Highlights

Tainted air: The link between pollution and Alzheimer’s disease

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ABSTRACT

In this issue of the Biomedical Journal, we learn how air pollution may contribute to cognitive decline and even increase risk for Alzheimer’s disease. We also highlight original research documenting the body’s response to infection with a common oral pathogen. Finally, we learn how a cellular antioxidant protein protects against mitochondrial dysfunction in Parkinson’s disease.

Spotlight on reviews

Tainted air: the link between air pollution and Alzheimer’s disease

Without prevention, the incidence of Alzheimer’s disease (AD) is expected to triple by the year 2050 [1]. The insurmountable pressure that this would place on health care systems has intensified the search for major preventable risk factors for AD. In this issue of the Biomedical Journal, Kilian and Kitazawa [2] discuss how one such risk factor for AD may literally be in the air we breathe, and describe the potential mechanisms linking pollution to decline in brain function and AD.

Chronic exposure to polluted air is a major health issue worldwide, with the WHO estimating that a staggering 91% of the world’s population lives in places where air pollution exceeds recommended health guidelines [3]. Polluted air contains particulate matter (PM) of various sizes along with noxious compounds such as nitrogen and sulfur oxide species, carbon monoxide, metals and other inorganic compounds. PM contains sulfates, nitrates, ammonium, chlorides, carbon, and other biological material and dust, and is divided according to size: “ultrafine” (PM < 100 nm, PM0.1), “fine” (PM < 2.5 μm, PM2.5), and “coarse” (PM < 10 μm, PM10) [4]. Once inhaled, fine and ultrafine PM are capable of crossing into the bloodstream where they are taken up by cells leading to oxidative stress and mitochondrial damage [5]. Ultrafine PM in particular is considered the most toxic form of PM, and may even be able to penetrate the brain directly through the olfactory nerve [6].

It has long been known that exposure to polluted air affects respiratory health [7], but could the effects be more widespread to extend even to the brain? Many epidemiological studies have focused on vehicle exhaust as a source of PM, measuring how urban vs. rural or distance to roadway affects

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brain function and cognition. Overall, these studies reveal an alarming trend: exposure to PM at any stage of life negatively impacts cognition. In childhood, exposure to PM negatively affects performance in multiple intelligence subscales [8] and is associated with poor motor coordination and response time [9]. Similarly, elderly residents living in highly polluted areas perform poorly in cognitive tests [10] and it is estimated that every 10 μg/m³ increase in exposure to black carbon is equivalent to 2 years cognitive decline by ageing [11]. Moreover, several studies support the notion that these cognitive effects translate into increased risk of dementia. For example, in a Canadian cohort of 2.2 million residents aged over 55 years, the incidence of dementia was significantly higher for those living within 50 m of a major roadway [12], and exposure to various pollutants including ozone [13] and nitrogen oxide [14] is associated with an increased risk of dementia. The exact risk seems to be modified by other environmental factors [15] along with genetic predisposition, since possessing copies of the APOE gene variant linked to dementia appears to exacerbate the effect of pollution [16].

These epidemiological studies are backed up by human pathological studies and animal studies revealing that PM exposure affects molecular pathways linked to AD. Individuals living in highly polluted areas accumulate higher amounts of Aβ42 [17], a toxic form of β-amyloid, and show hyperphosphorylated tau pre-tangles in the olfactory bulb and hippocampus [18]. Likewise, in a mouse model of AD, exposure to ultrafine PM led to an increase in the expected amount of Aβ plaques and reduced neuron density in the hippocampus [16]. These findings suggest that PM exposure can affect amyloid processing in the brain. The mechanism by which this occurs is likely to involve oxidative stress and an inflammatory response gone haywire, characterized by the massive release of pro-inflammatory cytokines from glial cells, generation of reactive oxygen species and increase in cellular antioxidant proteins [19] [Fig. 1].

Still, many questions remain unanswered regarding the link between PM and AD. The reliance of epidemiological studies on proxy measures of exposure like distance to roadway means that it is difficult to determine whether any specific air pollutants increase AD risk. If this were indeed the case, it would be arguably easier to enact policy to limit the release of these compounds into the environment. There is also a need for epidemiological studies over longer time courses, since the time window for the development of AD is still unknown. Despite these unknowns, evidence is mounting that make an “airtight” case against pollution in AD, with a call to clean up our cities and urban environments of the future.

