Pembrey, M., Saffery, R., & Bygren, L. O. (2014). Human transgenerational responses to early-life experience: potential impact on development, health and biomedical research. *Journal of Medical Genetics, 51*(9), 563-572. https://doi.org/10.1136/jmedgenet-2014-102577
Human transgenerational responses to early-life experience: potential impact on development, health and biomedical research

Marcus Pembrey,1,2 Richard Saffery,3,4 Lars Olov Bygren,5,6 Network in Epigenetic Epidemiology

ABSTRACT

Mammalian experiments provide clear evidence of male line transgenerational effects on health and development from paternal or ancestral early-life exposures such as diet or stress. The few human observational studies to date suggest (male line) transgenerational effects exist that cannot easily be attributed to cultural and/or genetic inheritance. Here we summarise relevant studies, drawing attention to exposure sensitive periods in early life and sex differences in transmission and offspring outcomes. Thus, variation, or changes, in the parental/ancestral environment may influence phenotypic variation for better or worse in the next generation(s), and so contribute to common, non-communicable disease risk including sex differences. We argue that life-course epidemiology should be reframed to include exposures from previous generations, keeping an open mind as to the mechanisms that transmit this information to offspring. Finally, we discuss animal experiments, including the role of epigenetic inheritance and non-coding RNAs, in terms of what lessons can be learnt for designing and interpreting human studies. This review was developed initially as a position paper by the multidisciplinary Network in Epigenetic Epidemiology to encourage transgenerational research in human cohorts.

BACKGROUND

The challenge of common chronic disease

The focus of public health is increasingly on chronic disorders, such as diabetes, cardiovascular disease, mental health and cancer manifesting in adulthood. Unlike monogenic (Mendelian) disorders or infectious disease, where the known primary cause permits an aetiological classification and rational approach to management, common diseases are generally multifactorial in origin with diverse genetic and environmental determinants. This contributes to the highly heterogenous nature of such conditions, which remain an aetiological and management challenge. A multidisciplinary perspective, which includes genetic analysis, is recognised as the most appropriate approach to studying such conditions.1–3 Genetic associations are now routinely integrated with long-established epidemiological analysis of nutrition, social and other exposures through which health and disease have been traditionally explored.

Extending epidemiology beyond genetics and a single generation

For many years, parental and ancestral experience has been regarded as contributing only indirectly across generations. Clearly, such ‘cultural transmission’, which includes parental nurturing behaviour and social patterning of all kinds, does play an important role in shaping the phenotype/trait of progeny. This is an area of active research integrating information on social circumstances, skills, health and mortality, sometimes across three generations of the same lineage.4–5

In contrast, biological inheritance of phenotype from one generation to the next is usually regarded as resulting from transmission of genetic material (DNA) from both parents, plus ‘maternal effects’ either (i) carried within the egg cytoplasm (such as in mitochondria, proteins and RNA molecules), (ii) experienced in utero, primarily through the transplacental passage of nutrients, metabolic signals and toxins, or (iii) experienced early postnatally, via breast feeding and/or the exchange of microbiota. Additionally, the maternal genotype also contributes to the early nutritional environment of progeny through its influence on maternal metabolism during pregnancy. This is best exemplified in studies on insulin sensing and fetal growth.6

It is plausible that a maternal exposure in pregnancy could induce a ‘metabolic cascade’ to subsequent generations, whereby fetal programming could alter later adult metabolism, which, in turn, changes the physiology of the uterus receiving and programming the early embryo of the next generation and/or the transplacental metabolic signals to the fetus. Theoretically, this process could be genotype independent. The challenges of unravelling the complexities associated with maternal transmission of exposure effects are manifold due to the multiple levels of interdependence (figure 1).

Until recently, paternal exposures prior to conception (other than those known to induce DNA mutations) were assumed to have little or no effect on offspring development, primarily due to the lack of a plausible biological mechanism. So ingrained was this view that confirming a lack of paternal exposure effect has been promoted as a useful ‘control’ for social confounders in the study of maternal exposure effects.8,9 However, several robust animal experiments exist showing transgenerational phenotypic effects of exposures via the male line.10–16 Human observational studies (reviewed in17) also support this type of non-genetic transmission.

In light of the emerging evidence for both maternal and paternal transmission of non-genetic, non-cultural effects to progeny, consideration must be given to extend traditional epidemiological
methodology to include not only early-life paternal and maternal exposures, but also ancestral exposure data. The consequences of such transmission are potentially profound, particularly in relation to past assumptions underpinning much of epidemiological and genetic analyses. In transgenerational epidemiology, assuming cultural inheritance automatically when results make genetic inheritance implausible (and vice versa) is no longer acceptable.

Framing future transgenerational epidemiological research

Although some have adopted the terms intergenerational, multigenerational and transgenerational to mean specific routes and numbers of generations of transmission, we use ‘transgenerational’ in a general sense to describe a measurable outcome in one generation (offspring) with exposures in either the mother or the father (prior to conception), or their respective parents or ancestors. ‘Transgenerational’ was first used in this context in the 1980s for the apparent transmission of the trauma suffered by Holocaust survivors to their unexposed offspring. This term was also adopted in early speculation about imprinted genes mediating the transmission of exposure information to the next generation and is used in recent reviews of the subject in mammals. Importantly, the mechanism of transmission must be primarily non-genetic, although mounting evidence implicates genetic factors as moderators of such effects (see below). In prenatal exposure via the mother, the fetus can be assumed to be directly exposed; and here the outcomes of interest in a ‘transgenerational’ context would be in that child’s own offspring.

A priority for transgenerational epidemiology is to describe the range of measurable exposures and outcomes mediating transgenerational effects in humans. That is, what exposure at which life stage in parents, grandparents or distant ancestors is associated with a measurable phenotypic outcome in the offspring or subsequent generations? We suggest this observational research is best done in a setting where the effects of social patterning can be accounted for, thereby distinguishing cultural inheritance from a more direct biological transmission across generations. Defining exposure-sensitive periods associated with transgenerational effects will be highly informative in this regard.

