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B-vitamin mediated gene regulation in vascular smooth muscle cells

Folate deficiency promotes differentiation of vascular smooth muscle cells without affecting the methylation status of regulated genes

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Abstract

Elevated serum homocysteine, an intermediate of cellular one-carbon metabolism, is an independent risk factor for cardiovascular disease (CVD). Folate deficiency increases serum homocysteine and may contribute to CVD progression. Vascular smooth muscle cells regulate vascular contractility, but also contribute to repair processes in response to vascular injury. Nutritional deficiencies, like folate deficiency, are thought to impact on this phenotypic plasticity, possibly by epigenetic mechanisms.

We have investigated the effect of folate deficiency on vascular smooth muscle cells in two cell culture systems representing early and late stages of smooth muscle cells differentiation. We find that folate deficiency promotes differentiation towards a more contractile phenotype as indicated by increased expression of respective marker genes. However, microarray analysis identified markers of striated muscle as the predominant gene expression change elicited by folate deficiency. These changes are not merely a reflection of cell cycle arrest, as foetal calf serum restriction or iron deficiency do not replicate the gene expression changes observed in response to folate deficiency. Folate deficiency only has a marginal effect on global DNA methylation. DNA methylation of CpG islands associated with genes regulated by folate deficiency remains unaffected. This supports our earlier findings in a mouse model system which also did not show any changes in global DNA methylation in response to folate and vitamin B6/B12 deficiency.

These data suggest that folate deficiency enhances the expression of smooth muscle marker gene expression, promotes a shift towards a skeletal muscle phenotype, and does not regulate gene expression via DNA methylation.

Keywords

B-vitamins, folate, vascular smooth muscle cells, DNA methylation, gene regulation

Abbreviations

cardiovascular disease: CVD, estrogen receptor alpha: ERα, Hcy: homocysteine, MTHFR: methylene-tetra hydro folate reductase, RIA: radio-immuno assay,

1. Introduction

Hyperhomocysteinemia is a risk factor for all facets of atherosclerotic disease including stroke, coronary heart disease and peripheral artery disease [1]. Homocysteine is generated as an intermediate of cellular one carbon metabolism. Folate (also known as vitamin B9), vitamin B6 and vitamin B12 are all essential components of cellular one carbon metabolism, either as substrate or as enzyme co-factors (**Fig. 1**). Single nucleotide polymorphisms or mutations of genes involved in one carbon metabolism are significant predictors of vascular disease. In general a lower activity of these enzymes (e.g. MTHFR, methionine synthase or cystathionine-b-synthase) is associated with a higher risk of vascular disease [2–5].

At present it is unclear whether homocysteine acts as a direct promoter of cardiovascular disease or whether it is mainly an indicator of low B-vitamin status. Numerous studies in animal model systems have assessed the potential mechanisms by which homocysteine might directly influence cardiovascular disease risk [6]. These experiments typically used diets deficient in folate, vitamin B6 and B12 and rich in methionine [7,8] or drinking water enriched with methionine or homocysteine [9]. Some of these studies have indeed demonstrated that indicators of vascular disease respond directly to serum homocysteine concentrations [10-12]. However, in other studies an increase in serum homocysteine was unrelated to the formation of cardiovascular lesions, whereas a reduction in serum B-vitamin levels was [7,8,13,14]. This suggests that B-vitamin deficiency may promote cardiovascular disease risk by mechanisms other than causing hyperhomocysteinemia. This is further supported by findings from several large-scale human intervention trials [15,16] using supplementation regimes containing folate, B6 and B12 [16]. Typically supplementation of patients with folate and B-vitamins was successful in reducing serum homocysteine, but ineffective in preventing cardiovascular mortality [15]. This suggests that the major value of homocysteine may be as a marker of B-vitamin deficiency, rather than a mechanistic agent in vascular disease development.

One alternative hypothesis explaining the role of B-vitamin deficiency in vascular disease suggests that a lack of methyl groups available for DNA and histone methylation may alter epigenetic gene regulation in the vascular system [17]. Support for this hypothesis comes from animal experiments which show that alterations in folate supply in pregnancy leads to changes in DNA methylation in "metastable" gene alleles and subsequent changes in gene expression and phenotype [18–20]. As these epigenetic mechanisms are important for the maintenance of differentiated phenotypes, it has been suggested that a shortage in dietary methyl donors may impact on the maintenance of the differentiation status of vascular smooth muscle cells and other cell types [21]. Patients with cardiovascular disease, who suffered from hyperhomocysteinemia and were given a folate supplement (15 mg oral methyltetrahydrofolate a day for 8 weeks) showed an increase in the methylation of imprinted genes suggesting that folate supply may influence the DNA methylation status of genes [22].

Vascular smooth muscle cells (VSMC) display a flexible phenotype in that they are, under normal circumstances, quiescent, contractile and do not synthesize extra-cellular matrix or matrix-modifying enzymes [10,23,24]. In the context of vascular injury VSMC become mobile, proliferative and secrete extra-cellular matrix proteins, possibly adopting a less differentiated phenotype. In VSMC isolated from New Zealand White rabbits a more proliferative phenotype is associated with a reduction in global DNA methylation [17,25]. In human atherosclerotic lesions global DNA methylation is also decreased relative to normal tissue [25]. However, it is unclear, whether this reduction in DNA methylation is a prerequisite for increased proliferation. Alternatively, the most proliferative (and by inference, least differentiated) cells in the population of isolated smooth muscle cells may be those with a reduced DNA methylation status.

We have used two in vitro systems of smooth muscle cell differentiation to determine the influence of B-vitamin supply on differentiation status. One model, A404, represents early developmental stages of smooth muscle differentiation and one model, A7r5, represents a partially differentiated state of vascular smooth muscle cell development. We find that folate deficiency impairs cell proliferation, increases homocysteine secretion and only leads to marginal changes in global DNA methylation. B-vitamin deficiency promotes, rather than prevents, the expression of smooth muscle cell marker genes. However, a microarray analysis of folate deficient VSCM suggest a phenotypic shift towards a skeletal muscle phenotype. The DNA methylation status of the deficiency responsive genes is not affected. This suggests that the methylation of CpG islands in the vicinity of normal genes (in contrast to metastable alleles) is not responsive to B-vitamin status.

2. Material & Methods

2.1. Cells

A404 cells [26], a gift of Prof Gary Owens, University of Virginia, are derivatives of the mouse embryonic carcinoma cell line P19. The cells can be differentiated into several lineages, including a smooth muscle cell lineage, in the presence of retinoic acid. The cell line also carries a puromycin resistance gene under the control of the smooth muscle specific myosin heavy chain 11 (MYH11) promoter. Selection of retinoic acid treated cells in medium containing puromycin permits survival of cells which have differentiated towards the smooth muscle cell lineage but removes cells which have differentiated into other lineages (e.g. neuronal). A404 cells were grown in alpha-MEM medium (Sigma, M8042) supplemented with 7.5% of foetal calf serum (Hyclone), 200µg/ml glutamine (Life Technologies) and 100U/ml penicillin/streptomycin (50U/ml penicillin and 50µg/ml streptomycin; Life Technologies) and 200µg/ml hygromycin B (Roche). Cells were split 1:50 every 2 or 3 days and used for experiments up to passage 20. Differentiation of cells was induced by the addition of 1µM all-trans retinoic acid (Sigma) for 3 days. In some experiments retinoic acid treated cells were additionally treated with $0.5\mu g/ml$ puromycin (Invivogen) for another 3 days. Rat A7r5 cells (ECACC catalogue number 86050803) were grown in DMEM (Sigma, D-6171) supplemented with 10% foetal calf serum (Sigma), 2mM glutamine and penicillin/streptomycin at 37°C and 5% CO₂ in a humidified incubator.

In order to generate folate deficient cells, custom made alpha-MEM and DMEM medium was used (BioConcept). The custom-made medium is equivalent to the normal medium but does not contain folic acid, vitamin B6 (pyridoxine-HCl; concentration 4µg/ml or 23µM in DMEM, 1µg/ml or 5.9µM in alpha-MEM), vitamin B12 (cobalamin; absent in DMEM, 1.36µg/ml or 3.4µM in alpha MEM) and methionine (30mg/l or 200µM in DMEM, 15mg/l or 100µM in alpha-MEM), which can be added separately. While normal alpha-MEM contains both, vitamin B6 and B12, DMEM does not contain vitamin B12. However, sufficient amounts of vitamin B12 are contained in foetal calf serum. A reduced folate medium (indicated as 10F, containing a folate concentration equivalent to human serum) contained 10ng/ml of folic acid. A folate deficient medium (indicated as F) contained no folic acid. A medium deficient in folic acid and vitamin B6 and B12 is indicated as FB. All media were used in conjunction with conventional (i.e. non-dialysed) FCS, which contains 0.6ng/ml of folate (as determined by radio-immunoassay). The deficient medium therefore allows for a limited level of cell proliferation. Cell numbers were determined by counting in a haemocytometer.

For treatment with azacytidine cells were seeded into 6cm dishes in conventional medium (DMEM for A7r5 cells, alpha-MEM for A404 cells). 24h later cells the medium was

changed to different concentrations of azacytidine (typically 0.1, 1, 5 or 10μ M) or control medium. 72h later the medium was changed back to control medium. After a further 72h cells were counted and genomic DNA was isolated.

