

**Male demography in East Asia: a north-south contrast in human
population expansion times**

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ABSTRACT

The human population has increased greatly in size in the last 100,000 years, but the initial stimuli to growth, the times when expansion started, and their variation between different parts of the world are poorly understood. We have investigated male demography in East Asia, applying a Bayesian full-likelihood analysis to data from 988 men representing 27 populations from China, Mongolia, Korea and Japan typed with 45 binary and 16 STR markers from the Y chromosome. According to our analysis, the northern populations examined all started to expand in number between 34 (18-68) and 22 (12-39) thousand years ago (KYA), before the Last Glacial Maximum at 21-18 KYA, while the southern populations all started to expand between 18 (6-47) and 12 (1-45) KYA, but then grew faster. We suggest that the northern populations expanded earlier because they could exploit the abundant megafauna of the 'Mammoth Steppe', while the southern populations could only increase in number when a warmer and more stable climate led to more plentiful plant resources such as tubers.

INTRODUCTION

Humans have expanded enormously in geographical range and numbers in the last 100,000 years, starting as a rare species confined to parts of Africa and ending with the current population of more than six billion distributed all over the world, but the details of these changes are poorly understood (JOBLING *et al.* 2004). Historical records document a substantial demographic expansion within historical times, and also the complexity of the changes that have occurred, but are only available for the last few thousand years. Before this time, the archaeological record indicates that humans increased substantially in number when Neolithic transitions led to greater and more reliable food production after ~10 thousand years ago (KYA), but provides only limited quantitative information. Genetic variation can also provide insights into past demography. Standard neutral models of evolution predict the extent and pattern of variation expected in a constant-sized population, but experimental data from the human population are often not consistent with such a model. For example, an overall excess of rare variants, reflected by negative values for Tajima's D (e.g. AKEY *et al.* 2004), is commonly interpreted as a signal of demographic expansion, although the details of such an expansion remain unclear (PTAK and PRZEWORSKI 2002; WALL and PRZEWORSKI 2000). Genomewide analyses of STRs have been interpreted as revealing an early expansion in Africa 49-640 KYA with no expansion outside Africa (REICH and GOLDSTEIN 1998), or alternatively a constant population size in Africa compared with expansions in Europeans and Africans (KIMMEL *et al.* 1998). A larger-scale study using 377 loci in 52 populations suggested expansion in African farmers starting ~35 KYA, Eurasians ~25 KYA and East Asians ~18 KYA, but found no significant signal of growth in African hunter-gatherers or populations from Oceania and America (ZHIVOTOVSKY *et al.* 2003). The conflicting conclusions may reflect, in

part, the complexity of the real events, so that it may not be useful to compare descriptions of demographic change summed over large geographical regions. Instead, studies at higher spatial resolution may be necessary to understand how the demography has changed in different ways at local levels.

Single loci can be influenced by stochastic variation and locus-specific selection, but nevertheless two of them, mitochondrial DNA (mtDNA) and the Y chromosome, are of particular interest because of the insights they can provide into female-specific and male-specific evolutionary patterns, respectively. Mismatch distributions of mtDNA sequences from populations around the world have suggested expansion, on average, ~40 KYA (SHERRY *et al.* 1994), while a phylogenetic star contraction method indicated expansion of the major Asian clades M and N ~30 KYA (FORSTER *et al.* 2001). Studies of the Y chromosome have shown a strong signal of expansion beginning in the Paleolithic ~18 KYA (7-41 KYA, PRITCHARD *et al.* 1999) or ~22 KYA (8.5-50 KYA, MACPHERSON *et al.* 2004) worldwide, with limited variation between continents. In contrast, a detailed study of one country, Armenia, suggested a start of expansion in the Neolithic ~4.8 KYA (2.0-11.1 KYA, WEALE *et al.* 2001).

We want to understand the history of East Asia, including its male demographic history. Modern humans were present in Australia at ~50 KYA and, despite a lack of direct archaeological evidence, may have reached the southern part of East Asia at about the same time (JOBLING *et al.* 2004). Classical marker studies reveal a genetic distinction between northern and southern China, with a boundary corresponding approximately to the Yangtze River (XIAO *et al.* 2000). Some authors have suggested that modern East Asian populations are derived largely from a northwards expansion

of southern populations after the Last Glacial Maximum (LGM) ~18-21 KYA (JIN and SU 2000), while others have suggested a significant male contribution from Central Asia (KARAFET *et al.* 2001). Despite these, and other (DENG *et al.* 2004; SU *et al.* 1999) surveys of Y-chromosomal haplogroup distributions, we know little about the detailed demography of the region, and how it compares between north and south. We now show that male demographic history differs substantially between the northern and southern parts of East Asia, and link this to ecological differences between the regions in the Paleolithic period.

MATERIALS AND METHODS

Dataset: 988 males belonging to 27 populations from China, Mongolia, Korea and Japan were included in this analysis. The samples, and their typing with 16 Y-specific binary and 16 short tandem repeat (STR) markers, have been described previously (XUE *et al.* 2005; ZERJAL *et al.* 2003). For this study, we typed hierarchically an additional 29 binary markers (M89, M8, M38, P33, M217, M93, M48, M61, M76, M147, M27, M214, M5, M128, M178, M119, M101, M50, M175, P31, M95, M88, M122, M121, M134, M164, M159, M113, M7) using multiplexed primer-extension reactions (PARACCHINI *et al.* 2002) adapted for the ABI Prism SNaPshot™ system (HURLES *et al.* 2005) according to the manufacturer's guidelines. As before, *DYS19* was excluded from most analyses because it is duplicated in some individuals.

