





Con l'intervento di

Ernesto Burgio · 'Il genoma minacciato'

Carine Brochier · 'Cure palliative ed eutanasia'

David Lana Tuñon, Fermín Jesús González Melado, Luis Torró Ferrero Tavola rotonda: 'Sfide attuali del transumanesimo'

Claudia Estela Vanney · 'Costruire una nuova cultura dall'interdisciplinarità'
Brad S. Gregory · 'Come la rivoluzione religiosa Ha secolarizzato la società'

18 - 19 Ottobre 2019



Pontificio Collegio Spagnolo

italiana

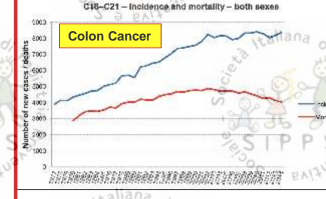
Via di Torre Rosa 2 00165 Roma (Italia) www.colegioespagnol.org

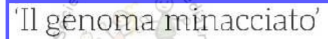
№ Santander

OBESITY IS NOW AS GLOBAL EPIDEMIC!

Obesity

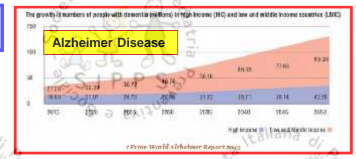
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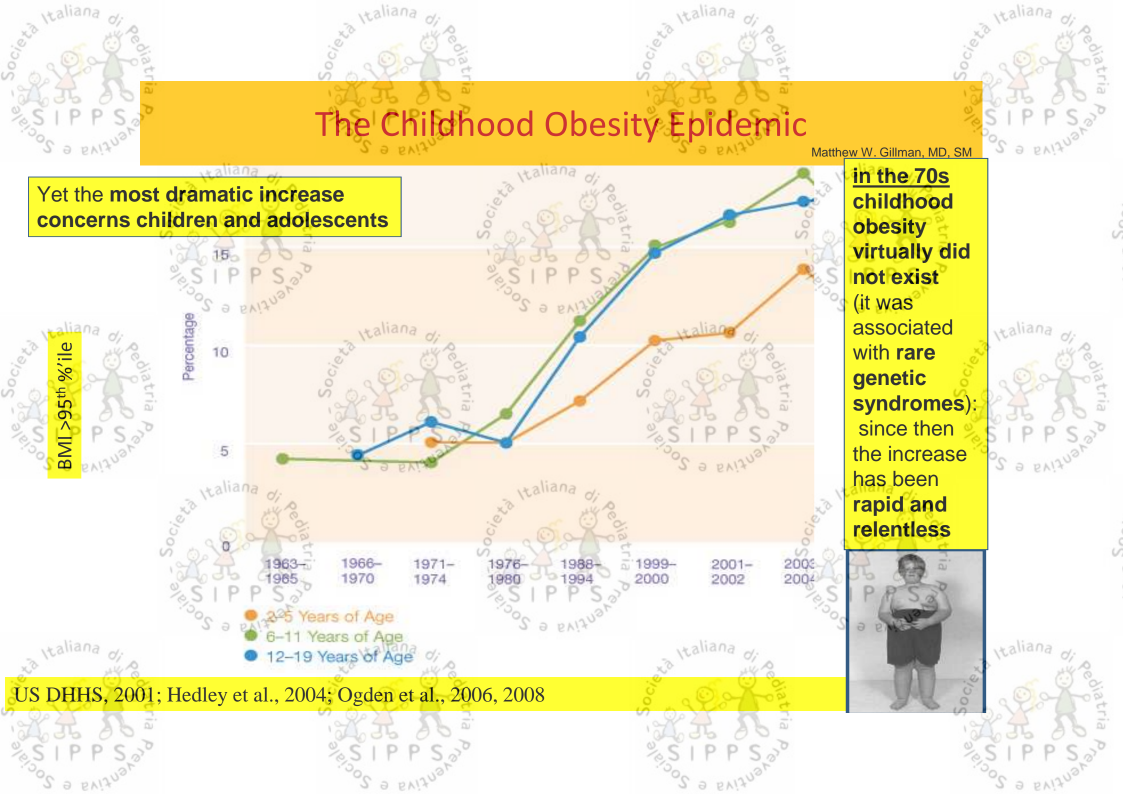


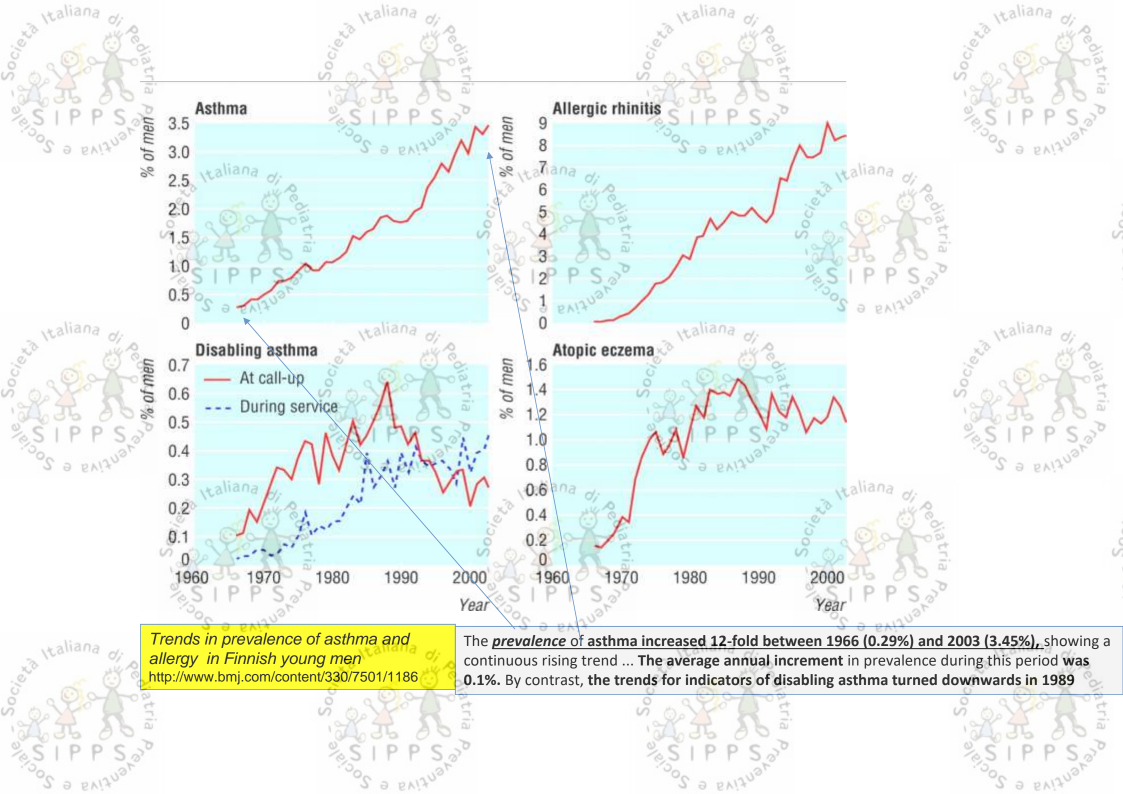


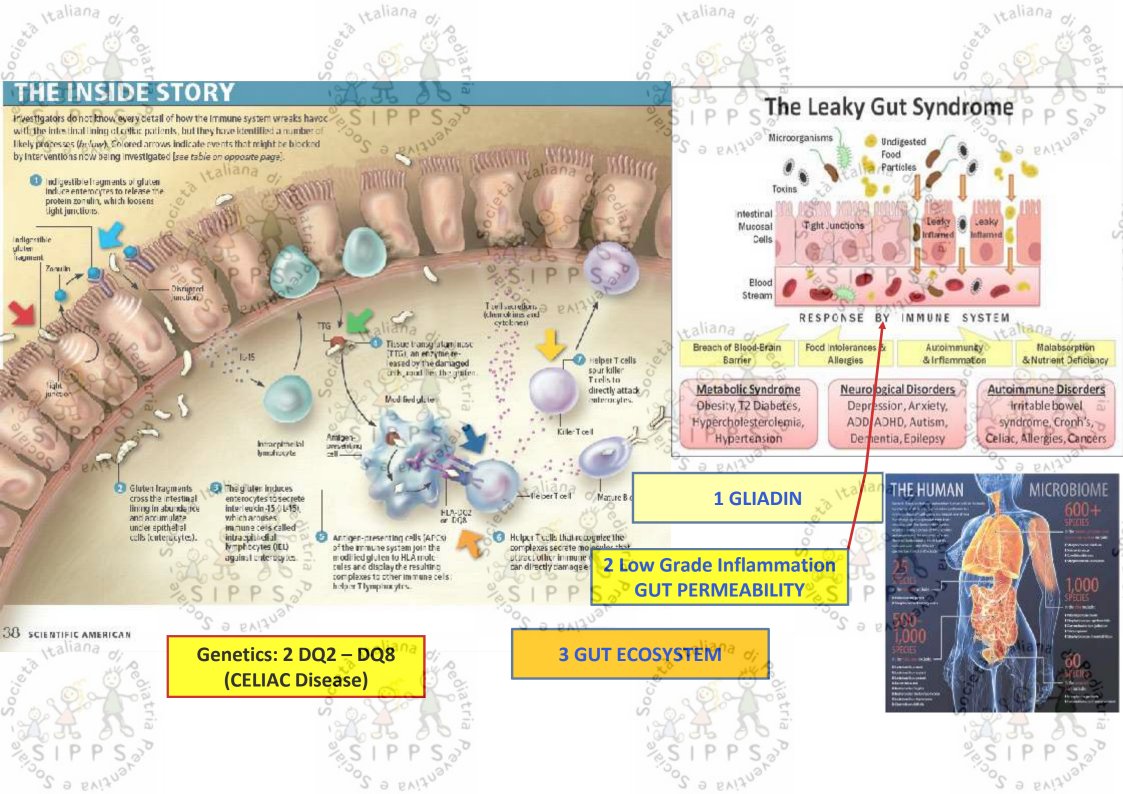
ERNESTO BURGIO
ECERI - European Cancer and
Environment Research Institute





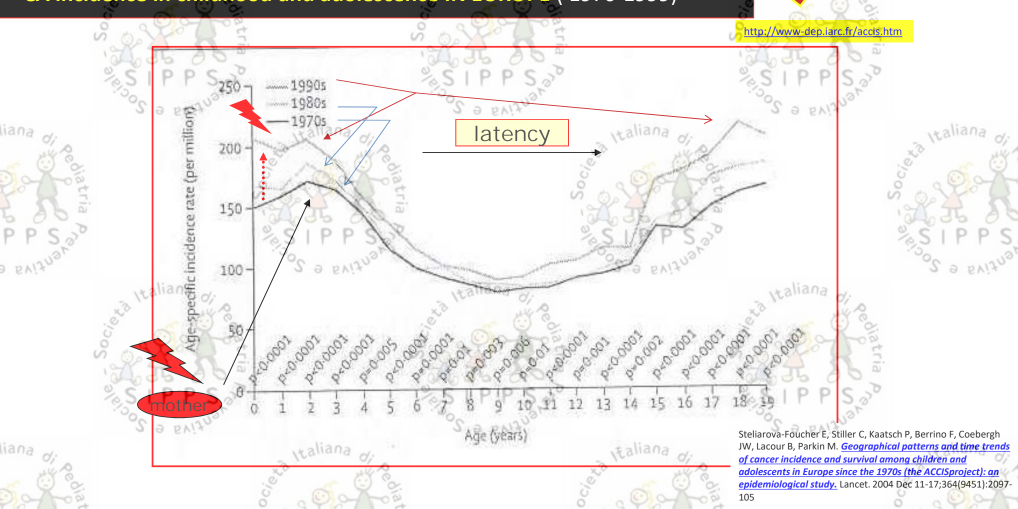






A first draft of the report, published on the Lancet in 2004, demonstrated an annual increase of 1-1,5% for all cancers (with more marked increases in lymphomas, soft tissue sarcomas, tumours of the nervous system...). But the most troubling was the increase - almost the double - for all cancers in the very first year of life (apparently due to transplacental or even trans-generational exposure)

CA incidence in childhood and adolescence IN EUROPE (1970-1999)





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A Silent Pandemic

Industrial Chemicals Are Impairing The Brain Development of Children Worldwide

For immediate release: Tuesday, November 7, 2006



Landrigan Ph

THE LANCET

Developmental neurotoxicity of industrial chemicals

P.Grandjean, P.J. andrigan

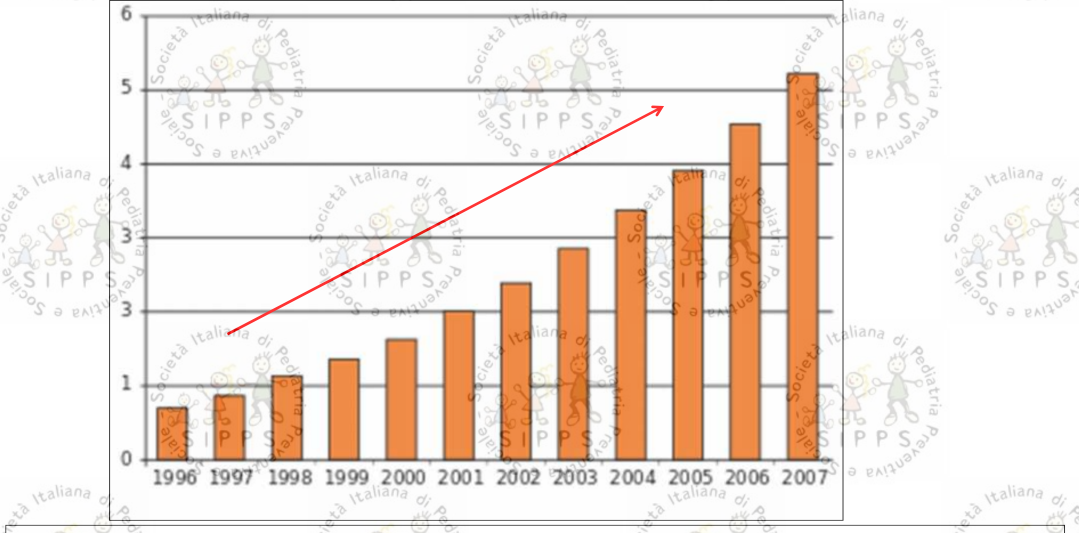
Neurodevelopmental disorders such as autism, attention deficit disorder, mental retardation, and cerebral palsy are common, costly, and can cause lifelong disability. Their causes are mostly unknown. A few industrial chemicals (eg. lead, methylmercury, polychlorinated biphenyls IPCBs], arsenic, and toluene) are recognised causes of neurodevelopmental disorders and subclinical brain dysfunction. Exposure to these chemicals during early fetal development can cause brain injury at doses much lower than those affecting adult brain function. Recognition of these risks has led to evidence-based programmes of prevention, such as elimination of lead additives in petrol. Although these prevention campaigns are highly successful, most were initiated only after substantial delays. Another 200 chemicals are known to cause clinical neurotoxic effects in adults. Despite an absence of systematic testing, many additional chemicals have been shown to be neurotoxic in laboratory models. The toxic effects of such chemicals in the developing human brain are not known and they are not regulated to protect children. The two main impediments to prevention of neurodevelopmental deficits of chemical origin are the great gaps in testing chemicals for developmental neurotoxicity and the high level of proof required for regulation. New, precatitionary approaches that recognise the unique vulnerability of the developing brain are needed for testing and control of chemicals.

A few industrial chemicals (eg, lead, methylmercury, polychlorinated biphenyls [PCBs], arsenic, and toluene) were recognized causes of neurodevelopmental disorders and subclinical brain dysfunction.

Twelve years ago two well-known experts in Environmental Health, a pediatrician and an epidemiologist, launched an alarm from the pages of the Lancet, affirming that a silent pandemic of neurodevelopmental disorders was spreading, also due to the shortage of funds in this area of research



In fact the reports of autism cases per 1,000 children had increased dramatically over the years in the U.S. from 1996 to 2007



Newschaffer CJ, Croen LA, Daniels J et al. *The epidemiology of autism spectrum disorders* Annu Rev Public Health. 2007;28:235–58.



AUTISM

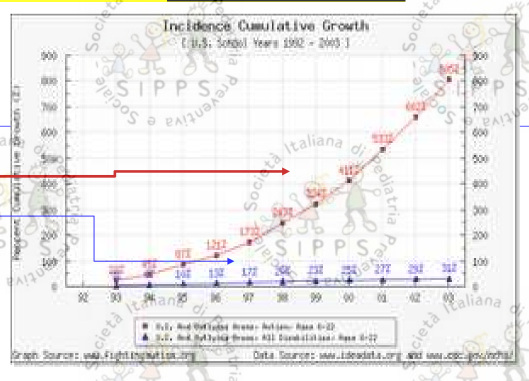
(ASD : Autism Spectrum Disorders)

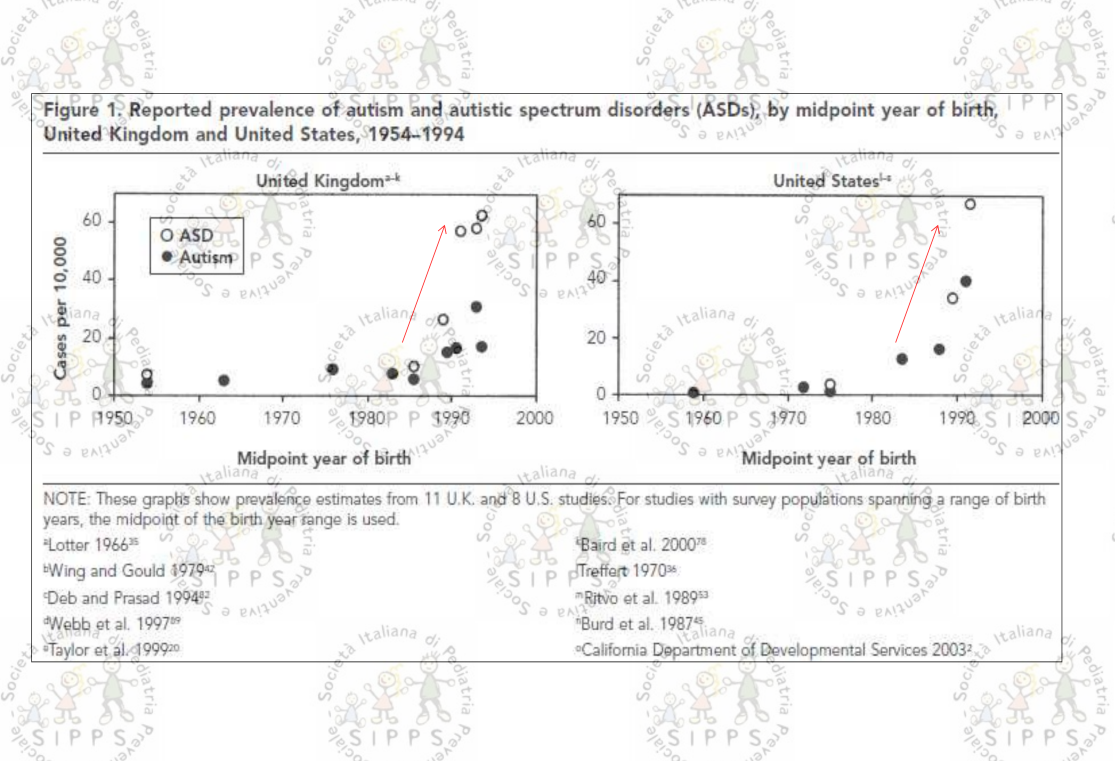
ASD is the fastest-growing developmental disorder in the world, the prevalence of diagnosis having increased by 600% over the last 20 years. New diagnosed

cases (incidence) in US increased from 15,580 in 1992 to 163.773 in 2003

The estimated <u>prevalence</u> was of 8-12 cases/1000 children in 2012...

Chart showing the increase in autism diagnosis (A) versus all disabilities (B) (statistics based on data from the National Center for Health Statistics)







autism the great modern health concern



Autism spectrum disorders (ASDs) are a group of developmental disabilities that can cause significant social, communication and behavioral challenges. People with ASDs handle information in their brain differently than other people. ASDs are spectrum disorders." That means ASDs affect each person in different ways, and can range from very mild to severe. There are three different types of ASDs: Autistic Disorder (also called classic autism). 1:1500 Asperger Syndrome and Pervasive Developmental Disorder - Not Otherwise Specified (PPD-NOS, also called atypical autism?)

Autistic Disorder

What most people think of when hearing the word "autism." People with autistic disorder usually have significant language delays, social and communication challenges and unusual behaviors and interests.

Asperger Syndrome

Usually have some milder symptoms of autistic disorder. They might have social challenges and unusual behaviors and interests. However, typically do not have problems with language or intellectual disability.

Pervasive Developmental Disorder

The symptoms might cause only social and communication challenges. People with PDD-NOSustally have fewer and milder symptoms than those with autistic disorder.

2002 1:150

ASD CDC estimated aver 2014 1:68

of the population of children Daged 3-V7 have an ASD

with

ASDs 4 to 7 times more likely to occur in BOYS than in GIRLS

2006 1: 110

There is no medical test to diagnose ASDs. doctors look at the child's behavior and development to make a diagnosis.

About half of parents of children with ASD notice their child's unusual behaviors by age 18 months



about four-fifths notice by age 24 months

A person with an ASD might:

Not respond to their name by 12 months. Applid eye contact and want to be alone | Have delayed speech and language skills Repeat words or phrases over and over (echolalia) | Give unrelated answers to questions | Get upon by minor changes

2008 1:88

ASDs are the fastest-growing developmental disability

annual growth

Reports of autism cases per 1,000 children

Lifetime cost to care for an Individual with an ASD

Estimated from regart studies

\$3.2m \$4,110-\$6,200 per year

of medical expenditures for an individual with an ASD than one without

20141:68

SOURCES: CDC | WWW.

http://arstechnica.com/science/2012/04/new-autism-studies-find-new-mutations-many-genes-behind-the-disorder/



Neurobehavioural effects of developmental toxicity

Philippe Grandjean, Philip J Landrigan

Neurodevelopmental disabilities, including autism, attention-deficit hyperactivity disorder, dyslexia, and other cognitive impairments, affect millions of children worldwide, and some diagnoses seem to be increasing in frequency. Industrial chemicals that injure the developing brain are among the known causes for this rise in prevalence. In 2006, we did a systematic review and identified five industrial chemicals as developmental neurotoxicants: lead, methylmercury, polychlorinated biphenyls, arsenic, and toluene. Since 2006, epidemiological studies have documented six additional developmental neurotoxicants—manganese, fluoride, chlorpyrifos, dichlorodiphenyltrichloroethane, tetrachloroethylene, and the polybrominated diphenyl ethers. We postulate that even more neurotoxicants remain undiscovered. To control the pandemic of developmental neurotoxicity, we propose a global prevention strategy. Untested chemicals should not be presumed to be safe to brain development, and chemicals in existing use and all new chemicals must therefore be tested for developmental neurotoxicity, To coordinate these efforts and to accelerate translation of science into prevention, we propose the urgent formation of a new international clearinghouse.

The same two authors returned to the problem seven years later, with a broad review published the Lancet Neurology (2014)

Since 2006, epidemiological studies have documented six additional developmental neurotoxicants — manganese, fluoride, chlorpyrifos, tetrachloroethylene, dichlorodiphenyltrichloroethane, and the polybrominated diphenyl ethers.

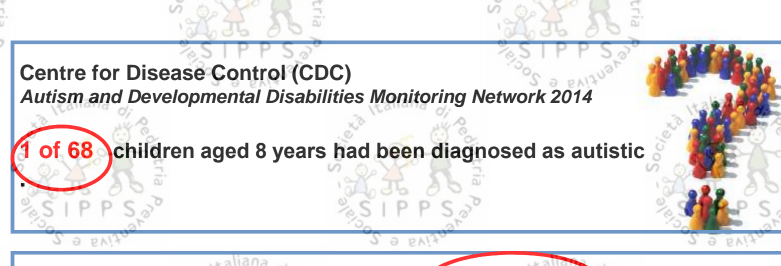
We postulate that even more neurotoxicants remain undiscovered

Lancet Neural 2014; 13: 330-38

Published Online February 15, 2014 http://dx.doi.org/10.1016/ 51474-4422(13)70278-3

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And it is increasingly evident that the increase continues unabated

Prevalence of Autism Spectrum Disorders in EU: 0,62 - 0,7%

Autism. Lai MC, Lombardo MV, Baron-Cohen S. Lancet. 2014 Mar.



Community Report on Autism 2018

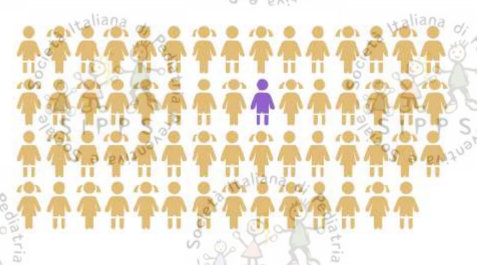
Centers for Disease Control and Prevention





Community Report from the Autism and Developmental Disabilities Monitoring (ADDM) Network

is the average percentage identified with ASD



1 in 5 9
8-year-old children
were identified with ASD
by ADDM in 2014

Why is this information important and how can it be used?

1. Lower the age of first evaluation by community providers

 Increase awareness of ASD among black and Hispanic families, and identify and address barriers in order to ensure that all children with ASD are evaluated, diagnosed, and connected to services.



OBJECTIVES: To estimate the national prevalence of parent-reported autism spectrum disorder (ASD) diagnosis among US children aged 3 to 17 years as well as their treatment and health care experiences using the 2016 National Survey of Children's Health (NSCH).

on the health and well-being of children aged 0 to 17 years. The NSCH collected parentreported information on whether children ever received an ASD diagnosis by a care
provider, current ASD status, health care use, access and challenges, and methods of
treatment. We calculated weighted prevalence estimates of ASD, compared health care
experiences of children with ASD to other children, and examined factors associated with
increased likelihood of medication and behavioral treatment.

that their child had ever received an ASD diagnosis and currently had the condition.

diagrosis is now 1 in 40, with rates of ASD-specific treatment usage varying by children's socio-sengraphic and exoccurring conditions.

American Academy of Pediatrics

Downloaded from www.aappublications.org/news by guest on December 3, 20

TELE THE POINTS PAR PUBLIC TO GOOD HIS CT 2018

New genetic risk factor for developing autism spectrum disorder identified

Date: August 31, 2017

Source: Oregon Health & Science University

Summary: A new systematic analysis has been applied to a cohort of 2,300 families who have a

single child affected with autism. The study focused on identifying and characterizing low-lying genetic mutations that may have been missed in previous research, given

these mutations are only present in a fraction of the bulk DNA of an individual.

tematic analysis to a cohort of 2,300 families who have a single child affected with autism. The study focused on identifying and characterizing low-lying genetic mutations that may have been missed in previous research, given these mutations are only

present in a fraction of the bulk DNA of an individual.

Known as postzygotic mosaic mutations, or PMMs, these genetic changes occur after the conception of the human zygote during the development cycle of a fetus. An individual will contain a mosaic — or assortment — of mutated and non-mutated cells with the level of mosaicism depending on the time and location of the mutation's occurrence. This emerging class of genetic risk factors has recently been implicated in various neurologic conditions, however,

.. yet many continue to define autism (and schizophrenia) as "genetic" diseases !!??!!

As in this case: The risk of autism connected to unexpected exonic mutations ...!!??!!

Deidre R. Krupp, Rebecca A. Barnard, Yannis Duffourd, Sara A. Evans, Ryan M. Mulqueen, Raphael Bernier, Jean-Baptiste Rivière, Eric Fombonne, Bran J. O'Roak. Exonic Mosaic Mutations Contribute Risk for Autism Spectrum Disorder.

The American Journal of Human Genetics, 2017, DOI: 10.

10.1016/j.ajhg.2017.07.016

Autism risk due to unexpected mosaic mutations

Whole genome sequencing identifies new genetic signature for autism

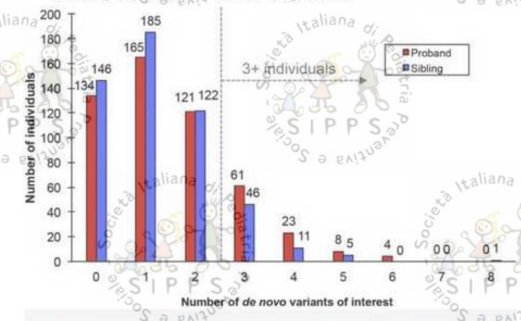
October 12, 2017 Date:

Source: Howard Hughes Medical Institute

Summary: An analysis of the complete genomes of 2,064 people reveals that multiple genetic vari-

ations could contribute to autism. The work suggests that scanning whole genomes may

one day be useful for clinical diagnostics.



Children with autism (red bars) were significantly more likely to have three or more genetic variations than their unaffected siblings (blue bars).

..or here: Autistic children have > 3 mutations if compared to unaffected siblings ...!!??!!

Tychele N. Turner, Bradley P. Cee, Diane E. Dickel, Kendra Hoekzema, Bradley J. Nelson, Michael C. Zody, Zev N. Kronenberg, Fereydoun Hormozdiari, Archana Raja, Len A. Pennacchio, Robert B. Darnel Evan E. Eichler. Genomic Patterns of De Novo Mutation in Simplex Autism Cell, 2017

DOI: 10.1016/j.cell.2017.08.047

Autism genetics study calls attention to motor skills, general cognitive impairment

Date: February 7, 2018

Source: Cold Spring Harbor Laboratory

Summary: A new study of the genetic factors involved in the causation of autism spectrum disor-

ders (ASD) draws fresh attention to the impact these illnesses frave on motor skills, and more broadly on cognitive function. Careful inference from the data suggests to researchers that the genetic factors causing ASD broadly diminish the brain's cognitive

functions.



Mutations that appear in a child which are not present in either parent — called de novo mutations — can be important in autism. Severe, gene-disrupting de novo mutations are thought to be capable of causing the disorder in certain instances. New research shows that diminished motor skills, like low non-verbal Q, correlate with the severity of de novo mutations. More broadly the study calls attention to role played by genetics in di-

...or in this case: new mutations disturbing motor functions could be important in autism ...!!??!!

Andreas Buja, Natalia Volfovsky, Abba M. Krieger, Catherine Lord, Blex E. Lash, Michael Wigler, Ivan Iossifov. Damaging de novo mutations diminish motor skills in children on the autism spectrum. Proceedings of the National Academy of Sciences, 2018; 201715427 DOI: 10.1073/pnas.1715427115

eSIPPS and Services

JAMA Psychiatry | Original Investigation

Association of Genetic and Environmental Factors
With Autism in a 5-Country Cohort

Dan Bal, MSc, Benjamin Hon, Kei Yip, PhD; Gayle C, Windham, PhD, MSPH, Andre Sourander, PhD, Richard Francis, PhD; Rinat Yoffe, MPH; Emma Glasson, PhD; Behrang Mahjani, PhD; Auli Suominen, MSe; Halen Leonard, MBChB, MPH; Mika Gissler, PhD; Joseph D, Buxbaum, PhD; Kingsley Weng, PhD; Diana Schendel, PhD, Arad Kodesh, MD; Michaeline Breshnahan, PhD, MPH; Stephen Z, Levine, PhD; Frik T, Pamer, PhD; Stefan N, Hansen, PhD; Christina Hultman, PhD; Abraham Reichenberg, PhD; Sven Sandin, PhD

IMPORTANCE The origins and development of autism spectrum disorder (ASD) remain unresolved. No individual-level study has provided estimates of additive genetic, maternal, and environmental effects in ASD across several countries.

OBJECTIVE To estimate the additive genetic, maternal, and environmental effects in ASD.

DESIGN. SETTING. AND PARTICIPANTS Population-based, multinational cohort study including full birth cohorts of children from Denmark, Finland, Sweden, Israel, and Western Australia born between January 1, 1998, and December 31, 2011, and followed up to age 16 years. Data were analyzed from September 23, 2016 through February 4, 2018.

MAIN OUTCOMES AND MEASURES Across 5 countries, models were fitted to estimate variance components describing the total variance in risk for ASD occurrence owing to additive genetics, maternal, and shared and nonshared environmental effects.

results. The analytic sample included 2 001 631 individuals, of whom 1 027 546 (51.3%) were male. Among the entire sample. 22 156 were diagnosed with ASD. The median (95% CI) ASD heritability was 80.8% (73.2%-85.5%) for country-specific point estimates, ranging from 50 9% (25.1%-75.5%) (Finland) to 86.8% (69.8%-100.0%) (Israel). For the Nordic countries combined heritability estimates ranged from 81.2% (73.9%-85.3%) to 82.7% (79.1%-86.0%). Maternal effect was estimated to range from 0.4% to 1.6%. Estimates of genetic, maternal, and environmental effects for autistic disorder were similar with ASD.

CONCLUSIONS AND RELEVANCE. Based on population data from 5 countries, the heritability of ASD was estimated to be approximately 80%, indicating that the variation in ASD occurrence in the population is mostly owing to inherited genetic influences, with no support for contribution from maternal effects. The results suggest possible modest differences in the sources of ASD risk between countries.

