At age 9, the methylome of assisted reproductive technology children that underwent embryo culture in different media is not significantly different on a genome-wide scale

Rebekka M. Koeck, Florence Busato, Jorg Tost, Heleen Zandstra, Sylvie Remy, Sabine Langie, Marij Gielen, Ron van Golde, John C.M. Dumoulin, Han Brunner, Masoud Zamani Esteki, and Aafke P.A. van Montfoort

1Department of Clinical Genetics, Maastricht University Medical Centre+, Maastricht, The Netherlands 2Department of Genetics and Cell Biology, GROW School for Oncology and Reproduction, Maastricht University, Maastricht, The Netherlands 3Laboratory for Epigenetics & Environment, CEA-Centre National de Recherche en Genomique Humaine, Evry, France 4Department of Obstetrics and Gynaecology, GROW School for Oncology and Reproduction, Maastricht University Medical Centre+, Maastricht, The Netherlands 5Health Unit, Flemish Institute for Technological Research (VITO), Mol, Belgium 4Department of Pharmacology & Toxicology, School for Nutrition and Translational Research in Metabolism (NUTRIM), Maastricht University, The Netherlands 7Department of Epidemiology and Nutrition and Translational Research in Metabolism (NUTRIM), Maastricht University Medical Centre, Maastricht, the Netherlands 8Department of Human Genetics, Radboud University Medical Center, Nijmegen, The Netherlands

*Correspondence address. Laboratory of Cellular Genomic Medicine, Department of Clinical Genetics, Maastricht UMC+, P. Debyelaan 25, 6229 HK Maastricht, The Netherlands. Tel: +31-43-38-75306; E-mail: masoud.zamaniesteki@mumc.nl

**https://orcid.org/0000-0003-3909-0050 (M.Z.E.); Department of Obstetrics and Gynaecology, IVF laboratory, Maastricht UMC+, PO Box 5800, 6202 AZ Maastricht, The Netherlands. E-mail: aafke.van.montfoort@mumc.nl

Submitted on May 14, 2022; resubmitted on September 5, 2022; editorial decision on September 14, 2022

**STUDY QUESTION:** Can we detect DNA methylation differences between ART children that underwent embryo culture in different media?

**SUMMARY ANSWER:** We identified no significant differences in site-specific or regional DNA methylation between the different culture medium groups.

**WHAT IS KNOWN ALREADY:** Embryo culture in G3 or K-SICM medium leads to differences in embryonic, neonatal and childhood outcomes, including growth and weight. The methylome may mediate this association as the period of in vitro culture of ART treatments coincides with epigenetic reprogramming.

**STUDY DESIGN, SIZE, DURATION:** This study was conducted as a follow-up to a previous culture medium comparison study in which couples were pseudo-randomized to embryo culture in G3 or K-SICM medium. Of the resultant singletons, 120 (n = 65 G3, n = 55 K-SICM), were recruited at age 9.

**PARTICIPANTS/MATERIALS, SETTING, METHODS:** The ART children provided a saliva sample from which the methylome was analysed using the Infinium MethylationEPIC array. After quality and context filtering, 106 (n = 57 G3, n = 49 K-SICM) samples and 659 708 sites were retained for the analyses. Differential methylation analyses were conducted using mixed effects linear models corrected for age, sex, sample plate and cell composition. These were applied to all cytosine-guanine dinucleotide (CpG) sites, various genomic regions (genes, promoters, CpG islands (CGIs)) and as a targeted analysis of imprinted genes and birth weight-associated CpG sites. Differential variance was assessed using the improved epigenetic variable outliers for risk prediction analysis (iEVORA) algorithm and methylation outliers were identified using a previously defined threshold (upper or lower quartile plus or minus three times the interquartile range, respectively).

---

1These authors jointly supervised this study.

© The Author(s) 2022. Published by Oxford University Press on behalf of European Society of Human Reproduction and Embryology. This is an Open Access article distributed under the terms of the Creative Commons Attribution-NonCommercial License (https://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com
Introduction

Year-on-year a worldwide increase in the number of ART procedures has led to the birth of more than 8 million babies (Adamson et al., 2019) and currently, ~3% of births in European countries are conceived through ART (Wyns et al., 2020). Even though most ART children appear healthy at birth, follow-up studies have reliably shown that, compared to their naturally conceived counterparts, ART offspring are at increased risk of adverse perinatal (Berntsen et al., 2019), childhood and later life outcomes (Ceelen et al., 2009; Hann et al., 2018). The perinatal risks include premature birth, low birth weight, being small for gestational age and perinatal mortality (Berntsen et al., 2019), while the later life outcomes relate mainly to cardiometabolic health, including weight (Ceelen et al., 2009; Guo et al., 2017; Hann et al., 2018; Bay et al., 2019). Specifically, the affected cardiometabolic parameters include an increase in both systolic and diastolic blood pressure (Guo et al., 2017) and features of cardiovascular dysfunction, such as suboptimal cardiac diastolic function and increased blood vessel thickness (Guo et al., 2017). Additionally, ART children have been shown to have significantly lower weights than their naturally conceived counterparts from birth until the age of 4 (Bay et al., 2019). These observations are in line with the developmental origins of health and disease (DOHaD) paradigm, which states that early life adversity predisposes an individual to disease in later life (Wadhwa et al., 2009), therefore raising concerns about the impact of ART procedures on the resultant children.

During ART treatments, embryos undergo in vitro culture for 2–5 days prior to intratumin transfer to establish a pregnancy. Throughout this time, embryos are exposed to an artificial in vitro environment consisting of the culture medium, the atmospheric conditions (oxygen levels) and laboratory-specific factors, such as laboratory plastics. Over time, a number of compositionally different culture media have been used for ART procedures (Mantikou et al., 2013; Morbeck et al., 2014; Sunde et al., 2016; Morbeck et al., 2017) and these have been associated with differences in short- and long-term outcomes of the resultant offspring both in animal and human studies (Fernández-González et al., 2004; Donjacour et al., 2014; Zandstra et al., 2015; Bouillon et al., 2016; Velazquez et al., 2018). We previously conducted a culture medium trial, in which couples undergoing ART treatments in Maastricht were pseudo-randomized, by strict alternation, to embryo culture in G3 (Vitrolife) or Sydney IVF cleavage medium (K-SICM, Cook). Although several components of these culture media are known, their precise concentrations and composition are not fully disclosed by the manufacturers. However, a possible difference between G3 and K-SICM is that the version of K-SICM used for the treatments described in these studies, contained an unstable form of the amino acid glutamine (according to product inserts from the media in this period), while G3 contained a more stable dipeptide form of the same amino acid. Several studies have shown that ammonium accumulation during storage of media containing the unstable form of glutamine is significantly higher (Kleijkers et al., 2016) and that an increased ammonium concentration in culture medium has an adverse effect on embryonic development (Virant-Klun et al., 2006; Hashimoto et al., 2008). Embryos cultured in G3 were found to have a greater number of cells, while their morphological grade was lower than K-SICM embryos (Dumoulin et al., 2010). Thereafter, implantation and pregnancy rates were higher in the G3 group (Dumoulin et al., 2010). Interestingly, growth differences, namely increased growth in the G3 group, could already be detected by ultrasound in the second trimester (Nelissen et al., 2013), were evident at birth (Dumoulin et al., 2010; Nelissen et al., 2012) and persisted at age 2 (weight) (Kleijkers et al., 2014) and age 9 (Zandstra et al., 2018a). At age 9, children from the G3 group remained heavier with higher waist circumferences and truncal adiposity than children from the K-SICM group (Zandstra et al., 2018a). Other markers of cardiovascular health, including blood pressure, lipid profile and endothelial function, were comparable between the culture medium groups (Zandstra et al., 2018a). Similarly, cognitive development was comparable between the culture medium groups (Zandstra et al., 2018b).