2.2. Biochemistry

Homocysteine was analysed using isotope dilution GC-MS (gas chromatography and mass spectrometry) largely as described [27]. Briefly, medium was harvested from treated cells. 100µl of medium was mixed with 50µl of a 50nM C13 labelled homocysteine standard solution and 25µl of a 0.1M DTT solution and incubated at room temperature for 20 min. After addition of a 2:1 acetonitrile/ethanol solution the reaction was centrifuged at 13000rpm for 5 min. The supernatant was absorbed to a Strata-X-C spe column (Phenomenex) activated with 1ml of methanol. The column was washed consecutively with 0.5ml of water and 0.5ml of methanol and dried. Homocysteine was eluted into a tube with 700µl of 5% NH₃ in 70% methanol and dried at 90°C under nitrogen. The pellet was treated with 10 μ l of 0.1M DTT and 70 μ l of 2M NH₃ for 30 min. at room temperature and again dried under nitrogen at 90°C. The dried material was then resuspended in 50µl of a 1:1 mix of MTBSTFA [N-Methyl-N-(tert-butyldimethylsilyl)-trifluoroacetamide] and acetonitrile and heated to 90°C for 20 min. The tertiarybutyldimethylsilyl (tBDMS) derivative of homocysteine was detected by GCMS analysis on a 30m x 0.32mm x 0.25 micron ZB-5MSi capillary column. Injections (1µl) were made in the split mode with a 25:1 split. The M-57 fragment ions at m/z 420 and 421 for unlabelled and 1-C13 labelled homocysteine-tBDMS are monitored under electron impact selective ion monitoring conditions (EI-SIM). Concentrations of homocysteine were calculated from peak area ratios obtained from EI-SIM fitted to a LREG curve generated from standards. Homocysteine concentrations are expressed in correlation with cell numbers (as measured by μg of DNA) to evaluate the secretion of homocysteine per cell. Typically, homocysteine concentrations in the medium of cells grown in full medium are higher (around 10μ M) than in the medium of cells grown in deficient medium (around 1-5µM). However, the number of cells in the deficient medium is substantially lower.

Folate concentrations were measured using a competitive radio-immuno-assay (RIA) assay (MP Biomedicals - SimulTRAC-S radioassay kit Vitamin Folate[¹²⁵I]; catalogue number: 06B 254932) following the supplier's instructions. Briefly, cultured cells were detached from plates using Trypsin-EDTA and an aliquot of the cells was counted in a haemocytometer. The remaining cells were pelleted by a brief centrifugation and the cell pellet was snap frozen in liquid nitrogen. For analysis the cells were thawed by addition of 500µl of standard A (blank), resuspended by vortexing and sonicated in an ultrasound water bath for 2 min. Cell culture supernatants were measured after diluting 100µl of medium with 400µl of standard A. 1ml of tracer solution containing DTT was added and the samples were incubated in a boiling water bath for 15min. The samples were then cooled to room temperature in a water bath. 200µl of antibody beads were added to each sample and incubated at room temperature for 1 hour in the dark. The beads were subsequently pelleted by centrifugation at 4°C for 10 min. and the supernatant discarded. The sample tubes were read on a gamma counter (Packard Instruments). The readings were correlated with a standard curve (range between: 1 and 20ng of folate per mI) and the folate concentration was correlated with cell number.

2.3. RNA and quantitative PCR

RNA was isolated using the Trizol reagent (Sigma) following the manufacturer's protocol (1ml of Trizol per 10^7 cells). Reverse transcription of RNA was done using MLV RNAse(-) reverse transcriptase (Promega) following the manufacturer's recommendations. 2µg of total RNA was used as template for the cDNA synthesis reaction. A one in 10 dilution of the cDNA synthesis reaction was subsequently used as template for quantitative PCR

(Applied Biosystems 7500 Fast System). At least three independent experiments were carried out for each data point and every sample was measured in triplicate.

Quantitative PCR amplifications were carried out with a final primer concentration of 0.5μ M. Oligonucleotide primers were designed using the Primer BLAST program (NCBI). The sequences, annealing temperatures and amplicon sizes of the oligonucleotides used in this study are provided in <u>table 1</u>. The quantitative PCR products were evaluated by melting point analysis and agarose gel electrophoresis. Amplifications were done at 40 cycles of 15s at 95°C, 15s at the indicated annealing temperature and 30s at 72°C. Data were collected at the end of each PCR cycle. Standard curves for all genes were generated from serial dilutions of a plasmid containing the cDNA for each gene. The crossing points obtained from the sample analysis was then correlated with the standard curves to provide a concentration of the individual PCR product. Expression of the genes was then correlated with expression of the reference genes (β -actin or GAPDH) in the same sample (expressed as pg of gene per pg of reference) and expressed as fold change from control treatments.

2.4. Microarray

RNA for microarrays was isolated from A7r5 cells grown in medium containing 2.5ng/ml, 10ng/ml or 100ng/ml of folic acid. The isolation was carried out in cells grown for 7, 14 or 21 days in the 3 different media (3 time points at 3 concentrations). 2.5ng/ml reflects a serum concentration typically seen in folate deficiency; 10ng/ml represents a folate sufficient status; 100ng/ml represents serum folate concentrations seen in response to folic acid supplement intake. 5 independent samples were analysed for each of the 9 experimental points. RNA was isolated using the Trizol reagent (Sigma) following the manufacturer's protocol, treated with RNAse-free DNAsel (Promega) for 30 min. at 37 C, extracted with phenol/chloroform and precipitated with ethanol. The quality of the RNA was assessed via the ratio of OD at 260nm and 280nm (only samples with an OD higher than 1.8 were used) and capillary electrophoresis analysis on Agilent Chips (only RNA with a RIN higher than 8 was used).

Microarray analysis was done on Affymetrix rat 230-2.0 Genechips by ServiceXS. Data from Service XS, in the CEL file format (containing the raw signal intensities) were transferred to the MadMax website (<u>https://madmax.bioinformatics.nl</u>) where data was analysed using the statistical programming language R [28] and R-libraries offered by the Bioconductor project [29]. The data were normalized using Bioconductor and a GCRMA method, a Robust Multiarray Analysis with correction for the G:C content of the oligos, and then statistically analysed using the Limma package [30], which allowed identification of the most differentially expressed genes between different conditions using a nominal *p*-value <0.05 to represent statistical significance. Microarray data were deposited at the NCBI Gene Expression Omnibus under accession number GSE125502.

2.5. DNA methylation

Genomic DNA isolation was done using the Promega Wizard kit following the supplier's instructions. For all DNA methylation analyses reference samples were generated which represent methylated and non-methylated DNA. Non-methylated control DNA was generated by whole genome amplification of genomic DNA using a Qiagen Repli-G kit. Methylated DNA was generated by treating isolated genomic DNA with SssI DNA methyl transferase and S-adenosyl methionine (SAM; as substrate) for 1h at 37 C. Additional enzyme and additional SAM were added after 30 min. The activity of SssI was monitored using bacteriophage lambda DNA as control (**suppl. Fig. 6**). BstUI restriction digestion of lambda DNA (which is not methylated) yields a total of 25 DNA fragments. Treatment of lambda DNA with SssI and SAM fully methylates the phage DNA and prevents digestion with BstUI (recognition sequence CGCG). Different relative amounts of methylated and non-

methylated DNA were mixed and assayed by LC-MS to establish a standard curve which correlates the percentage of SssI methylated DNA in the mixture and the experimentally determined % of methylated cytosine residues (**suppl. Fig. 8c**).

Global DNA methylation was measured using an LC-MS based method developed by Friso et al. [31]. Briefly, genomic DNA (1µg per reaction) was sequentially hydrolysed by digestion with nuclease P1 (2U per reaction), venom phosphodiesterase I (0.002 U per reaction) and alkaline phosphatase (0.5 U per reaction). The resulting hydrolysate was then separated by reverse phase HPLC. The 2'deoxycytidine and 5'methyl-2'deoxycytidine peaks were detected and quantified with reference to the internal standard ¹⁵N₃ 2'deoxycytidine.

Combined bisulfite restriction analysis (COBRA) was carried out as described [32]. Briefly, genomic DNA was treated with bisulfite reagent using the EZ DNA methylation kit (Zymo Research) and purified using the spin columns provided in the kit. The concentration of the converted DNA was measured using a Nanodrop device. 5 ng of bisulfite converted DNA was used as template for a PCR amplification using gene specific primer pairs. The primer pairs were designed using the Methyl Primer Express software (Applied Biosystems). Primer pairs for rat LINE1, rat estrogen receptor alpha exon 1 and rat 14-3-3 sigma (SFN1) were taken from the literature [33–35] (**table 2**). The amplified DNA was purified over a spin column (QIAGEN Qiaquick) and 100ng of PCR product were used for a digestion with a restriction enzyme whose recognition sequence carries a CG dinucleotide (including BstUI, Taq, SfaNI). Aliquots of the restriction reaction were separated on a 10% polyacrylamide gel and visualised by ethidium bromide staining. DNA bands were quantified using a Fuji Imaging station and the AIDA Imaging software.

Pyrosequencing primer combinations (PCR primer pairs and sequencing primer) were designed using the Pyromark software package (Qiagen) for the following genes: skeletal muscle myosin heavy chain (MYH2), skeletal muscle α -actin (ACTA1), prion protein dublet (PRND), thioredoxin interacting protein (TXNIP) and are shown in <u>table 3</u>. Analysis kits for the Insig1 (insulin induced gene 1; PM00547638), Id1 (inhibitor of DNA binding 1; PM00543697) and Idi1 (isopentenyl-diphosphate delta isomerase; PM00504119) genes were purchased from Qiagen. Bisulfite treated DNA was used for the PCR reaction at a concentration of 10ng/µl. Pyrosequencing was carried out using the Pyromark Gold Q24 reagents following the supplier's instructions.

2.6. Protein analysis

Cytoplasmic protein extracts for Western blot analysis were prepared as described [36]. Briefly, cells were lysed on ice in a buffer containing 25mM KPO₄ pH7.8, 8mM MgCl₂, 1mM EDTA, 1% Triton X-100 and 15% glycerol. The extracts were incubated on ice for 5 min. and then centrifuged at 4°C for 1 min. at 10000g. The supernatants were then aliquoted and stored at -80°C.

Western blot analyses were carried out after semi-dry transfer of the proteins to a nitrocellulose membrane as described [37]. Smooth muscle α -actin (ACTA2, 42kDa) and β -actin (ACTB, 42kDa) were detected using monoclonal antibodies (Sigma A2547 and A5441, respectively) and a horse-radish-peroxidase linked anti-mouse serum (Cell Signalling Technologies #7076). All antibodies were used at a dilution of 1:1000. Western blots were developed using a chemiluminescent detection kit (Pierce) and the light emission recorded on a Fuji Imaging Station.

