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1 2	Chronic Ill Effects of Air Pollution in Children: Neurological Derailment
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7 Abstract

8 Exposure to complex mixtures of air pollutants produces inflammation in the upper and lower

⁹ respiratory tract. Because the nasal cavity is a common portal of entry, respiratory and

¹⁰ olfactory epithelia are vulnerable targets for toxicological damage. The brain is a target for

¹¹ several environmental substances that may or may not be primarily airborne.

¹² Neurodevelopment and neuro-behaviour largely reflect brain development and its chemically

¹³ induced modification, with resulting delays or deficits in development. It is generally believed

14 that the developing brain is a particularly vulnerable target for chemical insult, and that such

¹⁵ insult may have long lasting or even irreversible developmental consequences. Environmental

¹⁶ exposures in uterus and during early life may permanently change the body?s structure,

17 physiology and metabolism, and lead to diseases in adult life. Infants are particularly

¹⁸ vulnerable because of their rapid growth and cell differentiation, immaturity of metabolic

¹⁹ pathways and development of vital organ systems. The central nervous system has

 $_{\rm 20}$ $\,$ unprotected barriers and a broad time window of conformation, leading to a long period of

 $_{\rm 21}$ $\,$ vulnerability in the developmental process and to susceptibility to any environmental insult

²² leading to neurological diseases.

23

24 Index terms— Air pollution, neurological derailment, children.

²⁵ 1 INTRODUCTION

26 I.

27 2 BACKROUND

M arch 2012 complex mixture of gases, particulate matter (PM), and chemicals present in outdoor and indoor 28 air produces adverse health effects. Because the nasal cavity is a common portal of entry for such pollutants, 29 the nasal olfactory and respiratory mucosa are vulnerable to damage and well-known targets for air pollutant-30 induced toxicity and carcinogenicity (Henriksson J & Tj"alve H, 2000 and Morgan KT & Monticello TM, 31 1990). The nose-brain barrier depends on intact epithelia, including tight junctions and an intact xenobiotic 32 33 metabolizing capacity (Dahl AR & Lewis JL, 1993). Olfactory receptor cell dendrites are in direct contact with 34 the environment, and, thus, pinocytosis and neuronal transport are likely routes of access to the central nervous 35 system (CNS) of potential toxins (Lin DM & Ngai J, 1999). Olfactory receptor neurons project from the sensory epithelium to targets within the olfactory. 36

A lead, mercury and (less frequently) manganese, whereas, the most extensively studied PHAH species include the polychlorinated biphenyls (PCBs).

Ambient air pollution is a global health problem and it is an important factor associated with morbidity and mortality worldwide (Narayan, Ali et al. 2010). In high income countries air pollution was associated with 2.5% of all deaths, making it the eighth leading risk factor for mortality (Narayan, Ali et al. 2010). Pollutants do not

5 A) POLLUTANTS OF INTEREST I. PARTICULATE MATTER

42 stay within the national borders. They are carried by winds, contaminating water and soil far from their origin.

43 The great London smog of 1952 focussed the world's attention on the problem of air pollution (LOGAN 1953).
44 Since then there has been an improvement in air quality in developed countries, but a in developing countries air

⁴⁵ pollution continues to be a big problem. The air quality in large cities in developing countries is remarkably poor

46 and large numbers of people living in these cities are exposed to levels of air pollutants well above the World

