Supplemental material

Limited Introgression Supports Division of Giraffe into Four Species

Authors:

Sven Winter^{1,2*}, Julian Fennessy³ & Axel Janke^{1,2}

Affiliations:

¹Senckenberg Biodiversity and Climate Research Centre, Senckenberganlage 25, 60325

Frankfurt am Main, Germany

²Goethe University, Institute for Ecology, Evolution & Diversity, Max-von-Laue-Str. 13, 60438

Frankfurt am Main, Germany

³Giraffe Conservation Foundation, PO Box 86099, Windhoek, Namibia

*corresponding author

sven.winter@senckenberg.de



Supplementary Fig. 1. MtDNA Neighbor-Joining tree of 217 giraffe individuals

Asterisks show major branches with a Bootstrap support > 80 % and the red frame shows potential reticulated giraffe / Nubian giraffe hybrids. Two different okapi individuals were used as outgroup.



Supplementary Fig. 2. Statistical calculations to evaluate the best fitting K in STRUCTURE (A) Mean likelihood (L(K)) and variance per K value from STRUCTURE according to (Pritchard, Wen, & Falush, 2010). (B) Delta K plot as per Evanno, Regnaut, & Goudet (2005) to find the best fitting number of populations (K) for the data. K = 4 has the highest Delta K. (C) Probability by K plot according to Pritchard, Stephens, & Donnelly (2000). K = 4 shows the highest probability.



axis 1 (31.64 % variance)

Supplementary Fig. 3. Additional PCAs based on mtDNA clades.

PCA axes 1-2 and axes 1-3 for seven giraffe mtDNA clades (1: West African; 2: Kordofan; 3: Nubian; 4: reticulated; 5: Masai; 6: Angolan; 7: South African). Colored as in Supplementary Fig. 1. The 95% confidential intervals are shown as grey colored oval outlines. Note that the confidence intervals in the PCA axes 1-2, as well as axes 1-3, indicate the same four significantly different clusters as seen in Fig. 2c.



Supplementary Fig. 4. Additional PCAs and STRUCTURE analyses for southern giraffe

(a) PCA axes 1-2 and axes 1-3 for the southern giraffe populations (1: South African giraffe; 2: Angolan giraffe). The 95% confidential intervals are shown as oval outlines. Note that the confidence intervals in the PCA axes 1-2, as well as axes 1-3, indicate no substructure within the southern giraffe.

(b) STRUCTURE analysis for the southern giraffe populations (CKGR and ENP are Angolan giraffe, the remaining populations are South African giraffe) does not show additional clustering but a high level of admixture, which contradicts the clear separation of subspecies by mtDNA. Note that detailed information about the populations are listed in Supplementary Table 1.



Supplementary Fig. 5. Additional PCAs and STRUCTURE analyses for reticulated giraffe

(a) PCA axes 1-2 and axes 1-3 for the sampled reticulated giraffe populations (1: RET; 2: RETRot; 3: RETWil; 4: ISC; 5: LWC). The 95% confidence intervals are shown as oval outlines. Note that the confidence intervals in the PCA axes 1-2, as well as axes 1-3, indicate no substructure within the reticulated giraffe.

(b) Separate STRUCTURE analysis for reticulated giraffe populations (RET, RETRot and RETWil: captive animals; ISC and LWC: wild populations) shows no additional clustering.

Note that detailed information about the populations are listed in Supplementary Table 1.



Supplementary Fig. 6. Additional PCAs and STRUCTURE analyses for northern giraffe

(a) PCA axes 1-2 and axes 1-3 for the sampled northern giraffe populations (1: West African giraffe; 2: Kordofan giraffe; 3: Nubian giraffe; 4: former Rothschild's giraffe). The 95% confidence intervals are shown as oval outlines. Note that the confidence intervals in the PCA axes 1-2, as well as axes 1-3, indicate some substructure within the northern giraffe. However, there is no clear distinction between Kordofan and Nubian giraffe.

