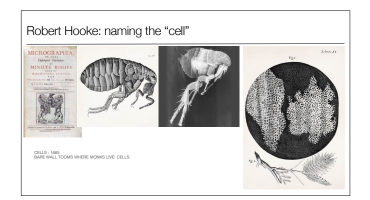
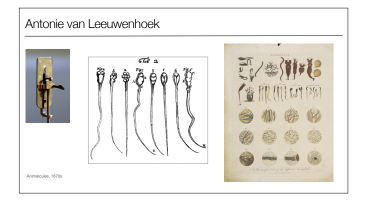


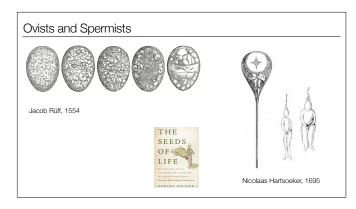
Robert Hooke's drying of cells in a sliver of cork (oak bark)



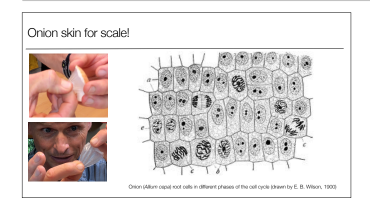
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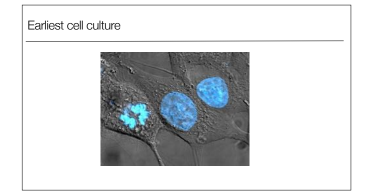
Antonie van Leeuwenhoek



preformationists, ovists, and spearmints. Lazzaro Spalanzani and frogs in pants



Robert Hooke's drying of cells in a sliver of cork (oak bark)



Keeping cells alive outside the body. How to keep dying tissue alive.

## Early cell culture





One stage in the preparation of the rabies vaccine: a rabbit brain on a square of muslin. Pasteur Institute, India, circa 1910. Wellcome Library, London

Pasteur and rabies vaccination. Illustration showing an anti-rabies vaccination being given at the Pasteur Institute in Paris, France. French chemist and microbiologist Louis Pasteur (1822-1895, standing at right) used rabbits to prepare a rabies virus which was milder and had a shorter incubation period than the wild virus. A person who has been bitten by a rabid animal is inoculated with the vaccine, which rapidly stimulates immunity to the wild strain. The first human patient was successfully treated in 1885. This engraving is based on a 1887 painting by Laurent Lucien Gsell (1860-1944). Titled 'La vaccine de la rage', the original is held at the Institute of Bacteriology at the Louis Pasteur University, Strasbourg, France.

Early polio vaccine experiments in monkey central nervous system







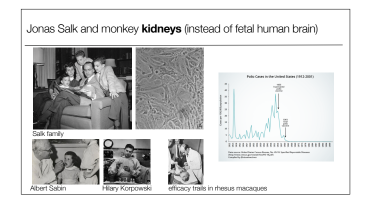


irus] which possess this limited irulence for monkeys by the spinal ite were found to be completely virulent when inoculated into the pinal cord of chimpanzees, producing neither paralysis nor lesions.

& brain tissue from human embryos

Semiannual Report, January 1-June 30, 1954. NFIP, Box 7, Folder 16. AS, WCHHP, UC, Ohi

In 1936, Albert Sabin and Peter Olitsky at the Rockefeller Institute successfully grew poliovirus in a culture of brain tissue from a human embryo. The virus grew guickly, which was promising, but Sabin and Olitsky were concerned about using this as starting material for a vaccine, fearing nervous system damage for vaccine recipients. They tried to grow poliovirus in cultures using tissue that had been taken from other sources, but were unsuccessful.



in the 1950s as the national effort to develop a polio vaccine required the importation of more than 200,000 rhesus monkeys annually for 6 years (Eudey and Mack 1984). Many of these imported NHPs were caught wild in their natural habitat (NAS 1970)

Dr. L. James Lewis, an employee of Dr. Jonas Salk, injects a rhesus monkey with the polio vaccine. At first, he anesthetized the monkey, shaved his leg and then disinfected the skin. He then injected the vaccine into the muscle tissue. The photo was taken in 1955, four days before the release of the evaluation report on the polio vaccine. Photo: Bettmann/ Corbis

In 1936, Albert Sabin and Peter Olitsky at the Rockefeller Institute successfully grew poliovirus in a culture of brain tissue from a human embryo. The virus grew quickly, which was promising, but Sabin and Olitsky were concerned about using this as starting material for a vaccine, fearing nervous system damage for vaccine recipients. They tried to grow poliovirus in cultures using tissue that had been taken from other sources, but were unsuccessful.





Indian rhesus monkeys (M. mulatta) were imported at a rate of 200,000 per year for at least six years and by the tens of thousands for the next 20 years...until the ban by India in 1978.

## Early cell culture: limited growth





#### Cutter Incident 1955





caused 40 000 cases of polio, leaching 200 children with varying degrees of paralysis and killing 10.

Fitzpatrick M. The Cutter Incident: How America's First Polio Vaccine Led to a Growing Vaccine Crisis. J R Soc Med. 2006;99(3):156

Years later, in a suit brought against Cutter, the firm was found not negligent in making its vaccine because it had done its best making a new drug that was complicated to produce.

But it was found financially liable for the calamity it had caused during that spring of 1955.

#### Hayflick Limit: ~ 50 cell doublings....









Over 750 million virus vaccine doses have been produced on WI-38 or similar diploid cell strains. Havflick established international standards for the production of human biologicals in passaged cells, which are still used today by the biotechnology industry

Robert Hooke's drying of cells in a sliver of cork (oak bark)

Long way from primary kidney cell culture to stable cell lines.

