**Environmentally sensitive hotspots in the methylome of the early human embryo**

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**Abstract**

In humans, DNA methylation marks inherited from gametes are largely erased following fertilisation, prior to construction of the embryonic methylome. Exploiting a natural experiment of seasonal variation including changes in diet and nutritional status in rural Gambia, we analysed three datasets covering two independent child cohorts and identified 259 CpGs showing consistent associations between season of conception (SoC) and DNA methylation. SoC effects were most apparent in early infancy, with evidence of attenuation by mid-childhood. SoC-associated CpGs were enriched for metastable epialleles, parent-of-origin specific methylation and germline DMRs, supporting a periconceptional environmental influence. Many SoC-associated CpGs overlapped enhancers or sites of active transcription in H1 ESCs and fetal tissues. Half were influenced but not determined by measured genetic variants that were independent of SoC. Environmental ‘hotspots’ providing a record of environmental influence at periconception constitute a valuable resource for investigating epigenetic mechanisms linking early exposures to lifelong health and disease.

**Introduction**

DNA methylation (DNAm) plays an important role in a diverse range of epigenetically regulated processes in mammals including cell differentiation, X-chromosome inactivation, genomic imprinting and the silencing of transposable elements1. DNAm can influence gene expression and can in turn be influenced by molecular processes including differential action of methyltransferases and transcription factor binding2,3.

The human methylome is extensively remodelled in the very early embryo when parental gametic methylation marks are largely erased before acquisition of lineage and tissue‑specific marks at implantation, gastrulation and beyond4. The days following conception may therefore offer a window of heightened sensitivity to external environmental influences, potentially stretching back to the period before conception coinciding with late maturation of oocytes and spermatozoa at loci that (partially) evade early embryonic reprogramming5.

The effects of early exposures on the mammalian methylome have been widely studied in animal models and periconceptional and early gestational factors including maternal folate and exposure to famine have been associated with DNAm changes in humans6–8. However, causal pathways are difficult to elucidate in human observational studies, and even randomised experimental designs are prone to confounding due to reverse causation from exposure-related effects9.

Here we address these limitations by exploiting a natural experiment in rural Gambia where conceptions occur against a background of repeating annual patterns of dry (‘harvest’) and rainy (‘hungry’) seasons with accompanying significant changes in energy balance, diet composition, nutrient status and rates of infection10,11. We assess the influence of seasonality on DNAm in two Gambian child cohorts12,13, one with longitudinal data, enabling robust identification of loci showing consistent effects at the ages of 2 years and in mid-childhood (Fig. 1). Through prospective study designs we capture conceptions throughout the year and, in contrast to previous analyses in this population14–16, we use statistical models that make no prior assumptions about specific seasonal windows driving DNAm changes in offspring.

We probe potential connections between season of conception (SoC)-associated loci and early embryonic events by leveraging published data on loci with evidence for the establishment of variable methylation states in the early embryo that persist in post-gastrulation and postnatal tissues; namely loci demonstrating systemic interindividual variation (SIV)17,18 and/or epigenetic supersimilarity (ESS)18 (Fig. 1). These loci bear the hallmarks of metastable epialleles (MEs), loci with methylation states that vary between individuals that were first identified in isogenic mice. MEs exhibit stable patterns of SIV indicating stochastic establishment of methylation marks prior to gastrulation when tissue differentiation begins19, and several MEs have been shown to be sensitive to periconceptional nutrition in mice20. These loci thus serve as useful tools for studying the effects of early environment on DNAm by enabling the use of accessible tissues (such as blood) that can serve as a proxy for systemic (cross-tissue) methylation, and by pinpointing the window of exposure to the periconceptional period21. We also investigate links with transposable elements and transcription factors associated with the establishment of methylation states in the early embryo, and assess the influence of genetic variation and gene-environment interactions. Finally, by comparing our results with public DNAm data obtained from sperm, oocytes and multi-stage human embryos, we investigate links between SoC-associated loci, histone marks, gametic and parent-of-origin specific methylation, and the establishment of DNAm states in early embryonic development.

Our identification of hotspots in the postnatal methylome that retain a record of environmental conditions during gametic maturation and/or in the very early embryo, provides a valuable resource for the investigation of epigenetic mechanisms linking early-life nutritional and other exposures to lifelong health and disease.

**Results**

*Identification of Gambian season of conception associated CpGs*

Key characteristics of DNAm datasets from the two Gambian cohorts analysed in this study are provided in Table 1. DNAm differences associated with season of conception are potentially confounded by season of sample collection effects in the ENID 2yr dataset (n=233) since all samples were collected at age 24 months (Fig. 2A top). This is not the case in the older EMPHASIS cohort (n=289; age 7-9 yr) where all samples were collected in the Gambian dry season (Fig. 2A bottom). To account for the potential influence of season of collection effects, we therefore compared year-round DNAm signatures across ENID (2yr) and EMPHASIS datasets by focussing on 391,814 autosomal CpGs (‘array background’) intersecting the Illumina HM450 and EPIC arrays used to measure DNAm in each dataset (Table 2). We modelled the effect of date of conception on DNAm using Fourier (or ‘cosinor’) regression22 which makes no prior assumptions about specific seasonal windows that might drive DNAm changes in offspring (see Methods).

We began by identifying 768 SoC-associated CpGs showing significant seasonal variation in 2-year olds from the ENID cohort with a false discovery rate (FDR)<5% (Supplementary File 1a; Methods). Fourier regression models revealed a heterogeneous distribution of year-round methylation peaks and nadirs at these loci (Fig. 2B). SoC-associated loci were highly enriched for loci exhibiting SIV/ESS previously identified in multi-tissue screens in adult Caucasians17,18, hereafter named ‘MEs’ for short (Table 3; Fig. 2B, 26 ME CpGs marked in red; enrichment p = 2.5x10‑14). More than twice as many of these loci were within 100bp of a putative ME (n=56; Supplementary File 1a). All identified loci showed increased seasonal amplitudes, defined as the distance between methylation peak and nadir, compared to matched and random controls (Supplementary File 1b; see Table 2 and Methods for justification and further details on selection of controls). Loci with the largest amplitudes tended to show increased methylation in conceptions in the Gambian rainy season (Fig. 2D left) in line with our previous observations in this population14,15.

Next, we analysed SoC effects at these 768 loci in 7-9-year olds from the EMPHASIS cohort. Mean methylation at SoC-associated loci was strongly correlated across cohorts (Appendix 1-Fig. 1), and we found evidence of a similar effect of increased methylation in conceptions in the Gambian rainy season (Fig. 2D right). 61 loci (including three ME CpGs) were also significantly associated with SoC (FDR<5%) in the older cohort (Fig. 2D right). Notably, the date of conception at methylation maximum was highly correlated across these two independent and different aged cohorts (Fig. 2C left; Spearman rho=0.7, p=1.0x10-8). No significant correlation was observed at matched controls with similar methylation distributions to SoC-associated loci (Fig. 2C right).

Given the strikingly similar seasonal patterns across the two cohorts, we next investigated reasons for the smaller number of SoC-associated loci (FDR<5%) in the older (EMPHASIS) cohort. Focussing on the 768 SoC-associated loci in the ENID cohort, we found significant and relatively large effect size (SoC amplitude) decreases in the older cohort, with SoC-effect attenuation most marked at loci that are also putative MEs (4.1% median methylation decrease, Wilcoxon p=2.8x10-6; non-ME CpGs: 1.9%, Wilcoxon p=1.5x10-77; Fig. 2E; Supplementary File 1c). Corresponding SoC amplitude changes in matched and random controls were much smaller (0.2%, p=2.0x10-8 and 0.05%, p=0.02 respectively; Supplementary File 1c).

Noting that loci with larger SoC amplitudes showed a more consistent pattern of seasonal variation, both within and between cohorts (Figs. 2C & 2D), and reasoning that even transient molecular biomarkers of periconceptional environment in postnatal tissues could have biological significance, we focussed on 259 SoC ‘hotspots’ or SoC-CpGs with FDR<5% and SoC amplitude ≥ 4% identified in the ENID 2yr cohort (Fig. 2D left, loci to the right of the dashed line; Table 2; Supplementary File 1d). Sensitivity analysis confirmed that SoC effect estimates at SoC-CpGs were robust to different modelling strategies with respect to estimated blood cell composition and batch effects (see Methods).

We tested the hypothesis that SoC amplitudes at SoC-CpGs decrease with age by generating EPIC array methylation data on blood samples in a subset of n=138 individuals from the ENID cohort at 5-7yrs (Table 1). This revealed a strongly consistent methylation signature at n=157 replicating SoC-CpGs (Fig. 3A; Supplementary File 1d; Spearman rho=0.7, p<2.2x10-16). This analysis also provided further evidence of SoC effect attenuation with age at SoC-CpGs (Fig. 3B; Supplementary File 1d; Wilcoxon signed rank test for difference in SoC amplitude p=6.7x10-12 for SoC-CpGs and p=1.0 for matched controls).). ENID longitudinal samples were collected in the rainy season (Appendix 1-Fig. 2), again exhibiting a different confounding structure with respect to season of sample collection compared to samples analysed at the earlier timepoint (ENID 2yr) and in EMPHASIS (Fig. 2A).

