South African developed meat type goats: A forgotten animal genetic resource?

A. Pieters¹, E. van Marle-Köster¹, C. Visser¹, & A. Kotze²

¹Department of Animal & Wildlife Sciences, University of Pretoria, Pretoria 0002, South Africa ²National Zoological Gardens, Pretoria 0002, South Africa

Summary

There are mainly three locally developed meat type goat breeds in South Africa namely the South African Boer goat, the Savanna and the Kalahari Red. In order to maintain the characteristics of these breeds and to ensure that their unique traits are not lost through continuous selection and cross-breeding, it has become important to revisit current breed standards and introduce genetic characterization to obtain genetic diversity parameters. Both the Boer goat and Savanna breeds have established breeders associations while a breeders club was founded for the Kalahari Red goats. These associations have set breed standards and the breeds' morphometric traits have recently been described. The differences in morphometric traits among the breeds are fairly insignificant. Preliminary data on genetic diversity report heterozygosity values above 50% for each of the three breeds and they can be distinguished as separate breeds based on genotyping results with eighteen ISAG recommended microsatellite markers. There is however a need for a genetic characterization of both meat type and indigenous goats to ensure sufficient diversity within the breeds for long term conservation of the unique genetic resource.

Resumé

Il y a en gros trois races de chèvres à viande développées en Afrique du Sud, à savoir la chèvre Boer sud-africaine, la Savanna et la Kalahari Rouge. Afin de conserver les caractéristiques de ces races et de faire en sorte que leurs traits uniques ne se perdent pas au cours des sélections et croisements continus, il est devenu nécessaire de revoir les normes des races actuelles et d'introduire des caractéristiques génétiques pour obtenir des paramètres de diversité génétique. La race de chèvre Boer et la Savanna ont des associations d'éleveurs établies, tandis qu'un club d'éleveurs a été créé pour les Kalahari Rouge. Ces associations ont des normes raciales bien définies et les traits morpho-métriques des races ont été décrits récemment. Les différences dans les traits morpho-métriques des races sont négligeables. Les données préliminaires sur la diversité génétique révèlent des valeurs hétérozygotes de plus de 50% pour chacune des trois races et elles peuvent être distinguées comme des races séparées en se basant sur les résultats génotypiques avec dix-huit marqueurs microsatellites recommandés par l'ISAG. Cependant, il faut faire une caractérisation génétique des deux types de viande et des chèvres indigènes pour garantir une diversité suffisante dans les races afin de conserver à long terme cette unique ressource génétique.

Resumen

Existen mayormente tres razas de cabras de carne en Sud Africa, la cabra Boer sudafricana, la Savanna y la Kalahari Roja. Con el fin de conservar las características de estas razas y conseguir que sus rasgos únicos no se pierdan a lo largo de selecciones y cruces continuos, se ha hecho necesario revisar las normas de las razas actuales e introducir características genéticas para obtener parámetros de diversidad genética. Las razas de cabra Boer y Savanna poseen asociaciones de ganaderos, mientras que un club de ganaderos ha sido creado para la Kalahari Roja. Estas asociaciones tienen normas bien definidas para las razas y los rasgos morfométricos han sido descritos recientemente. Las diferencias en los rasgos morfométricos entre las razas no son importantes. Los datos preliminares sobre la diversidad genética revelan valores de heterocigosis de más del 50% para cada una de las tres razas y pueden diferenciarse como razas separadas en base a los resultados genotípicos con dieciocho marcadores microsatelitares recomendados por ISAG. Sin

embargo, hay que hacer una caracterización genética de dos tipos de carne y de cabras indígenas para garantizar una diversidad suficiente entre las razas con el fin de conservar a largo plazo este recurso genético único.

Key words: Meat type, Phenotypic, Genetic characteristics, Utilization, Conservation.

Introduction

South Africa has a goat population of approximately 6.6 million of which 64% are kept and/or commercially farmed within the rural areas (Coetzee, 1998). The primary goat breeds consist of Angora for mohair production, three meat type goat breeds namely the South African Boer goat, the Savanna and the Kalahari Red and a number of indigenous types that are primarily classified according to the geographical area in which they reside. Apart from the Angora goats, these breeds are all believed to originate from indigenous types, which had migrated to Southern Africa by 2500 BC, due to increasing desertification of the southern regions of the Sahara (Maree and Plug, 1993). The local Khoisan shepherds travelled southwards from Northern Botswana down to the Orange River and later followed two additional routes to reach the Southern and Western Cape. One of the earliest references to the presence of goats in the Northern Cape was made by the missionary, J. Burrow, who described the goats kept by the local people as "handsome goats, speckled like the leopard" (Campbell, 2003). These goats, also referred to as 'skilder' goats or speckled goats, were most likely the resource, which led to the development of the current meat type goats.

These locally developed breeds have been subjected to artificial selection for improved production and growth. Although this has led to improved performance in the breeds, the question arises if specific genetic characteristics could be compromised by continuous selection for improvement of growth and meat characteristics. Often these breeds and our indigenous types are marketed as having special adaptive characteristics; local types often survive tick borne diseases better than commercial types (Malan, 2000; Erasmus, 2000).

According to some authors, true indigenous goat breeds in South Africa have been virtually cross-bred to extinction due to the development of the meat type goats such as the Boer goat (Campbell, 1995). Some researchers and farmers believe that the pure 'unimproved' indigenous goats possess important economic traits (including viability, good mothering ability, disease resistance and resistance against ticks), which should not be disregarded (Campbell, 1995; Van der Walt, 200, personal communication).

Recently a strategy for the conservation and utilization of South African genetic animal resources, including goat genetic resources, has been formulated. This policy aims at facilitating poverty alleviation through the strategic management of genetic resources, thereby improving the livelihoods of farmers and rural communities. The characterization and evaluation of genetic resources is one of the proposed ways in which this goal could be attained (National Department of Agriculture: Genetic resource management, 2008).

In order to maintain the characteristics of these locally developed breeds and ensure that their unique traits are not lost through continuous selection and cross-breeding, it has become important to revisit current breed standards and introduce genetic characterization to obtain genetic diversity parameters.

The objective of this paper was therefore to review the potential of locally developed meat type goats as a genetic resource with reference to their phenotypic and genetic characteristics.

Material and Methods

Phenotypic and genetic characteristics

Goats are well adapted to the drier western region of South Africa and northern regions where bush encroachment is a problem (Figure 1). It is also in these areas where the Boer, Kalahari Red and Savanna goats are primarily farmed in commercial systems for meat production.