1962 – Hayflick entwickelt den ersten menschlichen diploiden Zellstamm WI-38 aus dem Lungengewebe eines drei Monate alten weiblichen Fötus. Diese Zellen werden bis heute in der Herstellung von Impfstoffen eingesetzt [10]. https://www.atsjournals.org/doi/pdf/10.1164/arrd.1963.88.3P2.387

Hayflick führte einen sechs Jahre andauernden Streit mit den nationalen Gesundheitsbehörden um die Rechte an der daraus entwickelten Zelllinie – und gewann. Seither dürfen amerikanische Forscher die Verwertungsrechte für ihre

Entdeckungen behalten, auch wenn deren Forschung durch nationale Mittel finanziert wurde. Ein Kommentar von Hayflick hierzu wurde 2012 in Science veröffentlicht.

WI-38 is a diploid human cell line composed of fibroblasts derived from lung tissue of a 3-month-gestation female fetus. The fetus came from the elective abortion of a Swedish woman in 1962, and was used without her knowledge or permission.

#### Hayflick argued against the use of monkey cells....

A COMPARISON OF PRIMARY MONKEY KIDNEY, HETEROPLOID CELL LINES, AND HUMAN DIPLOID CELL STRAINS FOR HUMAN VIRUS VACCINE PREPARATION<sup>1,2</sup>

LEONARD HAVELICE

It has now been established that no less than serologically distinct serian viruses can be serologically distinct serian viruses can be serologically distinct serian viruses can be series and the series of the se

tection or this contaminating virus in routile vaccine after tests is relatively easy, the rerisk lies with those who work with peissary monkey kidney during vaccine manufacture Indeed, it is within this group that a number of I virus statistics have occurred.

The second similar virus that we shall conside briefly in the vaccolating agent or SV-40 (2). This

¹ From the Wistar Institute of Anatomy and Bielogy, Philadelphis, Pennsylvania.
¹ This work was supported (in part) by U. S. Public Health Service Contract No. TH41-60-187 and by Grant No. CAO1531-01 from the National Cancer Institute, National Institutes of Health, U. S. Public Health Service.

\*A Comparison of Primary Monkey Kidney, Heteroploid Cell Lines, and Human Diploid Cell Strains for Human Virus Vaccine Preparation1,2.\* American Review of Respiratory Disease, 88(3P2), pp. 387–393 uses of a majority of moskey kidacya, has nount he be cooperin for the haranter (4, 5), to recently this view has been demonstrated for the consequence of the consequence of a co-weekers (7) to cause alternations of acernal ann cells in without coells having attributes of cer cells. As if this indictment of moskey are for use in harant view succine production so not sufficient, it is also now well recognized 18-Vu io expalsh of surviving the usual for-

in which to prepare human virus was the property of the prope

primate tissue.

The use of heteroploid cell lines can be quickly rejected on the ground that these cell systems share many of the characteristics of neoplastic cells. The risk of using these heteroploid cell lines has been aptly put by Westwood and associates

Leonard Hayflick's warning about SV40 another viruses.

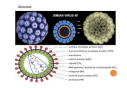
#### Simian vacuolating virus 40. SV40

Some of the polio vaccine administered from 1955–1963 was contaminated with a virus, called simian virus 40 (SV40) a macaque polyomavirus that can induce cancer in rodents.

An estimated 10-30% of polio vaccines administered in the US were contaminated with simian virus 40 (SV40).

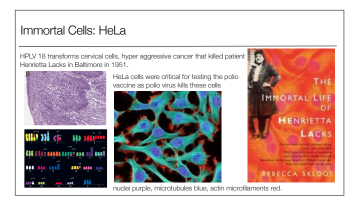
~30 million Americans were exposed to SV40 via contaminated vaccines.

The virus codes a protein known as the T-antigen, which regulates viral replication and inactivates tumor suppressor genes (p53)

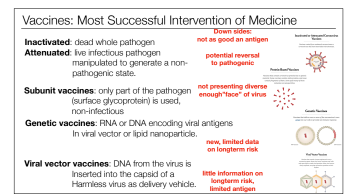


Rollison, D. E. M., and K. V. Shah. 2001. The epidemiology of SV40 infection due to contaminated polio vaccines: relation of the virus to human cancer, p. 561-584. In K. Khalili and G. L. Stoner (ed.), Human polyomaviruses: molecular and clinical perspectives. Wiley-Liss, Inc., New York, N.Y.

SV40: a stowaway passenger in the monkey kidneys.....that inadvertently was injected into ~30 million Americans.

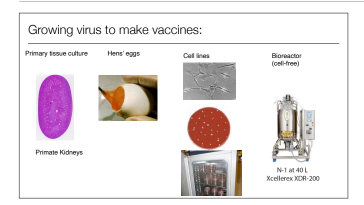


HeLa cells are rapidly dividing cancer cells, and the number of chromosomes varied during cancer formation and cell culture. The current estimate (excluding very tiny fragments) is a "hypertriploid chromosome number (3n+)" which means 76 to 80 total chromosomes (rather than the normal diploid number of 46) with 22–25 clonally abnormal chromosomes, known as "HeLa signature chromosomes."

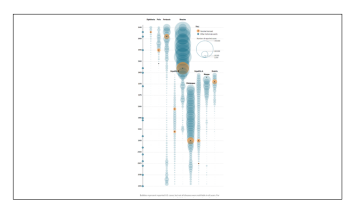


There are different ways of manufacturing vaccines.

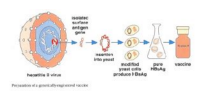
Vaccines can have risks, but more than half a century of studies have shown that overall the benefits of mass-immunization far outweigh the risks to the individuals.