To date, the molecular mechanisms mediating the relationship between culture medium composition and the observed outcomes are not fully understood. It has been suggested that the epigenome, and
specifically DNA methylation in which a methyl (-CH$_3$) group is added to the cytosine base of cytosine-guanine dinucleotides (CpG), is sensitive to environmental perturbations and subsequently "programs" an individual’s disease susceptibility (Felix and Cecil, 2019). Furthermore, pre-implantation human embryos are undergoing epigenetic reprogramming, consisting of virtually complete erasure and re-establishment of DNA methylation marks (Li et al., 2018; Hanna et al., 2018), during which they may be especially sensitive to environmental regulation of the epigenome. DNA methylation-associated imprinting disorders, although still rare, are also more common amongst IVF children (DeAngelis et al., 2018). Consequently, methylome profiling of ART offspring born after culture medium trials has been carried out. For instance, our group examined DNA methylation in tissues collected at birth, namely the placenta (Mulder et al., 2020) and umbilical cord blood (Koeck et al., 2022), from neonates born after a multicentre randomized controlled trial comparing G5 (Vitrolife) and human tubal fluid (HTF, Lonza) media. Neither study identified any significant DNA methylation differences between the culture medium groups and the group mean differences at the sites analysed were small (largely <10%) (Mulder et al., 2020; Koeck et al., 2022). Interestingly, we identified several CpG sites with differential variability between the culture medium groups (Koeck et al., 2022). These could relate to prenatal factors, such as the pregnancy complications, only experienced by some individuals in each group, alternatively, they could represent epigenetic marks indicative of disease that only some individuals will develop later in life. To investigate this further, later life or longitudinal methylome studies of ART offspring are required. Thus far, the methylome of ART children (age 7 or 8) has only been characterized in one cohort, in the context of a culture medium trial (global medium—LifeGlobal vs single-step medium—Irvine Scientific), finding no significant DNA methylation differences, and small intragroup differences (mostly <10%) (Barberet et al., 2021; Ducrœux et al., 2021).

Not in the context of culture media trials, the methylome has also been compared between ART and naturally conceived offspring and such studies were the focus of a recent meta-analysis (Barberet et al., 2022). In neonates, the meta-analysis only identified consistent differential methylation at the paternally expressed gene 1/mesoderm specific transcript (PEG1/MEST) imprinting gene locus when including all targeted methylome studies of placenta and cord blood samples (Barberet et al., 2022). The findings from genome-wide analyses conducted on samples collected during the neonatal period have yielded contradictory result, with some studies identifying differentially methylated sites, while other studies found no differences (El Hajj and Haff, 2013; Melamed et al., 2015; Barberet et al., 2022; Håberg et al., 2022). In children, on the other hand, the meta-analysis of targeted methylation studies on blood and saliva samples, identified no significant differences between ART and naturally conceived individuals (Barberet et al., 2022). Similarly, genome-wide studies comparing ART and naturally conceived children (Ducrœux et al., 2021; Yeung et al., 2021), adolescents (Penova-Veselinovic et al., 2021) or adults (Novakovic et al., 2019), found no or few significant differences. Although this suggests that any ART-associated methylation differences present at birth do not persist into adulthood, these results come from a small number of studies that have not been validated by other groups. Additionally, the studies lack a detailed description of the ART culture conditions that these individuals were exposed to, meaning that their effect on the methylome cannot be established. As already described, clinical differences are observed between ART offspring that were exposed to different culture environments, highlighting the need to specifically examine the methylomes of these ART sub-groups.

Here, we describe the saliva methylome of ART children (aged 9 or 10) that underwent embryo culture either in G3 (Vitrolife) or K-SICM (Cook) media. For this, we profiled DNA methylation on a genome-wide scale, using the EPIC array, in the largest cohort of its kind to date. Comparison to naturally conceived children was attempted using data from the Flemish Environment Health Study (FLEHS) (Fig. 1A).

**Materials and methods**

**Ethical approval**

This study was registered in the Dutch Trial register (trial number NL4083) and was approved by the ethical review board of the Maastricht University Medical Centre (MUMC+). Both parents of the children provided written, informed consent.

**Study population and sample collection**

Samples for methylome analysis were collected during medical follow-up of children born after a previously conducted culture medium comparison study (Sandstra et al., 2018a). Between July 2003 and December 2006, Vitrolife G1™ Version 3 (G3) (Göteborg, Sweden) and K-SICM from Cook (Brisbane, Australia) were used at MUMC+. Consecutive IVF treatments (with or without ICSI) were strictly alternated between the two media types, while all other ART procedures remained consistent (Dumoulin et al., 2010; Nelissen et al., 2012). Parents of all liveborn singletons from this study were approached for a follow-up investigation after the 9th birthday of the child. In addition to growth and cardiometabolic measurements, 2 ml of saliva were collected using the Saliva DNA Collection, Preservation and Isolation Kit (Norgen Biotek, Thorold, Canada). The saliva samples were collected after an overnight fast (necessary for blood glucose measurements) and after the children had rinsed their mouths with water. Subsequently, the preservation liquid was added to the tube and the samples were stored at room temperature, according to the manufacturer’s instructions.

**DNA extraction**

DNA was extracted using the Saliva DNA Collection, Preservation and Isolation Kit (Norgen Biotek, Thorold, Canada) according to the manufacturer’s instructions. Briefly, 2 ml of preserved saliva was mixed with 80 µl of proteinase K, 800 µl of Binding buffer B and 2.88 ml of isopropanol. After centrifugation, the resulting DNA pellet was washed in 70% ethanol and air-dried. The pellet was rehydrated in 300 µl of Tris–EDTA buffer. DNA quantity and quality were determined using a Nanodrop ND-1000 spectrophotometer (Nanodrop Technologies, Wilmington, USA).