Critically, the date of conception at methylation maximum at SoC-CpGs was highly consistent across all three datasets analysed, with a distinct pattern of methylation maxima for conceptions falling within the August-September period, most markedly at putative MEs with independent evidence of establishment in the early embryo (Fig. 4A). The August to September period corresponds to the peak of the Gambian rainy season, a strong validation of our previous studies in babies and infants that focussed on conceptions at peak seasons only, with similar observations of higher methylation in conceptions at the peak of the Gambian rainy season compared to peak dry season14–16,18. Methylation minima fall within the February-April period, corresponding to the peak of the dry season (Appendix 1-Fig. 3).

Our observation of a remarkably similar season of conception signature across two cohorts and three datasets with different confounding structures with respect to season of sample collection, batch and biological variables (Supp Tables 16-18), combined with evidence from cross-cohort and longitudinal analyses of SoC-effect attenuation with age led us to conclude that SoC-CpGs act as robust sentinels of SoC-associated effects persisting at least until the age of 2 years.

*Properties of SoC-CpGs*

SoC-CpGs are distributed throughout the genome and cluster together in several regions (Appendix 1-Fig. 4). Noting that the number of clusters is relatively insensitive to the inter-CpG distance used to define them (Appendix 1-Fig. 5), we identified 56 distinct SoC-CpG clusters and 105 ‘singletons’ (SoC-CpGs with no close neighbours) using a maximum inter-CpG distance of 5kbp (Supplementary File 1e). With this definition, 59% of SoC-CpGs fell within clusters (Supplementary File 1f). Of note, SoC effect amplitudes and cross-cohort correlations were greater at SoC-CpGs falling within clusters than with singletons (Appendix 1-Fig. 6).

Several SoC-CpG clusters extend over more than 500bp, notably a cluster mapping to *IGF1R* which spans 872bp and covers 7 CpGs (Supplementary File 1e, Appendix 1-Fig. 7). All but one of these 7 CpGs were also significantly associated with SoC (FDR<5%) in the older EMPHASIS cohort (Supplementary File 1d).

Compared to array background, SoC-CpGs are intermediately methylated, most notably at putative MEs (Fig. 4B), and they tend to fall within CpG islands compared to array background and matched controls (Fig. 4C). SoC-CpGs are also highly enriched for MEs (21-fold enrichment, p=3.0x10‑23, cluster-adjusted: 17-fold, p=3.1x10-11; Supplementary File 1g; see Methods for further details on cluster-based adjustments). The number of SoC-CpGs directly overlapping previously identified MEs is small (n=24), although 49 SoC-CpGs (19%) fall within 100bp of a ME (Supplementary File 1d). Further investigation revealed that a large majority (n=19/24) of overlapping MEs were identified in our previous screen for SIV, with a smaller number exhibiting ESS (n=7/24; Supplementary File 1g). No MEs overlap methylation distribution-matched controls.

Pairwise methylation states are highly correlated at a large majority of SoC-CpGs in both cohorts, so that the same individuals tend to have relatively high or low methylation at multiple SoC-CpGs (Fig. 4D). Pairwise correlations are not driven by increased correlation within SoC-CpG clusters (Appendix 1-Fig. 8), and methylation at matched and random controls is uncorrelated, thus reducing the possibility that these correlations are driven by statistical artefacts.

Finally, SoC-CpG reliability is classified as ‘excellent’ (median ICC=0.76) using probe reliability estimates from a recent repeated measures study23. Reliability of matched control CpGs is classified as ‘good’ (median ICC=0.68) using the same method.

*Early stage embryo, gametic and parent-of-origin specific methylation*

Given the strong enrichment for MEs within the set of SoC-CpGs, we next analysed links to methylation changes in early stage human embryos, as we have done previously for putative MEs identified in a whole-genome bisulfite-seq (WGBS) multi-tissue screen17. We aligned our data with public ﻿reduced representation bisulfite-seq (RRBS) data from human IVF embryos4 and obtained informative methylation calls for 112,380 array background CpGs covered at ≥ 10x read depth in inner cell mass (ICM, pre-gastrulation) and/or embryonic liver (post-gastrulation) tissues. As previously noted at putative MEs17, we found a distinctive pattern of increased incidence of intermediate methylation states at SoC-CpGs in post-gastrulation embryonic liver tissue, strongly contrasting with a general trend of genome-wide hyper- and hypo-methylation at loci mapping to array background (Fig. 5A). A similar pattern of increased incidence of intermediate methylation states was observed at distribution-matched controls.

We previously observed consistent hypomethylation at MEs across all gametic and early embryonic developmental stages, most notably in sperm17. We tested the latter observation at SoC-CpGs by aligning our data with public sperm WGBS data24, restricting our analysis to 294,240 CpGs mapping to array background that were covered at ≥ 10x. SoC-CpGs tended to be hypomethylated in sperm, compared to loci mapping to matched control CpGs and array background respectively (Figs. 5B & 5C). Intermediate methylation states at SoC-CpGs were preserved in both Gambian cohorts irrespective of sperm methylation states, in contrast to array background CpGs where methylation distributions strongly reflected sperm hypomethylation status (ENID cohort: Fig. 4D left; EMPHASIS cohort Appendix 1-Fig. 9 left).

Our observation of an increased incidence of sperm hypomethylation at SoC-associated loci, together with existing evidence that imprinted genes may be sensitive to prenatal exposures16,25,26, prompted us to investigate a potential link between SoC-CpGs and parent-of-origin specific methylation (PofOm). A recent study used phased WGBS methylomes to identify regions of PofOm in 200 Icelanders27. We analysed 699 of these PofOm CpGs overlapping array background (Table 3) and observed strong enrichment for PofOm CpGs at SoC-CpGs and at all MEs on the array (18- and 15-fold enrichment, p=3.0x10-8 and 1.8x10-36 respectively; Supplementary File 1h; Fig. 5D right, PofOm CpGs marked as green triangles). No enrichment was observed at distribution-matched controls (Supplementary File 1h). PofOm enrichment at SoC-CpGs is driven by a large PofOm region spanning 6 CpGs on chr15 at *IGF1R* (Supplementary File 1d); along with two singleton PofOm SoC-CpGs, one on chr18 close to *PARD6G*, and the other in the Prader-Willi syndrome-associated imprinted region neighbouring *MAGEL2*, also on chr15. All of these loci have increased methylation in the rainy season, with SoC effect sizes (methylation amplitudes) ranging from 4.1-8.4% (median 6.1%; Supplementary File 1d).

Regions of PofOm detected in postnatal samples tend to be differentially methylated in gametes27, and may thus have evaded epigenetic reprogramming in the pre-implantation embryo25. We tested this directly by interrogating data from a whole-genome screen for germline differentially methylated regions (gDMRs) that persist to the blastocyst stage and beyond28. In this analysis, gDMRs were defined as contiguous 25-CpG regions that were hypomethylated (mean DNAm < 25%) in one gamete and hypermethylated (mean DNAm > 75%) in the other. We began by observing strong enrichment for oocyte (maternally methylated), but not sperm gDMRs, at all PofOm loci identified by Zink *et al*27 (Supplementary File 1h), confirming previous observations of an excess of PofOm loci that are methylated in oocytes only27. This enrichment was particularly strong for oocyte gDMRs (oo-gDMRs) persisting in placenta (Supplementary File 1h). We next analysed SoC-CpGs and MEs and again found evidence for strong enrichment of oocyte, but not sperm gDMRs at these loci (6.2-fold oo-gDMR enrichment, p=2.3x10-16 at SoC-CpGs; 2.9-fold, p=1.2x10-24 at MEs), including after adjustment for CpG clustering (Supplementary File 1h). Of note, 14% (36/259) of SoC-CpGs overlapped oo-gDMRs, in strong contrast to matched and random controls (Fig. 5C). These clustered into 19 distinct oo-gDMR regions (Supplementary File 1i) - more than six times the number identified as exhibiting PofOm by Zink *et al.* (3 regions; Supplementary File 1d).

A large majority of SoC-CpGs that are hypomethylated in sperm are not oo-gDMRs (i.e. they are not hypermethylated in oocytes) (Fig. 5C & 5D bottom right), suggesting that factors associated with regional sperm hypomethylation rather than differential gametic methylation may be a key driver of sensitivity to periconceptional environment at these loci.

*SoC-CpG overlap with predicted chromatin states*

We assessed the overlap of SoC-CpGs with predicted chromatin states generated from histone marks in various cell lines and tissues by the Roadmap Epigenomics Consortium29. Given our interest in methylation states associated with periconceptional environment that persist into early postnatal life, we focussed on data from H1 embryonic stem cells (ESCs) and fetal tissues (fetal brain, muscle and small intestine) derived from all 3 germ layers, described as having the ‘highest quality’ epigenomes (see Fig. 2 in Kundaje et al29). Around half of all SoC-CpGs overlapped sites with predicted transcriptional or regulatory function, with relatively few overlapping constitutive heterochromatic regions (Fig. 6). Overlaps with specific histone marks in H1 ESCs are given in Appendix 1-Fig. 10. As expected, given predicted chromatin states, many show a predominance of overlapping H3K4me1 and H3K27me3 marks and combinations thereof, suggestive of active or poised enhancers.

*Enrichment of transposable elements and transcription factors associated with genomic imprinting*

Variable methylation states at MEs are associated with transposable elements (TEs) in murine models30,31, and we have previously observed enrichment for proximity to two classes of endogenous retroviruses, ERV1 and ERVK, at putative human MEs16,17. Here we found evidence that SoC-CpGs are enriched for proximity to ERV1 (Fig. 7A top) but not ERVK retroviral elements (Appendix 1-Fig. 11; Table 3).

