Environmentally induced epigenetic toxicity: potential public health concerns

Emma L. Marczylo, Miriam N. Jacobs and Timothy W. Gant

Toxicology Department, CRCE, PHE, Chilton, Oxfordshire, UK

ABSTRACT
Throughout our lives, epigenetic processes shape our development and enable us to adapt to a constantly changing environment. Identifying and understanding environmentally induced epigenetic change(s) that may lead to adverse outcomes is vital for protecting public health. This review, therefore, examines the present understanding of epigenetic mechanisms involved in the mammalian life cycle, evaluates the current evidence for environmentally induced epigenetic toxicity in human cohorts and rodent models and highlights the research considerations and implications of this emerging knowledge for public health and regulatory toxicology. Many hundreds of studies have investigated such toxicity, yet relatively few have demonstrated a mechanistic association among specific environmental exposures, epigenetic changes and adverse health outcomes in human epidemiological cohorts and/or rodent models. While this small body of evidence is largely composed of exploratory in vivo high-dose range studies, it does set a precedent for the existence of environmentally induced epigenetic toxicity. Consequently, there is worldwide recognition of this phenomenon, and discussion on how to both guide further scientific research towards a greater mechanistic understanding of environmentally induced epigenetic toxicity in humans, and translate relevant research outcomes into appropriate regulatory policies for effective public health protection.

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Introduction
The belief that the sequence of our DNA (our genome) is the sole determinant of our life-long development and health has long been regarded as too simplistic. Darwin’s ‘On the Origin of the Species’ highlighted the role of the environment in the process of evolution, the natural selection of advantageous genetic mutations through environmental pressures (Darwin 1859). It has also been recognized that the pace of such change is sometimes too rapid to be accounted for by natural selection. First Lamarck, and then Waddington, suggested that alternative mechanisms, now termed epigenetics, must also contribute to phenotypic outcomes (Noble 2006). Epigenetics encompasses the mechanisms that regulate gene expression without altering the underlying DNA sequence, and the complete epigenetic status of a cell at any given time is termed the epigenome. Epigenetic programming is fundamental for normal mammalian development, and provides a more subtle mechanism by which the environment can rapidly alter gene expression within single or multiple generations. It is the complex interaction among our genome, epigenome and environment that shapes our development.
into unique individuals, and thus influences our health and potentially the health of our future offspring.

The induction of adverse DNA mutations by environmental factors that lead to disease, particularly cancer, is well established and recognized by regulatory bodies worldwide. As such, testing procedures and safety guidelines are in place to protect human health against the adverse effects of environmentally induced genetic mutations. There are, however, no current established regulatory procedures or guidelines in chemical safety programs for determining environmentally induced epigenetic toxicity. We, therefore, wanted to examine both the potential public health concerns and the resulting regulatory implications of such toxicity. For this purpose, we searched the current literature on epigenetic mechanisms involved in the mammalian life cycle and environmentally induced epigenetic toxicity in human epidemiological cohorts and rodent models in PubMed using the terms in Supplementary Table 1; defining environmentally induced epigenetic toxicity as environmental factors (chemicals, radiation and lifestyle factors (including alcohol, nutrition and smoking)) that induce adverse effects (growth retardation, fertility problems, hormonal changes, reproductive and/or other organ- and system-specific abnormalities (including immune disorders, obesity, diabetes and cancer)) and are associated with epigenetic mechanisms (histone modifications, DNA methylation, non-coding RNAs (ncRNAs) and other epigenetic machinery (DNA/RNA binding and modifying proteins)). Relevant papers were selected based on the inclusion criteria in Supplementary Table 1. These were then further evaluated and collated to provide a critical review that summarizes the main epigenetic mechanisms important during the mammalian life cycle, evaluates the current evidence for environmentally induced epigenetic toxicity in human cohorts and rodents models, and highlights the research considerations and implications of this emerging knowledge for public health and regulatory toxicology.

Environmentally induced epigenetic toxicity is an issue that has been highlighted as a current research focus within Public Health England’s (PHE’s) Center for Radiation, Chemical and Environmental Hazards (CRCE), the wider PHE organization as part of the early life priority ‘ensuring every child has the best start in life’, and the international research and governmental communities. By identifying and contributing to the knowledge gaps PHE, and similar organizations worldwide, can ensure that they stay fully aware of the current situation and are in the best position to give advice and put forward any necessary recommendations and policies to protect public health against such toxicity.

**Epigenetic mechanisms**

Histones are a family of highly conserved, small, basic (positively charged) proteins around which (negatively charged) DNA winds to form nucleosomes, the main structural units of chromatin (the protein-DNA complex) that enable DNA to be packaged into the nucleus (Kornberg & Lorch 1999). Both the tails and globular domains of histones can undergo multiple posttranslational modifications, including acetylation, methylation, phosphorylation, ADP ribosylation, ubiquitination and sumoylation, giving rise to the so-called histone code (Strahl & Allis 2000; Turner 2000; Jenuwein & Allis 2001; Bhaumik et al. 2007; Kouzarides 2007; Suganuma & Workman 2011). The functional groups are added or removed by a range of enzymes: acetylases, methyltransferases, kinases, ADP ribosyltransferases, ubiquitin ligases, sumo ligases; and deacteylation, demethylases, phosphatases, ADP ribosyl hydrolases, deubiquitinases, sumo deconjugating enzymes, respectively (Wang & Dasso 2009; Khare et al. 2012). These modifications alter the interaction between DNA and DNA binding proteins (such as transcription factors and RNA polymerases) so that it is the overall combined effect of the histone code that determines an active or repressive chromatin structure and thus regulates gene expression. In general, histone acetylation and demethylation are associated with active chromatin (euchromatin), while histone deacteylation and methylation are associated with repressive chromatin (heterochromatin).

DNA methylation involves the addition of a methyl group, primarily to the C5 of cytosine within cytosine–phosphate–guanine dinucleotides (CpGs) to produce 5-methylcytosine (5mC). Due to the high mutagenic potential of 5mC to spontaneously deaminate to thymine, the mammalian genome is largely depleted of CpGs (Bird 1980; Bird & Taggart 1980), with the exception of distinct regions known as CpG islands (CGIs) (Bird et al. 1985). These CpG rich CGIs are non-randomly distributed throughout the mammalian genome, being predominantly associated with regulatory elements such as promoters (Deaton & Bird 2011). DNA methylation also alters the interaction between DNA and DNA binding proteins to regulate gene expression. Methylation at CGIs is generally associated with transcriptional repression, although the local CpG density within a specific promoter can contribute to the actual transcriptional effect (Hackett & Surani 2013; Messerschmidt et al. 2014). DNA is methylated by a family of DNA methyltransferases (DNMTs) (Bestor & Ingram 1983; Li et al. 1992, 1996; Okano et al. 1998, 1999; Aapola et al. 2000; Gowher et al. 2005). DNMT1, the maintenance DNMT, preferentially methylates...
hemimethylated DNA and thus maintains DNA methylation following DNA replication. De novo methylation is established by the de novo DNMTs 3A and 3B, which preferentially methylate unmethylated DNA. The third member of the DNMT family, DNMT3L, does not possess any DNMT activity, but can help to recruit, and stimulate the activity of, DNMT3A and 3B. Much less is known about the mechanisms of DNA demethylation. It has long been suggested that 5mC can be removed by both passive (through lack of maintenance during replication) and active (enzymatic) mechanisms. Yet, specific DNA demethylase enzyme(s) in mammals remained elusive until the discovery of the 10–11 translocation (TET) enzyme family. This family, which can oxidize 5mC to 5-hydroxymethylcytosine (5hmC) and further to 5-formylcytosine (5fC) and 5-carboxycytosine (5caC) (Tahiliani et al. 2009; Ito et al. 2011), has fueled research into indirect active DNA demethylation pathways. Consequently, a range of mechanisms for the demethylation of DNA have been proposed (described in relation to mammalian development in Dean 2014; Messerschmidt et al. 2014; Messerschmidt, 2016).