Unlike much of the research in the field to date, it is important that future transgenerational epidemiology should not be restricted to the inheritance of acquired characteristics, or ‘phenotypic transmission’ to the next generation. The exposure is key, not the founder, phenotype. In fact, there need not be a measurable phenotype in the exposed (or intermediate) generation. Similarly, phenotypic outcomes should not only be

**Figure 1** Schematic pedigree diagram showing the main routes for biological transmission of the effects of exposure to the next generation(s). Left, female line; right, male line. The exposure can potentially affect the germline, the reproductive system and the soma more generally. The traditional pedigree lines (blue) show chromosomal transmission, which, in addition to the DNA, can potentially include exposure-induced epigenetic marks that escape erasure and impact on offspring development. The germline can potentially transmit exposure-induced non-coding RNAs (ncRNAs) that influence offspring development. Exposure induced metabolic changes can set up a ‘metabolic cascade’ such that changes in the reproductive tract influence early embryo programming of the offspring or change metabolic signals across the placenta. An additional maternal transmission route is the influence of the mother’s microbiome on that of her child.
assumed to be detrimental. Although the usual research design in transgenerational epidemiology is to test associations between parental or ancestral exposure and adverse outcomes in unexposed offspring, transgenerational responses/effects can also be protective ‘adaptations’, potentially lessening the risk of adverse outcomes in successive exposed generations. The latter has been demonstrated in a remarkable male line rat experiment where ancestral carbon tetrachloride-induced liver damage led to inheritable, epigenetically mediated, reprogramming of hepatic healing, such that exposure in the third generation did not produce liver cirrhosis.\(^{21}\) Lastly, the exposures associated with transgenerational effects need not be adverse. Enriched environments can induce transgenerational responses in mice.\(^{22}\)

In summary, a transgenerational perspective is needed if we are to better understand the determinants of major public health problems and adequately assess the value and feasibility of potential interventions. Studies in the field should not be predicated upon a particular mode of transmission from one generation to the next and should make no assumptions about the relative contributions of cultural, genetic or other molecular mechanisms of inheritance. We need to keep an open mind and design appropriate experiments for testing each potential mode of transmission independently.

### HUMAN OBSERVATIONS

Below we summarise human studies by type of exposure and route (paternal or maternal) of transmission. Studies of exposures around the time of conception\(^{23–26}\) are not included.

#### Food supply

**Paternal line**

Compelling historical findings of paternal transgenerational effects in humans come from the Överkalix population in northern Sweden\(^{27–31}\) using samples of individuals born in specified years. Longevity and specific causes of death were linked to detailed historical records of harvests and food supply experienced by previous generations in early life. This series of studies is summarised in Table 1. The initial all-cause mortality study\(^{27}\) noted an exposure-sensitive period during mid-childhood (the ‘slow growth’ period during the few years leading up to the prepubertal growth spurt) but not later in childhood, which was confirmed in a subsequent analysis in 2 of 3 independent cohorts\(^{28}\) and then in a sex-specific analysis of mortality rate in the grandchildren.\(^{29}\) This latter analysis showed mortality rate point estimates of grandsons were 1.67 or 0.65 depending on good or poor food supply of the paternal grandfather during the slow growth period (compared with grandsons of grandfathers with moderate food supply). For granddaughters, the comparable figures were 2.13 and 0.72 for good or poor food supply of the paternal grandmother. This intriguing pattern of sex-specific association persisted when the grandchild’s early life circumstances were taken into account.\(^{30}\) A recent study\(^{31}\) shows that sudden change in food supply between 0 and 13 years of the paternal grandmother is linked to cardiac deaths in the granddaughters raising the possibility that this type of exposure might underlie some of the transgenerational responses reported earlier. Recently, a discussion paper has been posted\(^{32}\) that used data on the German famine of 1916–1918 to look at height, mental health and educational achievement in the descendants of boys exposed at 9–12 and girls at 8–10 years. Among the third generation, males tend to have better mental health scores if their paternal grandfather was exposed, thus providing support for the importance of exposures in the mid-childhood period.

| Reference | Överkalix cohorts by grandchild’s or proband birth year | Prior research question | Main findings | Comments |
|-----------|--------------------------------------------------------|-------------------------|--------------|----------|
| Bygren et al\(^{27}\) | 1905 (n=94) | Any link between ancestral food supply at two periods in childhood, the prepubertal spurt or the period just before and after proband longevity? | Paternal grandfather’s food supply just before prepubertal growth spurt inversely associated with proband longevity | This study defined the mid-childhood ‘slow growth period’ as an exposure period associated with transgenerational effects |
| Kaati et al\(^{28}\) | 1895 (n=107) | Any link between ancestral mid-childhood food supply and proband cardiovascular and diabetes mortality? | Father’s poor, mother’s good, food supply in mid-childhood linked to reduced proband cardiovascular mortality. Paternal grandfather’s good mid-childhood food supply linked to increased proband diabetic mortality | Diabetic mortality was included as a prior hypothesis based on possible role of imprint genes. Each diabetic proband had a different paternal grandfather (Bygren et al. 2006) |
| Pembrey et al\(^{29}\) | 1895 (n=107) | Any sex-specific link between (grand) parental mid-childhood food supply and proband mortality rate ratio? | Paternal grandfather’s food supply linked to grandson’s mortality; paternal grandmother’s food supply to granddaughter’s mortality | Stratification by sex of the proband suggested by early ALSPAC results of paternal smoking effects (Northstone et al. 2014\(^{34}\)). Exposure-sensitive period in mid childhood but not (pre)puberty confirmed |
| Kaati et al\(^{30}\) | 1895 (n=107) | Any link between (grand) parental mid-childhood food supply and proband early-life circumstances and sex-specific longevity? | Grandparental sex-specific transgenerational effects (as above) persisted. Parental effects now revealed as well | Taking proband’s early-life circumstances into account revealed a father to son effect on longevity |
| Bygren et al\(^{31}\) | 1895 (n=107) | Any link between sharp change in grandparental food supply in childhood (0–13ys) and proband cardiovascular mortality? | Sharp change in food supply of paternal grandfather linked to increased cardiovascular mortality in female probands | Prior hypothesis—transgenerational effects of change in supply as the demonstrated effects from gestation to adulthood (Bygren et al. 2000)\(^{32}\) |