Data analyses: Haplotypes for this haploid locus could be constructed simply from the combination of STRs and/or binary markers present in the same individual, and their frequencies determined by counting. Analysis of Molecular Variance (AMOVA) was performed using Arlequin 2.0 (SCHNEIDER *et al.* 2000) and Spatial AMOVA

(SAMOVA) analysis using SAMOVA1.0 (DUPANLOUP *et al.* 2002). Spatial autocorrelation was carried out using Autocorrelation Index for DNA Analysis (AIDA) (BERTORELLE and BARBUJANI 1995). Inferences about Y-chromosomal lineage histories and demographies were made using the program BATWING (Bayesian Analysis of Trees With Internal Node Generation) (WILSON *et al.* 2003). Populations (represented by 25 to 65 individuals) were analyzed individually using weakly informative prior distributions for N , the effective population size before expansion (gamma(1,0.0001): mean = 10,000, SD = 10,000), α , the rate of growth per generation (gamma(2,400): mean = 0.005, SD = 0.0035) and β , the time in coalescent units when exponential growth began (gamma(2,1): mean = 2, SD = 1.41) (WILSON *et al.* 2003). A calibrated ‘evolutionary’ mutation rate for Y-STRs (ZHIVOTOVSKY *et al.* 2004) was used as the basis for a per-locus mutation rate prior of gamma(1.47,2130) (mean = 0.00069, SD = 0.00057), and was allowed to vary independently for each locus. This mutation rate was calibrated against two historical events (the divergence of the Maoris and Cook Islanders in the Pacific, and the migration of the Bulgarian Roma from India to Europe), and thus our time estimates are also calibrated against these events, and do not depend on assumptions about generation time. Binary markers (Unique Event Polymorphisms or UEPs) were included under option 2, in which they only condition the tree structures possible. 10^4 samples of the program’s output representing 10^6 MCMC cycles were taken after discarding the first 3×10^3 samples as ‘burn-in’, and convergence was confirmed by examining longer runs of 10^8 MCMC cycles for four populations and finding the same posterior distributions. The influence of population sample size in the range 25-65 was investigated by randomly subsampling 25, 30, 40 or 50 individuals from the Outer Mongolian population with size 65, and found to be negligible. The 0.025, median and 0.975

percentiles of the output samples were recorded. Regression analyses were carried out using SPSS 14; the stepwise criteria in multiple linear regression were the defaults: probability of F to enter ≤ 0.05 , probability of F to remove ≥ 0.10 . A contour plot of expansion times was drawn using SigmaPlot version 9 with inverse square smoothing and a sampling proportion 0.5.

‘Expansion’ always refers to an increase in numbers rather than area, and ‘expansion time’ to the time when the increase *started*, unless otherwise stated.

RESULTS

Approximately one thousand males from 27 East Asian populations were typed with 61 Y-chromosomal markers, and we first describe the basic properties of this dataset. The 45 binary markers identified 31 haplogroups (including paragroups) in the sample, while the 15 STRs defined 730 different haplotypes (Figure 1, Table 1; see also Supplementary Table 1). Population diversities ranged from 0.60 to 0.94 for binary markers and from 0.84 to 1.00 for STRs (Table 2). There was considerable variation in the distribution of lineages between populations, but this did not correspond to the major ethnic distinction in the area, which is between the Han Chinese (>80% of the combined populations of China, Mongolia, Korea and Japan) and the other populations. AMOVA analysis showed that only 1.8% and 0.5% of variation lay between Han and non-Han populations using binary and STR markers, respectively, and neither of these values was significantly greater than zero. There were, however, major geographical differences. Figure 2 shows that, despite the overall predominance of haplogroup O (56%), specific haplogroups were concentrated in each geographical region: C and N in the north, P and J in the west,

O2b in the east and O1*, O2* and O3d in the south. We therefore wished to identify the most important elements of the geographical pattern in an objective way.

We based the subsequent analyses on the STR data unless otherwise indicated because of the problems in interpreting data from pre-ascertained binary markers. SAMOVA analysis (DUPANLOUP *et al.* 2002) identifies, for a pre-specified number of groups of populations, the geographical groups that are most differentiated from one another. Application of this method to the East Asian Y-STR dataset using two or three groups distinguished small numbers of unusual populations, a finding that is readily understood from the high frequencies of the ‘star cluster’ (ZERJAL *et al.* 2003) and ‘Manchu cluster’ (XUE *et al.* 2005) lineages in some northern populations, and reflects extreme expansions of individual patrilineages within historical times. The use of four groups provided the most informative subdivision, with a cluster of six southern populations distinguished in addition to some of the northern ones (Figure 3A). This pattern corresponds well to the north-south distinction seen with classical markers and shows that, in this respect, the Y-chromosomal variation is typical of that on other chromosomes. The division of the sample into more groups led to further subdivisions in the south (e.g. Figure 3B). Spatial autocorrelation analysis (BERTORELLE and BARBUJANI 1995), based on the binary marker variation, produced correlograms that indicated significant clinal patterns or long-distance differentiation (not shown). The north-south haplogroup structure is therefore a continuum rather than a sharp bipartite division. In order to understand it further, we have explored the characteristics of the populations in more detail, concentrating on the 22 non-Han populations because of the spread of the Han during historical times (WEN *et al.* 2004).

