

AIMS Genetics, 4 (3): 166-191 DOI: 10.3934/genet.2017.3.166 Received: 13 March 2017 Accepted: 30 June 2017 Published: 11 August 2017

http://www.aimspress.com/journal/Genetics

Review

# The use of genotoxicity biomarkers in molecular epidemiology:

# applications in environmental, occupational and dietary studies

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**Abstract:** Molecular epidemiology is an approach increasingly used in the establishment of associations between exposure to hazardous substances and development of disease, including the possible modulation by genetic susceptibility factors. Environmental chemicals and contaminants from anthropogenic pollution of air, water and soil, but also originating specifically in occupational contexts, are potential sources of risk of development of disease. Also, diet presents an important role in this process, with some well characterized associations existing between nutrition and some types of cancer. Genotoxicity biomarkers allow the detection of early effects that result from the interaction between the individual and the environment; they are therefore important tools in cancer epidemiology and are extensively used in human biomonitoring studies. This work intends to give an overview of the potential for genotoxic effects assessment, specifically with the cytokinesis blocked micronucleus assay and comet assay in environmental and occupational scenarios, including diet. The plasticity of these techniques allows their inclusion in human biomonitoring studies, adding important information with the ultimate aim of disease prevention, in particular cancer, and so it is important that they be included as genotoxicity assays in molecular epidemiology.

**Keywords:** molecular epidemiology; biomarkers; genotoxicity; micronuclei; comet assay; environment; occupation; diet

# 1. Introduction

Genetic factors are clearly important in terms of influencing individual susceptibility to carcinogens; however, external factors represent the greatest opportunity for primary prevention. By 'external factors' we mean those related with environment-a broad scope, including all non-genetic factors such as diet, lifestyle and infectious agents. In a more specific approach, environmental factors include natural or man-made agents encountered by humans in their daily life, upon which they have no or limited personal control. The most important 'environmental' exposures, defined in this strict sense, include outdoor and indoor air pollution and soil and drinking water contamination [1]. In a more specific environmental niche are the occupational settings. People who work in certain jobs may have a higher risk of cancer due to exposure to some chemicals, radiation, or other aspects of their work (ergonomics, complex networks of safety risks, and many and varied psychosocial factors). Activities such as agriculture, painting, and industry are examples where workers can handle certain chemicals or be exposed to hazardous agents that can increase the risk of developing cancer [2]. Diet is also included in environment, particularly in lifestyle, and recognition of its importance has increased in recent decades, since it is a factor linked to some types of cancer [3,4]. The molecular epidemiology approach, measuring molecular or cellular biomarkers as indicators of disease risk or exposure to causative or preventive factors, has applications in studies of environmental and occupational exposure, disease etiology, nutrition, lifestyle and others [5], particularly in biomonitoring of populations.

This review aims to demonstrate the importance of genotoxicity biomarkers, such as those provided by cytokinesis blocked micronucleus assay and comet assay, as molecular epidemiology tools in human biomonitoring studies. With this approach, it is possible to detect, and therefore, prevent disease, specifically cancer in a wide variety of exposures—environmental, occupational and from diet.

#### 2. Molecular Epidemiology

Classical epidemiology has historically been the hallmark approach to demonstrate associations between exposure to hazardous substances and development of disease; however, inter-individual variation, i.e., genetic/individual susceptibility, did not have a place in this equation. The development of molecular biology and its use as a potential tool in epidemiological studies strengthened the identification of diseases associated with environmental exposures related to lifestyle, occupation, or ambient pollution. In molecular epidemiology, laboratory methods are employed to document the molecular basis and preclinical effects of environmental carcinogenesis [6-9].

Molecular epidemiology has the advantage of being directly relevant to human risk, unlike animal or other experimental models that require extrapolation to humans. Moreover, biomarker data on the distribution of procarcinogenic changes and susceptibility factors in the population can improve the estimation of cancer risk from a given exposure [10]. Increasingly, molecular epidemiology studies are incorporating panels of biomarkers relevant to exposure, preclinical effects and susceptibility, using blood and exfoliated cells, tissues and body fluids. These biomarkers are now being widely used in cross-sectional, retrospective, prospective and nested case-control epidemiologic studies, with the aim of improving our understanding of the causes of specific human cancers [5,11].

It is well established that maintaining the integrity of the genome is essential for normal cell function and any disruption in the process can lead to either cell death or cancer development [12], and so the majority of the available biomarkers used in molecular epidemiology studies are related to agents that cause DNA damage and are mutagenic [5,13]. Major gains in cancer prevention should stem from theoretically important strategies, namely regulations, public education programs, health surveillance, behavior modification, and chemoprevention programs and other interventions that adequately protect these groups from environmental carcinogens [10,14].

### 3. Biomarkers of Genotoxicity

Traditionally, biomarkers are defined as biomarkers of exposure, effect and individual susceptibility. For the purpose of this review, we will focus on biomarkers of effect. A biomarker can be any substance, structure or process that can be monitored in tissues or fluids and that predicts or influences health; or that assesses the incidence or biological behavior of a disease, but is not a measure of disease, disorder or health condition itself [15,16]. Ideally, biomarkers should be accessible (non-invasive), non-destructive, easy and cheap to measure [17,18].

One of the criteria for establishing associations between an exposure and disease is biological plausibility. In this context, biomarkers may contribute by illuminating some of the carcinogenic steps linked to a particular risk factor. This is possibly an undervalued area where biomarkers can make significant contributions to cancer epidemiology. If a particular chemical exposure from ambient air is associated with increased risk, the additional information that exposed individuals have higher levels of DNA damage would add support to the exposure-disease association [19].