The substrate used for making vaccine contributes to certain risks of the vaccine, e.e. Influenza vaccine made in chicken eggs can cause reactions in people who have egg allergies. Animal or human cell lines each carry risks of disease transmission, plant cells are also used, latest technology uses cell-free reactors to synthesize viral RNA (e.g. Pfizer)



# Hepatitis B subunit vaccine

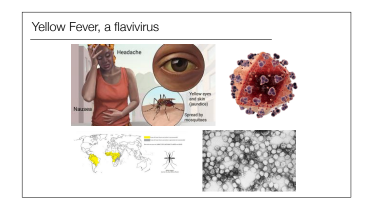


First successful anticancer vaccine

# Your Hepatitis B vaccine was tested for safety in chimpanzees!



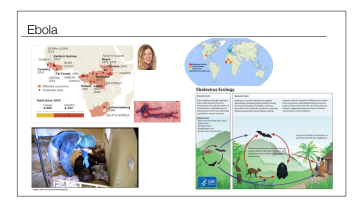
Studies by Alfred Prince and his team at the Vilab in Liberia have paved the way for a Hepatitis B vaccine. The vaccine is now produced in yeast cells.



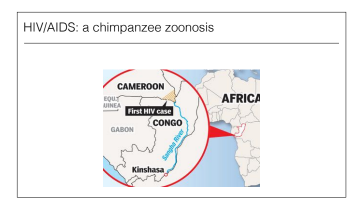
yellow fever is the only flavivirus that can be prevented with a very efficient vaccine.



A recently developed vaccine against ebola is a big hope for many. VSV-EBOV or rVSV-ZEBOV, sold under the brand name Ervebo, is a vaccine based on the vesicular stomatitis virus which was genetically modified to express a surface glycoprotein of Zaire Ebola virus



My friend and colleague was patient zero for the Ebola Ivory Coast outbreak in 1994. She infected herself while helping a veterinarian conduct an autopsy of a dead wild chimpanzee.

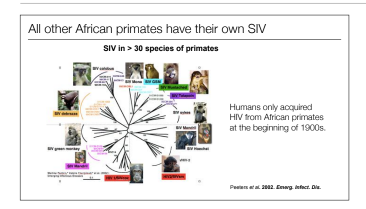


It is now clear that HIV/AIDS emerged as a zoonosis bin Central Africa around the turn of the the 1900s.

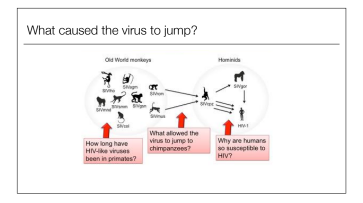


HIV infects T-lymphocytes in the blood stream, ultimately causing AIDS. Terese Winslow created this artwork to give scientists new insight into how HIV infects T-lymphocytes. The virion is shown in the first stage of infection, when the virion attaches to the surface of the T-cell. The molecules involved in this docking process are of particular interest to scientists, so she rendered them accurately according to the most up-to-date scientific information. These molecules include gp120 (the blue 'mushrooms' on the surface of the virus), CD4 (the long red molecules on the cell surface), and chemokine receptors (the groups of blue cylinders on the cell surface).

Again, no depiction of the many complex glycan molecules on both, the virus glycoprotein "mushrooms" or the host cell surface.



Most African non-human primates each have their own versions of HIV, named SIV (simian immunodeficiency virus, a misnomer, as most other African primate species do not get sick).



More than a million years in other African primates. Jump likely aided by bush meat hunting/butchering. The bases for human susceptibility are still being studied.



The convergence of colonial brutality, the first large urban centers (including sex workers), intercontinental medical aid, blood commerce (plasma pheresis businesses in Haiti), and sex tourism and IV drug use formed the perfect storm.

Practice question:

Which factors helped spark the HIV/AIDS pandemic? see above

#### Bush meat trade



Apes are still hunted for their meat throughout tropical Africa, even in the cities, bush meat is valued much more highly than farmed meat.

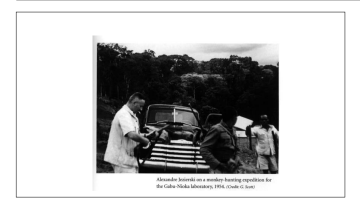


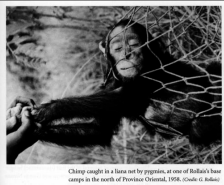
perfect opportunity fro cross-species infections.



Polio vaccine studies in the Belgian Congo used hundreds of wild caught chimpanzees and bonobos for testing the efficacy and safety of vaccine.

These studies could not have cause the HIV1 epidemic which was well underway by the late 1950s.







Two African assistants dismembering a dead chimp in the

# Mass vaccination in Belgian Congo 1959



Mass vaccination in Belgian Congo 1959: suspected by some as possible origin of HIV/ AIDS

#### BUT

clearly not the case rather HIV was already circulating at the time



#### The Alternative hypotheses about HIV origins:

- 1.Natural Transfer: infection by killing and butchering of apes for meat, more hunting in modern times, larger cities and more travel.
- 2. Natural Transfer & syringes (aided by rural clinics with rampant reuse of unsterilized hypodermic needles).
- 3.Oral Polio Vaccine (OPV), vaccine prepared on chimpanzee tissue cultures? infected with SIV and fed to ~1 million Africans in 1957-1960.
- # 3 has been proven wrong, so likely a combination of 1 and 2.

