gorontalo cattle showed the deviation from HWE. The presence of null alleles in the study was between 6.15% (BM1818) and 24.01% (HEL013). The  $F_{ST}$  value of 23.1% indicated that 23.1% of the total genetic variation was due to allele differentiation between breeds, while the remaining 76.9% was due to the difference among individuals within the breed across the 3 markers. The average  $F_{ST}$  value of 0.231 indicated that the genetic

difference in the population was high (0.15-0.25). Hartl and Clark (2007) state that the  $F_{ST}$  value of 0-0.05 (small), 0.05-0.15 (medium), 0.15-0.25 (high), and > 0.25 (very high). Obtained F<sub>ST</sub> values were higher than Egyptian cattle (El-Sayeed et al., 2016), Algerian cattle (Rahal et al., 2021), South African Nguni cattle (Sanarana et al., 2016), and cattle raised in Turkey (Demis and Balciogly, 2019), but these values were lower compared to previous research in Indonesian cattle (Agung *et al.*, 2019). The  $F_{IS}$  values were 0.412, 0.067, and -0.017 for ILSTS017, HEL013, and BM1818, respectively. The obtained  $F_{15}$  value was used to gain a better understanding of the degree of inbreeding and endangered potentiality. A  $\mathrm{F}_{\mathrm{IS}}$  value less than 0.05 means the breeds are not in danger, 0.05-0.15 means the breeds are potentially endangered, 0.15-0.25 means the breeds are minimally endangered, 0.25-0.40 means the breeds are endangered, and >0.40 means the breeds are critically endangered (Simon and Bchenauer, 1993; El-Sayeed et al., 2016).

PIC values were found to be polymorphic between 0.601 (HEL013) and 0.803 (BM1818). The PIC value based on each population ranged from 0.353 to 0.674, which indicated a moderately to highly informative value. In the present study, the PIC values of three markers were highly informative (> 0.5), whereas the PIC values for ILSTS017 loci in the Bali and PO populations had low informative values (0.25-0.5). The PIC values in previous research on Egyptian cattle were 0.45 and 0.64 (El-Sayeed et al., 2016); those of Taro White cattle were 0.536 (Hervani et al., 2019); those of Zimbabwean Sanga cattle were 0.666-0.682 (Gororo et al., 2018); and those of Eastern European cattle were 0.752 (Illie et al., 2015). The PIC result showed that the markers are highly informative for the characterization of the three populations used in the study, i.e., the Bonebolango, Gorontalo, and North Gorontalo populations. Botstein et al. (1980) state that the PIC value for identifying genetic diversity should be greater than 0.5.

#### **GENETIC STRUCTURE AND RELATIONSHIPS**

The result of genetic distance was used as basic data to create the visualization of a phylogeny tree (Table 6),

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which revealed the closest genetic distance between Gorontalo-Bonebolango (0.037), followed by Gorontalo – North Gorontalo (0.043) and Bonebolango-North Gorontalo (0.078), respectively, while the farthest genetic distance was between Bali – PO (2.109). Estimation of the pair-wise  $F_{ST}$  value revealed the highest genetic differentiation between PO and Bali (0.206) and the lowest genetic differentiation between Bonebolango-North Gorontalo (0.011), Gorontalo-North Gorontalo (0.012), Bonebolango-Gorontalo (0.013), and PO-North Gorontalo (0.029). Low genetic differentiation indicated high genetic similarity between populations.

**Table 6:** Population pair-wise Fst (bottom diagonal) and Nei's standard genetic distance (top diagonal).

Populasi	Bali	РО	Bolon- go	Goronta- lo	Gorontalo Utara
Bali	***	0.053	0.959	0.714	0.807
PO	0.015	***	1.233	0.931	1.030
Bolongo	0.134	0.174	***	0.037	0.078
Gorontalo	0.104	0.142	0.008	***	0.043
Gorontalo Utara	0.128	0.166	0.017	0.010	***



**Figure 1:** The reconstruction of the UPGMA phylogeny tree of Gorontalo cattle populations with Nei genetic distance.



**Figure 2:** Principal coordinate analysis (PCoA) based on 3 microsatellite loci from 95 individual cattle in Indonesia. Bali= ◆; PO= ■; Bolongo= ▲; Gorontalo = ×; Gorontalo Utara= \*.