---

Pembrey M, et al. *J Med Genet* 2014;51:563–572. doi:10.1136/jmedgenet-2014-102577 565
Maternal line
Human studies of ancestral famine or food supplements during pregnancy on fetal growth of the grandchildren have been recently reviewed. The Guatemalan follow-up studies of the descendants of women with nutrition intervention in pregnancy indicated a positive association between improved pregnancy and first-generation progeny nutrition with grandchild development indicators—primarily higher birth weight and increased length. However, the findings were complicated by supplement use in pregnancy and breast feeding. Cross-sectional studies of the famine in China showed women exposed in utero and early life had bigger babies. A far more precise definition of famine was available in the widely cited Dutch famine studies. At the end of World War II (1944–1945), acute food shortages caused widespread starvation (the Hunger Winter) in the occupied cities of Holland. The first study of multigenerational outcomes concerned birth weights of 1808 firstborn children to mothers, most of them exposed in utero during the famine. The birth weights of babies born to women who were exposed to famine in utero (during the first and second trimesters) were lower than babies of non-exposed women. Subsequent prospective studies revealed a more complicated situation; in essence, the first born of women exposed in utero were heavier than controls, while second and third born were lighter at birth, the reverse of what is normally observed. In other analyses, no association was found between prenatal famine exposure and grandchild birth weight, whereas associations were identified between exposure and childhood adiposity in grandchildren.

Smoking
Paternal line
The study of the age of onset of paternal smoking on offspring outcomes (study child), using the Avon Longitudinal Study of Parents and Children (ALSPAC), was designed to specifically test the Överkalix hypothesis that mid-childhood was an exposure-sensitive period with respect to transgenerational effects. The initial offspring outcomes were those relevant to the Swedish results such as birth characteristics and body mass index (BMI) at ages 7 and 9 years. ALSPAC data, adjusted for paternal smoking at conception, showed that the earlier the father started smoking, the greater the BMI at 9 years of sons, but not daughters. The greatest association was found in sons of men who commenced smoking before the age of 11. Follow-up to age 17 years has just been reported. The adjusted mean differences in BMI, waist circumference and total fat mass in the group of sons whose fathers started smoking <11 years, relative to others, increased with age, being significantly greater (eg, an extra 3–10 kg of fat mass) from 13 years onwards. There was no significant BMI association in daughters, but they showed a reduction in total lean mass.

Maternal line
An ALSPAC study of non-smoking ALSPAC mothers, who were exposed in utero to maternal smoking, showed their sons (but not daughters) to be, on average, 61 g bigger at birth compared with babies where neither the study mother or grandmother smoked. There were no fetal growth differences if the father’s mother had smoked in pregnancy. Misra and colleagues analysed data on maternal and grand-maternal smoking from the Baltimore branch of the US Collaborative Perinatal Project and after many adjustments found there was a 244 g increased weight in offspring of non-smoking mothers if their own mother had smoked prenatally. Similarly, in a study of births to non-smoking women participating in the British 1958 birth cohort, after adjustment for gestation, maternal birth weight, maternal height and BMI, there was a mean increase of 45 g in weight in children of mothers exposed in utero to maternal smoking, but no allowance was made for parity. Neither of these analyses assessed whether there were different effects according to gender of the offspring.

In addition to studies examining anthropometric measures, a few studies have also examined the transgenerational effects of smoking on asthma. One much-quoted study published in 2005 indicated that the risk of childhood asthma was not only influenced by prenatal maternal smoking, but also by the exposure of the mother in utero to her own mother’s smoking. The ALSPAC study did not find an effect of maternal exposure in utero on childhood asthma, although there was suggestive evidence of paternal prenatal exposure being associated with asthma and persistent wheezing in his daughters. Both findings therefore need to be replicated.

Other paternal line exposures
Betel nut
An experimental study of betel nut (Areca catechu) in CD1 mice indicated that paternal exposure transmitted an increased risk of hyperglycaemia and obesity to non-betel-fed first-generation offspring, especially males. This led to an observation study within the Keelung Community-based Integrated Screening Program in Taiwan studying the effect of paternal betel-quid (paan) chewing prior to offspring conception. Chewing usually began around 18 years (Boucher, personal communication) and interestingly, a similar association to that found in mice was observed, with a dose-dependent association of paternal betel-quid use with early metabolic syndrome in the adult offspring who had never chewed betel-quid themselves.

Conclusions from human observations to date
It is fair to say that publicity related to recent publications has drawn some statistical criticism on the issue of multiple testing. Our response points to there being clear prior hypotheses (see table 1) and coherence between different studies/populations. We recognise that research conclusions that extend the public health paradigm need a higher level of evidence than normal, and further replication is needed. However, taken together the above studies demonstrate transgenerational associations between grandpaternal/paternal exposures and health outcomes that are most unlikely to be due to ‘cultural’ inheritance/social patterning or genetic inheritance in isolation. The demonstration of an exposure-sensitive period in mid-childhood prior to puberty, but not in adolescence, provides compelling evidence for some other form of transmission between generations.

LESSONS FROM ANIMAL EXPERIMENTS
This is not the place to review the large number of mammalian experiments characterising transgenerational responses to ancestral exposures. Nevertheless, such experiments have revealed certain features that need to be considered in designing and interpreting human studies.

Sex differences in transgenerational effects
Sex differences occur in most non-communicable diseases, including metabolic diseases, diabetes, hypertension, cardiovascular disease, psychiatric and neurological disorders and cancer,
yet the potential role of non-genetic transgenerational effects in mediating this has rarely been investigated.

Mammalian experiments showing transgenerational responses have reported numerous sex-specific effects after exposure during pregnancy or on paternal exposure before breeding. These can affect offspring of both sexes, solely/predominantly males or solely/predominantly females. Thus, in line with the Överkalix results, the phenotypic outcomes are generally not sex-limited. The X and Y chromosomal difference between the sexes could potentially contribute to sex-specific transgenerational effects. This could include hormonal effects as well as effects that are independent of sex steroids. A recent review of sexual dimorphism emphasises the differential epigenetic processes in the placenta, noting the abundance of X-linked genes involved in placentogenesis and the early unequal gene expression by the sex chromosomes between males and females. Champagne’s comment on sex specificity in transgenerational responses highlights the differences in epigenetic plasticity during gamete maturation. The large number of rodent examples of sex differences in outcomes indicates that the sex differences in outcomes observed in the few human studies (summarised above) are not unexpected. Despite this, such differences tend to be viewed with suspicion, perhaps because of the dangers of post-birth exposure found both behavioural and metabolic disturbances in descendants.