(b) Separate STRUCTURE analysis for the northern giraffe populations (WA: West African giraffe; GNP, ZNP, and SNR: Kordofan giraffe; BaNP and ETH: Nubian giraffe; MF: former Rothschild's giraffe) shows additional substructure for up to four clusters, but also highlights admixture at K=3.

Note that detailed information about the populations are listed in Supplementary Table 1.



Supplementary Fig. 7. Additional PCAs and STRUCTURE analyses for Masai giraffe

(a) PCA axes 1-2 and axes 1-3 for the two sampled Masai giraffe populations (1: LVNP; 2: SGR). The 95% confidence intervals are shown as oval outlines. Note that the confidence intervals in the PCA axes 1-2, as well as axes 1-3, indicate some substructure within the Masai giraffe.

(b) Separate STRUCTURE analysis for the Masai giraffe populations shows additional clustering for two populations.

Note that detailed information about the populations are listed in Supplementary Table 1.

Supplementary Table 1. Location, abbreviation, number of individuals (n), (sub)species and source of analyzed giraffe and okapi sequences for mtDNA and nuclear analyses.

Location	Abbreviation	n (mtDNA)	n (nuclear DNA)	(Sub)species
Badingilo National Park, South Sudan	BaNP	2	1	G. c. camelopardalis
Basel Zoo, Switzerland	Okapi	1	1	O. johnstoni
Bwabwata National Park, Namibia	BNP	7	7	G. g. giraffa
Central Kalahari Game Reserve, Botswana	CKGR	7	4	G. g. angolensis
Chobe National Park, Botswana	CNP	11	3	G. g. giraffa
Etosha National Park, Namibia	ENP	17	14	G. g. angolensis
Gambella National Park, Ethiopia	ETH	3	3	G. c. camelopardalis
Garamba National Park, DR Congo	GNP	5	5	G. c. antiquorum
Ishqbini Conservancy, Kenya	ISC	4	3	G. reticulata
Khamab Kalahari Reserve, South Africa	KKR	6	7	G. g. giraffa
Koure, Niger	WA	18	16	G. c. peralta
Loisaba Wildlife Conservancy, Kenya	LWC	1	1	G. reticulata
Luangwa Valley National Park, Zambia	LVNP	11	12	G. tippelskirchi
Moremi Game Reserve, Botswana	MGR	16	5	G. g. giraffa
Mosi-oa-Tunya National Park, Zambia	MTNP	11	3	G. g. giraffa
Murchison Falls National Park, Uganda	MF	16	15	G. c. camelopardalis
Nürnberg Zoo, Germany	RET	5	5	G. reticulata
Nxai Pans, Botswana	NXP	1	-	G. g. giraffa
Rotterdam Zoo, Netherlands	RETRot	3	3	G. reticulata
Selous Game Reserve, Tanzania	SGR	6	7	G. tippelskirchi
Shambe National Park, South Sudan	SNR	2	2	G. c. antiquorum
Sioma Ngwezi National Park, Zambia	SNNP	1	1	G. g. giraffa
Sun hotel, Livingstone, Zambia	SUN	4	4	G. g. giraffa
Vumbura Concession, Botswana	V	11	13	G. g. giraffa
Wilhelma Stuttgart, Germany	RETWI	2	2	G. reticulata
Zakouma National Park, Chad	ZNP	1	1	G. c. antiquorum
Additional mtDNA sequences from GenBank	c			

Individual sample ID's see Figure 2 and Figure S1.

Accession No.	(Sub)species	Source
EF442263 - EF442274	Giraffa spp.	Hassanin, Ropiquet, Gourmand, Chardonnet, & Rigoulet, 2007
EU088317 - EU088320	Giraffa spp.	Brown et al., 2007
EU088322 - EU088351	Giraffa spp.	Brown et al., 2007
AP003424	G. g. angolensis	Yasue et al. unpublished
JN632674	O. johnstoni	Hassanin et al., 2012
NC_012100	G. g. angolensis	Yasue et al. unpublished

Supplementary Table 2. Giraffe and okapi specific primer sequences and PCR conditions for 14 new nuclear intron loci.