A simple property of a population is the variation it contains, and this can be expressed in a number of ways. A widely used measure, diversity, is so high when 15 STRs are used that the differences between populations are small (Table 2) and difficult to interpret. Reducing the number of STRs to an arbitrary four or three (Table 2) produces a wider range of diversity values, and these are noticeably higher in the north than the south. An alternative measure of variation within a population, ASD, shows a similar pattern. BATWING analysis allows demographic parameters of the populations to be explored. Using a model where the population size remains constant for a period and then begins to expand exponentially, we estimated, for each population, posterior values of (1) the effective population size during the constant period, $N_{\text{posterior}}$, (2) the time at which growth began, (3) the rate of growth per generation, α , and (4) the Time to the Most Recent Common Ancestor (TMRCA) of the population (Table 2). We again noticed substantial variation with latitude. Median $N_{\text{posterior}}$ was higher in the north, the expansion began earlier, the rate of growth was slower and the TMRCA was longer. Although all of these variables correlated significantly with latitude when examined individually in regression analyses (Table 3), the highest was with expansion time (adjusted $R^2 = 0.68$), compared with 0.40 for the next-highest, ASD. Unsurprisingly, a stepwise multiple regression analysis identified expansion time as the best predictor of north-south distance, and only α increased this significantly to reach an adjusted R^2 value of 0.75. Thus earlier expansion time in the north and, to a lesser extent, more rapid expansion in the south, account best for the observed north-south differences. We display the expansion times as a contour plot in Figure 4, where the consistent difference between north and south is apparent. Figure 4 suggests, however, that the highest correlation of population expansion may not be with distance due north-south, but with distance along an axis

tilted slightly northwest-southeast, and further examination showed that a tilt of about 10 degrees in fact gave the highest R^2 value (0.71 compared with 0.69).

The demographic model used is simple: it assumes that each population is independent and that a constant phase is followed by exponential growth. The other demographic models available in BATWING, constant population size or continuous expansion, are not informative about the expansion time. In order to explore one consequence of departure from the model used, we investigated artificial population mixtures constructed from combinations of the populations showing the earliest expansion (Inner Mongolians) and those showing the most recent (Yao (Bama) or Li). The artificial population mixtures showed an early expansion time equivalent to that of the Inner Mongolians (Figure 5), demonstrating that the signature of early expansion is not obliterated by admixture.

DISCUSSION

We will consider how our findings on East Asian male variation compare with previous studies, and the implications of our work for the understanding of the demographic history of the region.

The distribution of Y-chromosomal haplogroups in East Asia has been extensively documented (e.g. DENG *et al.* 2004; e.g. JIN and SU 2000; KARAFET *et al.* 2001), but these observations have raised questions about the relationship of northern and southern populations that remain unanswered. Su *et al.* (1999) typed 19 binary markers, 12 of which were chosen because they were already known to be variable in East Asia, and found higher diversity in the south than in the north, and that the

northern lineages were a subset of the southern ones, leading them to suggest that the northern populations were derived from the south by northwards migrations. In contrast, Karafet *et al.* (2001) used a larger set of 52 binary markers ascertained mainly because of their variation in worldwide populations, and discovered higher diversity (mean pairwise differences) in the north, and that the northern lineages were not a subset of the southern ones. They concluded that a contribution to the northern populations from Central Asia was likely. The use of pre-ascertained binary markers introduces a bias into estimates of diversity, but STRs are essentially free of this bias because they are variable in all populations. In our samples, STR diversity and ASD measurements were higher in the north than the south (Table 2), a finding that is not easily reconciled with a largely or exclusively southern origin for the northern populations. It has been suggested that some populations, such as Hui, Uygurs and Mongolians, have recent admixture with Central Asia and so reliance on them may give a false impression (SHI *et al.* 2005), but our findings are common to most populations from the north (Table 2).

Our most striking observation was the demographic contrast between north and south, which was explained largely by the variation in the start of population expansion (Tables 2, 3; Figure 4). Despite the simplified demographic model and wide confidence intervals in the BATWING estimates (Table 2), the median values exhibit a simple and striking pattern: all of the northern estimates lie between 22 and 34 KYA, while all of the southern estimates are between 12 and 18 KYA. These suggest that the northern populations started to expand before the LGM (~18-21 calendar KYA), while the southern populations started to expand after it. These time estimates are calibrated against historical events (ZHIVOTOVSKY *et al.* 2004), and so do not

depend on the assumption of a particular male generation time, but nevertheless are uncertain, and so any interpretation based on them must be regarded with caution. Importantly, however, they are little affected by extensive admixture (Figure 5), and in such a case reflect the earlier expansion time. While extreme northern latitudes were inhospitable to early humans, Siberia has an extensive Upper Paleolithic archaeological record (KUZMIN and ORLOVA 1998) and a highly-productive environment stretched across Asia. This showed an abundance of large animals and has been called the 'Mammoth Steppe' (GUTHRIE 1990). Expansion times calculated in the same way for the Central Asian populations described by Zerjal et al. (2002), excluding those showing recent severe bottlenecks, lay between 24 (13-45) and 36 (16-74) KYA, like the northern populations from East Asia. We therefore propose that this cold but rich environment allowed the demographic expansion of populations who learned to exploit the profuse animal resources, and these people contributed in sufficient numbers to the ancestry of the northern populations we have tested to leave a signature in their paternal lineages. In contrast, this environment did not extend to the southern region, and the populations based there only expanded after the end of the LGM as the climate became warmer and more stable. The large-scale use of underground tubers is thought to have begun in the south as early as 15 KYA (TONG 2004), and it is notable that population expansion was subsequently more rapid there. The survival of this distinct demographic signature provides further evidence for the genetic differentiation between north and south and lack of extensive gene flow, leading to a genetic boundary seen initially in classical marker studies (XIAO *et al.* 2000).

