## From discovery to technology explosion 1868: Discovery of DNA

(pyrosequencing)

(ligation sequencing)

- 1953: Watson and Crick propose double helix structure ٠
- 1977: Sanger sequencing ٠
- 1985: PCR
- 2000: Working draft human genome announced (Sanger method)
- 2005: 454 sequencer launch
- 2006: Genome Analyzer launched ٠
- 2007: SOLiD launched
- 2009: Whole human genome no longer merits Nature/Science ٠ paper
- 2010: "third-gen" systems



\$ human Genome

\$3 billion

\$2-3 million

\$250k \$50k \$20k <\$1k





Applied Biosystems ABI 3730XL 1 Mb / day



Roche / 454 Genome Sequencer FLX 100 Mb / run



Illumina / Solexa Genetic Analyzer 2000 Mb / run



Applied Biosystems SOLiD 3000 Mb / run



## 







Sequencer	454 GS FLX	HiSeq 2000	SOLiDv4	Sanger 3730xl
Sequencing mechanism	Pyrosequencing	Sequencing by synthesis	Ligation and two-base coding	Dideoxy chain termination
Read length	700 bp	50SE, 50PE, 101PE	50 + 35 bp or 50 + 50 bp	400~900 bp
Accuracy	99.9%*	98%, (100PE)	99.94% *raw data	99.999%
Reads	1 M	3 G	$1200 \sim 1400 \text{ M}$	—
Output data/run	0.7 Gb	600 Gb	120 Gb	1.9~84 Kb
Гime/run	24 Hours	3~10 Days	7 Days for SE 14 Days for PE	20 Mins~3 Hours
Advantage	Read length, fast	High throughput	Accuracy	High quality, long read length
Disadvantage	Error rate with polybase more than 6, high cost, low throughput	Short read assembly	Short read assembly	High cost low throughput

# **Oxford Nanopore**



# **DNA degradation**

Mechanical damage during tissue homogenization.

Wrong pH and ionic strength of extraction buffer.

Incomplete removal / contamination with nucleases.

**Phenol**: too old, or inappropriately buffered (**pH 7.8 – 8.0**); incomplete removal.

Wrong pH of **DNA solvent** (acidic water). *Recommended: 1:10 TE for short-term storage, or 1xTE for long-term storage.* 

Vigorous pipetting (wide-bore pipet tips).

**Vortexing** of DNA in high concentrations.

Too many **freeze-thaw** cycles (*we tested 5, still Ok*).

Debatable: sequence-dependent



Trade prices are not sourced from all markets

## Genome sequencing

Two strategies

- Whole genome shotgun (bottom-top)
- Clone-by-clone (top-bottom)



## Sequencing without a limit?

• A rapid progress in next generation sequencing technologies promises to provide complete (reference) DNA sequences



- The bottleneck:
  - NOT the sequencing capacity
  - BUT the ability to assemble many short reads with prevalence of repeated DNA (and polyploidy)

# Genome sequencing

**GenBank 1982 Los Alamos Sequence Database** 





Walter Goad

# **Frederick Sanger**

- 1958 Nobel prize insuline structure
- 1975 Dideoxy sequencing method
- 1977 Φ-X174 (5,368 bp) sequence
- 1980 second Nobel prize
- λ phage sequence shotgun method (48,502 bp)



# Genome sequencing

- **1986** Leroy Hood: automatic sequencing machine
- 1986 Human Genome Initiative