Primer sequences and PCR conditions of the remaining seven intron loci and the mtDNA loci can be found in Bock et al. (2014) and Fennessy et al. (2016).

Name & locus	Primer sequence 5'-3'	PCR conditions
Intron SW05 (RFC5, intron 9, chromosome 17)	for: GATCACTCTGGAACCTGCTCA rev: CATACCTGTGGTTCTGCGGT	TD-PCR (Ta = 70-60 °C; 10 cycles), standard PCR (Ta = 60 °C; 30 cycles)
Intron SW07 (USP33, intron 18, chromosome 3)	for: TGACGACCAGAGTGTCACTG rev: TCTTTTTGTGCTTCTTCACTGCT	TD-PCR (T _a = 67-57 °C; 10 cycles), standard PCR (T _a = 57 °C; 30 cycles)
Intron SW21 (MACF1, intron 46, chromosome 3)	for: GCTCACGACCTCATGGAAAT rev: GTTGGAAATGGCTGAGGATG	TD-PCR (T _a = 66-59 °C; 14 cycles), standard PCR (T _a = 59 °C; 26 cycles)
Intron SW40 (IGF2B1, intron 11, chromosome 19)	for: GGCAGCACATCAAACAGCTC rev: GGGGTCCAGTGATGATGACC	See Intron SW05
Intron SW43 (COL5A2, intron 32, chromosome 2)	for: AATGGCTGGAGGACATGGTC rev: GCCGGAAGTTCCTGCAATTC	See Intron SW05
Intron SW44 (CTAGE5, intron 8, chromosome 21)	for: CCCTCAAATCACAAGTAGCTGA rev: TCTGGCTTTCCTGAAGTTGAGA	See Intron SW05
Intron SW51 (NOTCH2, intron 33, chromosome 3)	for: AAAAATGGGGCCAACCGAGA rev: GGGCAGCAAGAAACAGAGGT	See Intron SW05
Intron SW68 (CCT2, intron 14, chromosome 5)	for: TGAAGGCAAAACAACCGCTG rev: CGCTTCACTTGGAAACTTTCTGT	See Intron SW05
Intron SW84 (DHX36, intron 4, chromosome 1)	for: CATCCTTCCCTGACCTCTCAG rev: TCCACAACCAGTTTCACCACT	See Intron SW05
Intron SW108 (C1orf74, intron 1, chromosome 16)	for: TCCAGTGTTGTTGCTGCTGA rev: TCTGGGAGGACCTCGTTTCT	See Intron SW05
Intron SW111 (SAP130, intron 5, chromosome 2)	for: CCCCCTCTTCACATTGGAGC rev: AACTGGACATCACTGCAGCA	See Intron SW05
Intron SW113 (PLCE1, intron 20, chromosome 26)	for: ACTCTGCTTGCAACAAAGGA rev: TCTGGTGCAATCTGTCTGCT	See Intron SW05
Intron SW117 (DDX1, intron 15, chromosome 11)	for: TTCCTGGTCCTGGATGAAGC rev: AACCTTGAGAAAGAAGCCCAT	See Intron SW05
Intron SW123 (USP54, intron 9, chromosome 28)	for: CCCCCAAGTTGAGTTCCAGT rev: CCGTTGAGGAATCGGTTCGA	See Intron SW05

Note – for: forward primer. rev: reverse primer. TD-PCR: touchdown PCR. T_a: primer annealing temperature. The locus is the gene name of the human orthologs, the respective intron, and the chromosome of *Bos taurus* it is located on.

	Northern	Reticulated	Masai	Southern
Northern	-	0.80374	0.18638	0.15276
Reticulated	0.23725**	-	0.18243	0.14427
Masai	0.57289**	0.57813**	-	0.11342
Southern	0.62072**	0.63408**	0.68790**	-

Supplementary Table 3. List of pairwise F_{st} values and resulting N_em for 21 nuclear loci of the four giraffe species.

Note – ** indicates significance of F_{st} values at p < 0.001. N_em values are shown in italics.