Our conclusions, of course, only refer to the time when expansion *began*, and do not conflict with the notion that population numbers increased much further during Neolithic and historical times. They do, however, illustrate the value of demographic studies at high spatial resolution: a similar analysis of a combined East Asian sample would lead to the conclusion that population growth began at about 30 KYA (in remarkable agreement with the mtDNA estimate, FORSTER *et al.* 2001), and miss an important distinction. Further detailed genetic studies of demography in other parts of the world are now needed.

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LITERATURE CITED

- AKEY, J. M., M. A. EBERLE, M. J. RIEDER, C. S. CARLSON, M. D. SHRIVER *et al.*, 2004 Population history and natural selection shape patterns of genetic variation in 132 genes. *PLoS Biol* **2**: e286.
- BERTORELLE, G., and G. BARBUJANI, 1995 Analysis of DNA diversity by spatial autocorrelation. *Genetics* **140**: 811-819.

- DENG, W., B. SHI, X. HE, Z. ZHANG, J. XU *et al.*, 2004 Evolution and migration history of the Chinese population inferred from Chinese Y-chromosome evidence. *J Hum Genet* **49**: 339-348.
- DUPANLOUP, I., S. SCHNEIDER and L. EXCOFFIER, 2002 A simulated annealing approach to define the genetic structure of populations. *Mol Ecol* **11**: 2571-2581.
- FORSTER, P., A. TORRONI, C. RENFREW and A. RÖHL, 2001 Phylogenetic star contraction applied to Asian and Papuan mtDNA evolution. *Mol Biol Evol* **18**: 1864-1881.
- GUTHRIE, R. D., 1990 *Frozen Fauna of the Mammoth Steppe*. University of Chicago Press, Chicago.
- HURLES, M. E., B. C. SYKES, M. A. JOBLING and P. FORSTER, 2005 The dual origin of the Malagasy in Island Southeast Asia and East Africa: evidence from maternal and paternal lineages. *Am J Hum Genet* **76**: 894-901.
- JIN, L., and B. SU, 2000 Natives or immigrants: modern human origin in East Asia. *Nature Reviews Genetics* **1**: 126-133.
- JOBLING, M. A., M. E. HURLES and C. TYLER-SMITH, 2004 *Human Evolutionary Genetics*. Garland Science, New York and Abingdon.
- KARAFET, T., L. XU, R. DU, W. WANG, S. FENG *et al.*, 2001 Paternal population history of East Asia: sources, patterns, and microevolutionary processes. *Am J Hum Genet* **69**: 615-628.
- KIMMEL, M., R. CHAKRABORTY, J. P. KING, M. BAMSHAD, W. S. WATKINS *et al.*, 1998 Signatures of population expansion in microsatellite repeat data. *Genetics* **148**: 1921-1930.

- KUZMIN, V. Y., and L. A. ORLOVA, 1998 Radiocarbon chronology of the Siberian Paleolithic. *J World Prehist* **12**: 1-53.
- MACPHERSON, J. M., S. RAMACHANDRAN, L. DIAMOND and M. W. FELDMAN, 2004 Demographic estimates from Y chromosome microsatellite polymorphisms: analysis of a worldwide sample. *Human Genomics* **1**: 345-354.
- PARACCHINI, S., B. ARREDI, R. CHALK and C. TYLER-SMITH, 2002 Hierarchical high-throughput SNP genotyping of the human Y chromosome using MALDI-TOF mass spectrometry. *Nucleic Acids Res* **30**: e27.
- PRITCHARD, J. K., M. T. SEIELSTAD, A. PEREZ-LEZAUN and M. W. FELDMAN, 1999 Population growth of human Y chromosomes: a study of Y chromosome microsatellites. *Mol Biol Evol* **16**: 1791-1798.
- PTAK, S. E., and M. PRZEWORSKI, 2002 Evidence for population growth in humans is confounded by fine-scale population structure. *Trends Genet* **18**: 559-563.
- REICH, D. E., and D. B. GOLDSTEIN, 1998 Genetic evidence for a Paleolithic human population expansion in Africa. *Proc Natl Acad Sci U S A* **95**: 8119-8123.
- SCHNEIDER, S., D. ROESSLI and L. EXCOFFIER, 2000 Arelquin: a software for population genetics data analysis, pp. *Genetics and Biometry Lab., Dept. of Anthropology, University of Geneva.*
- SHERRY, S. T., A. R. ROGERS, H. HARPENDING, H. SOODYALL, T. JENKINS *et al.*, 1994 Mismatch distributions of mtDNA reveal recent human population expansions. *Hum Biol* **66**: 761-775.
- SHI, H., Y. L. DONG, B. WEN, C. J. XIAO, P. A. UNDERHILL *et al.*, 2005 Y-chromosome evidence of southern origin of the East Asian-specific haplogroup O3-M122. *Am J Hum Genet* **77**: 408-419.