The ncRNA superfamily encompasses a number of families broadly classified according to their length: long non-coding RNAs (lncRNAs) (>200 nt) and short non-coding RNAs (sncRNAs) (<200 nt), which include microRNAs (miRNAs, single stranded, ~19–25 nt), piwi-interacting RNAs (piRNAs, single stranded, ~24–30 nt) and endogenous short interfering RNAs (esiRNAs, double stranded, ~21–22 nt). The vast majority of the mammalian genome is composed of so-called non-coding DNA (ncDNA), with only 1–5% coding for proteins. It was widely believed that these 20–30 000 protein-coding genes were the sole mediators and executors of all cellular functions. The remaining 95–99% of the genome was regarded as ‘junk’ DNA. However, a functional role for ncDNA was inferred from the strong correlation between increasing ncDNA abundance and increasing organism complexity (Mattick 2007). Prokaryote genomes contain only 10% ncDNA, more complex fungi and animals >50%, rising to >98% in complex mammals (including mice and humans) (Carey 2011). Indeed, over the last decade, the previous gene-centric dogma, central to molecular biology, has been shown to be incorrect. NcDNA is transcribed into ncRNAs, which play major roles in regulating gene expression. While lncRNAs do this in a variety of ways, including chromosome remodeling, and transcriptional or post-transcriptional regulation (Galupa & Heard 2015; Kanduri 2016; Taylor et al. 2015), sncRNAs predominantly mediate gene expression at the post-transcriptional level (Cook & Blelloch 2013; Hale et al. 2014). In general, microRNAs repress gene expression by binding to mRNAs in a sequence-specific manner and either inducing their degradation or inhibiting their translation (Ivey & Srivastava 2015), whereas piRNAs and esiRNAs bind to complementary transposable elements (TEs) and induce their degradation (Watanabe et al. 2006, 2008). Thus, microRNAs are involved in fine tuning gene expression, whilst piRNAs and esiRNAs play a primary role in maintaining genome stability.

All these mechanisms play critical roles throughout normal mammalian development, particularly during early embryo and germ cell development (Cook & Blelloch 2013; Beaujean 2014; Dean 2014; Hale et al. 2014; Luk et al. 2014; Messerschmidt et al. 2014; Mukherjee et al. 2014; O’Doherty & McGettigan 2014; Grote & Herrmann 2015; Hogg & Western 2015; Marcho et al. 2015). As with any rapidly developing field, there is a continuous generation of new information that must be incorporated as appropriate, such as novel histone or DNA modifications and ncRNAs families implicated in the epigenetic regulation of the mammalian life cycle (Dean 2014; Hale et al. 2014; Messerschmidt et al. 2014; Sun et al. 2014; Garcia-Lopez et al. 2015; Sharma et al. 2016); and new functions for existing epigenetic mechanisms within mammalian development (Gou et al. 2014; Kuramochi-Miyagawa et al. 2008; Watanabe et al. 2008, 2011; Li et al. 2015, Pantano et al. 2015; Williams et al. 2015; Zhang et al. 2015).

**Role of epigenetics in the mammalian life cycle**

Normal mammalian development is a cyclical series of events that enable the continued production of multiple generations (described in Gilbert 2014). A new generation begins with the fusion of the male (spermatozoa) and female (oocyte) gametes (fertilization) to produce a single cell (zygote) containing one haploid female pronucleus and one haploid male pronucleus. All the different cell types that comprise the new individual develop from this zygote and so contain the same unique genome. To enable de-differentiation of the highly specialized mature male and female gametes and subsequent generation of multiple specialized cell types capable of performing a wide range of diverse functions, these genomes require differential modifications so that functionally relevant genes can be switched on/off. This is achieved through epigenetic re-programming and programming. Thus, throughout development, epigenetic processes regulate the expression and silencing of specific genes to drive the mammalian life cycle. Two key global epigenetic re-programming and programming events occur, one after fertilization in the early embryo in all cells, and the second in germ cells only during primordial germ cell (PGC) development (Figure 1).
The first re-programing event immediately following fertilization de-differentiates the mature parental gametes so that the resulting zygote becomes totipotent, capable of producing all the different cell types required to form a new individual. Re-establishment of new epigenetic marks then redirects and drives development of the zygote through embryogenesis and ultimately into a mature adult. Primordial germ cells (PGCs) undergo a second epigenetic re-programing and programing to ensure correct imprinting and development of mature germ cells according to the gender of the embryo, and enable the successful production of subsequent generations in a continuous cycle. Such epigenetic processes can be perturbed by environmental factors, potentially leading to an adverse phenotypic outcome.

![Figure 1. Epigenetics and the mammalian life cycle. Following fertilization epigenetic marks are removed to generate a totipotent zygote, capable of producing all the different cell types required to form a new individual. Re-establishment of new epigenetic marks then redirects and drives development of the zygote through embryogenesis and ultimately into a mature adult. Primordial germ cells (PGCs) undergo a second epigenetic re-programing and programing to ensure correct imprinting and development of mature germ cells according to the gender of the embryo, and enable the successful production of subsequent generations in a continuous cycle. Such epigenetic processes can be perturbed by environmental factors, potentially leading to an adverse phenotypic outcome.](image)
genome stability (Watanabe et al. 2006, 2008). Once transcription within the embryo genome is activated, however, there is rapid cell proliferation and differentiation, processes for which miRNA-mediated regulation is essential (Ivey & Srivastava 2015), hence increased expression of miRNAs.

It is worth noting that some genomic regions escape this first global 'reset', including imprinted control regions (ICRs) and the most active TEs. The continued transcriptional silencing of such regions is essential for early embryogenesis and genome stability, respectively. Normal embryo development requires a maternal and paternal contribution, generation of uniparental embryos by pronuclear transplantation results in developmental failure (Barton et al. 1984; McGrath & Solter 1984). This is because the male and female pronuclei are differentially imprinted; that is certain genes carry a parent-of-origin epigenetic mark so that they are only expressed from a single parental allele, ensuring the correct dosage of maternally and paternally derived gene products (reviewed in Bartolomei & Ferguson-Smith 2011). Imprinted genes are generally widely and highly expressed during prenatal development and predominantly down-regulated after birth. The placenta and brain, in particular, express many of the imprinted genes, in keeping with experimentally identified roles for imprinted genes in prenatal nutrient acquisition and growth regulation, neurodevelopment, and postnatal energy homeostasis and behavior. It is, therefore, vital for normal embryo development that these imprints are maintained during the first global epigenetic re-programming event. Similarly, the activation of TEs would be deleterious for the embryonic genome and so the continued silencing of such loci through early embryogenesis is essential for genome stability and subsequent normal development.

The second re-programing and programing event involves PGCs only. It is essential for the generation of subsequent germ cells (and thus the successful production of future generations) that parental imprints within PGCs are removed and new imprints are re-established according to the gender of the embryo. A female must reprogram the paternally imprinted genes on the chromosomes inherited from her father into maternally imprinted genes and vice versa. Thus, PGCs undergo a second round of global demethylation and changes in histone modification and non-coding RNA expression, this time including removal of the parent-specific imprints. Once again some of the most active TEs also escape this second phase of epigenetic re-programing, thereby maintaining genome stability. The mechanisms protecting specific genomic regions from epigenetic re-programing are not yet well-understood. However, some recent insights are presented and discussed in Hatanaka et al. (2015) and Leseva et al. (2016). Following sex determination, epigenetic programing via DNA methylation, histone modifications and non-coding RNAs redirect and drive differentiation of the PGCs along the distinct spermatogenic or oocyte lineage. Like the early zygote, mature gametes are transcriptionally quiescent. Thus, a similar sncRNA switch, this time back to increased genome protection makes functional sense. Indeed, PGCs, which represent an interim stage of the developing germ line that requires miRNA-mediated proliferation and differentiation with some pi/esiRNA-mediated genome protection, express greater levels of miRNAs and reduced levels of piRNAs and esiRNAs compared with the mature gametes and early zygote (Garcia-Lopez et al. 2015). Ultimately, the mature gametes come together to enable the cycle to begin again and so on.

Finally, it is important to note that histone modifications, DNA methylation and ncRNAs are not separate processes; they are closely linked throughout mammalian development. The different epigenetic pathways converge and interact to regulate gene expression at multiple levels (Figure 2). Thus, all the machinery involved in these pathways, including DNA and RNA binding and modifying proteins, are epigenetically important. This combination of interactive pathways helps to confer efficiency, redundancy, robustness and fidelity onto epigenetic re-programing during development, and thus the successful and continued production of multiple generations.