Genetic distance analysis shows that the genetic distance between populations ranged from 0.037 to 2.109, while the pair-wise  $F_{\rm ST}$  value ranged from 0.011 to 0.206. The reconstruction of the phylogeny tree based on Nei's genetic distance is shown in Figure 1. The figure revealed

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two main clusters of cattle populations. The first cluster consisted of Bali cattle, and the second cluster consisted of PO cattle. Clusters of PO population were divided into two subpopulations, i.e., the North Gorontalo population and the Bonebolango–Gorontalo population. The principal coordinate analysis (PCoA) result supported the visualization of the phylogeny tree. It is shown that Bali cattle clearly separate from other cattle, while the population of PO cattle forms a cluster that contains local cattle in three different populations as subpopulations.

The reconstruction of a phylogeny tree using five cattle populations revealed a high gene share between Gorontalo and Bonebolango. The PCoA analysis confirmed that Bali populations are distinct from other populations studied, whereas Gorontalo-Bonebolango admixture and North Gorontalo were mixed with the PO population. This is due to the distribution pattern of the arrival of cattle to Gorontalo province, which starts from North Gorontalo, and then the incoming cattle will be selected. After that, the cattle will be distributed to other areas, such as Gorontalo and Bonebolango. Cattle that entered North Sulawesi province (before Gorontalo province separated from North Sulawesi province) were dominated by PO cattle. Then, in 2000 (UU No. 38 of 2000), when Gorontalo officially became a separate province from North Sulawesi, the province of Gorontalo was dominated by Bali cattle. This distribution pattern allows for differences between the cattle populations in North Gorontalo and other areas. The similarity in cattle between the populations of Gorontalo and Bonebolango can be due to the fact that the two regions are close together, which allows for inbreeding between livestock and causes the appearance of the same genetics.



**Figure 3:** Principal coordinate analysis (PCoA) based on each population.

The populations of Bali and PO cattle form a different cluster. This is because Bali and PO cattle are different species; Bali cattle are included in *Bos sondaicus*, while PO cattle are in *Bos indicus*. The existence of PO cattle for more than two decades can cause a high share of genetic pool between PO cattle and local cattle in Gorontalo province. Samples of cattle in the populations of North Gorontalo,

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Gorontalo, and Bonebolango are local cattle in each of these regions. The phenotypic appearance of the three cattle in the area has a dominant mixture of PO phenotype, a little Bali phenotype, and both cattle's phenotypes (Suyadi *et al.*, 2014; Laya *et al.*, 2020; Domili *et al.*, 2021; Gobel *et al.*, 2021). However, they are not said to be PO and Bali cattle (Figures 4, 5, and 6), but local cattle for each region. The study could serve as guidelines for future genetic studies on the development and maintenance of local cattle in Gorontalo Province, Indonesia.



Figure 4: Local cattle population of Gorontalo Regency.



Figure 5: Local cattle population of North Gorontalo.



Figure 6: Local cattle of the Bonebolango population.

### CONCLUSIONS AND RECOMMENDATIONS

Three markers used were highly informative and polymorphic in investigating genetic diversity in three cattle populations in Gorontalo Province, i.e., Bonebolango, Gorontalo, and North Gorontalo. There are two main clusters: (1) Bali cattle, which are distinct from all other cattle populations; and (2) the PO population, which includes the North Gorontalo, Bonebolango, and Gorontalo subpopulations and closely reflects the breeding process state in the research area. This information will be useful for the future development and maintenance of local cattle in Gorontalo Province, Indonesia, and it is recommended to standardize local Gorontalo cattle as typical local Gorontalo cattle.

### ACKNOWLEDGEMENTS

The authors thank the team of Department of Animal Husbandry, Faculty of Agriculture, Gorontalo University for supporting the data and the members of the Genomic and Proteomic Research Group, Faculty of Animal Science, Universitas Brawijaya for data analysis.

### **NOVELTY STATEMENT**

### **AUTHOR'S CONTRIBUTION**

SD, SS, VMAN and GC idea and design. SD, NKL, SIG, AA and DW material sample collection and lab analysis. DW, SD and NKL write the manuscript. SD and SS: Revision.

