Lecture 18. Ancient and Modern Humans

Michael Schatz

April 8, 2019

JHU 600.749: Applied Comparative Genomics



Preliminary Project Report

Assignment Date: April 8, 2019

Due Date: Monday, April 15, 2017 @ 11:59pm

Each team should submit a PDF of your preliminary project proposal (2 to 3 pages) to GradeScope by 11:59pm on Monday April 15.

The preliminary report should have at least:

- · Title of your project
- List of team members and email addresses
- 1 paragraph abstract summarizing the project
- 1+ paragraph of Introduction
- 1+ paragraph of Methods that you are using
- 1+ paragraph of Results, describing the data evaluated and any any preliminary results
- 1+ paragraph of Dicsussion (what you have seen or expect to see)
- 1+ figure showing a preliminary result
- 5+ References to relevant papers and data

The preliminary report should use the Bioinformatics style template. Word and LaTeX templates are available at https://academic.oup.com/bioinformatics/pages/submission_online

Later, you will present your project in class starting the week of April 24. You will also submit your final written report (5-7 pages) of your project by May 15

Please use Piazza if you have any general questions!

Midterm Exam Page 1 of 6

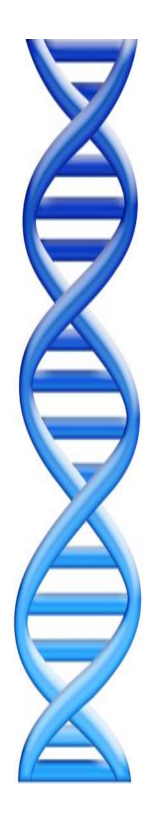
601.749 Computational Genomics: Applied Comparative Genomics Midterm Exam

Michael C. Schatz mschatz@cs.jhu.edu

April 3, 2019 Time: 75 Minutes

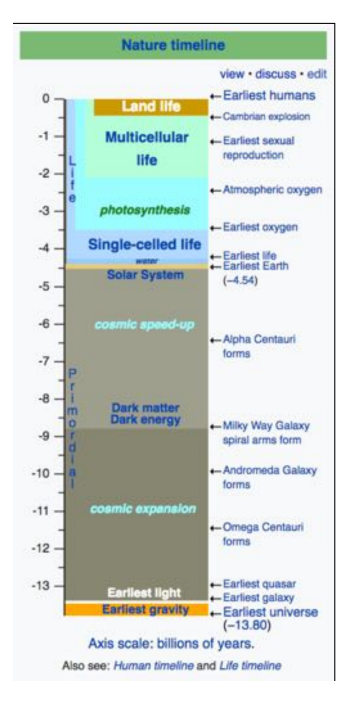
Start here: Please fill in the following important information using a permanent pen before you do anything else! Your exam will not be graded if you use a pencil or erasable ink on this page.

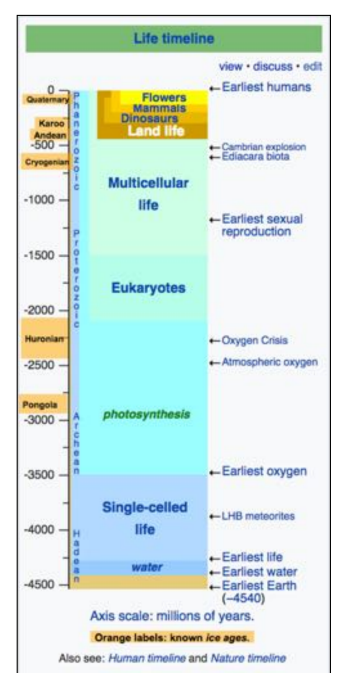
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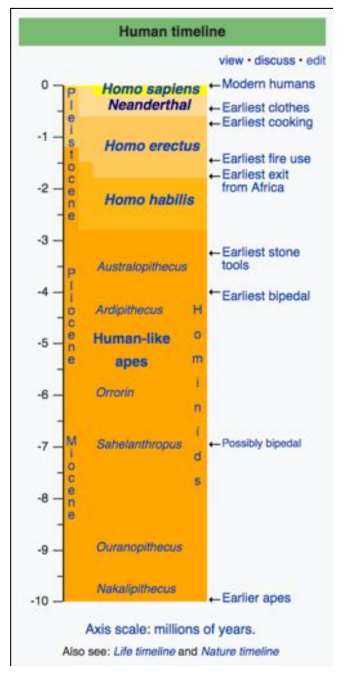


Part I: Ancient Hominds

Our Origins







Sequencing ancient genomes

Janet Kelso Max-Planck Institute





Homo neanderthalensis

- Proto-Neanderthals emerge around 600k years ago
- •"True" Neanderthals emerge around 200k years ago
- •Died out approximately 40,000 years ago
- •Known for their robust physique
- •Made advanced tools, probably had a language (the nature of which is debated and likely unknowable) and lived in complex social groups



Homo sapiens

- Apparently emerged from earlier hominids in Africa around 50k years ago
- Capable of amazing intellectual and social behaviors
- Mostly Harmless ☺



A Draft Sequence of the Neandertal Genome

Richard E. Green, et al. Science 328, 710 (2010);

DOI: 10.1126/science.1188021

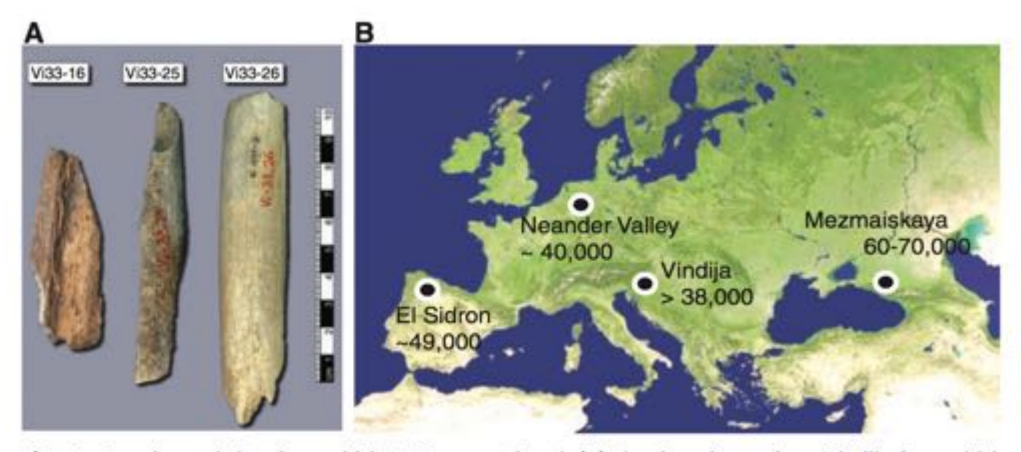
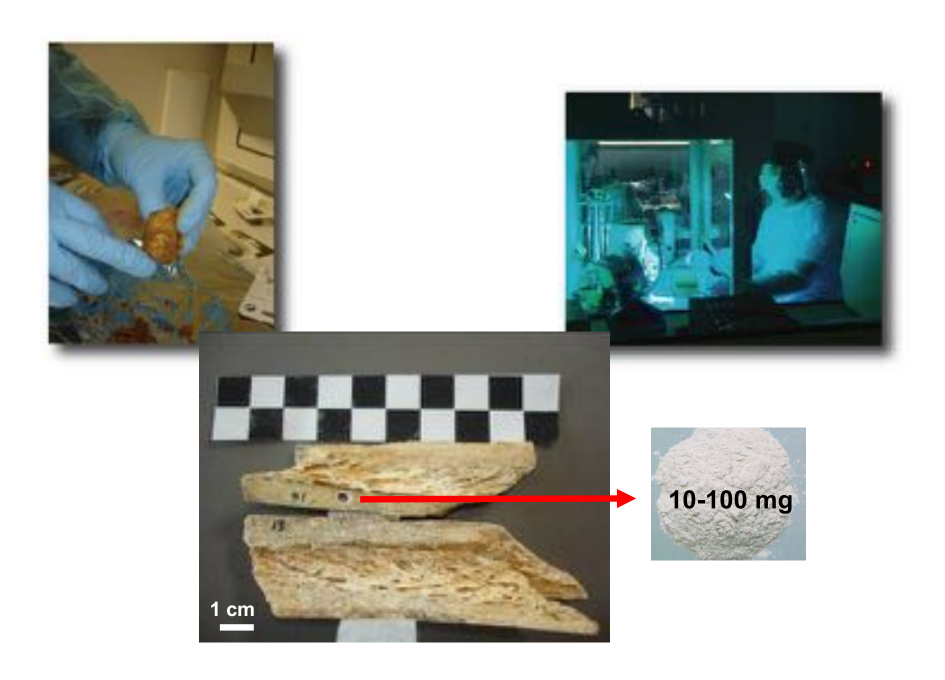
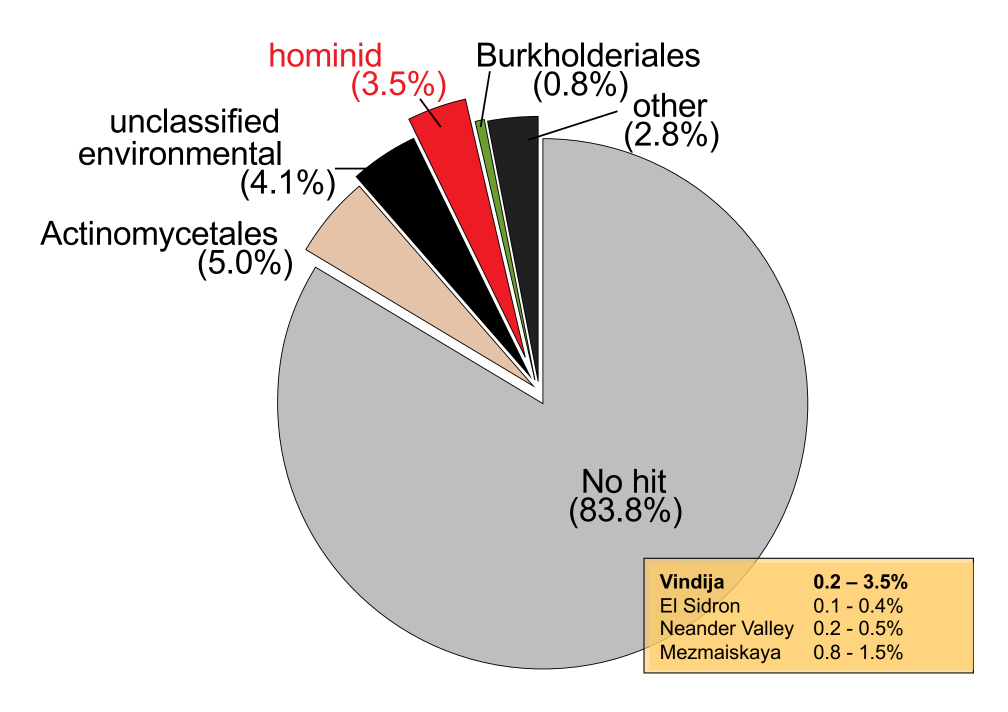