Enrichment for PofOm and gDMRs at SoC-CpGs suggests a potential link to mechanisms implicated in the maintenance of PofOm and genomic imprinting in the early embryo. Our previous analysis of MEs identified from WGBS data found enrichment for proximal binding sites for 3 transcription factors (TFs: CTCF, ZFP57 and TRIM28) identified through ChIP-seq of embryonic stem and kidney cells that are linked to maintenance of PofOm at imprints17. Here we found evidence for enrichment of proximal ZFP57 binding sites within 2kbp of a SoC-CpG as previously observed at MEs, but we found no evidence for enrichment of proximal CTCF or TRIM28 binding sites in this array-based study (Fig. 7A bottom; Appendix 1-Fig. 11; Table 3).

*Influence of genotype*

Genetic variation is a major driver of inter-individual variation in DNAm via methylation quantitative trait loci (mQTL)32. We explored the influence of mQTL on SoC-CpGs in the EMPHASIS cohort for which we had genotype data on 284 individuals measured at >2.6M SNPs after imputation from the Illumina Global Screening Array (GSA) and subsequent LD-pruning (see Methods). The majority of mQTL effects occur in *cis*32*.* In order to maximise power we therefore adopted a two-step approach where we performed separate screens for mQTLs in *cis* (defined as SNPs within 1Mb of an associated CpG32) and *trans* (all others), and compared our findings at SoC-CpGs with matched and random control CpGs (Methods). Half of SoC-CpGs had one or more associated mQTL compared with 78% and 19% of matched and random controls respectively. 92% of SoC-CpG mQTL were in *cis* (Table 4).

We next compared methylation variance explained by significant mQTL using adjusted R2 values for all SoC-CpGs and controls with at least one genome wide significant mQTL (FDR<5%; n=130, 201 and 50 CpGs for SoC-CpGs, matched and random controls respectively; Table 4)*.* These values were compared to a baseline model that included the same set of covariates (principal components, age and sex) used in Fourier regression models for the main seasonality analysis, in order to account for potential differences in additional variance explained by other covariates and unmeasured factors (see Methods). There was no difference in additional variance explained by significant mQTL between SoC-CpGs and both sets of control CpGs (Fig. 7B; Table 4).

To assess the potential for genetic confounding of SoC-associated DNAm signals at SoC-CpGs with associated mQTL, we tested all SoC-CpG-mQTL for association with season of conception using an allelic model. After accounting for multiple testing, no significant SoC-mQTL associations were identified (Supplementary File 1j and Methods). Our observations that i) SoC-CpGs are distributed throughout the genome; ii) SoC-CpG mQTL occur primarily in *cis*; and iii) none are associated with season of conception; strongly suggest that SoC-methylation associations at SoC-CpGs are not confounded by genetic variation.

Finally, we searched for gene-environment (SoC) interaction (GxE) effects, again performing separate tests for SNPs in *cis* and *trans.* No GxE associations were identified after correcting for multiple testing (see Methods).

*Influence of genetic ancestry*

Eighty percent of the population of the Kiang West region of The Gambia from which the cohorts analysed in this study are drawn are of Mandinka ethnicity, with the majority of the remainder Fula33. This is evident from a genome-wide principal component analysis (PCA) of genetic variation in the EMPHASIS cohort, where we observed a distinct cluster of 16 individuals from a single village which is predominantly Fula (Appendix 1-Fig. 12). Individuals in the main ENID cohort were drawn from the same Kiang West villages as the EMPHASIS study, but we were unable to directly adjust for potential confounding effects due to genetic ancestry since no genetic data was available for this cohort. Based upon the EMPHASIS cohort PCA and our knowledge of village population structures, we reasoned that village of origin is a useful proxy for genetic ancestry in the ENID cohort and performed a sensitivity analysis with an additional covariate dichotomised according to whether an individual came from the predominantly Fula village. The first two genetic principal components were used as adjustment covariates for the corresponding EMPHASIS analysis. Results from this ethnicity-adjusted sensitivity analysis were not materially different from those obtained for the main analysis (Appendix 1-Fig. 13; Supplementary File 1s).

*Overlap of SoC-CpGs with existing studies*

To place our findings in the context of existing literature on associations between DNAm and nutrition-related exposures, including exposure to famine conditions, folate supplementation in gestation and previous reported associations with Gambian SoC, we checked for overlaps between SoC-CpGs and loci identified in a recent review by James et al26. Many cited studies including the majority of previous work in The Gambia used pyrosequencing and other methylation platforms targeting loci not covered by Illumina arrays. However, a total of 57 previously identified loci did overlap or partially overlap array background. None of these overlapped a SoC-CpG within 1kbp. We also checked for overlaps with the larger set of SoC-associated CpGs not passing the 4% minimum effect size threshold, and found a single CpG (cg17434309) mapping to *IGF2* that was within 1kbp of two previously identified loci, one linking maternal plasma vitamin B12 with cord blood methylation34, and the second linking gestational famine to blood methylation in older adults35.

We next looked for overlaps between SoC-CpGs and CpGs identified in the EWAS Catalog (<http://ewascatalog.org/>), a manually curated database of significant results (p<1x10-4) from published epigenome-wide association studies (EWAS). This search produced published associations for 167 out of the 259 SoC-CpGs, mapping to 27 unique traits covering a range of pre- and post-natal exposures (Supplementary File 1l). Noteworthy amongst the most frequently reported associations with SoC-CpGs (Supplementary File 1m) in the context of our study were those with sex, gestational age, maternal smoking in pregnancy, maternal plasma folate levels and adult body mass index (BMI) with 109, 45, 16, 6 and 1 associated SoC-CpG(s) respectively.

We investigated some of these links using data from the 2yr ENID cohort considered in our main analysis and confirmed multiple significant associations with infant sex but not gestational age or maternal folate at conception (Supplementary File 1n). Links with adult BMI and maternal smoking were not considered as adult BMI was not available and the incidence of smoking is extremely low in our study population.

All Fourier regression models in our main season of conception analysis included sex as an adjustment covariate. The finding that multiple SoC-CpGs were associated with sex in the EWAS Catalog, with replication of this association in the ENID cohort, was therefore surprising. This prompted us to check for a residual confounding effect due to sex by repeating our analysis with methylation values pre-adjusted for sex using a regression model with sex as the only adjustment covariate, prior to running the full regression models. This produced near identical results to the main analysis without pre-adjustment. This, combined with our observation that sex is not associated with any tested batch or biological covariates (Supplementary Files 1p-1r), strongly suggests that the observed SoC associations were not driven by confounding due to sex.

Finally we searched for SoC-CpGs within 1kbp of SNP associations (p<1x10-5) in the GWAS Catalog36, since DNAm could mediate GWAS signals in genomic regions where functional effects are difficult to elucidate37. 11 SoC-CpGs mapped to a total of 12 SNPs associated with 8 unique traits in the GWAS Catalog (Supplementary File 1o). Notable traits from a developmental programming perspective were those linked to childhood obesity38 (2 SoC-CpGs) and QRS traits associated with cardiovascular mortality in adults (1 SoC-CpG)39,40.

**Discussion**

We have utilised a natural, seasonal experiment in rural Gambia whereby human conceptions are ‘randomised’ to contrasting environmental (especially dietary) conditions to examine whether this differential exposure leaves a discernible signature on the offspring methylome. We analysed methylation data from two independent, different-aged cohorts and identified 259 ‘SoC-CpGs’ with evidence of sensitivity to season of conception (SoC) in infancy. We found evidence of SoC effect attenuation with age, but SoC-related temporal patterns were nonetheless strikingly similar, suggestive of a common effect of periconceptional environment. Importantly, the analysed datasets have contrasting confounding structures, notably with regard to the timing of sample collection, the latter eliminating potential confounding due to seasonal differences in leukocyte composition. These results, derived from analysis of Illumina array data with limited coverage, suggest there may be many more hotspots sensitive to the periconceptional environment across the human methylome.

This analysis builds on previous epigenetic studies in this setting that have focussed on single cohorts and analysed methylation differences between individuals conceived at the peaks of the Gambian dry and rainy seasons only14–16,18,41. Increased methylation in offspring conceived at the peak of the Gambian rainy season is consistent with previous findings and this observation is now greatly strengthened by the application of Fourier regression to model year-round conceptions, an approach that makes no prior assumption of when methylation peaks and nadirs may occur. The number of identified SoC-CpGs is also substantially increased in this study. Comparisons with array background and control CpGs matching SoC-CpG methylation distributions increase confidence that these findings are not statistical artefacts.

Triangulation with other public data provides multiple lines of evidence supporting the notion that methylation states at these loci are established in the periconceptional period. First, they are highly enriched for putative MEs and related loci identified in other studies with characteristic methylation signatures suggestive of establishment early in embryonic development17,18. Second, like MEs, season-associated loci exhibit unusual methylation dynamics in early stage embryos17. Third, they have distinctive gametic methylation patterns, notably hypomethylation in sperm (in common with putative MEs17), and differential gametic and parent-of-origin specific methylation in a subset. Increased incidence of hypomethylated states in sperm at SoC-CpGs may reflect their enrichment at CpG islands42, sequence features that are largely refractory to protamine exchange, with the possibility for retaining epigenetic function associated with histone modifications into the early embryo43. Fourth, many overlap H3K4me1 and H3K27me3 marks which coordinate transient gene expression and early lineage commitment in human ESCs, in part through demarking poised enhancers44.