### **CONFLICT OF INTEREST**

The authors have declared no conflict of interest.

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### Genetic Diversity of Gorontalo Local Cattle Based on Microsatellite DNA

Genetic Diversity of Gorontalo Local Cattle Based on Microsatellite DNA

Journal	Advances in Animal and Veterinary Sciences				
Manuscript ID	MH20221207091228-R2				
Manuscript Type	Research Article				
Area of Interest	Husbandry				
Date Submitted by the Author	Thu, 26 Jan 2023, 09:34 AM				
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Microsatellites are one option for characterizing cattle populations; that information can be used as a basis for the development and maintenance of local cattle. This study aims to evaluate genetic diversity and relationships using microsatellite markers between the populations of Gorontalo cattle in Gorontalo Province, Indonesia, using 3 microsatellite markers. A total of 126 samples of local bovine blood were collected, which consisted of North Gorontalo (n=28), Gorontalo (n=52), Bonebolango (n=37), PO (n=3), and Bali (n=6), were used in this study. The Genomic DNA Mini Kits were used for DNA extraction for an analysis fragment in the microsatellite DNA region using ILSTS017, HEL13, and BM1818 primers. A total of 74 alleles were identified across entire populations. The expected heterozygosity ranged from 0.407±0.216 (Bali) to 0.716±0.050 (Gorontalo), and the observed heterozygosity ranged from 0.471±0.084 (Bonebolango) to  $0.778\pm0.222$  (PO). F statistical analysis includes F<sub>IS</sub> 0.038, F<sub>IT</sub> 0.248, and F<sub>ST</sub> 0.231. The three microsatellite markers were moderate (0.25-0.5) to highly informative (PIC>0.5). The research showed that Bali cattle were distinct from all other cattle populations, while Gorontalo-Bonebolango admixture and North Gorontalo cattle were mixed with the PO cattle population. In conclusion, markers used were highly informative and polymorphic in investigating genetic diversity in Bonebolango, Gorontalo, and North Gorontalo populations, while the genetic relationship among cattle populations was divided into two main clusters, i.e., the Bali and PO populations, which closely reflect the breeding process in the research area. These information will be useful for future development and maintenance of local cattle in Gorontalo Province, Indonesia.

The increased demand for livestock products as human food illustrates an increase in human quality of life but challenges the genetic availability of livestock. The development of local livestock as a provider of genetic resources needs to be preserved because local livestock have the ability to adapt to their own environment. This genetic ability is expected to produce more productive livestock (Gobel et al., 2021; Pribadi et al., 2015; Ilham et al., 2016).

Gorontalo Province is located in Indonesia, specifically between Central Sulawesi and North Sulawesi. The potential for cattle in Gorontalo Province is 192,229 heads (BPS, 2020). There are several types of cattle that are well developed in the Gorontalo region, such as Bali cattle, PO cattle, Brahman cattle, and other local cattle that have not been identified. Local cattle that have not been identified are often called Diiti/ordinary/Local cattle. The existence of local cattle has a different phenotypic appearance compared to Bali and PO cattle. Naturally, livestock that adapts to the environment will bring up different phenotype variations in the population. Muladno (1994) state that genetic change can be used for the process of evolution of livestock populations/breeds by looking at the genetic distance of a population. Adaptability of local livestock describes a relative ability of an individual to survive and reproduce next generation to ensure continued survival of the population. Adaptations are mutations or genetic changes that help an organism or animal survive in its environment. Genetic adaptation is a biological characteristic with a heritable basis that improves reproduction and/or survival and results from evolution by natural selection. Increased animal performance based on genetic improvement results in more product produced per animal. Genetic improvements made in one generation are passed on to the next (Naskar et al., 2012; Mueller and Eenennaam, 2022). Efforts to maintain the genetic abilities that are formed in local cattle require adequate information, including qualitative characteristics, morphometrics, productivity, and genetics information, so that these local livestock can be developed in the future.