Immunohistochemistry was carried out as previously described [37].

2.7. Statistical analysis

Statistical analysis of datasets was done using one or two-way ANOVA in Microsoft Office Excel 2016 and post-hoc analysis and t-tests were carried out Graph-Pad Prism. Error bars indicate standard deviations. The number of replicates is indicated for each figure. The significance levels indicated in the figures typically refer to the comparisons between the control medium and the modified media. Principal components analysis was done using the prcomp function in R applied to scaled data, i.e. using the correlation matrix rather than the covariance matrix. Pathway analysis was carried out using the Bioconductor-TopGo (<u>https://rdrr.io/bioc/topGO/</u>), DAVID (<u>https://david.ncifcrf.gov/</u>) and ENRICHR (<u>http://amp.pharm.mssm.edu/Enrichr/</u>) software packages.

3. Results

Vascular smooth muscle cells surround the endothelium of blood vessels and regulate vascular tone in response to exogenous and endogenous signals [23]. They also play a key role in vascular repair in response to injury. During this process vascular smooth muscle cells switch from a contractile, quiescent phenotype to a proliferative, synthetic phenotype in which the cells contribute to the repair process [10].

Cellular one carbon metabolism is fueled by dietary folate and methionine (Fig. 1). Increased serum levels of the central metabolite homocysteine have been associated with vascular disease including stroke, coronary heart disease and peripheral arterial disease [38]. This is particularly obvious in the genetic disease of cystathionine B-synthase deficiency which leads to extremely high levels of serum Hcy (up to 100 μ M compared to a level of 10 μ M in healthy individuals) and death as a consequence of vascular disease in early life [39]. In addition SNPs in enzymes involved in cellular one carbon metabolism (e.g. MTHFR or methionine synthase) are associated with increased risk of atherosclerosis [40]. These findings link B-vitamin supply with vascular disease. Therefore, we set out to assess whether B-vitamin deficiency affects the switch of vascular smooth muscle cells to a pro-atherogenic phenotype.

3.1. Differentiation induces a switch in DNA methyl transferase expression in A404 cells

The differentiation capacity of vascular smooth muscle cells cell culture models is typically characterised by an increase of smooth muscle cell marker gene expression (e.g. MYH11, ACTA2, calponin, SM22/transgelin). To assess the effect of B-vitamin deficiency on the process of smooth muscle cell differentiation we first used the A404 model. This cell line is derived from mouse PC19 embryonal carcinoma cells and can be differentiated towards a smooth muscle cell lineage using retinoic acid. The cells also contain a puromycin marker gene under the control of the MYH11 promoter. Therefore cells which have differentiated can be enriched by puromycin selection [26].

As a first step we differentiated A404 cells as described and monitored the ensuing gene expression changes in smooth muscle cell marker genes ACTA2, SM22, MYH11 and the DNA methyl-transferase genes Dnmt1, 3a and 3b. Retinoic acid treatment strongly increases the expression of the smooth muscle marker genes ACTA2 (smooth muscle α -actin), SM22 (transgelin 1) and MYH11 (smooth muscle myosin heavy chain) as measured by quantitative PCR (**Fig. 2a**). This is supported by Western blot analysis and immune-histochemistry analysis of ACTA2 protein expression (**Fig. 2c and d**). At the same time expression of the maintenance DNA methyl transferase 1 (Dnmt1) is increased significantly (9 fold) whereas expression of the de novo methyl transferase 3b is significantly decreased (**Fig. 2b**). As expected, the relative induction of differentiation process is further enhanced by additional treatment of the differentiated cells with puromycin (**suppl. Fig. 1**). The differentiation process is also accompanied by a drastic (more than 1000-fold) reduction in the expression of the stem cell marker Oct4 (**suppl. Fig. 1c**). Retinoic acid treatment of A404 cells for 1, 2 and 3 days reveals a gradual increase in smooth muscle marker gene expression over time (**suppl. Fig. 1d**).

These data are consistent with published microarray data analysing the retinoic acid/puromycin driven differentiation process (**suppl. Fig. 2**) [41].

3.2. Azacytidine treatment promotes smooth muscle cell differentiation in A404 cells

Differentiation of A404 cells towards a smooth muscle phenotype is associated with Dnmt1 activation. The effect of Dnmt1 activity on the differentiation process was further assessed by treating the A404 cells with different concentrations of azacytidine (an inhibitor of Dnmt1 activity). Azacytidine treatment was initiated 24h before the 3-day retinoic acid treatment.

Azacytidine treatment reduces cell viability at concentrations above 0.1µM. But cells remain viable even at 10µM, although they divide much more slowly (Fig. 3a). Global DNA methylation in A404 cells as measured by liquid chromatography-mass spectrometry (LCMS) is not significantly affected by the azacytidine treatment (Fig. 3b). Expression of the smooth muscle marker genes α -actin and SM22 is affected by azacytidine treatment at higher concentrations. 10µM azacytidine increases expression levels of both marker genes in both, the untreated and the retinoic acid treated cells (Fig. 3c and d). Lower concentrations of azacytidine also increase expression of α -actin and SM22 in retinoic acid treated A404 cells, albeit to a lesser degree (Fig. 3d). Azacytidine has modest effects on the expression of Dnmt1 and Dnmt3b at higher concentrations (Fig. 3c, d, e). Combined treatment of cells with 10µM azacytidine and retinoic acid increase the expression of the stem cell marker Oct4 (Fig. 3e). Treatment of A404 cells with aza-dCTP (rather than azacytidine) shows a similar increase in α -actin and SM22 expression in A404 cells in the absence of retinoic acid induction (data not shown). Aza-dCTP is more toxic than azacytidine and the effects of smooth muscle cell marker expression are seen at a lower concentration (0.5µM aza-dCTP, rather than 10µM azacytidine). These data suggest that although Dnmt1 expression in A404 cells is increased in response to retinoic acid treatment this does not increase global DNA methylation and does not appear to be a prerequisite for differentiation.

3.3. Folate deficiency promotes A404 smooth muscle cell differentiation

In order to assess the impact of folate, vitamin B6 and vitamin B12 supply on the phenotypic plasticity of smooth muscle cells, A404 cells were grown in media containing various concentrations of folate, either in the presence of vitamin B6 and B12 or in their absence. Cultured cells are typically grown in medium containing a vast excess of folate (4000ng/ml). In order to study the effect of folate and B-vitamin deficiency custom made medium with a maximum folate supplementation of 1000ng/ml was used.

Growth of A404 cells is impaired significantly in folate free medium containing vitamins B6 and B12 (**Fig. 4a**) or devoid of B6 and B12 (**Fig. 4b**). However, the cells still proliferate in this medium, albeit slowly, due to the presence of between 0.5 to 1ng/ml of folate in foetal calf serum (typical concentration 0.6ng/ml). The use of dialysed serum completely blocks cell proliferation in folate free medium (data not shown). Cellular folate stores are severely depleted after an incubation period of 6 days in folate free medium (**Fig. 4c**). Therefore, homocysteine secretion per cell is significantly elevated in response to folate restriction (homocysteine is an intermediate of cellular one-carbon metabolism which can be regenerated to methionine in the presence of folate; **Fig. 1**). However, overall homocysteine concentrations in the medium are higher (around 10 μ M) in cells grown in normal medium compared to cells grown in deficient medium (around 1 μ M). No significant impact on global DNA methylation can be detected under any of these conditions (**Fig. 4e**).

The expression of the smooth muscle cell marker genes ACTA2, SM22 and MYH11 is significantly increased in cells exposed to folate and B-vitamin deficiency, even in the absence of retinoic acid (**Fig. 4f**). The differentiation process induced by retinoic acid treatment is significantly enhanced in folate/B-vitamin deficient cells (**Fig. 4g**). Expression of

Dnmt genes and their activation by retinoic acid are not significantly affected by folate/Bvitamin deficiency (data not shown). These data suggest that B-vitamin deficiency enhances, rather than inhibits, the differentiation of A404 cells in response to retinoic acid treatment. This conclusion is valid irrespective of the absolute degree of A404 smooth muscle cell differentiation.

3.4. Folate deficiency promotes vascular smooth muscle cell differentiation in A7r5 cells

To analyse whether the responses seen in A404 cells, which represent an early stage of differentiation also apply to a more differentiated vascular smooth muscle cell line, we analysed the bona fide rat vascular smooth muscle cell line A7r5. A7r5 cells respond to extended cultivation at confluence with characteristic morphological changes (including cell elongation, Fig. 5a), and induction of several smooth muscle marker genes like smooth muscle α -actin (ACTA2) and smooth muscle myosin heavy chain (MYH11) (Fig. 5b, c and d). A7r5 cell grown under conditions of folate deficiency show a reduction in cell proliferation (Fig. 6a and b) even if folate levels are only reduced to the physiological concentration of 10ng/ml (Fig. 6b and c). Folate deficiency also leads to a decrease in the concentration of intracellular folate (Fig. 6d), and an increase in secretion of homocysteine per cell (Fig. 6e). The increase of homocysteine secretion is diminished under conditions of additional B6 deficiency in which cell proliferation and, presumably, overall cell metabolism is attenuated (suppl. Fig. 3a). Under conditions of folate deficiency an increase in global DNA methylation (as measured by LC-MS) is detectable (Fig. 6f). In conditions of combined folate and vitamin B6 deficiency this change in DNA global DNA methylation is not observed (Fig. 6f). This change is more pronounced in the presence of vitamin B6 sufficient medium and gradually lost in decreasing medium concentrations of B6 (suppl. Fig. 3b). As in A404 cells, the expression of the smooth muscle cell marker genes MYH11 and SM22 is increased significantly in response to combined folate and B6 deficiency (Fig. 6g). No significant changes can be detected in the expression of ACTA2. A trend towards an increase in gene expression is found for Dnmt3a (Fig. 6h) which was confirmed in later experiments. Dnmt1 expression is unchanged and Dnmt3b expression was undetectable by qPCR in A7r5 cells (using our experimental set-up). A similar set of responses is also seen if folate deficiency is combined with methionine deficiency (suppl. Fig. 4).

