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X-chromosome as a marker for population history: linkage disequilibrium and haplotype study in Eurasian populations

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Linkage disequilibrium (LD) structure is still unpredictable because the interplay of regional recombination rate and demographic history is poorly understood. We have compared the distribution of LD across two genomic regions differing in crossing-over activity – Xg13 (0.166 cM/Mb) and Xp22 (1.3 cM/Mb) – in 15 Eurasian populations. Demographic events predicted to increase the LD level – genetic drift, bottleneck and admixture – had a very strong impact on extent and patterns of regional LD across Xq13 compared to Xp22. The haplotype distribution of the DXS1225–DXS8082 microsatellites from Xq13 exhibiting strong association in all populations was remarkably influenced by population history. European populations shared one common haplotype with a frequency of 25–40%. The Volga-Ural populations studied, living at the geographic borderline of Europe, showed elevated LD as well as harboring a significant fraction of haplotypes originating from East Asia, thus reflecting their past migrations and admixture. In the young Kuusamo isolate from Finland, a bottleneck has led to allelic associations between loci and shifted the haplotype distribution, but has much less affected single microsatellite allele frequencies compared to the main Finnish population. The data show that the footprint of a demographic event is longer preserved in haplotype distribution within a region of low crossing-over rate, than in the information content of a single marker, or between actively recombining markers. As the knowledge of LD patterns is often chosen to assist association mapping of common disease, our conclusions emphasize the importance of understanding the history, structure and variation of a study population. European Journal of Human Genetics (2005) 13, 452–462. doi:10.1038/sj.ejhg.5201340 Published online 22 December 2004

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Introduction

More than 10 years ago linkage disequilibrium (LD), or allelic association mapping, was pioneered as a tool for fine-scale localization of genes responsible for rare monogenetic diseases,^{1,2} but has now come center-stage as the method of choice for finding the genes behind common diseases.^{3,4} Both simulated as well as empirical data have shown that population history, in terms of bottlenecks, genetic drift in small populations, and admixture, has an impact on population background LD level.⁵ The data gathered from across different genomic regions also suggest the unique locus history of every genomic segment is influenced by local mutation, recombination rates and selection shaping the regional LD patterns.^{5,6} Against the

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background of the recently initiated International Hap-Map project to create a genome-wide map of LD patterns in the human genome, there is still much debate as to whether this map would be applicable to association mapping in a population of interest or to the provision of detailed fine-scale structure for regions of interest.^{7,8} More empirical data on worldwide populations are needed to understand how population history in combination with regional crossover activity acts on local LD patterns.

The X chromosome is a useful marker for population genetic studies owing to its intrinsic properties: accessible haplotypes in males, lower recombination rate, lower mutation rate and faster genetic drift due to smaller effective population size.⁹ As a result, we can expect LD to be greater on the X chromosome, and population structure more pronounced, compared to autosomes. In order to study the interplay between population history and recombination rate shaping the local LD patterns, we have chosen two X-chromosomal regions with contrasting crossing-over activity. Xq13 can be defined as an LD desert (0.166 cM/Mb), whereas Xp22 represents a recombination rate close to the average in human genome (1.3 cM/Mb). We have compared allelic associations for an extensive data set of 14 Eurasian populations and a further isolated subpopulation (including nine novel and six previously published populations). The populations have been chosen to represent different demographic scenarios predicted to generate LD: (i) the Saami and the Evenki: small constant populations, whose genepool has been influenced by genetic drift;^{10,11} (ii) Kuusamo: a young, 300-years-old regional subpopulation of Finland, which has experienced multiple bottlenecks and is geographical isolated;^{12,13} (iii) Volga-Ural populations of Mari, Udmurt, Chuvash and Komi: complex population history and ethnic structure due to geographic location on the borders of Europe and Asia.¹⁴ All the populations of the Volga-Ural region are distributed in smaller subgroups, often speaking distinct dialects, and having a wide geographic range even today. Based on mtDNA analysis the major part of the genepool of these populations is European.^{14,15} The history of these populations, however, is rich in active contact with neighbors of different genetic background (East Asians, Turkic-speaking Bolgars, Tatars as well as Sub-Arctic groups).^{14,16} Specifically, the Chuvash population is known from history to have arisen from the descendents of Turkicspeaking Volga Bolgars and local Fenno-Ugric tribes (apparently the Mari) after the fall of the Bolgar Kingdom due Mongol-Tatar invasion in around 1230. The Komi tribe is known to have moved to its current territories rich in taiga and tundra only after 900 AD. The founder population mixed with local Nenents tribes, as well as adopting their economy based on reindeer-breeding, hunting and fishing. The Mari have two subpopulations (eastern or meadow, and western or mountain Mari), speaking distinct dialects, and even with distinct written languages. For comparison we studied LD patterns for outbred and expanded populations of different size from eastern (Russians, Mordvin), northern (Finns, Estonians and Swedes), western (Dutch, Germans) and southern (Italians) Europe.