**Bisulphite conversion and DNA methylation profiling by EPIC array**

Prior to DNA quantification, DNA samples containing precipitated material were heated to 65°C and centrifuged. Thereafter, 1 µg of
Figure 1. Genome-wide DNA methylation analysis of ART children that underwent embryo culture in different media revealed no significant differences. (A) Schematic overview showing sample collection/inclusion of ART-conceived and naturally conceived children, methylome data generation alongside data processing and analyses included in this study. (B) Principal component analysis (PCA) of all cytosine-guanine dinucleotide (CpG) sites passing our quality control criteria in data from saliva samples of ART children that underwent embryo culture in G3 (orange) or Sydney IVF cleavage medium (K-SICM, purple) medium. (C) Density plot showing the distribution of beta values from all sites and samples within each group (G3 = orange, K-SICM = purple).
DNA was bisulfite-treated using the EpiTect® Fast 96 DNA Bisulfite Kit (Qiagen Hilden, Germany) and analysed using the Infinium Human MethylationEPIC BeadChip Kit (Illumina, CA, USA) according to the manufacturer’s protocol.

Data analysis

All data analysis was conducted using R (version 3.6.3) (R Core Team, 2021) and visualized using the ggplot2 and ComplexHeatmap packages (Gu et al., 2016). Custom R code used for the processing and analysis of the data described in this article are available at https://github.com/CellularGenomicMedicine/saliva_methyolome.

Participant characteristics

The participant characteristics were compared for differences between the culture medium groups using Student’s t-tests for continuous variables and Pearson’s chi-squared tests for categorical variables.

Quality control and pre-processing

Data were pre-processed using pre-processing functions contained within the RnBeads package (Müller et al., 2019). Data were normalized using the subset-quantile within array normalization (SWAN) method (Maksimovic et al., 2012). Poor quality probes and samples were removed using the greedycut algorithm with a detection P-value threshold of 0.05. Sites were further filtered out based on the following criteria: (i) if they were located on the sex chromosomes, (ii) if they were in close proximity to single-nucleotide polymorphisms, (iii) if more than 5% of the samples contained missing values and (iv) if they were not in a CpG context. Sites containing any missing values (between 0% and 5% missing values) were only used to aggregate sites into regions, they were excluded for all single site-based analyses. Unless indicated otherwise, we used methylation beta values which represent the methylated signal divided by the sum of the methylated and unmethylated signal at each CpG site. Sample sex prediction was carried out as described by Jung et al. (2018) and implemented in the sEst package. Briefly, the X and Y chromosome beta value distributions of the test (ART) samples were combined with those of the reference male and female samples (Jung et al., 2018). Subsequently, a principal component analysis (PCA) was conducted separately for each sex chromosome (referred to as PCA.X and PCA.Y). Then, k-means clustering (number of groups = 2) was conducted on the results from PCA.X and PCA.Y to determine if the test (ART) samples cluster with the female or male samples of the reference samples, respectively. When the k-means clustering for both sex chromosomes assigned the same sex, samples were labelled as male or female, all other samples were labelled ‘not specified’. The X principal component (PC)1 threshold, to distinguishing between the presence of one or two X chromosomes, was set at 0.05. Principal component PC1 of the PCA.X and PCA.Y are shown in Fig. 2A and Supplementary Fig. S1A, without showing the reference samples that were analysed alongside the test (ART) samples.

Cellular deconvolution of saliva samples

Cell composition of the samples was estimated using the reference-based Houseman algorithm (Houseman et al., 2014) implemented using the easwastools package (Heiss and Just, 2018). Probes for cell-type deconvolution were identified using DNA methylation signatures from sorted saliva samples collected from children (Middleton et al., 2022) (available at Gene Expression Omnibus GSE147318) yielding an estimate of the proportion of leucocytes and epithelial cells.

Comparison of G3 and K-SICM children

We applied PCA on all high-quality CpG sites. The beta values were centred but not scaled for the PCA. Associations between the PCs and technical or demographic features of our samples were tested either (i) using permutation tests with 10,000 permutations to determine the significance of correlations (age, leucocytes, epithelial cells), (ii) using a two-sided Wilcoxon rank test for categorical variables creating two groups (sex, culture medium, sample plate) or (iii) using a Kruskal–Wallis one-way ANOVA for categorical variables generating three or more groups (Sentrix ID, Sentrix position—chip number and sample position, respectively).

Methylation M-values, representing the log2 ratio of the methylated probe intensity compared to the unmethylated probe intensity (Du et al., 2010), were used to test for associations between DNA methylation and culture medium using mixed effects linear models implemented with the variancePartition package (Hoffman and Schadt, 2016). The models were adjusted for a priori chosen potential confounders: age at sample collection, sex and cell composition as fixed effects alongside batch correction (sample plate) as a random effect. They were applied to individual CpG sites or aggregate values of multiple CpG sites to identify differentially methylated positions (DMPs) and differentially methylated regions (DMRs), respectively. To aggregate sites into regions, we calculated the mean of all the beta values from probes attributed to the same gene, promoter or CpG island (CGI). For the targeted analyses, the aforementioned models were applied to (sites within) imprinted genes (Ginjala, 2001) and sites associated with birth weight (Küppers et al., 2019). The Benjamini–Hochberg (Benjamini and Hochberg, 1995) method was used to correct all analyses for multiple testing, and an adjusted P-value of <0.1 was considered significant.

DNA methylation outliers were defined as previously described by Gentilini et al. (2015). Briefly, hypomethylation outliers were defined as beta values more than three interquartile ranges (IQRs) below the 25th percentile, while hypermethylation outliers were defined as beta values more than three IQRs above the 75th percentile. The IQR and percentile thresholds were calculated across all saliva samples. We tested for associations between the log10 transformed number of outliers and culture medium using the mixed effects linear models described above. The significance of associations between the number of outliers per sample and various clinical and technical features were assessed using permutation tests with 10,000 permutations for continuous variables (age and weight at sample collection, birth weight, leucocyte proportion, epithelial cell proportion) and with two-sided Wilcoxon rank tests for categorical variables (sample plate). To identify differentially variable sites, the improved epigenetic variable outliers for risk prediction analysis (iEVORA) algorithm (Teschendorff et al., 2016) was applied using the matrixTests package (Koncevicius, 2020). In brief, iEVORA applies Bartlett’s test, a parametric test for differential variance, and a Student’s t-test to each CpG site. Subsequently, sites reaching significance after multiple testing correction (false discovery rate (FDR) corrected P-value < 0.05) of the Bartlett’s test and reaching nominal significance (P-value < 0.05 without multiple
testing correction) of the t-test are considered significant. This approach regularizes the result of the Bartlett’s test which is usually overly sensitive to single outliers. The resulting differentially variable sites were used for Gene ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses using functionality from the missMethyl package (Phipson et al., 2016).