### Spotlight on original articles

**Tooth loss in periodontitis: the role of Fusobacterium and inflammation**

We all know the importance of a daily routine of brushing and flossing, or we may otherwise suffer the consequences of gingivitis, an inflammation of the gums that is caused by the build-up of bacteria. If left untreated, gingivitis can lead to periodontitis, a serious gum infection that damages soft tissue...
and destroys the bone supporting teeth. In this issue of the Biomedical Journal, Johnson et al. [20] investigate the in vivo responses to infection with one of the leading perpetrators of periodontitis, Fusobacterium nucleatum.

The epithelium surrounding our teeth is a haven for anaerobic bacteria like F. nucleatum, which can grow in biofilms to form dental plaque [21]. To defend against this colonization, the oral epithelium secretes antimicrobial peptides, which both penetrate the bacterial membrane and act as chemoattractants to recruit immune cells to the site of infection [22]. Gingival epithelial cells (GECs) themselves also alerted to the infection when bacterial molecules bind to host pathogen recognition receptors on their surface. As a result, GECs activate NF-κB and express cytokines and chemokines to call in for reinforcement innate and adaptive immune cells [23]. Some oral bacteria, like Porphyromonas gingivalis, are able to evade these host responses by negatively regulating the production of pro-inflammatory cytokines in GECs [24]. F. nucleatum on the other hand, triggers inflammatory responses [25], but infection with this bacteria in isolation had not been studied before in mice until now.

To investigate responses to F. nucleatum, Johnson et al. [20] infected BALB/c mice and collected maxillas at several time points after infection. Consistent with previous findings, F. nucleatum infection caused a strong pro-inflammatory response characterized by the upregulation of several cytokines, including IL-1β and IFN-γ. This response was accompanied by the infiltration of immune cells, notably macrophages, into the maxilla. Although the goal of inflammation is to resolve the infection, pro-inflammatory cytokines like IL-1β and IFN-γ can activate osteoclasts, which break down bone tissue [26]. Using immunohistochemistry, Johnson et al. detected bone resorption pits in the alveolar bone seven days after infection, which were accompanied by a striking loss in bone elasticity as measured with an atomic force microscope, as early as one day after infection.

Overall, these findings reveal that the oral cavity launches a strong pro-inflammatory attack when challenged with F. nucleatum. Unlike many other pathogenic oral bacteria, F. nucleatum does little to hide from these host defenses and as such, may inadvertently promote bone resorption and tooth loss through macrophage recruitment and osteoclast activation, very early during the infection process.

Also in this issue

Review

Autophagy inhibition drives liver cancer
In this short review, Chen et al. [27] discuss how the inhibition of autophagy drives the progression of liver cancer and describe the molecular mechanisms in the tumor microenvironment underpinning this effect.

Original articles

Nrf2 protects against mitochondrial dysfunction in Parkinson’s disease
It has been known for many years that defects in mitochondria play a pivotal role in Parkinson’s disease [28]. Here, Fu et al. [29] examine the relationship between mitochondrial dysfunction and the accumulation of α-synuclein (SNCA) into Lewy bodies, the pathological hallmark of Parkinson’s disease. Using a SNCA mutant that is prone to aggregation, they find that the accumulation of SNCA is accompanied by an increase in reactive oxygen species and decrease in both mitochondrial fitness and mitochondrial DNA copy number. Interestingly, these effects could be reserved by activating nuclear factor/erythroid-derived 2)-like 2 (Nrf2), a transcription factor that regulates many anti-oxidant proteins [30].

Antimicrobial activity discovered in medicinal plant
Recent findings of an entirely new class of antibiotics in soil [31] suggest that many more antimicrobials are awaiting discovery in the environment. Here, da Silva et al. [32] test the oral toxicity and antimicrobial activity of the South American medicinal plant Baccharis trimera. Not only was the plant well tolerated in rats, it also inhibited the growth of Gram positive and Gram negative bacteria and limited biofilm production. These promising findings suggest that the hunt for new antimicrobials may prove fruitful by taking a “leaf” out of the book of nature.

Risk factors for hip fracture among elderly Taiwanese women
Taiwan has one of the highest incidences of hip fracture in the world [33], in particular among postmenopausal women. In this case-control study, Chen et al. [34] investigate the factors that put elderly Taiwanese women at risk of hip fracture and find that education level and total hip bone mineral density were the strongest predictors of first incident hip fracture.

Correspondence

Kaposi sarcoma-like lesions found in other skin conditions
In Kaposi’s sarcoma, patches of abnormal tissue growing under the skin that appear as multi-colored areas or the so-called “rainbow pattern” are considered a hallmark of the disease. However, in this correspondence, Kelati and Mernissi [35] report finding these Kaposi-like lesions in various dermatological conditions, suggesting that they are more widespread than previously thought.