JAMA Psychiatry. doi:10.1001/jamapsychiatry.2019.1411 Published online July 17, 2019.

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Corresponding Author: Sven Sandin, PhD, Department of Medical Epidemiology and Biostatistics, Karolinska Institutet, Nobels väg 6, SE-17177 Stockholm, Sweden (sven sandin@ki.se).

Italiana

Heritability estimates ranged from **81.2%**(73.9%-85.3%) to 82.7% (79.1%-86.0%). **Maternal effect** was estimated to range from **0.4% to 1.6%**.

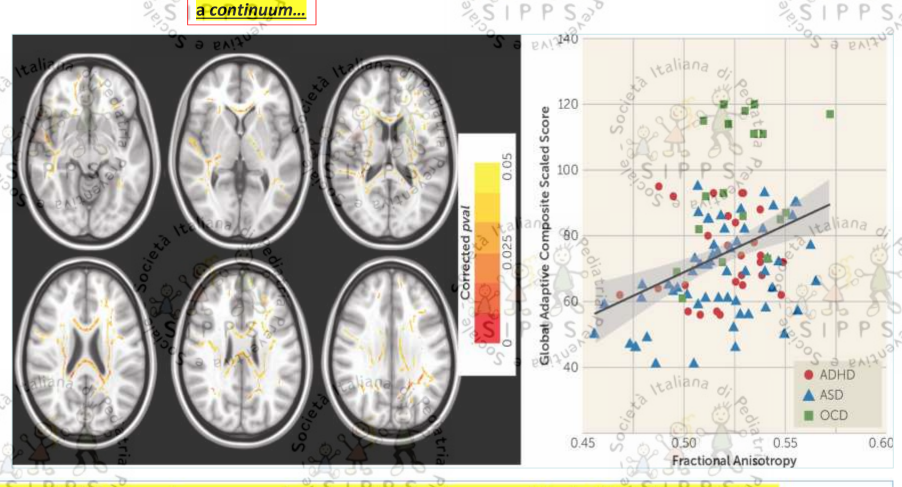
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Autism, ADHD and OCD have common symptoms and are linked by some of the same genes.

Yet they have always been considered as separate disorders

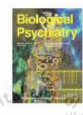


Children with <u>autism</u> and ADHD showed more severe impairments affecting more of the brain's white matter than those with OCD. This finding may reflect the fact that both <u>autism</u> and ADHD typically have an onset at a much younger age than OCD, and at a time when a number of different white matter tracts are going through rapid development,



Biological Psychiatry

Volume 49, Issue 12, 15 June 2001, Pages 1002-1014



The unmet needs in diagnosis and treatment of mood disorders in children and adolescents

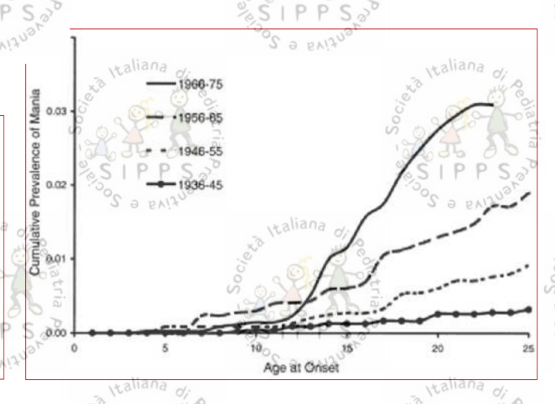
Mood disorders in children and adolescents: an epidemiologic perspective

Ronald C Kessler ^a A, Shelli Avenevoli ^b, Kathleen Ries Merikangas ^l

Adolescence is a time of increasing vulnerability for severe mental health disorders such as depression.

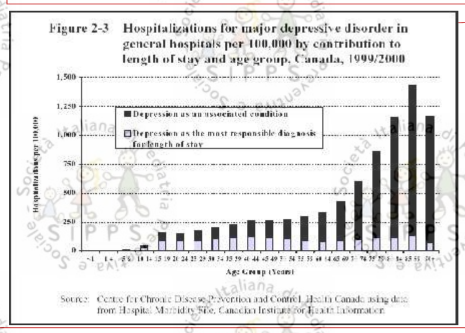
Epidemiological studies show that the incidence of new cases of depression drastically increases with puberty..

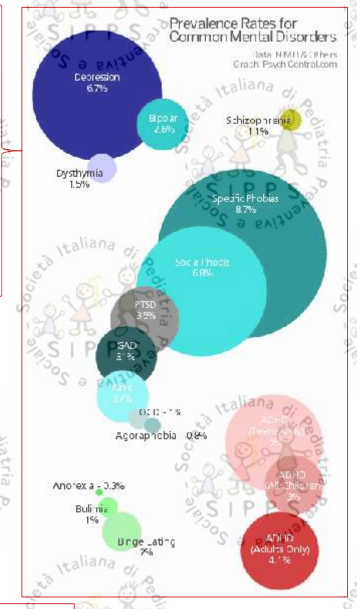
Importantly, there is growing evidence that sleep disturbance in adolescence may predict the development of depression.. In addition to the increase in the prevalence of depression with the transition from childhood to adolescence, there is also a secular trend of an increasing incidence of depression during adolescence since the 1960s



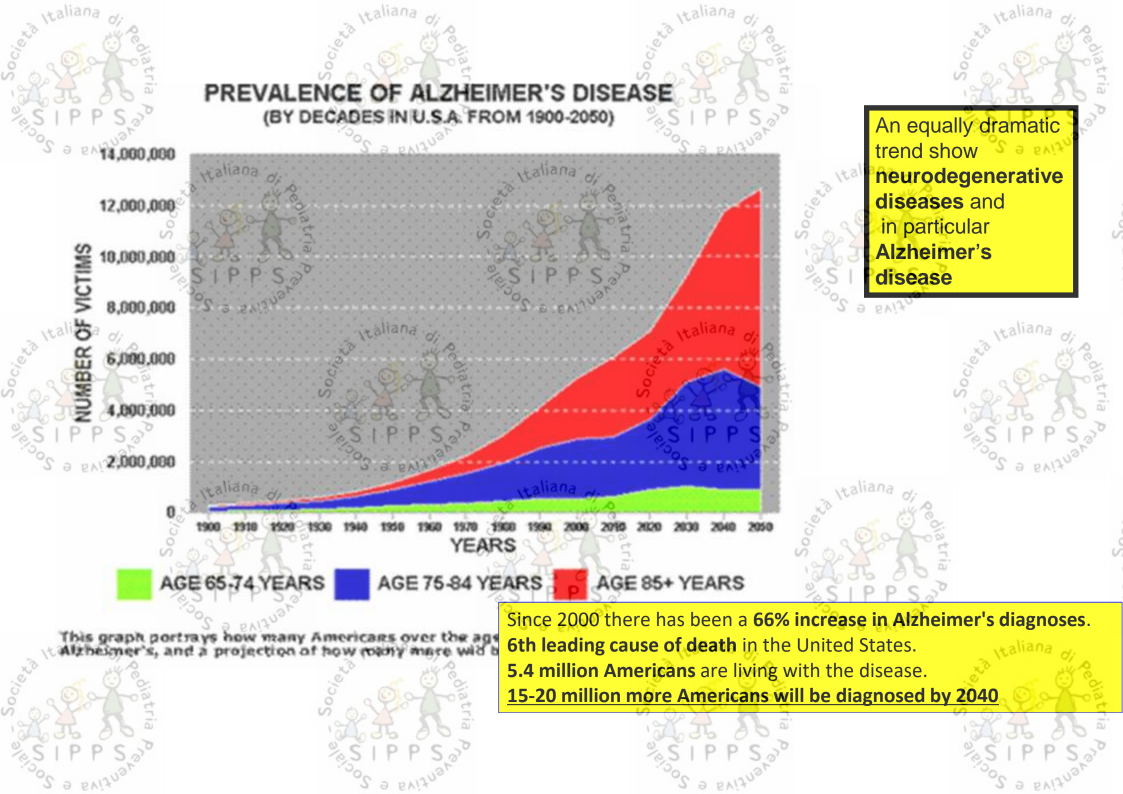
http://www.slideshare.net/CMoondog/depression-powerpoint-13945746

An estimated one in ten Americans suffer from depression, an illness that affects from depression, and mental well-being both physical and mental well-being chronic in nature, depression can be chronic in nature, depression mustances or chronic in nature, depression if requently, a chronic in nature, depression in triggered by adverse life circumstances or chronic in nature, depression in the blue. Frequently, a combination of genetic, psychological and combination of genetic psychological and

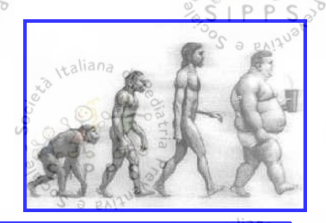




http://psychcentral.com/blog/archives/2009/10/05/prevalence-of-common-mental-disorders/

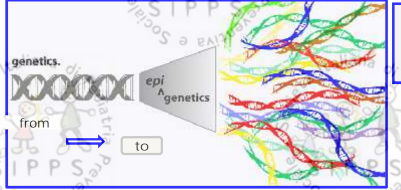








Evolution of DOHaD: the impact of environmental hazards on the origins of current "pandemics"

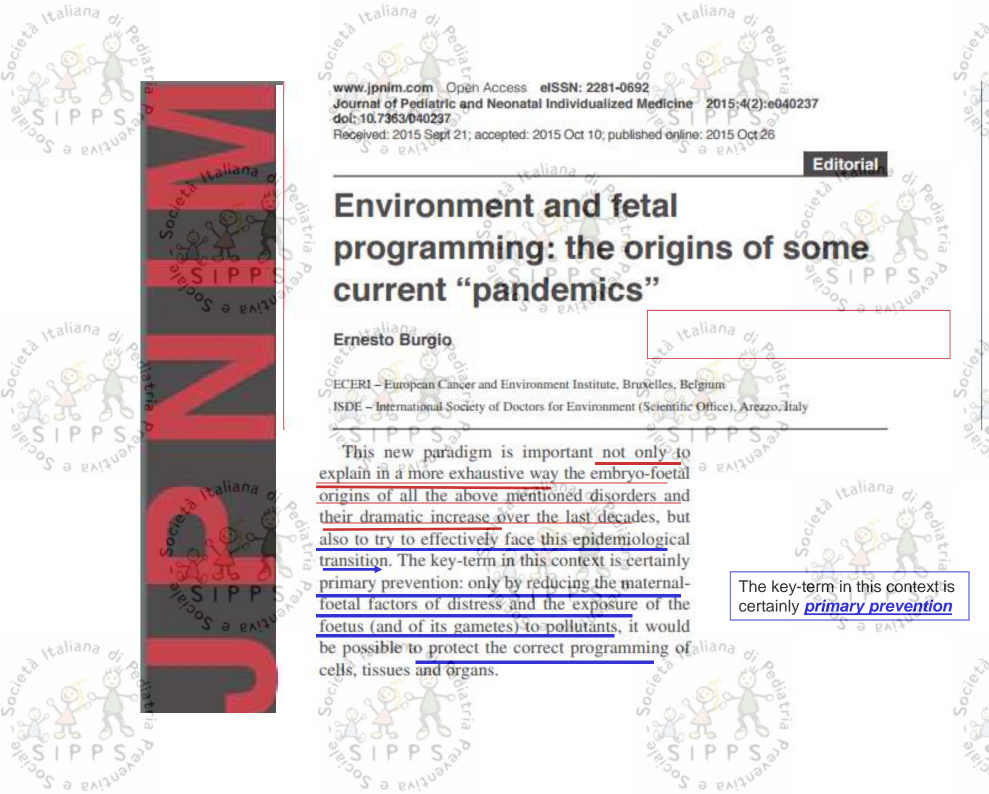


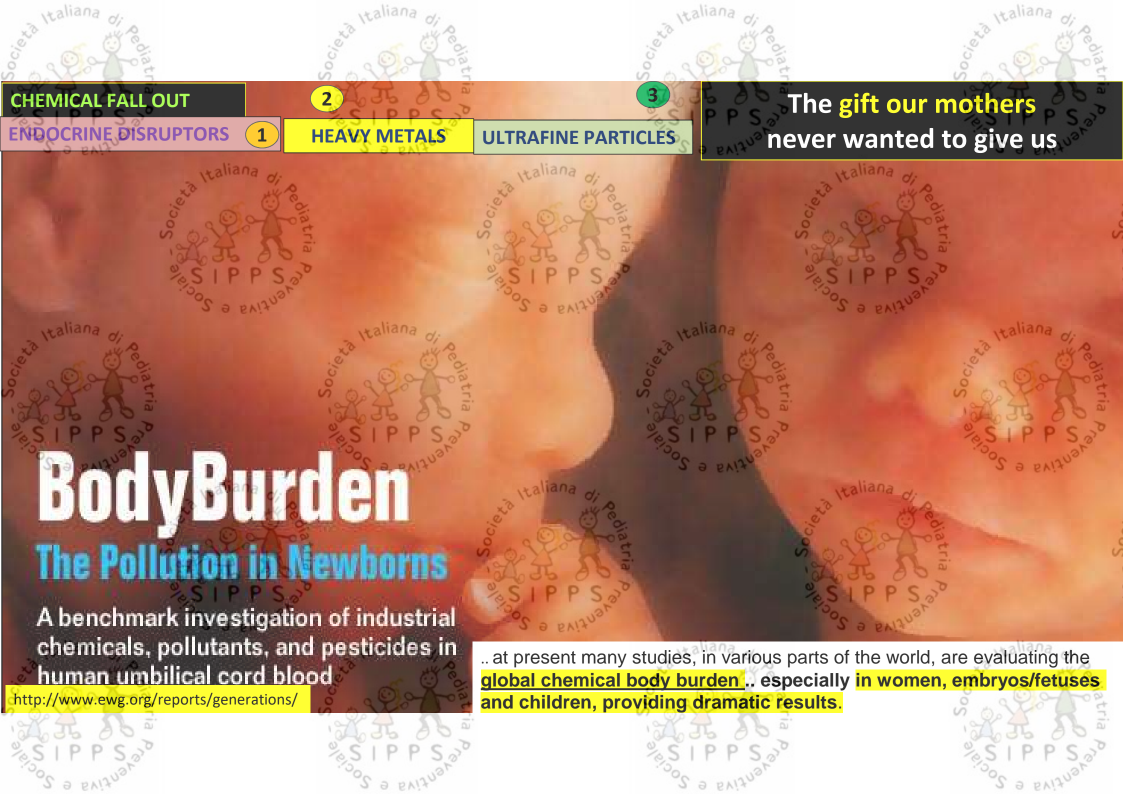
ERNESTO BURGIO

ECERI - European Cancer and Environment
Research Institute



It has been well known for many years that prenatal life is not fully protected in the uterine microenvironment. But only over the last decade we have been focusing on mechanisms and modalities of maternal and foetal exposure to an impressive range of chemicals (eg .: endocrine disruptors), physical factors (eg .:EMFs) and biological agents (eg .: viruses) able to induce potentially adaptive and predictive epigenetic changes in the embryo-fetal genome, thus interfering with the programming of tissues and organs in an often irreversible way.

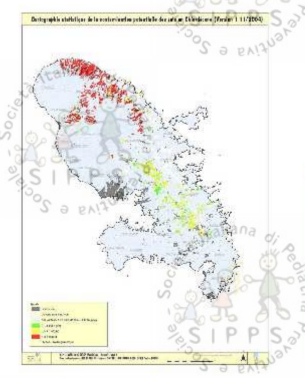




PCBs. RESULTS OF CONCERN **Monitoring Body-Burdens** 209 TESTED BDE-47 (Tetra) Test Result: 249 ppb* CCC Mean Ive HEALTH EFFECTS (SUSPECTED) - thyroid > 700 different synthetic chemicals or heavy metals Now being phased out this fire retaidant is in many products and are found in the cord blood and in the placenta. resists environmental degradation. Dieldrin Tost Result: 5,11 ppb CDC Mean: n/e - reurological NToxic Chemical Exposure A posticide once used to kill termites and other soil insects, it still lingers in **PBDEs** the environment. **POPs** p.p-DDE Test Result: 256 ppb CDC Mean: 295 ppb HEALTH EFFECTS (SUSPECTED) PESTICIDES 28 115710 A breakdown product of DET (now banned) that ingers in the body, it has health effects similar to those of the posticide. DIOXINS OFTEDTED Test Result: 34.8 ppb CDC Mean: 1.15 ppb HEALTH EFFECTS (SUSPECTED) PHTHALATES t's a member of a glass called phthalates, used to thisken letions and make plastics flexible. Marcury Tost 1: Ingestion PRODUCTS Test 2 12 mierogrems/I CDC Polioning Levet 10 Skin contact 3 DETECTED HEALTH EFFECTS METALS reurological
 reproductive Buncan's blood level of the fexic metal more than doubled after he ate two BISPHENOLS OO O DETECTED meats of swordfish and *PWRTS PER BILL ON Giuseppe Giordano











A significant, dramatic case: for some years I have been invited to

Martinique, a small paradise in the Atlantic Ocean, to investigate the

origins of the continuous increase of Cancer (in Martinique there is the

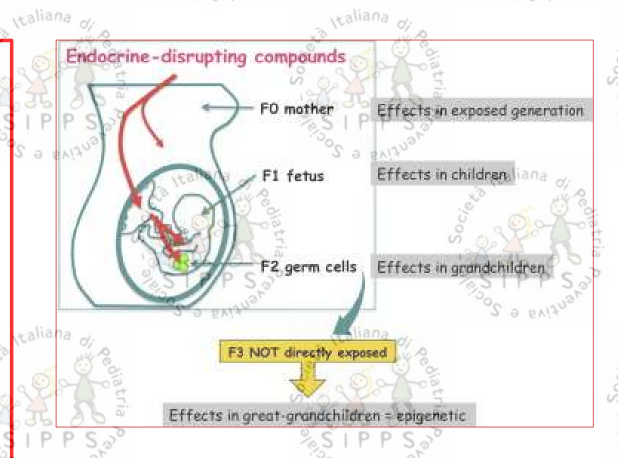
world record of prostate CA) and Autism in children...

Last year, at the last congress, I asked three questions:

SIPPS DE PAIR

Question 1

• To what extent the exposure
of moms and fetuses to
endocrine disruptors and other
epigenotoxic molecules that
interfere with fetal
programming represents
a serious threat to the health
of children and future
generations?



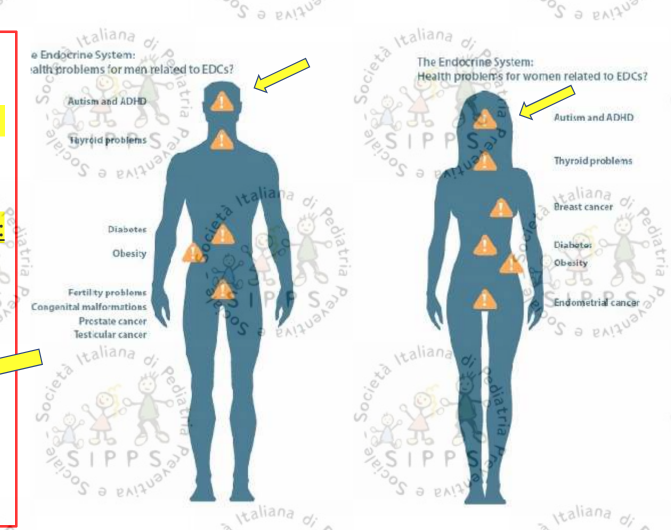
https://www.sciencedirect.com/science/article/pii/S030372071100635



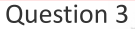
Question 2

What is the role of the ever increasing exposure of moms and fetuses to epigenotoxic molecules in the genesis of the current Epidemiological Transition:

Pandemics of obesity and juvenile diabetes 2, continuous increase in allergic and autoimmune diseases, neuro-developmental disorders, neurodegenerative diseases and cancer (especially in infants and young people)?

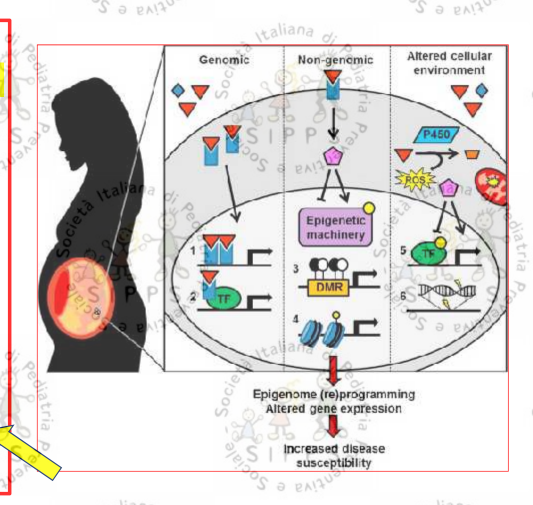


http://www.env-health.org/news/latest-news/article/health-costs-in-the-eu-how-much-is



Can we still doubt that the presence for many years of epi-genotoxic molecules such as dioxin in Seveso or Taranto and chlordecone in Martinique and Guadeloupe..

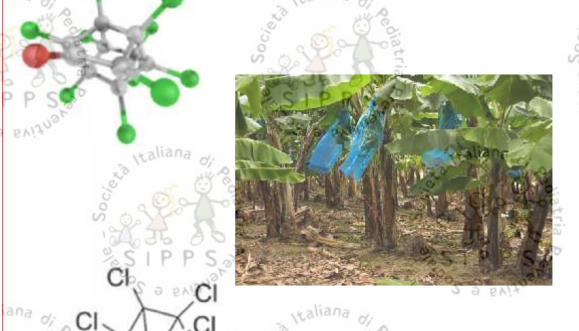
in the food chains and aquifers of a country and therefore in the organisms of young people at the age of procreating and in their gametes is a primary cause of poor fetal tissue and organ programming and thus of increasing tumors' rates (especially prostate cancer) and neurodevelopmental disorders?



https://www.sciencedirect.com/science/article/pii/S1084952115001056

• Kenone (Chlordecone) is an obsolete insecti

- Kepone (Chlordecone) is an obsolete insecticide related to Mirex and DDT: Martinique is heavily contaminated, following years of its unrestricted use in the banana plantations
- It is a known Persistent Organic Pollutant (POP), classified among the "dirty dozen": its use was so disastrous that it is now banned in the Western World by the Stockholm Convention on Persistent Organic Pollutants (2011) but only after many millions of kilograms had been produced
- Kepone bio-accumulates in animals and foodchains by factors up to a million-fold
- Workers with repeated exposure suffer severe convulsions resulting from degradation of the synaptic junctions.





https://cordis.europa.eu/result/rcn/84240_fr.html

CORDIS

Servizio Comunitario di Informazione in materia di Ricerca e Sviluppo

ommissione europea > CORDIS > Progetti e risultat > La placenta trasferisce i pesticici al feto...



ACTUALITÉS ET ÉVÈNEMENTS PROJETS

PROJETS ET RESULTATS

MAGAZINES RESEARCH*E

PLUTOCRACY - Résultat en bref

Project ID: QLK4-CT-2000-00279 taliana

Financé au titre de: FP5-LIFE QUALITY

Le placenta transmet les pesticides au fœtus

L'incidence des allergies comme l'asthme à augmenté au cours des dernières décennies. Dans le cadre des efforts menés pour en trouver la raison, les scientifiques ont étudié le transport des composés chimiques à travers le placenta, du milieu environnant vers le fœtus.

This is an official website of the European Community that lists many studies related to the problem of maternal-fetal exposure to pollutants and toxics (in particular to pesticides): scientists found that all xenobiotics cross the placental barrier by passive diffusion and reach the fetus..... In the main fetal organs (especially in the blood, spleen, bone marrow, brain and liver) the concentration of these pesticides is higher than in the corresponding maternal organs. The implications are of great significance: the accumulation of these compounds in the fetal tissues will have an impact on the development of the child's immune and nervous systems

msilieux des xemobioliques, la predisposition aux allergies.



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PL



In fact placental alterations are more and more frequent

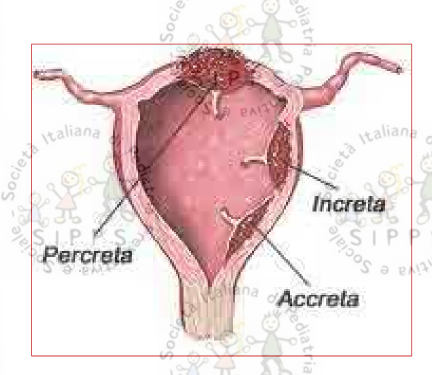
The *placenta accreta* is an insertion/invasion of/by the placenta **into maternal tissues**: there are three types according to the **insertion depth into the endo/myometrium**

- the proper placenta accreta : the villi penetrate more or less deeply into the myometrium;
- the placenta increta: the villi invade the whole myometrium;
- the placenta *percreta*: the villi go beyond the myometrium, sometimes invading neighboring organs (bladder) ...

... it is, in fact, as if the (immunological) mechanisms of maternal-fetal tolerance were weakening!

... we must not forget that the placenta is largely an embryofetal organ (that the embryo himself produces to connect to the mother to get oxygen, nutrition, information... certainly not to invade her)

(evolutionary mechanisms that are millions of years old)



Choriocarcinoma

even more common all over the world has become prematurity (today one child out of 10 is born prematurely ... which represents an increase of 30% over the last 35 years) .. another symptom of growing maternal-fetal intolerance that should not be underestimated..

L'INSERM today defines different stages of prematurity: extremely preterm (less than 28 weeks) very preterm (28 to 32 weeks) moderate to late preterm (32 to 37 weeks).

Epidémiologie [modifier | modifier le code]

En 2012, plus d'un bébé sur dix naît prématurément dans le monde sans évidence de décroissance avec le temps

Les naissances prématurées concernent 11 à 13 % des naissances aux États-Unis, soit près du double du taux des autres pays industrialisés et une augmentation de 30 % par rapport à 1981'. Plus du quart des décès néonataux seraient la conséquence de la prématurité 8.

Les données sont probablement assez solides et permettent d'avoir aujourd'hui un aperçu évolutif concernant les trois dernières décennies en France.

Évolution des taux d'incidence de la prématurité en France

S S S	1972	1981	1988	1995	2003
Très grande prématurité (de 22 à 27 SA)	a , p	-	-	0,4 %	0,5 %
Grande prématurité (de 28 à 32 SA)	1,3 %		1 %	1,2 %	1,3 %
Prématurité (de 33 à 37 SA)	8,2 %	5,7 %	4,8 %	5,9 %	7,2 %

L'incidence est donc en augmentation, ce que confirme les chiffres d'autres pays, en particulier américains '





American Journal of Epidemiology

Chlordecone Exposure, Length of Gestation, and Risk of Preterm Birth

Philippe Kadhel ➡, Christine Monfort, Nathalie Costet, Florence Rouget, Jean-Pierre Thomé, Luc Multigner, Sylvaine Cordier

American Journal of Epidemiology, Volume 179, Issue 5, 1 March 2014, Pages 536–544, https://doi.org/10.1093/aje/kwt313

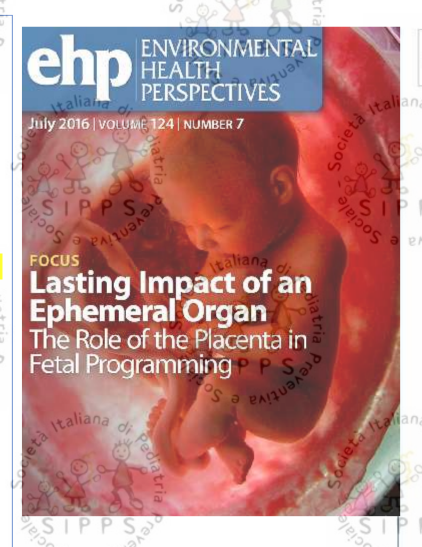


Volume 179, Issue 5

1 March 2014

Chlordecone is an organochlorine pesticide that has been widely used ... in the French West Indies. Data from the *Timoun Mother-Child Cohort Study* conducted in Guadeloupe between 2004 and 2007 examined combinations of chlordecone concentrations in maternal plasma with gestational duration and preterm birth rate in 818 pregnant women ... 1-log10 increase in chlordecone concentration was associated with decreased duration of pregnancy (-0.27 weeks, 95% confidence interval: -0.50, -0.03) and increased risk premature labor (60%; 130). ... These results are relevant to public health because of the prolonged persistence of Chlordecone in the environment and the high rate of preterm birth in this population.

In such a context, the organ that acquires a truly extraordinary importance is the **PLACENTA**: an organ that has been **poorly** studied until a few years ago and that emerges as a sort of "Black Box" for epigenetically programming fetal tissues and organs





HHS Public Access

Author manuscript

Am J Obsert Gynecol Abthor manuscript; available in PMC 2016 October 01

ublished in final edited form as

Am J Obstet Gynecol. 2015 October; 213(40): S14-S20. doi: 10.1016/j.ajog.2015.08.030.

THE PLACENTA IS THE CENTER OF THE CHRONIC DISEASE UNIVERSE

Kent L. Thornburg 1,2,3 and Nicole Marshall 2,3

Department of Medicine, School of Medicine, Oregon Health & Science University Portland, Oregon 97239

²Knight Cardiovascular Institute, Center for Developmental Health, School of Medicine, Oregon Health & Science University Portland, Oregon 97239

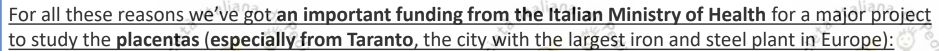
³Department of Obstetrics & Gynecology, Oregon Health & Science University Portland, Oregon 97239

Abstract

Over the past quarter century it has become clear that adult onset chronic diseases like heart disease and type 2 diabetes have their roots in early development. The report by David Barker and colleagues showing an inverse relationship between birthweight and mortality from ischemic heart disease was the first clear-cut demonstration of fetal programming. Because fetal growth depend: upon the placental capacity to transport nutrients from maternal blood, it has been a suspected . causative agent since the original Barker reports. Epidemiological studies have shown that placental size and shape have powerful associations with offspring disease. More recent studies have shown that maternal phenotypic characteristics, such as body mass index and height, interact ental size and shape to predict disease with much more precision than does birthweight Jone. For example, among people in the Helsinky Birth Cohort, who were born during 1924-1944, the risk for acquiring colorectal cancer increased as the placental surface became longer and more oval. Among people in whom the difference between the length and breadth of the surface exceeded 6 cm, the hazard ratio for the cancer was 2.3 (95% CF 1.2-4.7, p=0.003) compared with those in whom there was no difference. Among Finnish men, the hazard ratio for coronary heart disease was 1.07 (1.02-1.13, P =0.01) per 1% increase in the placental weight/hirthweight ratio. Thus, it appears that the ratio of birthweight to place that weight, known as placental efficiency, predicts cardiovascular risk as well. Babies born with placentas at the extremes of efficiency are more vulnerable for adult onset chronic diseases. Recent evidence suggests that placental growth patterns are sex specific. Boys, placentas are, in general, more efficient than those made by girls 2, 2111 Another recent discovery is that the size, shape and efficiencies of the placenta can change only years of time with very narrow confidence limits. This suggests that the growth of the placenta within a population of women is strongly affected by their nutritional environment. Even though it

19/3	PROGRAMMA CEM 2017— PROGETTI ESECUTIVI IN ORD VALUTAZIONE	INE DECRESCENT	EDIPU	N TEGGIO DI
N.	OJOTIT FING 6 2	PARTNER	ÎD	IMPORTO
1	URBAN HEALTH: BUONE PRATICHE PER LA VALUTAZIONE DI IMPALIO SULLA SALUTE DEGLI INTERVENTI DI RIQUATIFICAZIONE E RIGENERAZIONE URBANA E AMBIENTALE	LOMBARDIA	4	€ 450.000,00
2	SCEGLIERE LE PRIORITÀ DI SALUTE E SELEZIONARE GLI INTERVENTI EFFICACI PER PREVENIRE IL CARIÇO DELLE MALATTIE CRONICHE NON TRASMISSIBILI	PIEMONTE	6	€ 449.250,00
3	SVILUPPO E VALIDAZIONE DI UN SISTEMA DI MONTTORAGGIO EPIDEMIOLOGICO DELLE DEMENZE BASATO SUI DATI DEI SISTEMI INFORMATIVI SANITARI	campania	5	€ 450.000,00
4000	AMBIENTE, PROGRAMMAZIONE EPIGENETICA FETALE E PREVENZIONE DELLE PATOLOGIE CRONICHE	SARDEGNA	9 6	€ 448.000,00





- Mass spectrometry (IZS Bologna)
- Immunohistochemistry (University of Cagliari)
- *Epigenetics* (University of Pisa)
- Mitochondria (University of Milan)
- Metabolomics (University of Cagliari)
- follow-up of children at risk by the Italian Federation of Pediatricians (FIMP): early diagnosis!! personalized treatment!