While such complex networks exist to maintain cellular homeostasis, they are open to environmental perturbation (Figure 1). Indeed, numerous environmental factors have been shown to induce epigenetic changes via a variety of mechanisms. These include both direct and indirect processes that ultimately lead to alterations in DNA methylation, histone modification and/or ncRNA expression. For example, certain metals and environmental chemicals can directly bind and alter the activity of DNMTs, TETs and histone modifiers (Chervona & Costa 2012; Hou et al. 2012; Wang & Wang 2013; Ruiz-Hernandez et al. 2015). Both chemically induced DNA adducts and those generated as a consequence of reactive oxygen species damage resulting from chemical exposure can interfere with the ability of epigenetic machinery to interact with DNA (Baccarelli & Bollati 2009; Ruiz-Hernandez et al. 2015). Reactive oxygen species are also proposed to stimulate the production of alpha-ketoglutarate, a co-substrate for TETs and one of the two main classes of histone demethylases (Chervona & Costa 2012; Ruiz-Hernandez et al. 2015). Poor nutrition (reduced folate), detoxification of arsenic and consumption of
glutathione during the detoxification of reactive oxygen species can all lead to reduced levels of the DNMT co-substrate S-adenosylmethionine (Hou et al. 2012; Ruiz-Hernandez et al. 2015). Similarly, histone acetylases require the co-substrate acetyl-CoA, and altered glycolytic production of acetyl-CoA can lead to changes in histone acetylation (Wellen et al. 2009; Moussaieff et al. 2015). Endocrine disrupting chemicals can bind to nuclear receptors, and either directly induce activity altering modifications to the epigenetic machinery (such as histone methyl transferases); or act as transcription factors, binding to hormone-responsive promoter elements and promoting expression of ncRNAs and/or epigenetic machinery (Kabir et al. 2015; Trevino et al. 2015). Additional epigenetic mechanisms are likely to be elucidated as research continues. It is when these environmentally induced perturbations exceed the capacity of the cell to maintain homeostasis that epigenetic toxicity occurs, potentially leading to altered phenotypes and adversely impacting public health.

Figure 2. Epigenetic regulation of gene expression. A range of epigenetic processes including DNA methylation, histone modifications and non-coding RNAs (ncRNAs), (such as long ncRNAs (lncRNAs), microRNAs (miRNAs), piwi-interacting RNAs (piRNAs) and endogenous short interfering RNAs (esiRNAs)) form an efficient and robust complex interactive network that regulates gene expression at the transcriptional and post-transcriptional levels. Inhibitory processes are shown in red and positive in green. Both histone modifications and DNA methylation changes can inhibit or promote transcription of mRNAs and ncRNAs. ncRNAs can then inhibit the production of protein from mRNAs either via mRNA degradation and/or reduced translation, and/or protect genome stability via the degradation of transposable elements (retrotransposons). Some ncRNAs can also promote changes in DNA methylation, histone modifications, and/or the expression of other ncRNAs. To complete the network, DNA/RNA binding and modifying proteins can both inhibit or promote gene expression whilst their own expression is under the control of the DNA modifications and ncRNAs they promote and/or interact with.
Environmental factors and epigenetic perturbation

Epigenetic processes are susceptible to perturbation throughout an individual’s life (Figure 1). Even in adulthood the same epigenetic processes drive cell proliferation, differentiation, function and adaptation. Environmentally induced epigenetic changes later in life can, therefore, also influence an individual’s health. Indeed, numerous adult onset diseases have been associated with abnormal epigenetic changes, including cancer, diabetes, and neurological, renal, cardiac and respiratory conditions (Hamm & Costa 2015); and epigenetic processes play a key role in initiating the onset of puberty, changes to which can also increase the risk of some of these adult onset diseases (Rzeczowska et al. 2014). However, certain stages in development and cell types can be thought of as particularly sensitive to epigenetic change due to the resulting severity of the outcome for the individual or the potential for affecting multiple generations.

Early life

Early embryo and germ line development can be affected by in utero exposures. Environmentally induced epigenetic changes during the vital first developmental re-programing and programing event could result in incorrect erasure and/or re-establishment of the embryo epigenome. This could lead to activation of TEs, and abnormal imprinting and gene expression. Such changes could have far-reaching consequences on embryo viability and development (including early germ line formation), and thus subsequent future health and fertility. It is also important to consider, however, early ex utero exposures. Early childhood and adolescence are also periods of significant growth and development, so it is easy to envisage how environmentally induced epigenetic changes during these stages could have detrimental effects on future health and fertility.

Gametogenesis and pre-conception

Spermatogenesis occurs continuously in adult males and even though females are born with their full complement of oocytes, oocyte maturation occurs in a continuous cycle throughout the reproductive life of adult females. Thus, environmental exposures to those oocytes and spermatozoa that go onto produce an embryo (pre-conception exposures) are also an important consideration. This represents an adulthood exposure to the parent, but a very early life exposure to the offspring. Environmentally induced epigenetic changes in the sperm or egg prior to conception could, therefore, affect embryo viability and development and the subsequent future health and fertility of the next generation. Indeed, murine spermatozoal miRNAs and/or esiRNAs have been shown to function in fertilization and early embryo development (Liu et al. 2012, 2016; Yuan et al. 2016), and to influence phenotypic outcomes in subsequent progeny (Rassoulzadegan et al. 2006; Wagner et al. 2008; Grandjean et al. 2009). Moreover, subsequent studies have further demonstrated that early life trauma or parental stress can induce altered miRNA expression in sperm, which in turn can transmit behavioral and metabolic abnormalities through to the next murine generation (Rodgers et al. 2013; Gapp et al. 2014; Rodgers et al. 2015). It is also worth noting that some of these RNA-mediated phenotypes have been shown to be dependent upon the remaining member of the DNMT family, DNMT2 (Kiani et al. 2013). Although DNMT2 has demonstrated some weak DNMT activity, it is predominantly associated with RNA, particularly tRNA, methylation, potentially contributing to the regulation of RNA stability and translation (Motorin et al. 2010).

Somatic versus germ cell change

If an epigenetic change occurs in a somatic cell or in a germ cell that is correctly ‘reset’ during germ line development in the subsequent offspring, then only one generation will be affected. However, if an epigenetic change becomes permanently established in the germ line, it will be transmitted across future generations, even in the absence of the original stimulus. As mentioned earlier, there are a number of loci within PGCs that escape the two global DNA demethylation re-programing events. Recent data has extended this list to include specific regions not associated with TEs that are preferentially located within CGIs, enhancers, promoters and gene bodies (Seisenberger et al. 2012; Hackett et al. 2013; Tang et al. 2015). These both provide evidence that DNA methylation can be stably inherited, and highlight specific loci protected from global epigenetic erasure that could be involved in transmitting epigenetic information across generations. In addition, not all of the spermatozoal histones are replaced with protamines during spermiogenesis, up to 15% of histones within human spermatozoa and 1% in mouse spermatozoa are retained (Miller et al. 2010; Carrell 2012). These retained histones are not randomly distributed throughout the male genome, they are particularly enriched at the promoters of developmental genes. Thus, spermatozoal chromatin may also transmit unique epigenetic information that helps to activate and regulate early embryo
development and contribute to epigenetic inheritance across multiple generations.

This so-called transgenerational epigenetic inheritance is a hotly debated topic. Much of the controversy is concerned with multigenerational versus transgenerational epigenetic toxicity (or epigenetic effect(s) versus epigenetic inheritance) (Figure 3). If an environmental exposure is truly transgenerational, then: (1) following an in utero exposure the resulting change penetrates to at least the F_3 generation (great-great grandchildren), where F_0 is the exposed mother and F_3 represents the first non-exposed generation; or (2) following an ex utero exposure in childhood/adulthood the resulting change penetrates to at least the F_2 generation (the great-grandchildren), where F_0 is the exposed generation and F_2 represents the first non-exposed generation. A transgenerational change thus represents epigenetic inheritance. Conversely, a multigenerational change represents an epigenetic effect that affects multiple generations, all of which were exposed either directly or indirectly to the original factor represent epigenetic effect(s) and thus multigenerational toxicity.