- SU, B., J. XIAO, P. UNDERHILL, R. DEKA, W. ZHANG *et al.*, 1999 Y-Chromosome evidence for a northward migration of modern humans into Eastern Asia during the last Ice Age. *Am J Hum Genet* **65**: 1718-1724.
- TONG, E., 2004 *Southern Civilization (in Chinese)*. Chongqing Press, Chongqing.
- WALL, J. D., and M. PRZEWORSKI, 2000 When did the human population size start increasing? *Genetics* **155**: 1865-1874.
- WEALE, M. E., L. YEPISKOPOSYAN, R. F. JAGER, N. HOVHANNISYAN, A. KHUDOYAN *et al.*, 2001 Armenian Y chromosome haplotypes reveal strong regional structure within a single ethno-national group. *Hum Genet* **109**: 659-674.
- WEN, B., H. LI, D. LU, X. SONG, F. ZHANG *et al.*, 2004 Genetic evidence supports demic diffusion of Han culture. *Nature* **431**: 302-305.
- WILSON, I. J., M. E. WEALE and D. J. BALDING, 2003 Inferences from DNA data: population histories, evolutionary processes and forensic match probabilities. *Journal of the Royal Statistical Society: Series A (Statistics in Society)* **166**: 155-188.
- XIAO, C. J., L. L. CAVALLI-SFORZA, E. MINCH and R. F. DU, 2000 [Geographic distribution maps of human genes in China (in Chinese)]. *Yi Chuan Xue Bao* **27**: 1-6.
- XUE, Y., T. ZERJAL, W. BAO, S. ZHU, S.-K. LIM *et al.*, 2005 Recent spread of a Y-chromosomal lineage in Northern China and Mongolia. *Am J Hum Genet* **77**: 1112-1116.
- ZERJAL, T., R. S. WELLS, N. YULDASHEVA, R. RUZIBAKIEV and C. TYLER-SMITH, 2002 A genetic landscape reshaped by recent events: Y-chromosomal insights into central Asia. *Am J Hum Genet* **71**: 466-482.

ZERJAL, T., Y. XUE, G. BERTORELLE, R. S. WELLS, W. BAO *et al.*, 2003 The genetic legacy of the Mongols. *Am J Hum Genet* **72**: 717-721.

ZHIVOTOVSKY, L. A., N. A. ROSENBERG and M. W. FELDMAN, 2003 Features of evolution and expansion of modern humans, inferred from genomewide microsatellite markers. *Am J Hum Genet* **72**: 1171-1186.

ZHIVOTOVSKY, L. A., P. A. UNDERHILL, C. CINNIÖGLU, M. KAYSER, B. MORAR *et al.*, 2004 The effective mutation rate at Y chromosome short tandem repeats, with application to human population-divergence time. *Am J Hum Genet* **74**: 50-61.

FIGURE LEGENDS

Figure 1. Phylogeny of Y-chromosomal haplogroups detected in this study.

Figure 2. Geographical distributions of Y-chromosomal haplogroups. A. Populations sampled. B-F. Haplogroup frequencies; circle area is proportional to sample size and sector area to haplogroup frequency. In B-E, haplogroups are sorted into those showing predominantly northern (B), western (C), southern (D) and eastern (E) distributions. F shows the overall frequency of the most common haplogroup, O.

Figure 3. SAMOVA analysis illustrating the geographical divisions identified when four (A) or six (B) groups are specified.

Figure 4. Contour plot showing the distribution of expansion times. Demographic expansion began earlier in the north than the south.

Figure 5. Effect of artificial mixing of population data on estimated expansion time. Median values are plotted, together with their 95% confidence intervals.

Table 1. Haplogroup frequencies in East Asian populations

Haplogroup	Daur	Ewenki	Hezhe	Hui	Manchu	Inner Mongolian	Oroqen	Uyghur (Urumqi)	Uyghur (Yili)	Xibe	Han (Harbin)	Han (Yili)	Korean (China)	Buyi	Hani	Li	Qiang	She	Tibetans	Yao (Bama)	Yao (Liannan)	Han (Chengdu)	Han (Lanzhou)	Han (Meixian)	Japanese	Korean (Korea)	Outer Mongolian	TOTAL	
Y*(xA,CE,JR)	2			2		2		1	8	1		1	2		1												2	25	
C*(xC1,C3)	0	1	3	4				1	1											6									16
C1																									4			4	
C3*(xC3c)	11	7	5	4	8	17	6			9	5	2	3		6			1			1	4	6	3	1	7	22	128	
C3c	1	7	5		1	4	13		3	2																	13	49	
D				4	1				1	1		1					6		15	1			2		13	1	1	47	
E				1				2																				3	
J				3				8		3	1												3			1	2	21	
K*		1			1	2	1		5	2		3	1											1		1	1	19	
NO*/-	1	1	1	1										2						1					1	1		9	
N*					2			1	1	4	1		1		4				3			1	2	1	2	1		24	
N2			8		1		2	2																			2	15	
N1					2					1				2														5	
N3*(xN3a)																											1	1	
N3a	3					6			2	2	1																4	18	
O*	1						1									2							1					5	
O1*(xO1a,O1b)	2	2		3	1			1		3	1	3		1		9	5	5		1	2	5	2	7				53	