Conflicts of interest

The author declares no conflicts of interest.

References

[1] https://www.alzinfo.org/articles/alzheimers-cases-triple-2050/. [Accessed 25 June 2018].
[2] Kilian J, Kitazawa M. The emerging risk of exposure to air pollution on cognitive decline and Alzheimer’s disease – evidence from epidemiological and animal studies. Biomed J 2018;41:141–62.
[3] World Health Organization. Air pollution, http://www.who.int/airpollution/en/. [Accessed 25 June 2018].
particulate matter activates early markers of oxidative stress, inflammation and unfolded protein response in rat striatum. Toxicol Lett 2013;222:146–54.

[20] Johnson L, Almeida-da-Silva CL, Takiya CM, Filiguolo V, Rocha GM, Weissmüller G, et al. Oral infection of mice with Fusobacterium nucleatum results in macrophage recruitment to the dental pulp and bone resorption. Biomed J 2018;41:184–95.

[21] Hassan A, Palmer RM. A clinical guide to periodontology: pathology of periodontal disease. Br Dent J 2014;216:457–61.

[22] Diamond G, Ryan L. Beta-defensins: what are they really doing in the oral cavity? Oral Dis 2011;17:628–35.

[23] Laube DM, Dongari-Bagtzoglou A, Kashleva H, Eskdale J, Gallagher G, Diamond G. Differential regulation of innate immune response genes in gingival epithelial cells stimulated with Aggregatibacter actinomycetemcomitans. J Periodontal Res 2008;43:116–23.

[24] Johnson L, Atanasova KR, Bui PQ, Lee J, Hung SC, Yilmaz O, et al. Porphyrmonas gingivalis attenuates ATP-mediated inflammasome activation and HMGB1 release through expression of a nucleoside-diphosphate kinase. Microbiol Infect 2015;17:369–77.

[25] Bui PQ, Johnson L, Roberts J, Hung SC, Lee J, Atanasova KR, et al. Fusobacterium nucleatum infection of gingival epithelial cells leads to NLPR3 inflammasome-dependent secretion of IL-1beta and the danger signals ASC and HMGB1. Cell Microbiol 2015;18:970–81.

[26] Zupan J, Jeras M, Marc J. Osteoimmunology and the influence of pro-inflammatory cytokines on osteoclasts. Biochem Med 2013;23:43–63.

[27] Chen KD, Lin CC, Tsai MC, Huang KT, Chiw KW. Tumor microenvironment mediated by suppression of autophagic flux drives liver cancer malignancy. Biomed J 2018;41:163–8.

[28] Winkler K, Haass C. Mitochondrial dysfunction in Parkinson’s disease. Biochim Biophys Acta 2010;1802:29–44.

[29] Fu MH, Wu CW, Lee YC, Hung CY, Chen IC, Wu KLH. Nrf2 activation attenuates the early suppression of mitochondrial respiration due to the alpha-synuclein overexpression. Biomed J 2018;41:169–83.

[30] Ishii T, Itok K, Takahashi S, Sato H, Yanagawa T, Katoh Y, et al. Transcription factor Nrf2 coordinately regulates a group of oxidative stress-inducible genes in macrophages. J Biol Chem 2000;275:16023–7.

[31] Hover BM, Kim SH, Katz M, Charlop-Powers Z, Owen JG, Ternei MA, et al. Culture-independent discovery of the antimicrobial activity of leaf tincture Baccharis trimera (Less). Biomed J 2018;41:194–201.

[32] Da Silva ARH, Lopes LQS, Cassanego GB, de Jesus PR, Figueredo KC, Santos RCV, et al. Acute toxicity and antimicrobial activity of leaf tincture Baccharis trimera (Less). Biomed J 2018;41:194–201.

[33] Canis JA, Oden A, Mcloskey EV, Johansson H, Wahl DA, Cooper C, IOF Working Group on Epidemiology and Quality of Life. A systematic review of hip fracture incidence and probability of fracture worldwide. Osteoporos Int 2012;23:2239–56.

[34] Chen FP, Fu TS, Lin YC, Fan CM. Risk factors and quality of life for the occurrence of hip fracture in postmenopausal women. Biomed J 2018;41:202–8.

[35] Kellat A, Mernissi FZ. The rainbow pattern in dermatoscopy: a zoom on nonkaposi sarcoma skin diseases. Biomed J 2018;41:209–10.