Fig. 1. Samples and sites from which DNA was retrieved. (A) The three bones from Vindija from which Neandertal DNA was sequenced. (B) Map showing the four archaeological sites from which bones were used and their approximate dates (years B.P.).

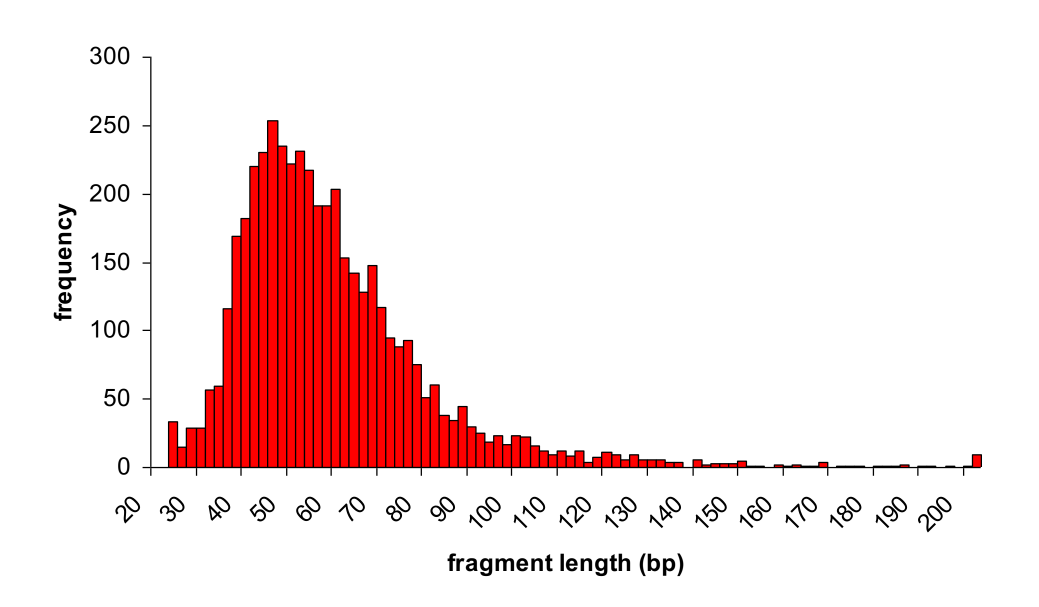
Extracting Ancient DNA



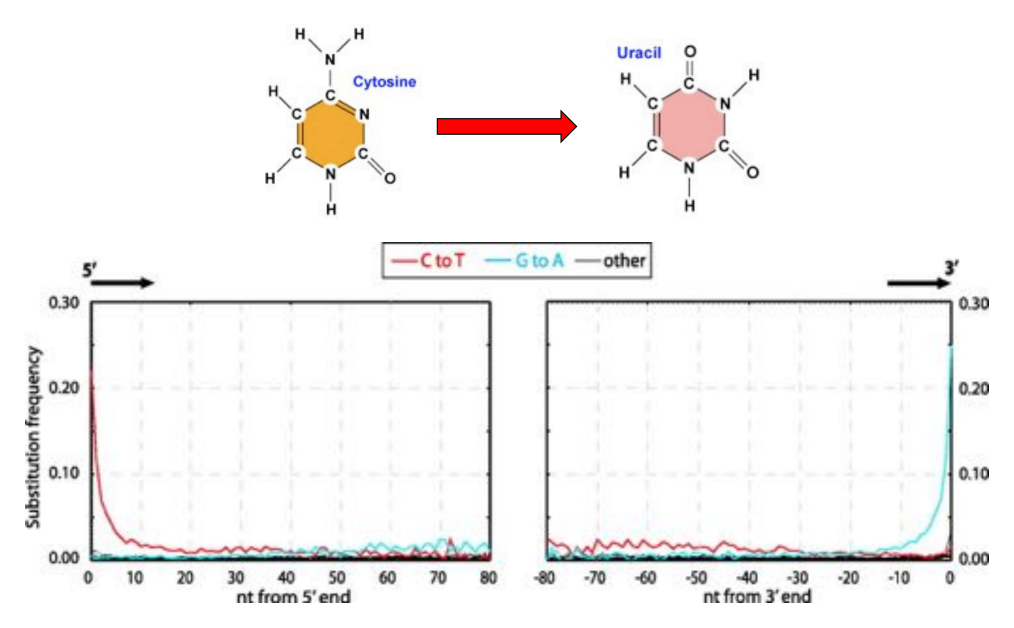
DNA is from mixed sources



DNA is degraded



DNA is chemically damaged





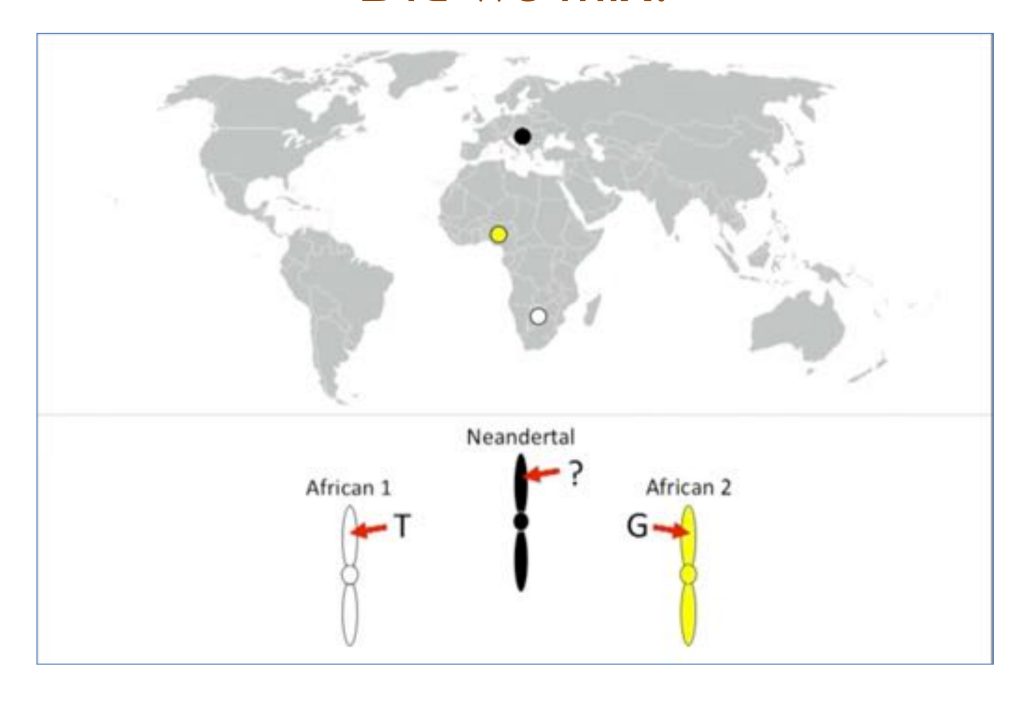
Green et al. 2010

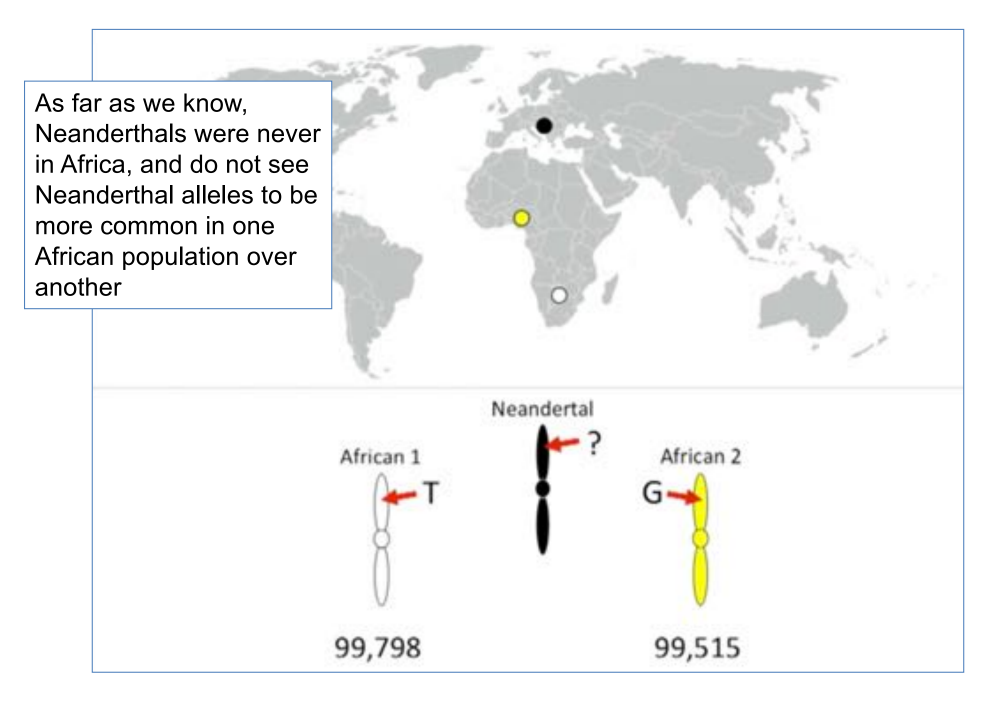
Vindija 33.16 ~1.2 Gb 33.25 ~1.3 Gb 33.26 ~1.5 Gb