A large majority of SoC-CpGs have not previously been identified as MEs, but given the supporting evidence described above, we speculate that many are likely to be so. Indeed, evidence of attenuation of SoC effects with age suggests that, to the extent that interindividual variation is driven by periconceptional environmental factors, screens for putative MEs (including ESS and SIV) in adult tissues used as a reference in this analysis may be missing metastable regions which are more pronounced earlier in the life course. Evidence of much larger SoC effect sizes at known MEs in both Gambian cohorts supports this. Furthermore, identification of putative human SIV-MEs requires analysis of datasets with tissues derived from different germ layers for each individual. Such datasets are expensive to acquire and rare, meaning that the few published ME screens are likely to be underpowered due to small sample sizes.

The observed attenuation of SoC effects with age has implications for detecting the effect of periconceptional exposures on DNAm in samples collected beyond the neonatal and early childhood periods, an important consideration for epigenetic epidemiological studies since non-persisting methylation differences could still have a significant impact on early developmental trajectories with life-long consequences45,46. The methylome changes with age, a phenomenon sometimes referred to as ‘epigenetic drift’47. Attenuation of SoC effects could be a consequence of localised or global epigenetic drift or other processes such as clonal selection48.

Interestingly, more SoC-CpGs replicated in the smaller ENID (5-7yr) dataset than in the older EMPHASIS (7-9yr) cohort (n=157 vs 61 respectively). This could be due to age or other cohort-specific differences, or it might reflect differences in seasonality such as secular changes in the intensity and duration of the rainy season.

Intra-individual methylation states at SoC-associated loci are highly correlated across loci despite being distributed throughout the genome, suggesting that a common mechanism is at play. MEs located within intracisternal A particle (IAP) transposable elements have been the focus of many studies of SIV in mice, the *Agouti* viable yellow locus being a paradigm example. While one study found that methylation does not co-vary across different murine IAP MEs in the absence of environmental perturbations31, another recent study identified several IAP MEs located on different chromosomes which are modified by the same cluster of KRAB zinc finger proteins49, demonstrating that genetic mechanisms for the simultaneous epigenetic regulation of multiple loci do exist.

Potential insights into mechanisms linking periconceptional environment to DNAm changes in postnatal tissues come from our investigations of the methylation status and genomic context of SoC-CpGs.

First, we observed strong enrichment for germline differentially methylated regions, with 14% of SoC-CpGs overlapping 19 gDMRs hypomethylated in sperm and hypermethylated in oocytes. A minority of these show evidence of PofOm persisting in postnatal tissues. This observation aligns with a growing body of evidence linking early environment, notably nutritional factors involved in one-carbon (C1) metabolism, with methylation at imprinted regions25,26. Indeed we have previously noted an association between season of conception and several C1 metabolites at a maternally imprinted region at the small non-coding RNA *VTRNA2-1*16, consistent with evidence of ‘polymorphic imprinting’ linked to prenatal environment at this locus27,50. Furthermore, we previously found strong enrichment for proximal binding sites of several transcription factors (TFs) associated with the maintenance of PofOm in the early embryo at MEs detected in a WGBS screen17. We were only able to replicate enrichment for one of these, ZFP57, at SoC-CpGs in this study. This may reflect the relatively small proportion of PofOm loci in the set of SoC-CpGs, or factors related to the limited methylome coverage of Illumina arrays. More targeted experimental work is required to determine the extent of SoC effects at imprinted loci, especially given our observation that SoC-CpGs are often proximal to ERV transposable elements that have recently been shown to drive the establishment of germline-derived maternal PofOm51. Hotspots with evidence of PofOm could be driven by an environmentally sensitive gain of methylation on the paternal allele that is propagated through development, incomplete reprogramming on the maternal allele leaving residual traces of methylated cytosines, or modest *de novo* methylation at some later point. A deeper understanding of mechanisms will require further investigation in cell and animal models.

Second, our observation of enrichment for proximity to ERV1 transposable elements at SoC-CpGs aligns with our previous finding at MEs16,17, and is notable since most environment-sensitive mouse MEs are associated with IAPs (which are rodent-specific ERVs)31. KRAB zinc-finger protein (KZFP)-mediated repression of transposable elements (TEs) including ERVs has also been proposed as a driver of the rapid evolution of gene regulation52. The KZFP ZFP57 is particularly interesting in this respect since its binding to DNA is linked both to repression of TEs and to the maintenance of genomic imprints in the pre-implantation embryo53,54. We previously identified a putative SoC-associated DMR in the *ZFP57* promoter in blood from younger Gambian infants (mean age 3.6 months)16. It is possible that non-replication of the SoC-association at *ZFP57* in the older Gambian cohort (and of the *VTRNA2-1* SoC-DMR mentioned above which was also identified in younger infants) reflects the more general attenuation of SoC effects described above. Interestingly there is some evidence that the *ZFP57* DMR, which lies 3kb upstream of the transcription start site, is established in the early embryo18,55, and that DNAm at this locus measured in neonatal blood is associated with maternal folate levels6,56,57. Given the important function of ZFP57 in pre-implantation methylation dynamics, its potential role as an environmentally sensitive regulator of multi-locus DNAm effects remains an open question.

Third, DNAm at SoC-CpGs is enriched for intermediate methylation states. Intermediate methylation has also been observed at MEs in Gambians and in non-Africans8,14,15,41, and this coincides with a similar observation at MEs in post-gastrulation embryonic tissues17. Intermediate methylation appears to be predominantly driven by variegated (intercellular) differences in methylation status within a sampled tissue, rather than by allele-specific methylation58. Observed differences in aggregate methylation at SoC-CpGs could thus reflect a direct influence of periconceptional environment on the establishment and maintenance of DNAm states in the early embryo at the cellular level, or ‘epigenetic selection’ whereby epigenetically distinct cells form a substrate for clonal selection during development, for example as a potential adaptation to differential metabolic exposures59. Stronger enrichment for putative MEs exhibiting SIV compared to those identified through ESS supports establishment in the post-implantation embryo, since methylation differences at ESS loci are presumed to originate in the pre-implantation embryo prior to embryo cleavage in MZ twins18.

The largest SoC-CpG cluster with 7 CpGs is on chromosome 15 and falls within the second intron of *IGF1R.* This bears the hallmarks of being a promoter or active/poised enhancer in multiple cell lines (Appendix 1-Fig. 14). Six CpGs in the *IGF1R* cluster have evidence of PofOm and show a relatively large SoC effect size (median SoC methylation amplitude) of 6.1%. Zink *et al*27 were unable to demonstrate PofO allele-specific expression in this region although others have found evidence of maternal imprinting of an intronic lncRNA at this gene in cancerous cells60,61. Loss of IGF1 receptors gives rise to a major decrease in expression at multiple imprinted genes in mice, suggesting a pathway by which *IGF1R* might regulate growth and metabolism during early development62. IGF1R signalling is implicated in fetal growth, glucose metabolism and cancer63–65, and DNAm differences at *IGF1R* have been observed in birthweight-discordant adult twins66. Another SoC-CpG with evidence of PofOm, also on chromosome 15, falls within the known Prader-Willi syndrome-associated paternally expressed imprinted region.

Our observation that 109 out of 259 SoC-CpGs have been associated with sex in previous studies is intriguing. Given the relatively small number (compared to array size) of autosomal sex-linked loci identified in large studies on the Illumina 450k array67,68, this represents a very strong enrichment. We replicated a significant sex association at these loci in the ENID cohort analysed in this study. Our regression analyses were adjusted for sex, and additional sensitivity analyses with DNAm pre-adjusted for sex strongly suggest that our main findings are not confounded by sex. Interestingly, sex-associated loci are enriched at imprinted regions and sex-discordance at autosomal CpGs has been linked to androgen exposures *in utero*67,69. There is also evidence of sex differences in methylation at *DNMT3A/B* and *TET1* genes involved in *de novo* methylation and de-methylation pathways67,69, suggesting a possible interaction between sex-linked epigenetic changes and periconceptional environment during reprogramming in the early embryo. A deeper understanding of potential mechanisms underpinning the observed enrichment of sex effects at loci associated with periconceptional environment requires further functional analysis in cell and/or animal models.

DNAm is influenced by genotype and the latter is therefore a potential confounder when studying the effects of environmental exposures in human populations. A strength of our quasi-randomised Gambian seasonal model is that it minimises the potential for genetic confounding of modelled seasonal DNAm patterns, on the assumption that the timing of conceptions is not linked to genetic variants influencing DNAm. However, it is still possible that such variants might confound our observations, for example if they promote embryo survival under conditions of environmental stress. We tested this possibility using genetic data available for the EMPHASIS cohort and found no evidence of SoC-associated genetic variants driving inter-individual methylation differences at SoC-associated loci in *cis* or *trans*.

However, we did find that half of SoC-CpGs had at least one associated mQTL, indicating the presence of independent additive effects of environment and genetics at these loci, as has been suggested previously at other loci sensitive to pre/periconceptional nutritional exposures35,70. We have previously argued that the definition of MEs should be extended to include genomic regions whose DNAm state is under partial but non-deterministic genetic influence in genetically heterogeneous human populations17, and we would argue that the above observations at SoC-CpGs that exhibit many of the characteristics of MEs support this. Further analysis in larger datasets with genome sequencing data combined with functional analysis using cell models will be required to fully understand the relative contributions of environment and genetics to DNAm variation at regions of the type highlighted in this study.