Biomarkers of effect offer the opportunity to provide scientific confirmation of proposed exposure-disease pathways in human populations, since they can be elicited as a result of interaction of the biological system with the environment [20,21]. The increasing demand for information about health risks derived from exposure to complex mixtures calls for the identification of biomarkers to evaluate genotoxic effects associated with occupational and environmental exposure to chemicals, and other potential sources of damage. An important group of effect biomarkers are genotoxicity biomarkers, which have been developed in vitro (cells and cell lines), in vivo (animals) and ex vivo (cells from humans). Cytogenetic biomarkers are the most frequently used endpoints in human biomonitoring studies, and are extensively used to assess the impact of environmental, occupational and other factors in genetic (in)stability [20-22]. Among the wide range of cytogenetic biomarkers, micronuclei in lymphocytes provide a promising approach to assess health risks [23].

The most used biological matrices for studying genotoxic effects in human biomonitoring are blood lymphocytes and exfoliated cells, both being easy to sample. Lymphocytes circulate throughout the body, have a reasonably long life span, and can therefore be damaged in any specific target tissue by a toxic substance [24]. Exfoliated buccal cells have been effective in showing the genotoxic effects of lifestyle factors such as tobacco smoking, alcohol, medical treatments, such as radiotherapy as well occupational and environmental exposure, namely exposure to potentially mutagenic and/or carcinogenic chemicals, and in studies of chemoprevention of cancer (antioxidants) and evaluation of malignant transformation of preneoplastic lesions associated with oral squamous cell carcinoma [25-33].

## 3.1. Cytokinesis Blocked Micronucleus (CBMN) Assay

Living organisms may be exposed to mutagenic substances that cause cellular damage, which may be induced by chemical, physical or biological agents that affect DNA, chromosome replication and gene transcription, causing abnormalities that may lead to cancer and cell death [34].

The cytokinesis-blocked micronucleus (CBMN) assay is a comprehensive system for measuring DNA damage, cytostasis and cytotoxicity-DNA damage events scored specifically in once-divided binucleated cells. It is a method for assessing DNA damage caused by xenobiotics, allowing detection of effects caused by clastogenic agents (that provoke chromosome breakage) and aneugenic agents (abnormal chromosome segregation associated with loss) [34-38]. Other endpoints that can be measured are nucleoplasmic bridges (NPB), a biomarker of DNA misrepair and/or telomere end-fusions, and nuclear buds (NBUD), a biomarker of elimination of amplified DNA and/or DNA repair complexes [29,39].

The CBMN assay is regularly used as an in vitro test in genotoxicity testing (OECD 487) and it is the preferred method in human biomonitoring studies to detect cytogenetic effects after exposure to genotoxic agents. It is regarded as an indicator of mutagen sensitivity, a biological dosimeter of ionizing radiation exposure, a measure of DNA-repair capacity and genomic stability, and a predictor of cancer susceptibility/risk [40,41]. In summary, it is defined as a robust assay for genetic damage with applications in ecotoxicology, nutrition, radiation sensitivity testing both for cancer risk assessment and optimization of radiotherapy; as well as these applications in biomonitoring of human populations, it is important for testing new pharmaceuticals and other chemicals. There are expectations regarding the future development of an automated system that can reliably score the various endpoints which are possible with the CBMN assay [29].

## 3.2. Comet Assay

The comet assay (otherwise called single-cell gel electrophoresis—SCGE) is a simple, sensitive method for detecting DNA-strand breaks. DNA strand breaks can originate from the direct modification of DNA by chemical agents or their metabolites; from the processes of DNA excision repair, replication, and recombination; or from the process of apoptosis. Direct breakage of the DNA strands occurs when reactive oxidative species (ROS) interact with DNA. Alkali-labile sites generated by loss of bases in the DNA, are converted to strand breaks by alkaline treatment (pH above 13.1) and so are also detected with the comet assay [42].

This assay was adapted to measure oxidized purines and oxidized pyrimidines by incubation of the nucleoids (the DNA structures remaining after lysis of agarose-embedded cells) with bacterial DNA repair enzymes [43] including formamidopyrimidine DNA glycosylase (Fpg), which recognizes the oxidized purine 8-oxoguanine, one of the most studied molecules regarding oxidative damage [34,43].

Comet assay has become one of the standard methods for assessing DNA damage, with a wide range of applications, namely in genotoxicity testing, human biomonitoring and molecular epidemiology, ecogenotoxicology (monitoring environmental pollution by studying sentinel organisms), research on oxidative damage as a factor in disease, monitoring oxidative stress in animals or human subjects resulting from exercise, or diet, or exposure to environmental agents as well as fundamental research in DNA damage and repair [9,44-46].

The congruence of results between the comet assay and other endpoints such as micronuclei or sister chromatid exchanges (SCE), has been one of the principal reasons to increase the use of the comet assay as a biomarker for hazard assessment, particularly in monitoring the effects of occupational hazards [47-52].

## 4. Human Genome-Environment Interaction—Biomonitoring as a Tool

The relative contribution of genetics versus the environment to human illness has been debated for decades, as the so-called gene-environment interaction. The importance of environmental exposures has been supported by geographic differences in incidence of disease, by variation in incidence trends over time, and by studies of disease patterns in immigrant populations [53].

Understanding risks to human health in the light of human genome-environment interactions is one of the most compelling challenges in environmental public health. With the sequencing of the human genome, renewed interest in understanding the role of the environment as a cause of human disease has emerged. Genes are expressed in response to the environment [54] and there are two kinds of susceptibility genes: those that predispose to disease without exposure to environmental factors and those that increase risk only by interaction with environmental agents [53]. Information about environmental risk factors should point to genes that might modify the risk, and identification of susceptibility genes should help identify previously unrecognized environmental risk factors [53].