**Comparison with naturally conceived children**

Saliva methylome data were obtained from children enrolled in the longitudinal birth cohort FLEHS; the cohort has previously been described in detail (Langie et al., 2018; Van Den Heuvel et al., 2020).
From these children, saliva samples were collected at the age of 10–11 and the methylome was profiled using the 450K array (Illumina, CA, USA). Data preprocessing and the analysis procedures were largely the same as those described above, for more details see the Supplementary materials and methods.

## Results

Of the 294 ART singletons from the culture medium study, 136 (48%) parent couples agreed for their child to participate. Three of the children failed or refused to provide 2 ml of saliva, a further 7 samples yielded insufficient DNA for processing and 6 samples did not meet quality criteria after bisulfite conversion. Therefore, methylome analysis by EPIC array was carried out on 120 saliva samples (n = 65 for G3, n = 55 for K-SICM). Quality control procedures on the generated data led to the exclusion of 5 poor quality samples (n = 3 for G3 and n = 2 for K-SICM) and 9 samples (n = 5 for G3 and n = 4 for K-SICM) with an undefined sex prediction (Fig. 2A). The characteristics of both groups were comparable (Table I). Although the weight of G3 offspring at birth (Dumoulin et al., 2010) and age 9 (Zandstra et al., 2018a) was reported to be higher than that of K-SICM offspring in the culture medium trial, these parameters were not significantly different in this sub-group of the original study. Nonetheless, the trend is the same.

Of ~850,000 profiled CpG sites, 666,262 were retained for analysis after the filtering procedures, of which 659,708 sites contained no missing values and were used for downstream analysis.

### Global analysis of DNA methylation

Global DNA methylation was first assessed by PCA (Fig. 1B), which did not show clear separation of the G3 and K-SICM groups within the first 4 PCs, which explain 49% of the variance in our data. Only PCs 5 and 6, representing 5.2% and 4.8% of the variance in our data, respectively, were significantly associated with the culture medium (Fig. 2B), thus suggesting that the culture medium is not the main source of variance within our data. Other significant associations were found between some of the first 8 PCs and certain technical or demographic factors, namely sample plate (PC1, PC3, PC5, PC7, PC8), sex (PC7), leucocytes (PC1-3) and epithelial cells (PC1, 2) (Fig. 2B and C). These factors are therefore corrected for in the subsequent analyses. Similarly, the distribution of beta values was very similar in both culture medium groups (Fig. 1C).

### Analysis of DNA methylation at individual CpG sites

We assessed the association of DNA methylation with culture medium at all individual CpG sites using mixed effects linear models corrected for potential confounders. We found no significant DMPs between the two culture medium groups after adjusting for multiple testing (adjusted P-value < 0.1) (Fig. 3A, Supplementary Figs S2A and S3). Additionally, the group mean differences at all sites were small, with <1% of sites (17 in total) having a group mean difference of more than 10% and the maximum group mean difference being 13.5% (Supplementary Table S1).

To reduce the number of comparisons in our analysis, we focused on genomic regions of potential relevance to our cohort, i.e. sites within imprinted genes (Ginjala, 2001) and sites associated with birth weight (Küpers et al., 2019). After our quality control and filtering procedure, 8940 sites within 207 imprinted genes were retained for the analysis. Among these sites, no significant DMPs were identified, and the maximum group mean was 7.4% (Fig. 3A, Supplementary Fig. S2A). Of the 914 previously identified birth weight-associated CpG sites (Küpers et al., 2019) 726 passed our quality control (QC) criteria and none were significantly differentially methylated with a maximal group mean difference of 3.0% (Fig. 3B, Supplementary Fig. S2B).

### Regional analysis of DNA methylation

We then analysed DNA methylation across larger regions of the genome, namely whole genes, promoters and CGIs. In total, 32,564 genes were included in the analysis. Of these, the maximal group difference was found to be 9.4% and no significantly differentially methylated genes were identified between the culture medium groups. A targeted analysis of only the imprinted genes showed that these had even lower group mean differences (maximum group mean difference 1.2%) than the individual sites within these genes (Fig. 3C, Supplementary Fig. S2C). Furthermore, no imprinted genes were found to be significantly differentially methylated between the G3 and K-SICM groups. A total of 41,519 promoters and 25,224 CGIs were included in our analysis. The maximal group mean differences were 9.4% and 8.4% for promoters and CGIs, respectively, and no promoters or CGIs were found to be significantly differentially methylated between the culture medium groups (Fig. 3D and E, Supplementary Fig. S2D and E).

### DNA-methylation variance in ART samples

To assess whether stochastic DNA methylation alterations contribute to the phenotypes observed in the culture medium trial, DNA methylation outliers were identified using previously defined thresholds (Gentilini et al., 2015) and differential variance was assessed using the iEVORA method (Teschendorff et al., 2016). Overall, we found a predominance of hypomethylation outliers compared to hypermethylation outliers (92,238 hypomethylation outliers vs 33,009 hypermethylation outliers). On average, we identified a total of 254 ± 485 (median ± IQR) outliers per G3 sample and 186 ± 153 (median ± IQR) outliers per K-SICM sample, which was not found to be significantly different (p = 0.368) (Fig. 4). Additionally, there was no significant difference between the culture medium groups when the numbers of hypomethylation outliers (P = 0.86) and hypermethylation outliers (P = 0.238) were analysed separately (Fig. 4). Outlier burden, i.e. the total number of outliers per sample, was not significantly associated with age at sample collection or weight at birth or follow-up. On the other hand, technical features, specifically leucocyte proportion and sample plate, were significantly associated with outlier burden (Supplementary Table SII). An association between the number of samples and the presence of common diseases (atopy, autism, urological problems) was not tested statistically, but one-third of the children with a diagnosis (8 out of 24) had a very high number of methylation outliers (more than the upper quartile).

Using iEVORA we identified 101 differentially variable CpG sites between the two groups, of which 94 sites were more variable in the G3 group than the K-SICM group. While 20 of the sites were unannotated, the remainder belonged to 80 unique genes. Two genes,
and PDZRN3, contained two differentially variable sites. Additionally, two of the sites were within imprinted genes, namely KCNQ1 and APBA1 (Supplementary Table SIII, sheet 1). The differentially methylated sites identified by iEVORA were not found to be significantly enriched (FDR < 0.05) in any pathways or ontologies according to the KEGG and GO analyses (Supplementary Table SIII, sheets 2–3).