POSSIBLE MEDIATING MOLECULAR MECHANISMS

A true transgenerational effect has several steps:—exposure; the biological embedding or ‘capture’ of the exposure (this may include DNA damage/mutation); transmission of this information between generations; and phenotypic change in (unexposed) offspring/descendants, potentially associated with fetal programming. The mediating mechanism may vary between steps/generations and therefore may require independent investigation as part of an epidemiological framework.

Transmission between generations

As noted above (figure 1), transmission of exposure information down the maternal line may have several different routes with potential to confound any specific mechanism. Thus, proof-of-principle studies often focus on the male line, even if the initial exposure is during fetal development via the pregnant mother/dam. In the latter situation, the exposed pregnant individual is usually termed F0.

Irrespective of the exposure type and timing, a prerequisite for transgenerational effects beyond the second generation is the involvement of germline transmission, which may occur in several ways. First, by directly impacting on the developing primordial germ cells early in utero, as has been reported in mice. In this instance, resulting gametes produced later in life might be anticipated to carry a transmissible ‘memory’ of the original exposure via the mother that can be passed on to subsequent generations. Alternatively, any effect transmitted to the F2 generation via F1 gametes might arise purely in response to the altered development or metabolism of the F1 generation arising from the F0 exposure (see Aiken and Ozanne for a discussion of such ‘propagational programming’). These are two distinct processes, each of which might mediate the effects of an F0 exposure on the F2 and subsequent generations, independently of non-gametic maternal effects. It is worth noting that even male gametic transmission does not necessarily require the molecular ‘memory’ of the original exposure to pass through meiosis. There are tight cell–cell junctions between Sertoli-cells and postmeiotic spermatids that theoretically could act as a route for ‘message delivery’ to the zygote and beyond, and recent work has shown an influence of seminal fluid in conjunction with uterine factors on offspring metabolic development.

A role for epigenetics in transgenerational responses

Epigenetic events are an essential mediator of cellular differentiation and therefore development in multicellular organisms. However, the underlying dynamism and responsiveness to subtle subcellular cues that are a hallmark of epigenetic change during development also make the developing epigenetic profile susceptible to external influence, with the potential to ‘program’
the underlying genome towards an altered phenotype in later life. The labile nature of some epigenetic marks to environmental influence has made epigenetic processes an appealing mechanism for transgenerational responses. In fact, transgenerational epigenetic inheritance has been widely documented in prokaryotes, fungi, plants and animals, via both the male and female germline. In at least some instances (eg, in Caenorhabditis elegans), initial transmission down the female line can ‘spread’ to encompass paternal transmission in subsequent generations.

In mammals, data demonstrating the erasure of nearly all epigenetic marks twice during the life course (early postzygotically and in the developing primordial germ cells in early development) have been difficult to reconcile with an epigenetic mode of transgenerational effects. Nevertheless, for decades it has been known that mammals have a set of imprinted genes; those that are selectively silenced in either the maternal or paternal germline during gamete formation, such that only one allele is expression following fertilisation. Genomic imprinting establishes the principle of transgenerational epigenetic inheritance—the gene is active or silent depending on epigenetic marks placed in the parental generation that survive erasure as they pass to the offspring. The parent-of-origin-dependent gene expression reflects a robust, evolved response to differences in cellular conditions between egg and sperm development. Imprinted genes escape one of the two phases of genome demethylation between generations, namely in the preimplantation embryo soon after fertilisation. Importantly, the epigenetic marks that regulate some imprinting regions have been demonstrated to show interindividual, tissue-specific, age-specific and environmentally sensitive variation.

Recent mouse experiments have also reported a small percentage of loci that escape demethylation at the second phase during primordial gonadal cell differentiation in the early embryo. These are largely associated with repeat sequences, often adjacent to transposable DNA elements. Interestingly, there is growing evidence that some repeat-based transposable elements have gene enhancer activity. It is also now clear that many non-imprinted, non-repetitive genomic loci remain subject to epigenetic control in both sperm and eggs. This applies to both DNA methylation and histone modification profiles. Further, emerging data exist that demonstrate intergenerational variation in epigenetic profile, supporting a role for environmental and/or genetic influence in determining the epigenetic state transmitted to the next generation.

Of particular note is a recent study in mice that provides a plausible mechanism for transgenerational inheritance of environmental information, linking behavioural, neuroanatomical and epigenetic processes. Using an olfactory fear conditioning protocol, wherein an odour (acetophenone) is paired with a mild foot-shock, Dias et al noted a behavioural sensitivity to acetophenone in the next two generations of animals, despite having no exposure to this odour. This was associated with structural changes in the olfactory nervous system accompanying the fear conditioning and distinct hypomethylation of the Olftr151 gene (known to be activated by acetophenone) in sperm of conditioned F0 males and F1-naive offspring. The extent to which other specific environmental exposures elicit similarly specific epigenetic changes in gametes, and the extent to which these are retained during postzygotic development, remains unclear but warrants further investigation.

Another recent study in a non-genetic prediabetes mouse model provides further compelling evidence for gametic transmission of environmentally induced epigenetic change. In this model, offspring of prediabetic fathers show glucose intolerance and insulin resistance in association with distinct changes in gene expression and DNA methylation patterns in pancreatic islets, including reproducible changes in methylation of insulin signalling genes. Interestingly, the sperm of prediabetic fathers have extensive changes in methylation patterns in sperm, many of which overlapped with that of pancreatic islets in offspring.

Non-coding RNAs (ncRNAs) as mediators of transgenerational responses

There is evidence for a role of sperm-derived RNAs in mediating paternal transgenerational effects, with several classes of RNA recently identified in sperm that are likely to be essential factors in male fertility, including many Y-linked ncRNAs. Such RNAs have the potential to play roles in early postfertilisation development.