3.5. Microarray analysis detects a shift towards skeletal muscle differentiation in response to folate deficiency in A7r5 cells

To characterise the gene expression changes in response to folate deficiency more fully, A7r5 cells were grown in medium containing 100ng/ml, 10ng/ml or 2.5ng/ml of folate and the transcriptome was analysed by an Affymetrix microarray. 100ng/ml of folate is representative of supra-physiological folate levels only detected in human serum in response to the consumption of folate supplements. 10ng/ml folate represents an average concentration of human serum folate derived from conventional dietary intake. 2.5ng/ml folate represents folate levels found in humans with folate deficiency [42].

RNA was harvested from the cells after 7, 14 and 21 days of cultivation and analysed using Affymetrix microarrays. A principal component analysis of the data demonstrates that the main differences in overall gene expression manifest themselves at day 14 and day 21, whereas fewer significant changes are observed at day 7 (Fig. 7). At day 21 there is significant variation between the transcriptome of cells grown at 2.5ng/ml and 100ng/ml, whereas the variation between cells grown at 2.5ng/ml and 10ng/ml is less pronounced. This is confirmed by Venn diagrams of the gene expression changes (for data with p<0.01 and a change bigger than 2-fold; Fig. 8). The top 20 genes regulated by folate availability (increased in 2.5ng/ml and increased at 100ng) are shown in table 4. A similar set of gene expression changes is found between cells grown at 2.5ng/ml and 10ng/ml, but the

amplitude of expression changes is much reduced. E.g. MYH2 expression is increased 11.5-fold in cells grown in 2.5ng/ml compared to cells grown in 100ng/ml. The fold change is only 6.5-fold if cell grown in 100 and 10ng/ml are compared (**Fig. 9**).

A pathway analysis was carried out using the TOP-GO, DAVID and Enrichr software suites. TOP-GO analysis ([43] evaluating the Gene Ontology-terms of the regulated genes) identifies pathways associated with cell division and cell proliferation, and cholesterol biosynthesis (GO:0006695) as processes which are significantly regulated by folate availability (<u>table 5</u>). These data confirm the key role folate plays as a one-carbon donor for DNA synthesis (<u>Fig. 1</u>).

The effect on cholesterol biosynthesis is also consistent with observations that folate supply and lipid metabolism are tightly linked [44–46]. Many of the genes which are significantly downregulated in response to folate deficiency (e.g. insulin induced gene 1 [Insig1], isopentenyl-diphosphate delta isomerase [Idi1], Steaoryl-CoA desaturase 2 [SCD2]) are involved in aspects of lipid and cholesterol biosynthesis [47]. Cholesterol biosynthesis is also the signalling pathway most affected by folate deficiency in a WikiPathways analysis (p=1.2 x 10^{-10}) in the Enrichr software suite (http://amp.pharm.mssm.edu/Enrichr/). The key role of folate supply for cell proliferation is supported by the identification of mitotic cell cycle as top term in a Reactome 2016 pathway analysis (p value: 3.2 x 10^{-21}) and PLK1 (polo-like kinase 1) signalling (a key pathway for mitosis in embryonic cells; p-value 2.4 x 10^{-12}).

A Go-term analysis of genes activated by folate deficiency using the DAVID software (https://david.ncifcrf.gov/) identifies muscle specific genes (GO:0043292 and related GOterms) with the highest enrichment score (p value 2.8 x 10^{-7}). This is consistent with the observation that 4 of the 8 genes which are most strongly induced by folate deficiency are skeletal muscle specific genes (table 4). Striated muscle contraction is also the most significant signalling pathway identified by REACTOME analysis in the ENRICHR software suite (**suppl. Fig. 5**) when a gene list with all genes responsive to folate deficiency (p<0.01; fold change >2) is analysed [48,49]. Cardiac myopathy is the disease network most closely associated with this gene list (suppl. Fig. 5b). Other genes which are activated by folate deficiency include prion protein dublet (Prnd) and inhibitor of DNA binding 1 (Id1). Both of these genes are associated with pro-angiogenic function [50–53]. Consistent with this observation the HIF-1a transcription factor network is the most significant NCI-Nature gene network activated in response to folate deficiency (ENRICHR) (suppl. Fig. 5a). Expression of several smooth muscle marker genes (including MYH11, ACTA2, SM22, MYL6L) and other muscle specific marker genes (including the transcription factors serum response factor and myocardin) are increased significantly (p<0.001), albeit by a small margin (up to around 1.5fold) in response to 2.5ng/ml folate treatment (data not shown). This suggests that the higher induction rates seen in deficient medium (0.6ng/ml, Fig. 6g) are elicited by the lower concentration of folate in the medium. Genes encoding enzymes of the cellular one carbon metabolism are only marginally changed by low folate. The biggest significant changes are found for dihydrofolate reductase (DHFR, down-regulated by 1.9-fold in 2.5ng/ml vs 100ng/ml of folate, p<0.001) and methylene-tetrahydrofolate dehydrogenase (MTHFD2, upregulated by 2-fold in 2.5ng/ml vs 100ng/ml of folate, p<0.001).

Taken together the microarray results confirm that smooth muscle cell markers are increased in response to folate deficiency, but that genes associated with a skeletal muscle phenotype are increased more strongly, together with genes associated with angiogenesis.

3.6. Folate deficiency induced gene expression changes are not caused by cell cycle arrest

The expression of the 8 most responsive genes (4 most highly activated in response to high folate and 4 most highly activated in response to low folate) was confirmed using quantitative PCR (data not shown). To assess whether the changes in gene expression were merely due to cell cycle arrest in response to folate deficiency, A7r5 cells were exposed to 2

other stimuli known to reduce cell proliferation. The cells were grown in medium containing 0.5% FCS or supplemented with 0.01µM of the iron chelator deferoxamine (DFO). Both treatments significantly reduce cell proliferation (**Fig. 10a and b**) similar to folate restriction (**Fig. 6c**). Expression of 7 genes regulated by folate deficiency (4 downregulated by low folate: TXNIP, IDI1, ID1 and INSIG1, 3 upregulated by folate deficiency: MYH2, ACTA1, and PRND) was measured in response to FCS reduction and iron deficiency using quantitative PCR. Expression of IDI1 and ID1 was significantly reduced in response to FCS reduction (as under conditions of folate deficiency) and INSIG1 expression was significantly reduced in response to FC deficiency (albeit much less than in response to F deficiency). In the case of TXNIP1, MYH2, ACTA1 and PRND only folate deficiency elicited the characteristic gene expression changes, whereas iron deficiency and low FCS did not. This suggests that most of the gene expression changes observed in the microarray are specific to folate deficiency rather than a reflection of cell cycle arrest. During this analysis it was also found that expression of Dnmt3a was increased in response to folate deficiency (but not iron deficiency or low serum, **Fig. 10d**) confirming the trend seen in **Fig. 6h**.

3.7. Folate deficiency induced gene expression changes are not associated with changes in gene specific DNA methylation

To assess whether the observed gene expression changes are correlated with gene specific changes in DNA methylation, A7r5 DNA derived from folate deficient and iron deficient cells and cells grown in low serum (0.5%) were analysed using combined bisulfite restriction analysis (COBRA) and pyrosequencing. Reference points for non-methylated DNA and methylated DNA were established by whole genome amplification and DNA methyltransferase SssI treatment, respectively, and used to establish standard curves for methylation frequency in all experiments (**suppl. Fig. 6a**).

Whole genome amplified A7r5 DNA only shows 0.2% of all cytosine residues being methylated when analysed by LC-MS [31] (**suppl. Fig. 6d**). In contrast in untreated A7r5 DNA 3.1% of all cytosine residues are methylated. Treatment of naïve A7r5 DNA with SssI and SAM increases the frequency of methylated cytosine residues to 5% (**suppl. Fig. 6d**). In contrast whole genome amplified DNA treated with SssI only contains 1.2% of cytosine residues in a methylated state. This suggests that SssI treatment has limited efficiency on non-methylated genomic DNA (**suppl. Fig. 6d**). Genomic DNA derived from azacytidine treated A7r5 cells show a progressive global demethylation as assessed by LC-MS (**suppl. Fig. 7**).

Gene specific methylation of LINE1 and estrogen receptor alpha was measured in genomic DNA derived from folate deficient (and control) A7r5 cells using COBRA (Fig. 11 and suppl. Fig. 8 and 9). Methylation of LINE1 was marginally increased in response to folate deficiency (Fig. 11e and suppl. Fig. 8c and d). It remained unchanged in response to cell cycle arrest inducing treatments of A7r5 cells in 0.5% FCS and DFO treatment (Fig. 11f). Similarly, methylation of ER α was marginally increased in response to folate deficiency (suppl. Fig. 9d). No significant change in DNA methylation was found for the tumour suppressor gene 14-3-3 sigma (SFN, data not shown) [35]. These marginal DNA methylation of the CMV promoter in A404 cells compared with HEK293 cells, which correlates with the expression rates of a CMV promoter driven nuclear GFP protein (suppl. Fig. 10). Methylation of a BstUI site in the Prnd gene, which is strongly activated in response to folate deficiency, is significantly decreased in response to folate deficiency, but not in response to reduced serum and iron deficiency (DFO treatment) (Fig. 12).

