Human Populations: History and Structure

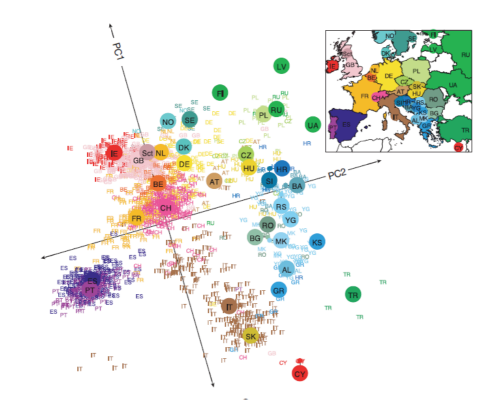
In the paper

Novembre J, Johnson, Bryc K, Kutalik Z, Boyko AR, Auton A, Indap A, King KS, Bergmann A, Nelson MB, Stephens M, Bustamante CD. 2008. Genes mirror geography within Europe. Nature 456:98

there is quite dramatic evidence that our genetic profiles contain information about where we live, suggesting that these profiles reflect the history of our populations.

The authors collected "SNP" (single nucleotide polymorphism) data on over people living in Europe. Either the country of origin of the people's grandparents or their own country of birth was known. On the next slide, these geographic locations were used to color the location of each of 1,387 people in "genetic space." Instead of latitude and longitude on a geographic map, their first two principal components were used: these components summarize the 500,000 SNPs typed for each person.

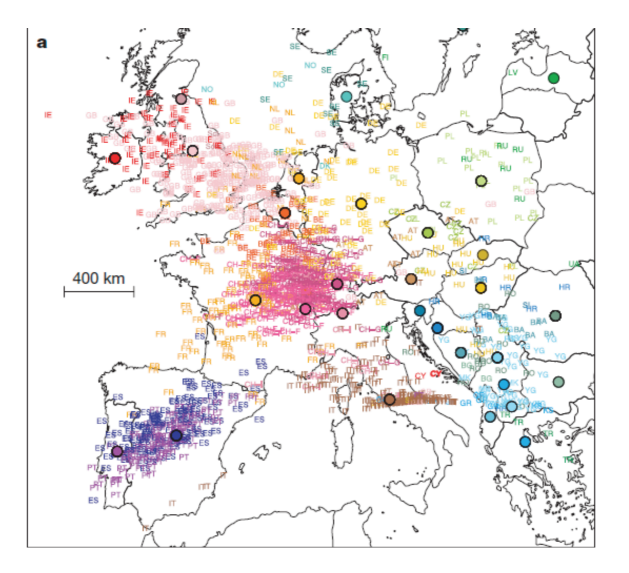
Novembre et al., 2008



Novembre et al., 2008

As a follow-up, the authors took the genetic profile of each person and used it to predict their latitude and longitude, and plotted these on a geographic map. These predicted positions are colored by the country of origin of each person.

Novembre et al., 2008



Y SNP Data Haplogroups

Another set of SNP data, this time from around the world, is available for the Y chromosome. These data were collected for the 1000 Genomes project (http://www.1000genomes.org/): there are 26 populations:

East Asia: CDX. Chinese Dai in Xishuangbanna; CHB. Han Chinese in Beijing; JPT. Japanese in Tokyo; KHV. Kinh in Ho Chi Minh City; CHS. Southern Han Chinese.

South Asian: BEB. Bengali in Bangladesh; GIH. Gujarati Indian in Houston; ITU. India Telugi in UK; PJL. Punjabi in Lahore; STU. Sri Lankan Tamil in UK.

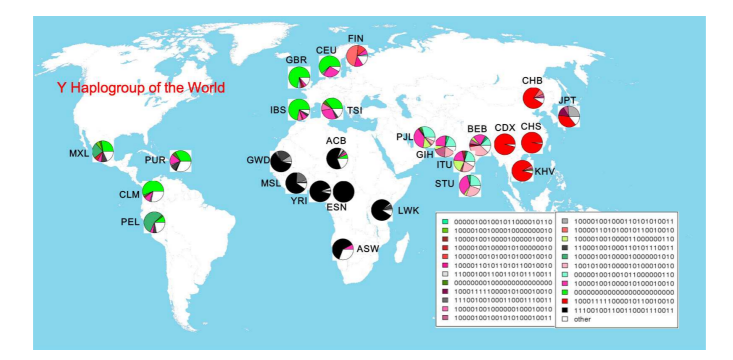
Y SNP Data Haplogroups

African: ASW. African Ancestry in Southwest US; ACB. African Caribbean in Barbados; ESN. Esan in Nigeria; GWD. Gambian in the Gambia; LWK. Luthya in Kenya; MSL. Mende in Sierra Leone; YRI. Yoruba in Nigeria.