Further work is required to investigate the functional relevance of DNAm changes at SoC-CpGs, some of which are relatively small (around 4% SoC amplitude). However, we can speculate that observed methylation changes, which may reflect changes in the chromatin landscape, have the potential to influence gene expression and early development, since our chromatin state analysis predicts that many overlap regions with functional significance in H1 ESCs and fetal tissues. A similarly modest DNAm change at a locus in the *POMC* gene that is associated with SoC in blood has been linked to obesity risk in German children and adults41,71, and to differential transcription factor binding and differences in *POMC* expression71. Furthermore, blood DNAm at a putative ME within the *PAX8* gene has been linked to SoC in Gambian infants15 and is associated with thyroid volume and function in Gambian children, and with certain maternal nutrition biomarkers at conception72. *PAX8* DNAm is also associated with expression of the anti-sense gene *PAX8-AS1* (alternatively known as *LOC654433*) in thyroid tissue72. No loci from either study overlap array background in this study. More generally, DNAm at SIV loci associated with periconceptional environment has been associated with a number of diseases including Alzheimer’s, cancer, rheumatoid arthritis and schizophrenia21.

We have previously shown that several metabolites involved in one-carbon methylation pathways show significant seasonal variation in maternal blood plasma in this population which is largely dependent on subsistence farming11,15. While we suspect that seasonal differences in nutrition are likely drivers of the effects that we have observed, analysis of the links between maternal nutritional biomarkers and DNAm is challenging due to the complex interdependence of one-carbon metabolites acting as substrates and cofactors within one-carbon pathways73 . Furthermore, it is possible that DNAm differences are linked to other aspects of Gambian seasonality, such as variation in pesticide use or infectious disease burden10,74.

Measured DNAm values at SoC-CpGs and matched controls have previously been reported to be reliable, in strong contrast to the majority of probes overlapping the 450k and EPIC arrays that have been found to have low test-retest reliability23. This likely reflects increased inter-individual variability and/or intermediate DNAm at SoC-CpGs and controls75,76.

Limitations of this analysis include a lack of genetic data for the ENID cohort, so that our genetic analyses are restricted to the older EMPHASIS cohort where SoC effects are smaller. Furthermore, comparator data on MEs and gamete, embryo and PofO methylation are derived from non-African individuals so we cannot assess the influence of ethnicity on these analyses. We also note that there is some inter-relatedness in this study population which practices polygamy. However, of the 199 individuals with available data on paternal identity in the ENID cohort, 2% have a shared father suggesting that this is unlikely to influence the main findings.

There is increasing interest in the phenomenon of methylation variability as a marker of disease and of prenatal adversity, and in genetic variation as a potential driver of methylation variance77. This raises the possibility that certain genetic variants could have been selected through their ability to enable graded, environmentally-responsive methylation patterns at MEs and SoC-associated loci that are able to sense the periconceptional environment, record the information, and adapt the phenotype accordingly. Our gene-environment interaction analysis likely lacked power to detect gene-environment (SoC) interactions, so that it was not possible to investigate the relative contributions of stochastic, environmental and genetically-mediated variance effects on the establishment of periconceptional SoC-associated methylation states in this study. Nonetheless, the proposition that environmentally sensitive epigenetic signals are selected through their ability to direct phenotypic development to better fit the anticipated future environment, leading to maladaptation and future disease if the environment changes, is intriguing and worthy of further investigation5.

**Materials and Methods**

*Gambian cohorts and sample processing*

Detailed descriptions of the Gambian cohorts analysed in the season of conception study are published elsewhere12,13. Briefly, ﻿for the ENID 2yr dataset, blood samples from 233 children aged 24 months (median[IQR]: 731[729,733] days old) were collected from participants in the **E**arly **N**utrition and **I**mmune **D**evelopment (“ENID”) study12, born in 2011 and 2012. DNA was extracted, bisulfite-converted and hybridised to Illumina HumanMethylation450 (hereafter “HM450”) arrays following standard protocols (see Van Baak *et al*18 for further details). N=138 ENID participants with 2yr HM450 data who were enrolled in a follow-up study with DNA collected at 5-7yr (6.2 yrs [5.7,6.6]) had DNA bisulfite-converted and hybridised to Illumina Infinium Methylation EPIC (hereafter “EPIC”) arrays. For the EMPHASIS cohort, DNA was extracted from blood samples from 289 Gambian children aged 7-9yr (9.0 [8.6,9.2] years) participating in the **﻿E**pigenetic **M**echanisms linking **P**re-conceptional nutrition and **H**ealth **As**sessed in **I**ndia and **S**ub-Saharan Africa (“EMPHASIS”) study13, born between 2006 and 2008. DNA was bisulfite-converted and hybridised to EPIC arrays, again using standard protocols.

For the ENID cohort, date of conception was calculated from fetal gestational age estimates obtained by ultrasound at the mother’s first ‘booking’ appointment. The same method was used for the EMPHASIS cohort, except for n=71 pregnancies that were > 24 weeks gestation at booking meaning that GA could not be accurately determined by ultrasound13,78. In this case date of conception was calculated as date of birth minus 280 days which is the average gestational length for this population.

Samples from the ENID 2yr dataset were processed in two batches with each batch covering conception dates throughout the year. Samples from the ENID (5-7yr) and EMHASIS (7-9yr) datasets were processed in single batches.

*Methylation array pre-processing and normalisation*

Raw intensity IDAT files from the HM450 and EPIC arrays were processed using the *meffil*79 package in R (v3.6.1) using standard *meffil* defaults. Briefly, this comprised probe and sample quality control steps (filtering on bisulfite conversion efficiency, low probe detection p-values and bead numbers, high number of failed samples per probe, high number of failed probes per sample); methylation-derived sex checks; removal of ambiguously mapping (i.e. cross-hybridising) probes80; removal of probes containing SNPs at the CpG site or at a single base extension; and removal of non-autosomal CpGs. Following filtering, methylation data was normalised with dye-bias and background correction using the *noob* method, followed by *Functional Normalisation* to reduce technical variation based on principal component analysis of control probes on the arrays81. After pre-processing and normalisation, methylation data comprised methylation Beta values for 421,026 CpGs on the HM450 array for the 233 individuals from the ENID 2yr cohort, and 802,283 CpGs on the EPIC array for 289 individuals from the EMPHASIS cohort. 391,814 CpGs intersecting both arrays were carried forward for statistical analysis of these two datasets. Finally, SoC-CpGs, matched and random controls, all of which were present in the ENID (5-7yr) EPIC dataset, were included in the longitudinal analysis.

*Statistical modelling*

Variation of DNAm with date of conception was modelled using Fourier regression22. This models the relationship between a response variable (here DNAm) and a cyclical predictor (date of conception). The effect of the latter is assumed to be cyclical due to annually varying seasonality patterns, so that the modelled effect for an individual conceived on the 31st December should be ‘close’ to that for an individual conceived on the 1st of January. This is achieved by deconvolving the conception date (predictor) into a series of pairs of sin and cosine terms, and obtaining estimates for the regression coefficients β and γ in the following model:

$$M\_{ij}=α\_{0j}+\sum\_{k=1}^{m}α\_{ik}+\sum\_{r=1}^{n}[β\_{rj}\sin((rθ\_{i}))+γ\_{rj}\cos((rθ\_{i})])+ε\_{ij}$$

Where, for individual *I* and CpG *j*:
*Mij* is the logit-transformed methylation Beta value82;
*α0j* is an intercept term;
*αik* is the *kth* of *m* adjustment covariates;
*θi* is the date of conception in radians in the interval [0, 2π], with 1st January = 0 and 31st
December = 2π, modelled as *n* pairs of Fourier terms, sin *θi* + cos *θi* + ... + sin *nθi* + cos *nθi*;
βr*j* and γr*j* are the estimated regression coefficients for the *r*th sin and cosine term respectively;
and ε*ij* is the error term.

With a single pair of Fourier terms (*n*=1), this gives a sinusoidal pattern of variation, with a single maximum and minimum whose phase (position in the year) and amplitude (distance between methylation maximum and minimum) is determined by β1 and γ1, with the constraint that the maximum and minimum are 6 months apart. More complex patterns of seasonal variation are afforded by higher frequency pairs of Fourier terms (*r*>1).

For this analysis we modelled the effect of date of conception using a single pair of Fourier terms (*n*=1) and assessed goodness of fit by comparing full and covariate-only models using likelihood ratio tests. For all datasets, covariates included child sex, and the first six principal components (PCs) obtained from unsupervised principal component analysis (PCA) of the normalised methylation M-values. The PCs were used to account for unmeasured and measured technical variation (due to bisulfite conversion sample plate, array slide etc) and cell composition effects (see Supplementary Files 1p-1r). Additional checks confirmed no seasonal variation in estimated white cell composition in the ENID 2yr and EMPHASIS 7-9yr cohort used in the main analysis (see below). 450k Sentrix Column was included as an additional adjustment covariate for the ENID 2yr cohort since this was not robustly captured by any of the first 6 PCs (Supplementary File 1p). Child age was included as an additional adjustment covariate for the ENID 5-7yr and EMPHASIS cohorts, as was maternal nutritional intervention group (see Chandak *et al*13 and Saffari *et al*70for further details).