Materials and methods Population samples

The sample sets of North-European DNA representing Finnish (n=80), Swedish (n=41), Estonian (n=45), the Saami, and the Evenki (n = 71) populations are reported elsewhere.^{10,11} The collection of the Volga-Ural population blood samples – Mari (n = 44), Komi (n = 46), Udmurt (n = 49) and Chuvash (n = 40) as well as Mordvin (n = 49) – was conducted with help of local Blood Centers of Volga-Ural region and is described in detail by Bermisheva et al.¹⁵ The Russian DNA samples (n = 66), collected in the framework of the International program INTAS Grant No. 93-0035, were kindly provided by Professor Voevoda (Institute of Internal Medicine, Novosibirsk, Russia). The German blood samples (n=41) represent the German population from the county of Munich (South-Bavaria) and were provided by Dr Weichhold (Institut für Rechtsmedicine, München). The Dutch blood samples (n = 70)originating from the rural county of Doetinchem (Mid-East of Holland) were shared by Dr de Knijff. The Italian DNA samples (n=92) were kindly provided by Dr Cristian Capelli (The Forensic Genetics Laboratory, Istituto di Medicina Legale, Università Cattolica di Roma) and originate from the populations of Rome and Genova. The Kuusamo represents a recent isolate of northeastern Finland, founded at the end of the 17th century by a nucleus of 39 families and now comprising 18000 inhabitants that remained isolated until World War II.¹² In this study the Kuusamo sample data were used as a model for recent bottleneck and rapid expansion. The East Asian Xq13 data set of Japanese and Mongolian populations described by Katoh *et al*¹⁷ was used as a reference source for east-Asian specific haplotypes. The Khalkh (1.8 million) represent the largest Mongolian population, whereas Khoton, Zahkchin and Uriankhai are young, isolated subpopulations (<25000 people).

Laboratory procedures

All samples were genotyped for eight dinucleotide microsatellites on Xq13 and six markers on Xp22 selected from the Genethon genetic map as previously described.^{10,11} All the genotypes were determined manually. The exact location (bp) on Human Genome Map of the microsatellites used is reported by Kaessmann *et al.*¹¹ The order and distances in Mb/cM (deCode map) between the markers are the following:

- (1) Xq13: DXS983 4.038/2.52 DXS8037 0.731/<0.2 -DXS8092 - 7.183/<0.2 - DXS1225 - 0.162/<0.19 -DXS8082 - 1.053/<0.19 - DXS986 - 3.359/<1.83 -DXS1066 - 4.442/<1.83 - DXS995;
- (2) Xp22: DXS987 1.627/<3.72 DXS1053 3.723/
 <3.72 DXS7163 0.814/<3.72 DXS999 1.142/2.9
 DXS1229 1.901/5.39 DXS989.

Data analysis

Arlequin version 2.0 software¹⁸ was used to estimate allele and haplotype frequencies, as well as calculate locus diversity (*d*) for each marker across both studied regions: $d = \frac{n}{n-1}(1 - \sum p_i^2)$, where p_i is the estimated frequency of the *i*th allele^{*i*} at the locus. For each microsatellite locus in studied samples, the population mutation parameter $\theta = 4N\mu$ (*N* is the effective population size; and μ the mutation rate) was estimated by maximum likelihood¹⁹ using MISAT software (http://www.bscb.cornell.edu/Homepages/Rasmus_Nielsen/files.html).

Using GOLD software,²⁰ we calculated multiallelic extension of the normalized association measure D' as

$$D_{m}^{'} = \sum_{i=1}^{k} \sum_{j=1}^{1} p_{i} q_{j} |D_{ij}^{'}|,$$

where *p* and *q* are observed allele frequencies at the two loci. As *D'* is sensitive to rare allele frequencies, alleles with frequencies <10% were pooled. First, to address the level of background LD between unlinked markers, 'baseline' distribution of *D'* values in each population sample was calculated for 48 possible pairs of unlinked microsatellites from Xq13 and Xp22. Second, we asked the question whether LD across studied regions is significantly different from the background LD between unlinked markers for each studied sample set. Mann–Whitney *U*-test was used to compare the distribution of *D'* values between unlinked markers with the distribution of *D'* values for marker-pairs across Xq13 or Xp22. Marker-pairs were grouped for the analysis according to distance separating them: (i) 0.1–2 Mb; (ii) 3–5 Mb; (iii) 5–10 Mb or (iv) 10–20 Mb.

Third, patterns of D' values were used to compare LD levels across Xq13 among populations. However, as D' estimates are strongly dependent on sample size, ^{21,22} we aimed to calculate D' values for Xq13 marker-pairs for equally sized sample sets (n=40) from each population. Equal sample sizes were obtained by sampling of 40 random individuals from the original data sets of >40 individuals. Small sample size for Kuusamo (n=39) and Chuvash (n=40) did not allow resampling procedure. Sampling was repeated 100 times and D' estimates were calculated as the mean of all replicates. Resampling in the present context was not meant as a traditional boot-strapping test to assess sample composition, but rather as

an approach to achieve equal number of representative individuals from each studied population.

Significance of the allelic association between all possible locus pairs was also estimated by the tail probability (*P*-value) of Fisher's exact test, computed by the Genepop 3.0 software.²³ For each pair of loci, $r \times c$ contingency table of gametes was formed and 1000 tables with the same marginal totals were generated based on a Markov chain algorithm. The procedure was repeated 500 times. The *P*-value is the mean fraction of such tables, which were equally or less likely than the observed table.