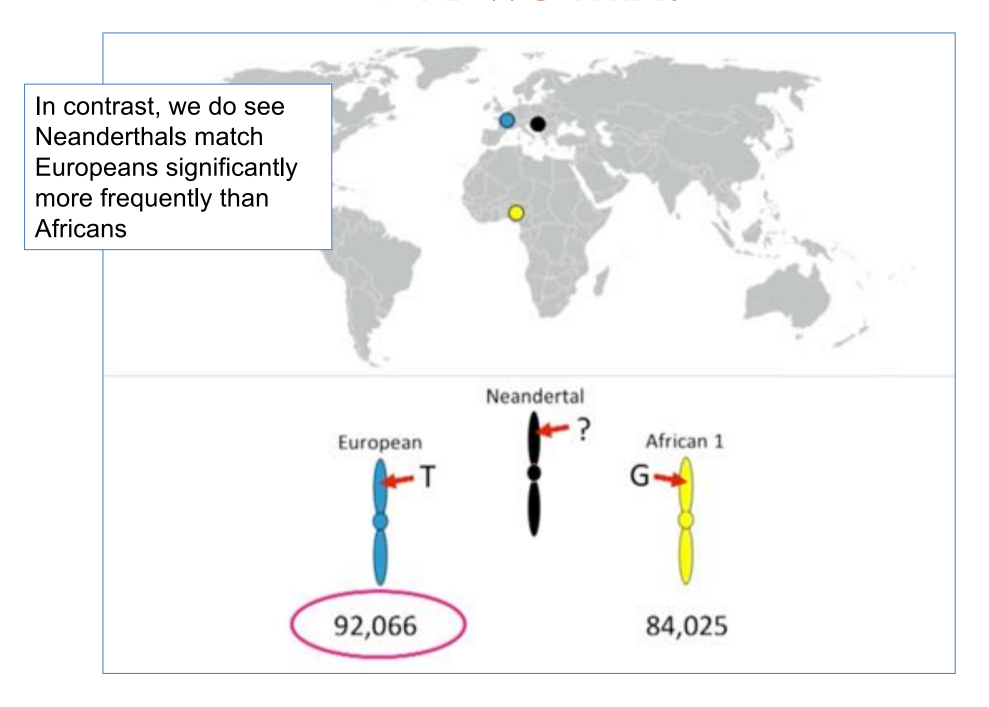
El Sidron (1253) ~2.2 Mb Feldhofer 1 ~2.2 Mb Mezmaiskaya 1 ~56.4 Mb

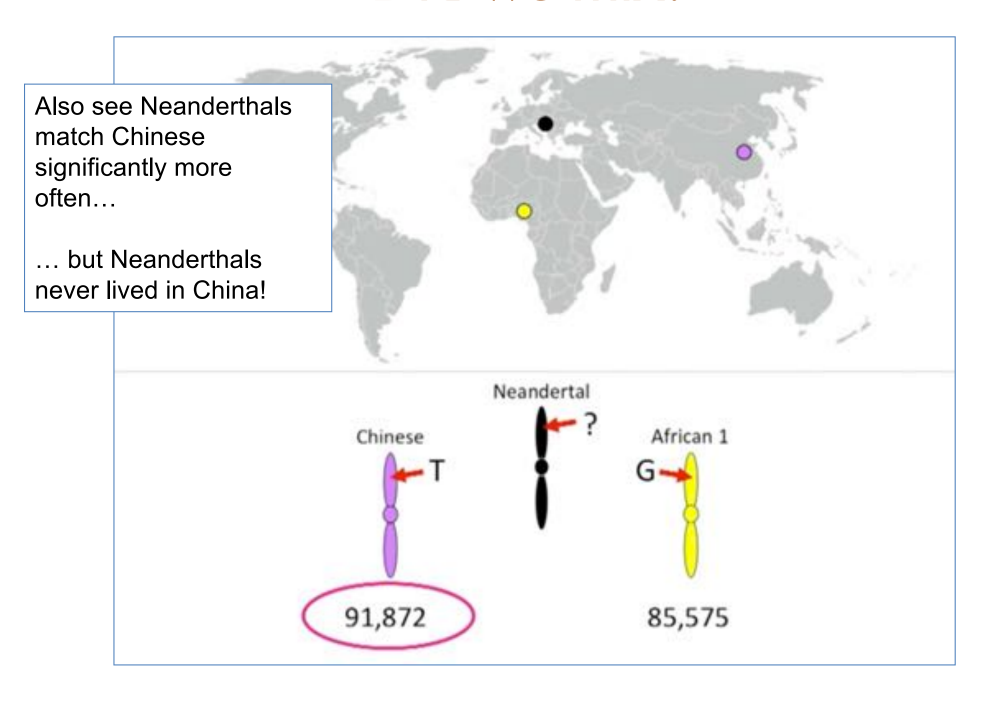
~35 Illumina flow cells

Genome coverage ~1.3 X

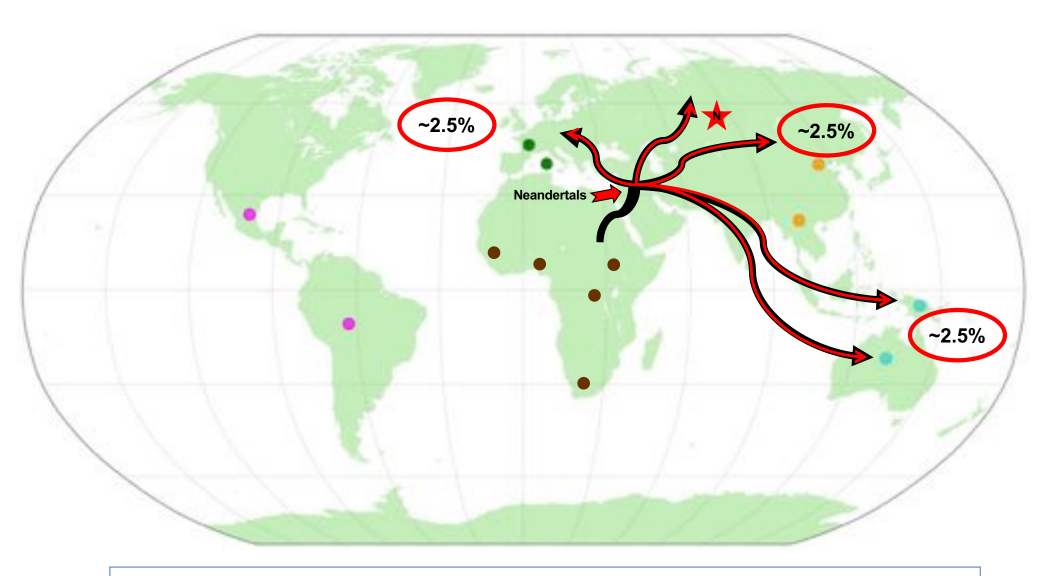








Neanderthal Interbreeding



As modern humans migrated out of Africa, they apparently interbred with Neanderthal's so we see their alleles across the rest of the world and carry about 2.5% of their genome with us!

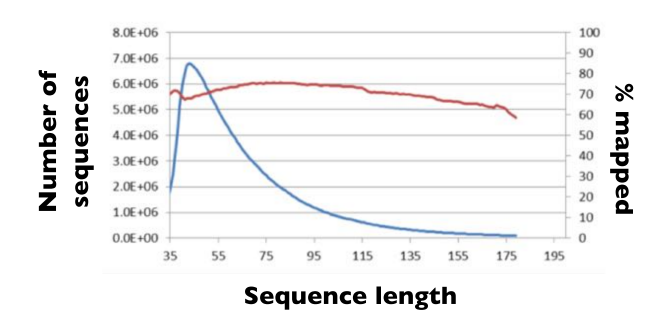
What about other ancient hominids?