But most importantly, it is becoming increasingly obvious that the most serious consequences of the increasing embryo-foetal exposure to toxics will become evident after decades (and sometimes only in the following generations)

Conséquences à long terme

(reconnaissables dans les Le tableau ci-dessous offre une vision gl premières années de la vie)

Données générales chez les nourrissons de moins de 32 SA et/ou moins de 1 500 a (en %)

	Séquelles majeures	ures Séquelles mineures			
Psychomotrices	17 °S & ENITY	28	45		
visuelles o	2	26 Ataliana	28		
Respiratoires	B .	26	270.		
Langage	20	20 0	40		
Auditive	2	1 300000	6		

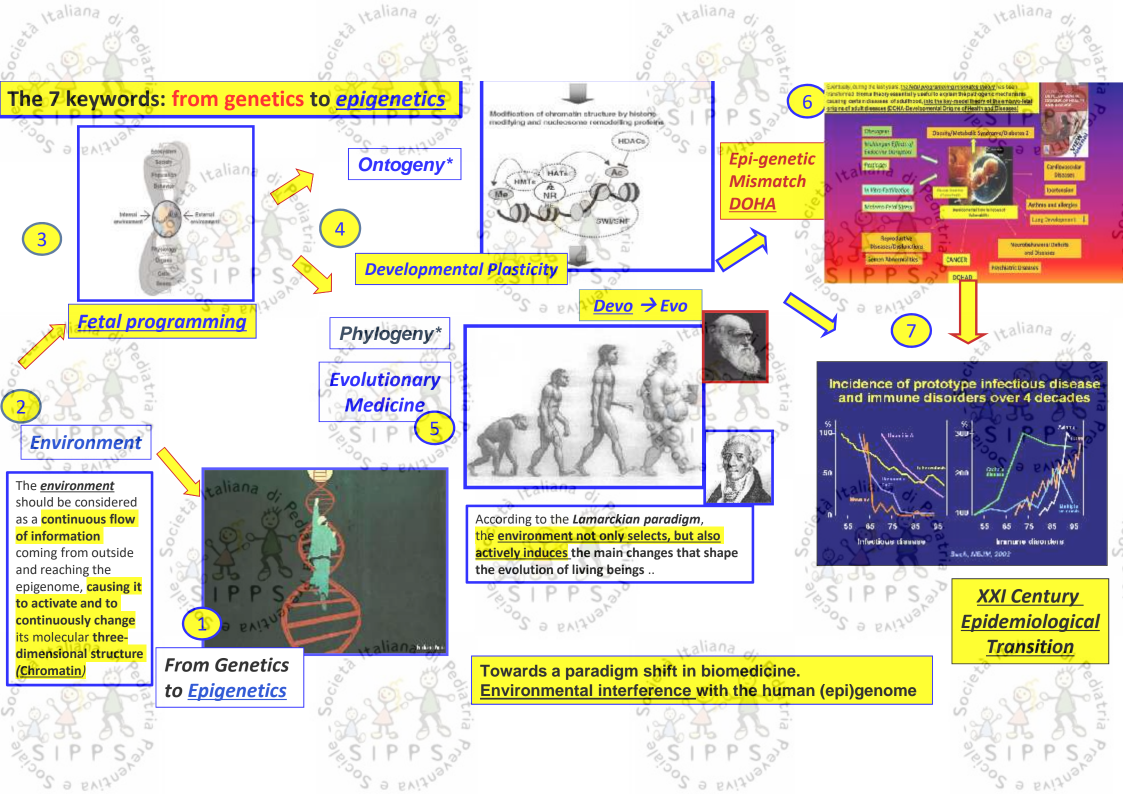


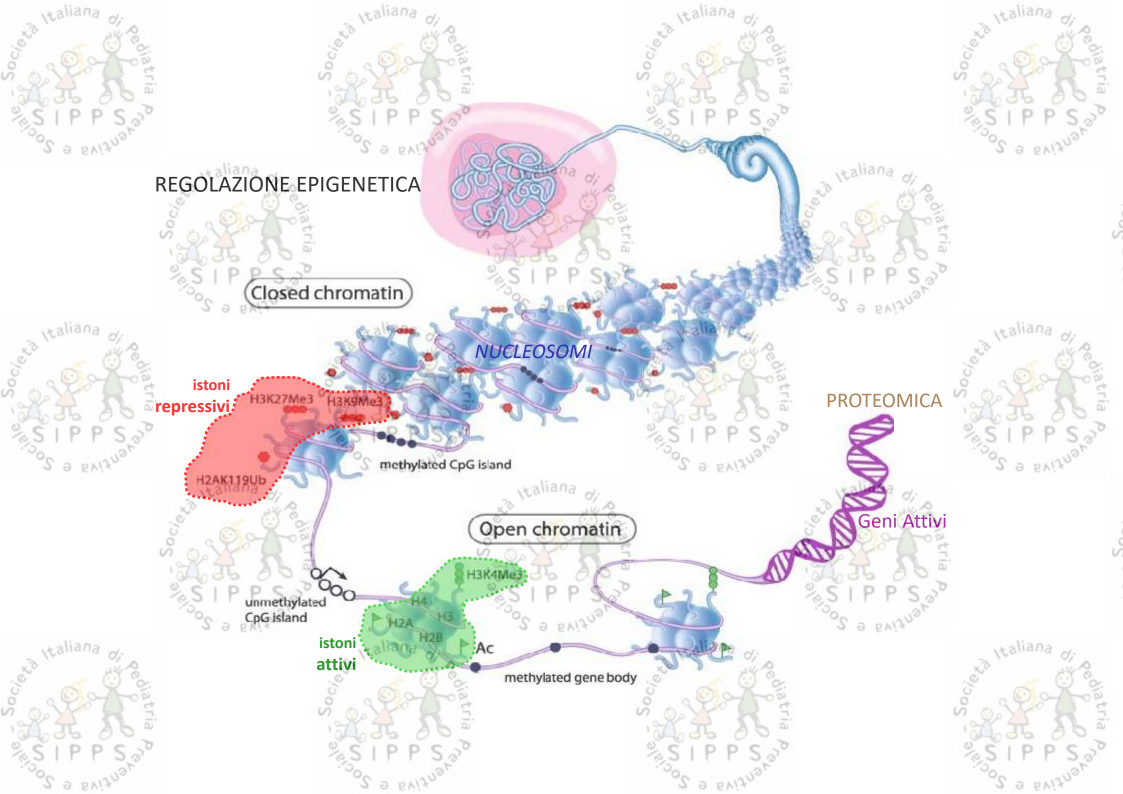
Les données de l'étude épidémiologique française ÉPIPAGE sur les petits âges gestationnels permettent de déceler un lien évident entre la survenue d'un handicap et l'importanç de la prématurité. Près de 40 % des grands prématurés présentent des séquelles - troubles moteurs, sensoriels ou cognitifs - à l'áge de 5 ans, sévères dans 5 % des cas, modérées pour 9 % des enfants, légères pour les autres 22. Ces données sont cohérentes avec celles issues d'autres études d'autres pays 2

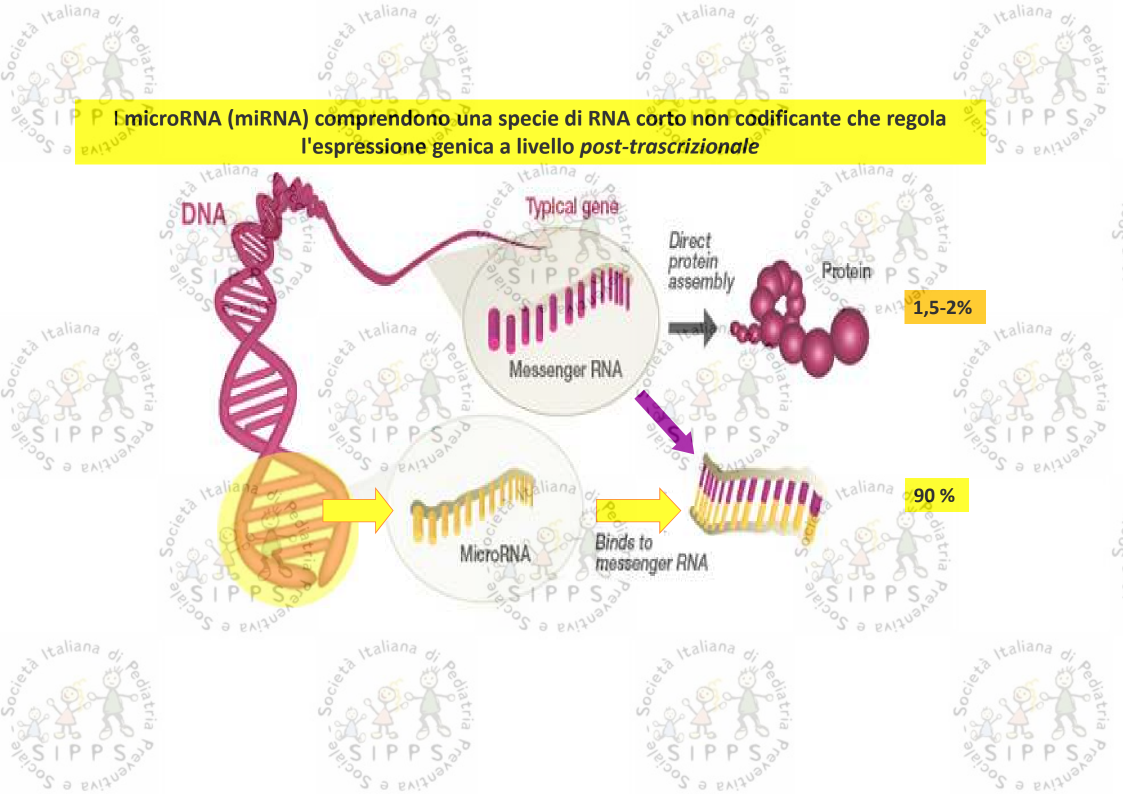
The Barker Hypothesis Fetal Origins of Adult Disease

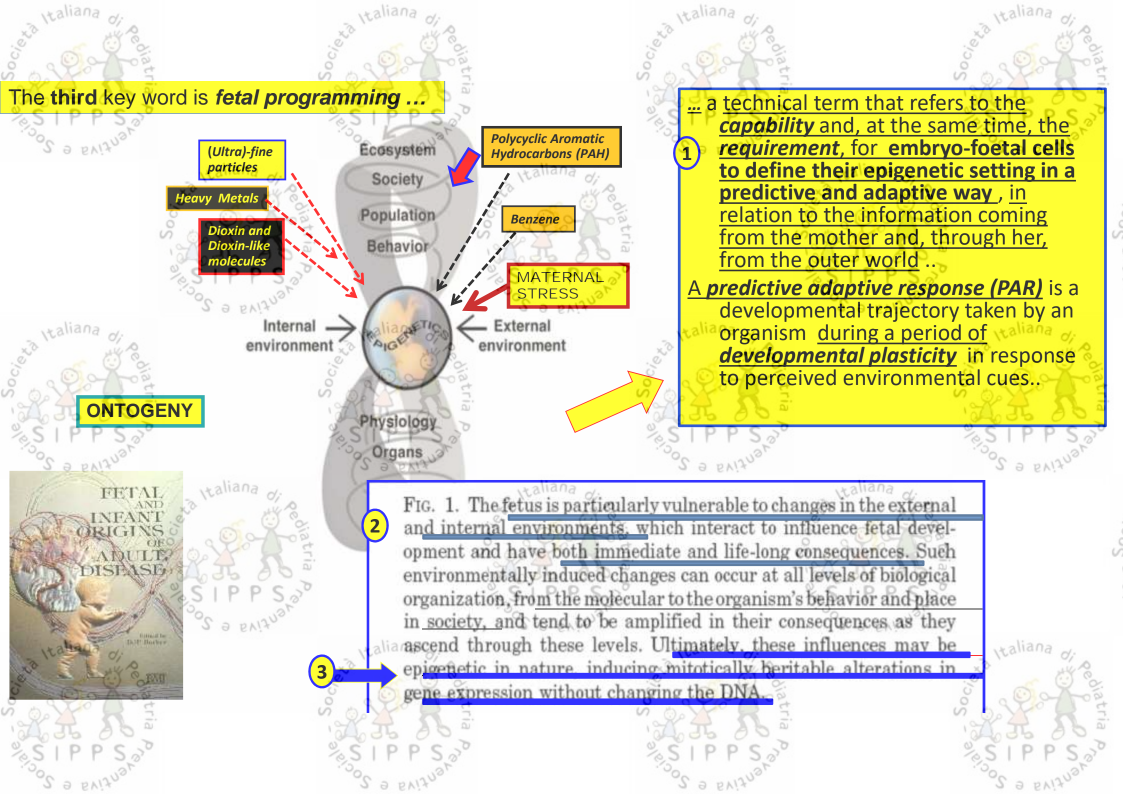
Adverse intrauterine events permanently "program" postnatal structure/function/homeostasis "Adapted Birth Phenotype" Better chance of fetal survival Increased risk of adult disease

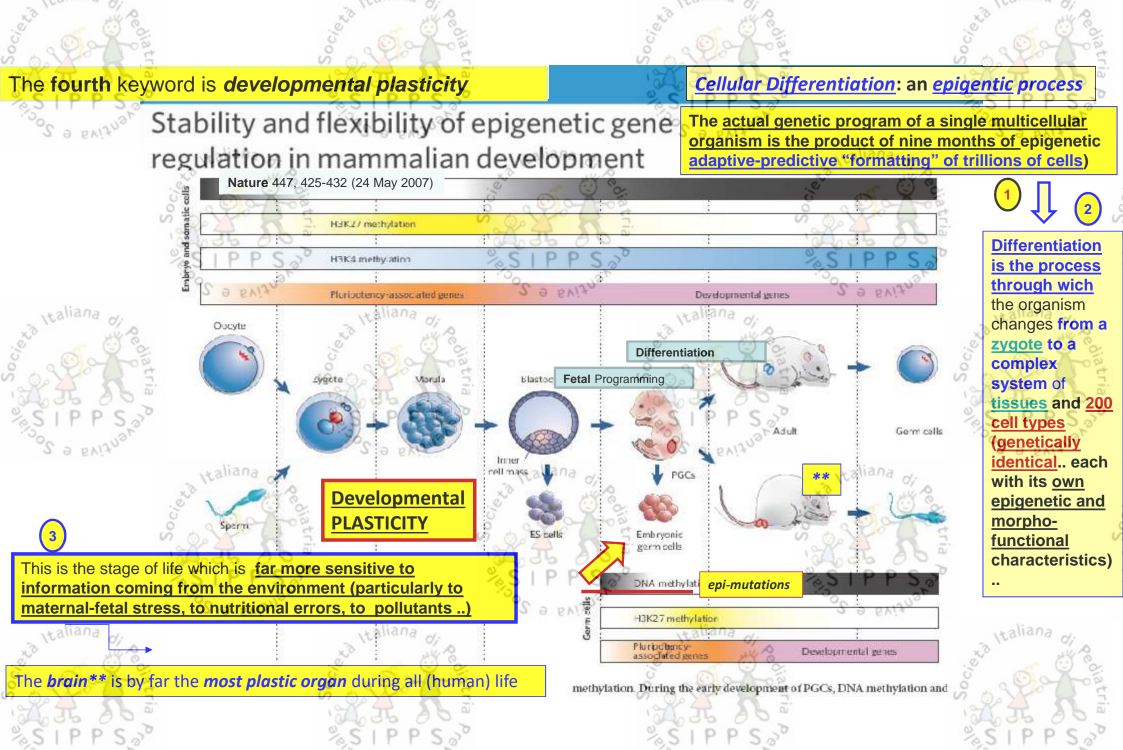
.. since every intrauterine adverse events might interfere permanently with the epigenetic programming of organs and tissues (DOHaD theory)

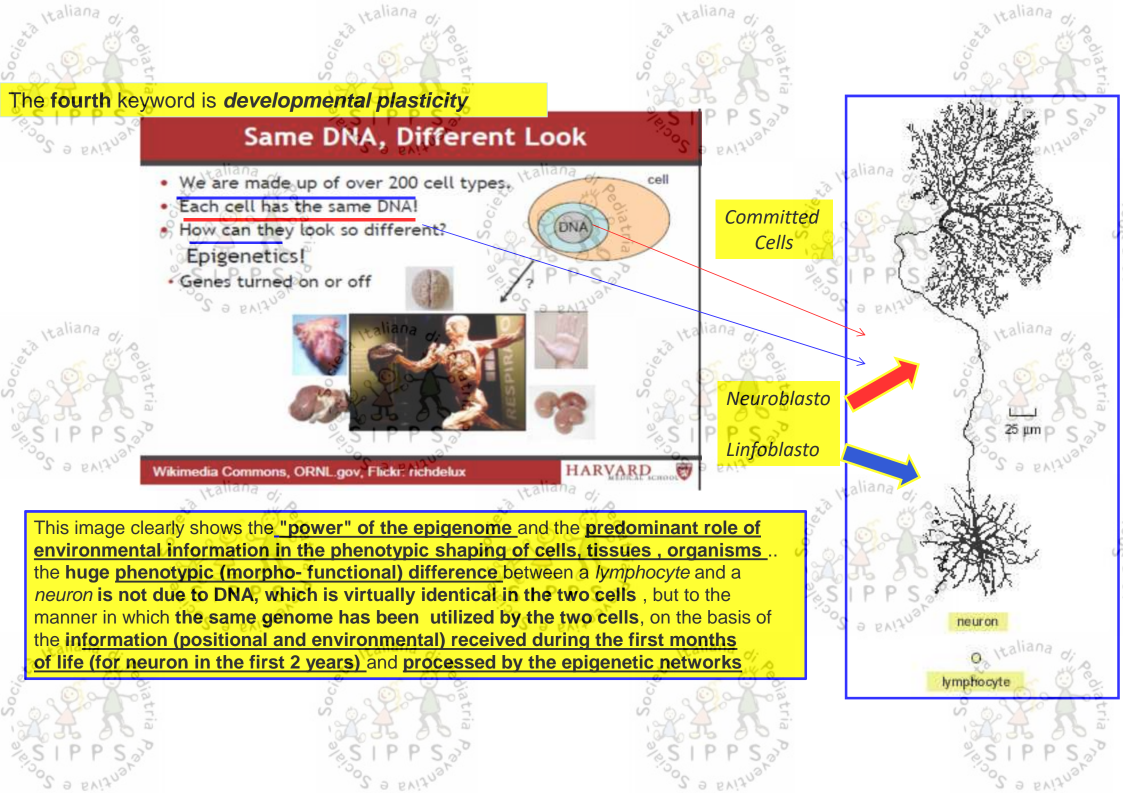








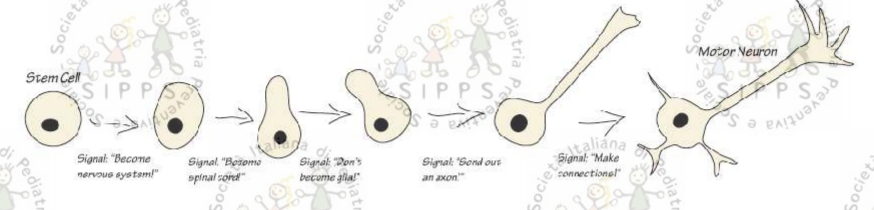


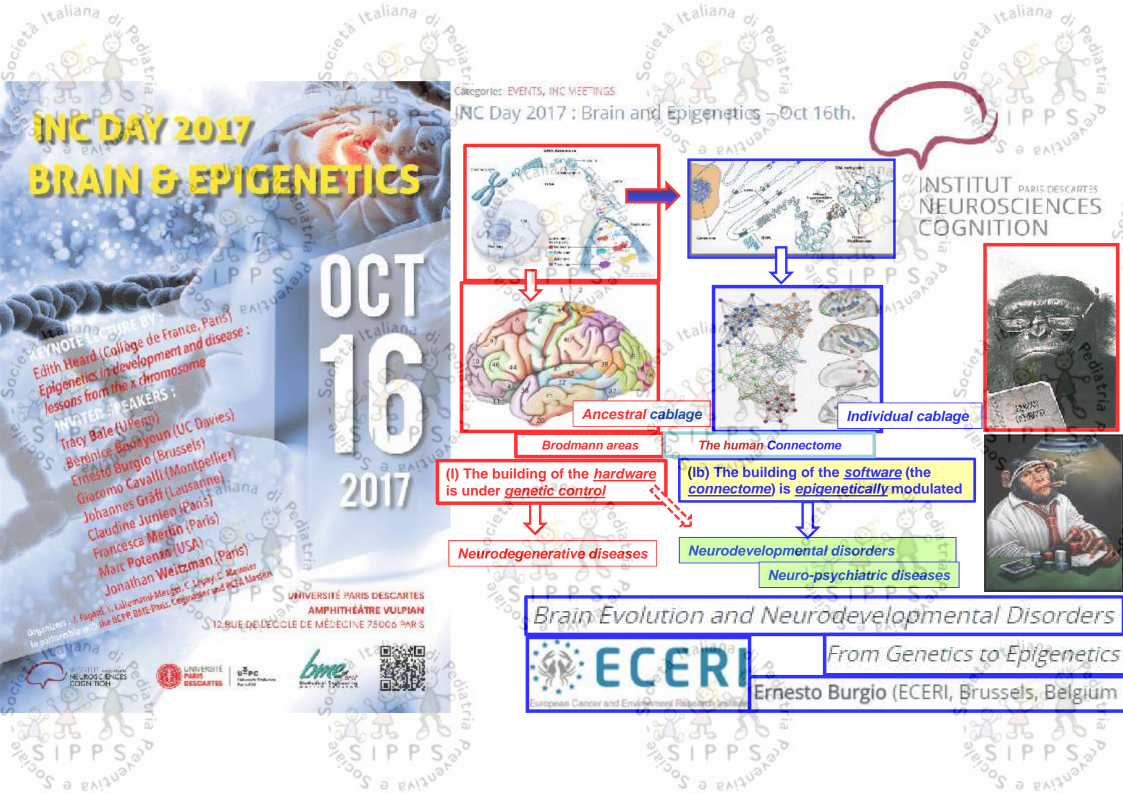


http://learn.genetics.utah.edu/content/epigenetics/epi_learns/

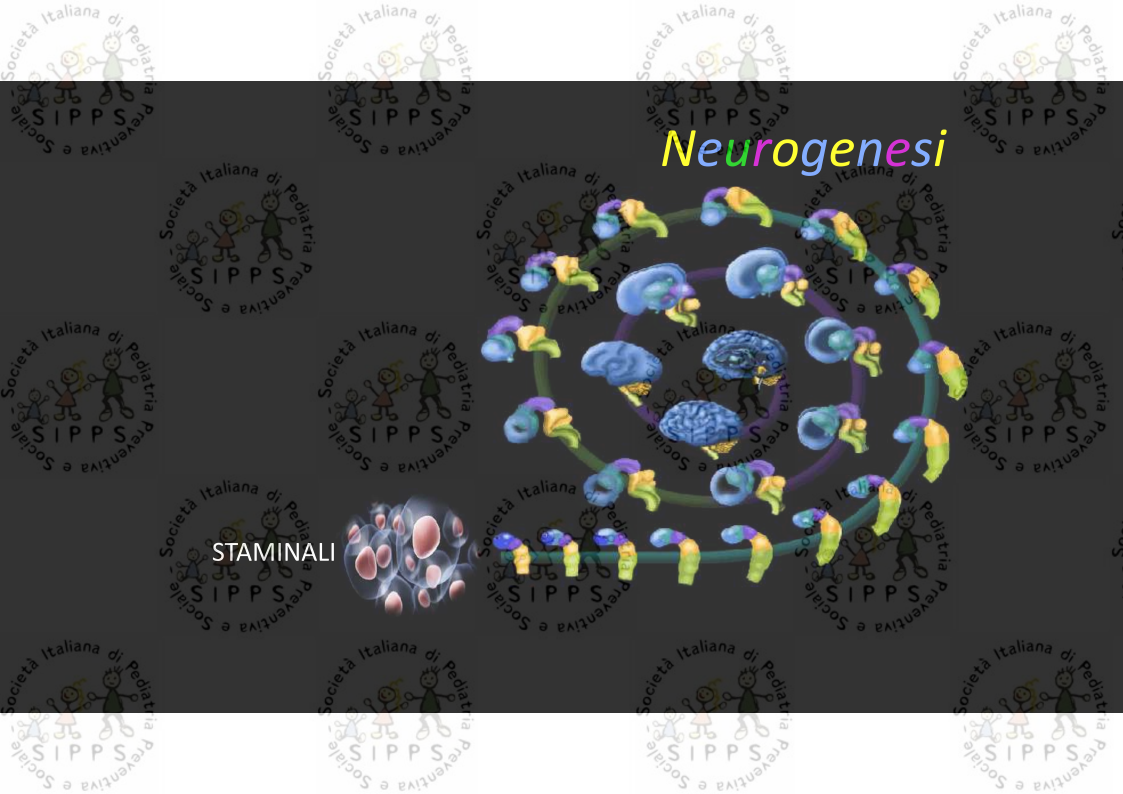
The Epigenome learns from its experiences

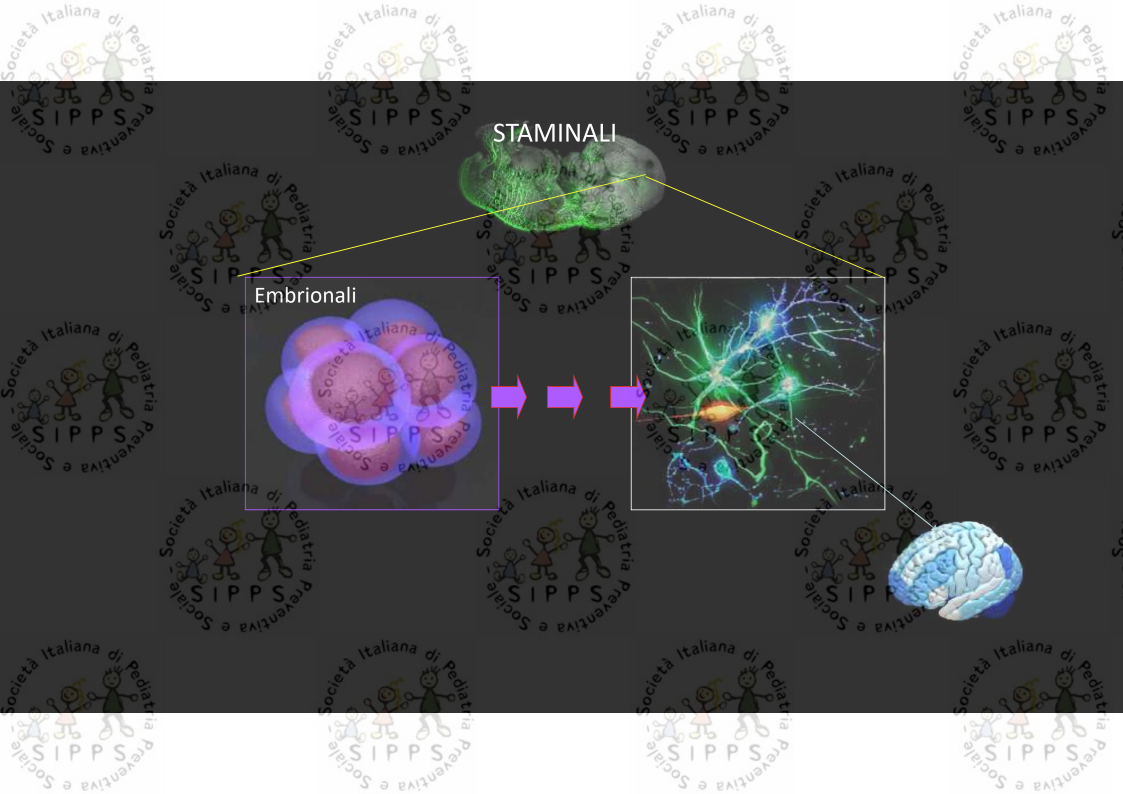
- Epigenetic tags act as a kind of cellular memory.
- A cell's epigenetic profile -- a collection of tags that tell genes whether to be on or off
 -- is the sum of the signals it has received during its lifetime