In order to compare the extent of overall LD across the studied regions, a multilocus LD statistic $r_{\rm d}$ was computed for 10 Mb regions of Xp22 (all markers) and Xq13 (from DXS8037 to DXS986) using Multilocus 1.2 software (Paul-Michael Agapow and Austin Burt, http://www.bio.ic.ac.uk/ evolve/software/multilocus). In essence, one is asking whether two individuals being the same at one locus makes them more likely to be the same at another. R_d is an extended statistic from traditional multilocus LD index of association, *I*_A,²⁴ correcting for the number of loci used in the analysis and thus making the comparison between different genomic regions possible. IA is based on comparing the variance of calculated pairwise differences between the haplotypes in the sample to the expected variance under the assumption of linkage equilibrium between the loci: $I_{\rm A} = V_{\rm D}/V_{\rm E} - 1$. To remove the dependency on number of loci, modified statistic r_d is used, where var (j, k) are the variance of single loci *j* and *k*, respectively:

$$r_d = (V_{\rm D} - \sum {\rm var}_j)/2 \sum \sum \sqrt{{\rm var}_j} \, {\rm var}_k$$

Exact test for locus differentiation between all pairs of populations was computed by Genepop 3.0 software.²² The threshold of significant differentiation was determined <0.01, more stringent that traditional <0.05 due to relatively small sample size.

Analysis of population structure and assignment of individuals into inferred population clusters was carried out using STRUCTURE version 2.0 software²⁵ (http:// pritch.bsd.uchicago.edu). We analyzed three alternative data sets: The data set A $(n_{\text{samples}} = 1241)$ consisted of seven Xq13 loci (DXS1066 excluded) for 21 population across from Europe (10), Volga Ural (4) and East-Asia (7). The reference populations of Asian origin included Evenki and Buriats as well as published genotypes from Japanese and four Mongolian populations.17 The data set B $(n_{\text{samples}} = 889)$ included eight Xq13 and six Xp22 microsatellites for 10 European, four Volga-Ural and two Siberian populations. The data set C used the same samples as B, but for analyzing only six markers of Xp22. Structure analysis was conducted under linkage model,²⁶ an extension to the original method for inteferring population structure from multilocus data,²⁵ but allowing for linkage between loci. Analysis was conducted with the following parameters: no prior population information, 30000 burn-in period and

1 000 000 run length. Multiple runs of each data set guaranteed the robustness of the analysis. The number of population clusters was estimated as the value of K that maximized estimated model log-likelihood, log(P(X|K).

Results

Locus diversities of Xq13 and Xp22 microsatellites

Microsatellites of both studied regions were characterized by locus diversity of similar magnitude, 0.62-0.72 (±s.d. 0.34-0.39) for Xq13 and 0.62-0.73 (±0.35-0.41) for Xp22 (Table 1). Consistent with population genetics theory, reduction in mean number of alleles per locus was found for the Saami and Evenki (genetic drift in small constant populations), as well as Kuusamo (extreme bottleneck) sample. For these populations the reduction in diversity level correlated with lower estimates of population genetics parameter θ : 3.84-5.60 averaged across Xq13 microsatellites and 3.74-5.81 for Xp22 loci compared to 8.42-11.15(Xq13) and 6.55-12.17 (Xp22) for other populations. In most of the populations, except for the Saami, Evenki, Kuusamo and Mari, every individual carried a unique haplotype constructed of 6 (Xp22) or 8 (Xq13) studied loci.

Background LD between unlinked markers varies among population samples

The 'baseline' LD for each population sample was estimated by computing the D'_{BASE} values for all possible pairs (n = 48) of unlinked microsatellite loci formed between the Xq13 and Xp22 markers. The mean 'baseline' D'_{BASE} varied two times across populations, ranging from $0.166\pm$ s.d. 0.052 for Italians to $0.331\pm$ s.d. 0.093 in Komi (Figure 1). The median D'_{BASE} follows tightly the mean values indicating the relatively even distribution of the D'_{BASE}

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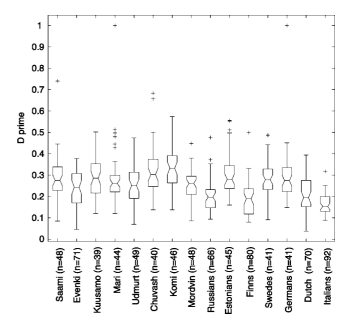


Figure 1 Notched boxplot for the distribution of D' values between 48 unlinked locus pairs (D'_{BASE}) formed between Xq13 and Xp22 microsatellite markers in studied populations. The boxes represent the 25th and 75th percentiles. The median D' is denoted as the line that bisects the boxes. Notches represent a robust estimate of the uncertainty about the medians for box to box comparison. The whiskers are lines extending from each end of the box covering the extent of the data on 15 X interquartile range. Crosses represent the outlier D'_{BASE} values.