Leroy Hood



# Genome sequencing

• **1995** John Craig Venter first bacterial genome



John Craig Venter

## **Craig Venter**

## Global Ocean Sampling Expedition



Synthetic genomics

Human Longevity Inc

http://www.youtube.com/watch?v=J0rDFbr hjtl

# Which applications are labs performing?



2010 Human genome reference

## 2010 Human genome reference



## 23andme (30% GSK)

welcome health ancestry research how it works buy help Q

HOME MY RESULTS

**Health Overview** 

Inherited Thrombophilia Variant Absent, Typical Risk

23andMe

**Genetic Risk Factors** 

Alzheimer's Disease

Factor XI Deficiency

Parkinson's Disease

**Bitter Taste Perception** 

REPORT

Traits

REPORT

Eye Color

FAMILY & FRIENDS

RESULT

RESULT

Unlike to Taste

Likely Brown

Variant Present, Higher Risk

Variant Absent, Typical Risk

Variant Absent, Typical Risk

See All Genetic Risk Factor Reports

**RESEARCH & COMMUNITY** 

**Inherited Conditions** 

Bloom Syndrome

Sickle Cell Anemia

Tay-Sachs Disease

**Drug Response** 

Clopidogrel (Plavix®) Efficacy

Proton Pump Inhibitor (PPI) M

REPORT

Se

Cystic Fibrosis

REPORT

#### <u>Anne Wojcicki</u> CEO - manželka spoluzakladatele Google Sergey Mikhaylovich Brin

Learn how your DNA may affect your health.

Our genes are a part of who we are, so naturally they impact our health. By knowing more about your DNA, you may be able to take steps towards living a healthier life.

Keep in mind that many conditions and traits are influenced by multiple factors. Our reports are intended for informational purposes only and do not diagnose disease or illness.

#### • Plan for the future.

23andMe

Learn if you are a carrier for certain inherited conditions, so you and your family can be prepared.

#### • Stay one step ahead.

Find out if you have certain genetic risk factors, so you can make better lifestyle choices and appropriately monitor your health.

#### Engage in your health care.

Understand how your DNA may affect your health and response to



#### HOME MY RESULTS FAMILY & FRIENDS RESEARCH & COMMUNITY

#### MATERNAL LINE: H1

Overview

History

Community

Locations of haplogroup H1 before the widespread migrations of the past few hundred years.

Haplogroup Tree



Haplogroup H1 is widespread in Europe, especially the western part of the continent. It originated about 13,000 years ago, not long after the Ice Age ended.

Maternal haplogroups are families of mitochondrial DNA types that all trace back to a single mutation at a specific place and time. By looking at the geographic distribution of mtDNA types, we learn how our ancient female ancestors migrated throughout the world.

Haplogroup: H1, a subgroup of H Age: 13,000 years Region: Europe, Near East, Central Asia, Northwestern Africa Example Populations: Spanish,Berbers,Lebanese Highlight: H1 appears to have been common in Doggerland, an ancient land now flooded by the North Sea.

HOME

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#### PATERNAL LINE: 11\*



Haplogroups of You and Your Connections

Roman Hobza

Haplogroup I1 can be found at levels of 10% and higher in many parts of Europe, due to its expansion with men who migrated northward after the end of the Ice Age about 12,000 years ago. It reaches its highest levels in Denmark and the southern parts of Sweden and Norway.

#### Haplogroups of Example Profiles

11\*

#### SHOW RESULTS FOR Roman Hobza 🔻

#### SEE NEW AND RECENTLY UPDATED REPORTS »

These reports provide information about your possible risk for developing certain health conditions based on genetics. Environmental and lifestyle factors also often play a large role in your risk for developing these conditions.

#### Elevated Risk 🕜

CONFIDENCE	YOUR RISK	AVG. RISK	COMPARED TO AVERAGE
****	41.8%	12.3%	3.39x 📕
****	35.7%	22.8%	1.57x 💻
****	4.0%	2.9%	1.38x <b>:</b>
****	2.5%	2.0%	1.25x 🚦
****	2.2%	0.7%	2.90x :
****	0.43%	0.36%	1.21x ¦
****	0.28%	0.23%	1.22x ¦
***	0.11%	0.08%	1.43x ¦
***	0.08%	0.07%	1.24x ¦
	**** **** **** **** **** **** ****	**** 41.8%   **** 35.7%   **** 4.0%   **** 2.5%   **** 2.2%   **** 0.43%   **** 0.28%   **** 0.11%	****   41.8%   12.3%     ****   35.7%   22.8%     ****   4.0%   2.9%     ****   2.5%   2.0%     ****   2.2%   0.7%     ****   0.43%   0.36%     ****   0.28%   0.23%

Show information for Roman Hobza

assuming European
ethnicity and an age range of 0-79

#### Roman Hobza 41.8 out of 100

Average

develop Venous

ages of 0 and 79.