European: GBR. British in UK; FIN. Finnish in Finland; IBS. Iberian in Spain; TSI. Toscani in Italy; CEU. Utah residents with European ancestry.

Americas: CLM. Columbian in Medellin; MXL. Mexican in Los Angeles; PEL. Peruvian in Lima, PUR. Puerto Rican in Puerto Rico.

Y SNP Data Haplogroups



Migration History of Early Humans

An interesting video of the migration of early humans is available at:

http://www.bradshawfoundation.com/journey/

Migration Map of Early Humans

https://genographic.nationalgeographic.com/human-journey/

This map summarizes the migration patterns of early humans.

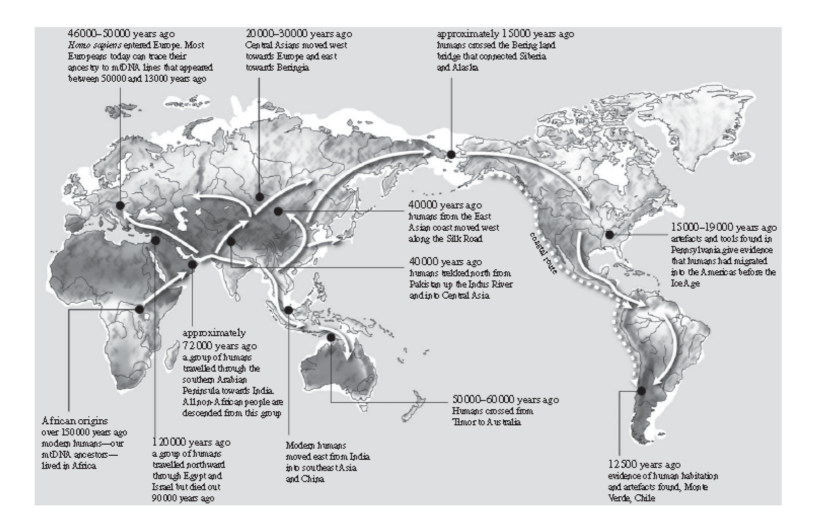
Migration Map of Early Humans

The map on the next slide, based on mitochondrial genetic profiles, is taken from:

Oppenheimer S. 2012. Out-of-Africa, the peopling of continents and islands: tracing uniparental gene trees across the map. Phil. Trans. R. Soc. B (2012) 367, 770-784 doi:10.1098/rstb.2011.0306.

The first two pages of this paper give a good overview, and they contain this quote: "The finding of a greater genetic diversity within Africa, when compared with outside, is now abundantly supported by many genetic markers; so Africa is the most likely geographic origin for a modern human dispersal."

Migration Map of Early Humans



Forensic Implications

What does the theory about the spread of modern humans tell us about how to interpret matching profiles?

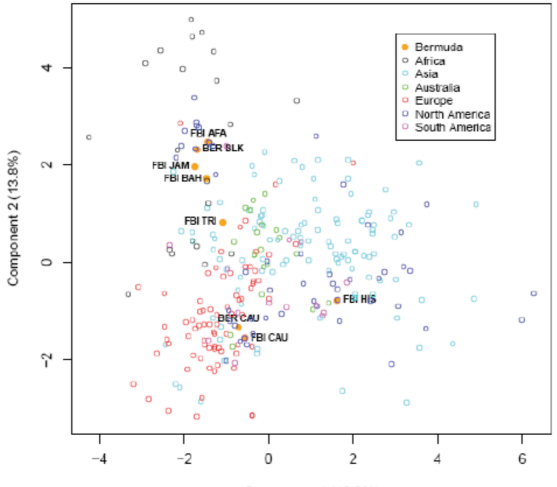
Matching probabilities should be bigger within populations, and more similar among populations that are closer together in time.

Forensic allele frequencies are consistent with the theory of human migration patterns.

Forensic STR PCA Map

A large collection of forensic STR allele frequencies was used to construct the principal component map on the next page. Also shown are some data collected by forensic agencies in the Caribbean, and by the FBI. The Bermuda police has been using FBI data - does this seem to be reasonable?

Forensic STR PCA Map



Component 1 (18.3%)

Forensic allele frequencies were collected from 21 populations. The next slides list the populations and show allele frequencies for the Gc marker. This has only three alleles, A, B, C.