#### Chimpanzee cells to Philadelphia

STUDIES OF LIVER FUNCTION TESTS IN CHIMPANZEES
AFTER INOCULATION WITH HUMAN INFECTIOUS
HEPATITIS VIRUS

FRIEDRICH DEINHARDT, GHEALAN COURTOIS, PAULETTE DHERTE,
PAUL, OSTERRIEPH, OASTON NINANE, GERTRUDE HENLE
AND WERNER HERLE
(Received for publication December 13, 1961)

Am.J.Hyo. 1962, Vol. 75: 311-321

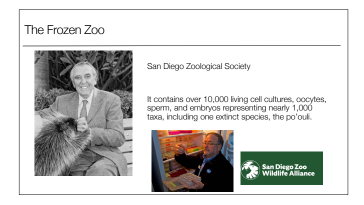


Tierac-culture studies. Additional efforts were made to isolate III virus in chinpanese hidron-lesses cultures. For a chinpanese hidron-lesses cultures. For a chinpanese idinops were sent by sir chinpanese lidneys were sent by sir chinpanese lidneys were trypholial of Philadelphia. These were trypholial of Philadelphia. These were trypholial control of the hidrogs and the preparation of the sidneys and the preparation of the sidneys and the preparation of the cultures were obtained from 3 or days and good cultures were obtained from 5 out for the cultures with the culture with

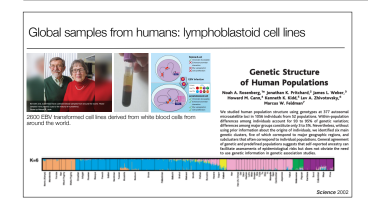


ATCC or the American Type Culture Collection is a nonprofit organization which collects, stores, and distributes standard reference microorganisms, cell lines and other materials for research and development

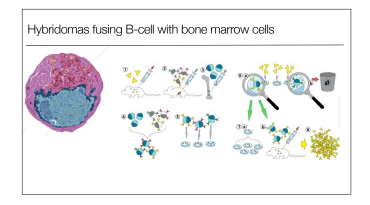
The organization holds a collection of more than 3,000 human and animal cell lines and an additional 1,200 hybridomas. ATCC's microorganism collection includes a collection of more than 18,000 strains of bacteria, as well as 3,000 different types of animal viruses and 1,000 plant viruses. In addition, ATCC maintains collections of protozoans, yeasts and fungi with over 7,500 yeast and fungus species and 1,000 strains of protists.



all primary cells, none of them transformed.

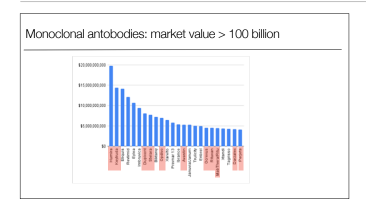


Kenn and Judy Kidd of Yale University have collected white blood cells from thousands of individuals from around the world.



B-cells from spleen of immunized animals fused with cancerous bone marrow cells (myeloma) generate immortal cells that produce monoclonal antibodies.

(1) Immunisation of a mouse (2) Isolation of B cells from the spleen (3) Cultivation of myeloma cells (4) Fusion of myeloma and B cells (5) Separation of cell lines (6) Screening of suitable cell lines (7) in vitro (a) or in vivo (b) multiplication (8) Harvesting



Humira (adalumimab) **AbbVie** anti-TNF, Crohns, RA, Psoriasis

Keytruda Merck: anti PD1 on T-cells, cancers

Dupixent (dupilimab) Sanofi: anti-IL4 receptor alpha, allergies, autoimmunity

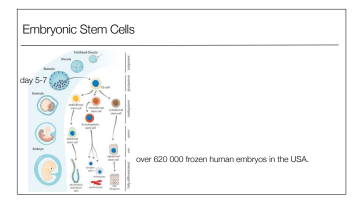
Stelara (ustekinumab), Jansen: IL12 & IL23 Crohns, Ulcer Col, Psoriasis

Opdivo (nivolumab), **Bristol Myers Squib**: anti-PD1, cancer Avastin (bevacizumab), **Roche**: anti-VEGF A, cancers, AMD

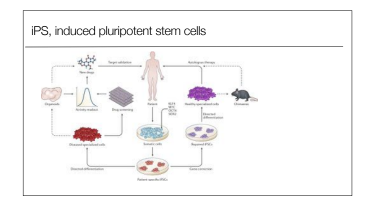
Ocrevus (ocrelizumab), **Roche**: anti-CD20, MS Rituxan (rituximab) **Roche**: antri-CD20, MS

Darzalex (daratumumab), Johnson & Johnson: anti-CD38, myleoma

Perjeta (pertuzumab), Roche, anti-HER2 breast cancer

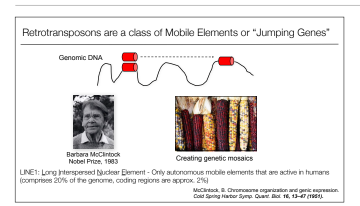


Tapping the "Germ Line"?
The inner cell mass day



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The organization holds a collection of more than 3,000 human and animal cell lines and an additional 1,200 hybridomas. ATCC's microorganism collection includes a collection of more than 18,000 strains of bacteria, as well as 3,000 different types of animal viruses and 1,000 plant viruses. In addition, ATCC maintains collections of protozoans, yeasts and fungi with over 7,500 yeast and fungus species and 1,000 strains of protists.



Retrotransposons are endogenous mobile elements or fragments of DNA that can copy themselves and insert into new chromosomal locations. That is the reason why transposons are also referred to as "jumping genes". Transposons have been discovered more then 50 years ago in maize by Barbara McClintock that won the Nobel prize for that discovery.

SHE COULD NOT EXPLAIN THE INHERITANCE OF MAIZE KERNEL COLORS BY MENDELIAN LAWS!