Table 1 Locus diversity in 15 studied populations based on eight markers on Xq13 and six markers on Xp22

			Xq13:8 mici	rosatellites		Xp22: 6 microso	ntellites	
Population	Census size ^a	Sample size Xq13/Xp22	Locus diversity Mean±SD	No. of alleles	No. of haplotypes	Locus diversity Mean±SD	No of alleles	No. of haplotypes
Saami ^b	<80000	54	0.62 ± 0.34	5.57±2.28	33	0.65 ± 0.37	5.50±2.26	43
Evenki ^b	50 000	71/70	0.63 ± 0.35	5.86 ± 3.11	56	0.62 ± 0.35	5.33 ± 2.94	53
Kuusamo ^c	~ 18000	39	0.64 ± 0.35	5.75 ± 1.70	34	0.71 ± 0.40	7.33 ± 3.50	35
Mari	~ 670000	44	0.69 ± 0.38	7.21 ± 3.07	37	0.71 ± 0.39	7.33 ± 3.01	43
Udmurts	\sim 746 000	49/42	0.70 ± 0.38	7.57 ± 3.48	48	0.72 ± 0.40	7.83 ± 4.49	42
Chuvash	\sim 1.8 million	40/38	0.71 ± 0.38	7.50 ± 3.18	38	0.73 ± 0.41	7.67 ± 4.08	37
Komi	\sim 350 000	46	0.72 ± 0.39	7.79 ± 3.12	45	0.71 ± 0.40	8.67 ± 3.78	46
Mordvin	\sim 1.2 million	48/49	0.71 ± 0.38	7.71 ± 3.12	47	0.70±0.39	8.50 ± 3.78	49
Russians	200 million	66	0.64 ± 0.35	8.29±3.29	66	0.70 ± 0.39	8.67±4.27	64
Estonians ^b	~ 900000	45/43	0.65 ± 0.36	7.00 ± 2.86	45	0.72 ± 0.41	7.83 ± 3.43	43
Finns ^b	\sim 5 million	80/79	0.69 ± 0.37	8.50 ± 3.11	77	0.67 ± 0.37	9.00 ± 3.79	77
Swedes ^b	9 million	41	0.67 ± 0.37	7.36 ± 2.41	41	0.72 ± 0.40	7.67 ± 2.80	41
Germans	80 million	41	0.66 ± 0.36	7.07 ± 3.10	41	0.72 ± 0.40	8.33 ± 3.98	41
Dutch	15 million	70/69	0.69 ± 0.37	8.07 ± 2.64	68	0.70 ± 0.40	8.83 ± 3.60	67
Italians	45 million	92/93	0.69 ± 0.37	8.71 ± 3.71	91	0.75 ± 0.41	10.33 ± 4.93	93

^aBertelsmann Lexikon Die Völker der Erde (1996); Laakso J Uralic populations (1991).

^bData for Xq13 from Laan and Pääbo (1997) and Kaessmann *et al* (2002).

^cData from Varilo *et al* (2000).

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by chance. The mean D'_{BASE} was found to be correlated with neither (I) population size - small vs large populations, Kuusamo mean $D'_{\text{BASE}} = 0.289 \pm \text{s.d.}$ 0.099 and the Saami 0.291±0.106 vs Swedes 0.286±0.079; (II) demographic history - constant vs expanded populations, the Saami and Evenki $D'_{\text{BASE}} = 0.291 \pm 0.106$ and 0.234 ± 0.086 vs Estonians and Russians $0.305\pm$ s.d. 0.098 and 0.203 ± 0.078); nor (III) sampling – from one county vs across population, Dutch $D'_{\text{BASE}} = 0.209 \pm 0.079$ vs Finns 0.191 ± 0.082 . Consistently with previous reports,^{21,22} we found negative correlation between the mean D'_{BASE} and sample size (Corr. Coef. = -0.911). Our *D'* values computed for unlinked microsatellite loci on X chromosome are higher than usually obtained for unlinked SNPs, where $D'_{\text{BASE}} < 0.2^{5,13}$ The higher baseline D' values could either result from different marker properties (SNPs vs microsatellites) and/or distinct LD patterns on X-chromosome due to smaller effective population size as well as two times reduced recombination events compared to autosomes. This indicates the importance of estimating the baseline LD for each population sample and marker set used in any particular study aiming to study LD patterns.

Different patterns of LD across Xp22 and Xq13

Allelic association between microsatellite loci across Xp22 and Xq13 in each population was studied by three statistics: (a) multiallelic extension of Lewontin's metric D', (b) Fisher's exact test for the significant departure from linkage equilibrium, (c) multilocus association parameter of r_d . Xp22 stands out as relatively LD – poor region, most of the populations exhibit 0–2 significant (0.01 < P < 0.05

from Fisher's exact test) associations for the 15 studied marker-pairs (data not shown). LD across Xp22 exceeds significantly X-chromosomal D'_{BASE} only between closely linked loci for the Saami, Mari and Udmurt, and for markers further apart in Kuusamo and Udmurt sample (Table 2). In the case of the Udmurt, apparently a recent mutation in DXS987 is responsible for creating LD as this marker was involved in three of four associations across Xp22. Consistently, the multilocus LD parameter r_d values (<0.05, except the Saami) across 10 Mb Xp22 region refer minimal association of markers (Figure 2).