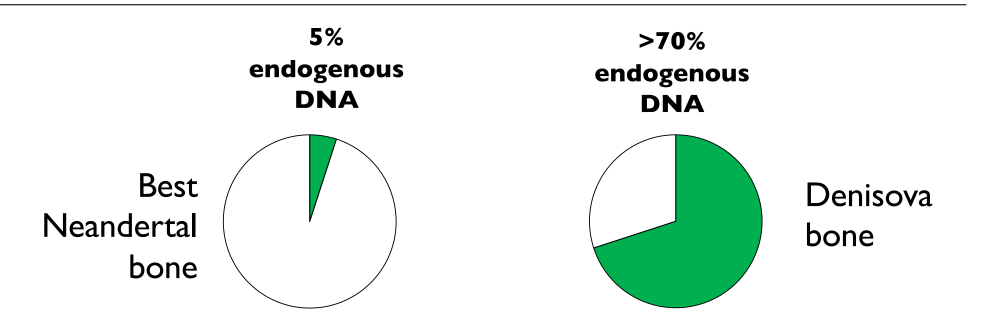




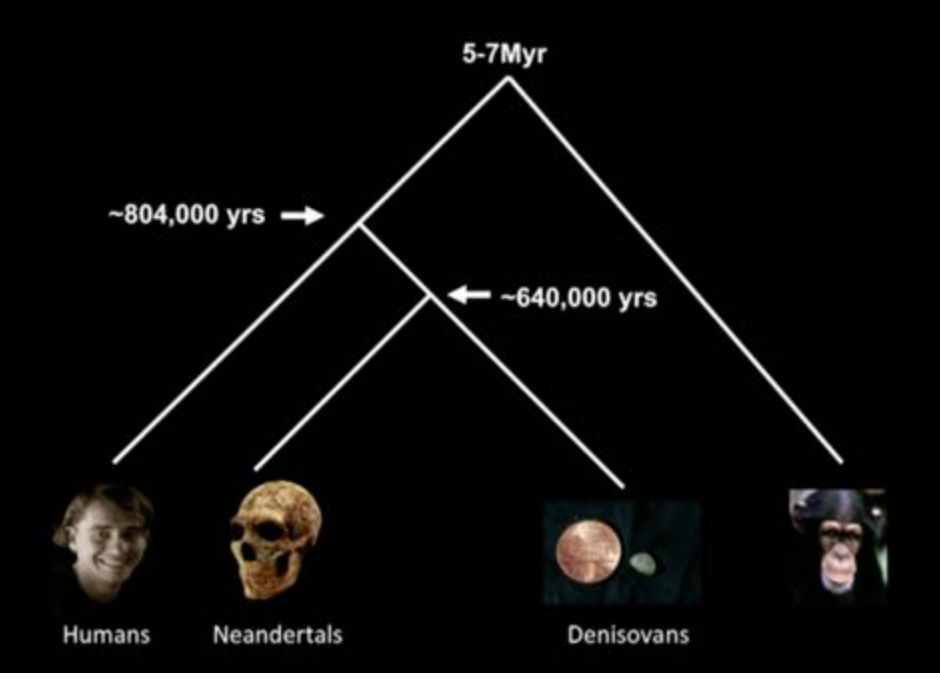


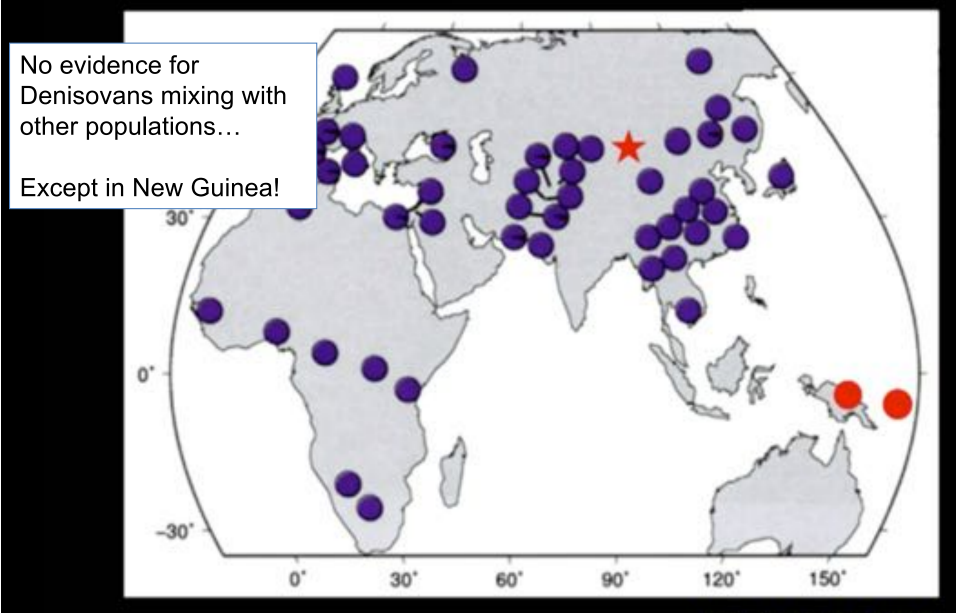
Extraordinary preservation

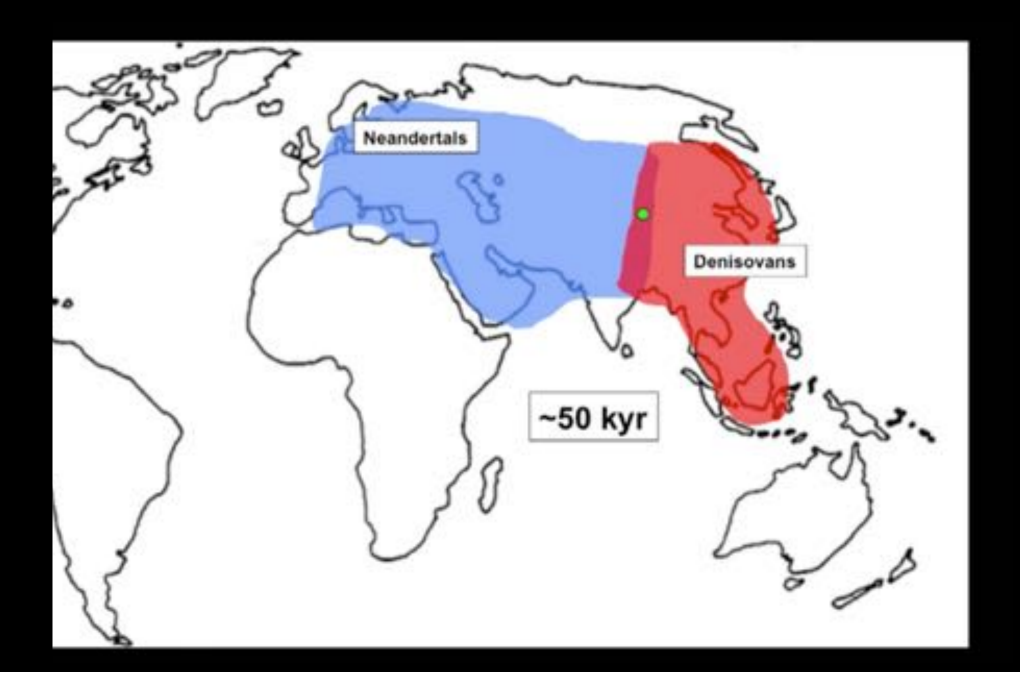


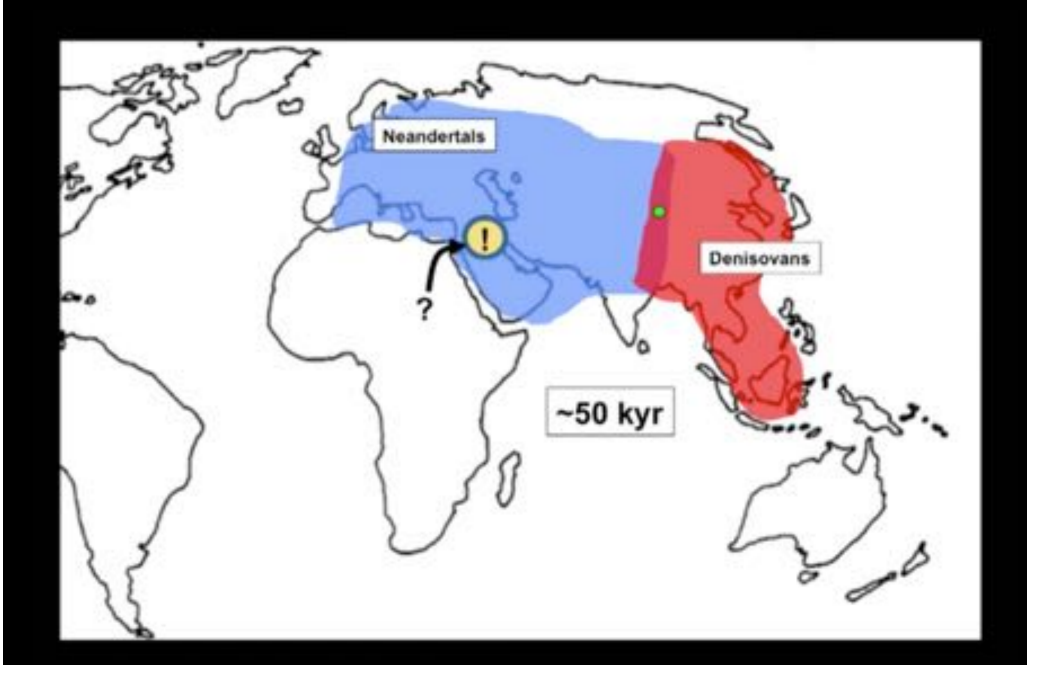


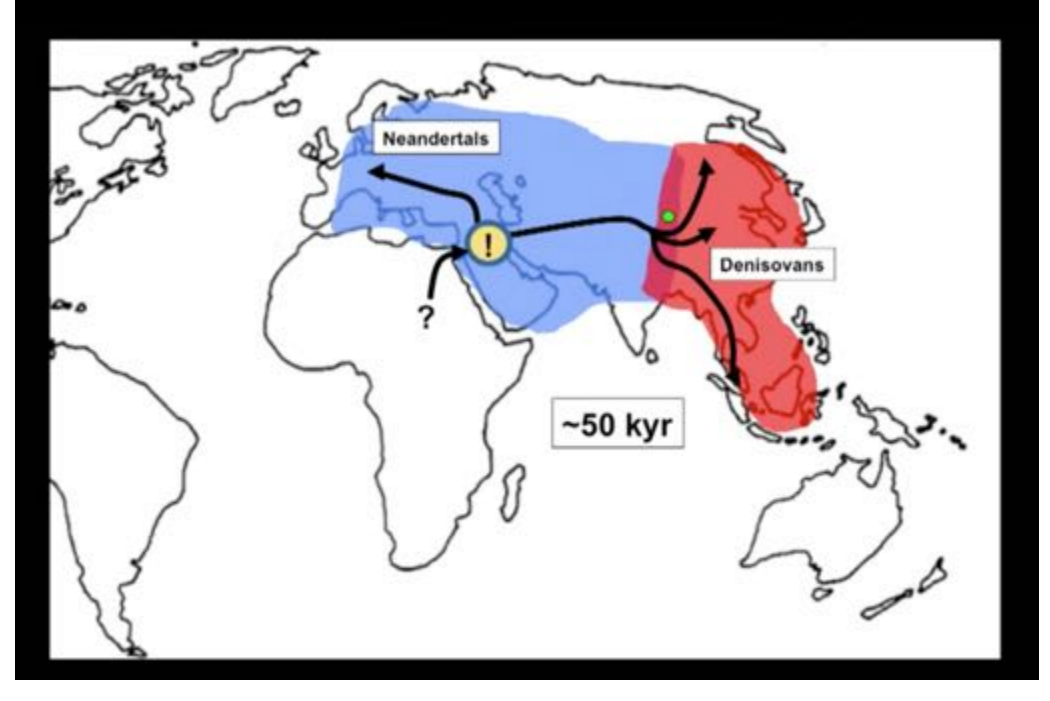
Denisovans & Neandertals

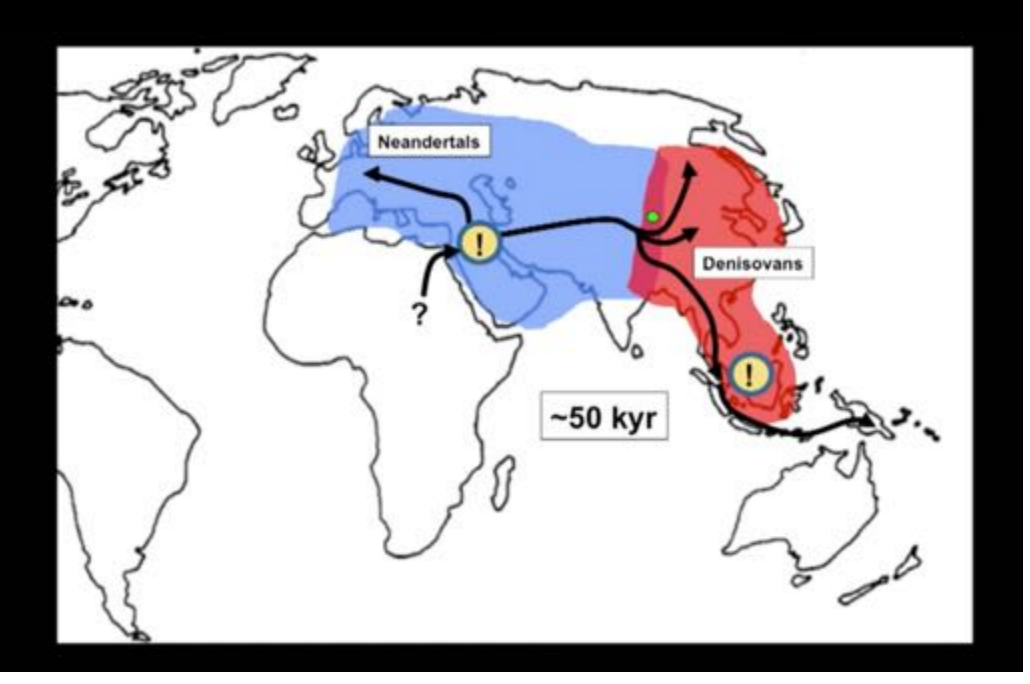


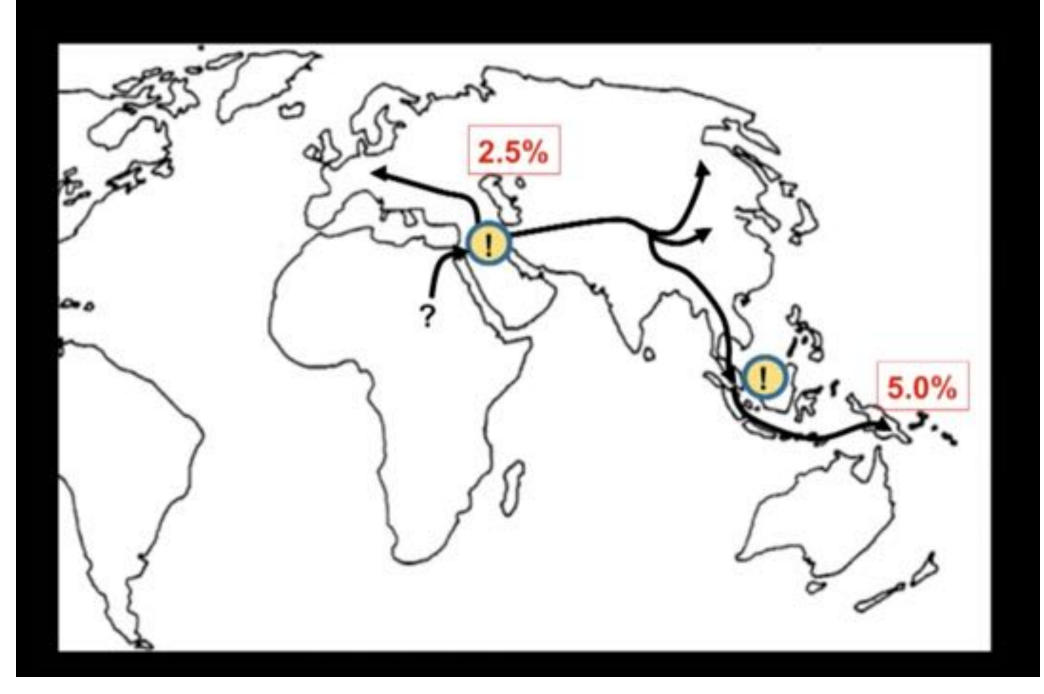












We have always mixed!



Cite as: B. Vernot et al., Science 10.1126/science.aad9416 (2016).

Excavating Neandertal and Denisovan DNA from the genomes of Melanesian individuals

Benjamin Vernot, Serena Tucci, Sanet Kelso, Joshua G. Schraiber, Aaron B. Wolf, Rachel M. Gittelman, Michael Dannemann, Steffi Grote, Rajiv C. McCoy, Heather Norton, Laura B. Scheinfeldt, David A. Merriwether, George Koki, Jonathan S. Friedlaender, Jon Wakefield, Svante Pääbo, Joshua M. Akey

Department of Genome Sciences, University of Washington, Seattle, Washington, USA. Department of Life Sciences and Biotechnology, University of Ferrara, Italy.

Department of Evolutionary Genetics, Max-Planck-Institute for Evolutionary Anthropology, Leipzig, Germany. Department of Anthropology, University of Cincinnati, Clincinnati, OH, USA. Coriell Institute for Medical Research, Camden, NJ, USA. Department of Anthropology, Binghamton University, Binghamton, NY, USA. Institute for Medical Research, Goroka, Eastern Highlands Province, Papua New Guinea. Department of Anthropology, Temple University, Philadelphia PA, USA. Department of Statistics, University of Washington, Seattle, Washington, USA.

*Corresponding author. E-mail: paabolileva.mpg.de (S.P.); akeyi@uw.edu (J.M.A.)

Although Neandertal sequences that persist in the genomes of modern humans have been identified in Eurasians, comparable studies in people whose ancestors hybridized with both Neandertals and Denisovans are lacking. We developed an approach to identify DNA inherited from multiple archaic hominin ancestors and applied it to whole-genome sequences from 1523 geographically diverse individuals, including 35 new Island Melanesian genomes. In aggregate, we recovered 1.34 Gb and 303 Mb of the Neandertal and Denisovan genome, respectively. We leverage these maps of archaic sequence to show that Neandertal admixture occurred multiple times in different non-African populations, characterize genomic regions that are significantly depleted of archaic sequence, and identify signatures of adaptive introgression.

Recipe for a modern human

109,295 single nucleotide changes (SNCs)