#### Consequences of Mosaicism after LINE1 Mobility: Mutations, Diversity and Disease

LINE1 (L1) mobility influence chromosome integrity and gene expression upon reinsertion causing genetic diversity that can generate changes in behavior and and population potentially diseases



Germline insertions can cause structural variants, deletions and sequence insertions within the human



First Evidence

Hemophilia A resulting from de novo insertion of LINE1 sequences. Kazazian et al Nature. 1988

To date, over 120 human diseases are associated with LINE1 events

Linker, Gage and Bedrosian, the Scientist 2017

Detecting recent (and relevant) events of LINE1 mobility in humans, prompted the field to look for when during development these insertions were happening and for many years it was thought that the insertions were only happening in the germline.

However, work from us and others have shown that new Line1 insertions happen during embryonic development and adulthood. Hence the idea that we are all walking mosaics.

WE ARE ALL "GENETIC MOSAICS" BUT ONLY TO LIMITD DEGRE AND ESPECILLY IN BRAIN AND TESTES... JUMPING GENES ARE A VERY DANGEROUS LIABILITY TO GENOMIC INTEGRITY ND SUCCESSFUL MULTICELLULARITY

In the following slides I will show you examples of studies lead by me and others that used reprograming technology to study retrotransposon mobility and we will also speculate on the implications of LINE1 mobility for disease and human evolution.

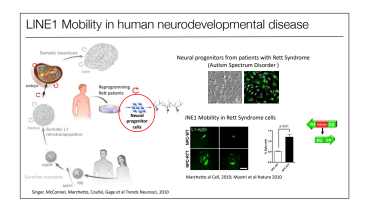
Germline retrotransposons are a major source of structural variants, deletions and sequence insertions within the human population 11-15. The vast majority of these germline variants have unknown functional effects. However, some variants are likely to have functional consequences for the individual. For example, although polymorphic insertions of retrotransposon sequences are abundant in the healthy human population, specific de novo retrotransposon insertions can cause haemophilia16, neurofibromatosis12 and other diseases. In addition to the insertion of the retrotransposon sequence, retrotransposition can mediate the deletion of the host DNA sequences. Furthermore, retrotransposon events can result in the presence of highly homologous sequences in different genomic locations. These sequences can then recombine, through nonallelic homologous recombination, to cause deletions, duplications, inversions

LINE1 mobility during human development How LINE1 mobility isn studied during development 1-Choose a cell line that represents human development During development, given the copy and paste-nature of the retrotransposon process LINE1-driven insertions accumulate progenito 2-Choose a system that allows for monitoring LINE1 mobility in real time Genetically Engineered LINE1 Element 1 Adapted from Moran et al Cell 1996 Singer, McConnel, Marchetto, Coufal, Gage et al Trends Neurosci. 2010

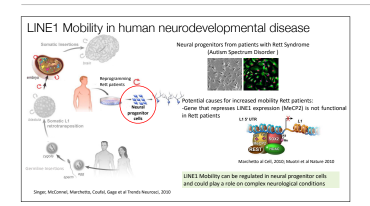
In the following slides I will show you examples of studies lead by me and others that used reprograming technology to study retrotransposon mobility and I will speculate on the implications of LINE1 mobility for disease and human evolution.

#### I WOULD STRESS: MOSAICISM MEANS THAT TWO NEIGHBROING NEURONS ARE NOT TOTALLY GENETICALLY IDENTICAL ANY MORE.

Putative implications of L1-mediated somatic mosaicism in the brain In a reversal of the commonly held belief that retrotransposition occurs primarily in the germline [83], it became clear that L1 elements are expressed in many somatic tissues, including the brain [7, 13, 84]. Recent evidence shows that L1 retrotransposition (curved red arrows) does not occur in the parental germline but in the soma during early embryonic development (colored dots), resulting in individuals that are genetically mosaic with respect to L1 composition [33]. It has been suggested, however, that L1 RNA may be transcribed in the parental germline and carried over in both male and female germ cells in the form of RNPs (black line with red dots) and integrated into the genome at the preimplantation stage [33] (colored spots); however, these events are probably rare, since retrotransposons are effectively silenced in the germline through a small RNA induced mechanism [78, 85]. Somatic L1 retrotranspositon events that occur during embryogenesis would result in clonal sectors of cells (colored patches) that carry the same insertion event. The size of clonal sectors depends on the developmental stage when the insertion occurred and the number of subsequent cell divisions. L1 insertion events that happen during embryonic brain development will be found in different brain regions (colored patches and dots), whereas events that happen during adult neurogenesis will be restricted to specific areas, such as the dentate gyrus (insert). According to our hypothesis, L1-induced mosaicism could increase variability in the brain (blue curve), which could have implications for behavioral phenotypes. The environment could influence regulation of somatic L1 retrotransposition in the brain and this influence could be mediated by epigenetic or hormonal mechanisms. Depending on its impact on the brain and the consequences, L1-induced somatic variability could either increase the risk for neurological disease or induce behavioral changes that could help the organism to better adapt to changing environments.