In contrast to the LD-poor Xp22 region, Xq13 shows a more diverse picture of the regional LD structure among populations both for overall association parameter $r_{\rm d}$ (Figure 2) as well as for pairwise LD patterns (Figure 3). For all studied populations, Xq13 exhibited stronger LD compared to Xp22 (Figure 2, Table 2). The Saami has the strongest multilocus LD ($r_d = 0.3$), the Evenki, Mari, Udmurt and Kuusamo show intermediate values $(r_d = 0.1 - 0.3)$ and for the rest of the populations $r_d < 0.1$. Also, the results from Mann-Whitney U-test comparing the distribution of D' values across Xq13 with D'_{BASE} estimates between unlinked markers and thus minimizing sample size effect, demonstrate 'useful', background-level exceeding LD for the above-mentioned populations (Table 2). Pairwise D' estimates for the microsatellites across Xq13 (Figure 3) correlated largely with the calculations for the significance of the association by Fisher's exact test (data not shown). Current extended data of the LD structure across Xq13 reveals that additionally to previously described isolates with distinct demographic

Population			Xq13 e class (Mb)	Xp22 Distance class (Mb)						
	0-15	3–5 Mb	7–10 Mb	>10 Mb	0–2 Mb	3–5 Mb	5–10 Mb			
Saami	0.001	0.018	0.000	0.002	0.005					
Evenki	0.001	0.001	0.000	0.001						
Kuusamo	0.023	0.012	0.022	0.014			0.019			
Mari	0.003			0.024	0.007					
Udmurt	0.005		0.000		0.015		0.033			
Chuvash										
Komi										
Mordvin	0.007									
Russians										
Estonians										
Finns	0.023									
Swedes							0.006 ^a			
Germans										
Dutch			0.022							
Italians	0.043									

Table 2 Mann–Whitney U-test for the detection of background-level exceeding LD applicable for mapping purposes

Listed are the significant *P*-values (P < 0.05) from Mann–Whitney *U*-test comparing the distribution of pairwise *D'* values within each distance class for Xq13 or Xp22 markers and the baseline distribution D'_{BASE} estimates calculated between unlinked loci for each population. Significant difference indicates that LD in the studied region is exceeding the background level LD between unlinked markers on X chromosome.

^aIn Swedes, the significant *P*-value for Xp22 markers 5–10 Mb apart arises from HIGHER distribution of D' values between UNLINKED markers compared to pairwise D' values in this distance class.

Blanks – nonsignificant *P*-values.

histories, $^{10-12,17,27}$ also the Volga-Ural populations of Mari, Udmurt, Komi and Chuvash harbor increased level of LD across Xq13 compared to other European populations (Figure 3). Consistently, in contrast to single strong association (*P*<0.05) among 28 studied locus pairs for the

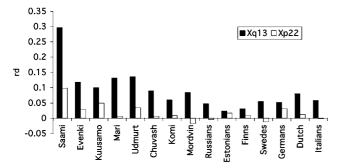


Figure 2 Comparative multilocus LD values (r_d) for the analyzed (1) a 10 kb subregion (five markers, DXS8037 to DXS986) from Xq13 (black columns); (2) 10 kb region (six markers) from Xp22 (white columns) in each studied population, calculated from the distribution of allelic mismatches between pairs of individuals over all loci. LD estimation is based on the variance of the number of pairwise differences among samples that have been subjected to genetic analysis at the multiple loci. Maximum value of $r_d = 1$ referring to absolute linkage disequilibrium across the whole region.

majority of populations, Fisher's exact test showed for the Mari 15, for Udmurts 11, for Chuvash 8 and Komi 7 loci in LD. This level of LD is comparable to the LD pattern of Xq13 from Kuusamo isolate (11/28 pairs P<0.05), where the increase in LD levels on X chromosome reflects a recent founder effect. As the current census sizes of these populations (500 000 to 1.8 million) exclude an extreme and recent bottleneck similar to Kuusamo, alternative scenarios could be considered responsible for increased LD.

European and East Asian populations form two clusters by STRUCTURE analysis: Volga-Ural populations fall to both clusters

In order to weigh the two alternative demographic scenarios – inner structuring into subgroups or admixture with Asian migrants – responsible for the elevated level of LD in Volga-Ural populations, the genetic structure of the study sample was analyzed by linkage-model based clustering method without prior assignment of individuals into populations.^{25,26} Multiple runs for data sets A ($n_{samples} = 1241$, $n_{pop} = 21$, $n_{loci} = 7$) and B ($n_{samples} = 889$, $n_{pop} = 16$, $n_{loci} = 14$) supported the estimate for K = 2, indicating two major population clusters among studied samples (Table 3). Data set C ($n_{samples} = 889$, $n_{pop} = 16$, $n_{loci} = 6$) did not resolve the population structure apparently due to low number of markers combined with the unbalanced sampling from East Asia (two populations)