To get a more comprehensive overview over gene specific methylation in A7r5 cells in response to folate deficiency, iron deficiency and restricted FCS, 7 of the genes strongly regulated by folate deficiency were assessed for gene specific DNA methylation using pyrosequencing. CpG islands were identified using the Methyl Primer Express software (Applied Biosystems) and confirmed using the CpG finder software. All 7 genes carry a CpG island in the vicinity of the gene (**suppl. Fig. 11**). Pyrosequencing assays for these areas were developed using the pyromark software (for Myh2, Acta1, Prnd, Txnip) or purchased as kits from QIAGEN (for Id1, Idi1 and Insig1). The overall methylation rate of the gene sequences is highly variable ranging from 5% for ID1 to 90% for PRND. This may reflect the position of the CpG islands relative to the transcriptional start site (table 3) which is a known determinant of genome methylation [54]. The results do not show a significant change in gene specific DNA methylation for any of the genes in response to the different treatments (Fig. 13). Control reactions using mixtures of whole genome amplified DNA and SssI treated DNA demonstrate that all sites tested are amenable to the presence and absence of DNA methylation (as shown for the Prnd gene in Fig. 13g). In contrast, a significant (albeit small) decrease in DNA methylation was detected in the Prnd gene in response to azacytidine treatment (suppl. Fig. 12). Taken together with the change in DNA methylation found in the COBRA analysis, this suggests that the Prnd gene may be most susceptible to gene regulation changes by DNA methylation.

4. Discussion

We have analysed the effects of folate deficiency on the phenotypic plasticity of vascular smooth muscle cells. Our results demonstrate, firstly, that folate deficiency induces significant changes in global gene expression patterns indicating that the cells adopt a more contractile phenotype. Secondly, the overall expression pattern suggests a transdifferentiation towards a skeletal muscle phenotype. Thirdly, folate deficiency mediated alterations of cellular one-carbon metabolism do not induce DNA methylation changes in CpG islands associated with folate responsive genes.

Folate deficiency is associated with several disease phenotypes. The most important of these is neural tube defects [55,56] which has led to the introduction of fortification of flour with folic acid in some countries [57]. The caveat to this approach is the promotion of cell proliferation which can have a small but measurable cancer promoting effect [58]. This is not surprising as many cancer drugs (e.g. methotrexate; see **Fig. 1**) act as anti-folates interfering with folate-supported DNA synthesis. Another well characterised disease outcome of folate deficiency is colon cancer which has been documented in animal studies and epidemiological observations in human populations [59]. Finally, folate deficiency (and one of its physiological outcomes, increased serum homocysteine levels) is associated with an increased risk of cardiovascular disease [60,61]. The mechanisms by which B-vitamin deficiency impacts on CVD risk, however, are unclear.

Numerous studies have investigated potential direct effects of homocysteine on vascular health. Several mechanistic explanations have been put forward including a promotion of inflammation, an increased proliferation of VSMC in the neointima, a toxic effect on the vascular endothelium, and cell signalling alterations [62]. Some studies in model systems seemed to support a mechanistic role for homocysteine in cardiovascular disease progression by promoting VSCM proliferation [11]. However other studies (including our own) failed to establish a conclusive link between serum homocysteine concentrations and vascular disease progression [7,8,13]. Specifically, our data demonstrate that plaque formation is not correlated with serum homocysteine levels. Under conditions of combined folate and B-vitamin deficiency serum homocysteine levels of around 50µM were detected, whereas folate deficiency alone only increased serum homocysteine to 15µM. In contrast, significant increases in plaque formation were detected in folate deficient, but not in

combined folate/B-vitamin deficient mice [8]. This is supported by another rodent model system, which specifically demonstrated that provision of high methionine seems to be the predominant driver of vascular disease and not high homocysteine levels [13]. These data argue against a direct link of serum homocysteine levels with the extent of vascular plaque formation.

Moreover, the finding that B-vitamin supplementation in cardiovascular patients reduces serum homocysteine, but appears to be of no clinical benefit has led to the suggestion that homocysteine may be a marker for B-vitamin deficiency, rather than an active agent in vascular disease progression [15,16]. These findings may be explained in two ways. Firstly, homocysteine may have no further effect on disease progression, once overt cardiovascular disease is present. Alternatively, homocysteine lowering may only be clinically effective in the small group of patients which suffer from severe hyperhomocysteinemia (i.e. serum homocysteine levels of higher than 100μ M; incidentally this is the concentration which is typically used in rodent and in vitro models of hyperhomocysteinemia) [62]. Therefore, other mechanistic links between B-vitamin deficiency and cardiovascular disease risk are being considered.

Hiltunen et al. observed a reduction in global DNA methylation of vascular smooth muscle cells in association with cardiovascular disease in a mouse/rat model [25,63]. This led to the hypothesis that B-vitamin deficiency may exert its effect on cardiovascular disease via epigenetic mechanism like DNA methylation or histone methylation. A potential link between folate supply, DNA methylation, and gene expression has been demonstrated e.g. for the agouti mouse model and related "metastable" alleles [64]. We have therefore studied the effects of folate, vitamins B6 and B12, and methionine deficiency in two complementary cell culture models representing early and late differentiation stages of vascular smooth muscle cells.

Our results allow two major conclusions. Firstly, folate deficiency does not prevent the establishment or maintenance of a differentiated vascular smooth muscle phenotype. In fact, the expression of smooth muscle cell marker genes is increased in response to folate deficiency in both, A404 and A7r5 cells. However, the microarray analysis of A7r5 cells suggests a shift of overall gene expression towards a skeletal muscle phenotype. Transdifferentiation on A7r5 cells into skeletal muscle has been observed before as a spontaneous outcome of cultivation but was only characterised by expression of two skeletal muscle proteins [65]. Our results suggest that an exogenous stressor like folate deficiency can promote this process. In addition, several genes involved in tissue repair are also activated in response to folate deficiency, while genes involved in cholesterol and lipid metabolism are downregulated. This is also reflected for the overall transcriptome by pathway analyses. The close link between B-vitamin/one carbon metabolism and lipid metabolism is well characterised and supports the findings of other researchers [44-46]. Trans-differentiation of VSMC in vivo has been documented [66,67] but is controversial [68,69]. The most relevant trans-differentiation processes observed in vivo appear to be towards an osteogenic and macrophage (rather than a skeletal muscle) lineage [66,67].

We have previously studied the effects of folate and vitamin B6 and B12 deficiency in the arterial tissue of folate deficient apoE-/- mice using proteomics [8,70]. Some of the changes seen in that analysis are also reflected in the A7r5 cells. E.g. the transferrin receptor, the electron-transfer flavoprotein, lactate dehydrogenase, glutathione-Stransferase mu1, several fibrinogen polypeptides are all significantly upregulated in conditions of folate deficiency in both analysis systems. The fact that the major changes seen in A7r5 cells are not seen in the whole aorta is presumably due to the different sensitivities of the two analyses (microarray vs proteomics, the latter only revealing changes in highly expressed proteins) and the modest contribution of smooth muscle cells to the overall tissue composition of a complete aorta.

The second conclusion from the data presented is that changes in gene expression are not correlated with significant changes in DNA methylation of CpG islands associated with genes regulated by folate supply. This is in accordance with recent findings by McKay et al. who found that gene expression and DNA methylation changes in the liver of folate deficient mice were not correlated for most genes [71]. In addition, Wang et al. investigated the role of DNA methylation in a model of neural tube defects, and found that the expression of very few genes was changed as a consequence of DNA methylation changes [72]. In contrast, Leclerc et al. found small but significant increases in the methylation of CpG islands associated with tumour suppressor genes in colon tumours induced by folate deficiency in C57B/6 mice [73]. However, even in that case the DNA methylation changes detected (by pyrosequencing) were in the range of a few percent; similar to the amplitude of changes we detect in some of the assays, especially for the Prnd gene. The available data suggest that while the methylation status of some genes is sensitive to environmental changes (like folate supply) for most genes no significant methylation changes are found. In order to conclusively assess the specific effect of environmental changes on DNA methylation a whole genome sequencing of bisulfite converted DNA would be necessary.

Recent technological advances have enabled the site directed modification of DNA methylation using fusion proteins of Cas9 with Dnmt3a or Tet1 [74–76] in which the methylating or demethylating activities are targeted to sites specified by guide RNAs interacting with Cas9. While these experiments clearly demonstrate a link between the methylation of target genes with their expression, it is possible that these effects only apply to a subset of genes. Our data also show profound impacts of DNA methylation on the expression of a CMV-GFP transgene stably inserted in HEK293 cells and A404 cells. Taken together these data suggest that a simple correlation between gene specific DNA methylation changes resulting in altered gene expression, as is the case for metastable alleles, is not generally applicable. Alternatively, DNA methylation changes in one genomic location may have long range effects on chromosome architecture resulting in changes of expression of distant genes with consequent changes in biological phenotypes.

The results in this study also raise the question as to how important folate supply is for DNA methylation. It is probable that DNA methylation of genes is mainly malleable during early stages of development (as reflected in the A404 cells). Effects of early life nutrition on cardiovascular disease risk have indeed been demonstrated [77]. We only detected small changes in global or gene specific DNA methylation in both vascular smooth muscle cell lines in response to folate deficiency, reflecting our findings in mice [8]. Human studies have demonstrated that folate and B-vitamin supply in utero can have differing effects on the methylation of different imprinted genes [78,79]. In addition, direct analysis of gene expression correlated with these methylation changes is restricted as very few tissues which are accessible for such studies in humans (mainly lymphocytes).

DNA methylation enzymes compete for methyl groups with other cellular processes like DNA synthesis [80]. In tumour cells an increased requirement for de novo nucleotide synthesis can therefore decrease overall levels of DNA methylation. However, even a dramatic increase in the DNA de novo synthesis rate of 36% (e.g. in tumour cells) only reduces the DNA methylation rate by 0.5% [81] suggesting that DNA methylation is not particularly responsive to one carbon unit supply. In addition, histone methylation, another mechanism of epigenetic gene control, may also provide a target for gene regulatory effects of folate deficiency. It has been suggested that THF (derived from dietary folate) is required to protect lysine demethylase enzymes from the formaldehyde generated as a consequence of histone demethylation [82]. Folate deficiency may therefore impair histone

demethylation thereby impacting on gene regulation. B-vitamin deficiency may also affect cardiovascular disease risk indirectly via other processes, including DNA repair, micro RNA stress responses, inflammation and energy sensors, like mTOR [83–86] either with or without epigenetic alterations.