Of the three breeds the Boer goat has the longest official history with the formulation of breed standards in 1959 when the South African Boer Goat Association was founded (Campbell, 2003). The breed standards specify a red head and the Boer goat replaced many unimproved local strains of varying colours with the strict selection for a white body and red head depicted in figure 2. Boer goats are large, long-legged goats with short, soft hair and long ears. These goats have a sturdy head with a compressed nose and strong horns that have a gradual backward curve. They have fleshy,



Figure 1. *Distribution of the meat type goat breeds.*

well-developed broad briskets, well-sprung ribs, broad backs and muscular legs (Sambraus, 1992) and well fleshed buttocks and thighs. The mature Boer goat buck weighs between 110 and 135 kg and does weigh between 90 and 100 kg (Malan, 2000). The Boer goat does are known for their good mothering ability (Figure 2) and can kid every seven to eight months. They have an exceptional ability to resist and survive diseases such as blue tongue, prussic acid poisoning and, to a lesser extent, enterotoxaemia (Malan, 2000; Erasmus, 2000).

The Kalahari Red (Figure 3) is believed to have originated from two lines, namely a line of red-headed Boer goat and another of the 'unimproved' local goats in South Africa (Campbell, 2003). The Kalahari Red have a distinct red colour, shown in Figure 3, and are often used in crossbreeding to produce goats with a uniform, solid, red colour. They are fully pigmented and are able to endure heat and intense sunlight, as their dark coats and long ears provide good heat resistance (*http://studbook.co.za*). The breed has excellent walking ability and good mothering ability and they can kid three times in two years.

The white Savanna goat (Figure 4), also known as the white Boer goat, was developed from indigenous goats of Southern Africa during the past few decades (Campbell, 2003). The breed standards allow limited red, blue and black hairs (http://ourfarmsite.com/web/goats/goatsavanna/ breedstandards.html). The Savanna goat has short kempy white hair with a black skin, horns, nose and udder, and during the winter the goats develop extra fluffy cashmere hair for protection. Their heads are fairly long and slightly curved with big, oval shaped ears. The forequarter is well muscled and of medium width, with a reasonably long neck for easy browsing. Does have excellent mothering traits and about 22% of their offspring are born as twins and triplets, under extensive conditions. The Savanna and Kalahari Red goats have been recognised in South Africa as official breeds since 1993 and 1990, respectively.

Morphometric traits

Morphometric data was collected from goats with an average age of 12 months in a recent study in 2007. The measurements taken and their full descriptions are reported in table 1. Goats were sampled from a number of stud herds in order to be representative of the different breeds (sample collection are indicated in figure 1) and linear traits were measured for a total of 42 Boer goats, 47 Kalahari Reds and 49 Savanna goats. The goats

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Figure 2. Typic South African Boer goats with a white body and red head and neck.



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Figure 3. Kalahari Red goats with a uniform red coat colour.



Figure 4. Typic flock of white Savanna goats.

Table 1. Description of morphometric measurements.

Measurement ¹	Description
Height (H)	Vertical distance to the ground, measured behind the whithers
Length (L)	From the anterior shoulder point to the posterior extremity of the pin bone
Loin width (LW)	Measured just in front of the haunches (os coxea)
Depth (D)	Between Corpus Sterni and the point between the shoulder blades
Heart girth (HG)	Measured just behind the shoulder
Hock length (HL)	From the front to back margin
Head width (HW)	Distance between eyes
Head length (HL)	Length of head
Neck circumference (N)	Circumference
Tail length (TL)	From the tail base to tip
Pelvic width (PW),	Between pelvis bones
Pelvic length (PL)	From pelvic bone to os coxea

¹A tape measure and measuring-rod were used to obtain various body measurements (in cm).

were also classified according to qualitative characteristics including beardedness (absence or presence of beard), incidence of horns (horned or polled) and coat colour. A General Linear Model (GLM) procedure of SAS (1992) was applied for analyses of the different traits measured.

Genetic characterization

A genetic characterization of the local meat type breeds has been undertaken by the Department of Animal and Wildlife Sciences at the University of Pretoria. The blood samples were collected from the

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same areas as described for the phenotypic measurements. For the genetic characterization additional flocks were included to ensure that unrelated samples are collected as far as possible. The Boer goats (62) represented three areas, two different flocks from the Northern Cape (31), and two experimental populations of Boer goats (31 Eastern Cape and 17 Limpopo provinces) not subjected to selection. The 60 blood samples for the Kalahari Red were collected from six flocks and the 55 Savannah samples from two different flocks in the Northern Cape. Eighteen microsatellite markers were selected from a panel of markers recommended by the International Society for Animal Genetics (ISAG) for application in diversity studies. These markers were selected based on the degree of polymorphism and genome coverage (Table 2) and available on web at: *lprdad.fao.org/cgi-bin/ getblob.cgi?sid=4d60c0a342d8413c49acc b8c1837ebf5,50005880* and background at *www.isag.org.uk/ISAG/all/ ISAG2004_ISAGFAO_AGAGD.pdf;ftp://ftp.fao.org/*

Table 2. Microsatellite marker information applied in this study.