Comparison of DNA methylation levels of ART children compared to naturally conceived individuals

All the saliva methylomes (n = 50) available from the FLEHS cohort passed our QC criteria (Supplementary Fig. S1) and the characteristics of these individuals are summarized in Supplementary Table SIV. Of note is that saliva samples were collected from significantly older naturally conceived children than ART children (ART age: 9.5 ± 0.3, naturally conceived age: 10.6 ± 0.3, mean ± SD, P < 0.001, Welch’s two-sample t-test) with a higher incidence of asthma (12% vs 6% in the ART and naturally conceived cohorts, respectively). PCA showed significant separation of the ART and naturally conceived samples on PCs 1, 3, 4 and 5 (Supplementary Fig. S4A and B). Seventy-two percent of CpG sites were found to be significantly differentially methylated (adjusted P-value < 0.05) with a strong predominance of sites that were hypomethylated in the ART samples compared to the naturally conceived samples (Supplementary Fig. S4C and D). Such marked methylation differences in phenotypically similar individuals suggest systematic differences in the data that could be due to technical or cohort differences, such as the array type (EPIC or 450K) used, differences induced by the array scanner, laboratory environmental factors or minor pipetting or laboratory protocol deviations. When the sample groups of interest are processed separately, these features align perfectly with the groups of interest and therefore cannot be differentiated from meaningful biological variation. Further analyses were therefore not carried out.

Discussion

Here, we present the largest study to date of the saliva methylome of ART children that underwent embryo culture in different media, namely G3 or K-SICM. At age 9, we have found no significant DNA methylation differences between the culture medium groups when considering individual CpG sites, multiple CpG sites aggregated into genomic regions or when conducting a targeted analysis of (sites within) imprinted genes or birth weight-associated CpG sites. The number of methylation outliers per sample was found to be comparable between the culture medium groups and was only significantly associated with

| Characteristic                                      | G3 (n = 57) | K-SICM (n = 49) | P-value |
|----------------------------------------------------|-------------|----------------|---------|
| Characteristics of the child                       |             |                |         |
| Sex (female)                                       | 29 (51)     | 26 (53)        | 0.977   |
| Age at sample collection (years)                   | 9.5 ± 0.3   | 9.5 ± 0.3      | 0.802   |
| Weight (kg)                                        | 34.1 ± 6.7  | 31.8 ± 5.5     | 0.054   |
| Height (cm)                                        | 139.0 ± 5.9 | 138.5 ± 7.5    | 0.706   |
| Medical diagnoses                                  |             |                |         |
| Urological                                          | 3 (5)       | 2 (4)          | 1.000   |
| Asthma/allergy                                     | 9 (16)      | 4 (8)          | 0.370   |
| Autism                                             | 4 (7)       | 2 (4)          | 0.818   |
| Pregnancy characteristics                          |             |                |         |
| Maternal age (years)                               | 32.8 ± 3.4  | 33.0 ± 3.4     | 0.772   |
| Paternal age (years)                               | 35.8 ± 4.5  | 35.6 ± 4.7     | 0.827   |
| Gestational age                                     | 39.7 ± 1.5  | 39.5 ± 2.2     | 0.583   |
| Birth weight (g)                                   | 3425.4 ± 486.4 | 3308.6 ± 537.0 | 0.247   |
| Fertility treatment                                |             |                |         |
| Fertilization method                               |             |                | 0.985   |
| IVF                                                | 18 (32)     | 17 (35)        |         |
| ICSI                                               | 39 (68)     | 32 (65)        |         |
| Treatment indication                               |             |                | 0.985   |
| Unknown                                            | 12 (21)     | 11 (22)        |         |
| Male factor                                        | 38 (67)     | 32 (65)        |         |
| Female factor                                      | 7 (12)      | 6 (12)         |         |

Continuous variables shown as mean ± SD, categorical variables shown as n (%). Maternal and paternal age at time of ovum pick-up is shown.

Table I Characteristics of the child, pregnancy and fertility treatment.
Interestingly, iEVORA identified that the majority of differentially variable sites between the G3 and K-SICM groups were more variable in the G3 group.

Our findings are consistent with those from the only other ART culture medium cohort in which the saliva methylome was investigated during childhood (ages 7–8) (Barberet et al., 2021; Ducreux et al., 2021). Although the ART children included in that study underwent embryo culture in other media than those compared in the present work, no significant DNA methylation differences were found at any of the sites or regions (Barberet et al., 2021; Ducreux et al., 2021).

**Figure 3.** Analysis of systematic methylation differences between G3 and K-SICM children: differentially methylated positions and regions. Volcano plots showing differential methylation between G3 and Sydney IVF cleavage medium (K-SICM) children where the grey dots represent all individual cytosine-guanine dinucleotide (CpG) sites (A, B) or multiple CpG sites aggregated into genomic regions, namely genes (C), promoters (D), CpG islands (E). Imprinted genes (C) and sites within them (A) are highlighted in purple while CpG sites associated with birth weight are shown in green (B). Unadjusted P-values are shown. These were generated using M-values for mixed effects linear models, while the group mean differences are shown as beta values for interpretability. No significant differences were found between the culture medium groups (FDR adjusted P-value < 0.1) (see also Supplementary Fig. S2).
The group mean differences that they observed between their culture medium groups were also similar to those observed in our cohort (Barberet et al., 2021). Due to the notably larger sample size of our study, we anticipate that we would have sufficient power (0.8) to detect small methylation differences compared to what was possible in the previously described cohort. The results from the current study were also comparable to those of the methylation studies conducted on placenta or umbilical cord blood samples collected from ART neonates cultured in G5 or HTF media (Mulder et al., 2020; Koeck et al., 2022). In addition to finding no significant DMPs or DMRs, these studies also reported that there was no difference in the number of outliers per sample between the culture medium groups (Mulder et al., 2020; Koeck et al., 2022).

Interestingly, in the recent umbilical cord blood methylation study (Koeck et al., 2022), 90% of the differentially variable sites identified using iEVORA were more variable in the G5 group than in the HTF culture medium group. Although the effect was less striking once data from neonates that had experienced pregnancy complications had been excluded, 56% of the identified sites were still more variable in the G5 group. In both studies, the culture medium from Vitrolife was associated with a higher number of more variable CpG sites amongst the differentially variable sites (Koeck et al., 2022). Several explanations have been proposed for the relevance of differentially variable sites. For instance, low methylation variance could result from environmental selection pressures that only facilitate the survival of individuals of certain methylation signatures (Tobi et al., 2018). Alternatively, in oncological samples, sites with differential variability in pre-cancerous tissues are commonly found to be differentially methylated in tumours, when compared with healthy tissues, thus suggesting that these epimutations play a role in disease pathogenesis (Teschendorff et al., 2016). Therefore, in our study, less variability in DNA methylation, as seen in the K-SiCM group, could indicate a greater environmental selection pressure which would be in concordance with the observed lower implantation and pregnancy rates in this group (Dumoulin et al., 2010). However, these sites with differentially variable methylation levels could also be indicative of (cardiometabolic) diseases that certain individuals may develop in later life, which would need to be investigated with further longitudinal follow-up studies of these individuals.