Conclusions

Taken together the data shown in here demonstrate that folate deficiency promotes a contractile phenotype in vascular smooth muscle cells, combined with a transdifferentiation towards a skeletal muscle gene expression profile. These expression changes are not accompanied by altered DNA methylation of the regulated genes. The data also clearly show that a simple correlation of gene specific DNA methylation changes and altered gene expression, as is the case for metastable alleles, is not generally applicable.

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Authors' contributions

Andreas Kolb designed and supervised the study, carried out experimental work, and wrote the manuscript, Linda Petrie and Lynn Pirie carried out experimental work, Claus Mayer analysed the microarray data, Susan Duthie co-designed and co-supervised the study and cowrote the manuscript.

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B-vitamin mediated gene regulation in vascular smooth muscle cells

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Figure legends:

Table 1: Oligonucleotides used for quantitative PCR. Primer sequences, annealingtemperatures and expected product sizes are given.

Table 2: Oligonucleotides used for COBRA analyses.

Table 3: Oligonucleotides used for pyrosequencing analyses and gene sequences analysed.

Table 4: Top 20 gene expression changes in response to folate deficiency. Fold change of gene expression between A7r5 cells grown in in folate sufficient (100ng/ml) and folate deficient (2.5ng/ml) medium

Table 5: TOP-GO pathway analysis of transcriptome changes in response to folate deficiency. elimKS here gives the p-values from topGO specific Kolmogorov-Smirnov Test with elimination of genes of significant GO-terms in parent terms.

Figure 1: Schematic representation of cellular one carbon metabolism. Relevant enzymes are shown: methylene tetrahydrofolate reductase (MTHFR), methionine synthase (MS), cystathionine-B-synthase (CBS), di-hydrofolate reductase (DHFR), betaine-homocysteine S-methyltransferase (BHMT). Enyzmes dependent on B-vitamins are indicated [B6], [B12]. Dietary folate is a precursor for both, DNA synthesis and DNA methylation. Methotrexate is a classic example of an "anti-folate" used in cancer therapy. It is a highly effective competitive inhibitor of DHFR.

Figure 2: Analysis of smooth muscle marker and Dnmt gene expression in A404 cells. A404 cells were treated with 1μ M all-trans retinoic acid (dissolved in DMSO; stock solution 1mM) or equivalent concentration of solvent for 3 days. RNA was isolated, reverse transcribed and analysed by quantitative real time PCR. Expression of the smooth muscle α -actin (ACTA2), SM22, myosin heavy chain (MYH11), Dnmt1, Dnmt3a and Dnmt3b genes was measured and correlated with expression of the reference gene GAPDH. Panel A: Fold induction of the smooth muscle cell marker genes ACTA2, SM22 and MYH11 in A404 cells treated with retinoic acid supplemented medium relative to control medium. Panel B: Fold induction of the DNA methyl-transferase genes Dnmt1, 3a and 3b in A404 cells treated with retinoic acid supplemented medium relative to control medium. Panel C: Western blot analysis of cytoplasmic extracts from A404 cells treated with retinoic acid supplemented medium ([+]RA) and control medium ([-]RA). Smooth muscle α -actin (ACTA2) and β -actin (ACTB) were detected using specific monoclonal antibodies. Statistical analysis was done by ANOVA (n=4). p<0.05: *, p<0.01: **, p< 0,001: ***. Panel D: Immunohistochemistry analysis of ACTA2 protein expression in untreated ([-]RA [-]puro), retinoic acid treated ([+]RA [-]puro) and retinoic acid and puromycin treated ([+]RA [+]puro) A404 cells.

Supplementary Figure 1: Changes in gene expression in response to retinoic acid treatment and combined retinoic acid/puromycin treatment in A404 cells. **Panel A:** Expression of smooth muscle marker genes smooth muscle α -actin (ACTA2), transgelin 1 (SM22) and smooth muscle myosin heavy chain (MYH11) in response to treatment with retinoic acid for 3 days (d3 [+]RA) and after treatment with retinoic acid for 3 days and treatment with puromycin for another 3 days (d6 [+]RA). Data are shown as fold change in expression from untreated cells at day 3 (d3 [-]RA). Expression data were correlated with the expression of the reference gene GAPDH. **Panel B:** Expression of ACTA2, SM22 and MYH11 correlated with the reference gene β -actin. **Panel C:** Expression of Dnmt1, 3a and 3b and Oct4 in response to

treatment with retinoic acid for 3 days (d3 [+]RA) and after treatment with retinoic acid for 3 days and treatment with puromycin for another 3 days (d6 [+]RA). Data are shown as fold change in expression from untreated cells at day 3 (d3 [-]RA). Expression data were correlated with the expression of the reference gene GAPDH. **Panel D:** Expression of ACTA2, SM22 and MYH11 in response to retinoic acid treatment for 1, 2, and 3 days. Data are shown as fold change in expression from untreated cells at day 3 (d3 [-]RA). Expression data were correlated with the expression of the reference gene GAPDH. **Panel D:** Expression of ACTA2, SM22 and MYH11 in response to retinoic acid treatment for 1, 2, and 3 days. Data are shown as fold change in expression from untreated cells at day 3 (d3 [-]RA). Expression data were correlated with the expression of the reference gene GAPDH.

Supplementary Figure 2: Expression data for the smooth muscle marker genes ACTA2 (panel A) and SM22 (panel B), the differentiation marker Oct4 (panel C) and the Dnmt1, 3a and 3b genes (panels D to F) in response to treatment of A404 cells with retinoic acid for 48h or 96h or treatment with retinoic acid for 96h with subsequent puromycin selection (96h-P). Data are derived from Spin et al. 2010 via the Gene Expression Omnibus at the NCBI. 3 different probe sets, representing variations in the expression of splice variants are shown for Dnmt3a.

Figure 3: Impact of azacytidine treatment on cell proliferation and global DNA methylation. A404 cells were grown in medium containing 0.1μ M, 1μ M or 10μ M of azacytidine or in normal medium for 6 days. 10^4 cells were seeded, and cell numbers were recorded on day 3 and 6. Genomic DNA was isolated at the end of the incubation and analysed by LC-MS. Panel A: Cell proliferation of cells grown in the different media. The number of cells was determined by trypan blue staining at the indicated time points using a haemocytometer (n=6). Panel B: Global DNA methylation. The percentage of methylated cytosine residues is shown (n=3). Panels C-E: Analysis of gene expression in azacytidine treated A404 cells. A404 cells were treated with azacytidine for 24h and then induced with all-trans retinoic acid or equivalent concentration of solvent for 3 days in the continued presence of azacytidine. RNA was isolated, reverse transcribed and analysed by quantitative real time PCR. Expression of the smooth muscle α -actin (ACTA2), SM22, Dnmt1, Dnmt3b and Oct4 genes was measured and correlated with expression of the reference gene GAPDH in A404 cells treated with different concentrations of azacytidine. Expression is show as fold change from control cells (not treated with azacytidine). Panel C: Fold induction of the ACTA2, SM22, Dnmt1, Dnmt3b and Oct4 genes in response to azacytidine treatment without retinoic acid. Panel D: Fold induction of the ACTA2, SM22 and Dnmt1 genes in response to combined retinoic acid and azacytidine treatment. Panel E: Expression of the Dnmt3b and Oct4 genes in response to combined retinoic acid and azacytidine treatment. Statistical analysis was done by ANOVA (n=3). p<0.05: *, p<0.01: **, p< 0,001: ***.

Figure 4. Impact of folate and vitamin B6/12 deficiency on A404 cells. **Panels A-B**: Proliferation of A404 cells in different cell culture media as determined by trypan blue staining at the indicated time points using a haemocytometer. Cells were grown in medium containing 100, 100, 10 or Ong/ml of folate either in the presence of B-vitamins (**Panel A**) or in their absence (**Panel B**). The folate deficient medium (Ong/ml) contains residual folate derived from foetal calf serum at a final folate concentration of 0.6ng/ml. 2*10⁴ A404 cells were seeded into 6 cm dishes and grown for 7 days. Cells were counted after 3, 5, and 7 days. The average of 4 independent experiments is shown. **Panel C**: Intracellular folate concentration in cells grown in folate deficient (F; Ong/ml) or folate and B-vitamin deficient (FB) media. Concentrations determined by radio-immunoassay are shown as pg folate per 10⁴ cells. **Panel D**: Homocysteine concentrations in cell culture medium as measured using a GC-MS method. Concentrations in nM per μ g of DNA (isolated from the underlying cell layer) were calculated and the results are shown as percentage of homocysteine in control medium. **Panel E:** Global DNA methylation. The percentage of methylated cytosine residues

is shown in medium containing 0, 10, 100 or 1000ng folic acid per ml either in the absence (FB) or presence of vitamins B6 and B12 (F). **Panel F:** Expression of the smooth muscle cell marker genes ACTA2, SM22 and MYH11 in response to F and FB media correlated to expression of the reference gene GAPDH. Results are shown as fold change of expression relative to cells in control medium (CON). **Panel G:** Expression of the smooth muscle cell marker genes ACTA2, SM22 and MYH11 in response to retinoic acid treatment in F and FB media relative to expression of the reference gene GAPDH. Results are shown as fold change of expression of the reference gene GAPDH. Results are shown as fold change of expression relative to untreated cells (i.e. in CON medium in the absence of retinoic acid). Measurements (n=4) were analysed by one-way ANOVA. p<0.05: *, p<0.01: **, p< 0,001: ***.