Human biomonitoring has tremendous utility providing an efficient and cost-effective means of measuring human exposure to hazardous substances establishing evidence that both exposure and uptake have been taking place [55,56]. This approach considers all routes of uptake and all sources which are relevant, making it an ideal instrument for risk assessment and risk management. It can identify new chemical exposures, trends and changes in exposure, establish distribution of exposure among the general population, identify vulnerable groups and populations with higher exposures, and identify environmental risks at specific contaminated sites with relatively low expenditure [56]. More attention should be given to monitoring populations which are known to be exposed to hazardous environmental contaminants and to providing reliable health risk evaluation, since that information is useful for supporting regulations on protection of the environment [57].

There are well-established national human biomonitoring survey programs worldwide, where a target population has been identified, questionnaires have been developed and sample collections have taken place. In Europe there are the German Environmental Survey (GerES, Germany), the Flemish Environment and Health Study (FLEHS, Belgium), the French National Survey on Nutrition and Health (ENNS, France), BIOAMBIENT.ES (Spain), Program for Biomonitoring the Italian Population Exposure (PROBE, Italy), Human Biomonitoring Project (CZ-HBM, Czech Republic). In America there are the Canada Health Measures Survey (NHANES), and the United States of America the National Health and Nutrition Examination Survey (NHANES), and in Asia, the Korea National Survey for Environmental Pollutans in the Human Body (KorSEP).

#### 5. Environmental Exposure

Nowadays people have to suffer the mutagenic and carcinogenic effects of many genotoxic agents in daily life and working environments due to changing lifestyles and innovations, for instance, chemical substances such as drugs, food additives, pesticides, and nanomaterials [58].

Anthropogenic pollution has become inherent to the modern environment. The global and rapid increase in technogenic stress in the biosphere raises the question about possible consequences for biota, including man, acknowledging that all forms of life are inter-connected and that human health is strongly linked to the ecosystem's health [59]. Environmental chemicals and contaminants are ubiquitous, occurring in water, air, food and soil. While some chemicals are short-lived in the environment and may elicit no harmful effects in humans, other chemicals bioaccumulate or persist for a long time in the environment or the human body due to frequent exposure, potentially leading to adverse health effects [60].

A more integrated approach is needed to deal with the fact that adverse biological effects induced by exposure to complex pollutant mixtures are not easily interpreted from a set of chemical analyses. The toxic effect of different interacting pollutants can be either additive, synergistic or antagonistic [61]. Molecular epidemiology studies on populations environmentally or occupationally exposed to high levels of complex mixtures of urban air pollutants have revealed genotoxic effects in terms of increased incidence of DNA damage [5,62]. Atmospheric pollutants, such as carbon monoxide, ozone, nitrogen oxides, sulfur dioxide, polycyclic aromatic hydrocarbons, and particulate matter are examples of chemical agents that may lead to DNA damage [34] and pose a serious threat to the health and the well-being of humans. According to their physicochemical properties, for instance, polycyclic aromatic hydrocarbons (PAHs) are released into the environment from both natural and anthropogenic sources, and are highly mobile in the environment, allowing them to distribute across air, soil, and water, becoming effectively ubiquitous [63,64]. It is also of great importance to assess the risk of future health effects from accidental or occupational radiation exposure to humans in order to be able to take appropriate measures to protect exposed individuals [65]. Multidisciplinary approaches combining chemical, ecotoxicological and ecological data have been undertaken to develop effective methods for assessing the quality of the environment, identifying the extent of genetic changes that occur when organisms are exposed to chronic, low-level, anthropogenic pollutants in selected species, such as protozoa, dicotyledonous plants [61], Scots pine [59], invertebrate and vertebrate native marine species [66], and others.

It is important to note that the genotoxicity biomarkers are applied in ecotoxicological studies; moreover, the application of early warning (sublethal) biomarkers in water-river quality monitoring programs is highly recommended since some of the pollutants are also relevant from a human health perspective—causing endocrine disruption, immune responses, or genotoxicity [61]. However this paper will cover just the effects in humans and human cells. Table 1 summarizes some studies regarding to environmental exposure, namely air pollutants [67-69], heavy metals [70,71], herbicides [72], mobile radiation [73], pesticides [74,75], pollution mixture [76], PAHs [77,78], and pyrethroids [79].

Risk factor/exposure	Studied population/number	Genotoxicity	Results			Refs.
Air pollutants (CO, NO <sub>2</sub> , SO <sub>2</sub> , benzene, $O_3$ , PM10 and PM2.5)	Children (Northen Italy)/N = 181/exfoliated buccal cells	MN assay	MN mean ±SD: MN mean frequ "reference" value	$0.29 \pm 0.13$ . ency of $0.29\%$ : 2–3-fold for children of this age.	l higher than that considered as a	[67]
Air pollutants: domestic heating (SO <sub>2</sub> and PM); traffic (NO <sub>x</sub> VOCs)	Children (suburban, urban- traffic sites in Turkey)/N = 1.841 summer; N = 1.497 winter/buccal epithelial cells	MN assay	Summer period Winter period <i>p</i> value No statistical dif children. Urban-traffic site Summer period Winter period <i>p</i> value MN frequencies than that of the w	MN (‰) (mean $\pm$ SD) 2.73 $\pm$ 1.98 1.87 $\pm$ 1.66 0.001 ferences between summer S MN (‰) (mean $\pm$ SD) 2.68 $\pm$ 1.99 1.64 $\pm$ 1.59 0.004 of urban-traffic children vinter ( $n < 0.05$ )	BEC with MN (‰) (mean $\pm$ SD)         2.28 $\pm$ 1.57         1.62 $\pm$ 1.33         0.003         and winter ( $p > 0.05$ ) in suburban         BEC with MN (‰) (mean $\pm$ SD)         2.68 $\pm$ 1.99         1.38 $\pm$ 1.15         0.005         significantly higher in the summer	[68]
Formaldehyde, nitrogen dioxide (NO2) in the air	Children 6–12 years old (living near chipboard- Viadana-Italy)/N = 413/oral mucosa cells	Comet assay MN assay	Children living exposure to form Comet assay Tail intensity (%) Tail lenght (µm) Tail moment	near (<2 km) the chipl aldehyde. ) 3.2 11 0.2	board industries — highest average ean 25 1.69 20	[69]