Microsatellite marker	Chromosome number	Fluorescent	Product size range	Sequence
SRCRSP24	Unknown	Fam	162 – 174	F 5'- AGC AAG AAG TGT CCA CTG ACA G-3'
				R 5' –TCT AGG TCC ATC TGT GTT ATT GC-3'
SRCRSP5	21	Tet	171 -183	F 5' -GGA CTC TAC CAA CTG AGC TAC AAG- 3'
				R 5 –TGA AAT GAA GCT AAA GCA ATG C-3'
SRCRSP8	Unknown	Tet	210 - 260	F 5' -TGC GGT CTG GTT CTG ATT TCA C-3'
				R 5' -CCT GCA TGA GAA AGT CGA TGC TTA G- 3'
MCM527	5	Hex	155 – 173	F 5' -GTC CAT TGC CTC AAA TCA ATT C- 3'
				R 5' -AAA CCA CTT GAC TAC TCC CCA A- 3'
INRA 23	3	Tet	208 - 214	F 5' -GAG TAG AGC TAC AAG ATAA AAC TTC- 3'
				R 5' –TAA CTA CAG GGT GTT AGA TGA ACT CA- 3'
BM1329	6 (sheep)	Tet	168 – 182	F 5'TTG TTT AGG CAA GTC CAA AGT C- 3'
				R 5' – AAC ACC GCA GCT TCA TCC- 3'
OARFCB20	2	Tet	99 – 125	F 5' – AAA TGT GTT TAA GAT TCC ATA CAG TG- 3'
				R 5' – GGA AAA CCC CCA TAT ATA CCT ATA C- 3'
CRSRD247	14	Fam	236 - 244	F 5' – GGA CTT GCC AGA ACT CTA CAA T- 3'
				R 5' -CAC TGT GGT TTG TAT TCA GG- 3'
ILST087	28	Fam	145 – 165	F 5' – AGC AGA CAT GAT GAC TCA GC- 3'
				R 5' -CTG CCT CTT TTC TTG AGA GC- 3'
SRCRSP23	Unknown	Fam	83 – 111	F 5' – TGA ACG GGT AAA GAT GTG -3'
				R 5' -TGT TTT TAA TGG CTG AGT AG- 3'
OARFCB11	2	Hex	142 - 150	F 5' -GGC CTG AAC TCA CAA GTT GAT ATA TCT ATC AC- 3'
				R 5' -GCA AGC AGG TTC TTT ACC ACT AGC ACC- 3'
ILST002	Unknown	Hex	118 – 127	F 5' -TCT ATA CAC ATG TGC TGT GC- 3'
				R 5' -CTT AGG GGT GAA GTG ACA CG- 3'
RM004	15	Tet	138 – 146	F 5' –AG CAA AAT ATC AGC AAA CCT- 3'
				R 5' -CCA CCT GGG AAG GCC TTT A- 3'
INRA63	18	Fam	174 – 190	F 5' –ATT TGC ACA AGC TAA ATC TAA CC- 3'
				R 5' –CCA CCT GGG AAG GCC TTT A- 3'
INRA006	3	Hex	109 – 123	F 5' –AGG AAT ATC TGT ATC AAC CTC AGT C- 3'
				R 5' -CTG AGC TGG GGT GGG AGC TAT AAA TA- 3'
MAF65	15	Tet	117 – 127	F 5' – AAA GGC CAG AGT ATG CAA TTA GGA G- 3'
				R 5' – CCA CTC CTC TGA GAA TAT AAC ATG- 3'
BM1258	23	Fam	102 – 106	F 5' -GTA TGT ATT TTT CCC ACC CTG C- 3'
				R 5' – GAG TCA GAC ATG ACT GAG CCT G(AT)- 3'

¹Dye colour of fluorescent label Fam = Blue; Tet = Green and Hex = Yellow.

docrep/fao/010/a1250e/annexes/

Reports%20from%20International%20Organizations/ISAG.pdf.

PCR reactions were carried out in a volume of 8.2 µl, containing 50 ng target DNA, 3.48 µM dNTP's, 10 mM TrisHCl (pH = 9.0), 1.5 mM MgCl₂, 1.74 unit Taq Polymerase enzyme (Goldstar) and 4.54pmol of each primer (microsatellite marker). Preparation of samples was followed by thermal cycling in a Thermal Controller (Perkin Elmer) using the following program: 12 minutes at 94°C followed by 33 cycles consisting of 45 sec at 94°C, 80 sec at 60 °C, 60 sec at 72°C and an extension step of 60 min at 72°C. GENESCAN-350 TAMRA was used as internal standard and samples were analyzed on an automated DNA-Sequencer (ABI 377). The Genescan version 2.0 and Genotyper for MacIntosch were used to determine the fragment sizes in base pairs. Hardy Weinberg Equilibrium, heterozygosity and effective number of alleles were calculated using POPGENE 3.2 (Yeh *et al.*, 1999) and Microsattellite Toolkit (Park, 2001). A factorial correspondence analysis was performed to

illustrate the relationship among the breeds using Genetix 4.03 (Belkhir *et al.*, 1996).

Results

Of the goats measured 13% of the Boer goats, 23% of the Kalahari Red and 25% of the Savanna goats were bearded. Seventy five percent of all the Boer goats had white bodies with red heads, while the remaining 25% had either speckled heads or a red spot on the body. The Kalahari Red goats were primarily red coated with a white or black spot appearing on the body of 15% of the population. Savanna goats were all white. Morphometric measurements analyzed were presented in table 3.

All the microsatellite markers tested were found to be in Hardy Weinberg Equilibrium, except for some markers in the Savanna goats.

The effective number of alleles showed little variation, from 2.9 in the Boer goat to 3.4 in the Kalahari Red goats. Preliminary heterozygosity



Figure 5 Assignment of individuals to the respective populations using a Factorial Correspondence Analyses.

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	, , ,	Body mea	asurements	Least square	Standard
Trait	Breed	Min.	Max.	means ¹	error (±)
Haiakt	Boer Goat	47.0	67.5	56.5ª	0.5
rieight (LI)	Kalahari Red	49.3	60.2	54.1ª	0.5
(П)	Savanna	47.8	61.9	55.7ª	0.5
Lonoth	Boer Goat	55.0	79.5	68.2 ^{ab}	0.8
(I)	Kalahari Red	60.6	83.2	69.8ª	0.8
(L)	Savanna	30.0	71.1	64.9 ^b	0.7
Donth	Boer Goat	22.2	33.0	26.4 ^{ab}	0.3
(D)	Kalahari Red	23.0	31.8	27 .1ª	0.3
(D)	Savanna	22.4	29.0	24.9 ^b	0.3
Hoort Cinth	Boer Goat	78.0	112.0	90.3 ^{ab}	1.0
HC)	Kalahari Red	65.6	114.0	95.3ª	1.0
(11G)	Savanna	72.6	100.2	86.5 ^b	1.0
Hock Longth	Boer Goat	16.5	34.0	28.3ª	0.5
(HI)	Kalahari Red	15.2	36.1	27.7ª	0.5
(11L)	Savanna	18.1	29.1	23.5 ^b	0.5
Hoad Width	Boer Goat	4.8	10.5	7.7ª	0.2
	Kalahari Red	4.8	8.9	6.6 ^b	0.3
(11)	Savanna	4.2	7.3	5.6°	0.2
Hoad Longth	Boer Goat	13.0	21.0	17.2ª	0.3
(HI)	Kalahari Red	9.8	21.6	15.7 ^b	0.2
(111)	Savanna	13.1	20.0	15.8 ^{ab}	0.2
Neck	Boer Goat	33.2	57.0	48.3ª	4.0
Circumference	Kalahari Red	33.7	58.5	42.5 ^{ab}	3.8
(N)	Savanna	30.4	65.4	37.7 ^{ab}	3.7
Tail Longth	Boer Goat	6.0	17.0	12.2 ^{ab}	0.3
(TI)	Kalahari Red	10.4	19.2	13.2ª	0.3
(1L)	Savanna	9.6	19.0	13.3ª	0.3
Pelvic Width	Boer Goat	7.8	21.4	13.8ª	0.3
(PW)	Kalahari Red	9.0	13.5	11.1 ^b	0.3
	Savanna	8.4	17.7	11.4 ^b	0.3
Polyic Longth	Boer Goat	13.5	26.5	19.7ª	0.4
(PI)	Kalahari Red	16.0	24.8	20.5ª	0.3
(1 L)	Savanna	16.7	27.4	19.1ª	0.3
Ear Length	Boer Goat	14.4	26.0	21.4ª	0.3
(EL)	Kalahari Red	14.0	18.5	19.2 ^b	0.2
	Savanna	16.3	22.7	19.5 ^b	0.2