The specific RNA content of sperm has been assessed in response to specific environmental exposures. Diet-induced paternal obesity in mice alters microRNA (miRNA) content and sperm methylation status in male mice in association with altered metabolic function in a second subsequent generations of progeny. Chronic stress in male mice results in altered miRNA content of sperm in association with reduced HPA stress axis responsiveness in the progeny. Recently, exploiting their transgenerational model of unpredictable maternal absence in neonatal mice as a stress-inducing exposure, Mansuy’s group have examined the associated miRNA changes in sperm. Injection of sperm RNAs from exposed males into fertilised wild-type oocytes reproduced the behavioural and metabolic alterations in the resulting offspring. Interestingly, affected F2 mice had normal sperm miRNAs, yet their F3 offspring still inherited the behavioural phenotype. In humans, the small RNA component of sperm is altered in male smokers, including miRNAs predicted to target the epigenetic regulators. As a whole, the emerging picture suggests that the RNA content of sperm is sensitive to environmental exposures, with the potential to influence early embryonic development, including the establishment of the postzygotic epigenome. It is speculated to be an initiating mechanism for transmission of impaired metabolic health to future generations via the male lineage.

A modifying role for genetics in mediating transgenerational effects

Genetic variation will clearly impact on a variable extent on the apparent heritability of some epigenetic marks as evidenced by multigenerational familial studies and exemplified at one extreme by MLH1 germline epimutation analysis. Originally thought to be solely due to inheritance of a ‘pure’ epigenetic variant (DNA hypermethylation) across generations, in some cases this is genetically driven, being erased in the germline but faithfully re-established early postzygotically in association with a proximal genetic variant or haplotype. Epigenetic responses to exposures and the propensity to be transmitted to the next generation(s) are likely to have evolved under selective pressure. As such one can expect variation in ‘responsiveness’ to a specific exposure to be underpinned by genetic differences. Compelling data in support of this have recently emerged, highlighting the potential of genetic variation in both cis (regions on the same chromosome) and trans (different chromosome) to mediate the effects of environmental exposures on the neonatal epigenetic profile in humans. As such, any study of environmentally driven epigenetic variation transmitted transgenerationally needs to be considered in a background of genetic variation as a modifier of this effect. For example, although...
phenotypic effects are likely to be completely independent of genetic effects. Alternatively, some transgenerational environmentally induced effects may be more nuanced on certain genetic backgrounds. A layer of epigenetic regulation has the potential to make such effects heritable in the absence of the genetic change, for example, by changing the nature of the environment, leading to growth defects in wild-type grandparents. Embryo transfer experiments revealed that epigenetic hypotheses that predict that a specific phenotype occurs through transgenerational epigenetic inheritance.94

Table 2

| Cohort name | Participant number* | General description | Reference |
|-------------|---------------------|---------------------|-----------|
| ABC         | 106 370             | The Aarhus Birth cohort Denmark has collected data during pregnancy and delivery for women since 1989. The associated biobank was established in 2008 to provide the opportunity to investigate the role of genetic factors, environmental exposures and lifestyle factors in the risk of disease in the offspring. | 102 |
| ABIS        | 17 045              | All Babies in Southeast Sweden (ABIS) is a prospective birth cohort of 17 055 children born October 1997–October 1999. Parental and child questionnaire follow-up. Extensive biobank with repeated sampling at follow up. | 103 |
| ALS PAC      | 14 541              | The Avon Longitudinal Study of Parents and Children recruited 14 541 pregnant women with due date April 1991–December 1992. Information on parents’ life taken during study pregnancy. Mothers and children have been followed using questionnaires. Children have been followed at regular clinical visit. Extensive biobank. | 104 |
| BCS 70      | 17 000              | The British Cohort Study follows people born in the UK in a single week of 1970. It has collected information on health, physical, educational and social development, among other factors. | 105 |
| BIB         | 13 776              | Born in Bradford is a longitudinal birth cohort study aimed at recruit a multiethnic cohort of babies born in Bradford (UK) and their parents in order to investigate fetal growth, birth and long-term outcomes by ethnic groups. | 106 |
| DNBC        | 94 837              | The Danish National Birth Cohort recruited pregnant women and their children from 1996 to 2002. Multiple interview data and biopspecimens were collected including food frequency questionnaire in gestational week 24 and periconceptional use of medicine and food supplements. Data for the cohort are collected at regular times from hospital discharge registry and other national registers. | 107 |
| MCS         | 19 000              | The Millennium Cohort Study follows the lives of children born in the UK in 2000–2001. It collects information on siblings and parents covering topics including socioeconomic variables, behaviour and cognitive development. | 108 |
| MOBA        | 108 500             | The Norwegian Mothers and Babies study aims to quantify the influence of various social, genetic, nutritional and environmental exposures on pregnancy outcomes and child health. Data and biopspecimens were collected in pregnancy and at birth. Fathers were also recruited and provided blood. Health outcomes were collected from hospital discharge registries as well as other health registries such as the Medical Birth Registry, the Cancer Registry and the Diabetes Registry. | 109 |
| NCDS        | 17 000              | The National Child Development Study (1958 birth cohort study) follows the lives of children born in the UK in a single week of 1958. It has collected information on physical and educational development, economic circumstances, employment, family life and health behaviour among other variables. DNA bank. | 110 |
| UBCoS Multigen | 7567                | Uppsala Birth Cohort Multigenerational Study exploring several issues highly relevant for health equity research. Life-course approach to analysis detailed biological and social data stretching from birth to old age with access to more than two successive generations. | 111 |

WHAT IS NEEDED FOR FUTURE HUMAN STUDIES?

Non-genetic biological transmission between generations, of which epigenetic hypotheses are an exemplar, spell out how exposures in one generation may lead to effects on health, behaviour or other personal characteristics in later generations. To date human studies of transgenerational responses have ranged from historical analysis of the Överkalix cohorts for which biological samples are not available (at least for all but the 1935 cohort) to the contemporary birth cohort ALS PAC where samples are available on the mother and offspring, plus limited paternal samples. The collection of blood or tissue samples is often seen as the next step to further our knowledge about transgenerational mechanisms in humans. Discoveries of epigenetic and other mechanisms in cells may indeed further our understanding of the aetiological relations. However, this does not mean that large population studies without direct information from human blood or tissue are redundant. On the contrary, epigenetic hypotheses that predict that a specific exposure in one generation will give rise to a specific outcome in a later generation(s) could and should be tested in large-scale population studies that hold information about such exposures and outcomes. Conventional non-experimental epidemiological studies...
studies can still provide an effective means to explain and potentially prevent (adverse) transgenerational responses. Public health interventions may be applied at many different entry points in the aetiological chain of events. Often one and the same condition may be causally attributed to different aetiological antecedents in the web of causation dependent on whether the focus of the study is primarily preventive, predictive, therapeutic or, more generally, explanatory.95 96

There are many birth cohorts and other general population samples worldwide (table 2), and current life-course epidemiology is well placed to extend its reach beyond a single generation and the DNA sequence. It is important to note that from the public health perspective, studying the effect of just early-life parental exposures is likely to increase our understanding of the phenotypic variation, health and well-being of the offspring generation, and perhaps point the way to intervention trials. Common parental exposure information is likely to be available, generation, and perhaps point the way to intervention trials. Phenotype variation, health and well-being of the offspring generation, and perhaps points the way to intervention trials. Standardisation of outcome measures across cohorts is important for many epidemiological studies, including transgenerational studies, and is being actively addressed.