7,944 insertions and deletions

Changes in protein coding genes

277 cause fixed amino acid substitutions

affect splice sites

Changes in Non-coding & regulatory sequences

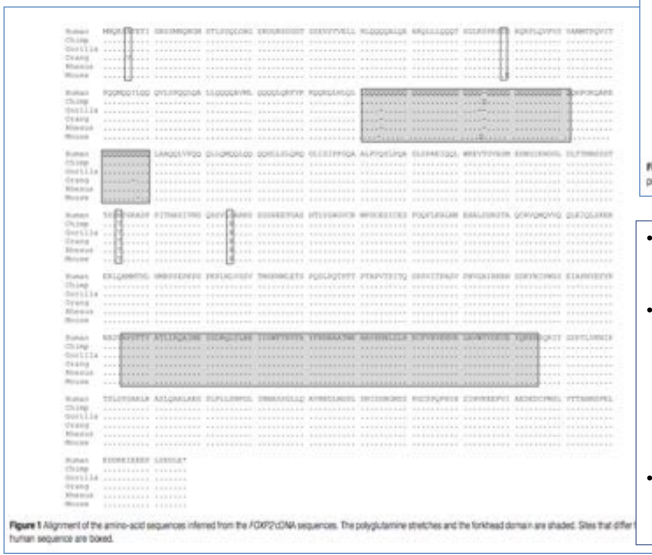
26 affect well-defined motifs inside

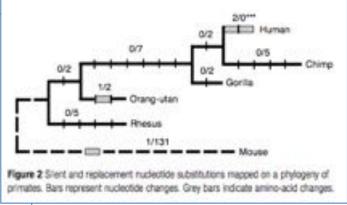
regulatory regions

Enrichment analysis

None	- Giant melanosomes in melanocytes (p-6.77e-6; FWER=0.091;
skin pigmentatio	n
None	 1-3 toe syndactyly (p=1.34288e-05; FWER=0.538; FDR=0.0887928) 1-5 toe syndactyly (p=1.34288e-05; FWER=0.538; FDR=0.0887928) Aplasia/Hypoplasia of the distal phalanx of the thumb (p=1.34288e-05; FWER=0.538; FDR=0.0887928) Bifid or hypoplastic epiglottis (p=1.34288e-05; FWER=0.538; FDR=0.0887928) Central polydactyly (feet) (p=1.34288e-05; FWER=0.538; FDR=0.0887928)
rphologies (lim	b length, digit development)
	FDR=0.0887928) - Dysplastic distal thumb phalanges with a central hole (p=1.34288e-05;
ies of the laryn	x and the epiglottis FWER-0.538;
	- Laryngeal cleft (p=1.34288e-05; FWER=0.538; FDR=0.0887928) - Midline facial capillary hemangioma (p=1.34288e-05; FWER=0.538; FDR=0.0887928)
	 Preductal coarctation of the aorta (p=1.34288e-05; FWER=0.538; FDR=0.0887928) Radial head subluxation (p=1.34288e-05; FWER=0.538; FDR=0.0887928) Short distal phalanx of the thumb (p=1.34288e-05; FWER=0.538; FDR=0.0887928)
	rphologies (lim

FOXP2 Analysis





- Mutations of FOXP2 cause a severe speech and language disorder in people
- Versions of FOXP2 exist in similar forms in distantly related vertebrates; functional studies of the gene in mice and in songbirds indicate that it is important for modulating plasticity of neural circuits.
- Outside the brain FOXP2 has also been implicated in development of other tissues such as the lung and gut.

Molecular evolution of FOXP2, a gene involved in speech and language

Enard et al (2002) Nature. doi:10.1038/nature01025



Part II: Modern Humans

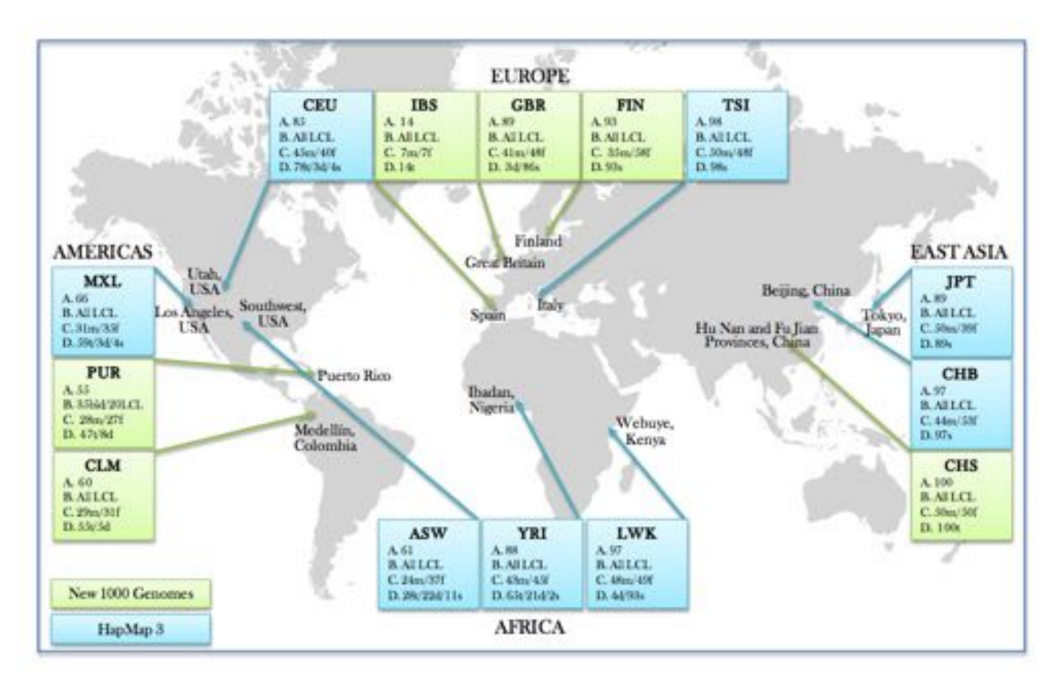


An integrated map of genetic variation from 1,092 human genomes

The 1000 Genomes Project Consortium*

By characterizing the geographic and functional spectrum of human genetic variation, the 1000 Genomes Project aims to build a resource to help to understand the genetic contribution to disease. Here we describe the genomes of 1,092 individuals from 14 populations, constructed using a combination of low-coverage whole-genome and exome sequencing. By developing methods to integrate information across several algorithms and diverse data sources, we provide a validated haplotype map of 38 million single nucleotide polymorphisms, 1.4 million short insertions and deletions, and more than 14,000 larger deletions. We show that individuals from different populations carry different profiles of rare and common variants, and that low-frequency variants show substantial geographic differentiation, which is further increased by the action of purifying selection. We show that evolutionary conservation and coding consequence are key determinants of the strength of purifying selection, that rare-variant load varies substantially across biological pathways, and that each individual contains hundreds of rare non-coding variants at conserved sites, such as motif-disrupting changes in transcription-factor-binding sites. This resource, which captures up to 98% of accessible single nucleotide polymorphisms at a frequency of 1% in related populations, enables analysis of common and low-frequency variants in individuals from diverse, including admixed, populations.

1000 Genomes Populations



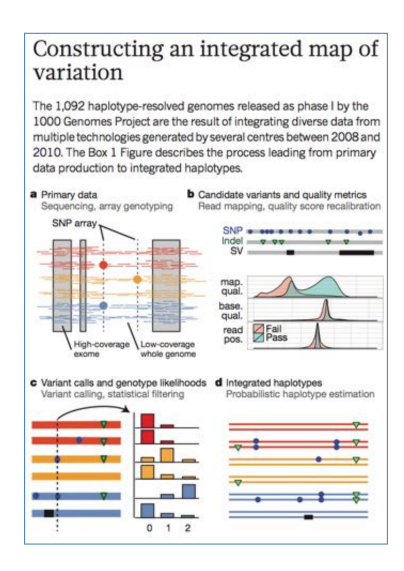
1000 Genomes Populations

Population	DNA sequenced from blood	Offspring Samples from Trios Available	Pilot Samples		Phase 1 Samples	Final Phas Discovery Sample	-	Final Release Sample		Total	B
Chinese Dai in Xishuanghanna, China(CDX)	mi	300		0	. 0		99		93		- 99
Han Chinese in Bejing, China (CHB)	.000	90	. 9	4	97		100		800		106
Japanese in Tokyo, Japan (JPT)	800	90		H	80		104		504		305
Kinh in Ho Chi Minh City, Vietnam (KHV)	369	306	2.9	0	0		101		90		301
Southern Han Chinese, China (CHS)	ine.	988	- 91	0	100		106		105		112
Total East Asian Ancestry (EAS)			185		284	515		594		523	
Bengali in Bangladesh (BEB)	80	968	126	ė	6		96		86		86
Gujarati Indian in Houston,TX (GIH)	100	398	100	0	0		106		103		106
Indian Teluga in the UK (ITU)	yes	969		0	0		105		107		100
Punjabi in Labore Pakistan (PJL)	2994	500	100	0	0		96		96		- 96
Sri Lankan Tamil in the UK (STU)	yes	244		0	0		103		1002		303
Total South Asian Ancestry (SAS)						494		400		494	
African Ancestry in Southwest US (ASW)	100	. 900		0	61		66		62		- 66
African Caribbean in Burbados (ACB)	705	508	33	0	0		96		96		96
Esan in Nigeria (ESN)	200	308	- 39	0	0		99		99		95
Gambian in Western Division, The Gambia (GWD)	100	bes.	10	0	0		113		113		113
Luhya in Webuye, Kenya (LWK)	ine .	300	10	0	W7		301		90		116
Mende in Sierra Leone (MSL)	100	366	- 1	0			85		85		83
Yoruba in Ibadan, Nigeria (YRI)	907	509	10	6	88		109		108		116
Total African Ancestry (AFR)			208		246	649		662		691	
British in England and Scotland (GBR)		988		o			12		90		94
Finnish in Finland (FIN)	769	200	56	0	93		99		99		300
Iberian populations in Spain (IBS)	800	396		0	14		107		100		103
Toscani in Italy (TSI)	966	80		6	56		108		1007		110
Utah residents with Northern and Western European	and .	500		4	85		99		98		303
ancestry (CEU) Total European Ancestry (EUR)	10010		160		379	508		.540	-	514	-
Colombian in Modellin, Colombia (CLM)	966	344		0	60		94		94		. 90
Mexican Ancestry in Los Angeles, California (MXL)	940	568	- 1	U.	66		67		85		86
Peruvian in Lima, Peru (PEL) Puerto Rican in Puerto Rico (PUR)	yes	368	- 3	o o	95		105		104		305
Total Americas Ancestry (AMR)	760	700			181	362	140	341	-	305	-
Trees Various Various Avenues Avenues					1000					-	
Total			A15		3992	200		2904		2817	