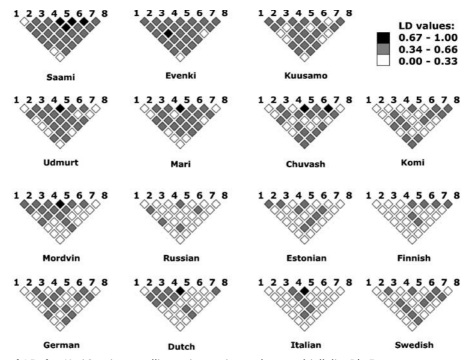


Figure 3 Patterns of LD for Xq13 microsatellite pairs, estimated as multiallelic *D'*. Every square represents the mean *D'* estimate between pairs of loci from random sampling of 40 individuals from each population sample over 100 replicates. The markers are indicated: 1 – DXS983, 2 – DXS8037, 3 – DXS8092, 4 – DXS1225, 5 – DXS8082, 6 – DXS986, 7 – DXS1066, 8 – DXS995.

	K – number of tested population clusters													
Log P(X K)	1	2	3	4	5									
$Log P(X_A K)$														
Run 1	-16126.0	-14688.0	-34335.2	-23891.5	-37730.7									
Run 2	-16125.8	-14685.1	-25566.5	-83364.7	-36122.0									
Run 3	-16125.9	-14690.8	-89021.3	-16694.5	-34648.7									
Run 4	-16124.9	-14689.1	-41689.0	-38380.4	-45344.6									
Run 5	ND	-14688.3	-26739.3	-18477.4	ND									
$Log P(X_B K)$														
Řun 1	-20778.1	-20269.8	-20319.5	-20677.4	ND									
Run 2	-20778.4	-20271.4	-20295.3	-20677.8	ND									
Run 3	-20778.2	-20270.2	-20284.1	-20679.1	ND									
Run 4	-20777.9	-20268.4	-20317.1	-20695.4	ND									
$Log P(X_C K)$														
Řun 1	-9088.8	-9880.1	-13246.7	-18025.6	ND									
Run 2	-9086.9	-9984.2	-14338.7	-17708.7	ND									
Run 3	-9085.7	-9853.9	-14879.9	-14698.5	ND									

 Table 3
 Inference for the number of populations (K) by STRUCTURE analysis

Estimates of model log-likelihood (log P(X|K)) for the data sets A (X_A) of seven markers from Xq13 for 21 population; data set B (X_B) of 14 markers from Xq13+Xp22 for 16 populations and data set C (X_C) of six markers from Xp22 for 16 populations. The highest log-likelihood in each row (bold) indicates the most supported K for each analysis of datasets A, B and C. ND- not done.

compared to Europe (14 populations). Based on data set A, for each individual the proportion of ancestry in both of the clusters was inferred (Figure 4). From the first glance it seems that one of the clusters is enriched in European and the other in Asian populations. Among the studied European populations almost all the individuals were assigned as most probably belonging to the 'European' cluster. On the other hand, almost all the Japanese and Evenki belong to the 'Asian' cluster. Consistent with known demographic history - admixture with Turkic tribes, as well as admixture LD in these populations shown by Katoh et al¹⁷ – the Mongolian populations of Zakhchin, Khoton, Uriankhai exhibit 2/3 of the 'Asian' and 1/3 'European' lineages. Minor European contribution was detected also for Buriats and Khalkh. Compared to other Europeans, the Volga-Ural populations of Mari, Chuvash, Udmurt and Komi, have a significant fraction of individuals belonging to the 'Asian' cluster. This strongly supports the interpretation of the increase of LD level in these populations owing to admixture with Asian migrants.

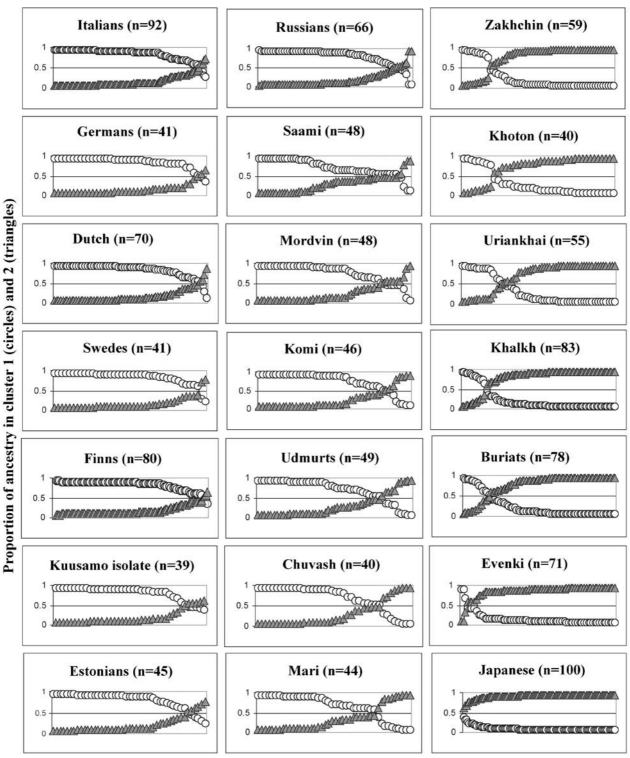
Haplotypes of nonrecombining loci DXS1225-DXS8082 support the hypothesis for admixture LD in Volga-Ural populations