47 Health Organization guidelines for air quality (Kim, 2004).

Traffic-related air pollution, basically urban outdoor pollution is a global public health problem. Cardio-48 respiratory effects and mechanisms have been fully investigated. In contrast, little is known regarding neurological 49 effects, with only some preliminary evidence. In rats, ultrafine carbon particles have been found in the olfactory 50 bulb and the cerebrum and cerebellum after inhalation exposure, his finding has been reproduced more recently 51 with manganese particles directly translocated to the olfactory nerve from the nose to the brain. In one study, 52 dogs living in a highly polluted region in Mexico City (Mexico) had an increase in brain inflammation compared 53 with animals living in a less polluted area (Calder 'on-Garcidue?nas L, Azzarelli B, Acuna H, et al. 2002). 54 The brain tissue of animals from Mexico City had higher levels of nuclear factor-k Bactivation and nitric oxide 55 production, as well as the principal pro-inflammatory cytokines interleukin (IL)-1 and tumour necrosis factor 56 57 (TNF)-a, compared with the animals from the non polluted area (Campbell A, Oldham M, Becaria A, et al. 58 2005). In a study on human autopsies in Mexico City, exposure to severe air pollution has been associated with 59 increased levels of cyclooxygenase (COX)-2 and accumulation of the 42amino-acid form of -amyloid, a cause of neuronal dysfunction (Caldero'n-Garciduen? as L, Reed W, Maronpot RR, et al. 2004). 60 Although urban air pollution crosses geographical and socio-economical boundaries, urban minority popula-61

tions are likely to be more exposed to indoor and outdoor air pollution than other populations (Breysse, Buckley
et al. 2005). Also, children may have increased exposures to air pollutants compared to adults, because of higher
minute ventilation and higher levels of physical activity. They are also more exposed to ambient air pollution,
because they usually spend much more time outdoors than adults (Kim, 2004).

66 **3** II.

67 4 AIR POLLUTION

68 Pollutants are substances in the air that can cause harm to humans and the environment. They arise in different 69 forms like gasses, solid particles and liquid droplets. They can also divided in pollutants from natural sources in the environment, like volcanism and forest fires (when started by lightning) and pollutants that are man-made or 70 anthropogenic, like fossil fuel combustion emitted by traffic, power generation and industry. Both the natural and 71 anthropogenic pollutants can also be classified as either primary or secondary pollutants. Primary pollutants 72 are directly formed and emitted, whereas secondary pollutants are formed in the atmosphere, when primary 73 pollutants react or interact. Since a lot of air pollutants are emitted by natural sources, industry and traffic, it 74 is logical that people are mainly focused on the outdoor air pollution. However, it is important to realize that 75 indoor air pollution (IAP) might also play a role in the origin of neurological effects. Especially in developing 76 countries, most people still rely on solid fuels for cooking and heating, which generates high levels of pollutants 77 like particulate matters and carbon-monoxide. Women and young children may be significantly more exposed 78 79 to this form of air pollution because of their traditional role in the house hold. But also in developing countries 80 IAP plays a role, since people tend to spend 80-90% of their time indoors and good ventilation is often lacking. 81 In this review we will mainly focus on the neurological effects of ambient air pollution. Toxicological studies will give an indication of the possible effects the air pollutants can have on the neurological development, whereas 82

 $\,$ case studies will give insight into the extension of these effects in children.

⁸⁴ 5 a) Pollutants of Interest i. Particulate matter

One of the most important groups of primary pollutants is the group of fine particles, also known as particulate 85 matter (PM). This PM contains particles from different sizes, sources and chemical composition. Based on size, 86 they are divided in three categories. Particles indicated by PM10 have a aerodynamic diameter of less than 10 87 m. PM2.5 are particles smaller than 2.5 m and PM0.1 are the particles smaller than 0.1 which are also called 88 ultra-fine particles ?? Miyata, van Eeden, 2011). Because of their size, the largest particles (2.5-10 m) are not 89 able to penetrate the respiratory tract very deeply. They are mainly deposited in the nasal cavity, larynx and 90 trachea ??Heyder, 2004). The particles that are between 0.1 and 2.5 m penetrate the bronchi and bronchioles. 91 However, the larger particles in this group will still deposit mostly in the upper airway, whereas the particles 92 93 between 0.1 and 1 m hardly deposit at all (NRPB, 2004). The ultrafine particles (<0.1 m) are able to penetrate 94 the alveoli and have a deposition rate of more than 80% (NRPB). The origin of particulate matter can be natural 95 (volcanoes, dust storms and forest fires), but mostly they are emitted as a result of fuel combustion from traffic 96 and industry or agriculture. ii. Nitrogen dioxide & Ozone Nitrogen dioxide (NO 2) is formed in most combustion processes that use oxygen 97 98