26 populations from 5 major population groups

1000 Genomes: Human Mutation Rate

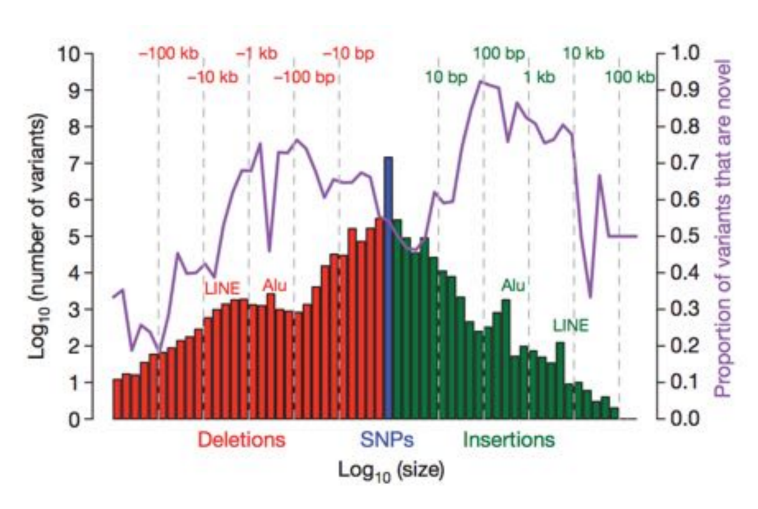
- Phase I Release
 - 1092 individuals from 14 populations
 - Combination of low coverage WGS, deep coverage WES, and SNP genotype data
- Overall SNP rate between any two people is ~1/1200bp to ~1/1300
 - ~3M SNPs between me and you (.1%)
 - ~30M SNPs between human to Chimpanzees (1%)
- De novo mutation rate ~1/100,000,000
 - ~100 de novo mutations from generation to generation
 - ~I-2 de novo mutations within the protein coding genes



An integrated map of genetic variation from 1,092 human genomes

1000 genomes project (2012) Nature. doi:10.1038/nature11632

Human Mutation Types



- Mutations follows a "log-normal" frequency distribution
 - Most mutations are SNPs followed by small indels followed by larger events

A map of human genome variation from population-scale sequencing 1000 genomes project (2010) *Nature*. doi:10.1038/nature09534

A Systematic Survey of Loss-of-Function Variants in Human Protein-Coding Genes

Daniel G. MacArthur, 1,2* Suganthi Balasubramanian, 3,4 Adam Frankish, 1 Ni Huang, 1
James Morris, 1 Klaudia Walter, 1 Luke Jostins, 1 Lukas Habegger, 3,4 Joseph K. Pickrell, 5
Stephen B. Montgomery, 6,7 Cornelis A. Albers, 1,8 Zhengdong D. Zhang, 9 Donald F. Conrad, 10
Gerton Lunter, 11 Hancheng Zheng, 12 Qasim Ayub, 1 Mark A. DePristo, 13 Eric Banks, 13
Min Hu, 1 Robert E. Handsaker, 13,14 Jeffrey A. Rosenfeld, 15 Menachem Fromer, 13 Mike Jin, 3
Xinmeng Jasmine Mu, 3,4 Ekta Khurana, 3,4 Kai Ye, 16 Mike Kay, 1 Gary Ian Saunders, 1
Marie-Marthe Suner, 1 Toby Hunt, 1 If H. A. Barnes, 1 Clara Amid, 1,17 Denise R. Carvalho-Silva, 1
Alexandra H. Bignell, 1 Catherine Snow, 1 Bryndis Yngvadottir, 1 Suzannah Bumpstead, 1
David N. Cooper, 18 Yali Xue, 1 Irene Gallego Romero, 1,5 1000 Genomes Project Consortium, 1
Jun Wang, 12 Yingrui Li, 12 Richard A. Gibbs, 19 Steven A. McCarroll, 13,14
Emmanouil T. Dermitzakis, 7 Jonathan K. Pritchard, 5,20 Jeffrey C. Barrett, 1 Jennifer Harrow, 1
Matthew E. Hurles, 1 Mark B. Gerstein, 3,4,21 † Chris Tyler-Smith 1†

Genome-sequencing studies indicate that all humans carry many genetic variants predicted to cause loss of function (LoF) of protein-coding genes, suggesting unexpected redundancy in the human genome. Here we apply stringent filters to 2951 putative LoF variants obtained from 185 human genomes to determine their true prevalence and properties. We estimate that human genomes typically contain ~100 genuine LoF variants with ~20 genes completely inactivated. We identify rare and likely deleterious LoF alleles, including 26 known and 21 predicted severe disease—causing variants, as well as common LoF variants in nonessential genes. We describe functional and evolutionary differences between LoF-tolerant and recessive disease genes and a method for using these differences to prioritize candidate genes found in clinical sequencing studies.

(2012) Science. doi: 10.1126/science.1215040

Homozygous LoF Mutations

LETTER

doi:10.1038/nature22034

Human knockouts and phenotypic analysis in a cohort with a high rate of consanguinity

Danish Saleheen^{1,7}*, Pradeep Natarajan^{3,6}*, Irina M. Armean^{4,5}, Wei Zhao³, Asif Rasheed², Surneet A. Khetarpal⁶, Hong-Hee Won⁷, Konrad J. Karczewski^{4,5}, Anne H. O'Donnell-Luria^{4,5,8}, Kaitlin E. Samocha^{4,5}, Benjamin Weisburd^{4,5}, Namrata Gupta⁴, Mozzam Zaidi², Maria Samuel², Atif Imran², Shahid Abbas⁹, Faisal Majeed², Madiha Ishaq², Saba Akhtar², Kevin Trindade⁶, Megan Mucksavage⁶, Nadeem Qamar¹⁰, Khan Shah Zaman¹⁰, Zia Yaqoob¹⁰, Tahir Saghir¹⁰, Syed Nadeem Hasan Rizvi²⁰, Anis Memon¹⁰, Nadeem Hayyat Mallick¹¹, Mohammad Ishaq¹², Syed Zahed Rasheed¹², Fazal-ur-Rehman Memon¹³, Khalid Mahmood¹⁴, Naveeduddin Ahmed¹⁵, Ron Do^{16,13}, Ronald M. Krauss¹⁸, Daniel G. MacArthur^{4,3}, Stacey Gabriel⁴, Eric S. Lander⁴, Mark J. Daly^{4,5}, Philippe Frossard²8, John Danesh^{19,20}8, Daniel J. Rader^{6,21}8 & Sekar Kathiresan^{3,4}8

A major goal of biomedicine is to understand the function of every across 14,345 autosomal genes were annotated as pLoF mutations (that gene in the human genome1. Loss-of-function mutations can disrupt both copies of a given gene in humans and phenotypic analysis of such 'human knockouts' can provide insight into gene function. Consanguineous unions are more likely to result in offspring carrying homozygous loss-of-function mutations. In Pakistan, consanguinity rates are notably high2. Here we sequence the proteincoding regions of 10,503 adult participants in the Pakistan Risk of Myocardial Infarction Study (PROMIS), designed to understand the determinants of cardiometabolic diseases in individuals from South Asia3. We identified individuals carrying homozygous predicted loss-of-function (pLoF) mutations, and performed phenotypic analysis involving more than 200 biochemical and disease traits. We enumerated 49,138 rare (<1% minor allele frequency) pLoF mutations. These pLoF mutations are estimated to knock out 1,317 genes, each in at least one participant. Homozygosity for pLoF mutations at PLA2G7 was associated with absent enzymatic activity of soluble lipoprotein-associated phospholipase A2; at CYP2F1, with higher plasma interleukin-8 concentrations; at TREH, with lower concentrations of apoB-containing lipoprotein subfractions; at either A3GALT2 or NRG4, with markedly reduced plasma insulin C-peptide concentrations; and at SLC9A3R1, with mediators of calcium and phosphate signalling. Heterozygous deficiency of APOC3 has been shown to protect against coronary heart disease (5); we identified APOC3 homozygous pLoF carriers in our cohort. We recruited these human knockouts and challenged them with an oral fat load. Compared with family members lacking the mutation, individuals with APOC3 knocked out displayed marked blunting of the usual post-prandial rise in plasma triglycerides. Overall, these observations provide a roadmap for a 'human knockout project', a systematic effort to understand the phenotypic consequences of complete disruption of genes in humans.

Across all participants (Table 1), exome sequencing yielded 1,639,223 exonic and splice-site sequence variants in 19,026 autosomal genes that passed initial quality control metrics. Of these, 57,137 mutations

is, nonsense, frameshift, or canonical splice-site mutations predicted to inactivate a gene). To increase the probability that mutations are correctly annotated as pLoF by automated algorithms, we removed nonsense and frameshift mutations occurring within the last 5% of the transcript and within exons flanked by non-canonical splice sites, splice-site mutations at small (<15 bp) introns, at non-canonical splice sites, and where the purported pLoF allele is observed across primates. Common pLoF alleles are less likely to exert strong functional effects as they are less constrained by purifying selection; thus, we define pLoF mutations in the rest of the manuscript as variants with a minor allele frequency (MAF) of <1% and passing the aforementioned bioinformatic filters. Applying these criteria, we generated a set of 49,138 pLoF mutations across 13,074 autosomal genes. The site-frequency spectrum for these pLoF mutations revealed that the majority was seen only in one or a few individuals (Extended Data Fig. 1).