The main aim of the study was to compare the methylomes of ART children who had undergone embryo culture in different media, therefore saliva samples from naturally conceived children were not collected and processed alongside the ART samples. Although a comparison with children from the FLEHS cohort was attempted, the results indicate a large contribution of technical or cohort differences that cannot be corrected in samples that are not processed simultaneously. As such, we recommend that future studies analysing the methylome of ART offspring also collect samples from naturally conceived individuals for simultaneous processing.

The strengths of our study lie in the use of a methylome analysis method that captures methylation status across the whole genome and the sample size which is the largest of any methylome study of ART children in the context of a culture medium study. Nonetheless, to reach sufficient power (0.8) to detect methylation differences of <10% between the culture medium groups a bigger sample size is likely required (Tsai and Bell, 2015; Saffari et al., 2018). On the other hand, it is unclear whether such small differences would represent clinically meaningful differences. Since methylation is a binary state (methylated or unmethylated) a 5% methylation increase in any given individual signifies that 5% more of their cells are methylated at the given position. Whether this contributes to pathology or phenotypic variance remains to be determined.

Overall, the findings of this study are reassuring. Although phenotypic differences are observed, even at the age of 9, between ART offspring that had undergone embryo culture in G3 or K-SiCM medium, there is no evidence that the epigenome of these individuals differs greatly. As the epigenome retains its plasticity throughout life it is possible that epigenetic dysregulation experienced during the pre-implantation period is no longer evident in ART offspring at the age of 9. Further research, profiling the epigenome of ART embryos, is required to understand how the ART culture environment modulates the epigenome during the period of in vitro culture and the significance of sites with differentially variable methylation for later life (disease) outcomes should be evaluated with longitudinal studies.

**Supplementary data**

Supplementary data are available at *Human Reproduction* online.
**Data availability**

The raw and processed array data from this study have been uploaded to the Gene Expression Omnibus (GEO) database and are available under the accession number GSE196432 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE196432). The raw array data from the FLEHS study are available within the GEO database under the accession number GSE110128.

**Acknowledgements**

We thank the ART couples who have agreed for their children to participate in this study. We thank the FLEHS Supervisory Board for the provision of data. The FLEHS studies were commissioned, financed and steered by the Flemish Government (Department of Economy, Science and Innovations, Agency for Care and Health and Department of Environment).

**Authors’ roles**

R.M.K., A.P.A.v.M. and M.Z.E. study design and conception. H.Z., R.V.G. and J.C.M.D. were involved in sample collection for the ART cohort. A.P.A.v.M. and F.B. carried out the lab work for the ART samples. S.R. and S.L. were involved in sample collection and processing for the naturally conceived cohorts. R.M.K., F.B., J.T., A.P.A.v.M. and M.Z.E were involved in the data analysis and interpretation. R.M.K. wrote the first draft of the manuscript. M.G., A.P.A.v.M. and M.Z.E. contributed to the writing of the manuscript. All authors provided textual comments and approved the manuscript. H.B., A.P.A.v.M. and M.Z.E supervised the study.

**Funding**

This study was funded by March of Dimes (6-FY13-153). Additionally, it was further supported by EVA (Erfelijkheid Voortplanting & Aanleg) specialty programme (grant no. KP111513) from Maastricht University Medical Centre (MUMC+) and the Horizon 2020 innovation (ERIN) (grant no. EU952516) from the European Commission. The FLEHS dataset has been generated by the Flemish Center of Expertise on Environment and Health (FLEHS 2016–2020), funded by the Environment, Nature and Energy Department of the Flemish government. The views expressed in this manuscript are those of the author(s) and are not necessarily endorsed by the Flemish government.

**Conflict of interest**

The authors do not report any conflicts of interest relevant to this study.

**References**

Adamson GD, Dyer S, Chambers GM, Ishihara O, Mansour R, Banker M, de Mouzon J, Zegers-Hochschild F. ICMART preliminary world report 2015. *Hum Reprod* 2019;**34**(Suppl_1):i65.

Barberet J, Binquet C, Guilleman M, Doukani A, Choux C, Bruno C, Bourredjem A, Chapusot C, Bourc‘his D, Duffourd Y et al. Do assisted reproductive technologies and *in vitro* embryo culture influence the epigenetic control of imprinted genes and transposable elements in children? *Hum Reprod* 2021;**36**:479–492.

Barberet J, Ducreux B, Guilleman M, Simon E, Bruno C, Fauche P. DNA methylation profiles after ART during human lifespan: a systematic review and meta-analysis. *Hum Reprod Update* 2022;**28**:629–655.

Bay B, Lyngsø J, Hohwü L, Kesmodel US. Childhood growth of singletons conceived following *in vitro* fertilisation or intracytoplasmic sperm injection: a systematic review and meta-analysis. *BJOG* 2019;**126**:158–166.

Benjamini Y, Hochberg Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. *J Roy Statist Soc* 1995;**57**:289–300.

Berntsen S, Söderström-Anttila V, Wennerholm U-B, Laivuori H, Loft A, Oldered NB, Romundstad LB, Bergh C, Pinborg A. The health of children conceived by ART: ‘the chicken or the egg?’. *Hum Reprod Update* 2019;**25**:137–158.

Bouillon C, Léandi R, Desch L, Ernst A, Bruno C, Cerf C, Chiron A, Souchay C, Burguet A, Jimenez C et al. Does embryo culture medium influence the health and development of children born after *in vitro* fertilization? *PLoS One* 2016;**11**:e0150857.

Ceelen M, van Weissenbruch MM, Prein J, Smit JJ, Vermeiden JP, Spreeuwenberg M, van Leeuwen FE, Delemarre-van de Waal HA. Growth during infancy and early childhood in relation to blood pressure and body fat measures at age 8-18 years of IVF children and spontaneously conceived controls born to subfertile parents. *Hum Reprod* 2009;**24**:2788–2795.

DeAngelis AM, Martini AE, Owen CM. Assisted reproductive technology and epigenetics. *Semin Reprod Med* 2018;**36**:221–232.

