Supporting Information

Population genetic assessment of study individuals

Our detailed knowledge of the orangutan population structure through previous studies [1-5] on both Borneo and Sumatra, allowed us to assign individuals to their population of origin. We genotyped the animals of the current study at the same 27 highly polymorphic microsatellite makers used in previous studies [6, 7]. We then used the program STRUCTURE v2.3.3 [8] to analyze the study animals for this study together with genotype data from 219 individuals from previous studies [1-5] that represent all major genetic clusters on both islands. For each individual we estimated the membership coefficient Q [8] of belonging to a particular cluster (Figure S1). Details on the analysis and the general observed population structure can be found in Nater [5]. The highest hierarchical level clearly separates Bornean and Sumatran individuals. Analyzing each island separately, three distinct clusters seem to best describe the structure observed in the variation analyzed on Sumatra and five clusters on Borneo. All individuals from this study show high membership to the South Kinabatangan and West Alas cluster, which had also been confirmed by phylogenetic analyses of mitochondrial DNA gene sequences in Nater et. al. [2].

SNP validation

All PCR reactions were performed in a 10 µl reaction volume containing 5 ng of template DNA, 0.1 µM of each primer, 0.2 mM dNTPs, 1 x Phire PCR Buffer and 0.1 units Phire Hot Start II DNA Polymerase (both Finnzymes). PCR conditions were as follows: initial denaturation at 98°C for 30 s, followed by 35 cycles of 98°C for 5 s, 62°C for 10 s, 72°C for 15 s, and final extension at 72°C for 2 min. Direct cycle sequencing was performed with 0.5 µl PCR product in a 10 µl reaction volume containing 1x sequencing buffer (400 mM Tris, 10 mM MgCl₂, pH 9.0), 0.4 µM forward primer and 0.3 µl BigDye Terminator v3.1 (Life Technologies). Cycle sequencing conditions were as follows: initial denaturation at 95°C for 45 s, 30 cycles of 95°C for 30 s, 52°C for 20 s, 60°C for 2 min. Samples were sequenced on a 3730 DNA Analyzer (Life Technologies). We aligned generated sequences with the reference genome sequence ponAbe2 [9] using the SeqMan program of the Lasergene 8 software package (DNASTAR) and visually called genotypes for the target SNP position based on the trace data.
References


Figure S1. STRUCTURE analyses to identify the population origin of study individuals. The analyses were based on 27 microsatellite markers. The membership coefficients $Q$ are average values over ten iterations with the same model parameters (admixture model with burn-in of $3\times10^5$ followed by $3\times10^6$ MCMC steps). Each bar represents a single individual. (A) Bornean and Sumatran individuals separate at the highest hierarchical level. (B) Within Borneo ($n=104$) we observe five clusters (most likely number of clusters as inferred by Arora et al. [7] and Nater [5]). Study individuals ($n=16$) were sampled from the South Kinabatangan cluster. CK: Central Kalimantan ($n=25$), WK: West Kalimantan ($n=20$), EK: East Kalimantan ($n=17$), SK: South Kinabatangan ($n=25$), NK: North Kinabatangan ($n=17$). (C) Within Sumatra ($n=115$) we observe three clusters (most likely number of clusters as inferred by Nater et al. [1] and Nater et al. [5]). Study individuals ($n=15$) were sampled from the West Alas population. NA: North Aceh and Langkat ($n=49$), WA: West Alas ($n=57$), BT: Batang Toru ($n=9$).
Figure S2. Example image of high precision excision of target fragments from Spreadex gel. HaeIII digested genomic DNA was separated on high-resolution Spreadex EL400 Wide Mini S-2x13 gels with M3 size marker. Bands at e.g. 101 bp, 136 bp, and 170 bp represent repetitive elements as predicted by the in-silico HaeIII digest of the orangutan reference genome (see Figure 2). (A) Image of a gel prior to fragment excision. (B) Image of the same gel after excision of DNA fragments in the target size range 104-123 bp.
Figure S3. Overlap of outlier regions among SNP datasets in genome-wide scans for positive selection. For all SNP datasets we performed sliding-window analyses (100 kb window, 25 kb step size) of the absolute allele-frequency differential \( (D) \) between the South Kinabatangan and West Alas population. All windows with an average window \( D > 0.95 \) were considered as outliers, i.e. candidate regions for selective sweeps. (A) Comparison of SAMtools and GATK with their intersect dataset. (B) Comparison of the three intersect datasets.