The matching proportions within each population, and between each pair of populations, were calculated. These allow distances ("theta" or β_{ij}) to be calculated for each pair of populations i, j: $\hat{\beta}_{ij} = (\tilde{M}_i + \tilde{M}_j - 2\tilde{M}_{Bij})/[2(1 - \tilde{M}_{Bij})].$

Gc Frequencies (1)

| Symbol | Description |
|--------|----------------------|
| AFA | FBI African-American |
| AL1 | North Slope Alaskan |
| AL2 | Bethel-Wade Alaskan |
| ARB | Arabic |
| CAU | FBI Caucaslan |
| CBA | Colmbran |
| DUT | Dutch Caucaslan |
| GAL | Gallclan |
| HN1 | Hungarlan |
| HN2 | Hungarlan |
| IT2 | Itallan |

Gc Frequencies (2)

| Symbol | Description |
|--------|---------------------------|
| IT4 | Itallan |
| KOR | Korean |
| NAV | NavaJo |
| NBA | North Bavarlan |
| PBL | Pueblo |
| SEH | FBI Southeastern Hispanic |
| SOU | Sloux |
| SPN | Spanlsh |
| SWH | FBI Southwestern Hispanic |
| SWI | Swlss Caucaslan |



Gc Frequencies (3)

| Popn. | Sample size | А | В | С |
|-------|-------------|------|------|------|
| AFA | 145 | .338 | .237 | .423 |
| AL1 | 96 | .177 | .489 | .334 |
| AL2 | 112 | .236 | .451 | .313 |
| ARB | 94 | .133 | .441 | .425 |
| CAU | 148 | .114 | .456 | .429 |
| CBA | 119 | .159 | .533 | .306 |
| DUT | 155 | .106 | .422 | .471 |
| GAL | 143 | .140 | .448 | .413 |
| HN1 | 345 | .106 | .457 | .438 |
| HN2 | 163 | .097 | .448 | .454 |
| IT2 | 374 | .139 | .454 | .408 |



Gc Frequencies (4)

| Popn. | Sample size | А | В | С |
|-------|-------------|------|------|------|
| IT4 | 200 | .302 | .163 | .535 |
| KOR | 116 | .310 | .422 | .267 |
| NAV | 81 | .105 | .240 | .654 |
| NBA | 150 | .133 | .383 | .484 |
| PBL | 103 | .102 | .374 | .524 |
| SEH | 94 | .165 | .447 | .389 |
| SOU | 64 | .055 | .422 | .524 |
| SPN | 132 | .118 | .474 | .409 |
| SWH | 96 | .156 | .437 | .407 |
| SWI | 100 | .135 | .465 | .400 |

Gc Theta Distances (1)

| | AFA | AL1 | AL2 | ARB | CAU | CBA | DUT | GAL | HN1 | HN2 |
|-----|------|------|------|------|------|------|------|------|------|------|
| AL1 | .201 | | | | | | | | | |
| AL2 | .163 | .000 | | | | | | | | |
| ARB | .224 | .002 | .016 | | | | | | | |
| CAU | .303 | .020 | .046 | .008 | | | | | | |
| CBA | .309 | .017 | .034 | .022 | .009 | | | | | |
| DUT | .341 | .039 | .070 | .021 | .000 | .017 | | | | |
| GAL | .295 | .015 | .037 | .007 | .000 | .004 | .002 | | | |
| HN1 | .339 | .040 | .072 | .025 | .001 | .013 | .000 | .002 | | |
| HN2 | .348 | .041 | .073 | .024 | .000 | .016 | .000 | .003 | .000 | |
| IT2 | .304 | .023 | .048 | .015 | .000 | .004 | .002 | .000 | .001 | .002 |

Gc Theta Distances (2)

| | AFA | AL1 | AL2 | ARB | CAU | CBA | DUT | GAL | HN1 | HN2 |
|-----|------|------|------|------|------|------|------|------|------|------|
| IT4 | .088 | .029 | .022 | .032 | .085 | .098 | .111 | .081 | .120 | .117 |
| KOR | .074 | .051 | .026 | .082 | .139 | .122 | .175 | .128 | .179 | .179 |
| NAV | .242 | .060 | .080 | .028 | .054 | .103 | .063 | .061 | .075 | .070 |
| NBA | .278 | .017 | .041 | .002 | .000 | .018 | .004 | .001 | .007 | .006 |
| PBL | .178 | .033 | .044 | .015 | .051 | .085 | .067 | .053 | .077 | .073 |
| SEH | .254 | .001 | .015 | .000 | .002 | .005 | .014 | .000 | .014 | .015 |
| SOU | .294 | .035 | .062 | .008 | .010 | .046 | .012 | .015 | .020 | .016 |
| SPN | .315 | .022 | .048 | .012 | .000 | .005 | .000 | .000 | .000 | .000 |
| SWH | .269 | .004 | .022 | .000 | .000 | .004 | .008 | .000 | .009 | .009 |
| SWI | .298 | .013 | .035 | .007 | .000 | .002 | .002 | .000 | .002 | .003 |