as an oxidant. NO2 is a highly reactive and nitrogen-centered free radical that can induce airway inflammation,
like asthma and acute bronchitis (Shima & Adachi, 2000). NO 2 is also a major precursor for a number of
harmful secondary pollutants. The most important form of a secondary pollutant is ozone (O 3). Ozone is
formed by photochemical reactions of sunlight on air containing hydrocarbons and nitrogen oxides. This newly

formed ozone reacts with UV light again, resulting in the production of hydroxyl radicals. These hydroxyl radicals are the first step in the creation of smog components, including peroxyacyl nitrates that can be powerful eye irritants. But also neuronal deficits have been shown after O 3 exposure. Upon O 3 inhalation, Gackiere et al. showed a time-and dose-dependant neuronal activation pattern similar to that induced by systemic stress. This chronic stress is known to disrupt sleeping patterns, anxiety, depression and social isolation (Gackiere, Saliba et al. 2011).

¹⁰⁸ 6 b) Air pollution levels

Air pollution is usually concentrated in densely populated areas, especially in developing countries where 109 environmental regulations are relatively lax or nonexistent, but also areas in the developed world can be highly 110 polluted. In many developing countries the absence of surface-based air pollution monitors makes it difficult and 111 sometimes impossible to get even a rough estimate of the abundance of a subcategory of airborne particles that 112 epidemiologists suspect contributes to millions of premature deaths each year. Therefore, van Donkelaar etal. 113 created a map (see figure 1) by blending total-column aerosol measurements with information about the vertical 114 115 distribution of aerosols from a computer model (van Donkelaar et al. 2010). Figure ?? clearly shows that the areas with the highest concentration of PM2.5 are concentrated around the equator and especially in Northern 116 Africa and the rapidly developing countries in Asia. 117

7 BIOLOGICAL CONSIDERATIONS OF HUMAN BRAIN DEVELOPMENT

A brief account of some basic principles of human brain development may help to better understand why and 120 how prenatal or early postnatal environmental exposure to chemicals may give rise to adverse neurobehavioural 121 alterations in infants and children postnatally. This brief overview can only be elementary; for a more detailed 122 account the reader is referred to standard textbooks. The maturation of the central nervous system (CNS) is 123 often described under the four headings of gross morphology, proliferation and migration of neurons and glial 124 cells, neuronal differentiation, and myelinization. By the end of the embryonic stage (12th week of gestation) 125 the organogenesis of the brain already shows division of the with a pronounced enlargement of the thalamus 126 127 and an initial formation of the cerebellum. Towards the end of the 12th week of gestation, separate ventricles occur but the brain surface is still smooth. Its structuring into lobes through the formation of primary sulci 128 (folds) occurs in the 4th month of gestation, such that the main lobes (frontal, parietal, occipital and temporal) 129 become discernible. Among the deeper structures the main hemispheric connections, namely the corpus callosum 130 and the commissurae anterior and posterior, also develop early. Brain damage during these early stages of CNS 131 development gives rise to gross structural anomalies. 132