12.3 out of 100 men of European ethnicity will

Thromboembolism between the

men of European ethnicity who share Roman Hobza's genotype will develop Venous Thromboembolism between the ages of 0 and 79.

#### What does the Odds Calculator show me?

Use the ethnicity and age range selectors above to see the estimated incidence of Venous Thromboembolism due to genetics for men with **Roman Hobza**'s genotype. The 23andMe Odds Calculator assumes that a person is free of the condition at the lower age in the range. You can use the name selector above to see the estimated incidence of Venous Thromboembolism for the genotypes of other people in your account.

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The 23andMe Odds Calculator only takes into account effects of markers with known associations that are also on our genotyping chip. Keep in mind that aside from genetics, environment and lifestyle may also contribute to one's risk for Venous Thromboembolism.

#### 55 % Attributable to Genetics

#### Understanding Your Results

The heritability of venous thromboembolism is estimated to be 55%. This means that genetics (including unknown factors and known ones such as the SNPs we describe here) and environment play nearly equal roles in this condition. There are a number of environmental factors of various strengths that contribute to venous thromboembolism. Strong risk factors include hip or leg fractures, hip or knee replacement, major surgery or trauma, and spinal cord injury or surgery. Moderate risk factors include arthroscopic knee surgery, having central venous lines, congestive heart or respiratory failure, hormone replacement or oral contraceptive use, cancer, pregnancy, paralytic stroke, previous venous thromboembolism, and thrombophilia. Weak risk factors include bed rest for more than three days, immobility due to sitting (such as a long car or plane trip), specific types of chemotherapy, increasing age, laparoscopic surgery, obesity, and varicose veins. (sources)

#### What You Can Do

Assuming the ethnicity setting above is correct, your test results indicate you are at increased risk for venous thromboembolism based on genetics. Note that family history and non-genetic factors can also influence your risk for venous thromboembolism. Below are some steps you can take to reduce your risk.

#### Gene or region: F5 SNP: rs6025

	SNP used	Genotype	Adjusted Odds Ratio*
Roman Hobza	rs6025	CT	European: 4.69
* Odds ratios are report	ted for all available ethi	nicities.	

Factor V is the last clotting factor in the pathway before the activation step that turns prothrombin into thrombin. Clotting is usually kept from spiraling out of control by a feedback loop, similar to the way a thermostat operates. Once enough thrombin has been activated, it binds to a protein called "protein C." Protein C then inactivates factor V, thus cutting off activation of prothrombin into thrombin.

The SNP in the F5 gene causes a change in the protein sequence of factor V that prevents protein C from inactivating it. Since this version of factor V can still participate in the activation of thrombin, a situation results in which thrombin can be turned on but cannot be turned off. Once the clotting cascade is set off (whether appropriately or not), the riskier version of the SNP makes it more difficult to shut it off.

The riskiness of the T version of this SNP is further increased for women who also take hormonal birth control.

(The riskier version of this gene is also sometimes called Factor V Leiden, after the city in the Netherlands where this SNP and its effects on factor V's role in clotting were first discovered.)

The studies whose data we report as applicable to those of "European" ancestry confirmed the association between this SNP and VTE in samples from the Netherlands, Sweden, the United Kingdom, Brazil, Italy, and France.

African and Asian populations appear to have only one version of the SNP, meaning that association studies are very difficult to perform.

#### Citations

Rosendaal et al. (1995). "High risk of thrombosis in patients homozygous for factor V Leiden (activated protein C resistance)." Blood 85(6):1504-8.

Smith et al. (2007). "Association of genetic variations with nonfatal venous thrombosis in postmenopausal women." JAMA 297(5):489-98.