Figure 3. Multigenerational versus transgenerational epigenetic toxicity following in utero or ex utero environmental exposure. Penetration of adverse effect(s) to the first non-exposed generation (the F_3 generation following in utero exposure, or the F_2 generation following ex utero exposure) represents true epigenetic inheritance and thus transgenerational toxicity. Adverse effect(s) that affect multiple generations, all of which were exposed either directly or indirectly to the original factor represent epigenetic effect(s) and thus multigenerational toxicity.
Current evidence for putative environmentally induced epigenetic toxicity

Human cohorts

For many decades, there have been epidemiological studies linking the risks of adverse phenotypes to a wide range of environmental factors. For example, pre-conception, in utero or childhood undernutrition have been associated with an increased risk of developing various adult-onset diseases, including psychiatric disorders, type 2 diabetes and obesity, in numerous famine cohorts such as the Dutch Hunger Winter, the Chinese Great Leap Forward and the Överkalix cohorts (Lumey et al. 2011; Pembrey et al. 2014). Likewise, metabolic syndromes, hormonal perturbation, reduced sperm quality and increased cancer risks have been associated with chemical and radiation exposures in the survivors of human disasters, including the Seveso dioxin release, the Hiroshima and Nagasaki atomic bombs and the Chernobyl nuclear accident (Warner et al. 2011, 2013; Foley et al. 2015; Kamiya et al. 2015). In addition, large human cohorts that have recruited participants at birth for long-term follow up with comprehensive data and sample collection, such as the Avon Longitudinal Study of Parents and Children (ALSPAC) cohort (Boyd et al. 2013), provide a wealth of information to investigate epidemiological links between specific environmental exposures and public health outcomes. There have been over 1200 publications using data from ALSPAC (ALSPAC 2016), many of which have linked pre-conception and in utero environmental factors (including chemicals, pharmaceuticals, nutrition, smoking, anxiety and depression) with adverse outcomes (such as asthma and other respiratory disorders, increased weight/BMI-related issues and behavioral problems) (Golding 2010; Boyd et al. 2013). In particular, studies examining the parental and grandparental smoking status have demonstrated an interesting gender-specific association between paternal pre-conception or grandmaternal in utero exposure to cigarette smoke and increased weight-related issues in the subsequent sons and grandsons, but not daughters or granddaughters (Pembrey et al. 2014). Gender-specific effects were also observed in a number of the famine cohorts, with an increased risk of obesity occurring in female descendants only (Lumey et al. 2011; Pembrey et al. 2014).

The association of later life health with pre-conception, in utero or childhood environmental factors has led to the “Developmental Origins of Health and Disease” (DOHaD) hypothesis (Wadhwa et al. 2009). The DOHaD hypothesis evolved from Barker’s “thrifty phenotype” hypothesis originally proposed over 25 years ago (reviewed in Barker 2007), and was formed “to recognise….the concept that the early life environment has widespread consequences for later health” (Wadhwa et al. 2009). This concept of phenotypic plasticity, particularly as a consequence of different environments in early life, implicates epigenetic mechanisms and has fueled research into environmentally induced epigenetic toxicity. We therefore sought to review the existing literature (up to 7 March 2016) to evaluate the current evidence for such toxicity.

We identified 76 human cohort studies within PubMed that associated adverse phenotype(s) (such as birth outcomes, organ-specific abnormalities/disorders, cancer or mortality) with epigenetic change(s) (mainly DNA methylation to-date, but also histone modification, epigenetic machinery and miRNAs) and exposure to environmental factor(s) (including air pollution, chemicals, fibers, radiation or lifestyle factors (alcohol, nutrition or smoking)) at various stages of development (pre-conception, in utero, neonatal, childhood, adolescence, adulthood and lifetime). These studies can be ranked into categories that demonstrate (1) associations between environmental exposure(s), global epigenetic change(s) and adverse phenotype(s); (2) associations between environmental exposure(s), specific epigenetic changes(s) and adverse phenotypes(s) (including further subcategories, such as size and nature of cohort, type of sample analyzed, use of case controls, dose- and/or time-dependent changes, single changes within relevant biological pathways, multiple changes within the same biological pathway, correlation of changes with gene expression at mRNA and/or protein level, use of additional mechanistically relevant controls, robust statistical analysis, validation in independent cohorts); or (3) associations between environmental exposure(s), specific epigenetic changes(s) and adverse phenotypes(s) that were confirmed in a relevant in vivo and/or in vitro system. Studies falling into category 3 provide more comprehensive mechanistic evidence for environmentally induced epigenetic toxicity and are shown in Table 1. Specifically, these studies provide mechanistic associations between bisphenol A (BPA), formaldehyde, arsenic, nickel, or cigarette smoke, epigenetic changes and adverse phenotypes, and potential mechanisms of epigenetic toxicity in humans. The smoking evidence is particularly compelling as it has been established by multiple independent groups investigating different epigenetic mechanisms. In addition, studies within categories 1 and 2 may also enable the identification of markers and/or mechanisms of environmentally induced epigenetic toxicity through a weight of
Only studies that measured both adverse phenotypes and associated epigenetic changes in response to an environmental exposure were reviewed. All the studies shown here demonstrate associations among environmental exposure(s), specific epigenetic change(s) and adverse phenotype(s) that were confirmed in a relevant in vivo system. AD: Alzheimer's disease; COPD: chronic obstructive pulmonary disease; ECAT: elemental carbon attributable to traffic; NSCLC: non-small cell lung cancer; PBL: peripheral blood lymphocyte.

*Studies directly sampling the target tissue or validating the same change in the blood and target tissue of an appropriate in vivo model.

Evidence approach (e.g. the same association(s) reported in multiple independent cohorts or multiple association(s) that contribute to the same biological pathway), and are, therefore, included in Supplementary Table 2.

Furthermore, while human epidemiological cohort studies can demonstrate association, they do not prove causality. They are also complicated by the fact that the biological material under study, often blood cells, is not from the target organ/tissue. Interpretation of the findings with respect to mechanistic connections between environmental factors, epigenetic changes and adverse effects in the target organ/tissue can therefore be challenging. However, as stated above, all 10 of the studies shown in Table 1 further confirmed identified associations between environmental exposures, specific epigenetic changes and adverse phenotypes in a relevant in vivo and/or in vitro system; and seven of the 10 studies either directly sampled the target tissue or validated the same change in the blood and target tissue of an appropriate in vivo model. As such, these studies are of greater mechanistic relevance, and provide more comprehensive evidence for putative causal relationships.

**Rodent models**

The deleterious effects of exposures to environmental factors in human cohorts have stimulated investigations of environmentally induced toxicity in rodent models. In the late 1990s/early 2000s, these studies began to associate adverse phenotypes resulting from environmental exposures with epigenetic mechanisms (Issa et al. 1996; Tao et al. 1998, 1999; Choi et al. 1999; Bielawski et al. 2002; Govindarajan et al. 2002; Chen et al. 2004; Marwick et al. 2004; Pulling et al. 2004; Vuillemenot et al. 2004; Wu et al. 2004). Then in 2005, Anway et al. demonstrated that in utero exposure to the environmental chemical vinclozolin (a fungicide used in the wine industry) reduced sperm quantity, quality and thus male fertility with a >90% penetrance through multiple generations (to at least the F2 generation). The effect appeared to be transmitted through the male germ line, with altered DNA methylation patterns detected in the testes of the F1 generation and the sperm of the F2 and F3 generations. This was the first study to show phenotypic and epigenetic changes in generations not directly exposed to the original stimulus. As expected, this study generated some controversy, particularly when subsequent groups failed to reproduce the same results (Gray & Furr 2008; Schneider et al. 2008; Inawaka et al. 2009; Schneider et al. 2013). These disparities may have been due to the use of different routes of administration and variations in rat strains/stocks. Nevertheless, over the ensuing decade the hundreds of subsequent studies have provided more evidence for environmentally induced epigenetic toxicity.