Microsatellites are an option for characterizing cattle breeds or populations (Metta et al., 2004; Rincon et al., 2007); they are also used to answer questions related to genetic diversity and genetic relationships among cattle populations (Rincon et al., 2007; Sun et al., 2008; Chaudhari et al., 2009). Microsatellite loci are widely used because of their high polymorphism, co-dominance, and relative abundance in the genome (Rincon et al., 2007; Karthickeyan et al., 2009). The study of microsatellite DNA diversity in local Indonesian cattle has been reported previously (Satriani et al., 2002; Sarbaini, 2004; Winaya et al., 2007; Abdullah, 2008; Sutarno et al., 2015; Septian and Sumantri, 2015; Agung et al., 2016; Agung et al., 2019), but has never been reported in local cattle in Gorontalo. It is necessary to have an overview of the genetic diversity analysis of local Gorontalo cattle based on microsatellite DNA analysis. The importance of this information can be used as a basis for the development and maintenance of local cattle. This study aims to evaluate genetic diversity and relationships using microsatellite markers between the populations of Gorontalo cattle in Gorontalo Province, Indonesia.

All procedures related to animal use in this study were approved by the Animal Care and Use Committee of Brawijaya University under regulation number 145-KEP-UB-2022 (Ethical Clearance). A total of 126 samples of local bovine blood were collected from 3 different regions: Gorontalo (n = 52), Bonebolango (n = 37), and North Gorontalo (n = 28), plus the PO (n = 3) and Bali (n = 6) population samples in the Gorontalo province, were used in this study. Meanwhile, blood samples were analyzed at the Biotechnology Laboratory, Faculty of Animal Science, Universitas Brawijaya, Malang. Blood was collected from the vena coxygealis of the cattle. The DNA was isolated using Genomic DNA Mini Kits. The DNA isolation procedures followed the protocol instructions.

A total of 3 bovine microsatellite markers were used in the PCR process. The markers consisted of ILSTS017, HEL13, and BM1818 (Table 1). The PCR reaction contained a mixture of 1  $\mu$ L DNA template (50–100 ng/ $\mu$ L) and 30  $\mu$ L PCR premix. The PCR premix consisted of 0.4  $\mu$ L primer (10 pmol/ $\mu$ L), 15  $\mu$ L Go Taq Green Master Mix (Promega, USA), and 14.2  $\mu$ L Nucleus Free Water (NFW). The PCR thermal cycler conditions are shown in Table 2. The PCR product visualization methods were based on Susilorini et al., (2022). Microsatellite fragment analysis was conducted at First Base Laboratory, Selangor, Malaysia.

The fragment analysis result was processed to get allele size data. Allele data was then converted using Convert version 1.31 for further analysis. The converted data were analyzed using Cervus version 3.0.7 to generate observed heterozygosity (HO), expected heterozygosity (HE), frequency/number of alleles, polymorphism information content (PIC) values, and Hardy–Weinberg (HW) equilibrium. GenAlEx 6.51 b2 was used to generate genetic differentiation (FST), the rate of inbreeding between populations (FIS), the rate of inbreeding in populations

(FIT), and principal coordinate analysis (PCoA) to determine the genetic relationship between livestock breeds. Genetic structures were analyzed using POPTREEW (POPTREEW website version) (Takezaki et al., 2014) to generate the reconstruction of phylogeny trees between populations and genetic distance.

#### **RESULT AND DISCUSSION**

#### **Microsatellite Polymorphisms and Genetic Diversity**

Studies on the determination of genetic diversity in Gorontalo cattle based on microsatellite markers aren't available, so this study was conducted to find information about its diversity. The polymorphism of molecular genetics and the F-statistic belonging to each locus are summarized in Table 3 and Table 4. A total of 74 alleles were detected through the 3 markers analyzed in entire populations: 8 alleles were detected in Bali cattle, 9 alleles in PO cattle, 20 alleles in Bonebolango cattle, 21 alleles in Gorontalo cattle, and 16 alleles in North Gorontalo cattle. The observed heterozygosity (OH) was 0.474, 0.407, 0.737, and the expected heterozygosity (EH) was 0.712, 0.658, and 0.829 for ILSTS017, HEL013, and BM1818, respectively. The genetic diversity belonging to each population was shown in Table 5. EH ranged from  $0.407\pm0.216$  (Bali) to  $0.716\pm0.050$  (Gorontalo), whereas OH ranged from  $0.471\pm0.084$  (Bonebolango) to  $0.778\pm0.222$  (PO). The EH value was higher than OH in all populations except in the PO and Bali populations.