Figure 5. Differentiation of A7r5 rat smooth muscle cells. **Panel A:** Micrographs of A7r5 cells at exponential growth and 3 days after reaching confluence. Cells were photographed at a magnification of 40x. Scale bar: 100um. **Panel B:** Western blot analysis of protein extracts from growing and confluent cells using antisera directed against β -actin (ACTB) and smooth muscle α -actin (ACTA2). Extracts from HEK 293 cells (which do not express ACTA2) were used as negative control. **Panel C:** Abundance of ACTA2 protein relative to ACTB protein in proliferating and confluent A7r5 cells as quantified by densitometry. **Panel D:** Expression of smooth muscle specific genes SM22, smooth muscle α -actin (ACTA2), smooth muscle myosin heavy chain (MYH11), brain specific creatine kinase (CKB) and muscle specific creatine kinase (CKM) in A7r5 cells 3 days and 10 days after reaching confluence. Expression is shown as percentage of expression in proliferating cells. Statistical analysis (n=3) was done by ANOVA. p<0.05: *, p<0.01: **, p< 0.001: ***.

Figure 6: Cellular responses to folate (F) and combined folate and B6 deficiency (FB) in A7r5 cells grown in the respective media for 21 days. Panel A: Representative microscopy pictures of A7r5 cells grown in full medium (CON), folate deficient medium (F; containing a residual concentration of 0.6ng folate per ml derived from serum) and medium deficient in folate and vitamins B6 (pyridoxal) and B12. Cells were photographed at a magnification of 100x. Scale bar: 40um. Panel B: Impact of different medium folate concentrations (from 0 to 100ng/ml) on cell growth over a period of 21 days (as determined by trypan blue staining at the end of the incubation period using a haemocytometer). Panel C: Growth of A7r5 cells in medium containing 10ng/ml of folate (10F), no folate (F), no folate and no vitamin B6 (FB), or control medium (CON; 100ng/ml of folate) over an 18-day period as determined by trypan blue staining at the indicated time points using a haemocytometer. Panel D: Cellular folate concentrations of cells grown in control or F or FB medium as measured by RIA. Panel E: Homocysteine concentration in medium of cells grown in control or F or FB medium (measured by LC-MS) expressed as nMol of homocysteine per μ g of DNA (derived from the underlying cells). Panel F: Global DNA methylation as measured by LC-MS. Results are shown as percentage of methylation in CON medium. Panel G: Expression of the smooth muscle cell marker genes ACTA2, SM22 and MYH11 in response to F and FB medium correlated with expression of the reference gene GAPDH. Results are shown as fold change of expression relative to control cells. Panel H: Expression of Dnmt1 and Dnmt3a. Measurements (n=4) were analysed by one-way ANOVA. p<0.05: *, p<0.01: **, p< 0,001: ***.

Supplementary Figure 3: Cellular responses to different combinations of folate and vitamin B6 content of cell culture medium. Folate concentrations were 4000ng/ml (F:4000), 10ng/ml (F:10) or 0ng/ml (F:0; equating to 0.6ng/ml when the residual folate in serum is taken into account). Pyridoxal/B6 concentrations were 23μ M (B6: 23, 0.1 μ M and 0 μ M. **Panel A:** Homocysteine content of medium derived from A7r5 cells grown in different combinations of folate and vitamin B6 as measured by LC-MS expressed as nMol of homocysteine per μ g

of DNA (derived from the underlying cells). **Panel B:** Global DNA methylation as measured by LC-MS. Measurements (n=4) were analysed by one-way ANOVA. p<0.05: *, p<0.01: ***, p< 0,001: ***.

Supplementary Figure 4: Cellular responses different combinations of folate and methionine content of cell culture medium. Folate concentrations were 4000ng/ml, 10ng/ml or 0ng/ml (equating to 0.6ng/ml when the residual folate in serum is taken into account). Methionine concentrations were 200μM, 20μM and 0μM. **Panel A:** Impact of different combinations of folate and methionine on cell growth over a period of 11 days. **Panel B:** Homocysteine content of medium derived from A7r5 cells grown in different combinations of folate and methionine as measured by LC-MS expressed as nMol of homocysteine per μg of DNA (derived from the underlying cells). **Panel C:** Cellular folate concentrations of cells as measured by RIA. **Panel D:** Global DNA methylation as measured by LC-MS. **Panel E:** Expression of the smooth muscle cell marker genes ACTA2, and MYH11 in response to F and FM (folate and methionine deficient) medium correlated to expression of the reference gene GAPDH. Results are shown as fold change of expression relative to control cells. Measurements (n=4) were analysed by one-way ANOVA. p<0.05: *, p<0.01: **, p< 0,001: ***.

Figure 7: Principal component analysis of a transcriptome analysis of A7r5 cells grown in medium containing 2.5ng/ml (shown in green), 10ng/ml (shown in orange) or 100ng/ml of folate (shown in red) for 7, 14 and 21 days. The first principal component separates the data by day of analysis; the second principal component separates the data by folate concentration.

Figure 8: Venn diagrams of the genes changed during the different incubation periods and the different folate concentrations. Genes included were significant with a p-value < 0.001 and a fold change of > 2. Note that very few gene changes are found at time points between cells grown in 2.5ng/ml and 10ng/ml.

Figure 9: Fold change of expression for 6 exemplary genes, which are highly responsive to modulations of folate concentrations in medium. Thioredoxin interacting protein (TXNIP), insulin induced gene 1 (INSIG1) and isopentenyl diphosphate isomerase (ID11) are highly expressed in 100ng/ml of folate but downregulated in response to folate deficiency (**panels A-C**). Prion protein dublet (PRND), skeletal muscle myosin heavy chain (MYH2) and skeletal muscle α -actin (ACTA1) are induced in response to low folate concentrations (i.e. 2.5ng/ml) (**panels D-F**). Measurements (n=5) were analysed by one-way ANOVA. p<0.05: *, p<0.01: ***, p<0.001: ***.

Supplementary Figure 5: Pathway analysis using the Enrichr software suite. **Panels A-C** show the most significantly activated pathways detected using a list of genes activated >2 fold with a p-value <0.01 in response to low folate supply (2.5ng/ml vs. 100ng/ml folate at day 21). Pathways detected in the NCI-Nature, OMIM disease and Reactome databases are shown. **Panels D-F** show the most significantly activated pathways detected using a list of genes activated >2 fold with a p-value <0.01 in response to high folate supply (100ng/ml vs. 2.5ng/ml folate at day 21). Pathways detected in the NCI-Nature, other the NCI-Nature, WikiPathways and Reactome databases are shown.

Figure 10: Panel A: Growth of A7r5 cells in medium containing 10%, 0.5% and 0.1% foetal calf serum over a period of 11 days. **Panel B:** Growth of A7r5 cells in control medium and medium containing 0.01µM of the iron chelator deferoxamine (DFO). **Panel C:** Expression of

the genes ID1, INSIG1, TXNIP1 and ID11 which were found to be downregulated in response to folate deficiency. Gene expression was measured in cells grown in medium containing 0.5% serum (FCS), 0.01 μ M DFO (FE) or 0.6ng/ml folate (F). Gene expression was correlated with expression of the reference gene β -actin. Results are shown as fold change of expression relative to control cells. **Panel D:** Expression of the genes MYH2, ACTA1, and PRND which were found to be upregulated in response to folate deficiency. In addition, expression of the Dnmt3a gene was measured. Measurements (n=3) were analysed by one-way ANOVA. p<0.05: *, p<0.01: **, p<0.001: ***.

Supplementary Figure 6: Control substrates for methylation analyses. **Panel A:** Schematic representation of the approaches used to generate non-methylated and fully methylated genomic DNA. **Panel B:** Analysis of DNA methylation efficiency of the SssI DNA methyl transferase using bacteriophage lambda DNA as template. Note that untreated lambda DNA is fully digested by BstUI (yielding 25 fragments), whereas SssI treated DNA is fully protected against digestion. **Panel C:** Untreated (unmethylated) and SssI treated lambda DNA was mixed in 6 different ratios (containing 100%, 50%, 25%, 12.5%, 6.25% or 3.1% of SssI treated DNA) and cytosine methylation was measured by LC-MS. The amount of SssI treated DNA added and percentage of methyl-cytosine measured is correlated. **Panel D:** Global DNA methylation as measured by LC-MS for methylated cytosine. The percentage of methyl-cytosine (% mC) is shown for DNA derived from A7r5 cells grown in control medium (CON), in medium supplemented with 10μM azacytidine, A7r5 DNA derived from whole genome amplification (WGA) or treatment with SssI (CON[+]) or DNA derived from whole genome amplification and subsequent SssI treatment (WGA[+]). Measurements (n=3) were analysed by one-way ANOVA. p<0.05: *, p<0.01: **, p<0,001: ***.

Supplementary Figure 7: Treatment of A7r5 cells with azacytidine. A7r5 cells were seeded into 6cm dishes (n=4) in normal medium. 24h later cells the medium was changed to 4 different concentrations of azacytidine (0.1, 1, 5 or 10 μ M) or control medium. 72h later the medium was changed back to control medium. After a further 72h cells were counted and genomic DNA was isolated. **Panel A:** Cell counts after azacytidine treatment. **Panel B:** Global DNA methylation as measured by LC-MS. Data were analysed by ANOVA. Error bars indicate standard deviation. p<0.05: *, p<0.001: ***.