Biological samples

Many cohorts are establishing DNA banks, but it is also worth bearing in mind that neonatal blood spots (Guthrie cards), collected and stored for many decades in some countries, can be a valuable resource for capturing neonatal epigenome and other data.97 In light of the role of miRNAs, discussed above, the collection of cell-free circulating miRNAs is likely to be valuable additional information. Standardisation of outcome measures across cohorts is important for many epidemiological studies, including transgenerational studies, and is being actively addressed.

Acknowledgements

This position paper was developed by the multidisciplinary Network in Epigenetic Epidemiology to encourage transgenerational research in human cohorts. This group (see ‘Contributors’) met four times at the Karolinska Institute since 2009.

Collaborators

LOB, MP, RS, John Carsensten, Department of Medical and Health Sciences, Linköping University, Linköping, Sweden. Sören Edvinsson, 1. Demographic Data Base, Umeå University, Umeå, Sweden, 2. Centre for Population Studies, Umeå University, Umeå, Sweden. Tomas Forsen, Department of Medicine and Health, Unit of Community Medicine/General Practice, Linköping University, Linköping, Sweden. Paul Franks, 1. Genetic and Molecular Epidemiology Unit, Lund University Diabetes Center, Department of Clinical Sciences, Skåne University Hospital, Lund University, Malmö, Sweden, 2. Department of Nutrition, Harvard School of Public Health, Boston, MA, USA, 3. Department of Public Health and Clinical Medicine, Section for Medicine, Umeå University, Umeå, Sweden. Jan-Ake Gustafsson, T. Department of Biosciences and Nutrition, Karolinska Institutet, Huddinge, Sweden, 2. Center for Nuclear Receptors and Cell Signaling, University of Houston, Texas, USA, Gunnar Kaelt, Department of Biosciences and Nutrition, Karolinska Institutet, Huddinge, Sweden. B.I.B. Lindahl, Department of Philosophy, Stockholm University, Stockholm, Sweden, Johnny Ludvigsson, Department of Paediatrics, Unit of Clinical and Experimental Medicine, Linköping University, Linköping, Sweden, L. H. Lunev, Department of Epidemiology, Mailman School of Public Health, Columbia University, New York, New York, USA. Bitte Modin, Centre for Health Equity Studies (CHESS), Stockholm University/Karolinska Institutet, Stockholm, Sweden. Hans Nilsson, Department for Studies of Social Change and Culture, Linköping University, Sweden. Michael Sjöström, Department of Biosciences and Nutrition, Karolinska Institutet, Huddinge, Sweden. Petter Tinghög, Department of Clinical Neurosciences, Karolinska Institutet, Solna, Sweden, Denny Vägerö, Centre of Health and Equity Studies (CHESS), Stockholm University/Karolinska Institutet, Stockholm, Sweden.

Contributors

LOB established and led The Network in Epigenetic Epidemiology, MP and RS drafted this review/position paper drawing on the discussions of the Network. These Network members scrutinised and commented on this final and earlier drafts.

Funding

The Network in Epigenetic Epidemiology was funded by the Swedish Research Council for Health, Working Life and Welfare. Dnr 2008-1153, 2009-2022 and 2010-1421; and also the UK Medical Research Council grant G1100226 to MP.

Competing interests

None.

Patient consent

No.

Provenance and peer review

Commissioned; internally peer reviewed.

Open Access

This is an Open Access article distributed in accordance with the terms of the Creative Commons Attribution (CC BY 4.0) license, which permits others to distribute, remix, adapt and build upon this work, for commercial use, provided the original work is properly cited. See: http://creativecommons.org/licenses/by/4.0/

REFERENCES

1 Forsdahl A. Are poor living conditions in childhood and adolescence an important risk factor for atherosclerotic heart disease? Br J Prev Soc Med 1977;31:91–5.
2 Barker DJ. The fetal and infant origins of adult disease. BMJ 1990;301:1111.
3 Barouki R, Gluckman PD, Grandjean P, Hanson M, Heindel JJ. Developmental origins of non-communicable disease: implications for research and public health. Environ Health 2012;11:42.
4 Modin B, Vägerö D, Hallqvist J, Koupil I. The contribution of parental and grandparental childhood social disadvantage to circulatory disease diagnosis in young Swedish men. Soc Sci Med 2008;66:822–34.
5 Modin B, Erickson RP, Vägerö D. Intergenerational continuity in school performance: do grandparents matter? Eur Soc Rev 2013;29:858–70.
6 Weedon MN, Clark VK, Qian Y, Ben-Shlomo Y, Timpson N, Ebrahim S, Lawlor DA, Pembrey ME, Ring S, Wilkin TJ, Voss LD, Jeffery A, Mielck A, Ferrari L, Cori AM, Murray A, Weber D, Knight B, Shields B, Smith GD, Hattersley AT, Di Rienzo A, Frayling TM. A common haplotype of the glucokinase gene alters fasting glucose and birth weight: association in six studies and population-genetics analyses. Am J Hum Genet 2006;79:991–1001.
7 Aiken CE, Ozanne SE. Transgenerational developmental programming. Hum Reprod Update 2014;20:63–75.
8 Smith GD. Assessing intrauterine influences on offspring health outcomes: can epidemiological studies yield robust findings? Basic Clin Pharmacol Toxicol 2008;102:245–56.
9 Macdonald-Wallis C, Tobias JH, Davey Smith G, Lawlor DA. Parental smoking during pregnancy and offspring bone mass at age 10 years: findings from a prospective birth cohort. Osteoporos Int 2011;22:1809–19.
10 Anway MD, Cupp AS, Uzumcu M, Skinner MK. Epigenetic transgenerational actions of endocrine disruptors and male fertility. Science (New York, NY) 2005;308:1466–9.
11 Franklin TB, Russig H, Weiss IC, Graff J, Linder N, Michalon A, Vizi S, Mansuy IM. Epigenetic Transmission of the Impact of Early Stress Across Generations. Biol Psychiatry 2010;68:408–15.
12 Ng SF, Lin RC, Laybutt DR, Barnes R, Owens JA, Morris MJ. Chronic high-fat diet expression in mammals. Nat Rev Endocrinol 2011;7:366–78.
13 Pembrey ME, Bygren LO, Golding J. The nature of human transgenerational responses. In: Jirtle RL, Tyler FL, ed. Environmental Epigenomics in Health and Disease Epigenetics and Disease Origins. Heidelberg: Springer, 2013:257–1.
18 Susser E, Kirkbride JB, Heijmans BT, Lumey LH, Stein AD. Maternal and grandmaternal smoking and offspring birth weights after maternal intrauterine exposure to smoking. *Cancer Causes Control* 2012;145:1213–18.