Following the formation of the primary sulci, secondary sulci are formed during the last three months of 133 gestation, whereas tertiary sulci develop postnatally until the end of the 2nd year of life. It is important to 134 note that the place of origin of neurons and glial cells is different from their final destination in the brain. 135 Cells migrate to form, for example, the six cortical layers and the architecture of other brain structures. 136 Besides migration, the maturation of neurons from their precursor cells, the neuroblasts, is another important 137 developmental principle. Maturation includes enlargement of the cell body, storage of Nissl substance, formation 138 of neurofibrils, arborization of axons and apical dendrites, and finally the increasing number of synaptic contacts 139 between neurons. Although neuronal differentiation begins prenatally (the six cortical layers, for example, being 140 already present around the 28th gestational week), much neuronal maturation extends into the first two postnatal 141 years, such as the arborization of dendrites and synapse formation. Also, much of the myelinization of fibres 142 occurs postnatally. Thus, both the prenatal and early postnatal phases of CNS development offer opportunities 143 for chemical insult. It must finally be pointed out that many of the abovementioned developmental processes 144 and their timely orchestration, namely proliferation, migration, differentiation and myelinization of neurons, 145 are partly under endocrine control. One prominent example is the hypothalamic-pituitary-thyroid (HPT) axis. 146 Clinical observations show that severe congenital or dietary hypothyroid conditions during pregnancy or in the 147 neonatal stage, if untreated, often result in cretinism associated with mental retardation. This is one mechanistic 148 possibility of how chemicals interacting with endocrine systems may also interfere with brain development and 149 associated dysfunctional neurobehavioural development postnatally (Porterfield S, 1994). 150

151 **8** a) Brain

152 The brain, or encephalon (derived from Greek), is the centre of the nervous system. It is the organ that 153 is responsible for sensing, controlling and processing signals and is therefore found close to primary sensory 154 apparatus like vision, hearing, balance, taste and smell. The brain consists of billions of neurons each connected via synapses. Neurons communicate via these synapses with chemical signals called neurotransmitters. Electrical 155 signals are transmitted through the axons that carry action potentials to distant parts of the brain or body 156 to target specific recipient cell. Every function will involve multiple brain regions and every brain region may 157 be involved in several other functions. Therefore, understanding the brain is not simple and straightforward 158 (Silverthorn, 2004). 159

¹⁶⁰ 9 b) Anatomy

In the early embryo, the cells that will form the nervous system are positioned in the neural plate. During the development of the embryo, neural crest cells migrate to the middle and thereby creating a neural tube. The anterior portion of the neural tube will specialize into the regions of the brain, being the forebrain, midbrain and hindbrain. The posterior part of the neural tube will form the spinal cord. Finally the six mainparts of the brain are formed: (1) the cerebrum, (2) the diencephalon, (3) the mesencephalon, (4) the cerebellum, (5) the pons and (??) the medulla oblongata (see figure ??). The cerebrum is the largest and most distinctive part of the human brain and fills most of the cranial cavity (Silverthorn, 2004).

¹⁶⁸ 10 M arch 2012 c) Gray and White Matter

The cerebrum has distinct regions of gray and white matter. The outer layer of the cerebrum, which is only a 169 few millimetres thick, forms the gray matter of the cerebral cortex. Gray matter consists of unmyelinated nerve 170 cell bodies, dendrites and axon terminals. The cell bodies form layers in some parts of the brain or cluster into 171 groups of neurons with a similar function. White matter in the cerebrum is found primarily in the interior. 172 White matter is made up mostly of myelinated axons and contains very few cell bodies. Its pale (white) colour 173 comes from the myelin sheaths that surround the axons. Bundles of fibres allow different regions of the cortex to 174 communicate with each other and transfer information from one hemisphere to the other primarily through the 175 corpus callosum (Purves et al. 2008). 176