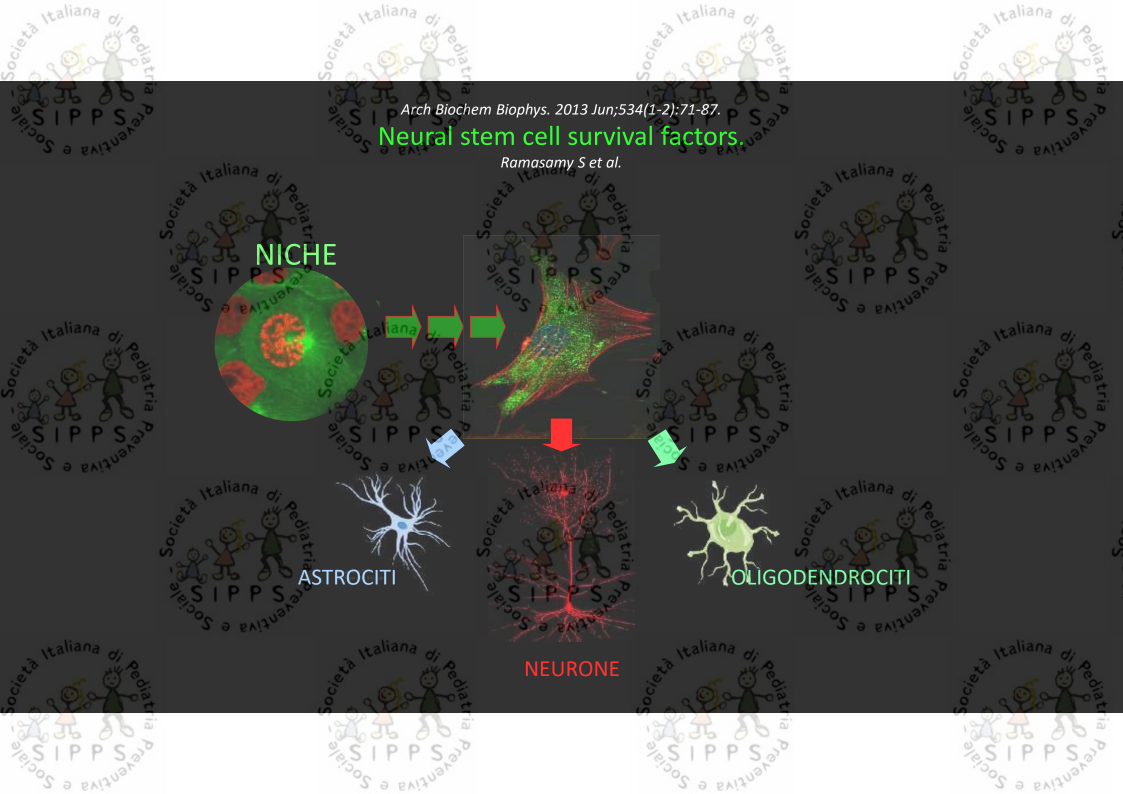


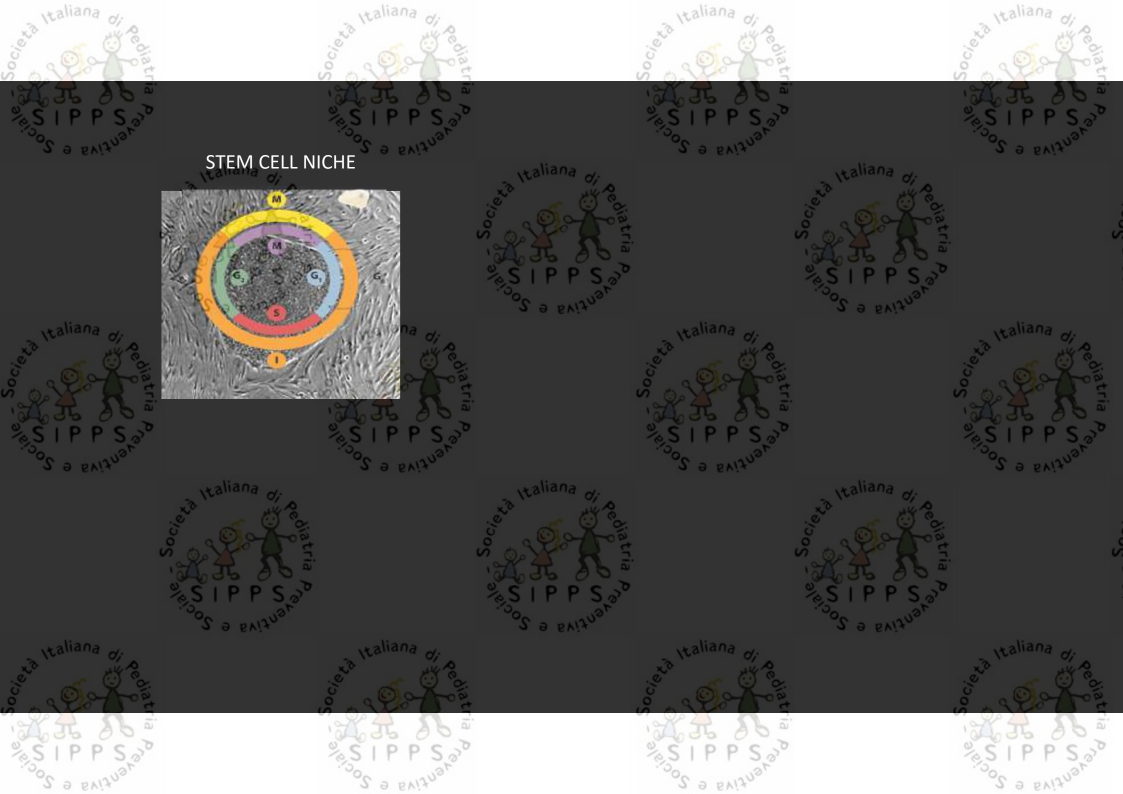






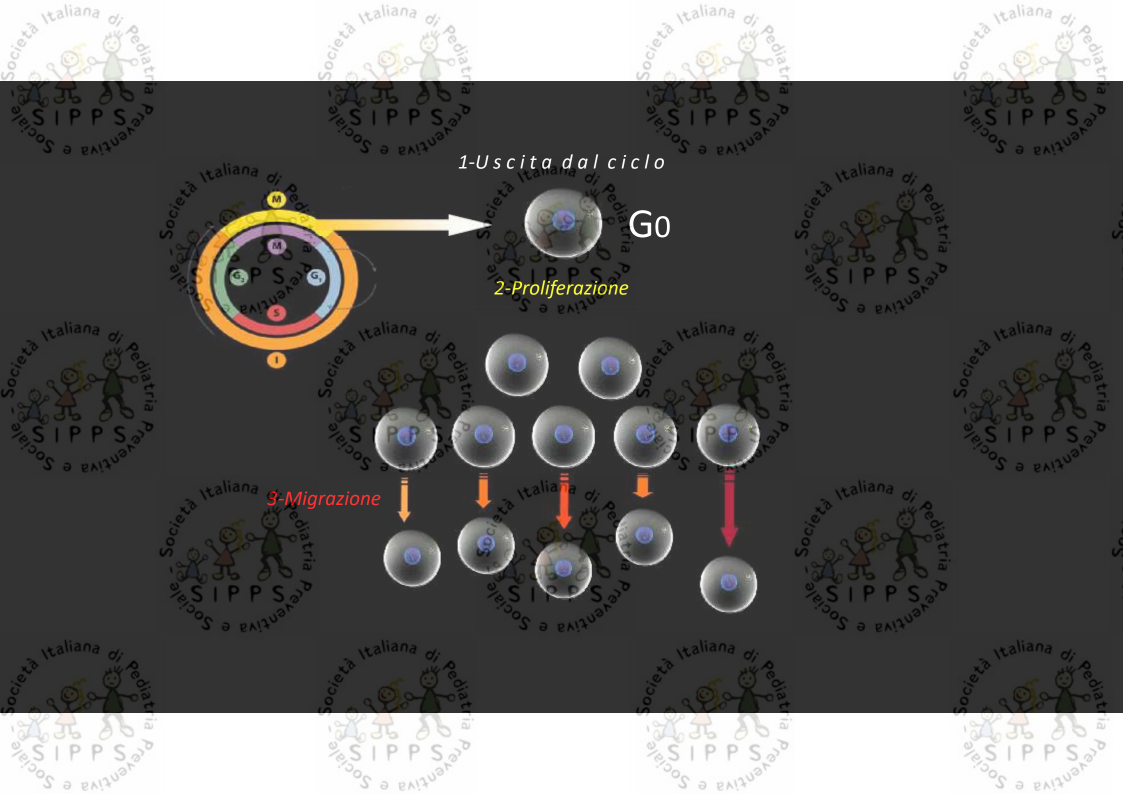


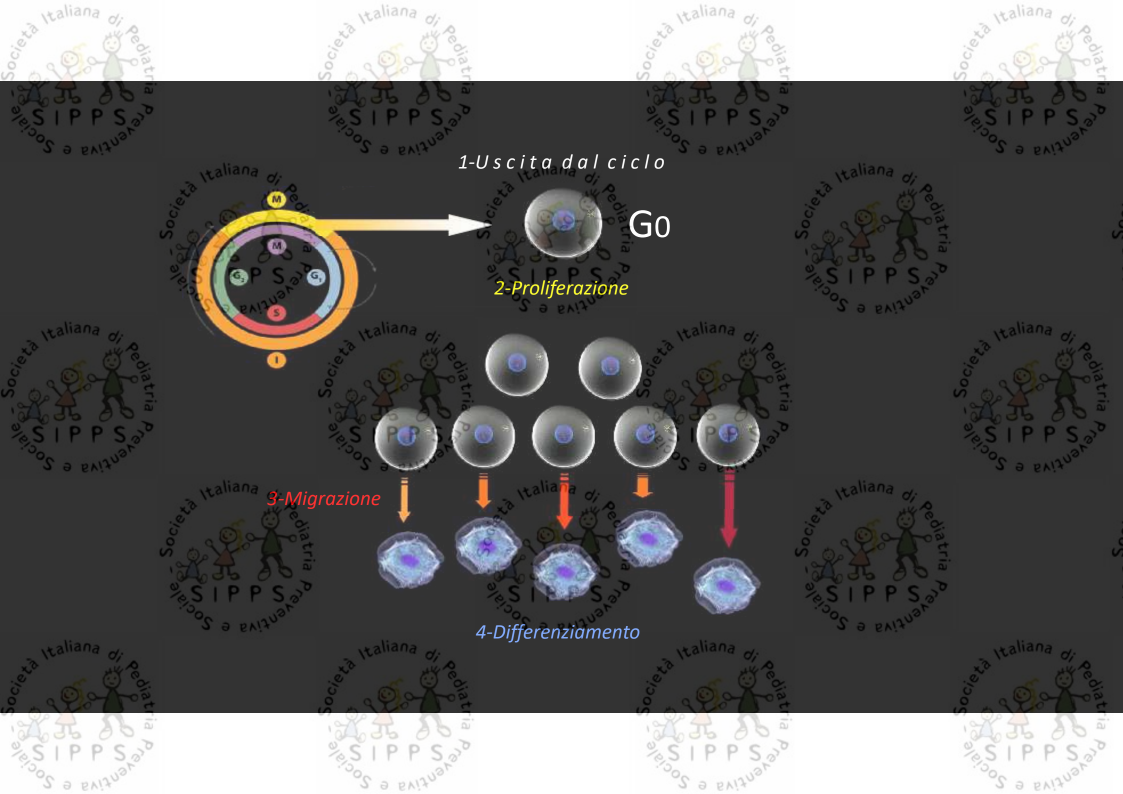


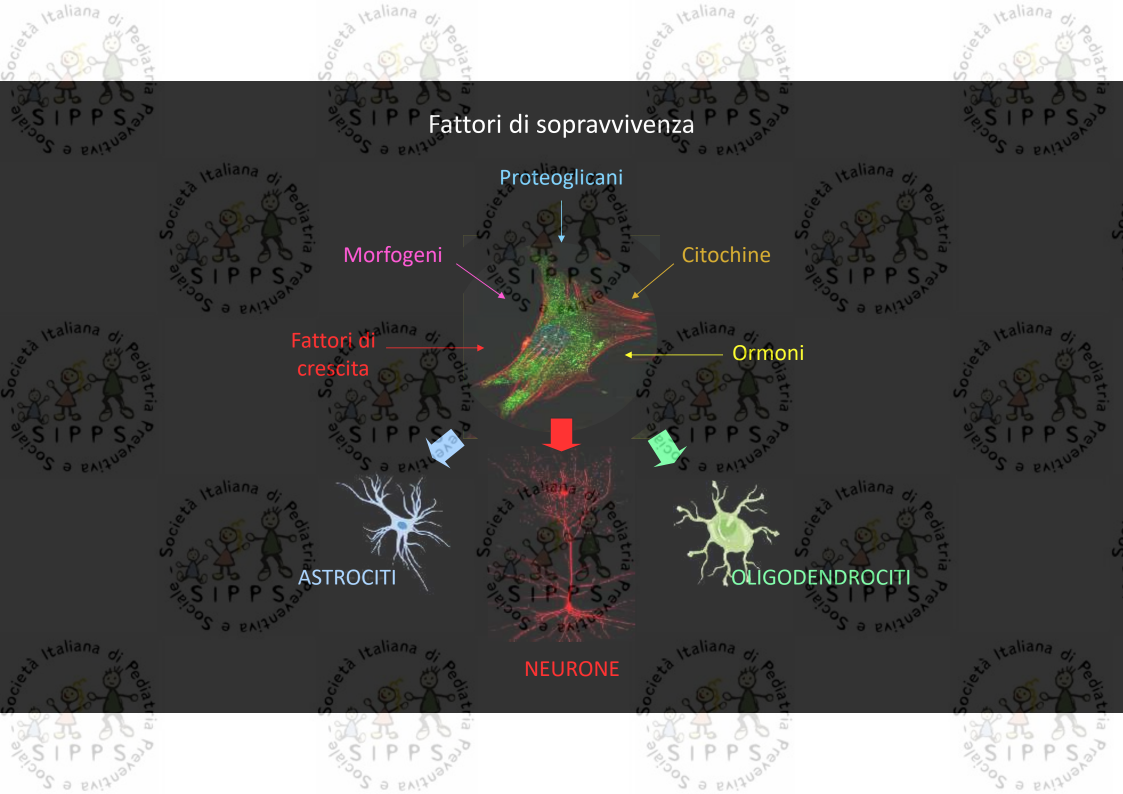


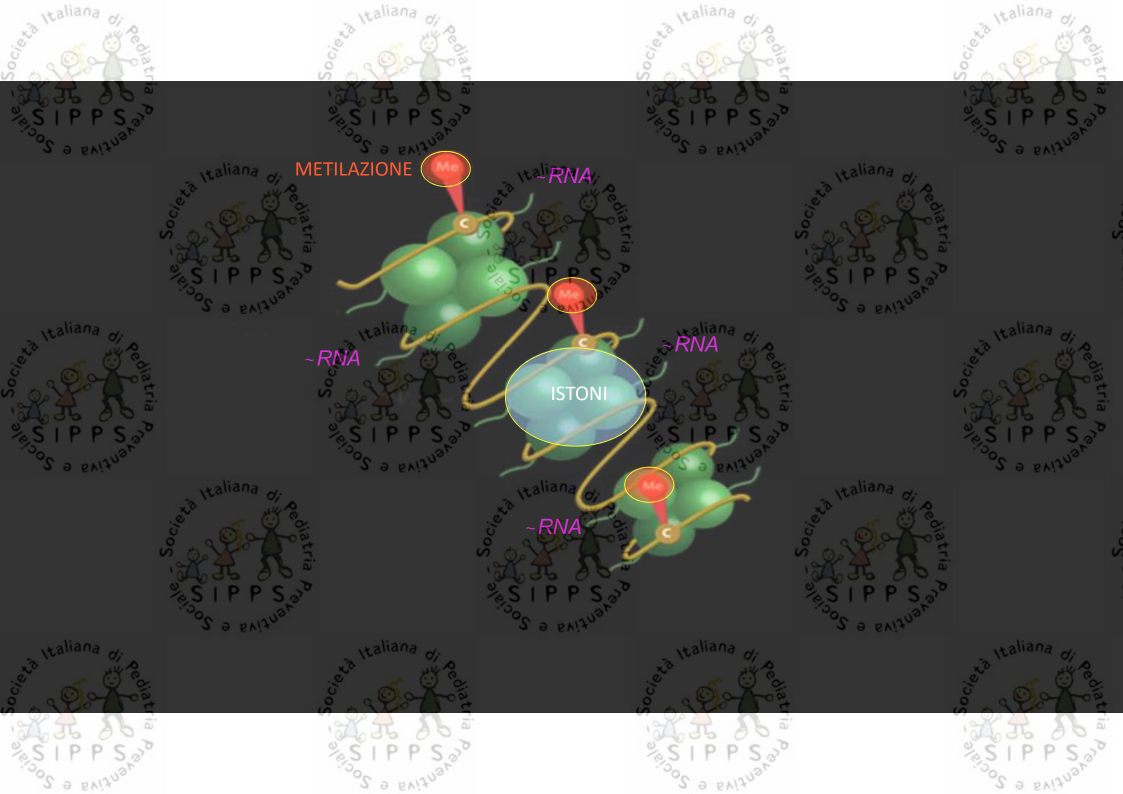


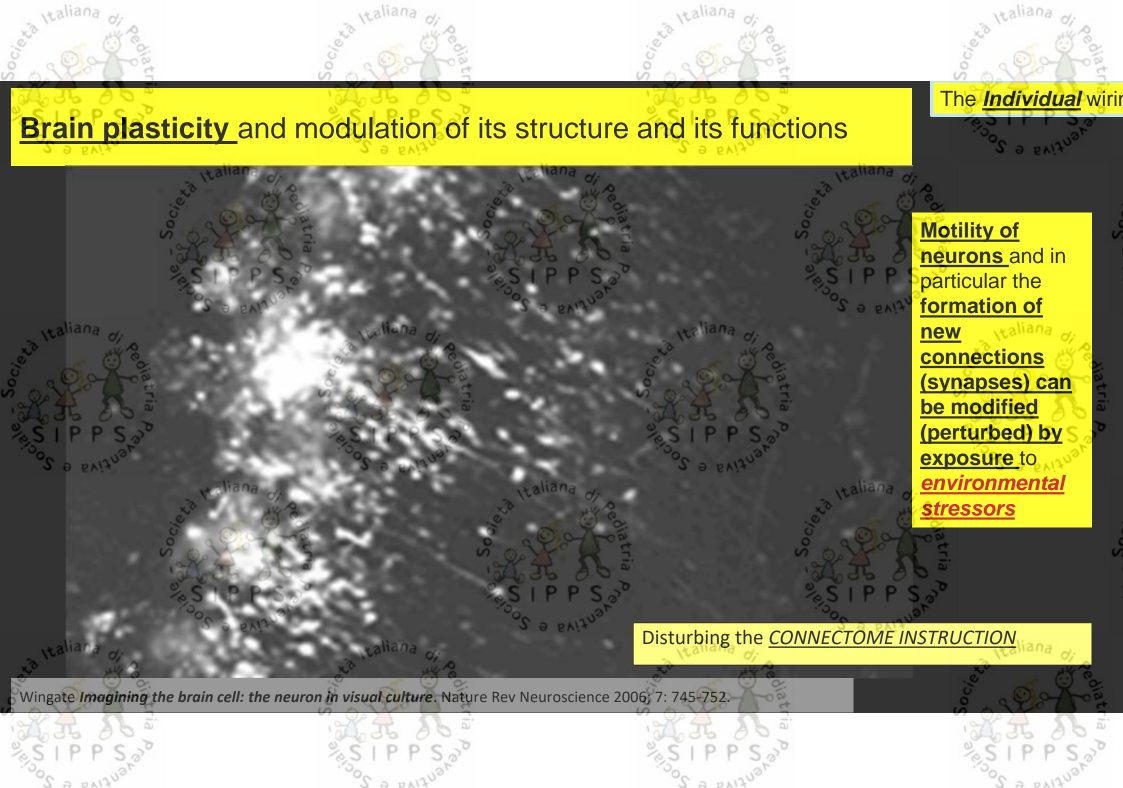


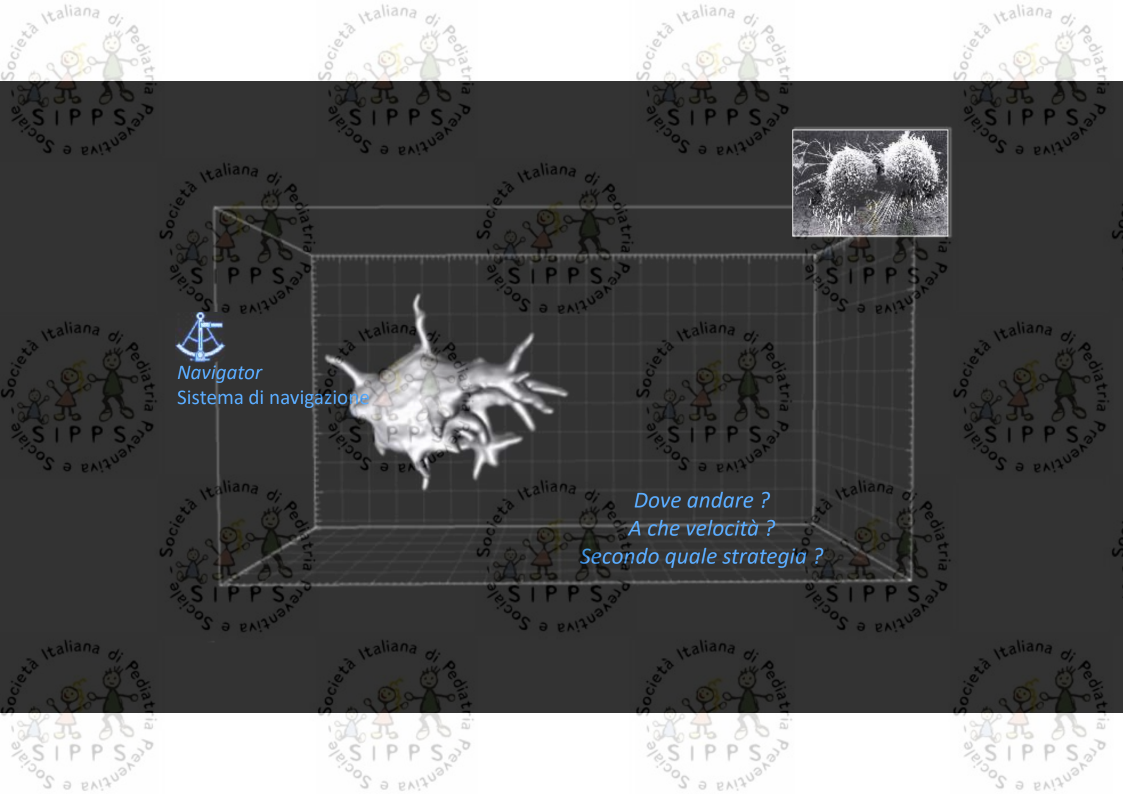


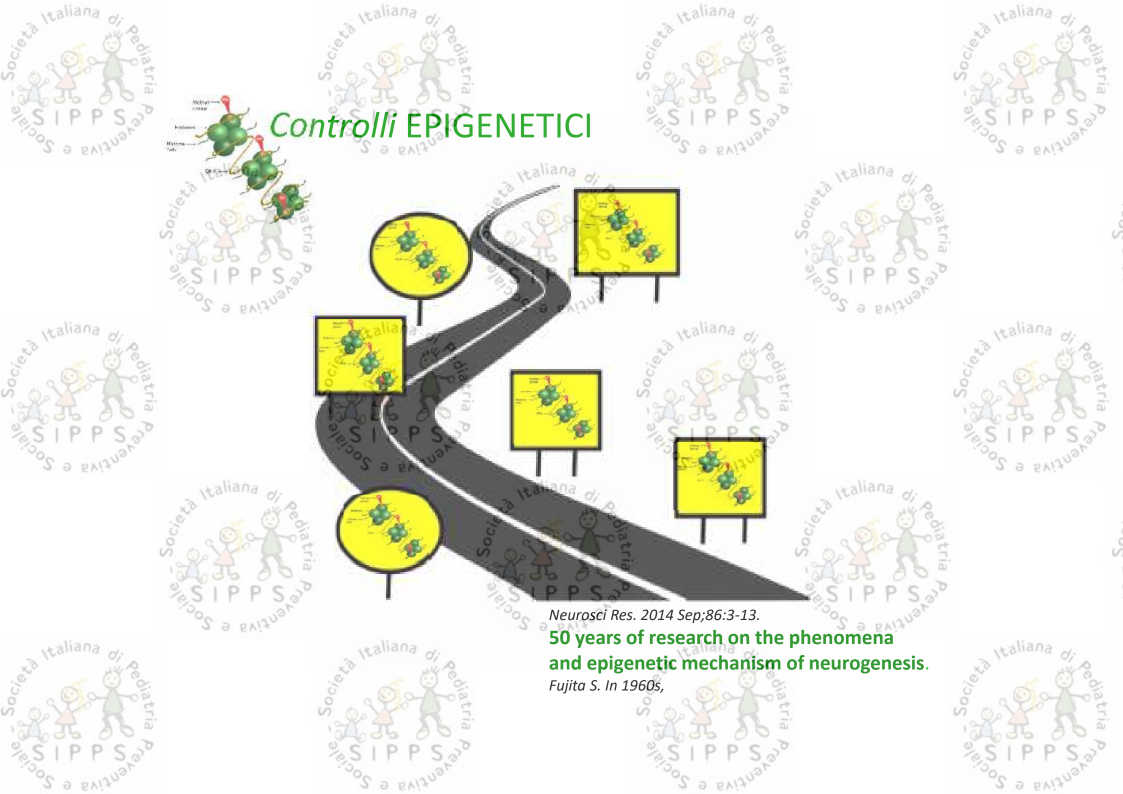


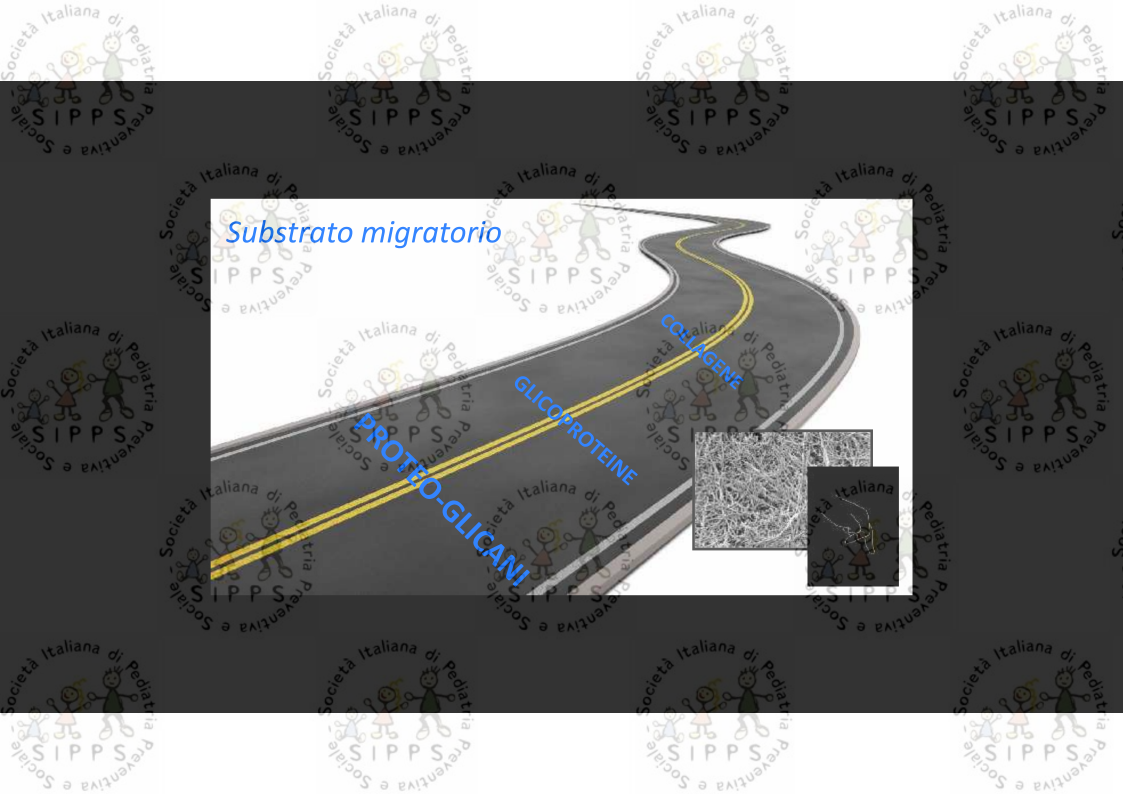


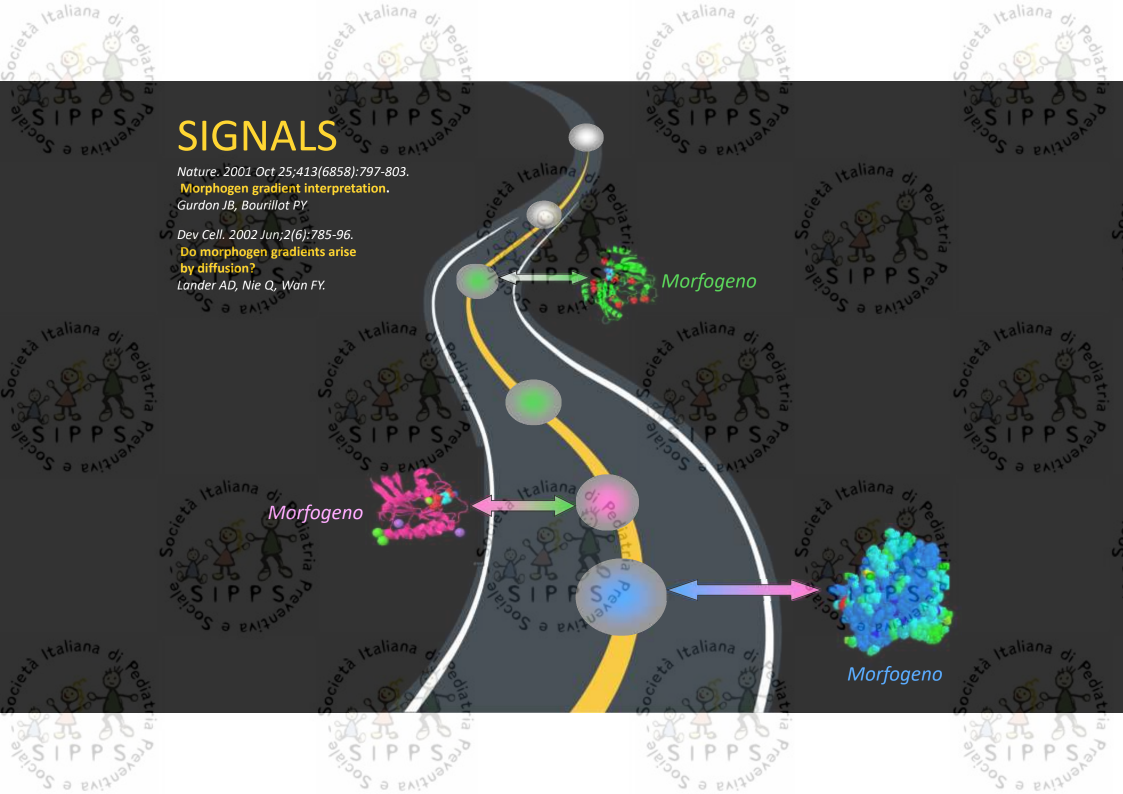




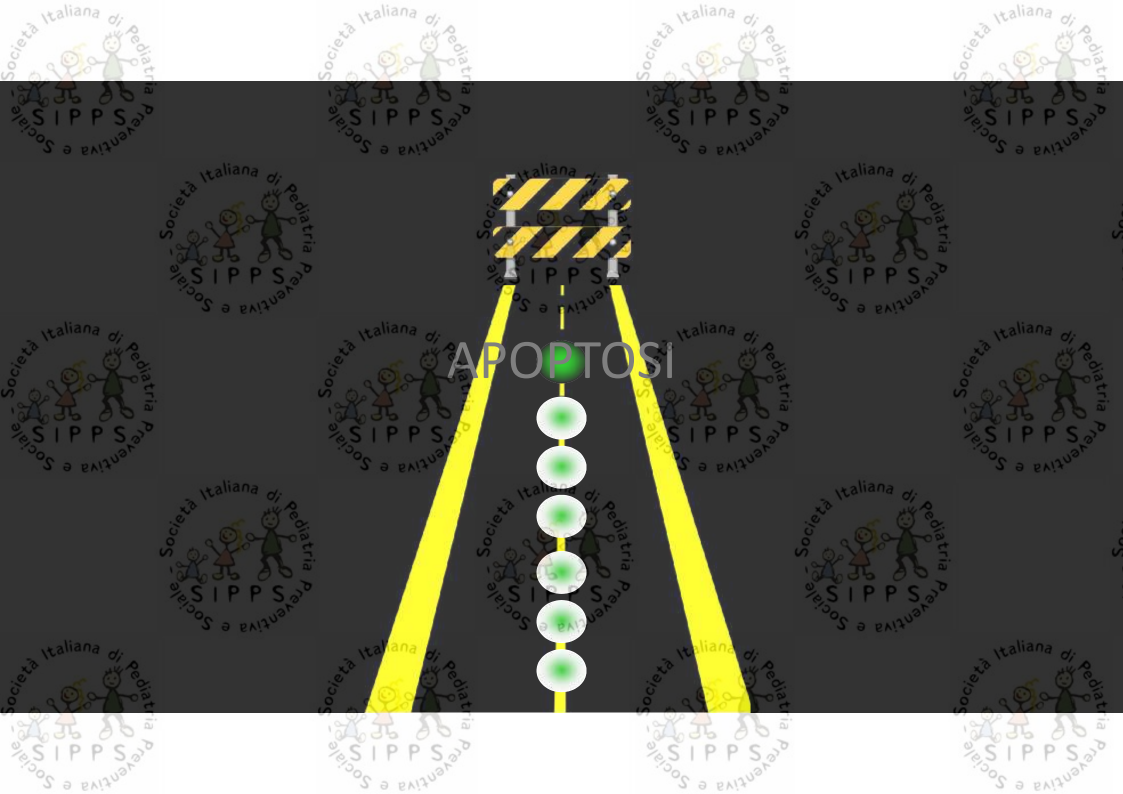


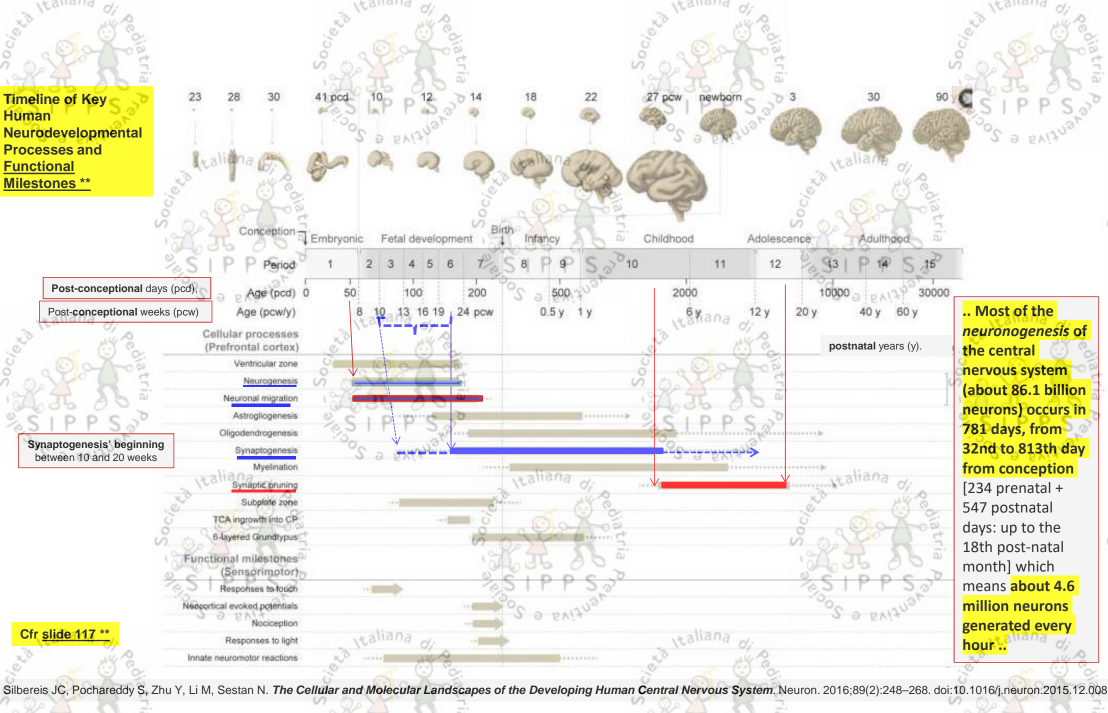










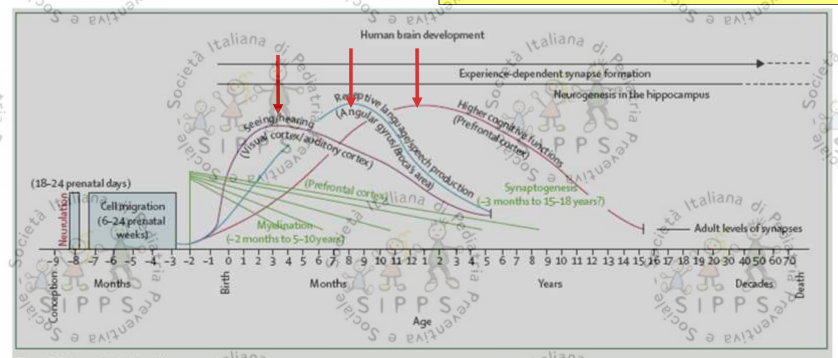


Early critical periods in the development of SYNAPTOGENESIS and brain functions

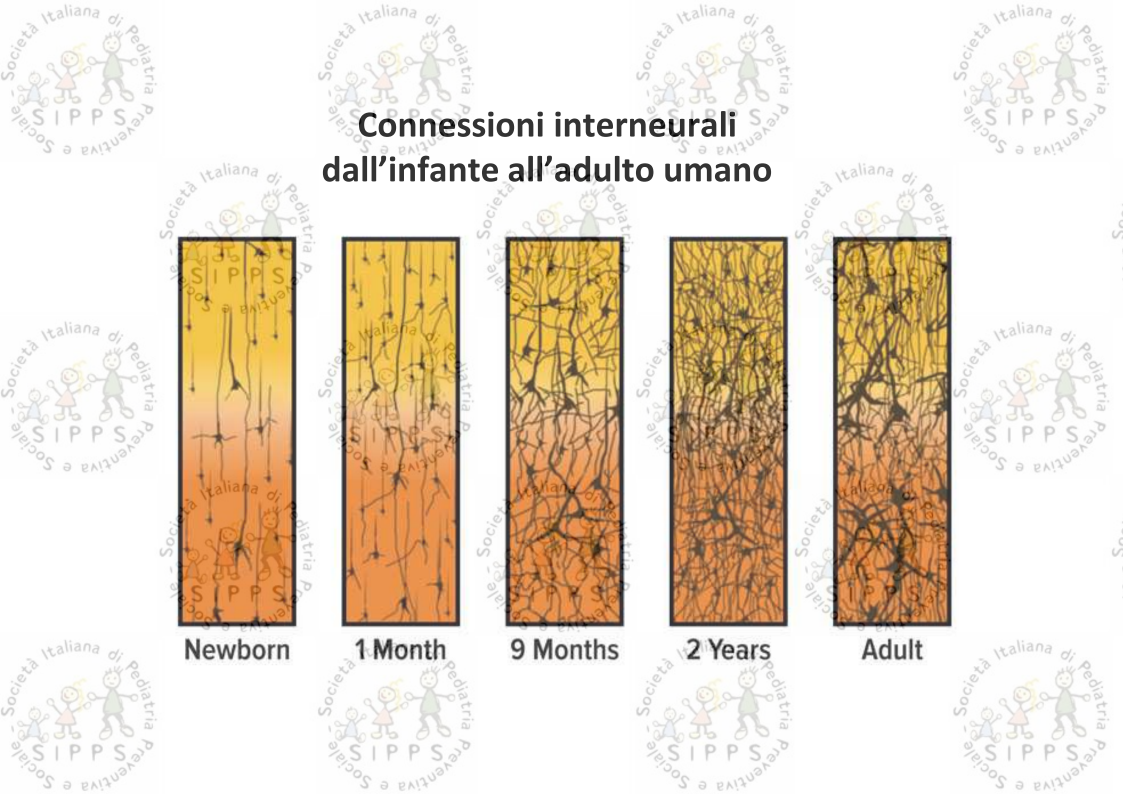
Formation of new synapses following stimulation..