O2*	1		1	1	3	1	2				3	3	1		1	1		13		1		5		1	1	39		
O2a*	6	1										1		13	2	20	3	3	1	2	1	1		2		56		
O2a1														6	15	1	1			1	4					28		
O2b*	1		2		2				1			5											2	6		19		
O2b1												2											11	6		19		
O3*(xO3a-O3e)	1		7	1	3	3		2	2	7	7	6		1		2	2			1	1	3	1	1	7	2	60	
O3/-cd*	6	1	4	3	3	2	2		2	3	3		1			1	5			3	7	3	9			1	59	
O3/-c																							1			1		
O3/-d							1					1		6			10		12	18			2			50		
O3/-d1							1					1														4		
O3e*(xO3e1)		1	2	1	2	3	1	1	2	5	8	1		3	3	1	4	2	1		2	5	2	1	2	5	1	59
O3e1*(xO3e1a)	3	4	7	1	5	5	1	1	2	2	5	2	4	1	2		3	6	13		5	5	3	4	8	5	3	100
P*(xR1a)				4				6	6	1		1				2									1	3	24	
R1a				2				7	6			2						3				2				6	28	
TOTAL	39	26	45	35	35	45	31	31	39	41	35	32	25	35	34	34	33	34	35	35	35	34	30	35	47	43	65	988

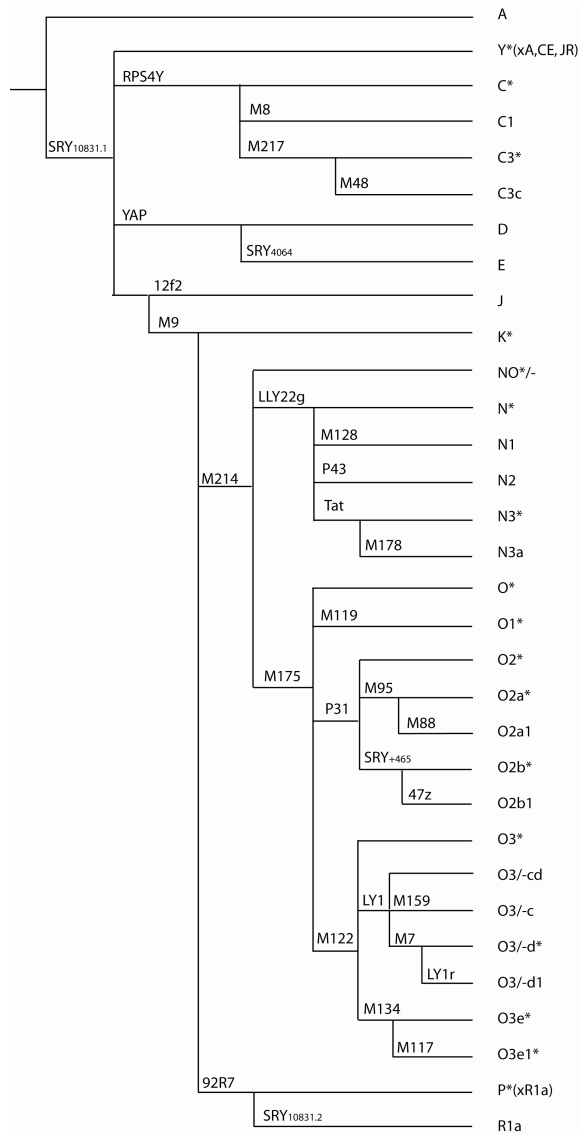
Table 2. Population statistics: observed variation and BATWING prior and posterior estimates

Population	Location		Y-SNP	Y-STR variation				BATWING output											
	Long	Lat	diversity	Diversity			ASD	Nposterior			Expansion time (KYA)			α posterior			TMRCA (KYA)		
	deg	deg		15	4	3		2.5%	Median	97.5%	2.5%	Median	97.5%	2.5%	Median	97.5%	2.5%	Median	97.5%
	E	N	STRs	STRs	STRs	2.5%	Median	97.5%	2.5%	Median	97.5%	2.5%	Median	97.5%	2.5%	Median	97.5%		
BATWING prior								254	6940	36900	1.5	291.5	5138.3	0.0006	0.0042	0.0140			
Buyi	105.7	26.0	0.810	0.988	0.958	0.866	64	203	585	1666	6.8	15.5	36.1	0.0011	0.0036	0.0090	18.7	34.6	70.4
Hani	101.9	23.5	0.767	0.995	0.865	0.825	75	168	453	1413	8.6	17.2	34.7	0.0018	0.0043	0.0098	18.1	31.6	61.8
Li	109.3	18.5	0.596	0.838	0.677	0.665	44	163	491	1462	4.6	12.4	35.5	0.0006	0.0036	0.0105	17.0	32.1	78.6
She	117.4	25.1	0.849	0.966	0.914	0.890	77	254	638	1773	6.0	17.6	46.6	0.0005	0.0024	0.0063	18.8	33.4	66.2
Tibet	91.1	29.6	0.682	0.990	0.956	0.936	145	298	868	2146	5.6	17.1	50.8	0.0007	0.0028	0.0103	24.7	44.4	85.2
Yao (Bama)	107.2	23.4	0.733	0.879	0.778	0.738	63	210	483	1072	1.1	11.9	44.6	0.0001	0.0011	0.0062	15.9	29.5	57.2
Yao (Liannan)	112.2	24.7	0.716	0.992	0.891	0.778	66	161	473	1571	5.7	15.6	36.4	0.0011	0.0039	0.0114	21.3	41.0	90.5
Daur	124.6	49.3	0.875	0.972	0.919	0.908	123	286	686	1791	13.4	26.4	59.4	0.0006	0.0019	0.0042	24.0	40.8	78.4
Ewenki	118.6	48.0	0.841	0.923	0.895	0.874	95	273	827	2127	4.2	22.7	57.7	0.0004	0.0017	0.0053	21.8	38.1	70.3
Hezhe	132.5	47.5	0.898	0.975	0.952	0.937	117	500	1181	2842	8.2	27.0	71.0	0.0003	0.0013	0.0034	25.4	43.7	80.2
Hui	105.5	36.6	0.941	0.985	0.958	0.881	115	197	452	1182	14.9	25.8	43.9	0.0017	0.0033	0.0063	20.8	34.5	58.1
Inner Mongolian	126.0	51.0	0.822	0.972	0.910	0.905	138	333	903	2029	17.7	34.3	68.3	0.0007	0.0017	0.0038	30.4	52.3	92.7
Japanese	134.0	34.2	0.842	0.997	0.938	0.909	117	224	632	1655	12.3	21.7	39.3	0.0019	0.0039	0.0074	24.7	40.2	69.4
Korean (China)	121.4	42.0	0.878	0.993	0.953	0.923	130	289	721	2657	8.8	26.5	56.8	0.0008	0.0024	0.0066	27.4	46.1	88.8