19 Nadler A, Kav-Venaki S, Gleitman B. Transgenerational effects of the holocaust: parental transgenerational responses. *Rev Anthropol* 2012;41:577–90.

20 Pembrey M, Stein AD. The induction of glucose intolerance in adult CD1 mice and in their F1 and F2 offspring. *Diabetologia* 2007;50:219–28.

21 Zeybel M, Hardy T, Wong YK, Mathers JC, Wilson CM, Anstee QM, Barter MJ, Mann DA, Stein AD, Pembrey ME. The importance of the intrauterine environment in shaping human growth as a model. *Acta Geneticae Medicae Et Gemellologiae* 1996;45:111–25.

22 Bygren LO, Tinghög P, Carstensen J, Edvinsson S, Kaati G, Pembrey ME, Sjöström M. Change in paternal grandmothers’ economic and health status influenced by the smoking of either parent of their grandchild. *Acta Geneticae Medicae Et Gemellologiae* 2006;55:170–46.

23 Helgason T, Jonasson MR. Evidence for a food additive as a cause of ketosis-prone diabetes. *Lancet* 2008;371:16–20.

24 Leary SD, Smith GD, Rogers IS, Reilly JJ, Wells JC, Hess AR. Smoking during pregnancy and offspring fat and lean mass in childhood. *Obesity (Silver Spring)* 2006;14:2284–93.

25 Proctor JT, Pembrey M, Bygren LO, Lumey LH, Stein AD, Pembrey M, Bygren LO, Pembrey M, Sjöström M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

26 Tobi EW, Lumey LH, Talens RP, Kremer D, Putter H, Stein AD, Slagboom PE, Mansuy IM. Implication of sperm RNAs in transgenerational inheritance of the origins of sexual dimorphism in health and diseases: sex chromosomes and epigenetics. *Bio Sex Differences* 2013;4:5.

27 Stein AD, Wang M, DiGioromo A, Grajeda R, Ramakrishnan U, Ramirez-Zea M, Yount K, Martorell R. Nutritional supplementation in early childhood, schooling, and intellectual functioning in adulthood: a prospective study in Guatemala. *Arch Pediatr Adolesc Med* 2008;162:612–8.

28 Pembrey ME, Bygren LO, Lumey LH, Stein AD, Pembrey M, Sjöström M, Dominici F, Wijmenga C, van Duijn CM. DNA methylation differences after exposure to prenatal famine are common and timing- and sex-specific. *Hum Mol Genet* 2009;18:4046–53.

29 Bygren LO, Kaati G, Edvinsson S. Longevity determined by paternal ancestors’ nutrition during their slow growth period. *Acta Biotheor* 2001;49:53–9.

30 Kaati G, Bygren LO, Edvinsson S. Cardiovascular and diabetes mortality determined by nutrition during parents’ and grandparents’ slow growth period. *Eur J Hum Genet* 2002;10:682–8.

31 Bygren LO, Pembrey M, Bygren LO, Kaati G, Edvinsson S, Northstone K, Sjostrom M, Pembrey M. Prepubertal start of fat deposition and offspring fat and lean mass in childhood. *Obesity (Silver Spring)* 2006;14:159–66.

32 van den Berg BJ, Pembrey M. A validation study fo transgenerational effects determined by nutrition during parents’ and grandparents’ slow growth period. *Eur J Hum Genet* 2002;10:682–8.

33 Pembrey ME, Bygren LO, Pembrey M, Bygren LO, Northstone K, Pembrey M, Bygren LO, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

34 Li YF, Langholtz B, Salam M, Gilland FD. Maternal and grandmaternal smoking patterns are associated with early childhood asthma. *Chest* 2005;127:1232–41.

35 Miller LL, Pembrey M, Northstone K, Pembrey M, Golding J. Do grandmaternal smoking patterns influence the aetiology of childhood asthma? *Chest* 2014;145:1213–18.

36 Boucher BJ, Even SW, Stowers JM. Betel nut (Areca catechu) consumption and the induction of glucose intolerance in adult CD1 mice and in their F1 and F2 offspring. *Diabetologia* 1994;37:49–55.

37 Pembrey ME, Bygren LO, Northstone K, Golding J, Team AS. Sex-specific, male-line transgenerational responses in humans. *Eur J Hum Genet* 2006;14:159–66.

38 Pembrey ME, Bygren LO, Pembrey M, Pembrey M, Northstone K, Golding J, Pembrey M, Bygren LO, Northstone K, Pembrey M. Male-line transgenerational responses in humans. *Hum Fertil (Camb)* 2010;13:268–71.

39 Northstone K, Golding J, Davey Smith G, Miller LL, Pembrey M. Prepubertal start of fat deposition and offspring fat and lean mass in childhood. *Obesity (Silver Spring)* 2006;14:159–66.

40 Pembrey M, Bygren LO, Northstone K, Golding J, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

41 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

42 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

43 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

44 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

45 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

46 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

47 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

48 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

49 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

50 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

51 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

52 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

53 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

54 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

55 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.