177 11 d) Blood Brain Barrier

The blood brain barrier is a separation of circulation blood and the brains extracellular fluid and prevents 178 179 potentially harmful particles from being delivered into the brain. It is formed by the brains capillary endothelium. These endothelial cells form tight junctions that are composed of transmembrane proteins that are anchored in 180 the endothelial cell (see figure ??). The capillary endothelium uses selected membrane carriers and channels to 181 move nutrients and other useful material from the blood to the brains interstitial fluid. Other transporters move 182 wastes from the interstitial fluid into the plasma. If a water soluble molecule is not transported on one of these 183 carriers, it cannot cross the blood brain barrier (Liddelow, 2011). Although the blood-brain barrier excludes 184 many water-soluble substances, smaller lipid-soluble molecules can simply diffuse through the cell membranes. 185 Because of the high demand of oxygen of the brain, oxygen can pass the blood-brain barrier freely. Neurons 186 consume oxygen at such high rates that interruption of the blood flow to the brain can have devastating effects 187 within only a few minutes (Pritchard, 1999). The brain receives sensory input from the internal and external 188 environments. After processing these incoming signals it creates a response. However, the brain is also able 189 to generate information and act without the external input of signals. There are three different systems that 190 influence the output signals, being (1) the sensory system that monitors the internal and external environment 191 192 and initiates reflex responses, (2) the cognitive system that is able to initiate voluntary responses and (3) a behavioural state system, which governs sleep-wake cycles and other behaviours. Cognition refers to mental 193 processes, which include: attention, memory, Understanding a language, solving problems and making decisions. 194 It can be natural or artificial and conscious or unconscious (Silverthorn 2004). 195

¹⁹⁶ 12 f) Toxicology Studies: i. Translocation of Particles to the ¹⁹⁷ Brain

Translocation to extra pulmonary sites after respiratory tract deposition is an important mechanism for particles 198 to cause effects in secondary organs. Whether this process occurs and to which extent, depends on several factors 199 including particle size, solubility, site of deposition and the integrity of the epithelial lining. Elder et al. showed 200 that ultra fine and fine particles can translocate from the lungs by penetrating pulmonary tissue and enter the 201 capillaries, reaching other organs, including the brain by circulation (Elder, Gelein et al. 2006). As mentioned 202 before, the blood brain barrier is supposed to inhibit harmful particles from entering the brain. However, some 203 particles are still able to cross the blood brain barrier either because they are small enough to leak through the 204 endothelial tight-junctions, or because they disrupt the blood brain barrier by inflammatory responses. Pollutants 205 can also enter the brain through direct translocation. Animal studies have shown that inhaled particles can be 206 translocated to the brain via the olfactory nerve that connects the nose and the braindirectly ?? 207

²⁰⁸ 13 ii. Neuro-inflammation and Degeneration

209 The first and main form of active immune defense in the central nervous system is formed by the action of 210 microglia. These microglia are a type of glial cells that reside in the brain and spinal cord. They respond to tissue 211 insult with a complex array of inflammatory cytokines and actions. They are recognized as the prime components of an intrinsic brain immune system. Before neuro-inflammation became a commonly used term, scientists used 212 the term 'reactive gliosis' (Streit, Mrak et al. 2004). This term specifically referred to the accumulation of 213 enlarged glial cells, notably microglia and astrocytes, immediately after CNS injury had occurred. Activation 214 of immune cells in the periphery leads to leukocyte infiltration of tissues, but this leukocyteinfiltration is absent 215 in the brain, unless there has been destruction of the blood brain barrier. Without breakdown of the blood 216