Table 3. Statistical description of body measurements (cm) recorded for 42 Boer goats , 47 Kalahari Red and 49 Savanna goats.

¹Superscripts differ significantly for P < 0.001.

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	Mean n of		Unbiased Hz	Observed H
Breeds	samples	Ne ¹	(SD)	(SD)
S A Boer goat	31	2.9	0.57 (0.19)	0.54 (0.22)
Boer goats EC ²	31	3.3	0.65 (0.15)	0.62 (0.20)
Boer goat ³	17	3.1	0.62 (0.17)	0.66 (0.35)
Kalahari-Red	60	3.3	0.68 (0.10)	0.63 (0.13)
Savanna	55	3.4	0.69 (0.10)	0.61 (0.08)

Table 4. Number of samples, effective number of alleles and expected and observed heterozygosity for the three breeds.

 1 Ne = Effective number of alleles.

² = Experimental population of Boer goats Eastern cape province.

³ = Experimental population of Boer goats Limpopo province.

values calculated for this study were all above 50% with the lowest value for the SA Boer goat (57%) and a slight difference between the Savanna (68%) and Kalahari Red (69%) (Table 4). Both the experimental flocks of SA Boer goat had higher heterozygosity values than the commercial flocks of the Northern Cape Province.

In figure 5 the results of a Factorial Correspondence Analysis are shown with the assignment of the individual goats analysed into three groups.

Discussion

Goats have been largely neglected as an animal genetic resource in South Africa when compared to cattle and sheep. Despite a number of research initiatives undertaken over the past 40 years on goat production, there are still aspects that require attention and a need for effective coordination of research programs to lead to the successful conservation and utilization of goats. The South African Boer goat, Savanna and Kalahari Red goats have only recently been classified as distinct breeds, while their phenotypic and genetic characterization are still not well defined.

Both the Boer goat and Savanna breeds have established breeders associations while a breeders club was founded for the Kalahari Red goats with set breed standards (*www.studbook.co.za/telers*). The breed standards consist of primarily phenotypic descriptions, colour and culling defects. Attention is also given to type traits related to functional efficiency. Except for the colour variation between the three breeds, the breed standards are quite similar and the aim is to select for a well adapted, functionally efficient meat type goat breed.

Morphometric measurements recorded for this study (reported in table 3) indicate a larger within breed variation (minimum and maximum) than between the breeds. Statistical differences were observed between the Savanna and the other two breeds for depth and length but not (P < 0.001) for height. The only trait where significant differences were observed among all three breeds was for head width, where the Boer goat had the broadest head and the Savanna the narrowest. The Boer goat had the largest pelvis with and the longest ears compared to the other two breeds (P < 0.001). The differences in morphometric traits among the breeds are fairly insignificant and highlight the need for a genetic characterization to be able to distinguish the breeds accurately on a genotypic level and to ensure sufficient diversity within the breeds for long term conservation. The heterozygosity values estimated for the three breeds are relatively high and correspond with the observations for local South African breeds by Visser *et al.* (2004) and values reported in studies on goats by Martinez et al., 2006; Els et al., 2004; Li et al., 2002, but higher than those reported by Kumar et al, (2005) for Marwari goats. It is important to note that both the experimental populations of Boer goats not subjected to selection for improved performance had a higher genetic variation than the Boer goats sampled from the breeders (Table 4). The sample size of the goats sampled from the Limpopo flock was small and more samples need to be tested for further analyses. According to the Factorial Correspondence Analysis (Figure 5) the three breeds assign to three groups. The Kalahari Red goats can be distinguished as a group, while the SA Boer goat and Savanna tend to overlap. More samples of indigenous goat types and Angora mohair goats are currently being added for further analyses of genetic diversity and relatedness for a more detailed characterization of all South African goats.

The meat goat breeds are often noted in literature as having adaptive characteristics in terms of heat tolerance and feeding behavior (Erasmus, 2000; Malan, 2000). There is an urgent need to investigate these traits to ensure conservation of the unique genetic resource. Shrestha and Fahmy (2005) reported that nearly 800 farm animal genetic resources have already been lost and about 30 percent of the remaining species are associated with some degree of risk. The FAO predicts that 20 percent of livestock breeds are at risk of extinction and more than half of these breeds are likely to be found in developing countries (FAO, 2007).

Conclusion

Goat meat production in South Africa is relatively low compared to beef and mutton, but it plays an important role in African cultural activities and is the preferred meat in certain communities (Webb *et al.*, 2005). In addition to the local production, both SA Boer goat and Kalahari Red have been exported to Europe and the USA. It is important that breed standards for the locally developed breeds be revised and genetic analyses included to ensure that superior genetics will be conserved. Adaptive phenotypic characteristics of both the meat type breeds and indigenous types require further investigation for the long term conservation of goats as a unique animal genetic resource in South Africa.