Donjacour A, Liu X, Lin W, Simbulan R, Rinaudo PF. *In vitro* fertilization affects growth and glucose metabolism in a sex-specific manner in an outbred mouse model. *Biol Reprod* 2014;**90**:80.

Du P, Zhang X, Huang CC, Jafari N, Kibbe WA, Hou L, Lin SM. Comparison of Beta-value and M-value methods for quantifying methylation levels by microarray analysis. *BMC Bioinformatics* 2010;**11**:587.

Ducreux B, Frappier J, Bruno C, Doukani A, Guilleman M, Simon E, Martinaud A, Bourc‘his D, Barberet J, Fauche P. Genome-wide analysis of DNA methylation in buccal cells of children conceived through IVF and ICSI. *Genes (Basel)* 2021;**12**:1912.

Dumoulin JC, Land JA, Van Montfoort AP, Nelissen EC, Coonen E, Derhaag JG, Schreurs IL, Dunselman GA, Kester AD, Geraeds JP et al. Effect of *in vitro* culture of human embryos on birthweight of newborns. *Hum Reprod* 2010;**25**:605–612.

El Hajj N, Haaf T. Epigenetic disturbances in *in vitro* cultured gametes and embryos: implications for human assisted reproduction. *Fertil Steril* 2013;**99**:632–641.

Felix JF, Cecil CAM. Population DNA methylation studies in the Developmental Origins of Health and Disease (DOHaD) framework. *J Dev Orig Health Dis* 2019;**10**:306–313.

Fernández-Gonzalez R, Moreira P, Bilbao A, Jiménez A, Pérez-Crespo M, Ramírez MA, Rodríguez De Fonseca F, Pintado B, Gutiérrez-Adán A. Long-term effect of *in vitro* culture of mouse embryos with serum on mRNA expression of imprinting genes, development, and behavior. *Proc Natl Acad Sci USA* 2004;**101**:5880–5885.
Gentilini D, Garagnani P, Pisoni S, Bacalini MG, Calzari L, Mari D, Vitale G, Franceschi C, Di Blasio AM. Stochastic epigenetic mutations (DNA methylation) increase exponentially in human aging and correlate with X chromosome inactivation skewing in females. *Aging (Albany NY)* 2015;7:568–578.

Ginjala V. Gene imprinting gateway. 2009. https://genomebiology.biomedcentral.com/articles/10.1186-gb-2001-2-8-reports (12 December 2021, date last accessed).

Gu Z, Eils R, Schlesner M. Complex heatmaps reveal patterns and correlations in multidimensional genomic data. *Bioinformatics* 2016;32:2847–2849.

Guo XY, Liu XM, Jin L, Wang TT, Ullah K, Sheng JZ, Huang HF. Cardiovascular and metabolic profiles of offspring conceived by assisted reproductive technologies: a systematic review and meta-analysis. *Fertil Steril* 2017;107:622–631.e5.

Hann M, Roberts SA, D’Souza SW, Clayton P, Macklon N, Brison DR. The growth of assisted reproductive treatment-conceived children from birth to 5 years: a national cohort study. *BMC Med* 2018;16:224.

Hanna CW, Demond H, Kelsey G. Epigenetic regulation in development: is the mouse a good model for the human? *Hum Reprod Update* 2018;24:556–576.

Hashimoto S, Nishihara T, Murata Y, Oku H, Nakaoka Y, Fukuda A, Morimoto Y. Medium without ammonium accumulation supports the developmental competence of human embryos. *J Reprod Dev* 2008;54:370–374.

Heiss JA, Just AC. Identifying mislabeled and contaminated DNA methylation microarray data: an extended quality control toolset with examples from GEO. *Clin Epigenetics* 2018;10:73.

Hoffman GE, Schadt EE. variancePartition: interpreting drivers of variation in complex gene expression studies. *BMC Bioinformatics* 2016;17:483.

Houseman EA, Molitor J, Marsit CJ. Reference-free cell mixture adjustments in analysis of DNA methylation data. *Bioinformatics* 2014;30:1431–1439.

Häberg SE, Page CM, Lee Y, Nustad HE, Magnus MC, Hafstrom KL, Carlsson E, Denault WRP, Bohlin J, Jugessur A et al. DNA methylation in newborns conceived by assisted reproductive technology. *Nat Commun* 2022;13:1896.

Jung CH, Park DJ, Georgeson P, Mahmood K, Milne RL, Southey MC, Pope BJ. sEst: accurate sex-estimation and abnormality detection in methylation microarray data. *Int J Mol Sci* 2018;19:3172–3183.

Kleijkers SH, van Montfoort AP, Bekers O, Coonen E, Derhaag JG, Evers JL, Dumoulin JC. Ammonium accumulation in commercially available embryo culture media and protein supplements during storage at 2-8°C and during incubation at 37°C. *Hum Reprod* 2016;31:1192–1199.

Kleijkers SH, van Montfoort AP, Smits LJ, Viechtbauer W, Roseboom TJ, Nelissen EC, Coonen E, Derhaag JG, Bastings L, Schreurs IE et al. IVF culture medium affects post-natal weight in humans during the first 2 years of life. *Hum Reprod* 2014;29:661–669.

Koeck RM, Busato F, Tost J, Consten D, van Echten-Arends J, Mastenbroek S, Wurth Y, Remy S, Langie S, Nawrot TS et al. Methylome-wide analysis of IVF neonates that underwent embryo culture in different media revealed no significant differences. *NPJ Genom Med* 2022;7:39.

Koncevicius K. matrixTests: Fast Statistical Hypothesis Tests on Rows and Columns of Matrices. 2020. https://cran.r-project.org/web/packages/matrixTests/index.html (23 February 2021, date last accessed).

Küppers LK, Monnerau C, Sharp GC, Yousefi P, Salas LA, Ghanotou A, Page CM, Reese SE, Wilcox AJ, Czamara D et al. Meta-analysis of epigenome-wide association studies in neonates reveals widespread differential DNA methylation associated with birthweight. *Nat Commun* 2019;10:1893.

Langie SAS, Moises M, Szarc Vel Szc K, Van Der Plas E, Koppen G, De Prins S, Louwies T, Nelen V, Van Camp G, Lambrechts D et al. GLI2 promoter hypermethylation in saliva of children with a respiratory allergy. *Clin Epigenet* 2018;10:50.

Li L, Guo F, Gao Y, Ren Y, Yuan P, Yan L, Li R, Lian Y, Li J, Hu B et al. Single-cell multi-omics sequencing of human early embryos. *Nat Cell Biol* 2018;20:847–858.

Maksimovic J, Gordon L, Oshlack A. SWAN: subset-quantile within array normalization for illumina infinium HumanMethylation450 BeadChips. *Genome Biol* 2012;13:R44.