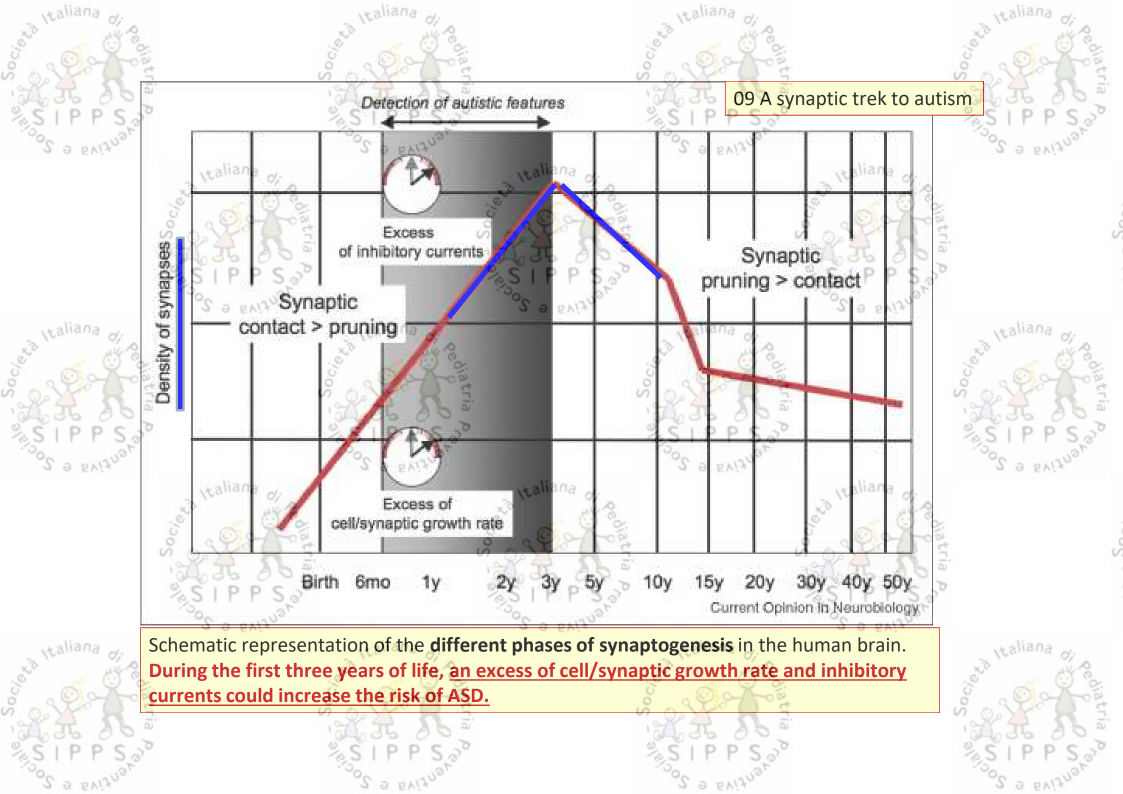
Disturbing the **CONNECTOME INSTRUCTION**

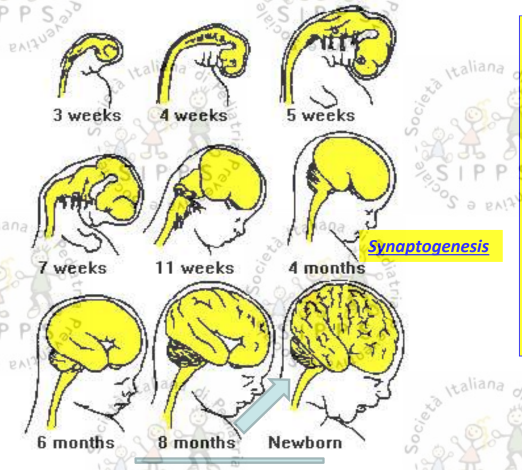
The *Individual* wirir



Reproduced with permission of authors and American Psychological Association® (Thompson RA, Nelson CA. Developmental science and the media: early brain development. Am Psychol 2001; 56: 5–15).







The brain grows at an amazing rate during development.

At times during brain development, 250,000 neurons are added every minute!

At birth, almost all the neurons that the brain will ever have are present.

However, the brain continues to grow for many years after birth.

By the age of 2 years old, the brain is about 80% of the adult size

2 9 BVis

A <u>stegosaurus dinosaur weighed approximately 1,600 kg but had a brain that weighed only approximately 70 grams (0.07 kg).</u> Therefore, <u>the brain was only 0.004% of its total body</u> weight. In contrast, an adult human weighs approximately 70 kg and has a brain that weighs approximately 1.4 kg. Therefore, <u>the human brain is about 2% of the total body weight</u>. This makes the brain to body ratio of the human <u>500 times greater than that of the stegosaurus</u>





Genes 2017, 8, 150; doi:10.3390/genes8060150



Review

Maternal Factors that Induce Epigenetic Changes Contribute to Neurological Disorders in Offspring

Avijit Banik ¹, Deepika Kandilya ¹, Seshadri Ramya ¹, Walter Stünkel ², Yap Seng Chong ³ and S. Thameem Dheen ^{1,*}

It is well established that the regulation of epigenetic factors, including chromatin reorganization, histone modifications, DNA methylation, and miRNA regulation, is critical for the normal development and functioning of the human brain.

There are a number of maternal factors influencing epigenetic pathways such as lifestyle, including diet, alcohol consumption, and smoking, as well as age and infections (viral or bacterial).

Genetic and metabolic alterations such as obesity, gestational diabetes mellitus (GDM), and thyroidism alter epigenetic mechanisms, thereby contributing to neurodevelopmental disorders (NDs) such as embryonic neural tube defects (NTDs), autism, Down's syndrome, Rett syndrome, and later onset of neuropsychological deficits.

This review comprehensively describes the recent findings in the epigenetic landscape contributing to altered molecular profiles resulting in NDs. Furthermore, we will discuss potential avenues for future research to identify diagnostic markers and therapeutic epi-drugs to reverse these abnormalities in the brain as epigenetic marks are plastic and reversible in nature.

Figure 1 Smoking in mothers alters neurodevelopmental processes in the fetus. <u>Maternal smoking alters the DNA methylation</u> of genes involved in placental and fetal development, leading to neurodevelopmental disorders in the offspring.

Maternal Smoking

Alteration in DNA methylation pattern of fetal gene pools

- Placental Function: LINE-1 [43], AluYb8 [9]
- Neurodevelopment: NR3C1 [50], HSD11B2 [51], GPR13, LRFN3 [53] Mana
- Neurotransmission: HTR2A, ADA [47,48]
- Immune development: ADA, PTPN22 [48]
- Transcriptome regulator: RUNX3 [46], PURA, GTF2H2, HKR1 [49]
- Calcium binding: GCA [45]
- Metabolism of aromatic hydrocarbon: CYP1A1 [49]
- · Placental abruption, Miscarriage, stillbirth, preterm delivery
- Neurobehavioral disorders: ADHD, Autism, Tourette's syndrome, Tic disorder, Obsessive-compulsive disorder

Exposure of the germline to nicotine produces epigenetic changes in the germline... they are permanent, and passed from one generation to the next

Mother - 1st general

Fetuse 2nd generatio

F2 Epigenetic targets of alcohol exposure in the fetus. Gestational alcohol exposure induces histone modification, alteration in DNA methylation pattern and miRNA targets, and expression of genes associated with fetal developmental process, leading to neurodevelopmental disorders. Gestational Alcohol Exposure

Susceptible targets in the fetus

· Developmental: Plune, Neurofilament, Pale ear [68],

Gene targets

Hoxa1 [87]

Oct4, Sox2, Nanog [72], Bub1, · Cell Proliferation: italiana

Cdc20, CcnB1, Plk1 [74]

Sox1, Zic1, Cxcl12, BMP8b, Dmrt1, · Cell Differentiation:

Meis1, Mef2c [72], Sh3bp2, Tnf,

Adrala, Pik3r1 [75]

Brain development, Pten, Otx2, Slitrk2, Nmnat1 [79]

H19 [76], POMC [80], Sfinbt2, Dlk1 · Imprinting:

Ube3a [79]

· Learning & Memory: PNOC, PDYN [82]

miRNA targets

miR-9, miR-21, miR-153, miR-335 [73]; miR-10a, miR-10b, miR-30a-3p, miR-145, miR-152, miR-29c, miR-30e-5p. miR-154, miR-200a.

miR-296, miR-339.

miR-362, miR-496 [87]

H3K9ac [81]

CBP [83]

H3K27me3 [82]

Histone modifying targets

DNMT, MeCP2 [67]

DNA methylation targets

Lamage to brain causes difficulty learning remembering. thirking things through and getting along with others

Vision problems

Heart, kidney, liver and other organ damage

Bones. limbs and fingers that are not formed properly

Be Safe: Have an alcohol-free pregnancy

talian

* I brinking alcohol darting pregnancy. can cause birth. defects and brain damage to your baby



drink any alcohol during pregnancy.

. In fact it is best to stop drinking before you get pregnant.

+ It is safest not to

Any kind of alcohol can harm your baby

Phenotypic outcomes in the offspring

Fetal alcohol spectrum disorder (FASD)

- Attention and memory deficit
- Craniofacial malformation
- Motor function abnormalities
- Auditory and language problem

F3 Effect of maternal dietary deficiency on fetal development.

The absence of essential dietary supplements in maternal diet during gestation leads to a disruption in metabolic pathways and several epigenetic alterations in the fetus, triggering abnormal uterine development and neurodevelopmental disorders.

AMaternal dietary deficiency taliana

Absence of dietary methyl group donors such as folate, choline, methionine, betain and methylcobalamine

- Imbalance in folate-mediated one-carbon metabolism (FOCM) pathway [98]
- Mutation in methionine synthase reductase (Mtrr) gene, essential for deployment of methyl groups from the folate cycle [104]
- Down-regulation of genes related to fetal brain development: BDNF, CREB, NGF and TrkB [105]
- · H3K9 and H4K20 methylation [114]
- Altered expression of miRNAs linked to FOCM pathway: miR-29c, miR-183, miR-422b, miR-189 [115]; miR-22, miR-24, miR-29b, miR-34a, miR-125, miR-344-5p/484, miR-488 [116-118]

Epigenetic changes 1
UNA Methyladical menteoding
RNA, Histone modifications

Maternal
Felic Acid

Altered Sene expression

Inserting Dens 7

Candidate
Sens 7

Developmental outcomes ?

Abnormal uterine development and congenital malformation [104]

F4 Effect of maternal metabolic conditions on fetal development.

Metabolic conditions at gestation such as GDM, obesity, and hypothyroidism induce epigenetic alterations in the fetus, leading to a series of metabolic and immunogenic changes triggering neuroanatomical and neuropsychological deficits in the developing brain.

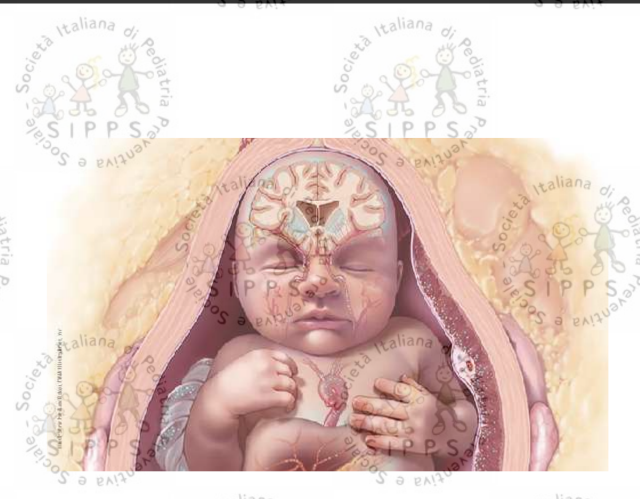
Maternal metabolic conditions

- · Gestational Diabetes Mellitus (GDM)
- · Maternal Obesity
- Maternal Hypothyroidism

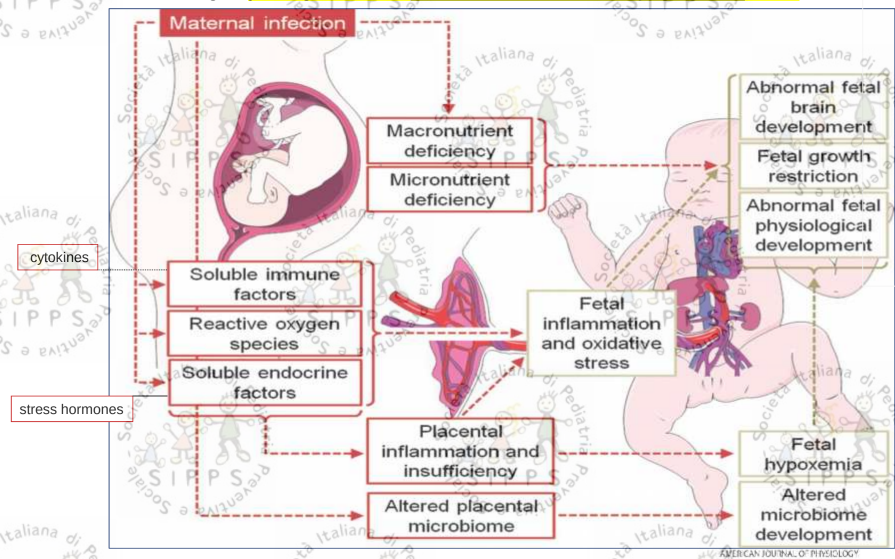
Trigger epigenetic imbalance in the fetus [149,150,157,158,172]

- Induces oxidative stress [148]
- ROS accumulation [148]
- Inflammatory response [155]
- · Cytokine production [156]
- Decreased T3 levels [169]
- · Altered levels of metabolic genes [172]

Neuroanatomical /neuropsychological deficits in developing brain

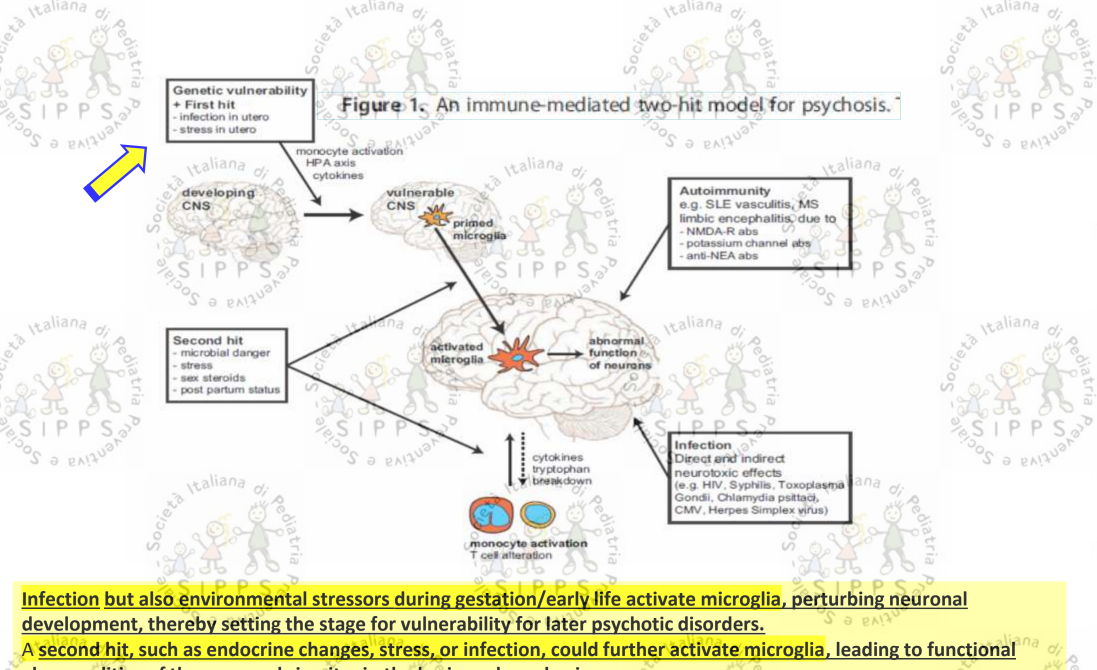


Possible mechanisms mediating the pathological effects of maternal infection on the developing organism in utero

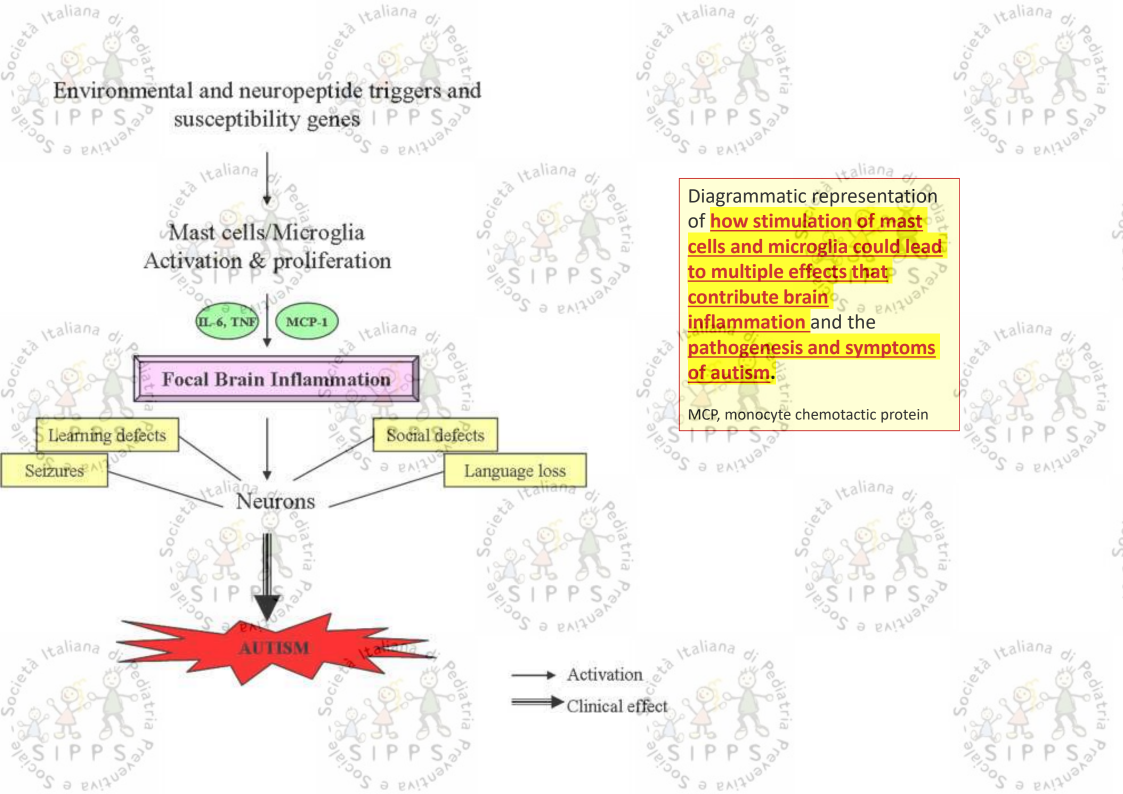


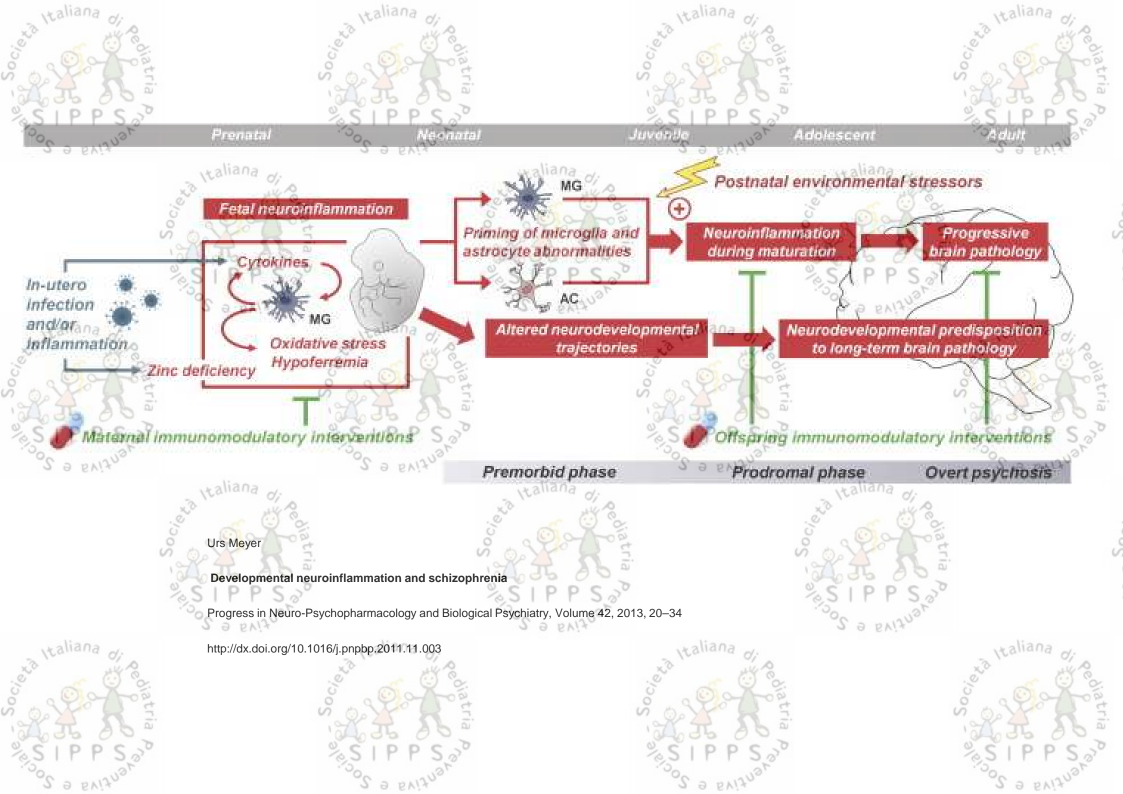
Regulatory, Integrative and Comparative Physiology

Marie A. Labouesse et al. Am J Physiol Regul Integr Comp Physiol 2015;309:R1-R12



abnormalities of the neuronal circuitry in the brain and psychosis





Pre or postnatal exposure?





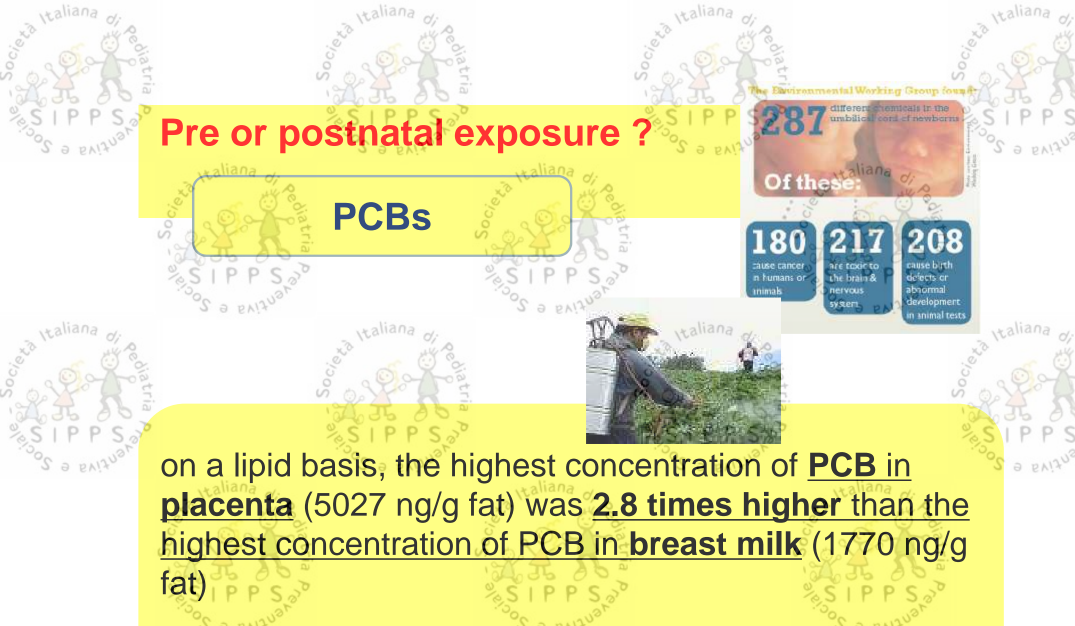


Incinerators, landfills.. primitive waste recycle, etc.

Higher PCDD/F levels were found in placenta (10.3 TEqpg/g lipid) and venous serum (9.1 TEq-pg/g lipid), compared to those in breast milk (7.6 TEq-pg/g lipid).

Chemosphere. 2004 Mar;54(10):1459-73. *Infant exposure to polychlorinated dibenzo-p-dioxins, dibenzofurans and biphenyls (PCDD/Fs, PCBs)--correlation between prenatal and postnatal exposure*. Wang SL, Lin CY, Guo YL, Lin LY, Chou WL, Chang LW.

Giuseppe Giordano ISDE Palermo



J Expo Anal Environ Epidemiol. 2000 May-Jun;10(3):285-93. PCB exposure in utero and via breast milk. A review. DeKoning EP, Karmaus W. Et al.

Giuseppe Giordano ISDE Palermo

Neurodevelopmental Disorders and Prenatal Residential Proximity to Agricultural Pesticides: The CHARGE Study

Janie F. Shelton, Estella M. Geraghty Environ Health Perspect; DOI:10.1289/ehp.1307044: 23 June 2014

970 participants, California Pesticide Use Report (1997-2008) linked to the addresses during pregnancy. Pounds of active ingredient ...
aggregated within 1.25km, 1.5km, and 1.75km buffer distances from the home







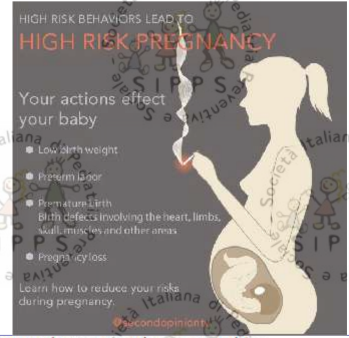
- •Organophosphates higher 3rd trimester expos: 60% increased risk ASD
- •Pyrethroid insecticide just prior to conception or for 3rd trimester at greater risk for both ASD and DD (developmental delay)
- •Carbamate: risk for DD increased (Arprocarb: Undene, Propoxur

 Baygon).

 Baygon

Giuseppe Giordano ISDE

"Tobacco smoke is
without a doubt
the most significant
environmental contaminant
to which children
are exposed indoors"



Children whose mothers smoke:

- 70% more respiratory problems
- Pneumonia and hospitalization in year 1 is 38% higher
- Infant mortality is 80% higher
- 20% of all infant deaths could be avoided if all pregnant smokers stopped by the 16th week of gestation

Environmental tobacco smoke (ETS)

Sudden infant death syndrome

Lower respiratory tract illness

Middle ear disease

Asthma

to secondhand smoke in homes



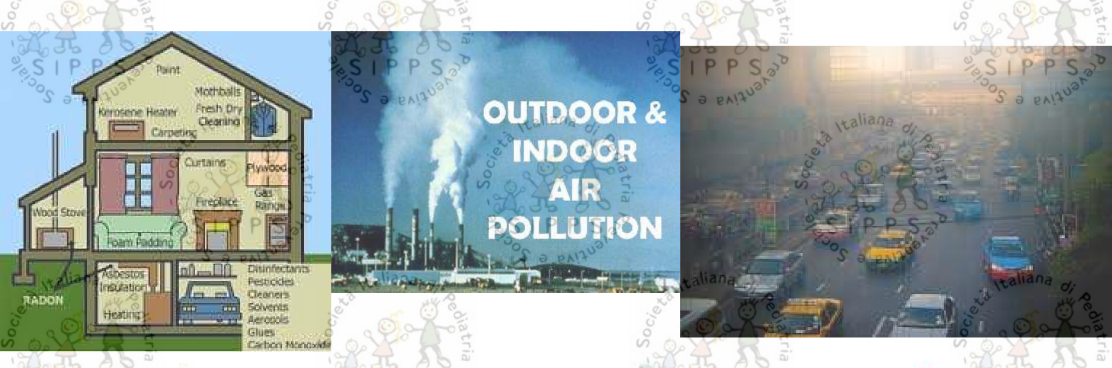


Exposure to environmental tobacco smoke (FTS) causes more than 35,000 deaths annually among non-smokers.

Smoking by pregnant women is responsible for about 1000 infant deaths each year in the U.S.

 Children exposed to ETS suffer higher rates of asthma, bronchitis, and pneumonia.

Smokeless tobacco use has tripled since 1972, and eigar use has increased 50% since 1993.



House dust mites

- House dust mites produce <u>Der phallergen</u>, a potent sensitizer
- Good evidence of increased risk of sensitization with increasing allergen exposure, but this does not necessarily lead to asthma
- Small reductions in exposure will not necessarily lead to reduced incidence and/or symptoms
- Indoor humidity is important

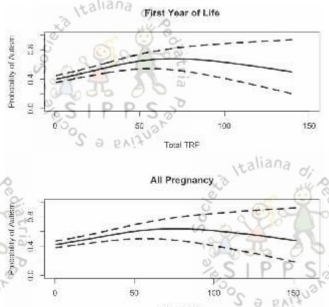


Living near a freeway, based on the location of the birth, and third trimester address, and **autism**

PM2.5, PM10, and NO2 at residences were higher in children with autism.