**Table 1.** Studies of human populations related environmental exposures.

Herease (2.15 pg/m ) was associated with a 10% relative increase (RK = 1.16; 95% CI: 1.06, 1.26) in NBUDs.Heavy Metals: arsenic, chromium, lead, manganese, in the same region)/N = 122/blood samplesComet assay (% DNAT) MN assayComet assay (% DNAT) MN assayControlsEnvironmentally exposed $p$ -value (70]Heavy metalsAdults (average age: 35.41) in 5 Bosnian regions with extensive mining, industrial activities/N = 104/blood samplesCBMN assay.Frequencies—range and mean $\pm$ SD[71]Heavy metalsMdults (average age: 35.41) in 5 Bosnian regions with extensive mining, industrial activities/N = 104/blood samplesCBMN assay.Frequencies—range and mean $\pm$ SD[71]Herbicide (alachlor)N = 1 male (age 43)/N = 1CBMN assayFrequencies—range and mean $\pm$ SD[71]Herbicide (alachlor)N = 1 male (age 43)/N = 1CBMN assayThe induction of MN-BN in isolated lymphocytes was not statistically [72]				Formaldehyde in 0.22%) higher c moment. Micronuclei assa MN: 0.12 NBUDs: 0.23	acrease (0.20 μg/n omet tail intensity y (%)	n <sup>3</sup> ) associated with a 0.1 y, 0.007 (95% CI: 0.001	3% (95% CI: 0.03, , 0.012) higher tail	
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$(age^{-1})$ was much higher than the other replicate leading to a higher but		30)/mononuclear isolated		signmean ( $p = 0.16$ ) annough one of the replicates at the highest concentration (20 µg mL <sup>-1</sup> ) was much higher than the other replicate, leading to a higher, but				
leukocytes not statistically significant difference.		leukocytes		not statistically s	ignificant differend	ce.	ang to a ingher, but	
Isolated blood lymphocytes				Isolated blood ly	mphocytes			

	Alachlor [µg/mL]	MN (per 1000)	
	0.0	6.0 ±0.0	
	2.5	6.0 ±2.1	
	5.0	5.5 ±0.7	
	10.0	6.8 ±0.4	
	20.0	10.3 ±4.6	
	Isolated human lymphocytes treated for	last 51 h of a 72 h culture period.	
	Isolated human lymphocytes		
	Alachlor [µg/mL]	MN in BN cells (per 1000)	
	0.0	3.8 ±0.4	
	2.5	4.8 ±3.2	
	5.0	4.5 ±0.7	
	10.0	4.8 ±1.8	
	20.0	Too few dividing cells	
	40.0	Too few dividing cells	
	4 h treatment with alachlor		
	Alachlor [µg/mL]	MN in BN cells (per 1000)	
	0.0	6.5 ±2.1	
	2.5	n.d.	
	5.0	n.d.	
	10.0	n.d.	
	20.0	4.5 ±0.7	
	40.0	13.5 ±3.5	
phone Male adults (age 20-30)/N MN assay	Group I mean $\pm$ SD (0.77 $\pm$ 0.815).		[73]
= 300 (150 high mobile	Group II mean $\pm$ SD (1.52 $\pm$ 1.176).		
users and 150 low mobile	Significant increase in the mean MN	count in group II in comparison to the	
users)/buccal epithelial	group I ( <i>p</i> -value < 0.0001).		
cells			

Mobile radiation

			In group II, the MN of statistically significant side $(0.90 \pm 0.3992)$ .	count in the side of m ly elevated $(1.52 \pm 1.1)$	obile phone use was found to 1 76) in comparison to the opposi	be te
			MN mean count was users $(2.08 \pm 1.291)$ in	found to be significate comparison to headpho	ntly increased in non-head photone users $(0.96 \pm 0.699)$ .	ne
Pesticides (complex	N = 239 agricultural	CBMN assay in		*	Mean ±SE	[74]
mixtures):	workers/N = $231$ unexposed	PBL	BNMN	Control	$12.25 \pm 0.60$	
carbamates,	controls/lymphocytes of	MN assay		Exposed	$11.40 \pm 0.49$	
organophosphates,	peripheral blood (PBL) and		MNL	Control	$13.82 \pm 0.69$	
pyrethroids	exfoliated cells of the oral			Exposed	$12.55 \pm 0.55$	
	mucosa		BCMN	Control	$1.06 \pm 0.10$	
				Exposed	$1.03 \pm 0.09$	
			MNBC	Control	$1.18 \pm 0.12$	
				Exposed	$1.12 \pm 0.10$	
Pesticides	Children (age $4-14$ )/N = 50	MN assay	MN mean per 1000 cel	lls		[75]
environmental exposure	pesticide spraying areas		Marcos Ju árez: 5.20 $\pm$	0.58		
(through inhalation):	$(C \circ rdoba)/N = 25$ children		R to Cuarto: 3.36 ±0.6.	3		
glyphosate, liquid	from the city of R b Cuarto		Genotoxicity is preser	nt in a group of child	ren in Marcos Juárez was high	er
formulations of cypermethrin, chlorpyrifos	(C órdoba), not exposed to pesticides/buccal mucosa cells		compared from to the I	R ó Cuarto.		
Pollution containing:	Adult residents (age 50-65)	MN assay		MN mean	% DNA mean	[76]
cadmium, lead, p,p'-	from 9 areas with different	Comet assay	Antwerp	7.30	1.69	_
DDE,	types of pollution/N =	(% DNA)	Antwerp port	6.65	1.23	
hexachlorobenzene,	1583/peripheral blood cells		Fruit area	6.00	1.35	
PCBs, dioxin-like			Olen	7.00	1.60	_
t,t'-muconic acid, 1-			Ghent	7.25	2.03	_
hydroxypyrene			Waste incinerators	8.60	2.24	