Emmerich et al. (2001). "Combined effect of factor V Leiden and prothrombin 20210A on the risk of venous thromboembolism--pooled analysis of 8 case-control studies including 2310 cases and 3204 controls. Study Group for Pooled-Analysis in Venous Thromboembolism." Thromb Haemost 86(3):809-16.

Bertina et al. (1994). "Mutation in blood coagulation factor V associated with resistance to activated protein C." Nature 369(6475):64-7.

Lane et al. (2000). "Role of hemostatic gene polymorphisms in venous and arterial thrombotic disease." Blood 95(5):1517-32.

Gene or region: F2 SNP: i3002432

### Decreased Risk 🖉

NAME	CONFIDENCE	YOUR RISK	AVG. RISK	COMPARED TO AVERAGE	
Type 2 Diabetes	****	17.7%	25.7%	0.69x 💻	
Alzheimer's Disease	****	4.3%	7.2%	0.60x	
Rheumatoid Arthritis	****	1.6%	2.4%	0.68x 🚦	
Parkinson's Disease	****	1.2%	1.6%	0.73x <b>:</b>	
Age-related Macular Degeneration	****	0.92%	6.55%	0.14x 🔒	
Crohn's Disease	****	0.31%	0.53%	0.58x ¦	
Multiple Sclerosis	****	0.24%	0.34%	0.69x ¦	
Type 1 Diabetes	****	0.12%	1.02%	0.12x ¦	
Celiac Disease	****	0.05%	0.12%	0.44x ¦	

BRCA Cancer Mutations (Selected)	****	Variant Absent
Beta Thalassemia	****	Variant Absent
Bloom's Syndrome	****	Variant Absent
Canavan Disease	****	Variant Absent
Congenital Disorder of Glycosylation Type 1a (PMM2-CDG)	****	Variant Absent
Connexin 26-Related Sensorineural Hearing Loss	****	Variant Absent
Cystic Fibrosis	****	Variant Absent
D-Bifunctional Protein Deficiency	****	Variant Absent
DPD Deficiency	****	Variant Absent
Dihydrolipoamide Dehydrogenase Deficiency	****	Variant Absent
Factor XI Deficiency	****	Variant Absent
Familial Dysautonomia	****	Variant Absent
Familial Hypercholesterolemia Type B	****	Variant Absent
Familial Hyperinsulinism (ABCC8-related)	****	Variant Absent
Familial Mediterranean Fever	****	Variant Absent
Fanconi Anemia (FANCC-related)	****	Variant Absent
G6PD Deficiency	****	Variant Absent

Reading Ability	***	Typical Nonword Reading Score
Response to Diet	***	See Report
Response to Exercise	***	See Report
Sex Hormone Regulation	***	See Report
Sweet Taste Preference 🔆	***	See Report
Tooth Development	***	See Report
Tuberculosis Susceptibility	***	See Report
Breast Morphology 🍳 🔆	***	Not Applicable
Menarche 9	***	Not Applicable
Menopause ♀	***	Not Applicable
Eating Behavior	**	Greater tendency to overeat
HIV Progression	**	See Report
Hair Thickness	**	Typical, if European or African
Longevity	**	See Report
Measures of Intelligence	**	Lower Non-Verbal IQ
Memory	**	Typical Episodic Memory
Odor Detection	**	Typical Sensitivity to Sweaty Odor
Pain Sensitivity	**	Increased
Avoidance of Errors	*	See Report

## Genome Sequencer 20 System 454 pyrosequencing (2005)

http://www.454.com





## **DNA** library preparation



# Fragmentace DNA



# Ligace adaptoru



# Vychytání DNA molekul



## denaturace





# emPCR


# Vznik emulze (olej)



## emPCR



## emPCR



# Vychytání kuliček



# Vychytání kuliček



### denaturace



## Sekvenační primer



## Disperze na sklíčko



## Disperze na sklíčko



# Parametry mikroreaktorů



**PicoTiterPlate device** 

## Parametry mikroreaktorů



















### **SOLID** (Sequencing by Oligonucleotide Ligation and Detection) 2-base encoding sequencing (2007)



#### Applied Biosystems

SOLiD<sup>™</sup> System Sequencing by Oligonucleotide Ligation and Detection

# Properties of the Probes Spatial separation among dye, ligation & cleavage sites Cleavage site, 3' Ligation site $3^{3'}$ Fluorescent dye