The magnitude of these <u>associations</u> appear to be <u>most pronounced during late</u> <u>gestation</u> (OR=1.98, 95%CI 1.20–3.31) <u>and early life / first year of life</u> (OR=1.98, 95%CI 1.20–3.31)

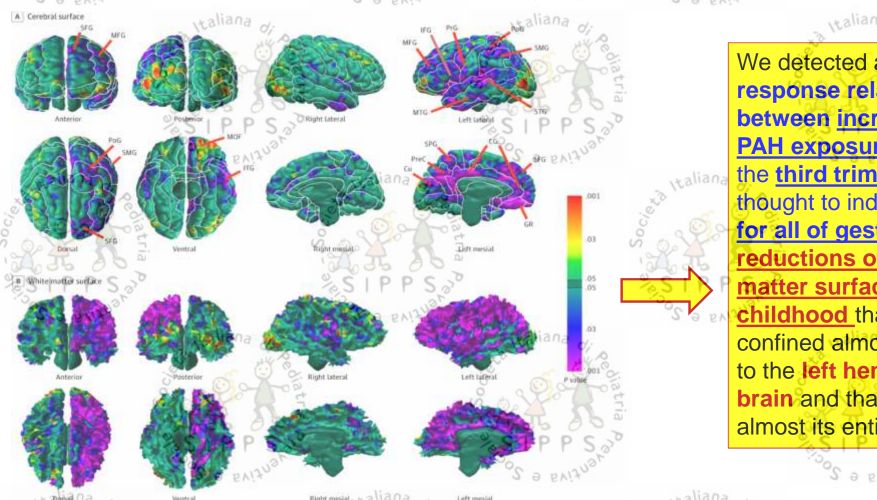




JAMA Psychiatry. 2013 January ; 70(1): 71–77. doi:10.1001/jamapsychiatry.2013.266

From: Effects of Prenatal Exposure to Air Pollutants (Polycyclic Aromatic Hydrocarbons) on the Development of Brain White Matter, Cognition, and Behavior in Later Childhood

JAMA Psychiatry. Published online March 25, 2015. doi:10.1001/jamapsychiatry.2015.57

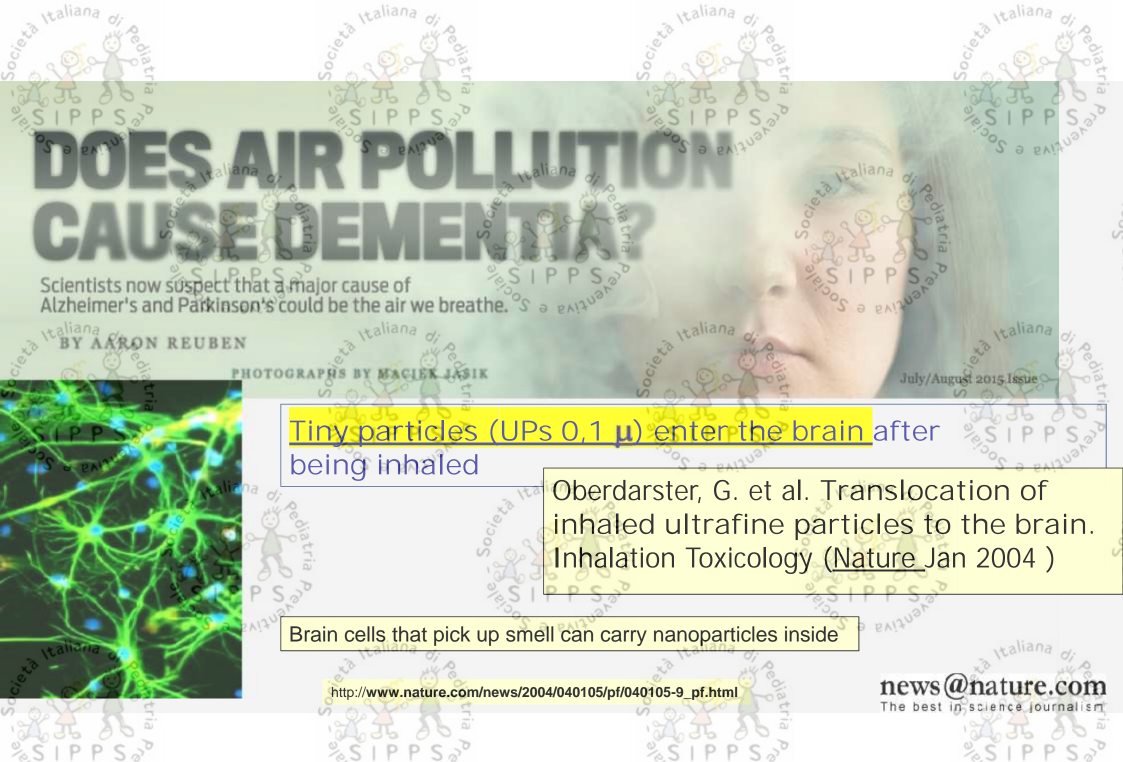


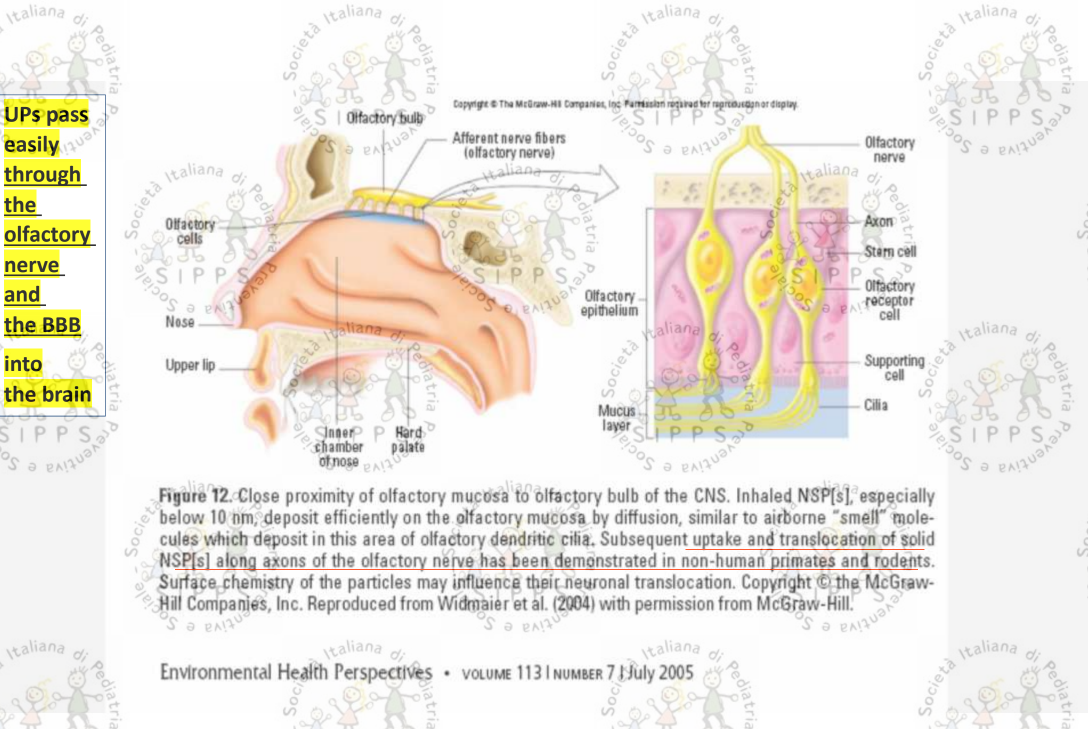
We detected a doseresponse relationship
between increased prenatal
PAH exposure (measured in
the third trimester but
thought to index exposure
for all of gestation) and
reductions of the white
matter surface in later
childhood that were
confined almost exclusively
to the left hemisphere of the
brain and that involved
almost its entire surface

Date of download: 4/6/2015

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In the most polluted cities even dogs have **Alzheimer's** disease

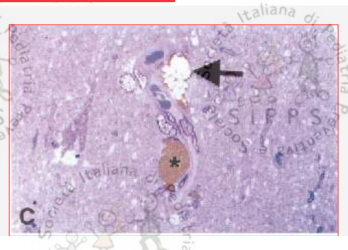
Toxicologic Pathology

http://tpx.sagepub.com

Air Pollution and Brain Damage
Lillan Calderon-Garcidueñas, Biagio Azzarelli, Hilda Acuna, Raquel Garcia, Todd M. Gambling, Norma Osnaya, Sylvia Monroy, Maria Del Rosario Tizapantzi, Johnny L. Carson, Anna Villarreal-Calderon and Barry Rewcastle. Toxicol Pathol 2002; 30: 373

Exposure to complex mixtures of air pollutants produces raffaringation in the upper and lower respiratory tract. Because the naval cavity is a common portal of entry, respiratory and olfactory epithelia are vulnerable targets for toxicological damage. This study has evaluated, by light and electron microscopy and immunohistochemical expression of nuclear factor-kappa beta (NF- κB) and inducible nitric oxide synthase (iNOS), the olfactory and respiratory nasal mucosae, olfactory bulb, and cortical and subcortical structures from 32 healthy mongrel caning residents in Southwest Metropolitan Mexico City (SWMMC), a highly polluted urban region. Findings were compared to those in 8 dogs from Tlaxcala, a less polluted, control city. In SWMMC dogs, expression of nuclear neuronal NF-kB and iNOS in cortical endothelial cells occurred at ages 2 and 4 weeks; subsequent damage included alterations of the blood-brain barrier (BBB), degenerating cortical neurons, apoptotic glial white matter cells, deposition of apolipoprotein E (apoE)-positive lipid droplets in smooth muscle cells and pericytes, nonneuritic plaques, and neurotibrillary tangles. Persistent pulmonary inflammation and deteriorating olfactory and respiratory barriers may play a role in the neuropathology observed in the brains of these highly exposed canines. Neurodegenerative disorders such as Alzheimer's may begin early in life with air pollutants playing a crucial role.





And a similar condition has been documented in the brain of young people dead for accidental causes...

Toxicologic Pathology

http://tpx.sagepub.com



Lilian Calderón-Garcidueñas, Maricela Franco-Lira, Ricardo Torres-Jardón, Carlos Henriquez-Roldán, Gerardo Barragán-Mejía, Cildardo Valencia-Salazar, Angelica Conzález-Maciel, Rafael Reynoso-Robles, Rafael Villarreal-Calderón and William Reed

Toxicol Pathol 2007; 35: 154

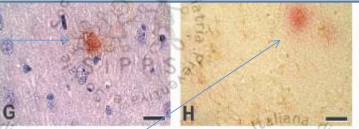
Exposures to particulate matter and gaseous air pollutants have been associated with respiratory tract inflammation, disruption of the nasal respiratory and olfactory barriers, systemic inflammation, production of mediators of inflammation capable

of <u>reaching the brain and systemic circulation of particulate matter</u>. Mexico City (MC) residents are exposed to significant amounts of *ozone, particulate matter* and associated *lipopolysaccharides*. MC dogs exhibit brain inflammation and an <u>acceleration of Alzheimer's-like pathology, suggesting that the brain is adversely</u> affected by air pollutants.

MC children, adolescents and adults have a significant upregulation of cyclooxygenase-2 (COX2) and interleukin-16 (IL-16) in olfactory bulb and frontal cortex, as well as neuronal and astrocytic accumulation of the 42 amino acid form of θ -amyloid peptide (A θ 42), including diffuse amyloid plaques in frontal cortex. The pathogenesis of Alzheimer's disease (AD) is characterized by brain inflammation and the accumulation of A θ 42, which precede the appearance of neuritic plaques and neurofibrillary tangles, the pathological hallmarks of AD.

Our findings of nasal barrier disruption, systemic inflammation, and the upregulation of COX2 and IL-16 expression and A642 accumulation in brain suggests that sustained exposures to significant concentrations of air pollutants such as particulate matter could be a risk factor for AD and other neurodegenerative diseases.

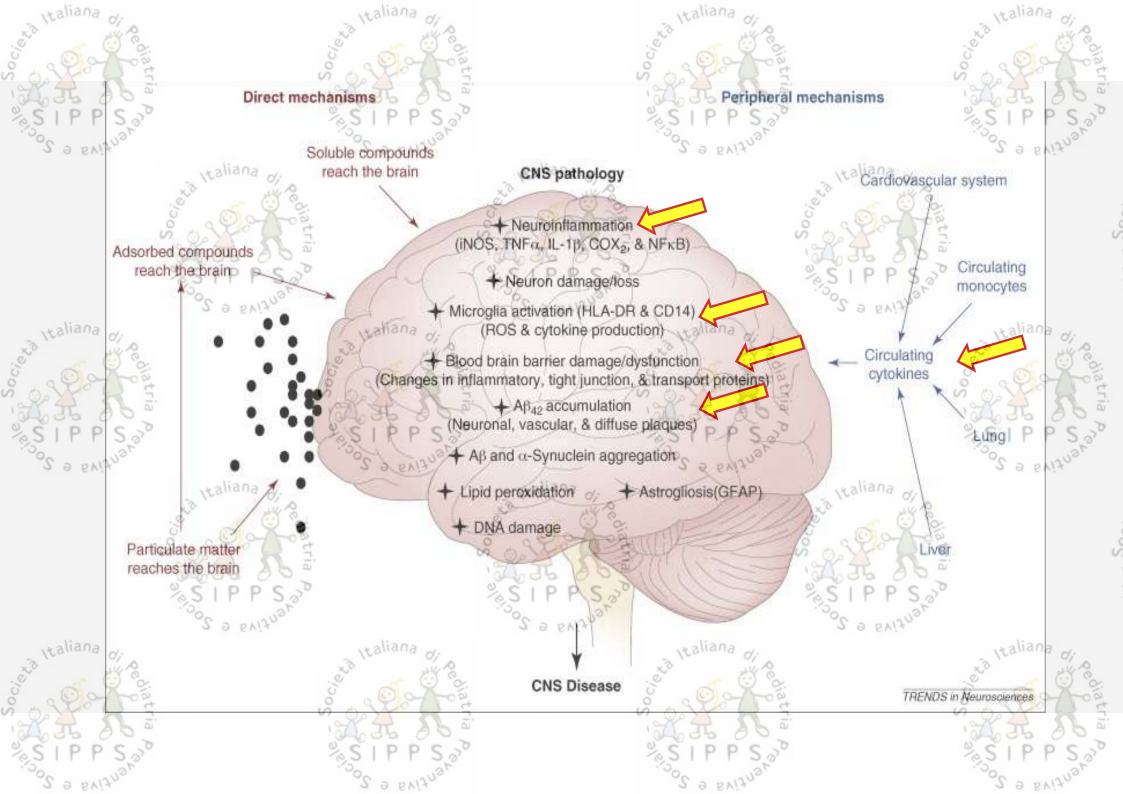
The frontal cortex of an 11-month-old healthy MC dog exhibits A 42 staining of a diffuse plaque, surrounded by a microglia-like nucleus

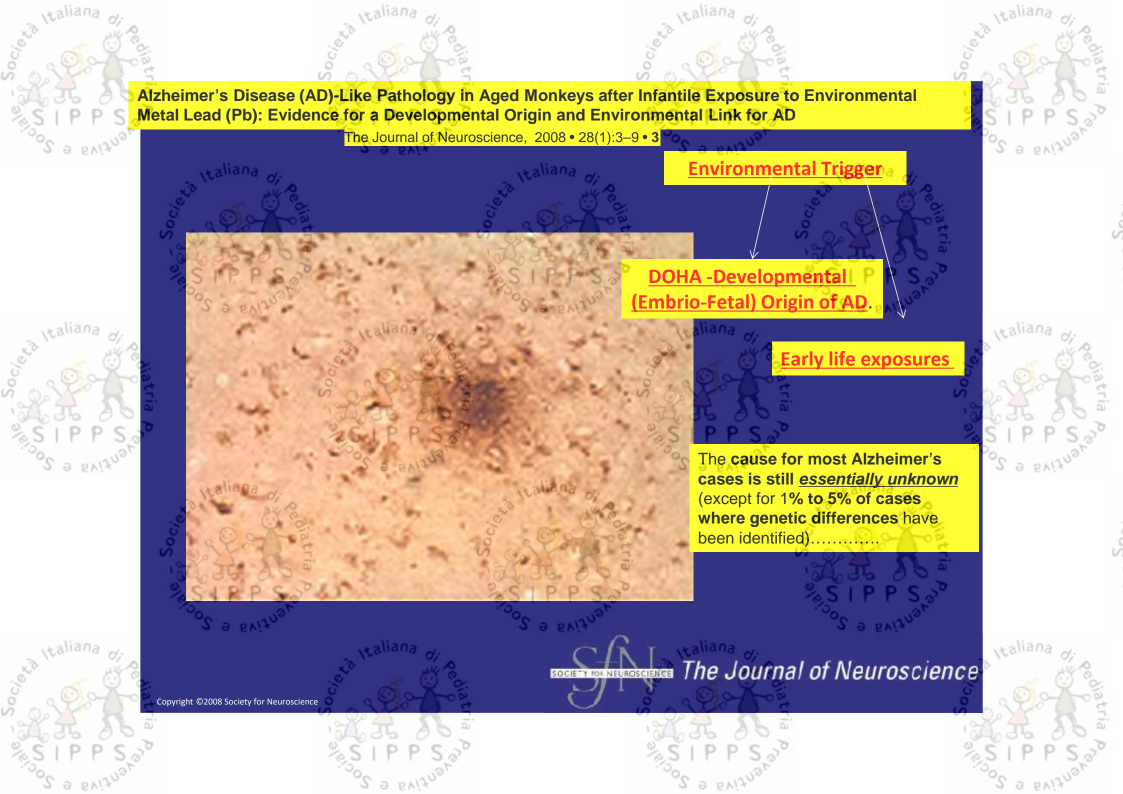


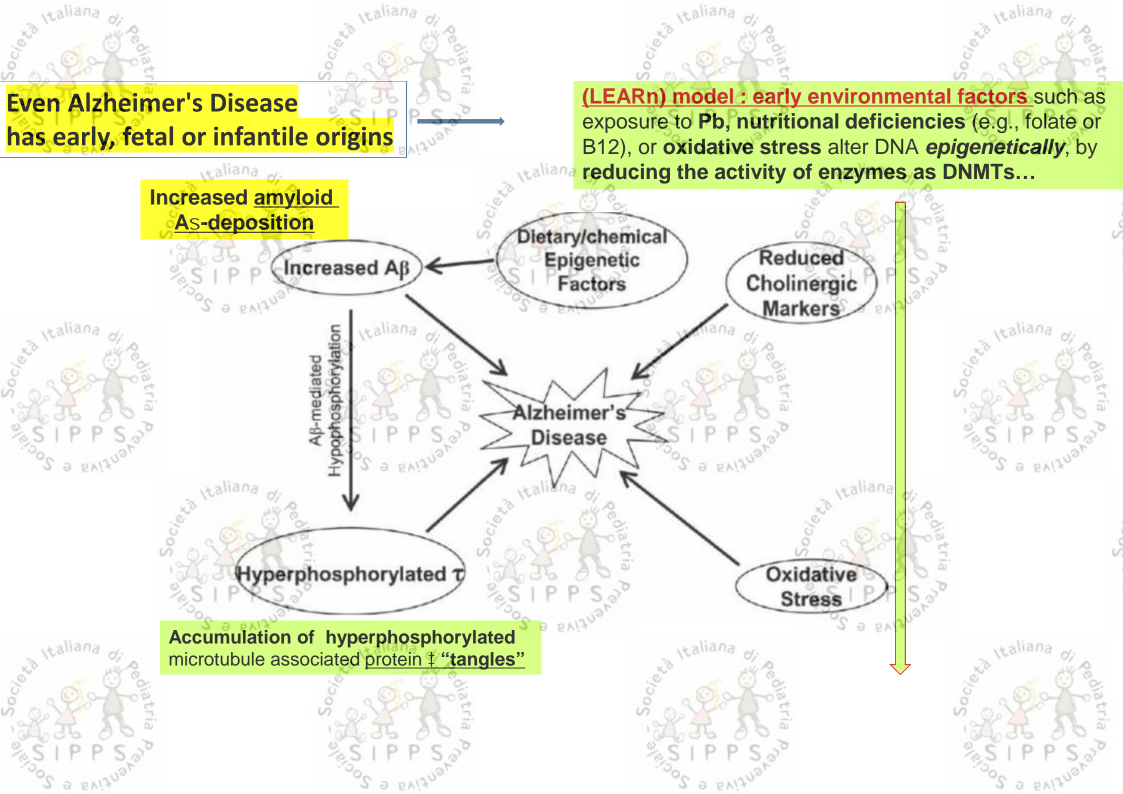


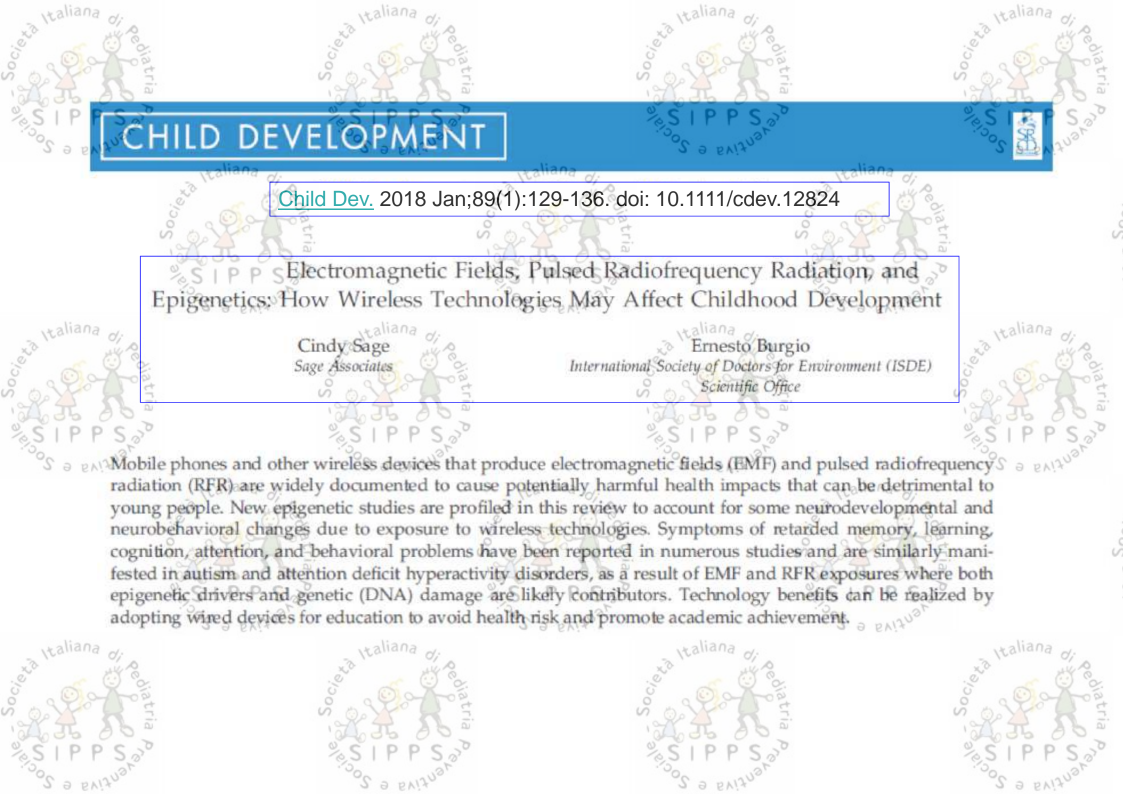
The frontal cortex of a **17-year-old MC boy**... shows a **diffuse**A **42 plaque** (red product) and GFAP-negative astrocytes

The frontal cortex of a 36-year-old MC male with an E3/E4 ApoE genotype .. shows abundant mature and diffuse A 42 plaques (red stain) along with GFAP-positive reactive astrocytosis

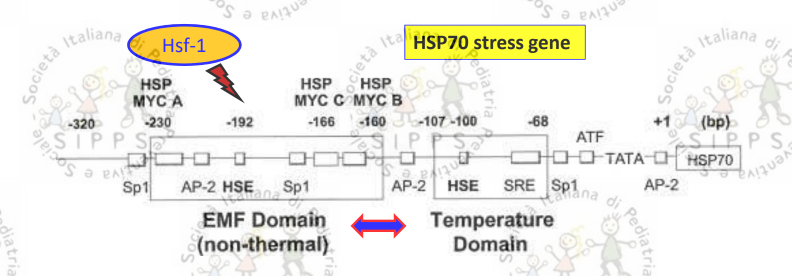








Specific DNA sequences on the promoter of the HSP70 stress gene are responsive to EMF...



Synthesis of this stress protein is initiated in a <u>region of the promoter</u> where a transcription factor known as <u>Heat Shock</u> <u>Factor 1 (HSF-1) binds to a Heat Shock Element (HSE).</u>

The <u>EMF sensitive region on HSP70 promoter</u> is <u>upstream from the thermal domain of the promoter and is not sensitive</u> to increased temperature. The binding of <u>HSF-1</u> to <u>HSE</u> occurs at **-192** in the HSP70 promoter relative to the transcription initiation site.

The EMF domain contains three nCTCTn myc-binding sites -230, -166 and -160 relative to the transcription initiation site and upstream of the binding sites for the heat shock (nGAAn) and serum responsive elements.... The electromagnetic response elements (EMREs) have also been identified on the c-myc promoter and are also responsive to EMF

Pathophysiology

Volume 16, Issues 2-3, August 2009, Pages 71-78

SCIENTIFIC REPORTS

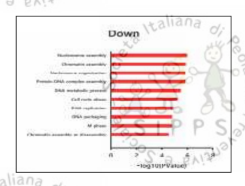
Effects of a hypomagnetic field on DNA methylation during the differentiation of embryonic stem cells

Soonbong Baek¹, Hwan Choi¹, Hanseul Park¹, Byunguk Cho¹, Siyoung Kim¹ & Jongpil Kim^{1,2}

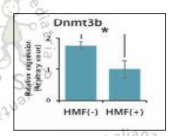
It has been reported that hypomagnetic fields (HMFs) have a negative influence on mammalian physiological functions. We previously reported that HMFs were detrimental to cell fate changes during reprogramming into pluripotency. These studies led us to investigate whether HMFs affect cell fate determination during direct differentiation. Here, we found that an HMF environment attenuates differentiation capacity and is detrimental to cell fate changes during the *invitro* differentiation of embryonic stem cells (ESCs). Moreover, HMF conditions cause abnormal DNA methylation through the dysregulation of DNA methyltransferase3b (Dnmt3b) expression, eventually resulting in incomplete DNA methylation during differentiation. Taken together, these results suggest that an appropriate electromagnetic field (EMF) environment may be essential for favorable epigenetic remodeling during cell fate determination via differentiation.

Published online: 04 February 2015





...campi ipomagnetici (HMF) influenzano la determinazione del destino cellulare... interferendo sulla differenziazione in vitro delle cellule staminali embrionali (ESC). ...attraverso la disregolazione dell'espressione di DNA metiltransferasi 3b (Dnmt3b), con conseguente metilazione incompleta del DNA



BIOPHYSICS

Weak magnetic fields alter stem cell-mediated growth

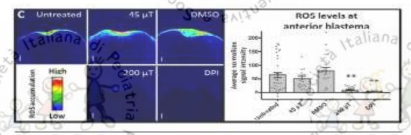
Alanna V. Van Huizen¹, Jacob M. Morton¹, Luke J. Kinsey¹,
Donald G. Von Kannon¹, Marwa A. Saad¹, Taylor R. Birkholz¹, Jordan M. Czajka¹,
Julian Cyrus², Frank S. Barnes², Wendy S. Beane¹*

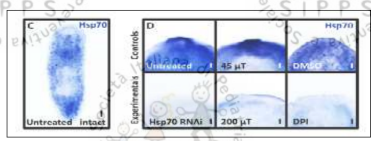
Biological systems are constantly exposed to electromagnetic fields (EMFs) in the form of natural geomagnetic fields and EMFs emitted from technology. While strong magnetic fields are known to change chemical reaction rates and free radical concentrations, the debate remains about whether static weak magnetic fields (WMFs; <1 mT) also produce biological effects. Using the planarian regeneration model, we show that WMFs altered stem cell proliferation and subsequent differentiation via changes in reactive oxygen species (ROS) accumulation and downstream heat shock protein 70 (Hsp70) expression. These data reveal that on the basis of field strength, WMF exposure can increase or decrease new tissue formation in vivo, suggesting WMFs as a potential therapeutic tool to manipulate mitotic activity.

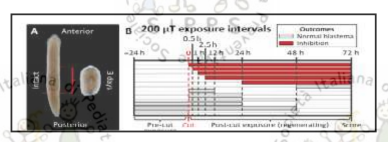
Campi magnetici statici deboli (WMF <1 mT) producono alterazioni della proliferazione delle cellule staminali e della successiva differenziazione attraverso cambiamenti nell'accumulo di specie reattive dell'ossigeno (ROS) e nell'espressione della proteina di shock termico 70 (Hsp70).

Questi dati rivelano che sulla base della forza del campo, l'esposizione al WMF può aumentare o diminuire la formazione di nuovo tessuto in vivo...

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American Association for the Advancement of Science. No claim to original BS Government Works. Distributed under a Creative
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Abuse Leaves Its Mark on the Brain

http://news.sciencemag.org/biology/2009/02/abuse-leaves-its-mark-brain



Francisco_de_Goya,_Saturno_devo rando_a_su_hijo_(1819-1823)



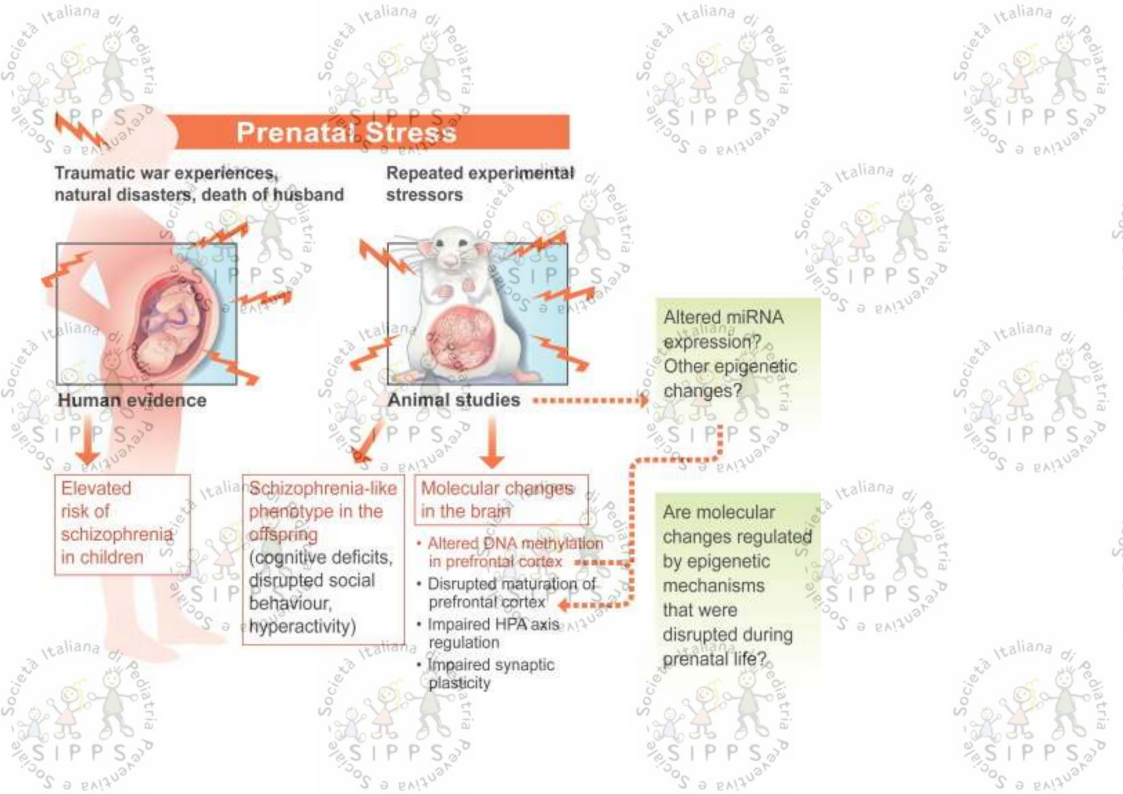
<u>Child abuse</u> is an environmental factor that leaves an <u>epigenetic mark on</u> the brain

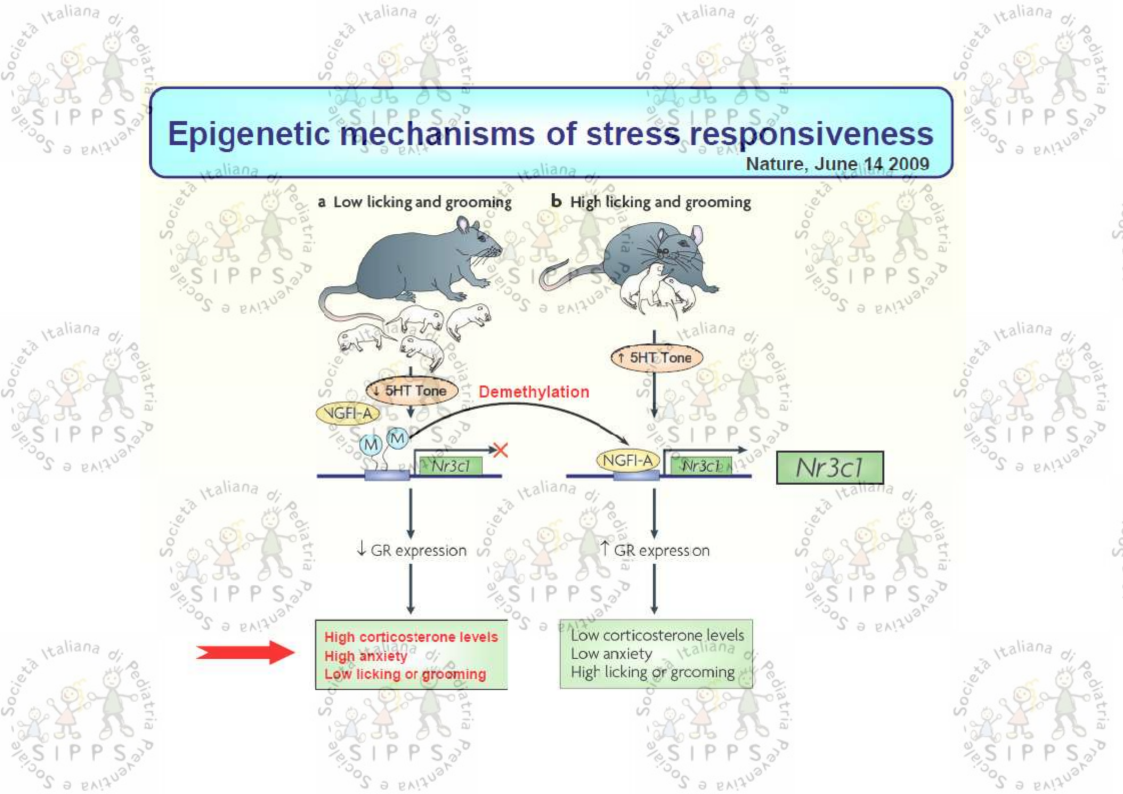
In a comparison of suicide victims who were abused or not, only the abused victims had an epigenetic tag on the GR gene

Interestingly, the GR gene receives a similar epigenetic tag in rat pups who receive low quality care from their mothers.



http://learn.genetics.utah.edu/content/epigenetics/brain/







Epigenetic regulation of the glucocorticoid receptor in human brain associates with childhood abuse

nature neuroscience

Patrick O McGowan^{1,2}, Aya Sasaki^{1,2}, Ana C D'Alessio³, Sergiy Dymoy³, Benoit Labonté^{1,4}, Moshe Szyf^{2,3}, Gustavo Turecki^{1,4} & Michael J Meaney^{1,2,5}

OLUME 12 NUMBER 3 MARCH 2009 NATURE NEUROSCIENCE

Maternal care influences hypothalamic-pituitary-adrenal (HPA) function in the rat through epigenetic programming of glucocorticoid receptor expression. In humans, childhood abuse alters HPA stress responses and increases the risk of suicide. We examined epigenetic differences in a neuron-specific glucocorticoid receptor (NR3C1) promoter between postmortem hippocampus obtained from suicide victims with a history of childhood abuse and those from either suicide victims with no childhood abuse or controls. We found decreased levels of glucocorticoid receptor mRNA, as well as mRNA transcripts bearing the glucocorticoid receptor 1st splice variant and increased cytosine methylation of an NR3C1 promoter. Patch-methylated NR3C1 promoter constructs that mimicked the methylation state in samples from abused suicide victims showed decreased NGFI-A transcription factor binding and NGFI-A-inducible gene transcription. These findings translate previous results from rat to human and suggest a common effect of parental care on the epigenetic regulation of hippocampal glucocorticoid receptor expression.