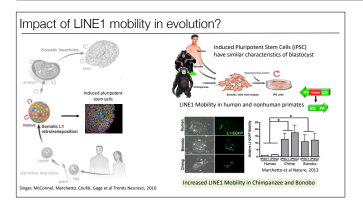


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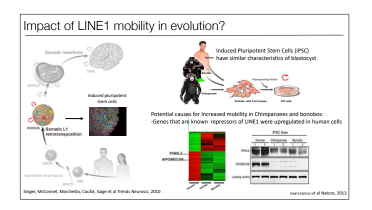


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RETT'S DEMONSTRATED THE HUGE DANGER OF UNCRONTROLLED, EXCESSIVE JUMPING

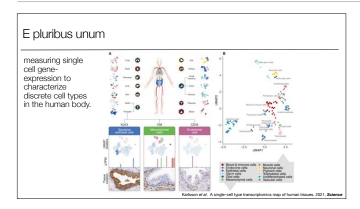


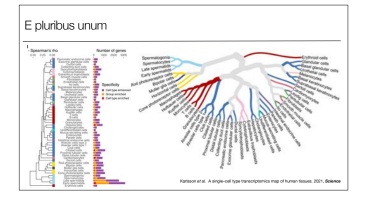
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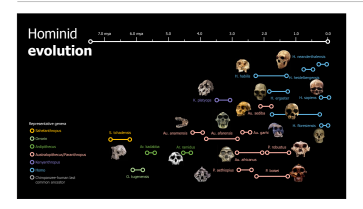
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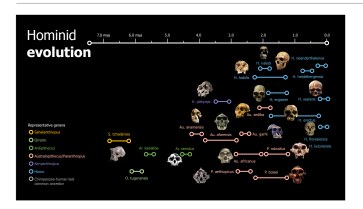
LINE1 ACTVITY (THEIR JUMPING) IS REPRESSED MOREN HUMANS THAN IN APES, IAM SURE THAT APES HAVE SOME LEVE OF REPRESSION AS WELL, OR THEIR GENOMES WOULD MELT....

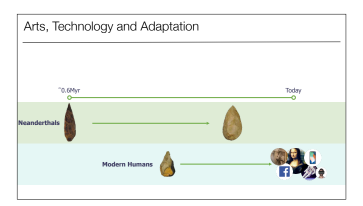


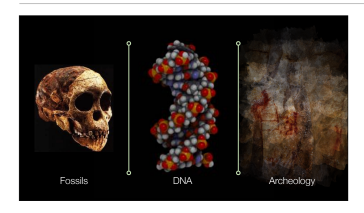


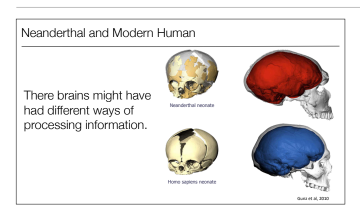


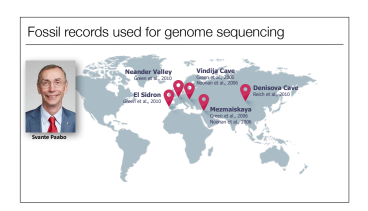


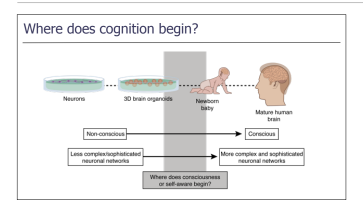


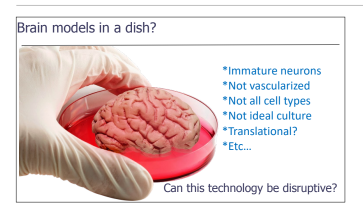


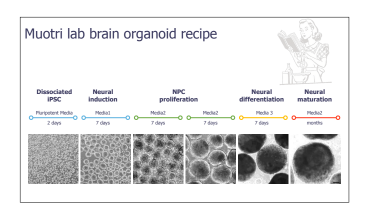






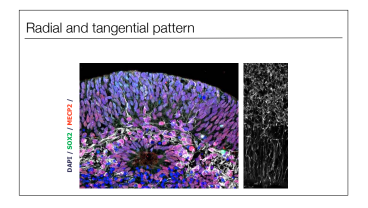








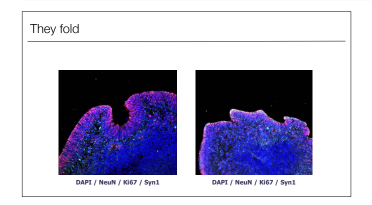




DAPI stains cell nuclei.

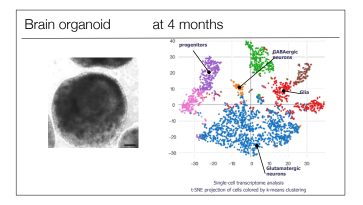
Staining for SOX2 in green: SOX2, is a transcription factor that is essential for maintaining self-renewal, or pluripotency, of undifferentiated embryonic stem cells. Sox2 has a critical role in maintenance of embryonic and neural stem cells.

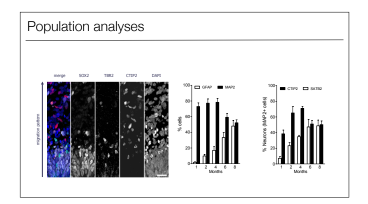
MECP2 (methyl CpG binding protein 2) is a gene that encodes the protein MECP2. MECP2 appears to be essential for the normal function of nerve cells.



NeuN also non as Fox 3 is a neuronal nuclear antigen that is commonly used as a biomarker for neurons.

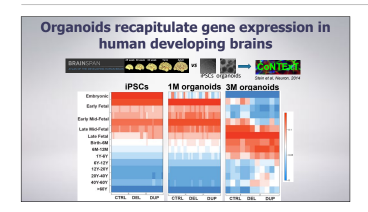
Antigen KI-67 is a nuclear protein that is associated with cellular proliferation. Syn1 (Syngap1) is aprotein that is critical for the development of cognition and proper synapse function. Mutations in humans can cause intellectual disability, epilepsy, autism and sensory processing deficits.