We identified 147 rodent studies within PubMed (up to 7 March 2016) demonstrating both adverse phenotype(s) (such as reduced fertility, reproductive, developmental or other organ-specific abnormalities, or adult onset diseases (cancer, diabetes, obesity or immune disorders)) and epigenetic change(s) (histone...
modification, DNA methylation, epigenetic machinery or miRNAs) following exposure to environmental factor(s) (including chemicals, radiation or lifestyle factors (alcohol, nutrition, or smoking)) at various stages of development (pre-conception, in utero, neonatal, lactation, childhood, adolescence and adulthood). These studies can be ranked into categories that demonstrate (1) environmentally induced adverse phenotype(s) and global epigenetic change(s); (2) environmentally induced adverse phenotype(s) and specific epigenetic change(s) (including further subcategories, such as dose- and/or time-dependent changes, single changes within relevant biological pathways, multiple changes within the same biological pathway, correlation of changes with gene expression at mRNA and/or protein level, use of additional mechanistically relevant controls, robust statistical analysis); (3) environmentally induced adverse phenotype(s) and specific epigenetic change(s) that were reversed by an inhibitor/treatment, absent in a knock out/down model, and/or mechanistically linked in a relevant in vitro system; or (4) the same environmentally induced adverse phenotype(s) and specific epigenetic change(s) in both rodent model(s) and human cohort(s). Studies falling into categories 3 and/or 4 provide more comprehensive mechanistic evidence for environmentally induced epigenetic toxicity and are shown in Table 2. In total, 18 of the 40 studies shown in Table 2 validated loss of both epigenetic change(s) and adverse effect(s) following appropriate inhibition, treatment or knock out/down; thereby demonstrating a definitive causal link between exposure (largely high dose) to BPA, formaldehyde, cadmium, methylmercury, 2,3,7,8-tetra-chloro-dibenzo-p-dioxin (TCDD), urethane, vinyl carbamate, alcohol, caffeine, overnutrition, cigarette smoke, nicotine or 4-(methyl-nitrosamino)-1-(3-pyridyl)-1-butanone (NNK), epigenetic changes and adverse phenotypes. While the remaining studies do not necessarily demonstrate causality, they do provide additional mechanistic associations between exposure (again largely high dose) to BPA, formaldehyde, nickel, phthalates, TCDD, alcohol, undernutrition, 2-amino-1-methyl-6-phenyl-imidazo[4,5-b]pyridine (PhIP), peanut, cigarette smoke, nicotine or NKK, epigenetic changes and adverse phenotypes. The BPA, TCDD, alcohol and smoking-related evidence is particularly compelling as it has been established by multiple independent groups investigating different epigenetic mechanisms. In addition, studies within categories 1 and 2 may also enable identification of markers and/or mechanisms of environmentally induced epigenetic toxicity through a weight of evidence approach (e.g. the same epigenetic change(s) reported in multiple independent studies or multiple epigenetic changes that contribute to the same biological pathway), and, therefore, are included in Supplementary Table 3. While much of the work to date has focused on rodent models, there is also some evidence for environmentally induced epigenetic toxicity in other mammalian species, including sheep and monkeys (Wu et al. 2008; Begum et al. 2013; Lie et al. 2014a, 2014b).

Other exposures of environmental origin that have been extensively investigated in both human cohorts and rodent models include stress, recreational drugs, maternal immune activation/infection and parental age. While this large body of work also provides some comprehensive evidence for, and specific mechanisms of, epigenetic toxicity, it was considered outside the scope of this review. Likewise, pharmaceutical drug-induced epigenetic toxicity in human cohorts and rodent models has also been reported. However, such drugs are not environmental factors and so were not considered any further. Moreover, there were additional studies that have investigated environmentally induced epigenetic changes without simultaneously assessing any phenotypic/toxicological endpoints and/or focused on acute exposures (less than 24 h). While such studies are more difficult to interpret with respect to epigenetic toxicity (and so were not included in this review), they may still provide useful mechanistic data.

It is important to note that not all environmentally induced adverse phenotypes have been associated with epigenetic changes. Indeed, 18 human cohort and 20 rodent studies within PubMed reported a lack of epigenetically related toxicity (Supplementary Table 4). These studies do not necessarily rule out an epigenetic mechanism of toxicity per se, but they do exclude a link between the specific epigenetic, exposure and/or adverse endpoints/markers investigated.

The literature also contains many in vitro studies investigating the role of specific epigenetic mechanisms in adverse outcomes induced by environmental exposures. In addition, concurrent in vivo and in vitro studies have improved our knowledge of the mechanisms involved in the mammalian life cycle, including recent insights into early human development (Laird 2013; Slier et al. 2015; Surani 2015; Williams et al. 2015), and epigenetic inheritance in simpler model systems such as Caenorhabditis elegans, Drosophila melanogaster and Danio rerio (zebrafish) (Padilla et al. 2014; Sommer & Thummel 2014; Williams et al. 2014). Thus, although comparison and integration of these different human cohort, animal, and in vitro studies is complicated by the different doses, routes of administration, timings and lengths of exposures, the various strains, species, cell types or statistical methods used, and/or the specific adverse outcome(s) or marker(s) selected, the current
| Environmental exposure | Species | Route of administration | Exposure | Stage of development | Effect | Adverse phenotype(s) | Epigenetic change(s) | Reference |
|------------------------|---------|-------------------------|----------|----------------------|--------|----------------------|---------------------|-----------|
| **Chemicals**          |         |                         |          |                      |        |                      |                     |           |
| Bisphenol A (BPA)      | Mouse   | Oral (diet) (50 μg/50 mg/kg) | In utero | In utero | Reproductive, neurological, behavioral, metabolic and cardiac abnormalities | DNA methylation | Chao et al. (2012)* |
|                        | Rat     | Oral (gavage) (40–200 μg/ kg/d) | Pre-conception (maternal) | Childhood |                      | Epigenetic machinery | Jiang et al. (2015)* |
|                        |         | Injection (not specified) (20 or 40 μg/kg/d) | Lactation | Neonatal |                      |                     | Kundakovic et al. (2015) |
|                        |         |                         | Adulthood | Childhood Adulthood | F<sub>2</sub>-F<sub>3</sub> generations |                      | Susaño et al. (2015)* |
|                        |         |                         | F<sub>0</sub>-F<sub>2</sub> generations |          |                      |                      | Zhou et al. (2015)* |
| Formaldehyde           | Rat     | Age-related accumulation (0.07–0.5 mM) | In utero | In utero | Behavioral abnormalities | DNA methylation | Chao et al. (2012)* |
|                        |         | Injection (i) (1 μl of 0.5 mM) | Adulthood | Adulthood |                      |                      | Jiang et al. (2015)* |
| Metals:                |         |                         |          |                      |        |                      |                     |           |
| Cadmium (Cd)           | Mouse   | Injection (i) (20 mmol/kg/2d CdCl<sub>2</sub>) | Adolescence/early adulthood | Later adulthood | Hepatic abnormalities | DNA methylation | Wang et al. (2012)* |
|                        | Rat     | Oral (water) (0.5 mg/kg) | In utero | Adulthood | Behavioral abnormalities | Histone modification DNA methylation miRNAs | Onishchenko et al. (2008)* |
| Methylmercury          | Mouse   | Injection (i) (10 mg/animal Ni<sub>3</sub>S<sub>2</sub>) | Early adulthood | Late adulthood | Muscle tumors | Histone modification DNA methylation miRNAs | Zhang et al. (2013)* |
| Nickel (Ni)            | Rat     | Oral (gavage) (1 mg/g/wk) | Late childhood/adolescence + early adulthood | Late adulthood | Hepatic tumors | Histone modification DNA methylation miRNAs | Pascual et al. (2015)* |
| Phthalates:            |         |                         |          |                      |        |                      |                     |           |
| Dibutylphthalate (DBP) | Rat     | Oral (gavage) (500 mg/kg/d) | In utero | In utero to early adulthood | F<sub>1</sub> male offspring | Histone modification DNA methylation | Kilcoyne et al. (2014)* |
|                        |         |                         |          |                      |        |                      |                     |           |
| Di(2-ethylhexyl)-phthalate (DEHP) | Mouse | Injection (i) (20 mmol/kg/2d CdCl<sub>2</sub>) | Adolescence/early adulthood | Later adulthood | Hepatic abnormalities | DNA methylation | Martinez-Arguelles et al. (2009)* |
|                        | Rat     | Oral (gavage) (100–950 mg/kg/d) | In utero | In utero to early adulthood | F<sub>1</sub> offspring | Histone modification DNA methylation miRNAs | Papoutsis et al. (2015)* |
|                        |         | Injection (i) (50 μg/kg) | In utero | Neonatal | Gonado-tropins and gonadal steroid-genesis | Mammary and hepatic abnormalities | Takeda et al. (2012)* |
|                        |         | Oral (gavage) (1 μg/kg) | Early adulthood | Neonatal |                      |                      | Yoshioka et al. (2011)* |
| Urethane               | Mouse   | Injection (i) (1 mg/g/wk) | Late childhood/adolescence + early adulthood | Early adulthood | F<sub>0</sub>-F<sub>1</sub> generations | DNA methylation miRNAs | Pandey et al. (2016)* |
|                        |         |                         |          |                      |        |                      |                     |           |
| Vinyl carbamate        | Mouse   | Injection (i) (0.32 mg/mouse) | Adolescence/early adulthood | Adult hood | Pulmonary carcinogenesis | Histone modification DNA methylation miRNAs | Pandey et al. (2014a)* |
|                        |         |                         |          |                      |        |                      |                     |           |
| Lifestyle factors      |         |                         |          |                      |        |                      |                     |           |
| Alcohol (ethanol)      | Mouse   | Oral (water) (10 g/kg) | In utero | In utero | Neurological, behavioral, cardiac and mesenchymal stem cell abnormalities | Histone modification miRNAs | Ignacio et al. (2014)* |
|                        | Rat     | Oral (gavage) (5.25 g/kg/d or 5% (v/v)/d) | Adulthood | Neonatal |                      | miRNAs | Middleton et al. (2012)* |
|                        |         | Injection (i) (2.9 g/kg + 1.45 g/kg 2 h later) | Adulthood | Neonatal |                      | miRNAs | Leu et al. (2014)* |
|                        |         |                         |          |                      |        |                      |                     |           |
| Caffeine               | Mouse   | Injection (i) (20 mg/kg) | In utero | Adulthood | Cardiac abnormalities | DNA methylation | Buscariollo et al. (2014)* |
|                        |         |                         |          |                      |        |                      |                     |           |
| Nutrition:             |         |                         |          |                      |        |                      |                     |           |
| High fat diet          | Mouse   | Oral (diet) (60% fat) | Pre-conception (maternal) to lactation | Adult hood | Weight-, glucose- and lipid-related metabolic and food preference abnormalities | DNA methylation miRNAs | Baseloga-Escudero et al. (2015)* |
|                        | Rat     | Oral (diet) (50 mg/kg) | Early adulthood | Early adulthood | F<sub>0</sub>-F<sub>1</sub> generations | miRNAs | Cañí et al. (2015)* |
| Under-nutrition        | Mouse   | Injection (i) (20 mg/kg) | Early adulthood | Later adulthood | Hepatic abnormalities | miRNAs | Vucetic et al. (2016)* |
|                        |         |                         |          |                      |        |                      |                     |           |