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Gc Theta Distances (3)

| | IT2 | IT4 | KOR | NAV | NBA | PBL | SEH | SOU | SPN | SWH |
|-----|------|------|------|------|------|------|------|------|------|------|
| IT4 | .098 | | | | | | | | | |
| KOR | .145 | .026 | | | | | | | | |
| NAV | .072 | .048 | .143 | | | | | | | |
| NBA | .005 | .067 | .127 | .034 | | | | | | |
| PBL | .066 | .016 | .088 | .003 | .032 | | | | | |
| SEH | .004 | .052 | .089 | .054 | .003 | .038 | | | | |
| SOU | .021 | .067 | .148 | .011 | .001 | .021 | .019 | | | |
| SPN | .000 | .093 | .144 | .066 | .002 | .061 | .003 | .016 | | |
| SWH | .001 | .060 | .102 | .053 | .000 | .040 | .000 | .014 | .000 | |
| SWI | .000 | .079 | .125 | .062 | .001 | .054 | .000 | .016 | .000 | .000 |

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Clustering Populations

Populations can be clustered on the basis of the genetic distances between them. For short-term evolution (among human populations) the simple UPGMA method performs satisfactorily. The closest pair of populations are clustered, and then distances recomputed from each other population to this cluster. Then the process continues.



Clustering 4 Populations

Look at four of the populations:

| | AFA | CAU | SEH | NAV |
|-----|-------|-------|-------|-----|
| AFA | - | | | |
| CAU | 0.303 | _ | | |
| SEH | 0.254 | 0.002 | _ | |
| NAV | 0.242 | 0.054 | 0.054 | - |

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Cluster Distances

The closest pair is CAU/SEH. Cluster them, and compute distances from the other two to this cluster:

AFA dlstance = (0.303+0.254)/2 = 0.278NAV dlstance = (0.054+0.054)/2 = 0.054



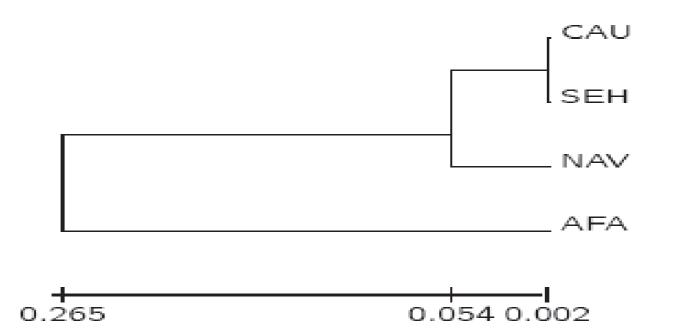
Cluster Distances (2)

The new distance matrix is

| | AFA | CAU/SEH | NAV |
|---------|-------|---------|-----|
| AFA | - | | |
| CAU/SEH | 0.278 | - | |
| NAV | 0.242 | 0.054 | - |

and the next shortest distance is between NAV and CAU/SEH.

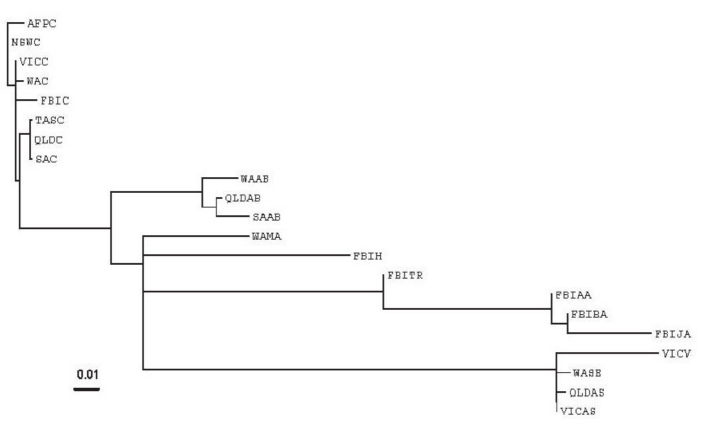






Australian STR Data

Australian Values





Worldwide Survey of STR Data

Published allele frequencies for 24 STR loci were obtained for 446 populations. For each population *i*, the withinpopulation matching proportion \tilde{M}_i was calculated. Also the average \tilde{M}_B of all the between-population matching proportions. The " θ " for each population is calculated as $\hat{\beta}_i =$ $(\tilde{M}_i - \tilde{M}_B)/(1 - \tilde{M}_B)$. These are shown on the next slide, ranked from smallest to largest and colored by continent.

Africa: black; America: red; South Asia: orange; East Asia: yellow; Europe: blue; Latino: turquoise; Middle East: grey; Oceania: green.

Worldwide Survey of STR Data