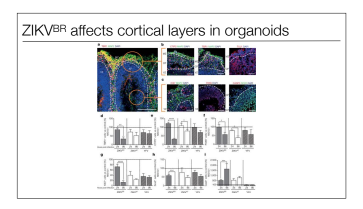


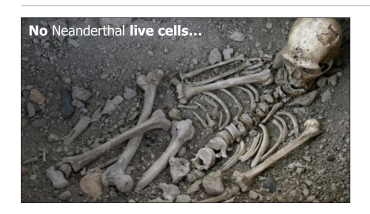
MAP2, microtubule also cited protein 2. This gene encodes a protein that belongs to the microtubule-associated protein family.

CTIP2 transcription factor expressed by subconical projecting neurons









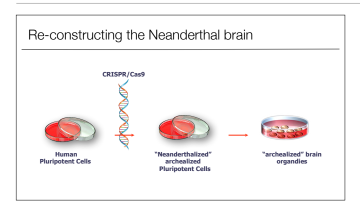
reconstruction of Neanderthal burial at La Chapelle aux Saints, Southern France (~50 000 year old)

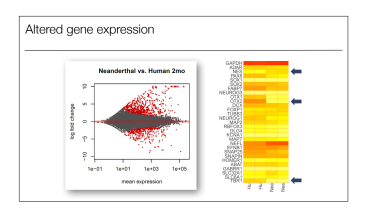


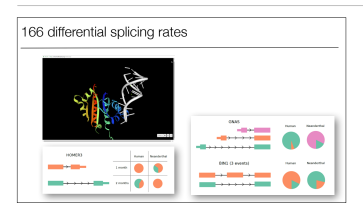
Reconstruction of a Neanderthal by Kennis brothers admired by modern human in London's Natural History museum.s

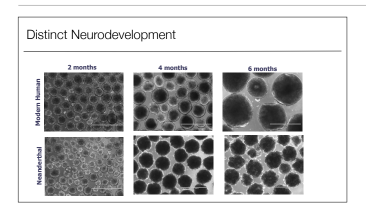
| Catalog of human-specific variant                |         |        |            |          |         |          |          |
|--|---------|--------|------------|----------|---------|----------|----------|
| DNA differences with effects on protein sequence |         |        |            |          |         |          |          |
|  | DDX53   | SLC8A1 | IRAK1BP1   | GLDC     | GPT     | STARD9   | KIF26B   |
|  | CXorf59 | NOTO   | MCHR2      | FRRS1L   | DCHS1   | SLC12A1  | GRB1L    |
|  | GAP43   | ANKMY1 | ZBTB24     | NEK6     | KIF18A  | KIAA1199 | LMNB2    |
|  | FRMD7   | SCAP   | KATNA1     | TTF1     | PLAC1L  | CDH16    | RASA1    |
|  | ZNF185  | ORSK4  | LRRD1      | FBXW5    | ZNHIT2  | PIEZ01   | MFSD12   |
|  | TKTL1   | NOP14  | KLF14      | FAM166A  | PRDM10  | SPAGS    | NCOA6    |
|  | IFI44L  | EVC2   | CALD1      | ARRDC1   | LRTM2   | SSH2     | LYPLA1   |
|  | VCAM1   | HERCS  | ERI1       | ANKRD30A | LAG3    | SYNRG    | TP53TG5  |
|  | SPAG17  | DHX29  | CSGALNACT1 | FAM14981 | SCAF11  | CD300LG  | C21orf62 |
|  | SLC27A3 | PTCD2  | GSR        | FAM178A  | SLITRK1 | TEX2     | UBQLN3   |
|  | SPTA1   | SV2C   | ADAM18     | CASCS    | NOVA1   | ITG84    | RSPH1    |
|  | NFASC   | VCAN   | RB1CC1     | PNLIP    | TTLLS   | RFNG     | ENTHD1   |
|  | GPR132  | DLGAP2 | ADSL       |          |         |          |          |