Expected heterozygosity in all loci showed a higher value than observed heterozygosity, indicating that all loci deviated from HWE. The deviation of loci from HWE can be caused by several factors, such as selection, genetic drift, inbreeding, the presence of null alleles, selection

in favor of homozygotes, and the Wahlund effect (Rahal et al., 2020). The EH value was lower compared to Algerian cattle (Rahal et al., 2020), but higher than the previous study in Indonesian cattle (Sutarno et al., 2015). The highest level of heterozygosity was in locus BM1818, which was higher than previous research in Red Steppe cattle using BM1818 loci, which obtained heterozygosity values of 0.692 (OH) and 0.701 (EH) (Kramarenko et al., 2018). The highest heterozygosity in the Gorontalo population ( $0.716\pm0.050$ ) could be explained by the lack of a breeding program. The EH > OH value indicates low heterozygosity. Kriswiyanti (2014) states that heterozygosity values range from 0 to 1. If heterozygosity equals 0 (zero), then the population being measured has a very close genetic relationship, and if the value is equal to 1 (one), then the population being measured has no genetic relationship.

The F-statistics consisting of  $F_{IS}$ ,  $F_{IT}$ , and  $F_{ST}$  were 0,038, 0248, and 0.231, respectively. All microsatellite loci were deviated from the Hardy-Weinberg Equilibrium (HWE), meanwhile for all loci in each population, only locus ILSTS and HEL013 in Gorontalo cattle showed the deviation from HWE. The presence of null alleles in the study was between 6.15% (BM1818) and 24.01% (HEL013). The  $F_{ST}$  value of 23.1% indicated that 23.1% of the total genetic variation was due to allele differentiation between breeds, while the remaining 76.9% was due to the difference among individuals within the breed across the 3 markers. The average  $F_{ST}$  value of 0.231 indicated that the genetic difference in the population was high (0.15-0.25). Hartl and Clark (2007) state that the  $F_{ST}$  value of 0-0.05 (small), 0.05-0.15 (medium), 0.15-0.25 (high), and > 0.25 (very high). Obtained  $F_{ST}$  values were higher than Egyptian cattle (El Sayeed et al., 2016), Algerian cattle (Rahal et al., 2020), South African Nguni cattle (Sanarana et al., 2016), and cattle raised in Turkey (Demis and Balciogly, 2019), but these values were lower compared to previous research in Indonesian cattle (Agung et al., 2019). The  $F_{IS}$  values were 0.412, 0.067, and -0.017 for ILSTS017, HEL013, and BM1818, respectively. The obtained  $F_{IS}$  value was used to gain a better understanding of the degree of inbreeding and endangered potentiality. A  $F_{IS}$  value less than 0.05 means the breeds are not in danger, 0.05-0.15 means the breeds are potentially endangered, 0.15-0.25 means the breeds are minimally endangered, 0.25-0.40 means the breeds are endangered, and >0.40 means the breeds are critically endangered (Simon and Bchenauer, 1993; El Sayeed et al., 2016).

PIC values were found to be polymorphic between 0.601 (HEL013) and 0.803 (BM1818). The PIC value based on each population ranged from 0.353 to 0.674, which indicated a moderately to highly informative value. In the present study, the PIC values of three markers were highly informative (> 0.5), whereas the PIC values for ILSTS017 loci in the Bali and PO populations had low informative values (0.25–0.5). The PIC values in previous research on Egyptian cattle were 0.45 and 0.64 (El Sayeed et al., 2016); those of Taro White cattle were 0.536 (Heryani et al., 2019); those of Zimbabwean Sanga cattle were 0.666–0.682 (Gororo et al., 2018); and those of Eastern European cattle were 0.752 (Illie et al., 2015). The PIC result showed that the markers are highly informative for the characterization of the three populations used in the study, i.e., the Bonebolango, Gorontalo, and North Gorontalo populations. Botstein et al., (1980) state that the PIC value for identifying genetic diversity should be greater than 0.5.