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Genetic diversity and zebu genes introgression in cattle population along the coastal region of the Bight of Benin

O. D. Koudandé¹, G. Dossou-Gbété¹, F. Mujibi², H. Kibogo², D. Mburu², G.A. Mensah¹, O. Hanotte² & J.A.M. van Arendonk³

¹Institut National des Recherches Agricoles du Bénin, 01 BP 884, Cotonou, Benin ²International Livestock Research Institute, (ILRI) P. O. Box 30709, Nairobi, Kenya ³Animal Breeding and Genetics Group, Wageningen Institute of Animal Sciences, Wageningen University, P.O. Box 338, 6700 AH Wageningen, The Netherlands

Summary

Genetic diversity and Zebu genetic introgression have been assessed in five subpopulations of cattle along the coastal region of Togo, Benin and Nigeria using 15 autosomal and one Y- specific microsatellite markers. Mean observed heterozygosity (Ho) ranges from 0.55 to 0.61 and the mean number of alleles (MNA) from 5.47 to 6.47. Genetic differentiation indexes (F_{st}) , were significant between the five subpopulations (P < 0.01). Some possible population diagnostic alleles are identified with allele 254 at locus ILSTS033 and allele 182 at locus ILSTS005 found only in the population from Togo with frequencies of 5.41% and 12.82% respectively. Allele 226 of locus ILSTS103 is fixed in the Togolese population (100%) and almost fixed (98.75%) in the Benin-Valley population. Y chromosome analysis reveals male Zebu introgression in all five populations with a frequency of indicine Y chromosome ranging from 37.5% in Benin-Valley and Benin Plateau East to 100% for Benin Plateau West. Admixture analysis using the programme STRUCTURE (k = 2) confirms phenotypic observations suggesting different level of taurine background and therefore Zebu introgression amongst the populations. Within populations, variations in levels of Zebu admixture between herds were also detected. Whereas the valley population from Benin shows low level of Zebu introgression, it is the population from Benin Plateau East which is the purest.

Résumé

La diversité génétique et l'introgression de gènes de zébu dans la population de bovins le long de la côte du golfe de Guinée allant du Togo au Nigeria ont été évaluées à partir de 15 marqueurs microsatellites autosomes et un marqueur spécifique du chromosome Y. La moyenne des hétérozygotes Ho varie de 0,55 à 0,61 et le nombre moyen d'allèles (NMA) de 5,47 à 6,47. Les indices de différenciation génétique (F_{-t}) sont différents entre les cinq sous populations (*P*<0,01). Des allèles spécifiques de population sont identifiés aux loci ILSTS033 (allèle 254) et ILSTS005 (allèle 182) dans la sous population du Togo avec des fréquences respectives de 5,41 et 12,82%. L'allèle 226 du locus ILSTS103 est totalement (100%) fixé dans la sous population togolaise et presque (98,75%) dans la sous population de la vallée au Bénin. L'évaluation du marqueur spécifique du chromosome Y révèle l'introgression de gènes de zébu dans les cinq sous populations allant de 37,5% dans la vallée du Bénin et sur le Plateau Est à 100% sur le Plateau Ouest. L'analyse de mélange utilisant le programme STRUCTURE (k = 2) confirme les observations phénotypiques concluant à l'introgression de gènes de zébu au sein des populations. Des variations de mélange sont observées entre troupeaux au sein d'une même sous population. Alors que les animaux dans la vallée au Bénin présentent un niveau faible d'introgression de gènes de zébu, c'est la sous population du Plateau Est qui paraît être la plus pure.

Resumen

La diversidad genética y la introgresión de genes de zebú en la población de bovinos a lo largo de la costa del golfo de Guinea que va desde Togo hasta Nigeria han sido evaluados a partir de 15 marcadores microsatelitares autosomos y un marcador específico del cromosoma Y. La media de heterocigosis Ho varía de 0,55 a 0,61 y el número medio de alelos (NMA) de 5,47 a 6,47. Los índices de diferenciación genética (F_{st}) son diferentes entre las cinco sub-poblaciones (P<0,01). Han sido identificados alelos específicos de poblaciones en los loci ILSTS033 (alelo 254) e ILSTS005 (alelo 182) en la sub-población del Togo con frecuencias respectivas de 5,41 y 12,82%. El alelo 226 del locus ILSTS103 está totalmente fijado (100%) en la sub-población togolesa y casi (98,75%) en la sub-población del valle del Benín. La evaluación del marcador específico del cromosoma Y revela la introgresión de genes de zebú en las cinco sub-poblaciones que van de 37,5% en el valle del Benín y en la Meseta Este, hasta 100% en la Meseta Oeste. El análisis de mezcla utilizando el programa STRUCTURE (k=2) confirma las observaciones fenotípicas que indican la introgresión de genes de zebú en el interior de las poblaciones. Variaciones de mezclas han sido observadas entre rebaños al interior de una misma población. Mientras que los animales del valle del Benín presentan un nivel de introgresión de genes de zebú bajo, la sub-población de la Meseta Este parece ser la más pura.

Key words: Cattle, Microsatellites, Genetic diversity, Zebu genes introgression.

Introduction

West African taurine cattle (B. taurus) breeds are under threat of extinction due to uncontrolled breeding practices by farmers aiming to enhance the size and the productivity of their animals. However, it is recognised that animals of these breeds are well adapted to their local environments and remain productive in areas where Zebu or crossbreeds are often unable to survive (Agyemang et al. 1991; Uza, 1997). The phenotypic traits of these animals are the result of hundreds of years of natural selection in relation to the local environments under permanent tsetse, ticks and helminth challenge. It is clear that uncontrolled crossbreeding with exotic breeds poses a threat to the adaptive traits of this unique animal genetic resource. This uncontrolled crossbreeding is the result of poor agricultural policies with no long term breeding goals, each farmer developing his own objective with emphasis on short-term results.

In the early eighties, the Food and Agriculture Organisation of the United Nations started alerting public opinion on the subject of animal genetic resources conservation and management (FAO, 1981). At the Earth Summit in 1992 in Rio de Janeiro, 157 countries signed the United Nations convention on biological diversity and subsequently FAO (1992) launched a special program for the global management of farm animal genetic resources. The aim of this program is to maintain in each species a maximum genetic diversity of the gene pool to allow for future unforeseen needs in the development of sustainable animal production systems. In addition the program aims at prompt actions to preserve animal breeds at risk of extinction.

Studies in the last decade have shown that African cattle were most likely domesticated within the African continent, separately from the other centres of domestication in the Fertile Crescent and in the Indian sub-continent, from the wild African auroch *B. primigenius* (Bradley *et al.*, 1996). Their separate origin indicates that they represent a unique set of genetic characteristics. Today, the remaining pure African taurine cattle are only found within the West African taurine cattle living in tsetse-infected areas. These populations, given their origins and their adaptation to local environmental conditions (e.g. disease resistance), represent a unique genetic resource.