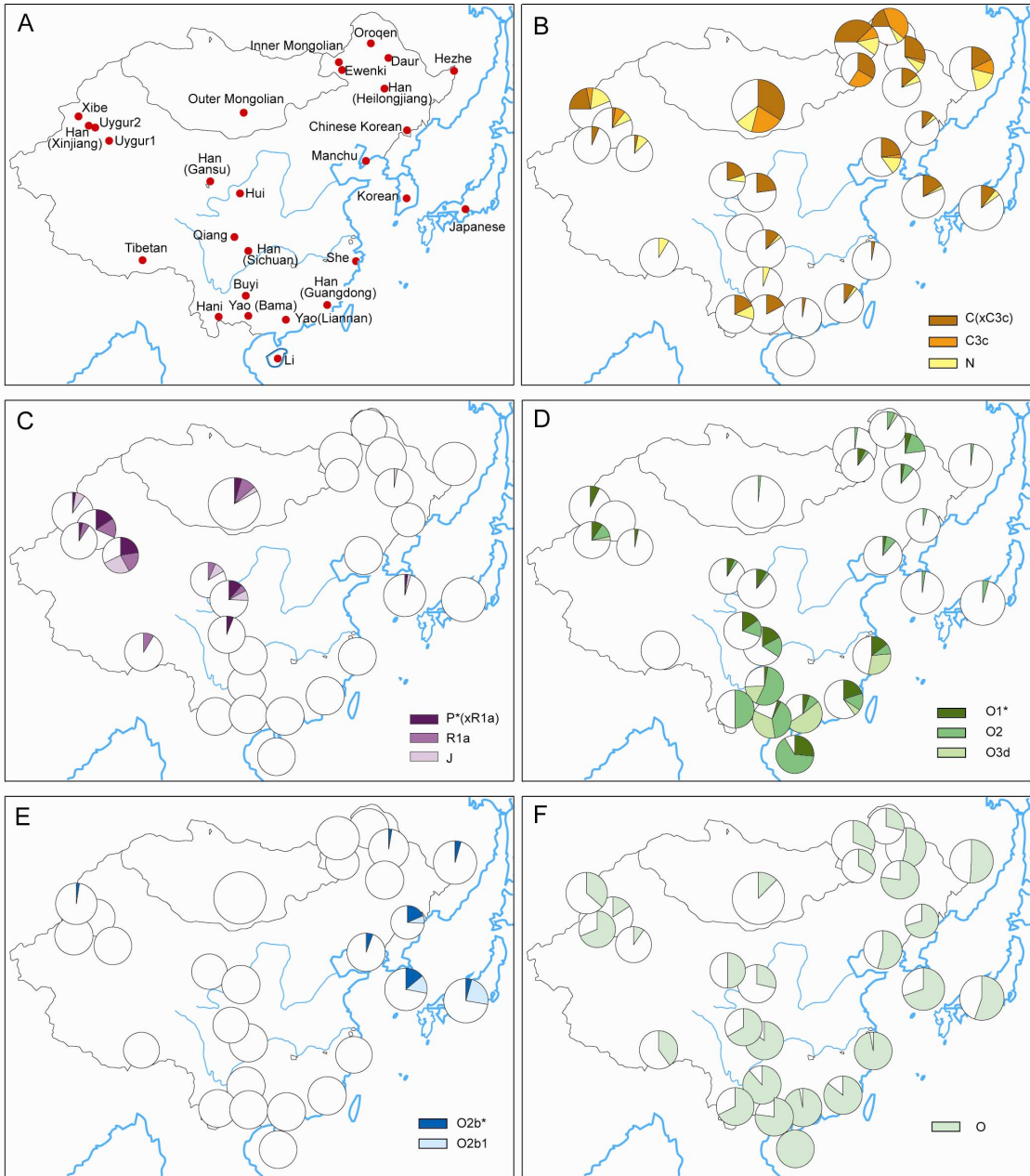
Korean (Korea)	127.0	37.5	0.898	0.991	0.956	0.932	123	240	602	1773	16.4	30.3	58.5	0.0010	0.0024	0.0048	28.5	48.3	86.6
Manchu	123.2	40.2	0.914	1.000	0.978	0.963	149	311	790	2361	11.8	22.5	45.1	0.0020	0.0046	0.0098	29.0	49.4	91.6
Oroqen	123.4	50.3	0.794	0.933	0.886	0.873	92	228	600	1600	8.9	22.7	53.4	0.0006	0.0022	0.0060	23.2	41.6	83.2
Outer Mongolian	107.0	48.2	0.837	0.987	0.929	0.904	114	316	617	1409	13.9	26.3	54.6	0.0009	0.0022	0.0045	23.5	39.4	74.4
Qiang	103.5	31.4	0.919	0.998	0.975	0.930	115	252	681	1632	12.0	22.9	44.6	0.0015	0.0035	0.0073	26.3	44.9	79.7
Uygur (Urumqi)	87.6	43.8	0.899	1.000	0.985	0.976	174	344	1336	3673	12.7	30.6	64.1	0.0009	0.0025	0.0068	38.1	65.0	117.2
Uygur (Yili)	87.3	43.5	0.858	0.997	0.976	0.961	142	578	970	1911	18.0	28.6	48.1	0.0018	0.0034	0.0060	30.8	48.3	81.6
Xibe	81.7	44.0	0.922	0.995	0.970	0.955	134	282	735	2432	16.0	30.0	55.6	0.0012	0.0027	0.0051	28.4	49.5	92.2