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Continued on next page

			Rural area Within an industrial area higher close to industrial industrial installations (p < Overall significant differe	7.00 DNA strand break leve installations than 5 kilor < 0.0001). ences between areas were	1.97 Is were almost three times metres upwind of the main still observed for oxidative	 -
			= 0.11).	and for DINA-straind break	s(p < 0.001) and for with (p	
Polycyclic aromatic	Children (age: 6–15)/5	Comet assay	,	Exposed children	Control group	[77]
hydrocarbons	groups of Tabasco-Mexico		Tail lenght	14.21–42.14	12.25	
(PAHs) in the air	5 groups/peripheral blood lymphocytes		Tail/head	0.97–2.83	0.63	•
PAHs and lead (Pb)	Children (age: 5–14), 2 most polluted cities-Katowice, Sosnowice/N = 74/peripheral blood lymphocytes	MN assay	MN mean: 4.44 Individual values reaching Positive significant corre 0.347, p < 0.05).	g 17 MN cells per 1000 bin lation was found betweer	ucleated cells. a PbB and MN levels ( $r =$	[78]
Pyrethroid insecticide	Males (age: 25–30)/N = 5/peripheral blood samples /human hepatoblastoma derived cell line HepG2	Alkaline comet assay with FPG	Dose dependent increas correlations between DNA 0.001 and tail lenght, tail 1 HepG2: tail DNA, $r = 0.84$	e of DNA damage in A damage in lymphocytes DNA, $r = 0.957$ , $p > 0.001$ 48, $p < 0.05$ and tail length	both cell types, positive (tail DNA, $r = 0.982$ , $p >$ , $r = 0.848$ , $p < 0.05$ .	[79]

## 6. Occupational Exposure

A wide range of chemicals that can act as environmental hazards, may also be exposure factors in specific occupational settings, and this is an extremely important consideration. For instance, besides the risks to the general public, atmospheric pollution can be considered an occupational health hazard to professional groups, such as traffic police or professional drivers working in urban areas [62], organic solvents [34, 80, 81], and others. Biomonitoring of exposure to toxic chemicals in the workplace is a fundamental tool to evaluate human health risks, supporting strategies to establish a safe work environment [82-85]. Table 2 summarizes some important occupational exposures, namely, antineoplastics [84], byproducts of petrol [85], formaldehyde [86], heavy metals [69,87,88], methyl bromide [89], organic solvents and smoke generated from biomass burning [34,80,81,90-92].

Occupational risk assessment may be defined as the qualitative and quantitative characterization of an occupational risk, i.e., the probability that an adverse health effect may result from human exposure to a toxic agent which is present in the occupational setting. It has three fundamental tools: environmental monitoring, health surveillance and biological monitoring. Risk assessment is meant to quantify the likelihood that a quantitatively defined occupational exposure of an individual (or group of individuals) to a chemical might result in adverse health effects [14,82].

National and international bodies set maximum allowable workplace concentrations for a wide range of substances. For instance, for airborne exposure to gases, vapors and particulates, recommended or mandatory occupational exposure limits (OELs) have been developed in many countries. The most widely used limits, called threshold limit values (TLVs), and are those issued in the United States of America by the American Conference of Governmental Industrial Hygienists (ACGIH). Specifically for airborne exposures, there are three other types of limit, namely the time-weighted average (TWA) exposure limit—the maximum average concentration of a chemical in air for a normal 8-hour working day and 40-hour week; the short-term exposure limit (STEL)—the maximum average concentration to which workers can be exposed for a short period (usually 15 minutes); and the ceiling value—the concentration that should not be exceeded at any time [83]. However, there is a need for revision of workplace limits to take also into account the levels of various agents that can cause allergies, for instance, in addition to occupational diseases. As new agents are identified they should be swiftly regulated.

Risk	Studied population/number	Genotoxicity	Results	Refs
factor/exposure	of samples/sample	biomarkers		
Antineoplastics	Occupationally exposed	CBMN assay	MN lymphocytes mean ±SE (range)	[84]
	nurses $N = 27/N = 111$ non-		Controls: 2.09 ±0.312 (0–15)	
	exposed subjects/peripheral	l	Exposed: $10.11 \pm 2.053 (1-58)$	
_	blood cells		The occupationally exposed group showed significantly higher MN mean ( $p$ value < 0.001 More Wikimen test)	
D		<u> </u>	0.001, Mann-whitney test).	<u></u>
Benzene	(GSA) N = $43$ /controls N	whole blood	DNA damage index, significant increase in the damage score in the GSA group compared to controls (Mann-Whitney test, $p < 0.001$ ).	[81]
	= 28/whole blood, buccal	MN assay in	3.8-fold higher in the GSA group compared to controls (Mann-Whitney test, $p < 0.001$ ).	
	exfoliated cells	buccal exfoliated		
		cells		
Benzene and	Gas station attendants (GSA	MN assay	Micronucleus assay	[34]
atmospheric	N = 43) taxi drivers (TD N	buccal cells	In the MN assay, no significant difference was observed among the groups ( $p > 0.05$ ).	-
pollutants	= 34)/persons without	Comet assay	Frequency of abnormal cells (MN/1000 cells):	_
	known occupational	blood	NE: 0.72	
	exposures (NE N =	lymphocytes	GSA: 2.70	_
	22)/buccal cells, blood		TD: 1.30	
			Comet assay	-
			Significant increase in DNA damage index (DI) in GSA and TD groups comparing to NE	-
			group ( <i>p</i> < 0.001).	
Byproducts of	Workers of car and battery	MN assay	MN mean (3000 cells per individual)	[85]
petrol and lead	repair garages N =		Exposed: 8.22	
	60/control group N = 80		Controls: 2.12	
	workers who were not		A significant difference ( $p < 0.001$ ) was found between the exposed and the control.	