Introgression with Zebu cattle in West African taurine cattle populations, however, is common (MacHugh et al., 1997; Hanotte et al., 2000) and is diluting progressively the African taurine genetic background of these breeds. Ultimately, this introgression will result in the loss of the unique genetic adaptation of these breeds. Currently, the extent and pattern of Zebu introgression into the indigenous taurine populations is well described by Hanotte et al. (2002). This study, however, did not include the Lagune breed. The current paper is therefore a complementary study to that of Hanotte et al. (2002), targeting the Lagune breed in the Bight of Benin. Results of this study can be used to identify suitable pure West African taurine populations for their inclusion in breeding programs that aim at conserving and utilising these unique indigenous genetic resources. These results could also be the starting point for the implementation of a sustainable breeding programme for livestock production in that region.

Material and Methods

Sampling

Blood samples were collected from different cattle subpopulations in Benin, Nigeria and Togo. Forty animals, of which eight were males, were sampled for each identified subpopulation. Three subpopulations have been sampled in Benin, which are Benin Plateau West (BPW), Benin Valley (BV) and Benin Plateau East (BPE). Two neighbouring regions of the target area in Benin have been considered (Figure 1) in Nigeria (NG) and Togo (TG) resulting in a maximum of 200 sampled cattle.

Appropriate materials such as Wattman filter paper, evacuated blood collection tubes (5 ml) and Eppendorf tubes for stocking buffy coats were used for handling samples. Blood samples were collected from each animal in two evacuated blood collection tubes containing 3.2% EDTA and the buffy coat (white cells) was taken after spinning the tubes at 2400 rpm at ambient temperature. Collected buffy coats were added with 8M urea solution and kept at ambient temperature. To be on the safe side, blood samples were also collected on Wattman paper and dried at 40°C in oven, then stored at ambient temperature. Sampling included as many animals from different herds as possible to avoid direct relationship between sampled individuals. In other terms, one male and four females were sampled per herd and a total of eight herds were visited for each identified subpopulation. All samples were conveyed to the laboratory of genetics at ILRI campus in Nairobi (Kenya) for processing.

Laboratory procedures

DNA was extracted from the buffy coats with standard procedures as described by Sambrook *et al.* (1999). Sixteen markers, of which one was on the Y chromosome, were used for genotyping. They were *TGLA126, TGLA122, ILSTS033, ILSTS013, ILSTS008, ILSTS005, ILSTS006, ILSTS036, ILSTS028, ILSTS023, ILSTS103, ETH152, BM2113, BM1824, AGLA293* and *INRA124*. Polymerase Chain Reaction (PCR) for amplification of microsatellite loci were performed as described in Hirano *et al.* (1996) and Kemp *et al.* (1995).

Genotyping and gel analysis were performed using automatic DNA sequencer ABI 377 and associated analysis software GenescanTM (version 3.1) and GenotyperTM version 2.0).

Data analysis

Marker allele frequencies have been analysed using the Microsatellite toolkit program (Park, 2001). The genetic diversity of each population was evaluated by computing the mean number of alleles per locus (MNA), the observed heterozygosity (H_{obs}), and the expected heterozygosity (H_{exp}) using the microsatellite toolkit (Park, 2001) as well as GENEPOP, version 3.3 (Raymond and Rousset, 1995). Subpopulations have been checked for HWE using the exact probability test with GENEPOP. The level of genetic differentiation among populations was examined using *F*-statistics (Weir and Cockerham, 1984). Genetic distances between populations D_A (Nei *et al.*, 1983) were calculated and used to construct a phylogenetic tree



Figure 1.Benin map showing the sampled populations locations.

with the neighbour joining algorithm (Saitou and Nei, 1987). The level of Zebu introgression was estimated with the programme STRUCTURE using the ADMIX 1.0 software (Bertorelle and Excoffier, 1998). Correlations between phenotype and populations structure results were performed using Microsoft Excel.

Results

Genetic variation

From the five subpopulations under study here, 130 alleles were found for all 15 autosomal marker loci that have been genotyped over the 200 sampled animals (Table 1). The total number of alleles detected per locus ranged from 4 to 16 resulting in a mean number of alleles (MNA) per locus of 8.67. The number of alleles summed over the 15 loci per subpopulation ranged from 82 in BPE to 97 in TG (Table 1) giving a MNA per locus for each subpopulation varying from 5.47 to 6.47 (Table 2). These variations are not strong enough (P>0.05) to differentiate current subpopulations. However, allele 254 of the locus ILSTS 033 and allele 182 of ILSTS_005 are found solely in TG with a frequency of 5.41% and 12.82% respectively. Alleles 171 and 175 of *ILSTS_023* are equally distributed (50%)

within TG and BV subpopulations. Further, allele 226 of locus *ILSTS_103* is fixed in TG (100%) whereas it is almost (98.75%) fixed in BV.

The expected heterozygosity computed over the 15 loci for each of the five subpopulations ranged from 0.593 (BV) to 0.673 (BPW) whereas the observed heterozygosity varied from 0.55 in BPE to 0.61 in BPW (Table 2).

None of the subpopulations deviated from HWE through all loci at 5 p. cent level. Out of the 15 loci under study, only loci *ILSTS006* (*P*<0.006), *ILSTS023* (*P*<0.005) and *ILSTS103* (*P*<0.05) deviated consistently from HWE for all five subpopulations (Table 3). Except those three loci, TG deviated (*P*<0.05) from equilibrium at two loci (*BM1824*, *BM2113*), BPW at four loci (*TGLA126*, *ILSTS033*, *ILSTS008*, *BM2113*), BV at two loci (*ILSTS008*, *ILSTS033*), BPE at three loci (*ILSTS033*, *ILSTS036*, *BM2113*), and NG at two loci (*ILSTS013*, *ILSTS036*).

Genetic distances and breed relationships

Table 4 summarizes genetic distances and gene differentiation indices between all pairs of subpopulations. The largest genetic distance (0.1183) was obtained between TG and BPE, whereas the smallest (0.0666) was obtained between

			Benin		_	
Locus	Togo	BPW	BV	BPE	Nigeria	Total
TGLA126	7	7	6	6	6	7
TGLA122	8	8	9	7	8	11
ILSTS033	7	5	4	7	6	9
ILSTS013	4	4	4	4	4	4
ILSTS008	4	5	4	3	3	5
ILSTS005	5	4	3	4	3	5
ILSTS006	8	5	8	5	5	9
ILSTS036	12	8	9	8	9	16
ILSTS028	6	8	6	6	7	10
ILSTS023	2	5	2	4	5	6
ILSTS103	1	5	2	4	5	7
ETH152	6	5	7	5	5	8
BM2113	12	10	7	6	9	12
BM1824	6	4	4	4	4	6
AGLA293	9	8	12	9	8	15
Total	97	91	87	82	87	130

Table 1. Total number of alleles per locus and per subpopulation.