20 - 0.5. Control Suicide nonabuser Suicide abuse 20 - 0.5. Suicide abuse 20 -

Figure 2 Methylation of the *NR3C1* promoter in the hippocampus. Twenty-clones were sequenced for each subject for methylation mapping percentage of methylated clones for suicide victims with a history of childhood abuse (n = 12), suicide victims without a history of childhood abuse on the number of clones with at least one methylated CpG site divides number of clones (* indicates $P \le 0.05$; n.s. indicates not statistically significant). (b) Methylation of the *NR3C1* promoter region, show of methylation observed at each CpG site for suicide victims with a history of childhood abuse, suicide victims with no history of childhood abuse. (posteron subjects (* $P \le 0.05$, * $P \le 0.001$, abused suicides versus controls; * $P \le 0.001$, non-abused suicides versus controls; * $P \le 0.001$, abused suicides versus non-abused suicides: Bonfarron post hoc comparisons).

Maternal care influences the programming of the hypothalamic-pituitary-adrenal Axis (HPA) through epigenetic programming of glucocorticoid receptors expression...

We found a greatly increased methylation of cytosine in the promoter of a gene codifying for a Glucocorticoids-Neuro-Receptor (NR3C1) in the hippocampus of suicide victims with a history of childhood abuse .. (post-mortem examinations)

ORIGINAL ARTICLE

Association of Maternal Exposure to Childhood Abuse With Elevated Risk for Autism in Offspring

Andrea L. Roberts, PhD; Kristen Lyall, ScD; Janet W. Rich-Edwards, ScD; Alberto Ascherio, DrPH; Marc G. Weisskopf, PhD, ScD

Importance: Adverse perinatal circumstances have been associated with increased risk for autism in offspring. Women exposed to childhood abuse experience more adverse perinatal circumstances than women unexposed, but whether maternal abuse is associated with autism in offspring is unknown.

Design and Setting: Nurses' Health Study II, a population-based longitudinal cohort of 116,430 women.

Conclusions and Relevance: We identify an intergenerational association between maternal exposure to childhood abuse and risk for autism in the subsequent generation. Adverse perinatal circumstances accounted for only a small portion of this increased risk.

JAMA Psychiatry. 2013;70(5):508-515. Published online March 20, 2013. doi:10.1001/jamapsychiatry.2013.447

Another transgenerational
effect, is based on a broad
longitudinal cohort study
(Nurses' Health Study II)
which identified maternal
exposure to abuse in early
childhood (!!) as a risk factor
for having a child with autism e
(Nurses 'Health Study II)



Isr J Psychiatry RelatSci - Vol. 50 - No I (2013)

Epigenetic Transmission of Holocaust Trauma: Can Nightmares Be Inherited?

Natan P.F. Kellermann

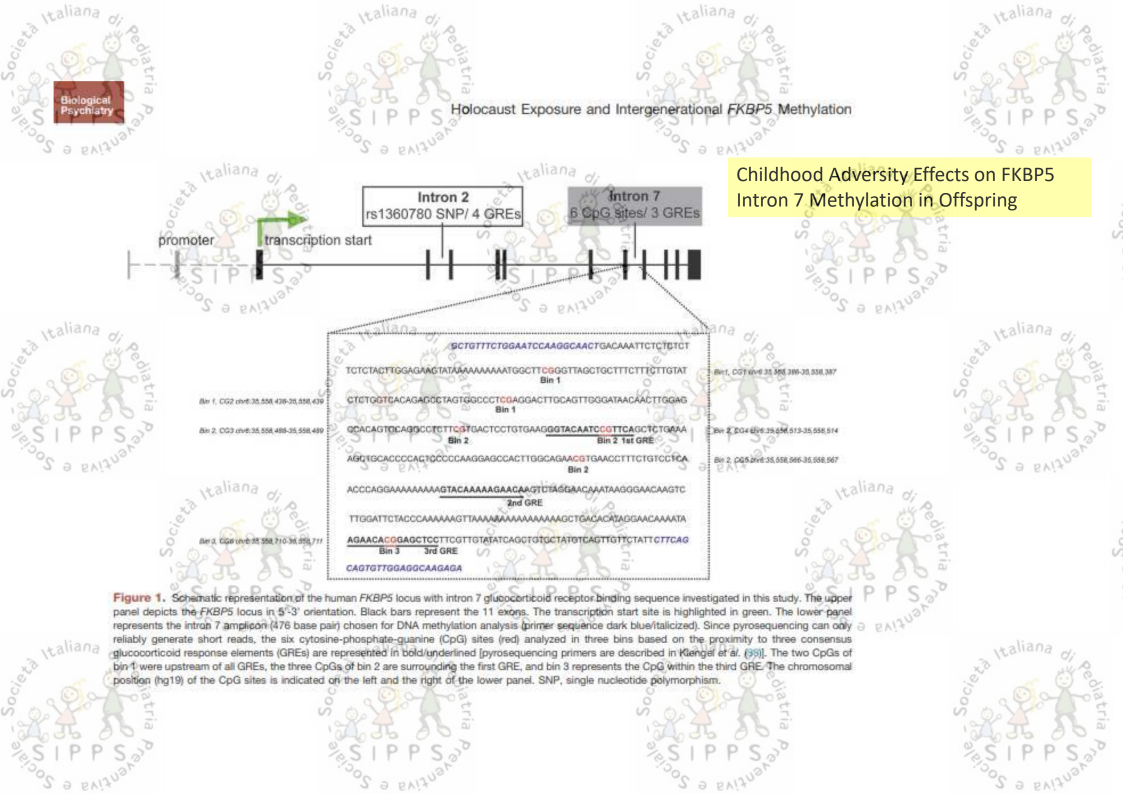
AMCHA, the National Israeli Center for Psychosocial Support of Survivors of the Holocoust and the Second Generation, Jerusalem, Israel

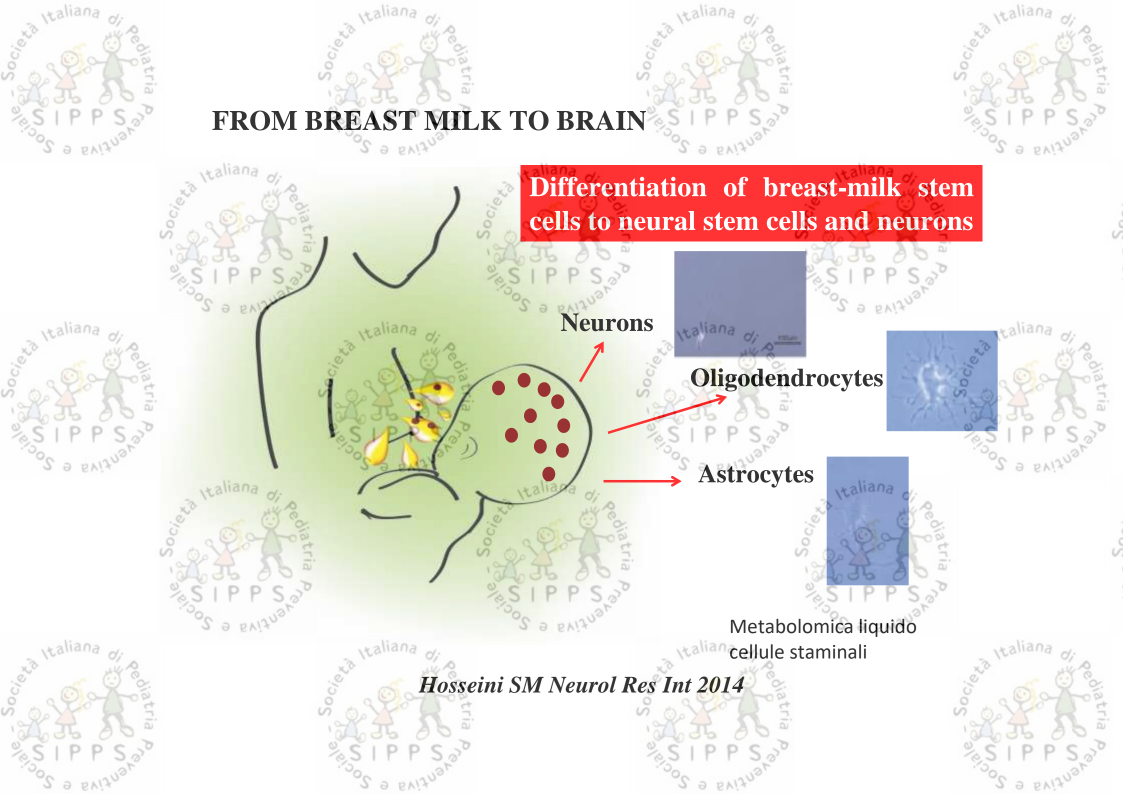
The Holocaust left its visible and invisible marks not only on the survivors, but also on their children. Instead of numbers tattooed on their forearms, however, they may have been marked epigenetically with a chemical coating upon their chromosomes, which would represent a kind of biological memory of what the parents experienced. As a result, some suffer from a general vulnerability to stress while others are more resilient. Previous research assumed that such transmission was caused by environmental factors, such as the parents' childrearing behavior. New research, however, indicates that these transgenerational effects may have been also (epi) genetically transmitted to their children Integrating both hereditary and environmental factors, epigenetics adds a new and more comprehensive psychobiological dimension to the explanation of transgenerational transmission of trauma. Specifically, epigenetics may explain why latent transmission becomes manifest under stress. A general theoretical overview of epigenetics and its relevance to research on trauma transmission is presented.

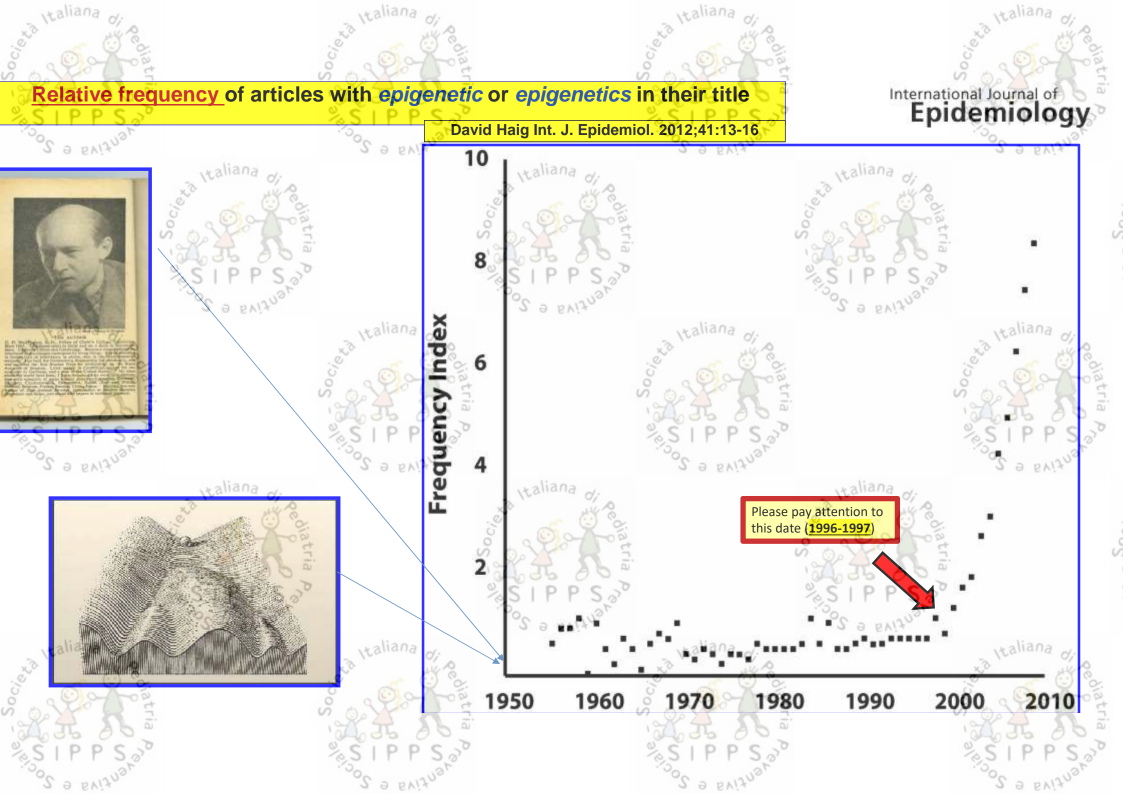
The Holocaust left its visible and invisible marks not only on the survivors, but also on their children. Instead of numbers tattooed on their forearms, however, they may have been marked epigenetically with a chemical coating upon their chromosomes, which would represent a kind of biological memory of what the parents experienced.

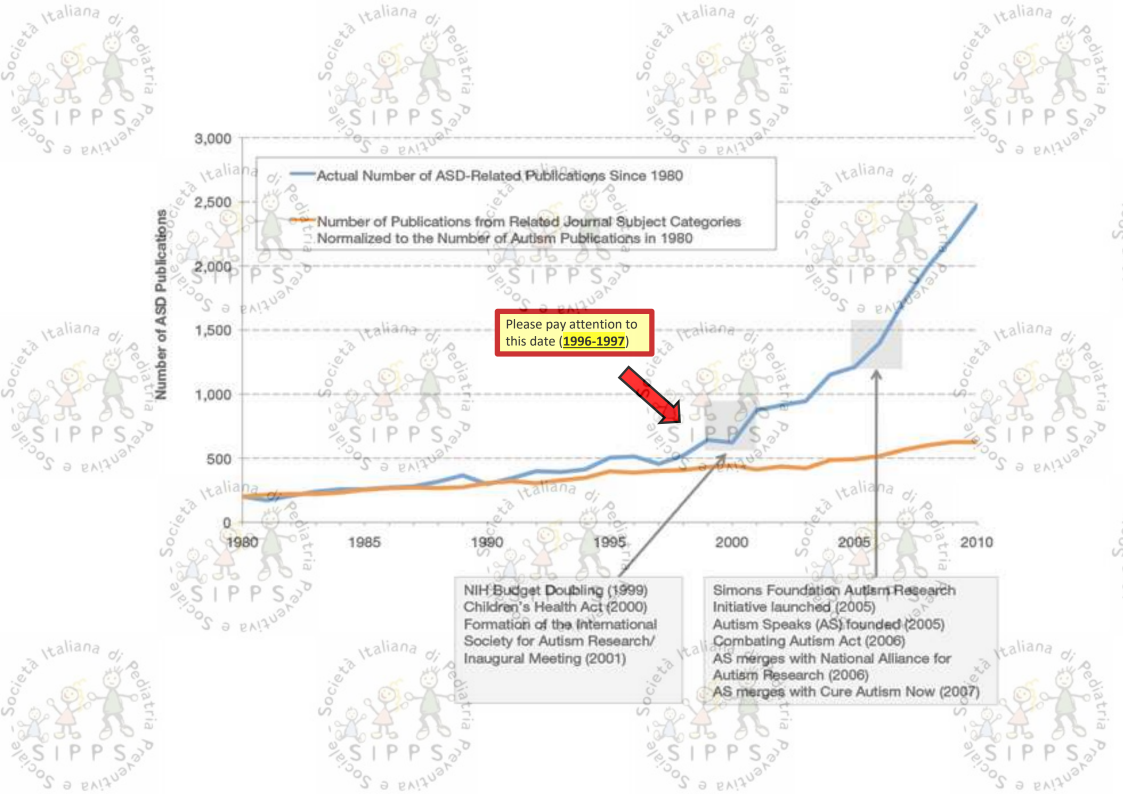




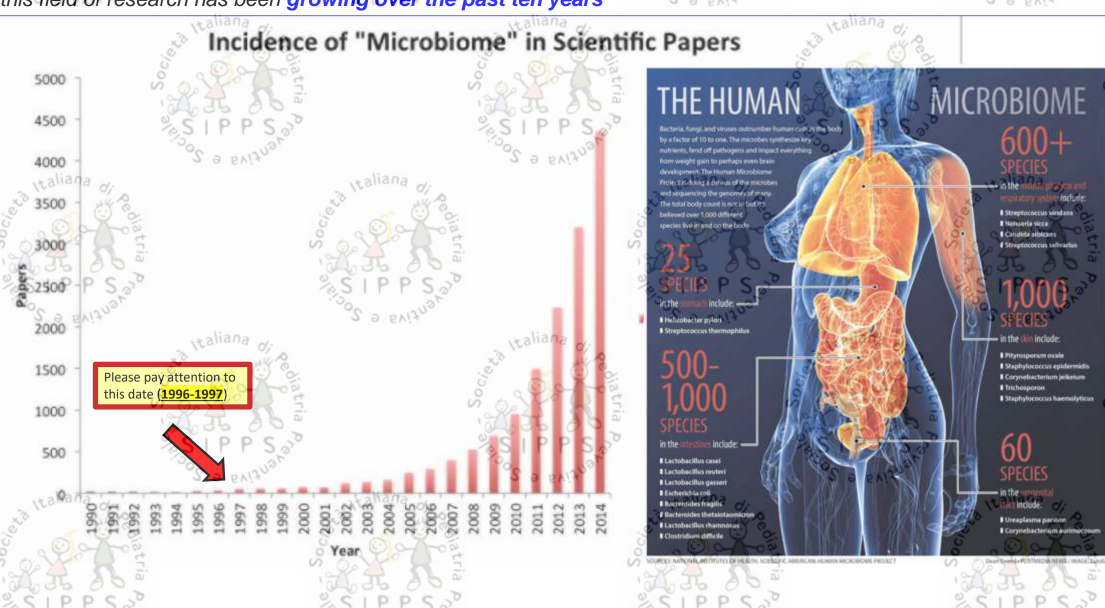


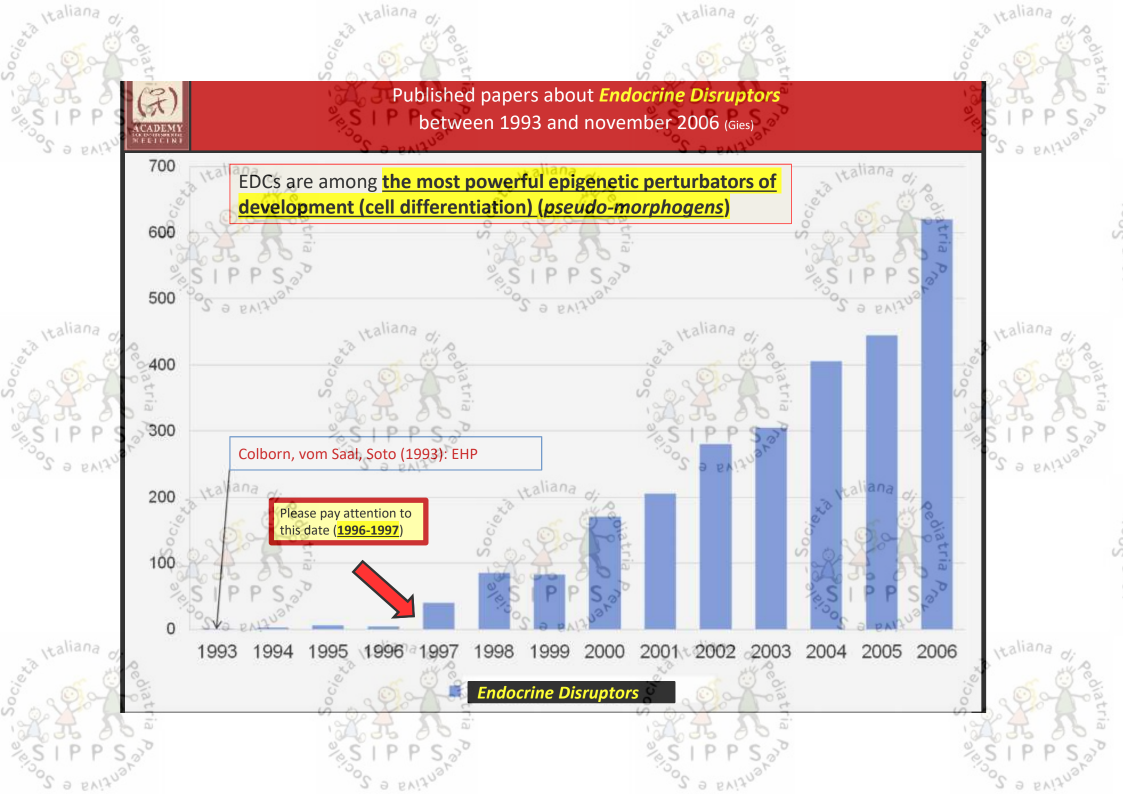


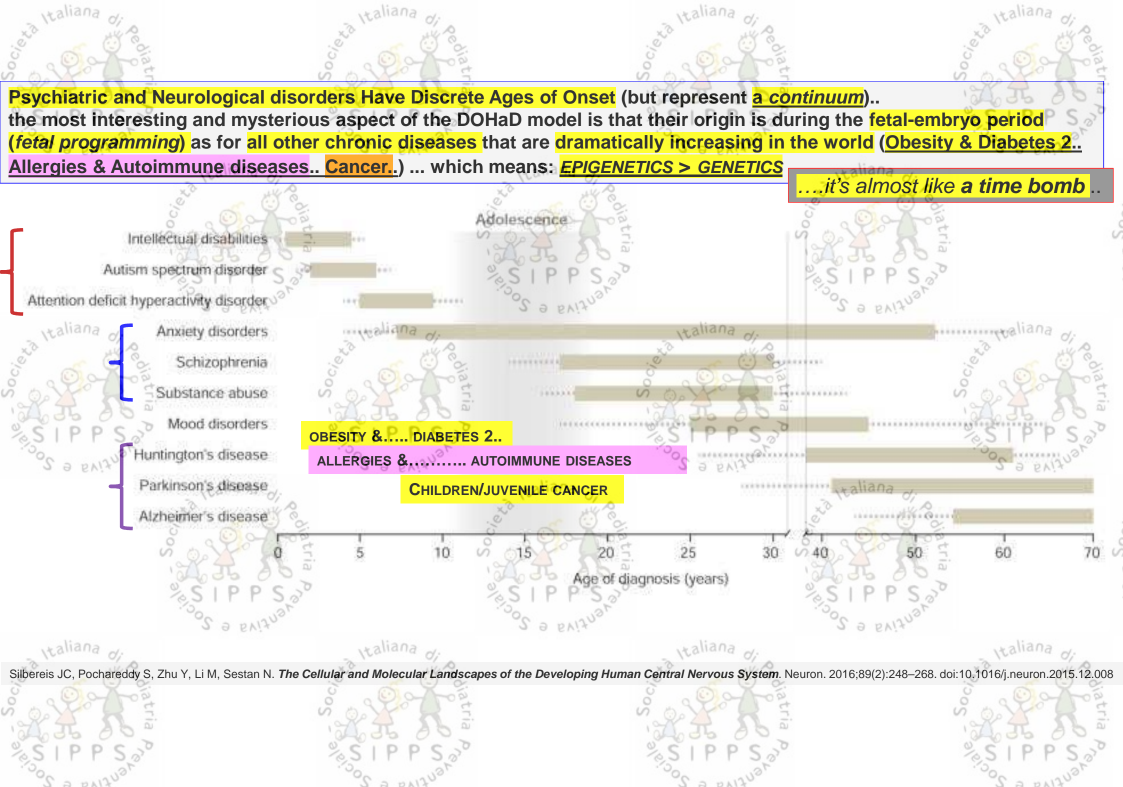




The microbiome is the most powerful "epigenetic internal modulator" of early childhood A quick search for "Microbiome" in scientific journals online demonstrates how significantly this field of research has been growing over the past ten years







The raise of Neurodevelopmental Disorders (NDS): from genetics to epigenetics

Ernesto Burgio, ECERI European Cancer and Environment Research Institute, Bruxelles

e mail erburg@libero.it

The NDS are a set of conditions with onset in the early stages of development and variously associated with cognitive and psychiatric dysfunction. The high heritability of these conditions argues in favor of a genetic component. On the other hand, the impressive increase of NDS calls into question environmental factors and epigenetic mechanisms.

From a neurobiological point of view autism involves early brain overgrowth and dysfunction that may be related to abnormal laminar development and cortical disorganization of neurons, in prefrontal and temporal cortical areas, where social, emotional, communication and language functions are located.

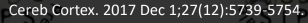
The **Human Connectome Project**



Autism and autism spectrum disorders (ADS) are <u>developmental</u> <u>disorders of neural connections</u> and of <u>synaptogenesis</u>

This affects <u>the way in which the brain "processes information"</u>

"We know that synapses are essential for learning, memory, and perception and suspect that imbalances in synapse formation impact disorders of the brain such as autism and schizophrenia," says Elva Diaz, assistant professor of pharmacology at UC Davis. "Our study is the first to identify SynDiG1 as a critical regulator of these important brain connections."



Dysregulation of Cortical Neuron DNA Methylation Profile in Autism Spectrum Disorder.

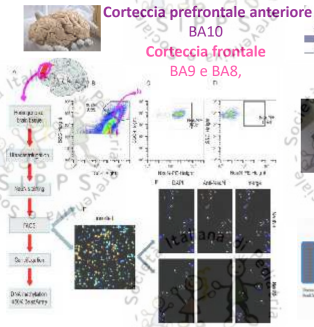
Nardone S et al.

Bar Ilan University Faculty of Medicine, Israel.

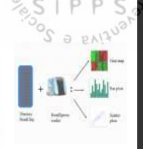
Department of Twin Research and Genetic Epidemiology, King's College London,

Campioni di cervello congelato da 15 casi di ASD e 16 controlli









Banca del cervello di Harvard

Illumina Infinium
HumanMethlation27
BeadChip
target> 450.000

siti di metilazione.

Misura dei livelli di metilazione a 27.578 dinucleotidi CpG in 14.495 geni.

Usando il 450 K BeadArray

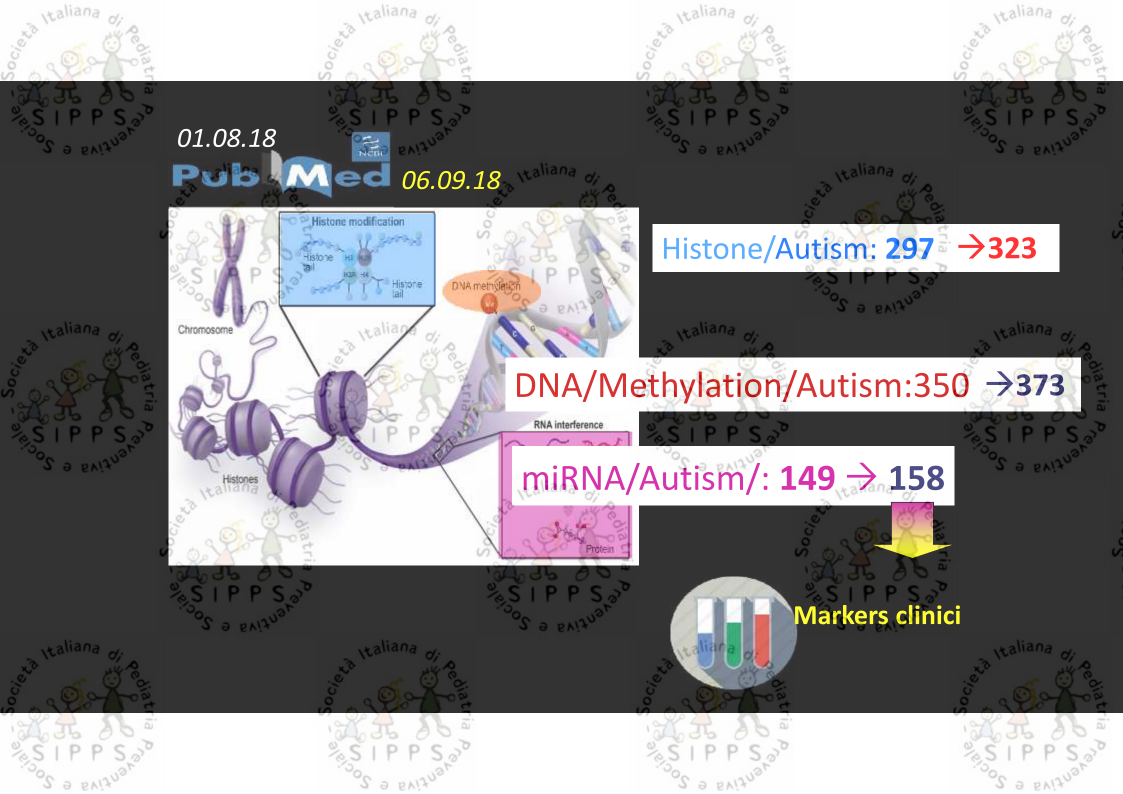
Sono state identificate

58 regioni differenzialmente metilate

che includevano loci associati ai geni del sistema GABAergic

ABAT e GABBR1

Maliana e MicroRNA specifici del cervello



Variation of global DNA methylation in autistic children. The box plot shows methylation levels in healthy children (black) autistic children (red). ... Here, we have demonstrated that the global methylation in autistic 100 children was increased compared to healthy children... Moreover, in comparison with the time profile for methylation, the higher methylation level is that expected of young to middle-aged 80 adults and this could be interpreted to suggest an abnormally Global DNA methylation (% advanced methylome in autistic children. This is reflected in that no significant difference in methylation was found between autistic children and their parents. (a) 20 Healthy Autistic Age groups (Years) children children

Transposable elements can be seen as a natural genetic engineering system capable of acting not just on one location at a time but on the genome as a whole ... This dynamic view of the genome has been illustrated most impressively by Shapiro who stated that the genome is composed of modular units arranged in a "Lego-like" manner that can be altered under circumstances

ELSEVIER

Available online at www.sciencedirect.com



Gene 345 (2005) 91-100



Review

A 21st century view of evolution: genome system architecture, repetitive DNΛ, and natural genetic engineering

James A. Shapiro a RA

Department of Biochemistry and Molecular Biology University of Chicago. 920 E. 58th Street, Chicago. IL 60037, United States

The last 50 years of molecular genetics have produced an abundance of new discoveries and data that make it useful to revisit some basic concepts and assumpt in our thinking about genomes and evolution. Chief among these observations are the complex modularity of genome organization, biological ubiquity of mobile and repetitive DNA sequences, and the fundamental importance of DNA rearrangements in the evolution of sequenced genomes. This review will take a broad overview of these developments and suggest some new ways of thinking about genomes as sophisticated informatic storage systems and about evolution as a systems engineering process.