Supplementary Table 4. Pairwise F_{st} values for 21 nuclear loci between subspecies / populations within the four giraffe species.

a. Pairwise F _{st} values among southern giraffe subspecies					
	South Afri	can giraffe	Angolan gir	affe	
South African giraffe	-				
Angolan giraffe	0.10199**		_		
b. Pairwise F _{st} values among reticulated giraffe populations					
	RET	ISC	LV	VC	
RET	-				
ISC	0.04000	-			
LWC	0.09065	0.10268	-		
c. Pairwise F _{st} values among northern giraffe subspecies (incl. former Rothschild's giraffe)					
	West African giraffe	Kordofan giraffe	Nubian giraffe	"Rothschild's giraffe"	
West African giraffe	-				
Kordofan giraffe	0.21326**	-			
Nubian giraffe	0.15513**	0.12841**	-		
"Rothschild's giraffe"	0.26777**	0.27702**	0.15245**	-	
d. Pairwise E., values among Masai giraffe populations					

SGR LVNP SGR LVNP 0.39517**

Note – ** indicates significance of F_{st} values at p < 0.001. Detailed information about the populations are listed in Supplementary Table 1.

Supplementary Table 5. Long- and short-term gene flow estimates among the four giraffe species.

a. Mutation-scaled effective population size derived from MIGRATE-N of the four giraffe species

Species	Estimated Θ (95 % conf. int.)
Northern giraffe	0.00137 (0.0 – 0.00287)
Reticulated giraffe	0.00110 (0.0 – 0.00267)
Masai giraffe	0.00003 (0.0 – 0.00153)
Southern giraffe	0.00003 (0.0 – 0.00193)

b. Estimates of gene flow derived from MIGRATE-N and BayesAss among the four giraffe species

Migration route	M (95 % conf. int.) [MIGRATE-N]	N _e m	m (+/- sdev.) [BayesAss]
Reticulated \rightarrow Northern	523.3 (253.3 – 793.3)	0.17923	0.0071 (0.0070)
Masai 🔿 Northern	190.0 (0.0 – 373.3)	0.06508	0.0071 (0.0069)
Southern $ ightarrow$ Northern	123.3 (0.0 – 306.7)	0.04223	0.0071 (0.0069)
Northern $ ightarrow$ Reticulated	3436.7 (3140.0 – 4246.7)	0.94509	0.0209 (0.0196)
Masai $ ightarrow$ Reticulated	390.0 (120.0 – 693.3)	0.10725	0.0208 (0.0195)
Southern $ ightarrow$ Reticulated	376.7 (0.0 – 660.0)	0.10359	0.0208 (0.0196)
Northern $ ightarrow$ Masai	276.7 (20.0 – 553.3)	0.00208	0.0144 (0.0138)
Reticulated $ ightarrow$ Masai	796.7 (366.7 – 1313.3)	0.00598	0.0145 (0.0139)
Southern $ ightarrow$ Masai	503.3 (186.7 – 786.7)	0.00377	0.0144 (0.0138)
Northern $ ightarrow$ Southern	176.7 (0.0 – 360.0)	0.00133	0.0052 (0.0051)
Reticulated $ ightarrow$ Southern	176.7 (0.0 – 366.7)	0.00133	0.0051 (0.0051)
Masai $ ightarrow$ Southern	243.3 (6.7 – 493.3)	0.00182	0.0051 (0.0050)

Supplementary Table 6. Long-term and short-term gene flow estimates between giraffe subspecies (populations) within species.

a. Mutation-scaled effective population size derived from MIGRATE-N for subspecies (populations) within giraffe species

Species	Estimated Θ (95 % conf. int.)
West African giraffe	0.00130 (0.0 – 0.00287)
Kordofan giraffe	0.00150 (0.0 – 0.00307)
Nubian giraffe	0.00110 (0.0 – 0.00273)
South African	0.00137 (0.0 – 0.00313)
Angolan	0.00050 (0.0 – 0.00213)
SGR	0.00143 (0.0 – 0.00300)
LVNP	0.00097 (0.0 – 0.00260)

b. Estimates of gene flow derived from MIGRATE-N and BayesAss among subspecies (populations) within species