(continued)
Table 2. Continued

| Environmental exposure | Species | Route of administration | Stage of development | Adverse phenotype(s) | Epigenetic change(s) | Reference |
|------------------------|---------|--------------------------|----------------------|----------------------|----------------------|-----------|
| Oral (diet) (folate and choline) | 2-amino-1-methyl-6-phenylimidazo[4,5-b]pyridine (PhIP) | Mouse (Human CYP1A2) | Oral (gavage) (200 mg/kg) | Adolescence/early adulthood | Later adulthood | Prostate abnormalities | DNA methylation | Epigenetic machinery | Li et al. (2012a)* |
| Smoke: Peanut | Mouse | Oral (gavage) (5 or 10 mg peanut + 10 or 20 μg CT/ mouse/d + 200 mg/mouse challenge) | Pre-conception (maternal) ± childhood | Childhood to adolescence | F1 offspring | Peanut-specific allergy and anaphylaxis | DNA methylation | Song et al. (2014)* |
| Smoking: Cigarette smoke | Mouse | Inhalation (exposure chamber) (1-2 h/d, 1 h/2 d or 8-10 cigarettes/d) | Combinations of: Neonatal, Adolescence, Early adulthood | Adolescence | Pulmonary inflammation and abnormalities | Histone modification | miRNAs | Halappanavar et al. (2013)*/y |
| Nicotine | Rat | Injection (sc) (1 mg/kg twice/d) | Combinations of: In utero, Neonatal | Neonatal | Growth retardation | Histone modification | DNA methylation | Epigenetic machinery | Li et al. (2012b)* |
| 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanone (NNK) | Mouse | Injection (ip) (2 mg/mouse or doses of 100 and 75 mg/kg a wk apart) | Late childhood/adolescence | Adolescence | Pulmonary tumors | DNA methylation | Epigenetic machinery | Lin et al. (2010)*/y |

Only studies assessing both adverse phenotypes and associated epigenetic changes in response to an environmental exposure were reviewed. CT: cholera toxin; ih: intrahippocampal; im: intramuscular; ip: intraperitoneal; sc: subcutaneous.

*Studies demonstrating environmentally-induced adverse phenotype(s) and specific epigenetic change(s) that were reversed by an inhibitor/treatment, absent in a knock out/down model, and/or mechanistically linked in a relevant in vitro system.

/y Studies validating loss of both epigenetic change(s) and adverse effect(s) following appropriate inhibition, treatment or knock out/down.

/z Studies demonstrating the same environmentally induced adverse phenotype(s) and specific epigenetic change(s) in both animal model(s) and human cohort(s).

* Studies in which environmentally induced adverse phenotype(s) and epigenetic change(s) were reported in separate publications but were performed within the same experimental animals.
evidence demonstrates a mechanistic association between specific environmental exposures, epigenetic changes and adverse health outcomes in human epidemiological cohorts and model systems. More studies that incorporate some form of inhibition/treatment/cessation into the experimental design are vital for establishing whether or not an exposure truly represents an epigenetic toxicant. Although potentially complex, the use of a systems biology approach, incorporating genome-wide analysis at multiple levels (histone code, methylome, non-coding RNA expression, transcriptome, proteome and metabolome) would provide comprehensive analysis of epigenetic changes and associated phenotypic outcomes. Subsequent identification of human relevant mechanisms of epigenetic toxicity can then be further validated as appropriate in smaller more focused studies. The challenge now is to fully identify and investigate the specific functional epigenetic mechanisms and biological pathways relevant to humans so that the risk(s) of environmentally induced epigenetic toxicity to public health can be properly assessed and addressed.

**Research considerations for public health**

There is no doubt that humans are exposed to a range of environmental factors during everyday life. However, to determine whether, or not, environmentally induced epigenetic toxicity is a real concern for public health there are a number of important considerations.

**Mechanisms and model systems**

The nature of the research means that studies in humans are mainly restricted to epidemiological cohorts. These cohorts often have limited availability of appropriate biosample at appropriate collection windows (Bakulski & Fallin 2014). *Ex vivo or in vitro* experiments on human embryonic tissue and cells are quite rightly restricted by ethics, and can only model a small part of the *in vivo* situation. Thus, the majority of whole system studies have been performed in animal models, predominantly rodents. While the rodent and human life cycles share many commonalities they do exhibit subtle differences, both with each other and other mammalian species (Kristensen et al. 2013; Laird 2013; Leitch et al. 2013; Surani 2015). Such differences are of critical importance when considering mechanistic relevance to humans.

Nevertheless, such work provides valuable data. The key is integrating and interpreting it correctly, in terms of differences/similarities at both the system (*in vitro versus in vivo*) and species (human versus non-human) levels. It is important to continue supplementing non-human models with human-derived data, incorporating new technologies to maximize data collection, and updating and re-evaluating current conclusions as novel information becomes available.

**Priority questions:** What are the detailed mechanism(s) that link a particular exposure to a specific adverse effect? Do these mechanism(s) function in humans? What are the potential human disease outcome(s)?

**Dose, route, metabolism, timing and mixtures of exposure**

Whether, or not, an environmental exposure elicits a response depends on a range of interacting variables, including dose, route of administration, metabolism and additional/pre-existing exposures (mixtures). Much of the animal studies to date on environmental chemicals have involved administration of high doses, many orders of magnitude above likely exposures in real human populations. Similarly, the first animal study to demonstrate that an *in utero* exposure could induce adverse phenotypes in subsequent generations administered the environmental chemical vinclozolin via the intraperitoneal route (Anway et al. 2005). Not only is it unlikely that a human would receive this, or any, environmental exposure via the same route of administration, but also the way in which an exposure is administered determines its overall effect. Indeed, oral administration of vinclozolin to pregnant females (the likely route of administration for humans given that vinclozolin is a fungicide used in the wine industry) failed to induce any adverse phenotypes in subsequent generations (Schneider et al. 2008). Thus, while it is important to fully characterize the hazard, further studies at doses and routes of administration similar to those found in humans are necessary for determining relevance to public health. It is worth noting, however, that establishing the actual level of human exposure to a particular factor and mimicking it in experimental models is not always easy.

Metabolism is another major factor that can determine the overall effect of an environmental exposure. Distinct species and model systems metabolize xenobiotics differently depending on their specific enzymatic profiles. Thus, metabolism of the parent environmental compound can vary greatly between models, resulting in the production of different types and levels of metabolites. It is, therefore, important to assess if the model system being used accurately replicates the *in vivo* human metabolic environment, and if not to conduct further studies using any additional appropriate metabolites.