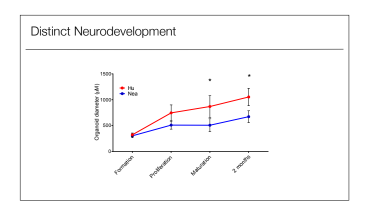


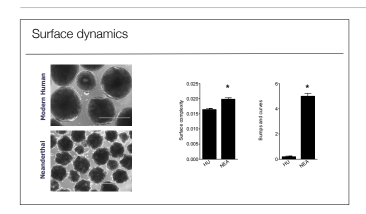


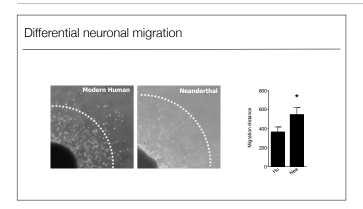


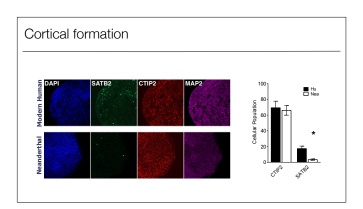


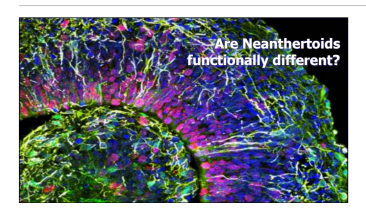


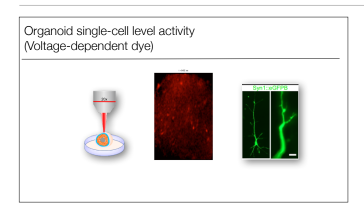


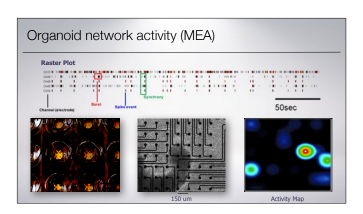


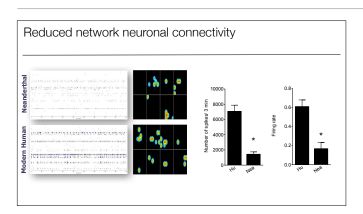


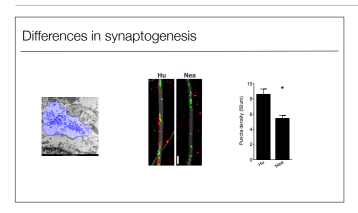


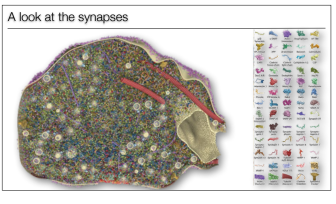










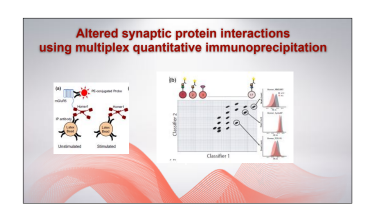


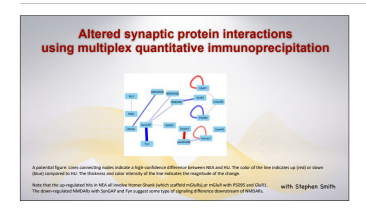






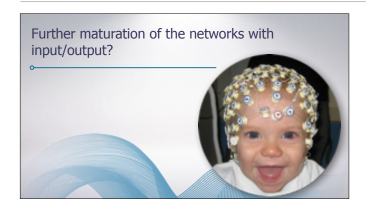








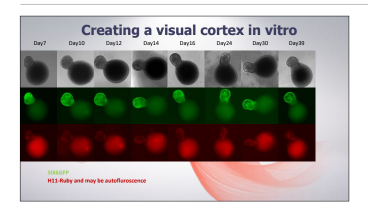


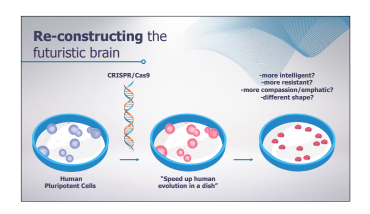










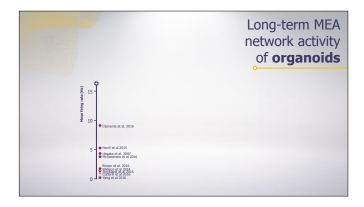


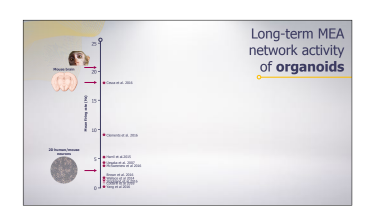


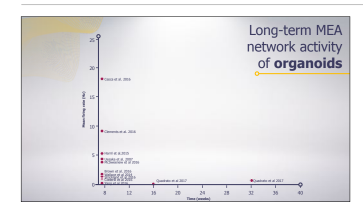


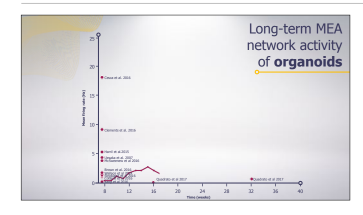


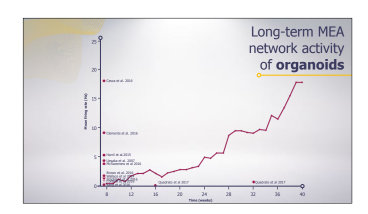


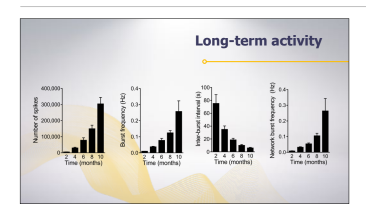


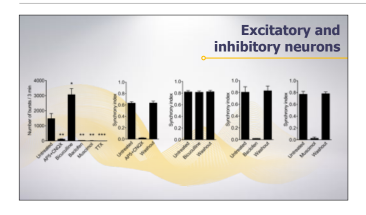


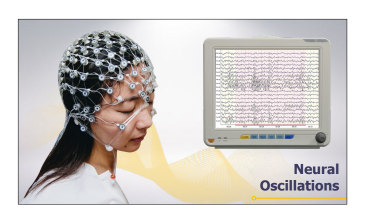




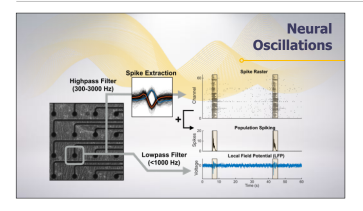


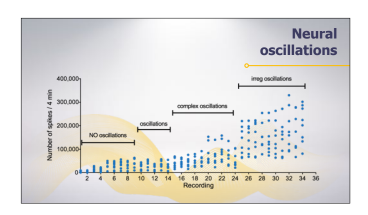


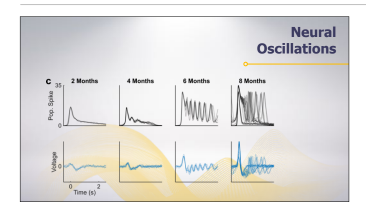


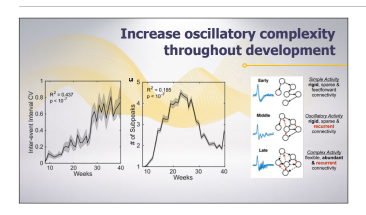












Spontaneous activity transients