At Xq13, the closest marker pair (162 kb apart) DXS1225– DXS8082 exhibited strong LD (P<0.000001) in all populations, irrespective of population structure or history, studied by us as well as other authors.^{17,27} Thus, as we can assume that recombination events between DXS1225 and DXS8082 are extremely rare, new variants are mostly created by mutation in one of the two loci. Table 4 summarizes the frequencies of 13 common haplotypes, present in one or more populations with frequencies exceeding 10%. Total number of DXS1225–DXS8082 haplotypes detected for a population sample ranged 10–27. Number of common haplotypes (>10% frequency) in each studied population ranged from 1 to 5, the haplotype diversities ranged from 0.19 in Japanese and the Saami to 0.44 in a Mongolian tribe of Uriankhai.

At first glance, there is a clear difference between European and Asian haplotype distributions. In European populations, except the Saami, across the vast area from West-Europe to the Urals one major haplotype (210–219) predominates (17 - 40%). Furthermore, in several populations this haplotype extends to neighboring DXS986 across 1.215 Mb. In East Asia, this haplotype is present at low frequency ($\leq 10\%$). On the contrary, the common haplotype detected in Asia (202-217) ranging from 16% in Uriankhai to 34% in Japanese, is almost absent in Europe. However, the exception here is the Volga-Ural region, where this East Asian haplotype is quite common among Udmurts (15%), Chuvash (10%) and Komi (11%). Overall, the distribution of common haplotypes of DXS1225-DXS8082 in Volga-Ural region is more complex compared to the vast area of the rest of Europe. Consistent with the STRUCTURE results it suggests admixture of mainly European gene pool with East Asians combined with influence of genetic drift. Notably, the allelic distributions of single microsatellites or haplotypes of weakly associated markers are mostly shared between the Volga-Ural and European populations, and have not preserved a footprint of East Asian migration (data not shown).

Comparing DXS1225–DXS8082 haplotype distribution in Finns and the Kuusamo isolate provides vivid evidence





Individuals in each populations

Figure 4 Summary of the clustering results (K = 2) for the data set A consisting of seven Xq13 microsatellite markers. In 21 studied populations, for each individual (*x*-axis of each graph) the mean value of the proportion of the ancestry in cluster 1 (white circles) and cluster 2 (gray triangles) was computed (*y*-axis of each graph). Based on the assignment of the majority of individuals of European origin into cluster 1 and of Asian origin into cluster 2, the two clusters are denoted in the text as 'European' and 'Asian'.

		Saa	Кии	lta	Du	Ger	Swe	Fin	Est	Ru	Mor	Ко	Chu	Udm	Ма	Eve	Bur	Zah ^a	Kho ^a	Ка ^а	Ur ^a	Jap ^a
Total haplotype no.		10	13	21	19	15	17	19	17	21	14	18	16	17	16	15	23	22	14	27	24	19
Diversity		0.19	0.33	0.23	0.28	0.37	0.41	0.24	0.38	0.32	0.29	0.39	0.40	0.35	0.36	0.21	0.29	0.37	0.35	0.33	0.44	0.19
Common haplotype no.		4	3	2	3	2	1	1	2	1	2	3	5	3	2	3	3	3	3	2	2	3
Fraction of carriers		0.74	0.59	0.42	0.60	0.39	0.27	0.31	0.36	0.40	0.52	0.39	0.61	0.58	0.32	0.54	0.55	0.40	0.40	0.41	0.38	0.70
Distribution of comm																						
DXS1225 allele DX	S8082 allel	e					Eur	ope					Volga	a-Ural					East As	ia		
192	227			*	*			*	*		*	*	**	0.13			*			*		
	229	*	*	*	0.10	0.12	**	*	0.11	**	**	**	**	**	*			*	*	*		
198	225					*		*	*	*	*				*	**	0.15	**	**	**	0.22 ^c	0.20
	227	**	**	*	*	*	*	*	*	**	*	**	0.10	*	**	0.10	*		0.10 ^b	*	*	**
	229	0.20 ^c	**	**	*	**	**	**	*	*		*	*	*	*		*	**	**	*		*
200	225									*			*			*	0.11	*	*	0.11	*	0.16
	229	_														*			0.15			
202	211	0.20 ^c		0.13	0.14	*	**	**	**	*	0.17	0.11	0.10	*	0.16 ^c	*	*		*	*	*	**
	217	0.13 ^b	**							*	*	0.11	0.10	0.15		0.23	0.29	0.20 ^c	0.15 ^c	0.30 ^c	0.16	0.34
	219	_			_			_	*		_				*	0.21 ^c	*	0.10			*	*
206	219	_		*	*	*	**	**	**	*	**	**	0.13 ^c		0.16	*		*	**			
210	219	**	0.28	0.29 ^c	0.36 ^c	0.27 ^c	0.27	0.31 ^c	0.25	0.40 ^c	0.35 ^c	0.17 ^c				*	**	0.10	**	*	*	*
212	219	0.20 ^b				*	*	*	*			*	**			_		*	*	*		

Table 4 Common haplotypes of microsatellite loci DXS1225–DXS8082 with frequency >0.1 for one or more populations

Haplotype frequency: *<0.05; **0.05-0.1; — missing. ^aData from Katoh *et al* (2002): Mongolian populations of Zahkchin, Khoton, Khalkh and Uriankhai, and Japanese population sample. ^bHaplotype associated with a single allele at DXS986 extending to a three-locus haplotype across 1.215 Mb. ^cHaplotype associated with reduced number of alleles at DXS986, presence of a common (>10% frequency) three-locus haplotype.