56 Pembrey ME, Bygren LO, Northstone K, Pembrey M, Bygren LO, Northstone K, Pembrey M, Sjostrom M. Transgenerational response to nutrition, early life experiences and longevity. *Eur J Hum Genet* 2007;15:784–90.
maternal environmental and nutrient supply line factors with DNA methylation at the imprinted IGF2/H19 locus in multiple tissues of newborn twins. Epigenetics 2013;8:1069–79.

73 Hackett JA, Sengupta R, Zylicz JJ, Murakami K, Lee C, Down TA, Surani MA. Germline DNA demethylation dynamics and imprint erasure through 5-hydroxymethylcytosine. Science (New York, NY) 2013;339:448–52.

74 Mukamel Z, Tanay A. Hypomethylation marks enhancers within transposable elements. Nat Genet 2013;45:717–8.

75 Smallwood SA, Tomizawa S, Krueger F, Ruf N, Carli N, Segonds-Pichon A, Sato S, Hata K, Andrews SR, Kelsey G. Dynamic CpG island methylation landscape in oocytes and preimplantation embryos. Nat Genet 2011;43:811–4.

76 Tomizawa S, Nowacka-Woszuk J, Kelsey G. DNA methylation establishment during oocyte growth: mechanisms and significance. Int J Dev Biol 2012;56:867–75.

77 Hammond SS, Nix DA, Zhang H, Punvar J, Carroll DT, Cairns BR. Distinctive chromatin in human sperm packages genes for embryo development. Nature 2009;460:473–8.

78 Bryckaerts SA, Hisano M, Eriks S, Ramos L, Daleke EJ, Roloff TC, Beisel C, Schubeler D, Stadler MB, Peters AH. Repressive and active histone methylation mark distinct promoters in human and mouse spermatozoa. Nat Struct Mol Biol 2010;17:679–87.

79 Carroll DT, Hammond SS. The human sperm epigenome and its potential role in embryonic development. Mol Hum Reprod 2010;16:37–47.

80 Gannon JR, Emery BR, Jenkins TG, Carrell DT. The sperm epigenome: implications for the embryo. Adv Exp Med Biol 2014;791:53–66.

81 Wei Y, Yang CR, Wei YP, Zhao ZA, Hou Y, Schatten H, Sun QY. Paternally induced transgenerational inheritance of susceptibility to diabetes in mammals. Proc Natl Acad Sci USA 2014.

82 Hitchins MP, Wong JJ, Suthers G, Suter CM, Martin DI, Hawkins NJ, Ward RL. Germline DNA demethylation dynamics and imprint erasure through 5-hydroxymethylcytosine. Science (New York, NY) 2013;339:448–52.

83 Rodgers AB, Morgan CP, Bronson SL, Revello S, Bale TL. Paternal stress exposure alters sperm microRNA content and reprograms offspring HPA stress axis regulation. J Neurosci 2013;33:9003–12.

84 Maszyczuk E1, Amoco AA, Konje IC, Gant TW, Maszyczuk TH. Smoking induces differential miRNA expression in human spermatozoa: a potential transgenerational epigenetic concern? Epigenetics 2012;7:432–9.

85 Gertz J, Varley KE, Surani MA. Germline DNA demethylation dynamics and imprint erasure through 5-hydroxymethylcytosine. Science (New York, NY) 2013;339:448–52.

86 Hitchins MP, Wong JJ, Suthers G, Suter CM, Martin DI, Hawkins NJ, Ward RL. Inheritance of a cancer-associated MLH1 germ-line epimutation. N Engl J Med 2007;356:697–705.

87 Hitchins MP, Rapkins RW, Kwok CT, Sirivastava S, Wong JJ, Khashigian LM, Polly P, Goldblatt J, Ward RL. Dominantly inherited constitutional epigenetic silencing of MLH1 in a cancer-affected family is linked to a single nucleotide variant within the 5′UTR. Cancer Cell 2011;20:200–13.

88 Kwok CT, Vogelaar IP, van Zelst-Stams WA, Mensenkamp AR, Ligtenberg MJ, Rapkins RW, Ward RL, Chun N, Ford J, Ladaabu U, McKinnon WC, Greenblatt MS, Hitchins MP. The MLH1 c.27C>T-A and c.853G>T variants are linked to dominantly inherited MLH1 epimutation and are borne on a European ancestral haplotype. Eur J Hum Genet 2014;22:617–24.

89 Teh AL, Pan H, Chen L, Ong ML, Dogra S, Wong J, Maciasac JL, Mah SM, McEvon LM, Saw SM, Godfrey KM, Chong YS, Kwok K, Kwoh CK, Soh SE, Chong MF, Barton S, Karmani N, Cheong CY, Buschdorf JP, Stunkel W, Kobor MS, Mearney MJ, Gluckman PD, Holford JT. The effect of genotype and in utero environment on inter-individual variation in neonate DNA methylomes. Genome Research 2014;24:1064–74.

90 Yazbek SN, Spiezo SN, Nadeau JH, Buchner DA. Maternal epigenetic and growth controls body weight and food intake for multiple generations. Hum Mol Genet 2010;19:434–44.

91 Hara MR, Kovacs JJ, Whelan EJ, Rajagopal S, Strachan RT, Grant W, Towers AJ, Williams B, Lam CM, Xiao K, Shenvy SK, Gregory SG, Ahn S, Duckett BR, Lefkowitz RJ. A stress response pathway regulates DNA damage through beta2-adrenergoreceptors and beta-arrestin-1. Nature 2011;477:349–53.

92 Yazbek SN, Spiezo SN, Nadeau JH, Buchner DA. Maternal epigenetic and growth controls body weight and food intake for multiple generations. Hum Mol Genet 2010;19:434–44.

93 Bygren LO, Nordinfelt L. Causation: an essay. Stockholm: Akademilitteratur, 1981.

94 Lindahl BB. On causal attribution. Stockholm: Stockholm University (Acta Universitatis Stockholmiensis), 2009.

95 Nordenfelt L. On causal attribution. Stockholm: Stockholm University (Acta Universitatis Stockholmiensis), 2009.

96 Beyan H, Down T, Rapkins RW, Chun N, Ford JM, Ladabaum U, McKinnon WC, Rapkins RW, Chun N, Ford JM, Ladabaum U, McKinnon WC, Heshmati A, Mishra G, Koupil I. Childhood and adulthood socio-economic position and hypertensive disorders in pregnancy: the Uppsala Birth Cohort Study. J Epidemiol Community Health 2013;67:939–46.