For each CpG *j*, coefficient estimates βj , γj were determined by fitting regression models using *lm()* in R. Model goodness-of-fit was determined by likelihood ratio test (LRT) using *lrtest*() in R, comparing the full model including Fourier terms, with a baseline covariates-only model. A model p-value, p*j* was then derived from the corresponding LRT chi-squared statistic. Thus for a given threshold, , pj< supports rejection of the null hypothesis that for CpG *j*, the full model including the effect of seasonality modelled by one pair of Fourier terms, fits no better than the covariate-only model at the  level.

To reduce the influence of methylation outliers, methylation values greater than three times the methylation inter-quartile range (IQR) beyond the 25th or 75th centiles were excluded prior to fitting the models. This resulted in 382,095 (267,369) outliers, or an average of 0.98 (0.68) outliers per CpG being removed from the ENID and EMPHASIS methylation datasets respectively.

Note that our observation of attenuation of SoC effect sizes with age at SoC-CpGs when comparing DNAm data at aged 2yr (ENID) and 7-9yr (EMPHASIS), supported by analysis of longitudinal data (ENID 5-7yr), suggested that a combined meta-analysis of ENID and EMPHASIS datasets would have reduced power to detect SoC effects at 2 years of age.

*Sensitivity analyses*

We performed sensitivity analyses following the same Fourier regression modelling strategy as outlined above, but with i) methylation estimated cell counts (using *estimateCellCounts*() from *minfi* (v1.30.0)); ii) known batch and technical covariates; and iii) village ID included as additional covariates. See Supplementary File 1s for further details on regression models used in the sensitivity analyses. For each sensitivity analysis we tested to see if Fourier regression coefficients estimated in the main analysis fell within 95% CIs in the adjusted models.

*Inflation of test statistics*

The concept of genomic inflation rests on the assumption that a relatively small number of loci will be associated with the exposure (or disease/outcome) of interest. Test statistics for the ENID (2yr) cohort did show signs of genomic inflation (lambda = 1.33), suggesting a potential effect of season of conception on global methylation levels83. A similar level of inflation was observed before in a study looking at the effect of periconceptional folate on DNAm6. There is also evidence of global and/or multi-locus effects of folate in other studies, including an RCT of folate supplementation in pregnancy57, and there are many other examples including studies investigating the effect of mutations in the *MTHFR* gene (see review by Crider et al84). While we do not know if the SoC associations we observe in our cohorts are driven by seasonal differences in folate, we do observe significant seasonal differences in multiple one-carbon (C1) metabolites including folate in our population11,15, so that SoC may serve as a proxy for multi-locus C1 metabolite effects in our analyses.

We tested for potential SoC effects on global methylation by analysing methylation differences at LINE1 and Alu elements using REMP (v1.16.0), a recently published method to predict methylation at repetitive elements from Illumina array data83. We used default cutoffs for reliability suggested by the authors. This analysis confirmed a small but significant effect of increased methylation in rainy vs dry conceptions at LINE1 and Alu elements (mean rainy season increase 0.02% and 0.01% respectively, both Wilcoxon p<2.2x10-16) suggesting a SoC effect on global methylation levels. This contrasted with a significant but extremely small effect in the opposite direction across array background (-5x10-4 %; p<2.2x10-16; see Appendix 1-Fig. 15), supporting evidence from previous studies of a potential effect of peri/preconception nutrition-related exposures on global methylation levels.

*Identification of SoC-CpGs*

For the ENID 2yr cohort, p-values, pj, were used to compute a false discovery rate for each CpG accounting for multiple testing (assuming 391,814 independent tests corresponding to the number of loci in array background) using *p.adjust()* in R with method= ‘fdr’. Following the rationale described in the main text, SoC-associated CpGs with a SoC amplitude < 4% in the ENID cohort were then excluded to form the final set of 259 ‘**SoC-CpGs**’ (see Table 2).

*Selection of control CpGs*

SoC-CpGs are enriched for intermediate methylation states (Fig. 4B), so that there is a risk that some downstream analyses reflect the distributional properties of these loci, rather than factors associated with their putative establishment at periconception. For this reason we identified a set of ‘**matched control**’ CpGs that were selected to have similar methylation Beta distributions to SoC-CpGs (and additional SoC-associated CpGs with amplitude <4%) in the ENID 2yr dataset. Matched controls were drawn from array background (excluding SoC-associated CpGs and known MEs/ESS/SIV CpGs), with one matched control identified for each of the 768 SoC-associated CpGs. Alignment of control and SoC-CpG methylation distributions was achieved using a two-sided Kolmogorov-Smirnov test for divergence of cumulative distribution functions (*ks.test()* in R) with a p-value threshold p>0.1. Examples are given in Appendix 1-Fig. 16, along with a comparison of sample mean distributions.

An additional 768 **random control** CpGs were randomly sampled from array background, again excluding SoC-CpGs and known MEs/ESS/SIV CpGs.

*CpG sets considered in analyses*

Summary information on external datasets considered in the analyses is provided in Table 3. Further information on these is provided below.

1. *1,881 putative ME* CpGs overlapping array background from one or more of the following curated sets of loci: putative MEs exhibiting SIV identified in a multi-tissue WGBS screen in individuals of European and African-American decent described in Kessler *et al*17; and CpGs exhibiting ‘epigenetic supersimilarity’ and/or SIV derived from analysis of 450k data from individuals of European decent, as described in Van Baak *et al*18.
2. *699 parent-of-origin-specific CpGs (PofOm CpGs)* overlapping array background from 229 regions with PofOm identified in Supplementary Table 1 from Zink et al27. PofOm identified in peripheral blood from Icelandic individuals.
3. *RRBS early stage embryo data* from Chinese embryos described in Guo *et al*4 downloaded from ﻿GEO (accession number GSE49828). Only CpGs covered at ≥ 10x in pre-gastrulation inner cell mass and/or post-gastrulation embryonic liver were considered in this analysis. Further details are provided in Kessler *et al*17.
4. *Sperm methylation data* from Japanese donors described in Okae *et al*24 downloaded from the Japanese Genotype-phenotype Archive (accession number S00000000006). Only CpGs covered at ≥ 10x were considered in this analysis.
5. *Germline gDMRs* (gDMRs), defined as contiguous 25 CpG regions that were hypomethylated (DNAm mean +1SD < 25%) in one gamete and hypermethylated (DNAm mean-1SD > 75%) in the other, were previously identified from WGBS data by Sanchez-Delgado *et al*28. Persistence of PofOm to the blastocyst and placental stages was established by identifying overlapping intermediately methylated regions in the relevant embryonic tissues, with confirmation of PofOm expression at multiple DMRs28. Japanese and US donors. See Sanchez-Delgado *et al*28 for further details.
6. *Transposable elements (ERVs)* determined by RepeatMasker were downloaded from the UCSC hg19 annotations repository.
7. ZFP57, TRIM28 and CTCF transcription factor binding sites identified from ChIP-seq in human embryonic kidney and human embryonic stem cells used in this analysis are described in Kessler *et al*17.

*Cluster-based adjustments*

Many SoC-CpGs cluster together and this could influence some analyses. For example, methylation at CpGs may be highly correlated, which could influence comparisons of inter-CpG correlations between SoC-CpGs and controls (Fig. 3D). Also enrichment tests are likely to be influenced by neighbouring CpGs that together constitute a single ‘enrichment signal’ proximal to a particular genomic feature (e.g. a transcription factor binding site).

To account for this, cluster-adjusted analyses used ‘de-clustered’ CpG sets that were constructed as follows:

1. Create CpG clusters formed from adjacent CpGs where each CpG is within 5kbp of the nearest neighbouring CpG (see Appendix 1-Fig. 5 for a justification of this threshold);
2. Construct de-clustered test set by randomly sampling a single CpG from each cluster; non-clustered ‘singleton’ CpGs are always selected.

In the case of SoC-CpGs, the set of 259 non-clustered CpGs were reduced to 161 CpGs after de-clustering.

*Chromatin state and histone (H3K) mark analysis*

Data on chromatin states predicted by the ChromHMM 15-state model85 in H1 ESCs, and 3 fetal tissues (brain, muscle and small intestine) derived from 3 different germ layers, generated by the Roadmap Epigenomics Consortium29 was downloaded from the Washington University Roadmap Epigenomics repository. 15-state model predictions were collapsed to 8 states for visualisation purposes, with sub-classifications described in the Fig. 5 caption.

Overlaps with H3K marks for each of the above tissues were assessed using the *annotatr (v1.10.0)* package in R to interrogate the same Roadmap Epigenomics ChIP-seq data used by ChromHMM for chromatin state prediction.

*Additional modelling of seasonal variation in blood cell composition*

Cell count estimates using the Houseman method86 were obtained using the *estimateCellCounts*() from *minfi* (v1.30.0) in R. Seasonal variation in blood cell composition was then modelled by Fourier regression with one pair of Fourier terms and sex (ENID+EMPHASIS) and age (EMPHASIS only) as adjustment covariates. Fitted models indicated no marked seasonal differences within and between cohorts (Appendix 1-Fig. 17).