Bdh <u>1</u>60

for the impact of genetic drift in changing the allele and haplotype frequencies within a short period of time. Although the isolate shares the major haplotype with Finns (28 vs 31%), there are two other enriched haplotypes (both 15%). As one of them, 212–219, is also common (20%) among the Saami living originally at the Kuusamo area, and rare in the rest of Europe, it could also reflect admixture with local Saami people during the establishment of the population.

Discussion

Demographic history has the strongest impact on LD patterns of recombination-poor regions

This study shows that demographic history has strong impact on local LD and haplotype patterns across a 20 Mb, but only 4.74 cM, genomic segment at Xq13. No such strong LD generating effect was detected across a 9.207 Mb region at Xp22 corresponding to 12.01 cM. As microsatellites genotyped for Xq13 and Xp22 were characterized by similar locus diversity, mean number of alleles and population diversity parameter θ , we can leave aside a scenario of higher mutation rate of Xp22 markers responsible for diversifying haplotype and LD patterns. Our result is concordant with the recent simulation study by Stumpf and Goldstein,²⁸ which demonstrated that following the LD-generating event, the differences among genomic regions in preserving a block-like structure depend on recombination rate. In our data set outbred and expanded populations, independent of their size, exhibited significant LD at Xq13 between only one single locus pair DXS1225-DXS8082. The reason for the strong LD between these markers, located 162 kb apart and apparently within an LD desert, is still to be studied. In our data set LD-generating events across Xq13 included not only genetic drift in a small population or severe foundereffect, but also admixture with genetically different migrants. Volga-Ural populations, distributed at the geographic borderline of Europe and Asia, have apparently historically lived in close contact with their East Asian neighbors. Both, population structure and haplotype analysis supported the hypothesis that the increased level of LD in these populations is due to admixture of mostly European genepool with East Asians. This level of LD is similar to the extent observed for X-chromosomal loci in a Bantu-Semidic hybrid population of Lemba.²⁹ For Lemba, similar to Komi and Chuvash, the elevated background LD was observed also for unlinked markers on X-chromosome. The strongest LD across Xq13 region was detected for Udmurt and Mari. There the inner structuring of the population could additionally contribute to the creation of nonrandom allelic associations.

Implications for mapping using LD

Two recent extensive scans for the landscape of LD and haplotype variation across human genome in distant

population groups point out that there is a lot of heterogeneity in the LD map as well as haplotype frequencies among populations.^{30,31} There is also evidence that the intervals across which LD is detectable depend on marker properties. Varilo *et al*¹³ showed that single informative microsatellites provide more power to detect long-range LD than did single SNPs or even 3-5 SNP haplotypes, It has been shown that long-range microsatellite data can be used to predict short-range LD between SNPs and thus assist in initial association analysis.^{32,33} In addition, our study emphasizes the importance of calculating the baseline LD between unlinked markers for each data set used in the study. The true indication of the increase of LD in a particular genomic region is in comparison to the baseline LD.

Data on X-chromosomal microsatellites show that the footprint of a demographic event persists longer in haplotype distribution within a region of low crossingover rate than in the information content of a single marker or between the actively recombining markers. The distribution of the haplotypes of strongly associated DXS1225-DXS8082 markers varies between populations, memorizing the demographic events of a population. For example, when bottleneck is accompanied with low level admixture, the few migrant haplotypes might drift frequent in the descendant population as proved by the 212-219 haplotype distribution in the Kuusamo and Saami (Table 4). Also, inner structuring within populations can lead to additional haplotype frequency differences. There are more and more data, which highlight the importance of taking into account the particular population history and its impact on regional LD patterns. Laan and Pääbo³⁴ have compared the allelic associations around renin-binding protein RnBP, a component of the reninangiotensin system, in the Saami and Finns. The minor allele of an SNP within the gene, T61C, present both on the Saami (21%) as well as Finns (19%) as common mutation, was associated with different alleles and haplotypes of flanking microsatellites in two populations. The recently described association between SNPs of the CARD15 (NOD2) gene and Crohn's disease³⁵ did not find any support in the respective study with Korean patients.³⁶ The three disease-associated SNPs sharing a common haplotypic background were absent in the Korean sample and the LD pattern across CARD15 differed between the studied populations.

In conclusion, our study demonstrates that demographic events leave their prolonged imprint on LD patterns across recombination-poor genomic regions. Consequently, as the haplotype distribution within LDrich blocks might exhibit much more variability among populations than previously expected, the key for successful gene mapping studies is detailed understanding of the history, structure and variation of the study population.

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