#### **Genetic Structure and Relationships**

The result of genetic distance was used as basic data to create the visualization of a phylogeny tree (Table 6), which revealed the closest genetic distance between Gorontalo – Bonebolango (0.037), followed by Gorontalo – North Gorontalo (0.043) and Bonebolango –

North Gorontalo (0.078), respectively, while the farthest genetic distance was between Bali – PO (2.109). Estimation of the pair-wise  $F_{ST}$  value revealed the highest genetic differentiation between PO and Bali (0.206) and the lowest genetic differentiation between Bonebolango – North Gorontalo (0.011), Gorontalo – North Gorontalo (0.012), Bonebolango – Gorontalo (0.013), and PO – North Gorontalo (0.029). Low genetic differentiation indicated high genetic similarity between populations.

Genetic distance analysis shows that the genetic distance between populations ranged from 0.037 to 2.109, while the pair-wise  $F_{ST}$  value ranged from 0.011 to 0.206. The reconstruction of the phylogeny tree based on Nei's genetic distance is shown in Figure 1. The figure revealed two main clusters of cattle populations. The first cluster consisted of Bali cattle, and the second cluster consisted of PO cattle. Clusters of PO population were divided into two subpopulations, i.e., the North Gorontalo population and the Bonebolango–Gorontalo population. The principal coordinate analysis (PCoA) result supported the visualization of the phylogeny tree. It is shown that Bali cattle clearly separate from other cattle, while the population of PO cattle forms a cluster that contains local cattle in three different populations as subpopulations.

The reconstruction of a phylogeny tree using five cattle populations revealed a high gene share between Gorontalo and Bonebolango. The PCoA analysis confirmed that Bali populations are distinct from other populations studied, whereas Gorontalo-Bonebolango admixture and North Gorontalo were mixed with the PO population. This is due to the distribution pattern of the arrival of cattle to Gorontalo province, which starts from North Gorontalo, and then the incoming cattle will be selected. After that, the cattle will be distributed to other areas, such as Gorontalo and Bonebolango. Cattle that entered North Sulawesi province (before Gorontalo province separated from North Sulawesi province) were dominated by PO cattle. Then, in 2000 (UU No. 38 of 2000), when Gorontalo officially became a separate province from North Sulawesi, the province of Gorontalo was dominated by Bali cattle. This distribution pattern allows for differences between the cattle populations in North Gorontalo and other areas. The similarity in cattle between the populations of Gorontalo and Bonebolango can be due to the fact that the two regions are close together, which allows for interbreeding between livestock and causes the appearance of the same genetics.

The populations of Bali and PO cattle form a different cluster. This is because Bali and PO cattle are different species; Bali cattle are included in *Bos sondaicus*, while PO cattle are in *Bos indicus*. The existence of PO cattle for more than two decades can cause a high share of genetic pool between PO cattle and local cattle in Gorontalo province. Samples of cattle in the populations of North Gorontalo, Gorontalo, and Bonebolango are local cattle in each of these regions. The phenotypic appearance of the three cattle in the area has a dominant mixture of PO phenotype, a little Bali phenotype, and both cattle's phenotypes (Suyadi et al., 2014; Laya et al., 2020; Domili et al., 2021; Gobel et al., 2021). However, they are not said to be PO and Bali cattle (Figures 4, 5, and 6), but local cattle for each region. The study could serve as guidelines for future genetic studies on the development and maintenance of local cattle in Gorontalo Province, Indonesia.

#### CONCLUSION

Three markers used were highly informative and polymorphic in investigating genetic diversity in three cattle populations in Gorontalo Province, i.e., Bonebolango, Gorontalo, and North Gorontalo. There are two main clusters: (1) Bali cattle, which are distinct from all other

cattle populations; and (2) the PO population, which includes the North Gorontalo, Bonebolango, and Gorontalo subpopulations and closely reflects the breeding process state in the research area. These information will be useful for the future development and maintenance of local cattle in Gorontalo Province, Indonesia, and it is recommended to standardize local Gorontalo cattle as typical local Gorontalo cattle.

The authors thank the team of Department of Animal Husbandry, Faculty of Agriculture, Gorontalo University for supporting the data and the members of the Genomic and Proteomic Research Group, Faculty of Animal Science, Universitas Brawijaya for data analysis.