*Genetic association analyses*

Genotype data

mQTL and related SoC-association analyses were performed on all 284 individuals from the EMPHASIS (7-9yr) cohort for which we had QC’d genotype data. 259 SoC-CpGs, plus sets of 259 matched and random control CpGs were considered in this analysis. Subjects were genotyped using the Illumina Infinium Global Screening Array-24 v1.0 Beadchip (Illumina, California, U.S.) following standard protocols. Array-derived genotypes were pre-phased using SHAPEITv2 and imputation was performed using IMPUTEv2.3.2 on 1000 genomes phase 3 data. Further details are provided in Saffari et al70. SNPs with a MAF ≤ 10% were excluded, along with those with an IMPUTE ‘info’ metric ≤ 0.9, a stringent threshold to ensure maximum confidence in imputation quality. Imputed SNPs were then pruned (using plink v1.90 -indep-pairwise with window size 50, step size 5 and r2 threshold of 0.8) to remove SNPs in strong LD. Finally, to minimise the influence of low frequency homozygous variants in linear models, analysis was restricted to SNPs with 10 or more homozygous variants, resulting in a final dataset comprising 2,609,310 SNPs.

Identification of mQTLs and ‘GxE’ SNPs

mQTL analysis was performed using the *GEM* package (v1.10.0) from R Bioconductor87.

SNP effects on methylation were modelled as follows*:*

M*j* ~ covs + G

Where M*j* is the methylation M-value for CpG *j*, G is the SNP genotype coded as allelic dosage (0,1,2) and covs correspond to the adjustment covariates used in the main EMPHASIS analysis (PCs 1-6, child age, sex and intervention status).

GxE (SoC) effects were modelled as follows:

M*j* ~ covs + sinθ+ G + G x sinθ

when sinθ is the most significant Fourier term in the main SoC analysis,

OR

M*j* ~ covs + cosθ + G + G x cosθ

when cosθ is the most significant Fourier term in the main SoC analysis.

Here, G and covs are as described above.

Since most mQTL effects are known to act in *cis*32, in order to maximise power a two-step procedure was used to identify significant mQTLs:

1. *Identification of cis-mQTLs* passing FDR<5% considering the reduced set of SNPs within 1Mbp of a CpG in the set to be analysed (SoC-CpGs, matched or random controls);
2. *Identification of trans-mQTLs* passing FDR<5%, considering the full set of 2.6M SNPs.

Calculation of methylation variance explained by mQTLs

For each CpG *j*, total methylation variance explained was calculated for the following model:

*Mj* ~ covs + mQTL*j*1 + mQTL*j*2 + ... mQTL*j*n

where covs are as defined above, and mQTL*jn* is the genotype (coded 0,1,2) of the *n*th mQTL mapping to CpG *j.* Methylation variance explained was calculated from the model adjusted R2 value, adjR2mQTL , to account for different levels of model complexity due to the differing number of mQTL identified for each CpG.

A final estimate of mQTL methylation variance explained was obtained by subtracting variance explained by the covariate-only model:

ΔadjR2 = adjR2mQTL – adjR2cov

where adjR2cov is the adjusted R2 for the covariate-only model,

*Mj* ~ covs.

SoC association analysis

Potential confounding of SoC-DNAm signals by SoC-associated genetic variants was assessed by analysing SoC associations with all 2,771 SoC-CpG-associated mQTLs identified in the mQTL analysis (see Table 4). SoC association analysis was performed under an allelic model using ­‑assoc with *plink* v1.90. Significant mQTL-SoC associations were identified using FDR<5% which assumes independence of all mQTL-SoC associations. We also considered a more liberal multiple testing correction threshold: *pBonf* = 0.05/130, assuming complete dependence of all *cis-*mQTL mapping to each of the 130 SoC-CpG with an associated mQTL.

Genetic ancestry sensitivity analysis

An investigation of population structure in the EMPHASIS cohort was conducted by first performing a principal component analysis (PCA) using ‑pca in plink v1.90. The PCA was performed on non-imputed data, with LD pruning using ‑indep-pairwise 50 5 0.2 in plink v1.90, corresponding to an r2 threshold of 0.288. Evidence for population structure was then obtained by plotting the first four principal components (Appendix 1-Fig. 12A). Confirmation of a likely link to Gambian ethnic ancestry in this largely ethnically Mandinka region of Gambia followed our observation that a distinct cluster (Appendix 1-Fig. 12B) was primarily made up of individuals from a single, predominantly ethnic Fula village.

A sensitivity analysis to check the effect of accounting for ancestry differences was performed by repeating the main analysis with the following additional covariates:

1. EMPHASIS (7-9yr) cohort

We adjusted for genetic ancestry directly using the first two genetic principal components identified in the genetic PCA.

1. ENID (2yr) cohort

Since no genetic data is available for this cohort, and since individuals from this cohort are drawn from the same villages as the EMPHASIS cohort on which we did the genetic PCA, we reasoned that village of origin is a useful proxy for genetic ancestry in our population. We therefore included an additional covariate dichotomised according to whether or not an individual was one of the 9 who came from the predominantly Fula village identified as genetic outlier in the EMPHASIS PCA analysis.

Sensitivity analysis to investigate confounding by infant sex

Our finding that multiple SoC-CpGs were associated with infant sex in previous EWAS prompted us to perform a sensitivity analysis checking for the possibility of a residual confounding effect due to sex. To do this we regressed out the effect of infant sex at each CpG in the M-value methylation matrix, prior to the main regression analysis. We then re-ran the Fourier regression analysis with and without an additional adjustment for infant sex in Fourier regression models. As expected, given that we adjusted for infant sex in the main analysis, this produced near identical results, suggesting that the main analysis was not confounded by infant sex.

*All bootstrapped confidence intervals presented in this paper use 1,000 bootstrap samples.*

Ethics approval and consent to participate

Ethics approval for the Gambian ENID and EMPHASIS trials was obtained from the joint Gambia Government/MRC Unit The Gambia’s Ethics Committee (ENID: ﻿SCC1126v2; EMPHASIS: SCC1441). The ENID study is registered as ﻿ISRCTN49285450. The EMPHASIS study is registered as ISRCTN14266771. Signed informed consent for both studies was obtained from parents, and verbal assent was additionally obtained from the older children who participated in the EMPHASIS study.

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**Competing interests:** Authors declare that they have no competing interests.

**Data and materials availability:**

Data and code availability

Illumina 450k methylation array data generated from Gambian 2 year olds from the ENID trial is deposited in GEO (GSE99863). Requests to access and analyse the other Gambian methylation datasets (ENID 5-7yr and EMPHASIS 7-9yr) should be submitted to the corresponding author in the first instance. An application would then need to be made to MRC Unit The Gambia’s Scientific Coordinating Committee and the Joint MRC/Gambia Government Ethics Committee. Sources and locations of other publicly available data used in this analysis are described in Methods. Bespoke code used in the analysis is available at <https://zenodo.org/record/5801480>89.

**Figures captions**

**Figure 1. Study design.** DNAm: DNA methylation; EPIC: Illumina Infinium MethylationEPIC BeadChip; Epig. Roadmap: Roadmap Epigenomics Consortium; ESC: embryonic stem cell; ESS: epigenetic supersimilarity; GSA: Global Screening Array; HM450: Illumina Infinium HumanMethylation450 BeadChip; IVF: in vitro fertilisation; MEs: metastable epialleles; MZ/DZ: monozygotic / dizygotic twins; PofO: parent of origin; RRBS: reduced representation bisulfite-seq; SIV: systemic interindividual variation; SoC: season of conception; WGBS: whole genome bisulfite-seq. See Tables 1 and 3 for further details of Gambian and public datasets used in this analysis.

**Figure 2. Identification of Gambian season of conception associated CpGs: ENID (2yr) vs EMPHASIS (7-9yr) cross-cohort analysis.** **(A)** Relationship between date of conception and date of sample collection for ENID (top) and EMPHASIS (bottom) cohorts. **(B)** Modelled seasonal change in methylation for 768 SoC-associated loci (FDR<5%) in the ENID cohort. 26 ME CpGs are marked in red. **(C)** Conception date of modelled methylation maximum in each cohort for 61 CpGs significantly associated with SoC in both cohorts (left) and 61 matched controls (right). **(D)** (Left) Date of modelled DNAm maximum vs seasonal amplitude in each cohort for 768 CpGs significantly associated with SoC in the ENID cohort. MEs are marked in red. Dashed line indicates SoC amplitude threshold used to identify SoC-CpGs. (Right) same CpGs as left, but in the older EMPHASIS cohort. Significant SoC association for this cohort are marked in a darker colour. **(E)** (Left) Seasonal amplitudes for SoC-associated CpGs that are (red) or are not (blue) MEs; along with amplitudes for 768 matched and random controls (light/dark grey respectively). (Right) as left but in the older EMPHASIS cohort. For EMPHASIS significant SoC associations are marked in a darker colour. Boxes represent the middle 50% of the data (interquartile range, IQR); the line inside the box is the median, and whiskers represent values lying within 1.5 times the IQR.

**Figure 3. Identification of Gambian season of conception associated CpGs: ENID 2yr vs 5-7yr longitudinal analysis. (A**) Relationship between conception date of modelled methylation maximum measured at 2yr and 5-7yr in the same n=138 individuals from the ENID cohort. n=157 SoC-CpGs with a significant SoC association (FDR<5%) at 5-7yr are plotted. (**B)** Change in seasonal amplitude between 2yr and 5-7yr for n=259 SoC-CpGs (left) and matched controls (right) with DNAm measured in the same n=138 individuals from the ENID cohort. There is evidence of SoC effect attenuation with age at SoC-CpGs, but not at matched controls (Wilcoxon signed rank test for difference in SoC amplitude p=6.7x10-12 and p=1.0 respectively).