Mantikou E, Youssef MA, van Wely M, van der Veen F, Al-Inany HG, Repping S, Mastenbroek S. Embryo culture media and IVF/ICSI success rates: a systematic review. *Hum Reprod Update* 2013;19:210–220.

Melamed N, Choufani S, Wilkins-Haug LE, Koren G, Wexberg R. Comparison of genome-wide and gene-specific DNA methylation between ART and naturally conceived pregnancies. *Epigenetics* 2015;10:474–483.

Middleton LYM, Dou J, Fisher J, Heiss JA, Nguyen VK, Just AC, Faul J, Ware EB, Mitchell C, Colacino JA et al. Saliva cell type DNA methylation reference panel for epidemiological studies in children. *Epigenetics* 2022;17:161–177.

Morbeck DE, Baumann NA, Oglesbee D. Composition of single-step media used for human embryo culture. *Fertil Steril* 2017;107:1055–1060.e1.

Morbeck DE, Krisher RL, Herrick JR, Baumann NA, Matern D, Moyer T. Composition of commercial media used for human embryo culture. *Fertil Steril* 2014;102:759–766.e9.

Mulder CL, Wattimury TM, Jongejan A, de Winter-Korver CM, van Daalen SKM, Struijk RB, Borgman SCM, Wurth Y, Consten D, van Echten-Arends J et al. Comparison of DNA methylation patterns of parentally imprinted genes in placenta derived from IVF conceptions in two different culture media. *Hum Reprod* 2020;35:516–528.

Müller F, Scherer M, Assenov Y, Lutsik P, Walter J, Lengauer T, Bock C. RnBeads 2.0: comprehensive analysis of DNA methylation data. *Genome Biol* 2019;20:55.

Nelissen EC, Van Montfoort AP, Coonen E, Derhaag JG, Geraedts JP, Smits LJ, Land JA, Evers JL, Dumoulin JC. Further evidence that culture media affect perinatal outcome: findings after transfer of fresh and cryopreserved embryos. *Hum Reprod* 2012;27:1966–1976.

Nelissen EC, Van Montfoort AP, Smits LJ, Meneheere PP, Evers JL, Coonen E, Derhaag JG, Peeters LL, Coumans AB, Dumoulin JC. IVF culture medium affects human intrauterine growth as early as...
the second trimester of pregnancy. *Hum Reprod* 2013;28:2067–2074.

Novakovic B, Lewis S, Halliday J, Kennedy J, Burgner DP, Czajko A, Kim B, Sexton-Oates A, Juonala M, Hammarberg K et al. Assisted reproductive technologies are associated with limited epigenetic variation at birth that largely resolves by adulthood. *Nat Commun* 2019;10:3922.

Penova-Veselinovic B, Melton PE, Huang RC, Yovich JL, Burton P, Wijs LA, Hart RJ. DNA methylation patterns within whole blood of adolescents born from assisted reproductive technology are not different from adolescents born from natural conception. *Hum Reprod* 2021;36:2035–2049.

Phipson B, Maksimovic J, Oshlack A. missMethyl: an R package for analyzing data from Illumina's HumanMethylation450 platform. *Bioinformatics* 2016;32:286–288.

R Core Team. *R: A Language and Environment for Statistical Computing*. Vienna, Austria: R Foundation for Statistical Computing, 2021. https://www.R-project.org/.

Ritchie ME, Phipson B, Wu D, Hu Y, Law CW, Shi W, Smyth GK. limma powers differential analyses for RNA-sequencing and microarray studies. *Nucleic Acids Res* 2015;43:e47.

Saffari A, Silver MJ, Zavattari P, Moi L, Columbano A, Meaburn EL, Dudbridge F. Estimation of a significance threshold for epigenome-wide association studies. *Genet Epidemiol* 2018;42:20–33.

Sunde A, Brison D, Dumoulin J, Harper J, Lundin K, Magli MC, Van den Abbeel E, Veiga A. Time to take human embryo culture seriously. *Hum Reprod* 2016;31:2174–2182.

Teschendorff AE, Jones A, Widschwendter M. Stochastic epigenetic outliers can define field defects in cancer. *BMC Bioinformatics* 2016;17:178.

Tobi EW, van den Heuvel J, Zwaan BJ, Lumey LH, Heijmans BT, Uller T. Selective survival of embryos can explain DNA methylation signatures of adverse prenatal environments. *Cell Rep* 2018;25:2660–2667.e4.

Tsai PC, Bell JT. Power and sample size estimation for epigenome-wide association scans to detect differential DNA methylation. *Int J Epidemiol* 2015;44:1429–1441.

Van Den Heuvel R, Den Hond E, Colles A, Nelen V, Van Campenhout K, Schoeters G. Biobank@VITO: biobanking the general population in Flanders. *Front Med (Lausanne)* 2020;7:37.

Velazquez MA, Sheth B, Smith SJ, Eckert JJ, Osmond C, Fleming TP. Insulin and branched-chain amino acid depletion during mouse pre-implantation embryo culture programmes body weight gain and raised blood pressure during early postnatal life. *Biochim Biophys Acta Mol Basis Dis* 2018;1864:590–600.

Virant-Klin I, Tomazevic T, Vratcnic-Bokal E, Vogler A, Krsnik M, Meden-Vrtovec H. Increased ammonium in culture medium reduces the development of human embryos to the blastocyst stage. *Fertil Steril* 2006;85:526–528.

Wadhwa PD, Buss C, Entringer S, Swanson JM. Developmental origins of health and disease: brief history of the approach and current focus on epigenetic mechanisms. *Semin Reprod Med* 2009;27:358–368.

Yeung EH, Mendola P, Sundaram R, Zeng X, Guan W, Tsai MY, Robinson SL, Stern JE, Ghassabian A, Lawrence D et al. Conception by fertility treatment and offspring deoxyribonucleic acid methylation. *Fertil Steril* 2021;116:493–504.

Zandstra H, Brentjens L, Spauwen BH, Bons JAP, Mulder AL, Smits LJM, van der Hoeven M, van Golde RJT, Evers JLM et al. Association of culture medium with growth, weight and cardiovascular development of IVF children at the age of 9 years. *Hum Reprod* 2018a;33:1645–1656.

Zandstra H, Smits LJM, van Kuijk SMJ, van Golde RJT, Evers JLMH, Dumoulin JCM, van Montfoort APA. No effect of IVF culture medium on cognitive development of 9-year-old children. *Hum Reprod* 2018b;33:690–698.

Zandstra H, Van Montfoort AP, Dumoulin JC. Does the type of culture medium used influence birthweight of children born after IVF? *Hum Reprod* 2015;30:530–542.