4

5



Current Opinion in Genetics & Development

Volume 23, Issue 3, June 2013, Fages 264-270



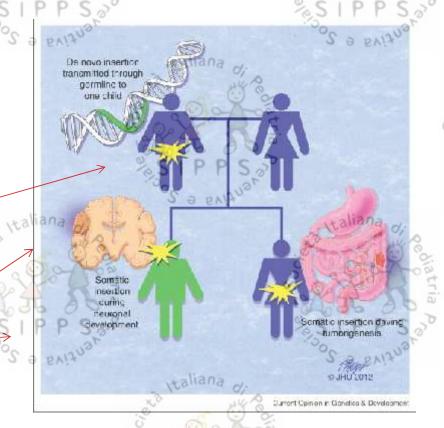
Functional impact of the human mobilome

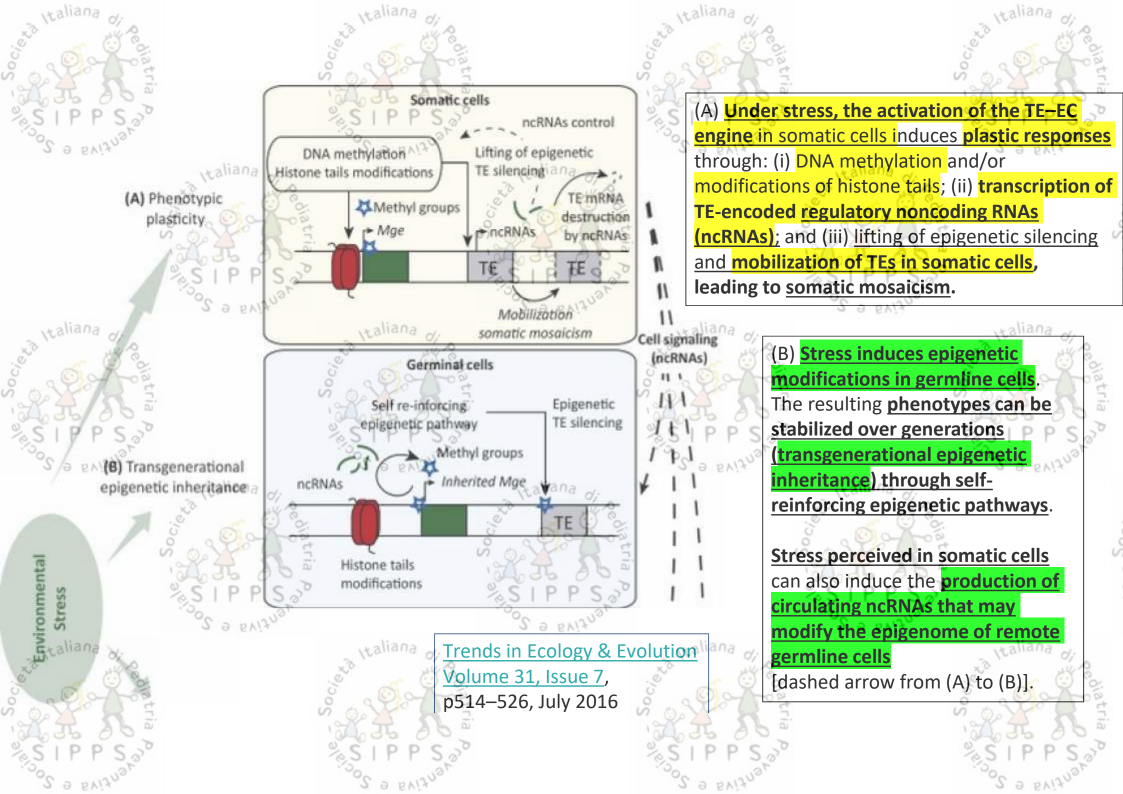
Timothy D Babatz 1, 2, Kathleen H Burns 1, 2, 3, 1

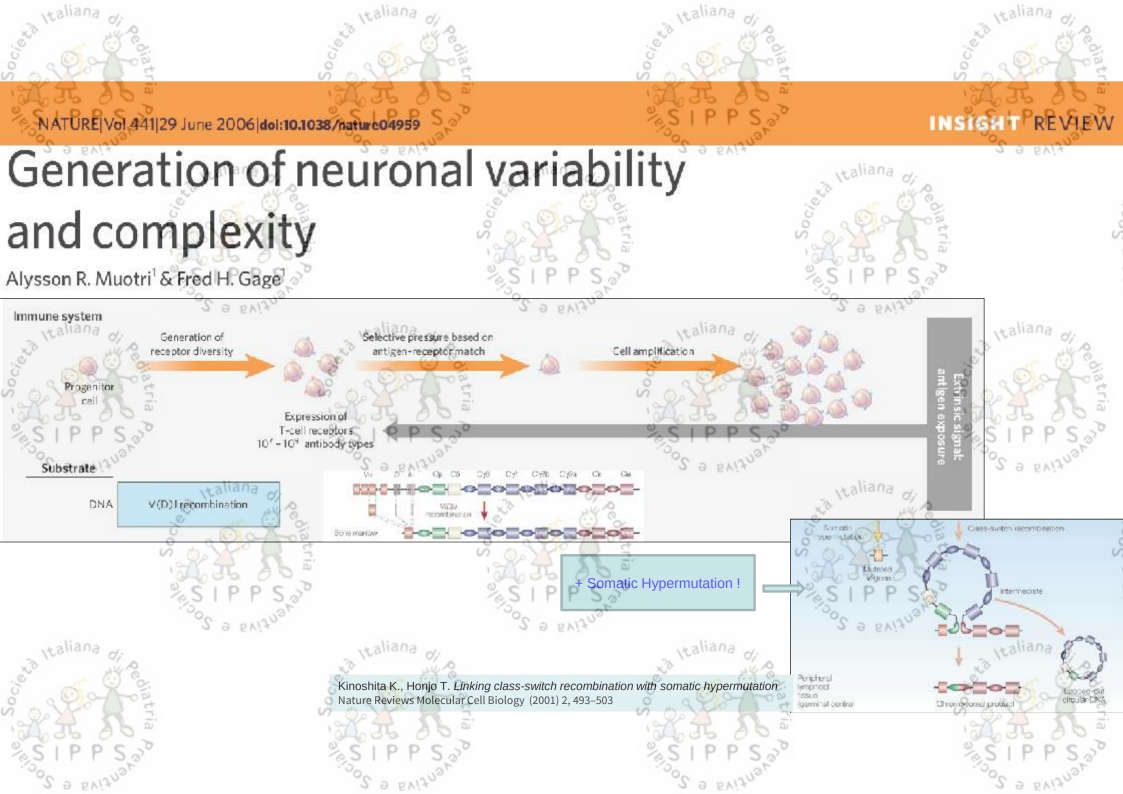
Three families of human retrotransposons remain active today: LINE1, Alu, and SVA elements. Since 1988, de novo insertions at previously recognized disease loci have been shown to generate highly penetrant alleles in Mendelian disorders. Only recently has the extent of germline-transmitted retrotransposon insertion polymorphism (RIP) in human populations been fully realized. Also exciting are recent studies of somatic retrotransposition in human tissues and reports of tumor-specific insertions

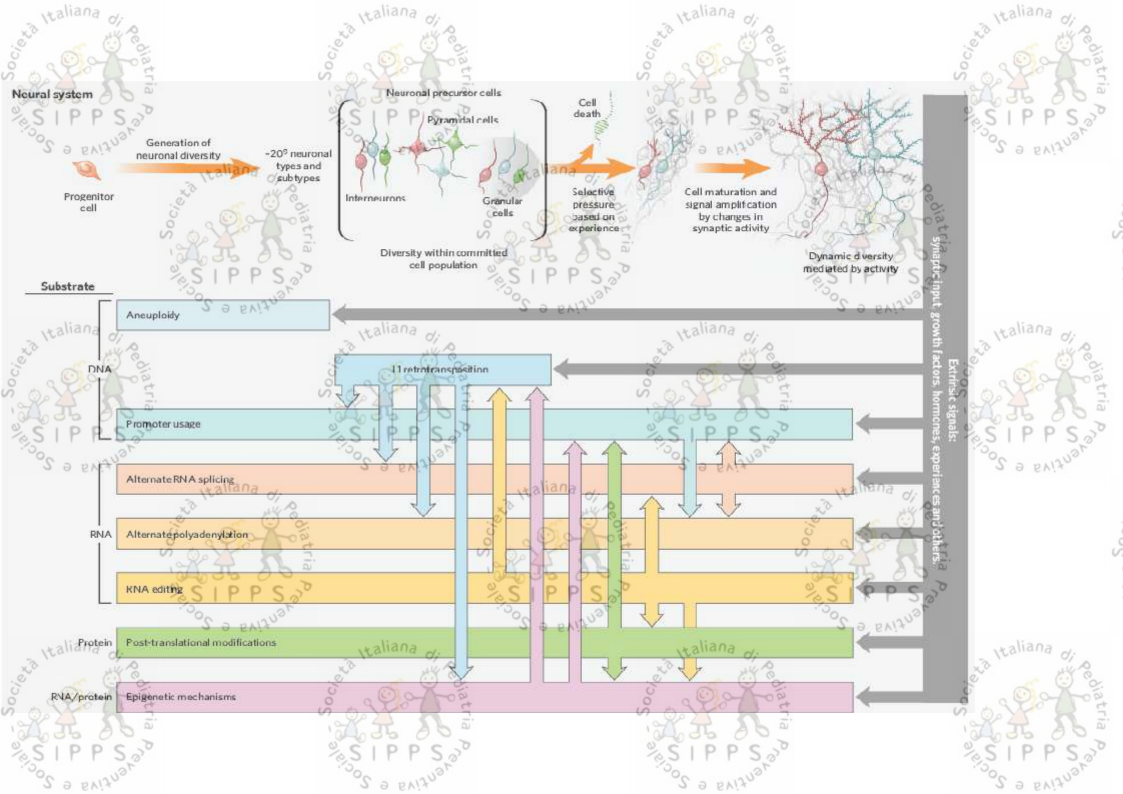
(Stochastic versus Active/Reactive or even Pro-evolutive)

taliana,









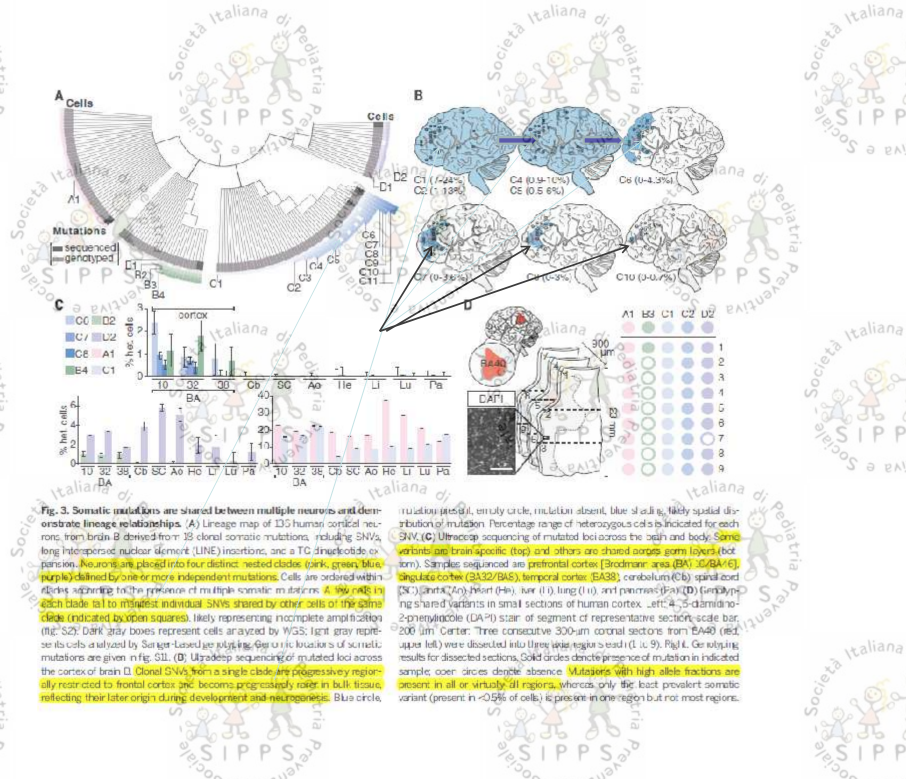
NT S PAIN POCTOBER 2015 - VOL 350 ISSUE 5256

Somatic mutation in single human neurons tracks developmental and transcriptional history

Michael A. Lodato, ** Mollie B. Woodworth, ** Semin Lee, ** Gilad D. Evrony, *
Bhaven K. Mehta, ** Amir Karger, ** Soohyun Lee, ** Thomas W. Chittenden, ***, **
Alissa M. D'Gama, ** Xuyu Cai, ** Lovelace J. Luquette, ** Eunjung Lee, **.

Peter J. Park, **, ** Christopher A. Walsh ** S

Neurons live for decades in a postmitotic state, their genomes susceptible to DNA damage. Here we survey the landscape of somatic single-nucleotide variants (SNVs) in the human brain. We identified thousands of somatic SNVs by single-cell sequencing of 36 neurons from the cerebral cortex of three normal individuals. Unlike germline and cancer SNVs, which are often caused by errors in DNA replication, neuronal mutations appear to reflect damage during active transcription. Somatic mutations create nested lineage trees, allowing them to be dated relative to developmental landmarks and revealing a polyclonal architecture of the human cerebral cortex. Thus, somatic mutations in the brain represent a durable and ongoing record of neuronal life history, from development through postmitotic function.



S I P P S



A Mechanism for Somatic Brain Mosaicism

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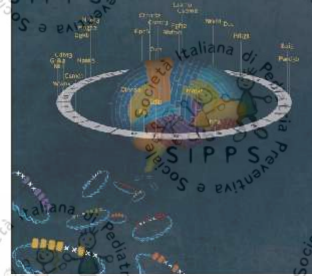
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Double-strand break repair is required for neural development, and brain cells contain somatic genomic variations. Now, Wei et al. demonstrate that neural stem and progenitor cells undergo very frequent DNA breaks in a very restricted set of genes involved in neural cell adhesion and synapse function.

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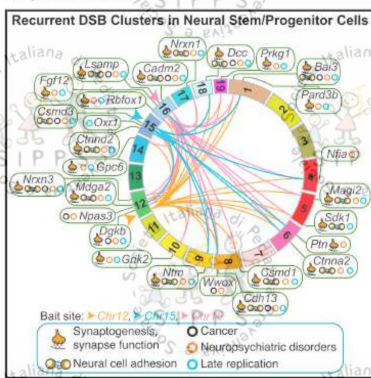
Many of the identified genes are expressed in NSPCs located in the brain regions responsible for higher functions such as short-term learning, and mutations in these genes in humans are associated with (and maybe predispose to) psychiatric and neurological disorders manifested in mind functions—autism, manic depressive and depressive disorders, schizophrenia, and others





Long Neural Genes Harbor Recurrent DNA Break Clusters in Neural Stem/Progenitor Cells

Graphical Abstract



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In Brief

Neural stem and progenitor cells undergo massive genomic alterations in a very restricted set of genes involved in synapse function and neural cell adhesion, processes that are likely to govern the special behavior of brain cells. Many of these genes have also been implicated in mental disorders.

Highlights

- 1) 27 Recurrent DSB clusters (RDCs) are identified in neural stem/progenitor cells
- 2) All RDCs are within genes, most of which are long, transcribed, and late replicating
- 3) Most RDC genes are involved in synapse function and/or neural cell adhesion
- 4) A nucleotide-resolution view of replication stress-associated fragile sites is provided

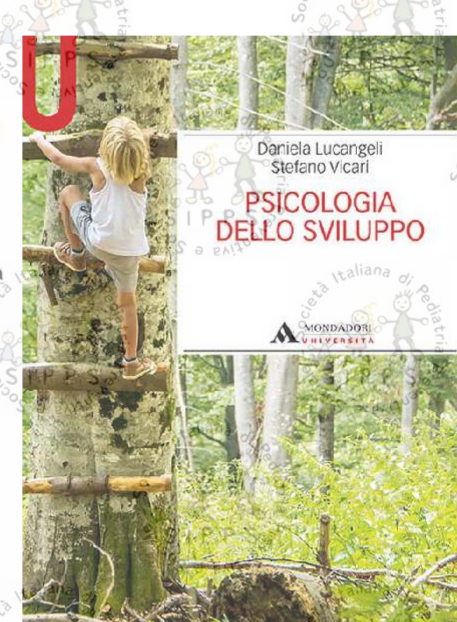
Capitolo 3. Disturbi del neurosviluppo: dalla genetica all'epigenetica, di Ernesto Burgio, Daniela Lucangeli e Maria Antonietta De Gennaro

- 1. I disturbi dello spettro autistico nell'ambito dei disturbi del neurosviluppo
- 2. Dati epidemiologici: aumento reale o semplice incremento di diagnosi?
- aliana 3. Verso un nuovo paradigma: dalla genetica lineare alla genomica sistemica (epigenetica, metagenomica, ologenomica)
 - Narture e Nature
 - 5. Filogenesi e ontogenesi: genetica ed epigenetica
 - 6. I fattori di rischio
 - 7. Il cervello nell'adolescente
 - 8. Epigenetica vs genetica

In sintesi

Domande per l'autoverifica

Bibliografia



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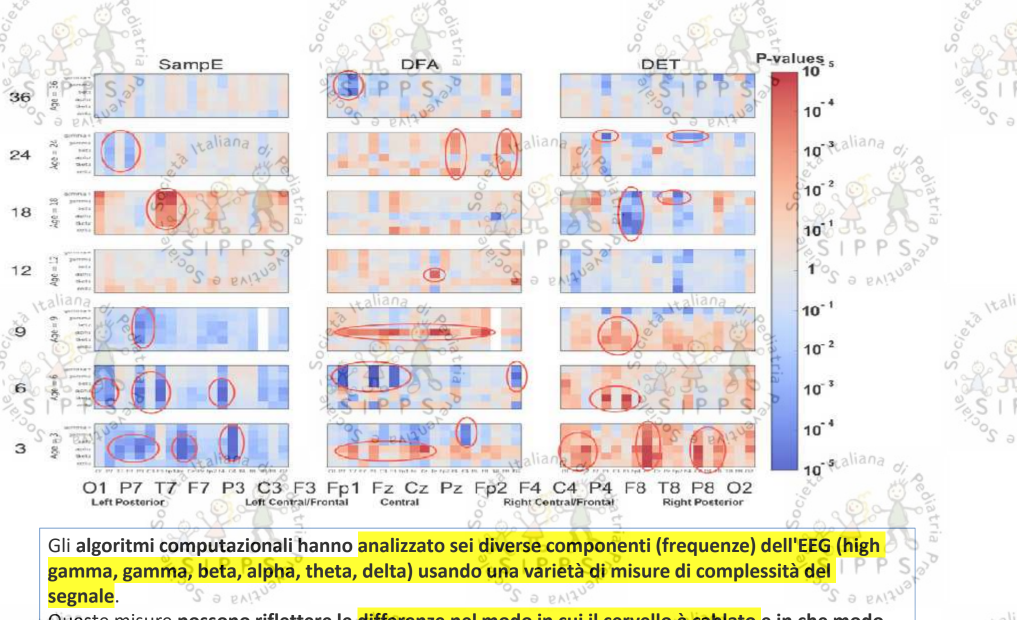
EEG Analytics for Early Detection of Autism Spectrum Disorder: A data-driven approach

William J. Bosl^{1,2,3}, Helen Tager-Flusberg⁴ & Charles A. Nelson^{1,2,5}

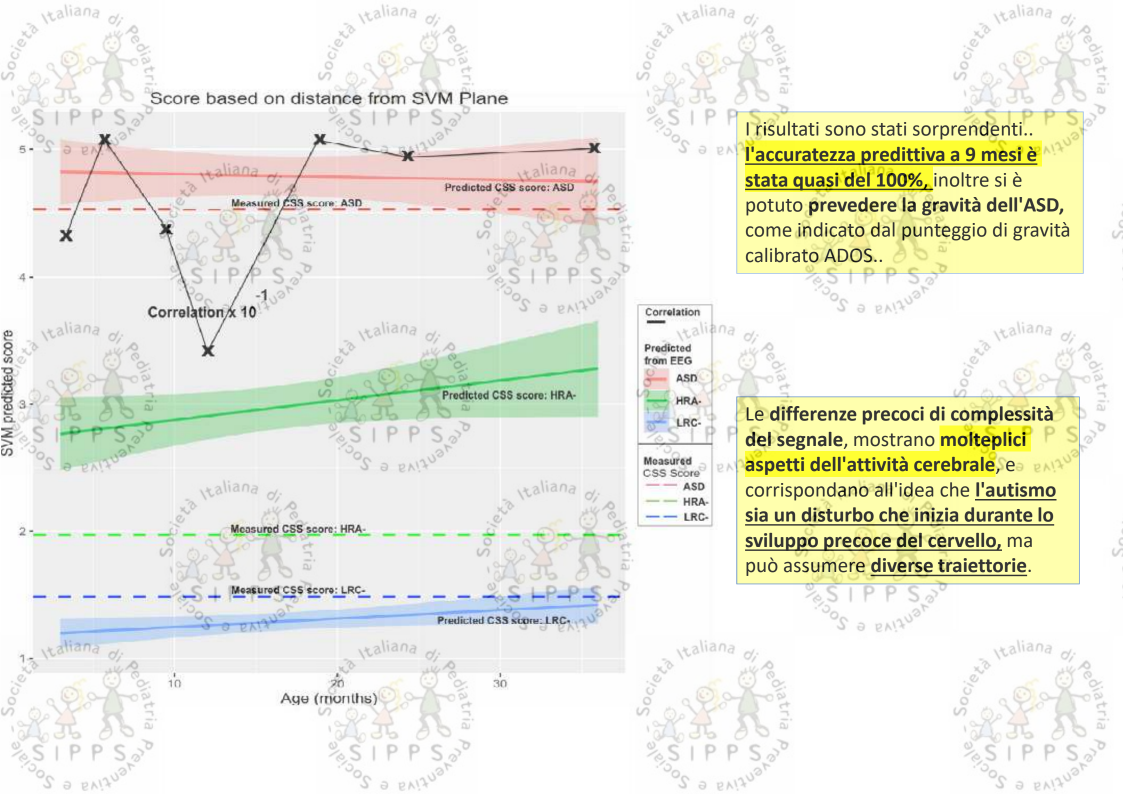
Received: 15 August 2017 Accepted: 28 March 2018 Published online: 01 May 2018

Autism spectrum disorder (ASD) is a complex and heterogeneous disorder, diagnosed on the basis of behavioral symptoms during the second year of life or later. Finding scalable biomarkers for early detection is challenging because of the variability in presentation of the disorder and the need for simple measurements that could be implemented routinely during well-baby checkups. EEG is a relatively easy-to-use, low cost brain measurement tool that is being increasingly explored as a potential clinical tool for monitoring atypical brain development. EEG measurements were collected from 99 infants with an older sibling diagnosed with ASD, and 89 low risk controls, beginning at 3 months of age and continuing until 36 months of age. Nonlinear features were computed from EEG signals and used as input to statistical learning methods. Prediction of the clinical diagnostic outcome of ASD or not ASD was highly accurate when using EEG measurements from as early as 3 months of age. Specificity, sensitivity and PPV were high, exceeding 95% at some ages. Prediction of ADOS calibrated severity scores for all infants in the study using only EEG data taken as early as 3 months of age was strongly correlated with the actual measured scores. This suggests that useful digital biomarkers might be extracted from EEG measurements.

L'autismo è difficile da diagnosticare, soprattutto all'inizio della vita. Un nuovo studio su *Scientific Reports* mostra che **EEG** (oltretutto poco costosi) predicono accuratamente o escludono il disturbo dello spettro autistico (ASD) in neonati di appena 3 mesi.



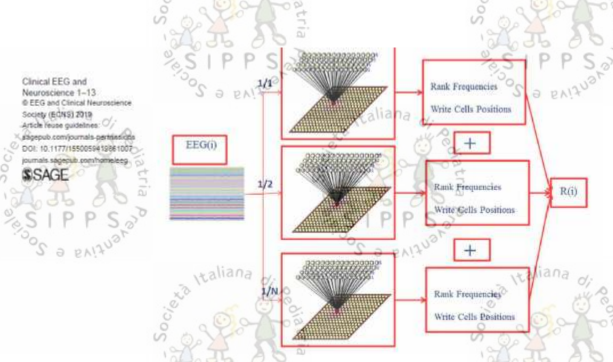
Queste misure possono riflettere le differenze nel modo in cui il cervello è cablato e in che modo elabora e integra le informazioni



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The "MS-ROM/IFAST" Model, a Novel
Parallel Nonlinear EEG Analysis Technique,
Distinguishes ASD Subjects From Children
Affected With Other Neuropsychiatric
Disorders With High Degree of Accuracy

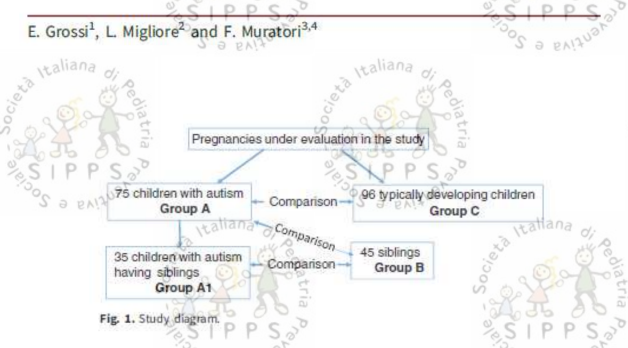
Enzo Grossi 10, Massimo Buscema 2,3, Francesca Della Torre 2, and Ronald J. Swatzyna 4

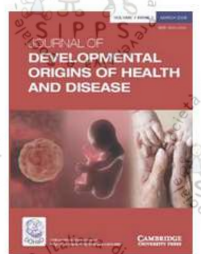


Abstract

Background and Objective. In a previous study, we showed a new EEG processing methodology called Multi-Scale Ranked Organizing Map/Implicit Function As Squashing Time (MS ROM/IFAST) performing an almost perfect distinction between computerized EEG of Italian children with autism spectrum disorder (ASD) and typically developing children. In this study, we assessed this system in distinguishing ASD subjects from children affected with other neuropsychiatric disorders (NPD). Methods. At a psychiatric practice in Texas, 20 children diagnosed with ASD and 20 children diagnosed with NPD were entered into the study. Continuous segments of artifact free EEG data lasting 10 minutes were entered in MS-ROM/IFAST. From the new venables created by MS-ROM/IFAST only 12 has been selected according to a correlation criterion. The selected features represent the input on which supervised machine learning systems (MLS) acted as blind classifiers. Results. The overall predictive capability in distinguishing ASD from other NFD cases ranged from 93% to 97.5%. The results were confirmed in further experiments in which Italian and US data have been combined, in this analysis, the best MLS reached 95.0% global accuracy in 1 out of 3 classes distriction (ASD, NPD, controls). This study demonstrates the value of EEG processing with advanced MLS in the differential diagnosis between ASD and NPD cases. The results were not affected by age, ethnicity and technicalities of EEG acquisition, confirming the existence of a specific EEG signature in ASD cases. To further support these findings, it was decided to test the behavior of already trained neural networks on 10 Italian very young ASD children (25-37 months). In this test, 9 out of 10 cases have been correctly recognized as ASD subjects in the best case. Conclusions. These results confirm the possibility of an early automatic autism detection based on standard EEG.

Pregnancy risk factors related to autism: an Italian case-control study in mothers of children with autism spectrum disorders (ASD), their siblings and of typically developing children





Journal of Developmental Origins of Health and Disease

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Original Article

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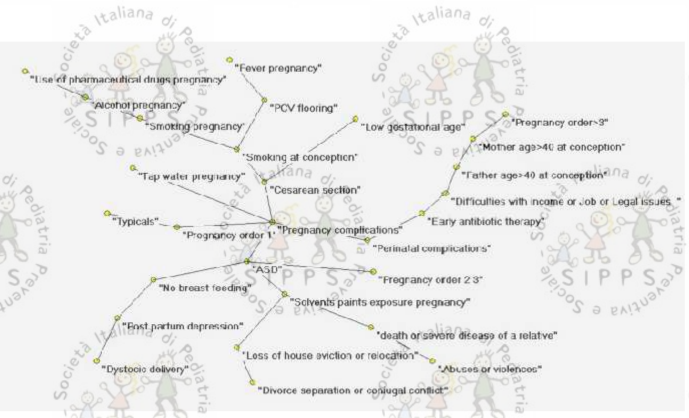
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SIPF	2 0'	100	F 30	Odds ratio	P.value	95% CI	SIPPS		SIPPS
.000	Pregnancy order 1	45.83%	39.13%	1.31	0.390	0.7-2.45	.30C - 1120St		.00C - 1130St
2 9 EV	Pregnancy order 2-3	50.00%	∂2939%\ ¹	2.4	0.010	1.26-4.58	2 9 BVit	line.	2 9 BVit
	Pregnancy order >3 taliana of	4.17%	0.00%	n.d.	redian			a Italiana of	
	Father age > 40 at conception	9.72%	7,61%	1.31 × 0	0.630	0.A3-3.51		in the state of th	
	Mother age > 40 at conception	2.78%	1.09%	2.60	0.440	0.23-29.25		So de diat	
	Smoking at conception	22.22%	15.22%	1,59	0.250	0.51-3.52	60%		- (
:	Smoking pregnaricy) 833%	4.35%	2.01	0.300	0.54-7,37		cally developing group	*
	Alcohol pregnancy S P P S	2.78%	2.17%	1.28	0.800	F0.17-9.35	50% Syb	lings group 45	.1
	Solvents/paints expositing pregnancy	23.61%	2.17%	13.91	0.001	3.09-62.52	40%	S S BVIJAS,	1 + Y
taliana	PVC flooring	18.06%	25.00%	0.66	0,290	0.3-1.41		ism group 73	. sasilem.
-3 Callaria	Tap water pregnancy	23.61%	18.48% 0/	1.36	0,420	0.63-2.9	30%/48/1003		3 010
Salar	Number of stressful events per mother	0.44	0.13	t-test.	0.002		. 020%		
	Death or severe disease of a relative	5.56%	3.26%	1.74	0.470	0.37-8.05	S 10%		
V 20 25	Divorce, separation or conjugal conflict	8.33%	0.00%	,n.d.	n.d.	1	S 10%		
0000	Loss of house, eviction or relocation	11.11%	3,26%	3.7	0.060	0.94-14.52	1000		A G I B BAC A
631 P	Abuses or violences	1.39%	T0.00% 30	n.d.	n.d.		Salar Pena grand	and letter white water defer to	The state of the s
SURN	Officulties with income or job	13.89%	3.26% 200	4.78	0.020	1.26-18.09	reales metal market in part	and a religion white the feet could	of the state of th
	Postpartum depression	15.28%	7.61%	2.19	0.120	0.8-5.97	Sign Siege, Stop, Story 64	god more marking march	Secretary Supplies Secretary Secreta
	Fever pregnancy	11.11%	10.87%	1.02	0.960	0.38-2.74	odine , see	Scale of State of Sta	d _n
	Use of drugs pregrancy	0.00%	2.17%	nd.	n.d.	000	spages, squee, there		
	Pregnancy complications	48,61%	14.13%	5.73	<0.0001	2,72-12.01	40. 0 Gin	SS Solo Total	
	Dystocic delivery	556%	3.26%	1.74	0.470	0.37-8.05	<u>.</u>	10 A A A	
	Cesarean delivery	35,94%	9.78%	4.32	0.001	1.85-10.1	7	00000	
	Perinatal complications	237.50%	13.04%	4 %	0.000	1.85-8.65		621120	
	Low gestational age S a PNIZ	16.67%	6.52%	2.87	S0.040 E	A 1.02-8.06		SIPPS S	
Raliana	No breastfeeding	31.94%	aliterry	3.85	0.001	1.69-8.76	taliana o		italiana ~.
× N	Early attribiotic therapy	16.67%	7.61%	2.43	0.08	0.9-6.52	10 M		13 HK 0
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Demographics	Abuses or violences
Pregnancy order 1	Job strain
Pregnancy order 2-3	Average number of stressful a PAIR
Pregnancy order >3	Health problems during pregnancy
Father age at conception	Fever
Mother age at conception	Use of drugs a "Lis
Behavior/environment	Pregnancy complications
Smoking at conception	Delivery problems
Smoking during pregnancy	Dystocic delivery
Alcohol during pregnancy	Cesarean section
Occupational exposure to solvents/paints	Perinatal complications
Drinking tap water	Postpartum 35 35
SPVC flooring at home	Low gestational age STPPS
Stressful events	Breastfeeding S a PAIZUS
Death or severe disease of a relativ	Early antibiotic therapy
Divorce, separation or conjugal conflict	Postpartum depression
Loss of house, evicted or relocation	A 5



La <u>Fig. 3</u> mostra la mappa di connettività semantica dei fattori in studio ottenuti con la rete neurale Auto-CM dai dati utilizzati per generare la <u>Tabella 2</u>. Il nodo di autismo, alla varianza del nodo tipico, funge da hub (variabile con tre o più collegamenti) che riceve la convergenza da più fattori, suggerendo l'esistenza di un effetto cumulativo multi-causale.

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