 $^{2}\text{H}_{obs}$ (±SE) Populations 1MNA (±SE) $3H_{exp}$ (±SE) Togo 6.47 (±3.14) 0.5905 (±0.0202) 0.6410 (±0.0539) Benin Plateau West 6.07 (±1.91) 0.6083 (±0.0200) 0.6725 (±0.0322) Benin Valley 5.80 (±2.88) $0.5604 (\pm 0.0204)$ $0.5930 (\pm 0.0503)$ Benin Plateau East 0.6194 (±0.0296) 5.47 (±1.73) $0.5504 (\pm 0.0204)$ Nigeria 5.80 (±2.01) 0.5603 (±0.0203) 0.6338 (±0.0383)

Table 2. Comparison of genetic variability of five populations of cattle along the coastal region of Togo, Benin and Nigeria.

¹Mean number of alleles per locus.

²Observed heterozygosity.

³Expected heterozygosity.

Table 3. Hardy-Weinberg exact probability test by population.

			Benin		
Locus	Togo	BPW	BV	BPE	Nigeria
TGLA126	0.1266 (0.0094)	0.0324 (0.0058)	0.9689 (0.0032)	0.3372 (0.0138)	0.1626 (0.0086)
TGLA122	0.0918 (0.0143)	0.1416 (0.0109)	0.5619 (0.0241)	0.3201 (0.0204)	0.2617 (0.0191)
ILSTS033	0.0122 (0.0035)	0.0000 (0.0000)	0.0302 (0.0026)	0.0018 (0.0009)	0.2594 (0.0157)
ILSTS013	0.3394 (0.0082)	0.2777 (0.0082)	0.9153 (0.0024)	0.5509 (0.0065)	0.0219 (0.0018)
ILSTS008	0.3769 (0.0107)	0.0052 (0.0014)	0.0024 (0.0007)	0.0842 (0.0049)	0.8024 (0.0036)
ILSTS005	0.1651 (0.0080)	0.0052 (0.0071)	0.3149 (0.0082)	0.7353 (0.0076)	0.6628 (0.0049)
ILSTS006	0.0000 (0.0000)	0.0000 (0.0000)	0.0000 (0.0000)	0.0057 (0.0018)	0.0000 (0.0000)
ILSTS036	0.0797 (0.0167)	0.1021 (0.0100)	0.1006 (0.0160)	0.0294 (0.0053)	0.0001 (0.0001)
ILSTS028	0.0805 (0.0088)	0.1186 (0.0129)	0.2908 (0.0173)	0.2782 (0.0152)	0.3055 (0.0202)
ILSTS023	0.0000 (0.0000)	0.0001 (0.0001)	0.0000 (0.0000)	0.0000 (0.0000)	0.0000 (0.0000)
ILSTS103	-	0.0003 (0.0003)	-	0.0000 (0.0000)	0.0183 (0.0035)
ETH152	0.0672 (0.0065)	0.4927 (0.0091)	0.3976 (0.0174)	0.9354 (0.0049)	0.3382 (0.0115)
BM2113	0.0396 (0.0076)	0.0110 (0.0028)	0.2903 (0.0158)	0.0089 (0.0016)	0.3507 (0.0163)
BM1824	0.0414 (0.0064)	0.9355 (0.0020)	1.0000 (0.0000)	0.8824 (0.0045)	0.0616 (0.0055)
AGLA293	0.5448 (0.0222)	0.6256 (0.0170)	0.2169 (0.0274)	0.0676 (0.0097)	0.0907 (0.0103)

NG and BPW. This result is in contrast to what would be expected given the geographic location of these subpopulations (Figure 1). The smallest coefficient of differentiation was obtained between TG and BPW, and the highest was between TG and BPE which is in agreement with the largest genetic distance obtained. The F-statistics results showed that the five populations were significantly different from each other (*P*=0.0001). Estimates of F_{sT} for each locus and for each population pair are summarized in Table 5.

The neighbour-joining tree based on genetic distance D_A is represented in figure 2. It shows the existing relationships among the five subpopulations according to studied loci. As it

would be expected, BV and BPE are close subpopulations as shown in this figure.

Population structure (admixture)

Genetic admixture analysis shows that all five subpopulations are introgressed with Zebu genes but to different degrees (Figure 3), whereas subpopulations from TG and BPW harbour high levels of introgression at 0.753 and 0.651, respectively. BPE is the least introgressed with introgression coefficients equal to 0.188. BV and NG have intermediate coefficient, 0.309 and 0.390, respectively. On individual basis, the level of

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Table 4. Genetic distance between five cattle subpopulations along the coastal region of Togo, Benin and Nigeria calculated using D_A (Nei et al., 1983) below diagonal and Weir and Cockerham (1984) F_{ST} above the diagonal.

ан- С	Togo	Benin Plateau West	Benin Vallev	Benin Plateau East	Nigeria
Togo	-	0.0205	0.0341	0.0564	0.0493
Benin	0.0795	-	0.0349	0.0310	0.0258
Plateau West					
Benin Valley	0.0866	0.0917	-	0.0335	0.0463
Benin	0.1183	0.0740	0.0809	-	0.0377
Plateau East					
Nigeria	0.1103	0.0666	0.0914	0.0688	-



Figure 2. Neighbor joining dendogram showing the genetic relationship among the five sampled subpopulations along the coastal region from Togo to Nigeria.



Figure 3. Admixture analysis showing zebu introgression in sampled subpopulations.