brain barrier, leukocytes are not able to crossthis barrier and there is a much subtler response of the brains own 217 immune system. Although these specific responses might be included in the term neuro-inflammation, this term 218 generally applies to a more chronic, sustained cycles of injury and response. This chronic microglia activation 219 likely contributes to injury, loss of neurons and neuronal dysfunction (Bellucci, Westwood et al. 2004). Neuro-220 degeneration is the overall term for progressive loss of structures or function of neurons and axons in the central 221 nervous system. Immune activation in the CNS is a classical feature of neuro-degeneration iii. IL-1 and COX-2 222 Among the pro-inflammatory molecules, cytokines are thought to play a central role in the selfpropagation of 223 neuro-inflammation, with a prominent function for interleukin-1(IL-1). IL-1 is family of three related proteins, 224 being Il-1a, Il-1b and Il-1ra. Normally, Il-1 is expressed at low levels, but is upregulated rapidly in response to 225 local orperipheral insults. The specific cellular source of these proteins is unclear, butmicroglia cells appear to 226 be the early primary source. Astrocytes and neurons have also been reported to express II-1. (Pearson, Rothwell 227 et al. 1999) It remains 16 uncertain whether IL-1 plays a major role in the normal, healthy brain, because the 228 expression is barely detectable. However, IL-1 has been shown to act by increasingfever, hypophagia, slow-wave 229 sleep, sickness behaviour and neuro-endocrinechanges. It's expression is also increased in human degenerative 230 conditions and inhibition of IL-1 in rodents reduced neuronal loss dramatically. (Rothwell & Luheshi, 2000). 231 Cyclooxygenase (COX) generates reactive oxygen species (ROS) as a byproduct of the conversion of prostaglandin 232 233 G2 to prostaglandin H2 in the synthetic pathway of prostaglandins and thromboxanes. There are two different 234 isoforms of COX. COX-1 is expressed and predominant in peripheral tissues, while COX-2 has been shown tobe 235 expressed at high levels in the CNS and is induced by a variety of stimuli (Yamagata, Andreasson et al. 1993). It is rapidly up regulated at sites of inflammation and it is primarily expressed by neurons, whereas microglia 236 and astrocytes are almost unlabelled. Given the fact that oxidative stress is involved in neurodegeneration, it is 237 likely that the induction of COX-2 and the generation of freeradicals by this protein are related to the underlying 238 mechanism ??Oka & Takashima,1997). 239

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M arch 2012 iv. White Matter Lesions White matter lesions are small area's of disrupted neurons in the white 242 matter, commonly seen in older people, since it is a normal result of aging (Sierra, 2001). However, aging is not 243 the only factor that induces these lesions. They also appear in the brains of people who have suffered stroke or 244 have progressive neurological diseases and they can be induced by exposure to toxicants. While it is not clear 245 that white matter lesions directly cause brain dysfunction, they are seen as good indicators. Namely there is 246 a clear connection between lesions and decreases in brain volume, loss of memory and vision For this study, 247 248 children from two different primary schools were selected to explore the association between traffic-related air 249 pollution and neurobehavioral function in children (Wang, Zhang et al. 2009). The first school was located 3.5km 250 away from primary traffic roads in the north of Quanzhou and had low traffic density. The second was located on a threeway-intersection in the centre of the city. Levels of ambient air PM10 and NO2 were measured for 251 two days at five different school sites. The children were 8-10 years old and selected based on a questionnaire 252 253 about their socioeconomic status and neurobehaviour was tested based on nine standardized tests. The NO2 levels in the polluted area where significantly higher than in the clear area. However, levels of PM10 did not 254 show any differences. Children that were going to school in a polluted area had significantly lower scores on all 255 nine tests, compared to the children that were going to school in de clean area. Therefore, Wang et al. state 256 suggests that there is a significant relationship between traffic related air pollution and neurobehavioral functions 257 258 in children. Similar findings were obtained by Kumar (2011) (Suglia, Gryparis et al. 2008). Pregnant woman 259 over 18 years, receiving prenatal care at an urban community health center in Boston were fitted for enrolment. At each clinic visit during their pregnancy the women were asked about their smoking status and the smoking 260 habits of other members of the household. Also a urine sample was taken to determine the cotinine level. The 261 postnatal exposure of the child to second hand smoke was obtained by a questionnaire every month for the first 262 26 months, twice a year for the period between 26 months to 4 years and annually for the remaining years. 263 To estimate the residential black carbon levels, data was used from pollution measurements at more than 80 264 sites performed at more than 2000 different days. For the follow-up study, children were selected based on their 265 birth weight, blood-leadlevel and ETS exposure. When the children were aged 8-11 years, cognitive tests were 266 administered, including the Kaufman Brief Intelligence Test (K-BIT) for verbal and non verbal intelligence and 267 the Wide Range Assessment of Memory and Learning (WRAML) for verbal and visual memory and learning 268 269 (Putzke, ??illiams et al. 2001). In this study, long-term concentration of black carbon particles from mobile 270 sources was associated with decreases in cognitive test scores, both on verbal and non-verbal intelligence and on 271 memory. Socioeconomic status could have been a confounder, since it can be a determinant of cognitive abilities 272 during childhood. Also it can determine whether a family lives close to traffic areas. However, since all families were recruited from one neighbourhood, the variability in socioeconomic status was restricted and therefore the 273 potential of confounding was reduced. (Suglia, Gryparis et al. 2008). Another study conducted by Kumar and 274 Sharma (2011) on the neurological effects of chronic exposure to black carbon in the children (N=150) of labor 275 class in the Greater Noida, district of U.P. (India) also found a significant deterioration in their cognitive system 276