**Table 2.** Studies of human populations related occupational exposures.

	exposed to byproducts of petrol and lead/exfoliated cells of buccal mucosa							
Formaldehyde	N = 46 workers occupationally exposed to	CBMN assay in peripheral blood		MN lymphocyte	in NPB es	NBUD	MN in buccal cells	[86]
	formaldehyde (20-61 years	lymphocytes		Mean	Mean	Mean	Mean	
	old)/N = 85 unexposed	MN assay in	Controls	0.81	0.18	0.07	0.16	
	individuals (20-53 years old)	buccal cells	Exposed	3.96	3.04	0.98	0.96	
			All genotoxicity comparison with	biomarkers controls (Ma	showed signi nn-Whitney test	ficant increases $(p < 0.002)$ .	in exposed workers in	
Heavy metals:	Adults (workers in the	Comet assay		Controls	Occup	pationaly exposed	<i>p</i> -value	[69]
arsenic, lead,	Panasqueira/N = 122/blood	(% DNA)		Mean	Mean			
chromium, ma-	samples	MN assay	% DNA	12.40	18.73		< 0.001	
nganese, moly-			MN (‰)	6.45	4.98		0.002	
bdenum, zinc			The occupational	ly exposed g	roup showed sig	nificantly higher	% DNA.	
Heavy metals	N = 90 male Pb recovery	Comet assay	Comet assay					[87]
lead (Pb)	unit workers/N = 90	in PBL			Come	t tail lengh (μm)		
	matched controls/peripheral	MN assay	Controls		8.15			
	blood lymphocytes, buccal	in buccal	Exposed		17.86			
	exfoliated cells	exfoliated	The results indic	ated that the	exposed worke	ers had a significa	ntly higher mean comet	
		cells and PBL	tail length than that of controls ( $p < 0.05$ ).					_
			Micronucleus ass	ay				
			MN frequency (%	bo) Bucc	cal cells	Lymphoc	ytes	
			Controls	2.97		3.17		
			Exposed	4.66		6.46		
			Increased MN fre	equency in ex	posed subjects t	than in controls (p	< 0.05).	
Heavy metals:	N = 204 male subjects (age:	Comet assay		Basal DN	A damage (µm)	MN frequency	(%)	[88]
nickel	18–50) in India/N = $102$	MN assay		Mean	Range	Mean	Range	

chromium	welders employed in		Control	8.94	4.14–17.10	0.32	0.00–0.80	
	welding plants, durations		Welders	23.05	17.24-35.62	1.30	0.12-2.89	
	of exposure (1–24		The results indicate	ated that the	e welders had a la	rger mean comet ta	il length than that of	
	years)/N = $102$ subjects-		the controls ( $p < 0$	0.001).				
	control group/blood		Welders showed	a significan	t increase in micr	onucleated cells co	mpared with controls	
	lymphocytes, buccal		(p < 0.001).					
	epithelial cells							
Methyl	N = 31 Methyl bromide-	Oropharyngeal	MN assay (MN/1	000 buccal	cells) mean:			[89]
bromide	exposed fumigation	MN assay	Workers: 2.00					_
	workers/n = $27$	(buccal cells)	Referents: 1.31					_
	referents/blood	lymphocyte	Two-sided <i>p</i> -valu	ue = 0.08.				_
	lymphocytes and	MN assay	Kinetochore-nega	ative micron	ucleated cells/100	0 lymphocytes mean	1:	_
	oropharyngeal cells	(blood	Workers: 10.48					_
		lymphocytes)	Referents: 10.41					_
			Kinetochore-posi	tive micron	cleated cells/1000	) lymphocytes mean		_
			Workers: 10.81					_
			Referents: 10.44					_
			No statistically s	ignificant d	ifferences were of	oserved between wo	orkers and referents for	
			mean kinetochore	e-negative ly	mphocyte MN.			
Organic	N = 45 footwear industry	Comet assay			Control	WBA	SBA	[90]
solvent	workers: solvent based	CBMN assay	Comet assay (blo	od)				_
mixtures:	adhesive (SBA N =		Damage index		$3.44 \pm 3.24$	$2.13 \pm 2.45$	$8.35 \pm 7.85$	_
acetone, 1-	29)/water solvent based		Damage frequence	cy (%)	$1.52 \pm 1.31$	$0.78 \pm 0.91$	$2.76 \pm 1.99$	
hexane,	adhesive (WSA N = $16$ )/N		Micronucleus test	t				
toluene,	= 25 controls/blood, buccal		MN (lymphocyte	s)	$5.20 \pm 2.33$	$3.88 \pm 1.93$	$4.90 \pm 2.34$	
methylethylket	cells		NPB (lymphocyte	es)	$3.00 \pm 1.97$	2.56 ±2.53	3.69 ±2.49	
one			MN (exfoliated b	uccal cells)	$0.62 \pm 0.73$	$0.69 \pm 0.87$	$1.15 \pm 1.45$	