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Loci	TG/BV	TG/BPE	TG/NG	TG/BPW	BV/BPE	BV/NG	BV/BPW	BPE/NG	BPE/BPW	NG/BPW
TGLA126	0.0407	0.0369	0.0170	0.0060	0.0006	0.0006	0.0375	0.0538	0.0346	0.0118
TGLA122	0.0303	0.1558	0.0162	0.0340	0.0498	0.0256	0.0099	0.1476	0.0879	0.0096
ILSTS033	0.0302	0.0583	0.0703	0.0208	-0.0015	0.0268	0.0213	0.0659	0.0757	0.0190
ILSTS013	0.0049	0.0095	0.0432	-0.0055	-0.0007	0.0009	0.0316	0.0280	0.0346	0.0847
ILSTS008	0.0061	0.0561	0.0333	-0.0032	0.0053	-0.0101	0.0024	-0.0023	0.0607	0.0256
ILSTS005	0.0627	0.0315	0.0523	0.0442	-0.0043	-0.0120	0.0072	-0.0076	-0.0033	0.0080
ILSTS006	0.0007	0.0079	0.0923	-0.0032	0.0363	0.0924	0.0056	0.1066	-0.0056	0.1163
ILSTS036	-0.0052	0.0009	0.0036	0.0006	-0.0047	0.0050	0.0053	-0.0004	-0.0012	-0.0041
ILSTS028	0.0363	0.0478	0.0132	0.0149	-0.0120	-0.0032	0.0065	0.0086	0.0062	0.0146
ILSTS023	0.0000	0.1077	0.1950	0.1281	0.1081	0.1955	0.1285	0.0183	-0.0050	0.0041
ILSTS103	0.0000	0.2046	0.1727	0.1606	0.1813	0.1465	0.1391	-0.0009	-0.0101	0.0020
ETH152	0.0193	0.0695	0.0320	-0.0119	0.0339	0.0156	0.0125	-0.0012	0.0617	0.0248
BM2113	0.0713	0.0317	0.0107	0.0032	0.0419	0.0609	0.0383	0.0234	0.0333	0.0106
BM1824	0.0846	0.0485	0.0714	-0.0037	0.0309	-0.0113	0.0562	0.0181	0.0478	0.0494
AGLA293	0.0765	0.0618	0.0026	0.0022	0.0777	0.0644	0.0688	0.0523	0.0397	0.0162
All loci	0.0341	0.0564	0.0493	0.0205	0.0335	0.0463	0.0349	0.0377	0.0310	0.0258
TG=Togo; BPW:	=Benin Plateau	West; BV=Beni	n Valley; BPE=	Benin Plateau E	last; NG=Niger	ia.				

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Figure 4. Typical Lagune cattle from Benin Plateau East (height is 81 cm; girth circumference is 112 cm).

Zebu introgression varies from 0.028 in TG to 0.967 in BPE.

Analysis of the Y chromosome reveals male Zebu introgression in males of all five subpopulations with a frequency of indicine Y chromosome ranging from 37.5% in BV and BPE to 100% for BPW. TG and NG have respectively 87.5 and 62.5%. The correlation coefficient between the admixture results and those of the Y chromosome analysis is 0.93.

Discussion

With the upgrade breeding performed by livestock producers along the coastal regions of the three countries involved in this study, the observed variation between populations was expected. Indeed, looking for taller animals, mainly for market purposes, Borgou breed and Zebu have been introduced in diverse herds (Figure 5) resulting in the higher MNA for these populations. This is the case in TG and BPW (Table 2). This result is supported by the admixture analysis which gives the higher proportion of Zebu genes to subpopulations TG and BPW. In contrast, the subpopulation BPE has the least genetic variation in terms of MNA associated with a low heterozygosity. The low genetic variation in this subpopulation could be explained by the higher level of inbreeding in this subpopulation. This explanation is supported by the management system of these animals. Farmers used to keep their animals under their own habitat during night, then they are tethered to trees and moved from time to time to access forage. Bulls are owned by few people that hire them to non-owners for breeding. This means that breeding is under control, so as to ensure these animals are kept as pure as possible for their use in certain ritual ceremonies and as a dowry. Subpopulations TG and BV, however, have in common some alleles, that is allele 171 and 175 from *ILSTS023* at 50%, allele 226 of locus *ILSTS103* at the rate of 100% for the former and 98.75% for the latter. This result is surprising given that they are separated by subpopulation BPW which has lower frequency for these alleles (results not shown).

Genetic distances between pairs of subpopulations, coefficients of differentiation and

Koudandé et al.



Figure 5. Zebu breeds that have been introduced in herds in the coastal regions.

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 F_{st} confirm that the five subpopulations are different, although some results showed patterns that were not expected. We would expect the largest genetic distance to be between TG and BPE on one side and between NG and BPW on the other. The latter showed the least distance which is in contrast to what would be expected given the geographic location of these subpopulations (Figure 1). According to the admixture analysis, TG and NG showing the highest introgression of Zebu genes would be closer - that is not the case in current results. This could be explained by different sources of gene pools that have been used to perform the introgression. The existing relationships among subpopulations through the neighbour-joining tree clearly show that BV and BPE are close subpopulations as one could predict from figure 1.

Concurrently to blood sampling, height at wither and girth circumference were measured to relate the phenotype with genotype. There is quite an agreement between the results of the admixture model analysis and the phenotype of animals (correlation coefficient equals to 0.59), that is the higher the percent of Zebu genes, the higher is the height at wither. The availability of data on control animals (pure Zebu and pure Lagune) would have better strengthened current results.

The hypothesis that animals in the valley would be less introgressed by Zebu is true, but the population of Benin Plateau East is even purer than the one targeted in our study.

Conservation futures

The major objective of this study was to determine the level of Zebu introgression within cattle populations along the coastal regions of the Bight of Benin. This region is known to originally be the cradle of the Lagune cattle breed (Figure 4), animals of small size with a wither height varying from 90 to 100 cm and a heart girth from 130 to 137 cm. As this study has demonstrated, there is a high tendency for Zebu or other breeds to be introgressed in this breed. Some herds and to some extent at least two populations, have been shown to be less polluted with Zebu blood. Throughout this study we placed an emphasis on naming cattle populations instead of breeds given the extent of crossbreeding, and current results support this point of view.

The hypothesis of the current study was to encounter cattle with low level of Zebu introgression that could be used for genetic resources conservation and it was expected to meet those animals in BV. Results of this study have shown that the purest taurine cattle were instead located in BPE. If such a conservation programme was to be undertaken, we would advise on the use of animals from BPE.

The status of BPE animals is the result of the breeding system in force in this area and for which objectives are clearly defined at the farmers' level. Therefore a sustainable way of handling the market for these animals remains to be identified. Indeed, there should be a strong marketing study that could identify market problems and define likely solutions. There should also be a sensitisation programme to make stakeholders be aware of different changes that might intervene. A contest and show program with good prizes could be a way of attracting farmers in that region to recognise the effort they are deploying on this breed. Any breeding programme, whatever is its goal, should integrate the indigenous knowledge and use of these animals. Throughout these actions, studies of the biochemical as well as the culinary quality of meat and milk produced from these animals should be undertaken to support the promotion of this breed.

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