Migration route	M (95 % conf. int.) [MIGRATE-N]	N _e m	m (+/- sdev.) [BayesAss]
Kordofan $ ightarrow$ West African	1611.7 (873.3 – 2563.3)	0.52380	0.0211 (0.0168)
Nubian $ ightarrow$ West African	2631.7 (1756.7 – 3520.0)	0.85530	0.0150 (0.0141)
West African $ ightarrow$ Kordofan	3285.0 (2413.3 – 4186.7)	1.23188	0.0636 (0.0364)
Nubian → Kordofan	4215.0 (3523.3 – 4990.0)	1.58063	0.0386 (0.0344)
West African $ ightarrow$ Nubian	2301.7 (1516.7 – 3193.3)	0.63298	0.0168 (0.0158)
Kordofan $ ightarrow$ Nubian	1438.3 (863.3 – 2276.7)	0.39553	0.0347 (0.0228)
South African $ ightarrow$ Angolan	8130.0 (6773.3 – 9420.0)	1.01625	0.0521 (0.0328)
Angolan $ ightarrow$ South African	8456.7 (7913.3 – 9446.7)	2.89642	0.0223 (0.0154)
Masai			
SGR \rightarrow LVNP	1610.0 (940.0 – 2793.3)	0.39043	0.0239 (0.0221)
LVNP \rightarrow SGR	8250.0 (6853.3 – 9746.7)	2.94938	0.0405 (0.0362)

Supplemental references

- Bock, F., Fennessy, J., Bidon, T., Tutchings, A., Marais, A., Deacon, F., & Janke, A. (2014). Mitochondrial sequences reveal a clear separation between Angolan and South African giraffe along a cryptic rift valley. *BMC Evolutionary Biology*, 14(1), 1–12. doi:10.1186/s12862-014-0219-7
- Brown, D. M., Brenneman, R. A., Koepfli, K.-P., Pollinger, J. P., Milá, B., Georgiadis, N. J., ...
 Wayne, R. K. (2007). Extensive population genetic structure in the giraffe. *BMC Biology*, 5(1), 1–13. doi:10.1186/1741-7007-5-57
- Evanno, G., Regnaut, S., & Goudet, J. (2005). Detecting the number of clusters of individuals using the software structure: a simulation study. *Molecular Ecology*, *14*(8), 2611– 2620. doi:10.1111/j.1365-294X.2005.02553.x
- Fennessy, J., Bidon, T., Reuss, F., Kumar, V., Elkan, P., Nilsson, M. A., ... Janke, A. (2016). Multi-locus Analyses Reveal Four Giraffe Species Instead of One. *Current Biology*, 26(18), 2543–2549. doi:10.1016/j.cub.2016.07.036
- Hassanin, A., Delsuc, F., Ropiquet, A., Hammer, C., Jansen Van Vuuren, V., Matthee, C., ...
 Couloux, A. (2012). Pattern and timing of diversification of Cetartiodactyla (Mammalia, Laurasitheria), as revealed by a comprehensive analysis of mitochondrial genomes. *C R Biol*, 335. doi:10.1016/j.crvi.2011.11.002
- Hassanin, A., Ropiquet, A., Gourmand, A. L., Chardonnet, B., & Rigoulet, J. (2007).
 Mitochondrial DNA variability in Giraffa camelopardalis: consequences for taxonomy, phylogeography and conservation of giraffes in West and central Africa. *C R Biol*, 330. doi:10.1016/j.crvi.2007.02.008
- Pritchard, J. K., Stephens, M., & Donnelly, P. (2000). Inference of Population Structure Using Multilocus Genotype Data. *Genetics*, 155(2), 945–959.
- Pritchard, J., Wen, X., & Falush, D. (2010). Documentation for STRUCTURE software, version 2.3. University of Chicago, Chicago, IL.