			The Comet assay results shindex for the SBA ( $p < 0.00$	nowed that there was a signification of the W	ant increase in the mean damage BA group and control ( $p < 0.05$ ).	
			For the MN test in binuclea	ated lymphocytes and exfoliate	d buccal cells, the 3 groups were	
			not statistically different.			
Smoke	N = 23 sugar cane N	MN assay	Micronucleus assay (MN/1	000 cells)		[91]
generated by	workers/N = 30 control			MN mean (lymphocytes)	MN mean (buccal cells)	
biomass	group/blood lymphocytes,		Controls	1.27	9.70	_
burning	buccal exfoliated cells		Cutters	8.22	22.75	
			The MN frequencies in ly compared with the control	mphocytes were higher $(p < 0)$ group.	0.001) in the sugar cane workers	_
			A higher MN frequency in	exfoliated cells was obtained i	n the group of sugar cane cutters	
			compared with the controls	s ( <i>p</i> < 0.001).		
Toluene	N = 34 male industrial $Q$	Comet assay	Comet assay (DNA damage	e index):		[80]
	painters, occupationally M	MN assay	Controls: 39.4			-
	exposed to toluene/ $N = 27$		Painters: 60.4			-
	control group subjects with		Significant increase in DNA	A damage index between painter	s and controls ( $p < 0.001$ ).	_
	no history of occupational		Micronucleus assay (MN/1	000 cells)		-
	exposure/blood		Controls: 2.24			-
	lymphocytes, buccal cells		Painters: 2.74			-
			No significant difference be	etween painters and controls (p	> 0.05).	
	N = 34 women from $Q$	Comet assay		TM	% TDNA	[92]
	shoemaking plants ( $n = 16$		Controls	$5.37 \pm 2.48$	$18.18 \pm 6.26$	_
	plant $A + n = 18$ plant $B)/N =$		Workers plant A	5.85 ±2.43	$19.49 \pm 5.80$	-
	19controls/bloodmononuclear lymphocytes		Workers plant B	6.09 ±1.91	20.26 ±4.35	
Vehicle	N = 49 traffic police with $C$	CBMN assay		Mean $\pm$ S.D.	95% CI	[62]
exhaust	outdoor activities		Controls	$4.83 \pm 1.84$	4.20–5.46	_
	N = 36 indoor workers from		Traffic police	$7.06 \pm 2.87$	6.23–7.89	_
	university/lymphocytes		(p = 0.001, Wilcoxon test).			

# 7. Diet

Dietary habits are recognized to be an important modifiable environmental factor influencing cancer risk and tumor development, and other diseases. Although some studies have estimated that about 30–40% of all cancers are related to dietary habits, the actual percentage is highly dependent on the foods consumed and the specific type of cancer [18,93,94]. Epidemiological studies on the role of environmental exposure to carcinogens in diet have identified specific cancers whose incidence is known to vary considerably among countries [89]; substantial increases in the risk of certain cancers are observed in populations migrating from low- to high-risk areas, and this suggests that international differences in cancer incidence can be attributed primarily to environmental or lifestyle rather than genetic factors [93,95]. Diet can influence cancer development in several ways, namely by direct action of carcinogens in food that can damage DNA, by dietary components that can change enzyme activity, or by inadequate intake of molecules involved in antioxidant protection, DNA synthesis, repair or methylation that can influence mutation rate or changes in gene expression [96], and others. It is important to note, however, that the role of dietary components with potential cancer chemopreventive activity is not the subject of this review [3].

Another perspective of diet related to cancer risk is unintended contamination, which can result from compounds used in agriculture (e.g., pesticides and herbicides in plant-based foods, and growth hormones or antibiotics used in animal farming), or food processing (e.g., preservatives, smoking) and food packaging (e.g., bisphenol A or phthalates). The latter are not known to directly cause cancer, but they may influence cancer risk in other ways—for example, by acting as hormone-like substances in the body [97]. Is important to note that heavy metals, such as cadmium or mercury, may enter the food chain, such as in fish, or they may enter through contamination or their natural presence in soil or water.

Many substances are added to foods to prolong shelf and storage life and to enhance color, flavor, and texture. The possible role of food additives in cancer risk is an area of great public interest [97]. Briefly, food additive is a substance not normally consumed as food by itself and not normally used as a typical ingredient of the food, whether or not it has nutritive value [98].

The presence of such chemical contaminants or other unwanted substances in food and feed is often unavoidable as some of these substances are ubiquitous in the environment. However, the collection of dietary intake data along with chemical analysis of biological samples allows human biomonitoring programs to identify chemical exposures that might be associated with diet [60].

The European Food Safety Authority (EFSA)—commissioned project to review the state of the art of human biomonitoring for chemical substances and its application to human exposure assessment for food safety, facilitated the identification of vulnerable populations (e.g., by age, sex, socioeconomic status, etc.) as well as chemical exposure associated with food intake [60]. An important and specific context where the studies in diet have been raising more attention and concerns are maternal diet during pregnancy, this being the main source of essential nutrients that are needed for optimal fetal and child development. This applies no just to diet itself but also to prenatal exposure to several environmental pollutants which enter the mother's body as food contaminants, such as dioxins, PAHs and polychlorinated biphenyls [99,100].