The timing of an environmental exposure can also be a major contributing factor to the significance of the
Overall outcome. As discussed earlier, some life stages may be more susceptible than others to a particular exposure. There was the assumption that in utero exposure represents the most vulnerable period in an individual’s life with respect to their future development. However, it is now becoming clear that pre-conception and postnatal exposures can also be detrimental to future health across multiple generations.

Finally, humans are continuously exposed to a mixture of different environmental factors. Multiple exposures at multiple life stages could be additive and/or cumulative. While a particular low-level exposure may have no adverse effects, multiple low-level exposures over multiple stages of life may induce adverse phenotypes. Thus, once individual exposures have been characterized, it may also be relevant to assess the effect of (1) mixtures of environmental exposures, (2) multiple lifetime exposures of single environmental exposures, and (3) multiple lifetime exposures of mixtures of environmental exposures.

Priority questions: What dose(s) via what route(s) of administration are human populations actually exposed to, and what metabolite(s) are formed? Do low-level exposures induce epigenetic toxicity? Is any part of the life cycle particularly more sensitive to an exposure than others? Are multiple exposures additive and/or cumulative?

Human variability

It is known that different strains of laboratory animals can respond very differently to the same environmental exposure. Such differences are largely driven by diverse genetic backgrounds. Individual humans demonstrate substantial genetic differences and so an individual’s underlying genetics could have a huge effect on his/her susceptibility to a particular environmental exposure, depending on his/her metabolism, adaptability, gender, age, previous exposures, etc. Thus, not all humans will respond to an environmental exposure in the same way. Indeed, there are many examples in the literature of how single polymorphisms (SNPs) can affect the response of individuals and populations to different environmental exposures. These include not only mutations that alter the function of gene products but also those that lead to changes in DNA methylation or miRNA binding sites.

Priority question: How can we identify particularly susceptible individuals or populations?

Adaptive versus adverse epigenetic change

Epigenetic change offers a much more versatile, reversible and rapid way for adapting to changes in our environment compared with genetic mutation. This view of epigenetically mediated phenotypic plasticity falls into the realms of Lamarckian inheritance, the idea that an organism can pass on acquired characteristics to future offspring. The benefits of such inheritance are obvious and even Darwin himself proposed a hypothetical mechanism for the transmission of somatic tissue characteristics via the germ cells to subsequent generations, which he termed pangenesis (Darwin 1868). Indeed, not all environmentally induced epigenetic changes have proved detrimental. Hughes et al. (2009) reported that calorie restriction during adolescence and young adulthood as a result of the Dutch Hunger Winter was associated with altered DNA methylation and reduced risk of colorectal cancer in later life. Tyagi et al. (2015) described a protective effect of early (in utero through lactation) exposure to dietary omega-3 fatty acids on neuroplasticity in rats that involved specific changes in DNA methylation. Such benefits are not just limited to nutrition; advantageous chemically induced epigenetic changes have also been reported. Zeybel et al. (2012) demonstrated a beneficial heritable histone modification-mediated reprogramming of hepatic wound healing in male rats that reduced fibrotic scarring in the subsequent offspring of rats treated with the hepatotoxin carbon tetrachloride. In addition, it is possible to have adverse outcomes that are reversible. Indeed, Nishihara et al. (2013) provided epidemiological data to support the benefits of smoking cessation, with DNA methylation-related colon cancer risk reverting back to that of non-smokers within 10 years of cessation. Thus, the reversibility of environmentally induced changes is also critically important when determining the true potential adversity of an environmental exposure. Such findings, both positive and negative, have been incorporated into UK governmental advice and recommendations (Boyd et al. 2013).

Furthermore, epigenetic changes may themselves have no consequences on gene expression (as demonstrated in a recent in vitro study by Ramos et al. (2015). Thus, environmentally induced epigenetic changes may not always be mechanistically involved in adverse effects; instead, they may act as markers of past exposures (Bakulski & Fallin 2014), or represent adaptations involved in maintaining cellular homeostasis. Such changes may still prove useful for regulatory purposes and also do not rule out the possibility of later onset effects. For example, robust epigenetic changes induced by environmental exposures in the absence of a phenotypic consequence essentially become the ‘new normal’ epigenome, which could have greater susceptibility to disease development and/or future exposures (as demonstrated in two in vivo rat models (Greathouse...
Characterizing normal variation within the epigenome between non-disease states would provide valuable data to begin establishing limits of epigenetic change above which cellular homeostasis fails and epigenetic toxicity is induced. The considerable inter- and intra-human/mammalian variation combined with the vast number of potential epigenetic changes make this a very challenging task. However, defining the limits of normal variation for specific epigenetic changes of relevance would be a sensible and pragmatic starting point.

Priority questions: What is the ‘normal’ epigenome? What are the real short- and/or long-term consequence(s) of environmentally induced epigenetic change(s)?

**Implications for regulatory toxicology and intervention**

All the previous considerations are vital for effective regulatory toxicology and potential intervention. A robust, dose-dependent, causal relationship among a specific environmental exposure, an epigenetic change and an adverse public health outcome is required to classify a chemical as an epigenetic toxicant. While a similar robust, dose-dependent relationship is also crucial to facilitate the development of testing methods for hazard assessment, relevant environmental regulation and/or appropriate medical intervention, establishing causality is not necessarily essential. Epigenetic changes that act as markers of exposure and/or predictors of future toxicity, but do not in themselves directly induce an adverse effect, may also be useful in risk assessment and intervention. As yet there is no precedent for epigenetically mediated toxicity in chemical regulation. However, environmental exposures during critical windows of human development and subsequent non-genotoxic/epigenetic changes that may lead to later life adverse health outcomes (such as obesity, diabetes and cancer) across multiple generations continues to be of high concern for all the Organization for Economic Co-operation and Development (OECD) member countries, the International Agency on Research for Cancer (IARC) and the European Commission (EFSA 2013; Greally & Jacobs 2013; Herceg et al. 2013; OECD 2014). Epigenetics thus offers an opportunity for improved chemical regulation and public health protection. For example, epigenetic changes have the potential to provide (1) more sensitive/earlier end points of toxicity within existing regulatory test guidelines (TGs), (2) novel markers of exposure/predictors of future toxicity for the development of *in vitro* chemical screening assays and (3) novel end points of toxicities for which we are currently rarely able to conduct definitive regulatory testing due to animal welfare, cost and time considerations (such as non-genotoxic carcinogens, multigenerational toxicity and chronic low-dose exposures that impact health in later life). As such, epigenetics is high on the agenda of the OECD expert working group on an integrated approach to testing and assessment (IATA) for non-genotoxic carcinogens (NGTxC) and the OECD endocrine disrupters testing and assessment advisory group (EDTA AG) (Greally & Jacobs 2013; Jacobs et al. 2015). The interest of OECD member countries in epigenetics within the former expert group is particularly strong, where epigenetic mechanisms are playing a pivotal role in the development of tools for improved chemical safety assessment (Jacobs et al. 2015). Thus, the incorporation of epigenetics within chemical regulation could ultimately reduce animal usage and the time and cost of chemical research and development, while simultaneously improving the protection of public health worldwide.