and brain impairment. 277

²⁷⁸ 15 V. OXIDATIVE STRESS INFLAMMATION OF LUNGS ²⁷⁹ AND BRAIN IMPAIRMENT

Oxidative stress, changes in autonomous function and progression of atherosclerosis have been hypothesised to 280 be mechanisms of the neurological effect of urban air pollution in humans at any age (Peters A, Veronesi B, 281 Caldero'n-Garciduen? as L, et al, 2006). Among them, inflammation secondary to oxidative stress appears 282 to be the major suspected culprit for delay in conformation and maturation during developmental steps. Even 283 though most of the available research about the inflammatory effects of air pollution refers to the lungs, there is 284 evidence that the oxidative stress and inflammation induced by particles translates systemically beyond the lungs 285 286 (Hirano S, Furuyama A, Koike E & Kobayashi T, 2006). For example, we found in an international longitudinal 287 study of 1,003 adult subjects that particle count increased markers of systemic inflammation (IL-6 and fibrinogen peripheral levels) (Ru["] ckerl R, Greven S, Ljungman P, et al. 2007). The major underlying hypothesis is that 288 chronic respiratory tract inflammation may lead to brain inflammation by altering levels of circulating cytokines, 289 such as TNF-a and IL-1. These cytokines have the ability to up regulate COX-2, a potent active mediator of 290 inflammation, in capillary brain endothelium (Campbell A, 2004). Changes in brain cytokine and chemokine 291 expression in mice have been directly linked to intranasal exposure to ultra fine carbon (Tin-Tin-Win-Shwe, 292 Yamamoto S, Ahmed S, Kakeyama M, Kobayashi T, Fujimaki H, 2006). Carbon particles themselves generally 293 adsorb transition metals (including antimony, barium, copper, ironand zinc) emitted from traffic exhaust and 294 also from tyres and brake wear. These metals, which are mainly generated by traffic in the current urban 295 atmospheres (Steiner M, Boller M, Schulz T, Pronk W, 2007), have been show no induce oxidative stress in the 296 lung An alternative hypothetical mechanism of the neurological effect of air pollution is based on the observation 297 that ultra fine particles containing metals translocate directly to the brain without entering the lung (Elder A, 298 Gelein R, Silva V, et al., 2006). Changes in cognitive function in children have been shown to be associated with 299 300 relatively low internal doses of lead (Lanphear BP, Hornung R, Khoury J, et al., 2005) and mercury. In addition to being linked to cognitive deficits in children, lead has been related to a diversity of behavioural problems 301 (reading problems, school failure and delinquent behaviour), with a high social impact. In experimental studies, 302 some metals, such as mercury and lead, inhibit neuronal differentiation, myelinisation and synaptogenesis (303 Johansson C, Castoldi AF, Onishchenko N, Manzo L, Vahter M, Ceccatelli S, 2007), but the specific mechanisms 304 for lead induced intellectual deficits have not been fully elucidated. A well-known constellation of factors related 305 to neurodevelopment could all play a confounding role or they could explain differences in vulnerability of the 306 dose-response relationship between air pollution and neurodevelopment (Bellinger DC, 2008). These factors 307 must be considered and include, for example, the social environment (including parental psychological status), 308 breast feeding, diet, maternal smoking, birth -weight and noise (Clark C, Martin R, van Kempen E, et al. 2006) 309 along with other pollutants such as lead, mercury, DDT and indoor air pollutants (those originating from indoor 310 sources, such as heating and cooking, or from microbial contaminants, such as endotoxins). Endotoxins are the 311 common structural component of Gram-negative bacteria in indoor air that induced chronic inflammation in the 312 rat brain (Lehnardt S, Massillon L, Follett P, et al., 2003). Therefore, it is important to examine diet, since it is 313 a major source of antioxidants. Antioxidant defense mechanisms could be increased by dietary means (vitamins, 314 omega-3 (docosahexaenoic acid) and omega-6 (arachidonic acid) fatty acids, and other micronutrients (zinc and 315 folic acid)) to protect against air pollutants (Kelly FJ., 2004). Antioxidants in the lung are the first line of defense 316 against oxygen free radicals. All of these antioxidants are free radical scavengers and they react rapidly to limit 317 interaction with lung fluid lipids Global Journal of Human Social Science Volume XII Issue VI Version I 318