Table 3. Regression analysis

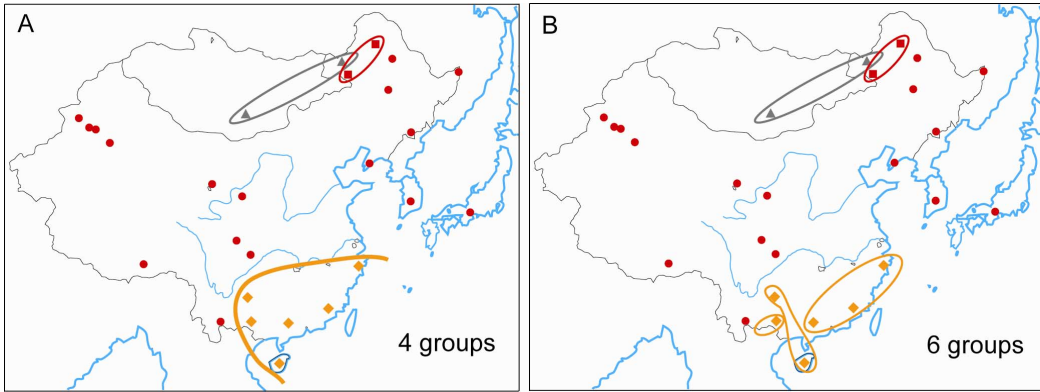
Simple regression (dependant variable: latitude)			
Independent variable	R ²	Adjusted R ²	<i>P</i> value
3 STRs	0.41	0.38	0.001
ASD	0.43	0.40	0.001
N _{posterior}	0.30	0.27	0.008
Expansion time	0.69	0.68	0.000
α	0.24	0.20	0.022
TMRCA	0.33	0.29	0.005
Stepwise multiple regression			
Model 1. Expansion time	0.69	0.68	0.000
Model 2. Expansion time + α	0.87	0.75	0.000



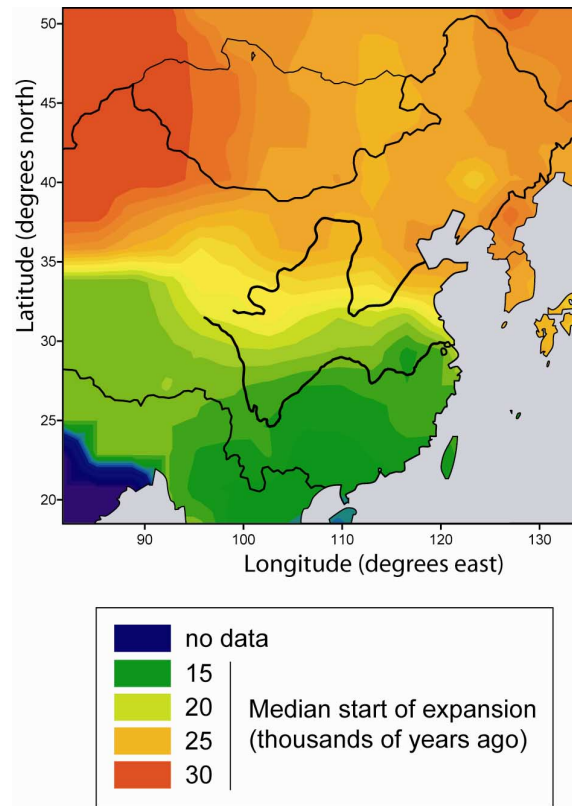
Xue *et al.* Figure 1



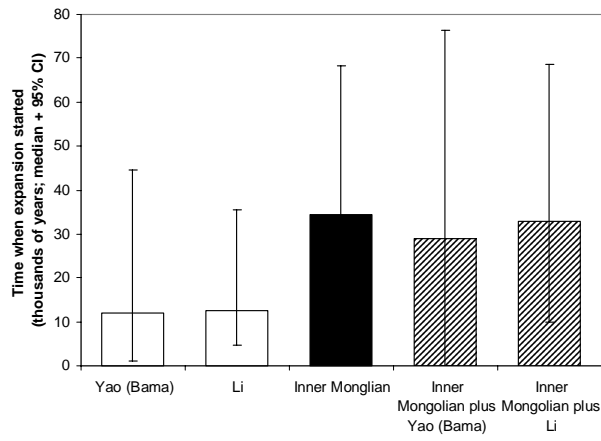
Xue *et al.* Figure 2



Xue *et al.* Figure 3



Xue *et al.* Figure 4



Xue *et al.* Figure 5