1,024 Octamer Probes (4<sup>5</sup>) 4 Dyes, 4 dinucleotides, 256 probes per dye N= degenerate bases Z= Universal bases

#### SOLiD Chemistry System 4-color ligation Ligation reaction



#### SOLiD Chemistry System 4-color ligation Ligation reaction



### **SOLiD Chemistry System 4-color ligation De-Phosphorylation**



### SOLiD Chemistry System 4-color ligation Visualization



### SOLiD Chemistry System 4-color ligation Cleavage



### SOLiD Chemistry System 4-color ligation Ligation (2<sup>nd</sup> cycle)



### SOLiD Chemistry System 4-color ligation Visualization (2<sup>nd</sup> cycle)



### SOLiD Chemistry System 4-color ligation Cleavage (2<sup>nd</sup> cycle)



# SOLiD Chemistry System 4-color ligation interrogates every 5<sup>th</sup> base



### SOLiD Chemistry System 4-color ligation Reset



# SOLiD Chemistry System 4-color ligation (1<sup>st</sup> cycle after reset)



# SOLiD Chemistry System 4-color ligation (1<sup>st</sup> cycle after reset)



# SOLiD Chemistry System 4-color ligation (2<sup>nd</sup> Round)



### Sequential rounds of sequencing Multiple cycles per round



#### Paired End two sequences generated Sequential rounds of sequencing Multiple cycles per round




## Solexa (2007)



Randomly fragment genomic DNA and ligate adapters to both ends of the fragments.

Bind single-stranded fragments randomly to the inside surface of the flow cell channels.

Add unlabeled nucleotides and enzyme to initiate solid-phase bridge amplification.



The enzyme incorporates nucleotides to build double-stranded bridges on the solidphase substrate.

Denaturation leaves single-stranded templates anchored to the substrate.

Several million dense clusters of doublestranded DNA are generated in each channel of the flow cell.



First chemistry cycle: to initiate the first sequencing cycle, add all four labeled reversible terminators, primers and DNA polymerase enzyme to the flow cell.

After laser excitation, capture the image of emitted fluorescence from each cluster on the flow cell. Record the identity of the first base for each cluster.

Second chemistry cycle: to initiate the next sequencing cycle, add all four labeled reversible terminators and enzyme to the flow cell.





True Single Molecule Sequencing (tSMS)







#### Single Molecule Real-Time (SMRT)



**Pacific Biosciences** 





#### **Oxford** nanopore





# Další technologie

- Mikroelektroforéza
- Sekvenování na bázi microarray

### CHALLENGES IN GENOME SEQUENCING

*De novo* genome assemblies using only short read data of NGS technologies are generally incomplete and highly fragmented due to

- Large duplications chromosomal approach, BAC-by-BAC sequencing
- High proportion of repetitive DNA challenge!



### **BAC-BY-BAC SEQUENCING**



- Physical map is composed of contigs of overlapping BAC clones
- BAC contigs are landed on the chromosome through markers comprised in the contigs



#### SOLUTIONS FOR THE REPEATS

- Long mate-pair reads > 10 kb
- Long read technologies PacBio, Oxford Nanopore
- Optical mapping
  - Single-molecule mapping of genomic DNA hundreds of kilobases to several megabases in size
  - Creates sequence-motif maps, which provide long-range template for ordering genomic sequences
  - Visualisation of reality "Seeing is Believing"

#### Three enzymatic approaches

 restriction enzymes: sequence-specifically cleave DNA immobilized on a surface



 nicking enzymes: fluorescent labelling of the nicking site in solution (BioNano Genomics - Irys)



 methyltransferase enzymes: labelling with ultra-high density

## **BIONANO GENOME MAPPING ON NANOCHANEL ARRAYS**