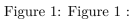
319 16 CONCLUSION

To conclude, the study of chronic effects of air pollution should incorporate subtle health effects, such as functional delays in brain maturation and impairment of neurobehavioural competences, from early life exposures. The longterm consequences of these effects in the co-causation of neurodegenerative diseases have proved that our urban air is neurotoxic and deadly for our children and the chronic inflammatory process elicited within the respiratory tract upon exposure to outdoor and indoor air pollutants could serve as the trigger for a chain of events involving the brain. Hence, necessary steps have to be taken to put a check on this disastrous chain of air pollution.

 $^{^{1}}$ © 2012 Global Journals Inc. (US) pros encephalon into two hemispheres occurs together

 $^{^2 \}odot$ 2012 Global Journals Inc. (US) Global Journal of Human Social Science Volume XII Issue VI Version I 2





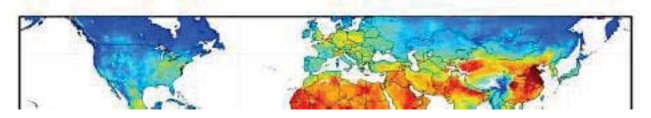


Figure 2: Chronic

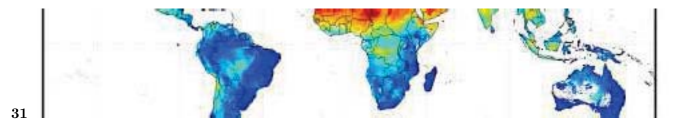


Figure 3: Figure 3 : Figure 1 :

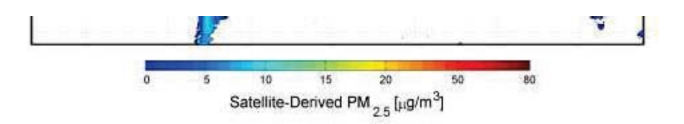


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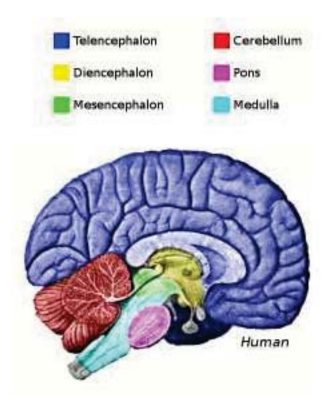


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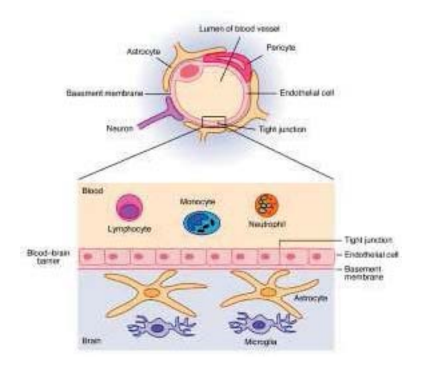


Figure 6:



Figure 7: Chronic

16 CONCLUSION

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