The current body of evidence for environmentally induced epigenetic toxicity is predominantly a collection of human epidemiological data and exploratory *in vivo* high (often single) dose range studies, performed, not for regulatory purposes, but to investigate the theoretical potential and putative mechanisms of epigenetic toxicity in biological systems. Nevertheless, such studies have helped to identify putative markers and assays that may be useful for biomonitoring such exposures and toxicities, and testing the epigenetic toxicity potential of environmental exposures, respectively. Those that may be of particular interest for development into regulatory strategies/procedures are presented in Tables 1 and 2. A number of reporter mouse models, including the Agouti \( A^{\nu} \), \( \text{Cabp}^{\text{Ap}} \) and \( \text{Auxin}^{\text{Fu}} \), have also been used to investigate environmentally induced epigenetic changes *in vivo* exploratory research (Dolinoy et al. 2007; Waterland et al. 2006; Rosenfeld 2010). The \( A^{\nu} \) model in particular has been put forward as a potential epigenetic toxicological tool (Dolinoy 2008; Rosenfeld 2010; Faulk et al. 2013). However, the suitability of the \( A^{\nu} \) mouse as a true reporter model has since been questioned following both lack of reproducibility (Rosenfeld et al. 2013), and the suggestion that it may possess a “thrifty” genotype, naturally predisposing offspring to a metabolic syndrome when food is plentiful, thus confounding results following environmental exposures (Greally & Jacobs 2013). The \( \text{Auxin}^{\text{Fu}} \) mouse may, therefore, represent a more promising reporter model in the development of regulatory toxicological assays for potential environmentally induced epigenetic toxicity.
In addition, current OECD human health-related TGs have the potential for adaptation to incorporate adverse epigenetic endpoints (Greally & Jacobs 2013). Inclusion of such endpoints as non-compulsory measurements would (1) encourage better use of existing studies by adding a valuable non-genotoxic/epigenetic component that is not currently included, (2) provide an extensive sample resource and (3) initiate the collection of epigenetic data in a regulatory context. This would begin to address regulatory concerns and help to develop formal epigenetically relevant toxicity TGs. Such an approach would also respect the 3Rs. An overview of the relevant TGs that could be explored for adaptation (updated from Greally & Jacobs 2013), is provided in Table 3. These include not only the in vitro endocrine TGs and in vivo reproductive and developmental TGs (Greally & Jacobs 2013) but also a newly revised genotoxicity TG (TG 483) where spermatogonial samples could be further utilized for RNA analyzes (Linschooten et al. 2009; Marczylo et al. 2012; Metzler-Guillemain et al. 2015). The potential for such adaptations is already acknowledged at the OECD, but more applied development of key markers and subsequent validation work is needed to formally incorporate epigenetic endpoints into TGs. Examples of such proposed adaptations are provided within the OECD Endocrine Disruptor Conceptual Framework (Supplementary Table 5; OECD 2012). Similar discussions on the incorporation of epigenetic toxicity into the regulatory framework are also a current focus of industry supported workshops (van Ravenzwaay et al. 2014; ECETOC 2016; Miousse et al. 2015).

When considering temporal aspects to reproductive, developmental and endocrine disrupting effects, the TGs currently in use may not incorporate either the most relevant life stage(s), the most sensitive endpoints for epigenetic perturbation and/or sufficient duration of observation to detect later-onset effects. Thus, it is likely that the full spectrum of potential effects will not be identified. While some tests in the OECD Conceptual Framework do cover exposure during critical periods of development in utero, to date, they may not address effects that might be induced by exposure during neonatal or pubertal development, and/or emerge during later life stages.

For regulatory purposes, there are major limitations to simply extending lifetime rodent studies beyond the standard 2 years to examine later-onset epigenetic adverse outcomes, including financial, ethical and scientific considerations. In addition to the increased costs and animal welfare issues associated with housing and monitoring up to the natural death of the animals, there is also the confounding factor of increased background pathology with age. Infectious pathologies and increased incidences of certain tumor types, such as pituitary tumors or lymphomas/leukaemias, are known to occur with increasing age, and the high spontaneous incidences of leukemias/lymphomas and mammary gland tumors in laboratory animals from inbred colonies are known to vary considerably between studies (Innes et al. 1967; Greaves 2000). Thus, an overview of non-neoplastic, pre-neoplastic, neoplastic and benign pathologies within each model system is essential for the accurate interpretation of life-long rodent studies. Such overviews are, however, rarely included in the relevant literature associated with extended lifetime studies. Consequently, these studies are rarely considered acceptable to many regulatory authorities and are not specifically requested for ethical reasons, the numbers of animals needed to ensure appropriate statistical power would be both unworkable, and unethical. There are other more time and financially economical ways in

| Type of study               | TGs                           | Description                                                                                           |
|----------------------------|-------------------------------|-------------------------------------------------------------------------------------------------------|
| General exposure studies   | TG 451                        | Carcinogenicity studies                                                                               |
|                            | TG 452                        | Chronic toxicity studies                                                                             |
|                            | TG 453                        | Combined chronic toxicity/carcinogenicity studies                                                    |
| Post-mitotic cell studies  | TG 424                        | Neurotoxicity study in rodents                                                                       |
| Prenatal effects           | TG 414                        | Prenatal development toxicity study                                                                   |
|                            | TG 426                        | Developmental neurotoxicity study                                                                    |
| Reproductive effects       | TGs 415, 416                  | One- and two-generation reproduction toxicity                                                        |
|                            | TG 421                        | (Revised) Reproduction/developmental toxicity screening test                                           |
|                            | TG 422                        | (Revised) Combined repeated dose toxicity study with the reproduction/developmental toxicity screening test |
|                            | TG 443                        | Extended one-generation reproductive toxicity study                                                  |
| Genotoxicity tests         | TG 483                        | (Revised 2015) Mammalian spermatogonial chromosome aberration test                                    |
| Alternative models integrating multiple mechanisms of toxicity | TG 236                        | Zebrafish embryo epigenetic toxicity assay                                                            |
|                            |                               | Xenopus Embryo Thyroid Assay (XETA Assay) (in validation)                                             |

Updated from Greally and Jacobs (2013). All TGs cited are publically available from the OECD website (http://www.oecd-ilibrary.org/environment/oecd-guidelines-for-the-testing-of-chemicals-section-4-health-effects_20745788).
which lifetime, multigenerational and/or transgenerational toxicity could be assessed, for example using zebrafish models, both respecting the 3Rs and satisfying sufficient statistical power. Already widely used in the pharmaceutical industry for developmental toxicity screening, zebrafish models appear to be a promising alternative to mice. The main epigenetic pathways, including histone modifications, DNA methylation and snRNAs, are also conserved in zebrafish (Mishima 2012; Kamstra et al. 2015; Williams et al. 2014).

Furthermore, it is also important to consider the broader repercussions of regulatory toxicology. Of course, it is vital that any environmental toxicant is carefully regulated or even banned from use. However, the situation is often not as straightforward as this. For example, chemicals can have far-reaching benefits to society, from agriculture to medicine and food/product safety to the development of new technologies. Regulations are frequently a fine balance between maximizing the ‘modern life’ benefits of an environmental chemical while minimizing any potential adverse effects. This is where dose/level of exposure and relevance to humans is crucial, but considerations of alternative replacements are also important. Do we run the risk of banning a chemical that does not demonstrate any epigenetic toxicity at the levels found within the environment or human populations only to replace it with an alternative whose characteristics and potential effects we know even less about? Surely, it is better for both society and public health to fully assess the potential epigenetic effects of environmental exposures and introduce regulations and TGs based on relevant data and proven mechanisms of toxicity.

Conclusions

Despite the fact that the seminal work on environmentally induced epigenetic toxicity was published over a decade ago, this area of research largely remains in its infancy, striving to elucidate putative mechanisms of epigenetic toxicity and identify mechanistic associations in human cohorts. This might well be expected since the fundamental idea that environmental factors can induce phenotypic changes across multiple generations challenges the central dogma of genetic inheritance, moving into what was a controversial area of genetics, but is now better accepted and established, Lamarckism. While many hundreds of studies have investigated environmentally induced epigenetic toxicity, relatively few (47 to-date) have demonstrated a mechanistic association among specific environmental exposures, epigenetic changes and adverse phenotypes. Even fewer (18 of the 47 to-date) further established causality, all of which were studies in rodent models. Moreover, the majority of these 47 studies represent exploratory in vivo high (often single) dose range experiments, many orders of magnitude above likely human exposures. Nevertheless, they do set a precedent for the existence of environmentally induced epigenetic toxicity, and provide tools for linking environmental exposures to adverse health outcomes. Thus, although the current literature remains incomplete regarding specific mechanisms of epigenetically mediated environmentally induced toxicity in humans at doses relevant to human exposures, there is sufficient information to perform retrospective epigenetic analysis of existing regulatory studies and identify future research needs. For example, epigenetic mechanisms and markers identified within the 47 studies highlighted in this review could be further examined and potentially validated in existing regulatory studies. Improved human exposure data could be collected and used to facilitate the use of more relevant doses, routes of administration, metabolites and mixtures. Guidelines could be developed for improved molecular (in vivo, in vitro and epidemiological) and bioinformatic study designs that incorporate suitable new technologies (such as omics platforms) and appropriate inhibitors/treatments/cessations (for exposure/change/effect validation) to definitively identify detailed epigenetic mechanisms that link, or act as markers of, environmental exposures and adverse health outcomes. Collaboration between scientists from academia, industry, and governmental and regulatory bodies will promote further research within a regulatory context, and drive the development and implementation of epigenetically relevant integrated testing strategies or policies for the continued protection of public health.

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