**Figure 4. Properties of SoC-CpGs.** (**A)** Date of conception at modelled methylation maxima for 259 SoC-CpGs and 259 corresponding matched and random controls across all 3 analysed datasets. Green and yellow bands indicate the extent of the rainy and dry seasons respectively. (**B)** SoC-CpGs show increased intermediate methylation compared to array background (data from ENID 2yr and EMPHASIS cohorts combined)**. (C)** Distribution of 259 SoC-CpGs (including MEs), 259 matched controls and array background with respect to CpG islands. Error bars are bootstrapped 95% CIs. N / S Shore / Shelf: North / South Shore / Shelf respectively (regions proximal to CpG Islands defined in Illumina manifest). (**D)** Distribution of pairwise Spearman correlations for CpG sets in the ENID (left) and EMPHASIS (right) cohorts. Boxplot elements as described in Fig. 2E.

**Figure 5. Early stage embryo, gametic and parent-of-origin specific methylation.** **(A)** Methylation distribution of SoC-CpGs, matched controls and array background in pre-gastrulation inner cell mass (ICM) and post-gastrulation embryonic liver, measured in RRBS embryo methylation data from Guo *et al*4. Data comprises 112,380 CpGs covered at ≥10x in ICM and/or embryonic liver that overlap array background, including 118 SoC-CpGs and 51 matched controls. **(B)** Methylation distribution of SoC-CpGs, matched controls and array background in sperm WGBS data from from Okae *et al*24. Data comprises 294,240 CpGs covered at ≥ 10x including 196 SoC-CpGs and 207 matched controls. **(C)** Sperm methylation and oocyte gDMR (oo-gDMR) status at 196 SoC-CpGs covered at ≥ 10x in Okae *et al* sperm WGBS data. Sperm hypomethylation is defined as methylation ≤ 25%. oo-gDMRs defined as sperm methylation <25% and oocyte methylation >75% in WGBS analysis by Sanchez-Delgado. *et al*28.**(D)** Mean methylation at SoC-CpGs and array background measured in n=233 individuals in the ENID (2yr) cohort stratified by sperm and oocyte methylation status. Left: Methylation stratified according to sperm hypomethylation status (n=175 SoC-CpGs hypomethylated in sperm, n=21 not hypomethylated). Right: as left but with loci hypomethylated in sperm further stratified according to oo-gDMR status (n=26 SoC-CpGs hypermethylated in oocytes / oo-gDMR, n=149 not hypermethylated). Sperm / oo-gDMR status thresholds as for Fig. 4C. 8 PofOm CpGs (green triangles) are those identified in Zink *et al*27.

**Figure 6. SoC-CpG overlap with predicted chromatin states.** Chromatin states predicted by ChromHMM85 from chromatin marks in four cell lines and tissues generated by the Roadmap Epigenomics Consortium29. Predicted states for all 259 SoC-CpGs are shown. Predictions from the ChromHMM 15-state model are collapsed to 8 states for clarity. TSS: active transcription start site/flanking active TSS/bivalent or poisedTSS; TSS/Enhancer: flanking bivalent TSS/enhancer; Enhancer: Enhancer/bivalent enhancer/Genic enhancer; Transcription: Strong/weak transcription/ Transcription at gene 5’ and 3’; ZNF/repeat: zinc finger genes & repeats; Polycomb: Repressed/ weak repressed polycomb; Heterochromatin; Low signal: low signal in all marks states used as inputs to ChromHMM.

**Figure 7. Links between endogenous retroviruses (ERVs), ZFP57 binding sites, genetic variation and DNAm at SoC-associated loci. (A)** Proportion of SoC-CpGs, matched controls and array background CpGs proximal to ERV1 endogenous retroviral elements (top) and ZFP57 binding sites (bottom), within the specified distance. CpG clustering effects are removed by sampling a single CpG from each cluster (see Methods). Error bars are bootstrapped 95% CIs. (**B)** Proportion of methylation variance explained by mQTL for matched controls, SoC-CpGs and random controls. Only CpGs with at least one significant mQTL are plotted (n=201, 130 and 50 respectively; see Methods for further details). Boxplot elements as described in Fig. 2E.

**Tables**

**Table 1. Gambian seasonality-methylation analysis: cohort characteristics**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Cohort | sample size | Age  | % male | Tissue | methylation array |
| ENID (2yr)12ENID (5-7yr) | 233138 | 2yr5-7yr | 50.656.5 | peripheral blood | Illumina Infinium HM450Illumina Infinium MethylationEPIC |
| EMPHASIS13 | 289 | 7-9yr | 54.3 | peripheral blood | Illumina Infinium MethylationEPIC |

*Note: Individuals with ENID longitudinal (5-7yr) methylation data are a subset of those with methylation at 2yr. There is no overlap between individuals included in the ENID and EMPHASIS cohorts.*

**Table 2. CpG sets considered in this analysis.**

|  |  |  |
| --- | --- | --- |
| CpG set  | Number of CpGs  | Notes |
| Array background | 391,814 | Intersection of CpGs on Illumina HM450 (ENID 2yr) and EPIC (EMPHASIS) cohort arrays, post QC |
| SoC-CpGs | 259 | SoC-associated CpGs with SoC effect size (SoC methylation amplitude) > 4% in the ENID 2yr dataset |
| Matched controls | 259 | CpGs with similar methylation distributions to SoC-CpGs in the ENID 2yr dataset\* |
| Random controls | 259 | Random sample from array background |

\*Matching methylation distributions determined by Kolmogorov-Smirnov tests (see Appendix 1-Fig. 16). QC: quality control; LRT: likelihood ratio test. See Methods for further details.

**Table 3. External datasets considered in this analysis.**

|  |  |
| --- | --- |
| CpG set  | Notes |
| Putative metastable epialleles (**MEs**) | 1881 **ME/SIV/ESS** CpGs overlapping array background identified in multi-tissue and MZ/DZ screens in Van Baak *et al*18 and Kessler *et al*17. |
| Parent-of-origin specific methylation (**PofOm**) | 699 Parent-of-origin specific methylation loci identified in peripheral blood in Zink *et al*27 overlapping array background. |
| Embryo DNAm data | RRBS data for inner cell mass and embryonic liver (<10 weeks gestation) from Guo *et al*4. |
| Sperm DNAm data | WGBS data from Okae *et al*24. |
| Germline DMRs (**gDMRs**) | Regions differentially methylated in sperm and oocytes identified in WGBS data by Sanchez-Delgado *et al*28. |
| Transposons (**ERVs**) | ERVs determined by RepeatMasker were downloaded from the UCSC h19 annotations repository. |
| Transcription factor ChIP-seq | ZFP57, TRIM28 and CTCF transcription factor binding sites identified from ChIP-seq in human embryonic kidney and hESCs are described in Kessler *et al*17. |
| Chromatin state predictions and histone 3 marks | Chromatin state predictions for H1 ESCs, fetal brain, fetal muscle and fetal small intestine generated using ChromHMM85 from Roadmap Epigenomic Consortium data29. Histone mark data are from the same source. |

ME: metastable epiallele; SIV: systemic interindividual variation; ESS: epigenetic supersimilarity; MZ/DZ: monozygotic/dizygotic twins; PofOm: parent-of-origin methylation; RRBS: reduced-representation bisulfite-seq; DMR: differentially methylated region; ERV: endogenous retrovirus; ESCs: embryonic stem cells. See Methods for further details.

**Table 4. Methylation quantitative trait loci (mQTL) associated with SoC-CpGs and controls**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| CpG set | Number of CpGs with mQTL | Number of mQTL (*cis*/*trans*) | Median number of mQTL per CpG (IQR) | Methylation variance explained1 |
| SoC-CpGs | 130 (50%) | 2771 (2549/222) | 6 (2-30) | 0.09 (0.08-0.15) |
| Matchedcontrols | 201 (78%) | 7886 (7417/469) | 15 (4-50) | 0.09 (0.06-0.21) |
| Randomcontrols | 50 (19%) | 1512 (1476/36) | 7 (2-35) | 0.1 (0.08-0.18) |

1 delta adjusted R2 (see Methods); IQR: inter-quartile range.

**Supplementary Files**

Appendix 1 contains Supplementary Figures 1-17

Supplementary File1 contains Supplementary Tables 1a-1s

1a ENID SoC-associated CpGs (no amplitude filter applied)

1b Seasonal amplitude tests

1c Inter-cohort change in SoC amplitude at ENID SoC-associated CpGs

1d SoC-CpGs (ENID SoC-associated CpGs with SoC methylation amplitude >=4%)

1e SoC-CpG clusters and singletons

1f SoC-CpG cluster sizes

1g SoC-CpG enrichment for MEs and overlap with ME sub-classes

1h SoC-CpG enrichment for gDMRs and parent-of-origin specific methylation

1i SoC-CpGs overlapping maternally methylated germline DMRs

1j SoC-CpGs mQTL and their association with SoC

1k Candidate genes previously associated with periconception / gestational exposures

1l Look-up of SoC-CpGs in the EWAS Catalog

1m EWAS Catalog data grouped by trait

1n SoC-CpG associations with selected variables measured in the ENID 2yr dataset

1o Look-up of SoC-CpGs in the GWAS Catalog

1p ENID (2yr): Association test p-values to detect potential residual confounding

1q EMPHASIS (7-9yr): Association test p-values to detect potential residual confounding

1r ENID 5-7yr: Association test p-values to detect potential residual confounding

1s ENID (2yr): Cell count, batch, village sensitivity analyses