Risk	Studied population/number	Genotoxicity	Results			Refs
factor/exposure	of samples/sample	biomarkers	Results			Reis.
Arsenic	Adults not significantly	MN assay		MN range	MN mean	[101]
Cooked rice	exposed to arsenic through	5	Whole cohort cooked rice arsenic (µg/kg)	0.50-4.98	2.12	
with > 200	drinking water		Lowest cooked rise arsenic group $\leq 100$		1.85	-
µg/kg	(west Bengal-India)/N =		Highest cooked rice arsenic group > 300		3.23	
	400/urothelial cells		Groups with mean cooked rice arsenic $> 200$	) μg have signific	cantly higher $(p < 0.05)$	
			induction of genetic damage compared to ea	ch of the groups	with mean cooked rice	
			arsenic $\leq 200 \ \mu g/kg$ .			
Beauvericin	N = 1 female (age:	Comet assay	BEA (0.5 $\mu M)$ and OTA (1 and 5 $\mu M)$ as w	ell as all toxin co	ombinations produced a	[102]
and ochratoxin	50)/human leukocytes		significant increase in tail moment compared	to control cells (	p < 0.05). BEA alone at	
А	PK15 cells		either concentration had a significantly low	wer DNA damag	e than BEA and OTA	
			combinations ( $p < 0.05$ ).			
Food additive	N = 2 adults (age: 24-	MN assay	Benzoic acid significantly increased micronuc	leus frequency (20	00 and 500 $\mu$ g/mL). This	[103]
benzoic acid	25)/human peripheral blood		increase was dose-dependent ( $r = 0.79$ ).			
	lymphocytes					
Monosodium	N = 3 adults (age: 23-	CBMN	MN assay:			[58]
glutamate	26)/peripheral blood samples	assay	Increase dose dependent ( $r = 0.96$ ).			
(MSG)		Comet assay	Comet assay:			
			% Tail intensity: $r = 0.60$ .			
			Mean tail lenght (mm): $r = 0.59$ .			
			Tail moment: $r = 0.71$ .			
			Increase dose dependent.			
Sodium sorbate	N = 2 adults (age: 24-	MN assay	SS increased SCEs/cell and MN frequency at	400 $\mu$ g/mL and 80	00 μg/mL concentrations	[104]
(SS)	25)/peripheral blood	Comet assay	at both 24 h and 48 h compared to negative con	ntrol.		

**Table 3.** Studies of human populations related dietary exposures.

			Comet assay	Average tail intensity (%)	
			Negative control ( $c = 0 \mu g/mL$ )	2.73	-
			SS (c = $400 \ \mu g/mL$ )	10.91	-
			SS (c = $8000 \mu g/mL$ )	5.97	_
			SS is genotoxic to the human periphera	l blood lymphocytes in vitro at the highest	
			concentrations.		
Synthetic food	N = 10 adults/blood samples.	MN assay	MN frequency was increased with increased brilliant blue	easing concentrations of sunset yellow and	[105]
Sunset vellow			Sunset vellow, significant increases in the	e MN rates were detected 30 mg/mL and 40	-
FCF and			mg/mL of the concentrations ( $p < 0.05$ ).	6	
brilliant blue			Brilliant blue, significant increases in the	MN rates were detected 30 mg/mL and 40	-
FCF			mg/mL of the concentrations ( $p < 0.05$ ).	C C	
Erythrosine	N = 1 adult/blood samples.	CBMN	Statistically significant increase in MN	N means induced by various food colors	[98]
(E127),		assay	(multivariate analysis, $p = 0.001$ and pairw	ise comparisons, $p < 0.05$ ).	
tartrazine			Control = 10		
(E102),			$100\mu g/mL = 12\pm 0.7$		
ponceau 4R			$200\mu g/mL = 12.8 \pm 0.8$		
(E124), sunset			$300\mu g/mL = 13.7 \pm 0.7$		
yellow (E110),					
brilliant blue					
FCF (E133),					
fast green					
(E143),					
carmoisine					
(E122), and					
indigo carmine					
(E132)					

Table 3 summarizes some important studies in diet field, namely the exposure to arsenic [101], mycotoxins as contaminants in food items [102], food additives [103,104], flavor enhancers [58], and synthetic food colorants [98,105].

For many other compounds for which the effects on cancer risk are not clear, there may be other good reasons to limit exposure. But at the levels that these are found in the food supply, lowering cancer risk is unlikely to be a major reason to justify this. There are moves to redefine maximum permissible limits for food colorants, instead of setting arbitrary limits for food additives in general; for instance in the case of colorants, each dye should have an individual limit based on well controlled genetic studies [98].

# 8. Conclusions

Human biomonitoring is a scientifically-developed approach for assessing human exposures to natural and synthetic compounds from the environment, occupation, and lifestyle, including diet [56]. It is the only available tool to integrate exposures from all sources and provide data for epidemiological studies of strengths of associations, dose response relations, etc.; however, it does not differentiate the exposure by source. Furthermore, human biomonitoring alone cannot provide information on how long a chemical has been in the body. Additional data collected from questionnaires, interviews and exposure assessment, combined with background knowledge, may provide valuable information regarding sources [21,60].

Although there has been growing recognition for the need to incorporate complex interactions between environmental exposures together with genetic factors, in order to fully understand cancer and diseases causation, since genetic instability is the startup point of carcinogenesis, there is growing recognition that environmental challenges not only interact with genes but may also modulate genetic effects and influence phenotypes [106]. An optimistic message is the fact that cancer development is not an inevitable consequence of the aging process *per se*, although there is a partly avoidable increased likelihood of the requisite number of mutations occurred, and the human species is not inevitably destined to suffer a high incidence of cancer. This awareness has lent greater urgency to the search for more powerful tools for primary prevention, for early warning systems to identify causal environmental agents and flag risks well before a disease condition develops [5].

In conclusion, the potential benefits of biomarkers and molecular epidemiology in illness prevention justify a major commitment to the further development of human biomonitoring programs, the only available tool that combines exposure assessment from different sources and relates their effects, together with individual susceptibility, to the risk of disease.

#### Acknowledgements

The authors would like to acknowledge Professor Susana Viegas and Professor Carla Viegas for their contribution in conceiving the idea of this review and the CA15132 hCOMET COST Action– European Cooperation in Science and Technology.

# **Conflict of interest**

The authors declare no conflict of interests.

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