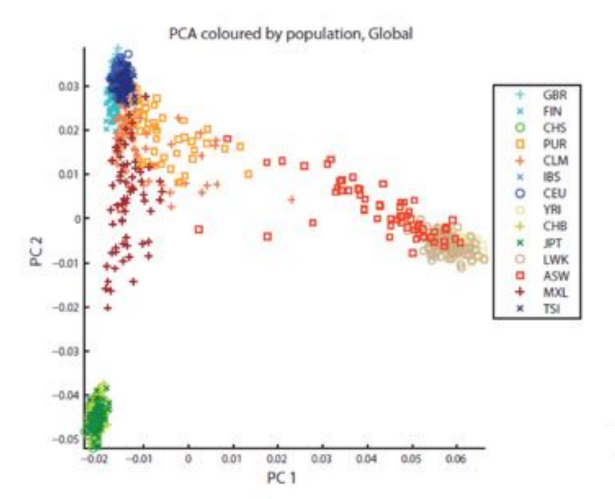
Across all 10,503 PROMIS participants, both copies of 1,317 distinct genes were predicted to be inactivated owing to pLoF mutations. A full listing of all 1,317 genes knocked out, the number of knockout participants for each gene, and the specific pLoF mutation(s) are provided in Supplementary Table 1. 891 (67.7%) of the genes were knocked out only in one participant (Fig. 1a). Nearly 1 in 5 of the participants that were sequenced (1,843 individuals, 17.5%) had at least one gene knocked out by a homozygous pLoF mutation. 1,504 of these 1,843 individuals (81.6%) were homozygous pLoF carriers for just one gene, but the minority of participants had more than one gene knocked out and one participant had six genes with homozygous pLoF genotypes.

We compared the coefficient of inbreeding (F coefficient) in PROMIS participants with that of 15,249 individuals from outbred populations of European or African American ancestry. The F coefficient estimates the excess homozygosity compared with an outbred ancestor. PROMIS participants had a fourfold higher median inbreeding coefficient compared to outbred populations (0.016 versus 0.0041; $P < 2 \times 10^{-16}$) (Fig. 1b). Additionally, those in PROMIS who reported that their parents were closely related had even higher median inbreeding coefficients than

- Homozygous LoF mutations are rare in most people, but enriched in people born from consanguineous relationships
- Sequence the exomes of many such people, find their homozygous LoFs, relate to 200 biochemical or disease traits
- A "natural" experiment to understand what genes do: people with both copies of APOC3 disabled can clear fat from their bloodstream much faster than others, suggests we should develop compounds to prevent heart attacks

(2017) Nature. doi:10.1038/nature22034

Variation across populations



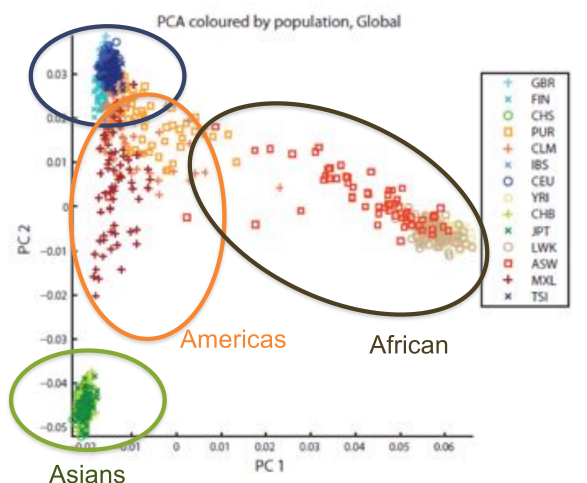
LEVEL	POP_PAIR	# of Highly differentiated SNPs	% in transcribed regions*		
AFR	ASW-LWK	258			
AFR	LWK-YRI	251	50.2		
AFR	ASW-YRI	213	45.8		
ASN	CHS-JPT	275	48.1		
ASN	CHB-JPT	176	43.7		
ASN	CHB-CHS	79	38.7		
EUR	FIN-TSI	343	42.6		
EUR	CEU-FIN	201	40.7		
EUR	FIN-GBR	197	43.2		
EUR	GBR-TSI	100	38.9		
EUR	CEU-TSI	57	53.8		
EUR	CEU-GBR	17	14.3		
CON	AFR-EUR	348	52.2		
CON	AFR-ASN	317	52.6		
CON	ASN-EUR	190	53.4		

Table S12A Summary of sites showing high levels of population differentiation

- Not a single variant 100% unique to a given population
- 17% of low-frequency variants (.5-5% pop. freq) observed in a single ancestry group
- 50% of rare variants (<.5%) observed in a single population

Variation across populations

Europeans

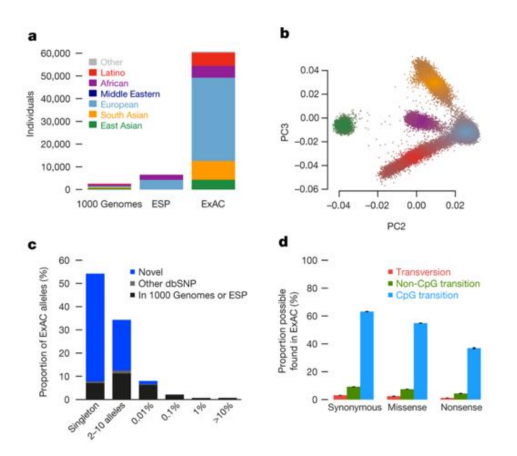


LEVEL	POP_PAIR	# of Highly differentiated SNPs	% in transcribed regions*		
AFR	ASW-LWK	258	46.8		
AFR	LWK-YRI	251	50.2		
AFR	ASW-YRI	213	45.8		
ASN	CHS-JPT	275	48.1		
ASN	CHB-JPT	176	43.7		
ASN	CHB-CHS	79	38.7		
EUR	FIN-TSI	343	42.6		
EUR	CEU-FIN	201	40.7		
EUR	FIN-GBR	197	43.2		
EUR	GBR-TSI	100	38.9		
EUR	CEU-TSI	57	53.8		
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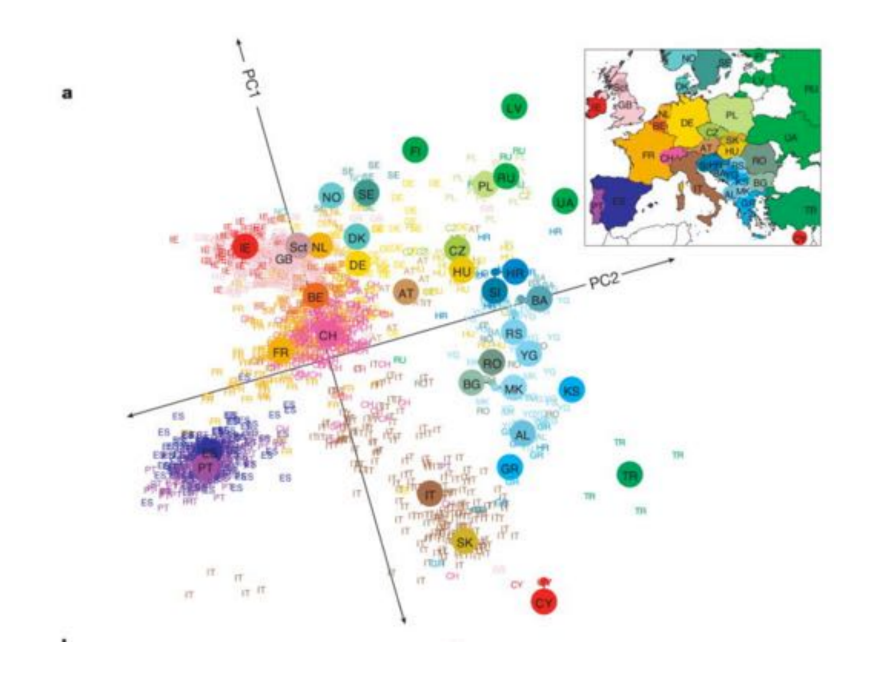
ExAC: Exome Aggregation Consortium



- The aggregation and analysis of highquality exome (protein-coding region) DNA sequence data for 60,706 individuals
- This catalogue of human genetic diversity contains an average of one variant every eight bases of the exome
- We have used this catalogue to calculate objective metrics of pathogenicity for sequence variants, and to identify genes subject to strong selection against various classes of mutation; identifying 3,230 genes with near-complete depletion of predicted protein-truncating

Analysis of protein-coding genetic variation in 60,706 humans

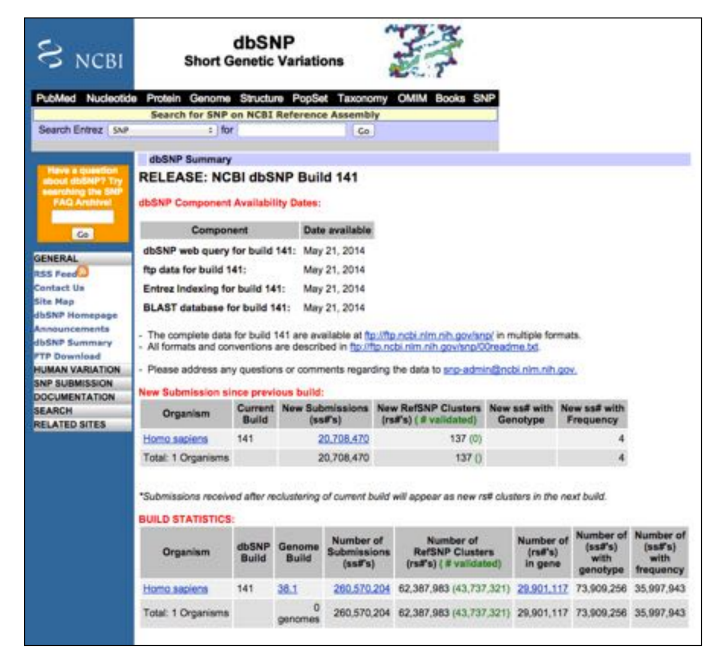
Lek et al (2016) Nature. doi:10.1038/nature19057



Genes mirror geography within Europe

Novembre et al (2008) Nature. doi: 10.1038/nature07331

dbSNP



- Periodic release of databases of known variants and their population frequencies
- Generally assumed to be non-disease related
- However, as catalog grows, almost certainly to contain some medically relevant SNPs.