Figure 11: COBRA analysis of DNA methylation at LINE1 (a repetitive mobile genetic element which serves as a proxy for global DNA methylation). Panel A: Schematic representation of the LINE1 section used for DNA amplification with the primer pair LINE-1s/as (from Pogribny et al. 2006). The location of the primer binding sites, the expected PCR product and its BstUI restriction digestion products are indicated, as are the CpG nucleotides in the amplified DNA region. A BstUI site (CGCG) is maintained in methylated but not in non-methylated DNA, where it is converted to TGTG. The rate of digestion is therefore reflective of the methylation rate of both CG dinucleotides (i.e. increases in digestion signify increases in methylation). Panel B: Polyacrylamide gel analysis of BstUI digested of LINE-1 PCR product. Different ratios of non-methylated (WGA: whole genome amplification) and untreated A7r5 DNA were mixed prior to bisulfite conversion and PCR amplification (from 100:0 to 0:100). The PCR products were purified and digested with BstUI and separated on a 10% polyacrylamide gel alongside a molecular weight marker (NEB PCR marker). The sizes of the relevant DNA fragments are indicated. Panel C: Polyacrylamide gel with COBRA samples derived from A7r5 cells grown in control medium (100ng/ml folate: CON), medium containing 10ng/ml folate (10F), medium without added folate (0.6ng/ml: F), and medium containing no added folate or vitamin B6 (FB). Panel D: Quantification of DNA methylation after image analysis of the gel shown in panel B. The percentage of methylated input DNA is correlated with DNA methylation measured by BstUI digestion. **Panel E:** Quantification of DNA methylation after image analysis of the gel shown in panel C. **Panel F:** Quantification of DNA methylation in A7r5 cells treated with folate deficient medium (F), iron deficient medium (FE) or medium with 0.5% FCS (FCS) as measured by COBRA analysis of the LINE1 element. Measurements (n=3) were analysed by one-way ANOVA. p<0.05: *.

Supplementary Figure 8: COBRA analysis of DNA methylation at LINE1. Panel A: Schematic representation of the LINE1 section used for DNA amplification with the primer pair LINE-1s/as (from Pogribny et al. 2006; table 2). The location of the primer binding sites, the expected PCR product and its Rsal restriction digestion products are indicated, as are the CpG nucleotides in the amplified DNA region. The Rsal (recognition sequence GTAC) site is generated by bisulfite conversion of the original sequence GCACG to a GTACG, but only maintained if the CG dinucleotide at the end of the recognition site is maintained. Therefore, digestion of the Rsal site is reflective of an increase of methylation at this CG site. Panel B: Polyacrylamide gel analysis of Rsal digested of LINE-1 PCR product. Different ratios of nonmethylated (WGA: whole genome amplification) and untreated A7r5 DNA were mixed prior to bisulfite conversion and PCR amplification (from 100:0 to 0:100). The PCR products were purified and digested with Rsal and separated on a 10% polyacrylamide gel alongside a molecular weight marker (NEB PCR marker). The sizes of the relevant DNA fragments are indicated. Panel C: Polyacrylamide gel with COBRA samples derived from A7r5 cells grown in control medium (100ng/ml folate: CON), medium containing 10ng/ml folate (10F), medium without added folate (0.6ng/ml: F), and medium containing no added folate or vitamin B6 (FB). Panel D: Quantification of DNA methylation after image analysis of the gel shown in panel C. The percentage of methylated input DNA is correlated with DNA methylation measured by Rsal digestion. Measurements (n=3) were analysed by one-way ANOVA. p<0.05: *.

Supplementary Figure 9: COBRA analysis of DNA methylation at the first exon of the estrogen receptor alpha (ERa) gene. Panel A: Schematic representation of the ERa gene used for DNA amplification with the primer pair ERa-e1 s/as (from Monje et al. 2007, table 2) [34]. The location of the primer binding sites, the expected PCR product and its BstUI restriction digestion products are indicated, as are the CpG nucleotides in the amplified DNA region. Panel B: Polyacrylamide gel analysis of BstUI digested ERa PCR product. Different ratios of non-methylated (WGA: whole genome amplification) and untreated A7r5 DNA were mixed prior to bisulfite conversion and PCR amplification (from 100:0 to 0:100). The PCR products were purified and digested with BstUI and separated on a 10% polyacrylamide gel alongside a molecular weight marker (NEB PCR marker). The sizes of the relevant DNA fragments are indicated. Panel C: Polyacrylamide gel with COBRA samples derived from A7r5 cells grown in control medium (100ng/ml folate: CON), medium containing 10ng/ml folate (10F), medium without added folate (0.6ng/ml: F), and medium containing no added folate or vitamin B6 (FB). Panel D: Quantification of DNA methylation after image analysis of the gel shown in panel C. The percentage of methylated input DNA is correlated with DNA methylation measured by BstUI digestion. Measurements (n=3) were analysed by one-way ANOVA. p<0.05: *.

Supplementary Figure 10: DNA methylation and activity of the CMV promoter. **Panel A:** Fluorescent microscope picture of A404 cells stably transfected with the plasmid CMV-nGFP carrying a GFP gene linked to a nuclear localisation signal under the control of the CMV promoter. **Panel B:** Fluorescent microscope picture of HEK293 cells stably transfected with the plasmid CMV-nGFP. **Panel C:** Schematic representation of the CMV promoter region used for DNA amplification with the primer pair CMV-BSP-s and CMV-BSP-as. The location of the primer binding sites, the expected PCR product and its SfaNI restriction digestion products are indicated. **Panel D:** Polyacrylamide gel analysis of SfaNI digested CMV PCR product. Different ratios of non-methylated (WGA: whole genome amplification) and untreated genomic DNA (derived from transfected HEK293 cells) were mixed prior to bisulfite conversion and PCR amplification (from 100:0 to 0:100). The PCR products were purified and digested with SfaNI and separated on a 10% polyacrylamide gel alongside a molecular weight marker (NEB PCR marker). The sizes of the relevant DNA fragments are indicated. **Panel E:** Polyacrylamide gel analysis of SfaNI digested CMV PCR product amplified from genomic DNA of 4 independent stable CMV-nGFP cell clones derived from HEK293 and A404 cells.

Figure 12: COBRA analysis of DNA methylation at the rat PRND gene. **Panel A:** Schematic representation of the PRND gene section used for DNA amplification with the primer pair PP2-s and PP2-as. The location of the primer binding sites, the expected PCR product and its BstUI restriction digestion products are indicated, as are the CpG nucleotides in the amplified DNA region. **Panel B:** Polyacrylamide gel analysis of BstUI digested PRND PCR product amplified from A7r5 cells grown in control medium (100ng/ml folate: CON), folate deficient medium (F), iron deficient medium (FE) or medium with 0.5% FCS (FCS). The PCR products were purified and digested with BstUI and separated on a 10% polyacrylamide gel. The sizes of the relevant DNA fragments are indicated. **Panel C:** Quantification of DNA methylation after image analysis of the gel shown in panel B. The percentage of methylated input DNA is correlated with DNA methylation measured by BstUI digestion. Measurements (n=3) were analysed by one-way ANOVA. p<0.01: **.

Supplementary Figure 11: Schematic representation of the MYH2, ACTA1, ID1, PRND, INSIG1, ID11 and TXNIP genes regulated by folate supply in A7r5 cells. The mRNA, coding sequences (CDS), the predicted CpG islands and the regions used for pyrosequencing analysis are indicated.

Figure 13: Pyrosequencing analysis of CpG islands of the ACTA1 (panel A), MYH2 (panel B), INSIG1 (panel C), TXNIP (panel D), ID1 (panel E), IDI1 (panel F) and PRND genes (panels G and H). Genomic DNA was derived from A7r5 cells grown in low serum (0.5% FCS; indicated as FCS), iron deficiency (10 μ M DFO; indicated as FE) or folate deficiency (0.6nM folate; indicated as F). The percentage of methylated Cs at each of the polymorphism positions is indicated. Panel G: Quality control of the pyrosequencing analysis for the Prnd gene. Different ratios of non-methylated (WGA: whole genome amplification) and untreated A7r5 DNA were mixed prior to bisulfite conversion and PCR amplification (from 100:0 to 0:100) before being analysed by pyrosequencing at the Prnd gene. Data are the average of 3 independent replicates and were analysed by ANOVA. p < 0.05: *.

Supplementary Figure 12: Pyrosequencing analysis of CpG rich regions of the MYH2 (**panel A**), PRND (**panel B**), TXNIP (**panel C**), and IDI1 (**panel D**) genes. Genomic DNA was derived from A7r5 cells grown in folate and vitamin B6 deficient medium (FB) or 10mM azacytidine (as outlined in suppl. Fig. 7; AZA). The percentage of methylated Cs at each of the polymorphism positions is indicated. Data are the average of 3 independent replicates and were analysed by ANOVA. p < 0.05: *, p < 0.01: ***, p < 0.001: ***.

Table 1: Oligonucleotides used for quantitative PCR. Primer sequences, annealing temperatures and expected product sizes are given.

	primer				
gene ID	name	primer sequence	annealing	product	Genbank
rat MYH11	rsmMYH3	CTGGGGAGCTGCGTGTC	57.4°C	530bp	NM_001170600.1
	rsmMYH4	AGCGGCCTTCTCCTCATACTG			
rat CKB	rbCK1	AGAAAGGGGGCAACATGAAGGAAG	58.3°C	482bp	NM_012529.3
	rbCK2	GCACTGCCCGGGTAATAAG			
rat CKM	rmCK1	GCGTGGGCCTGCAGAAGATTGA	62.9°C	441bp	NM_012530.2
	rmCK2	TCCAGGGGGCGGGGCTCCAG			
rat TAGLN	rSM22-1	ATGGCGTGATTCTGAGCAAGTT	57°C	415bp	NM_031549.2
	rSM22-2	GGTCGCCCATAGCCTGTC			
rat ACTA2	rsm-a-actin1	CTGGCCGAGATCTCACCGACTACC	59.6°C	490bp	NM_031004.2
	rsm-a-actin2	GAGCCGCCGATCCAGACAGAATA			
rat GAPDH	rGAPDH3	GCTTTCCAGAGGGGCCATCCACA	59°C	426bp	NM_017008.4
	rGAPDH4	ACGGCAAGTTCAACGGCACAGTCA			
rat DNMT1	rDnmt1-3	GTGCCTGCCAGCTGAGTGTTGTG	60°C	483bp	NM_053354.3
	rDnmt1-4	ACCAGGGGATGAGCGTGTTGAAT			
rat DNMT3a	rDnmt3a-1	GCACGGGCCGCCTCTTCTT	60°C	502bp	NM_001003957.1
	rDnmt3a-2	CACGATCGGCCCAGCAGTCTCT			
rat DNMT3b	rDnmt3b-1	AGCCAGGAGACGCGAGAACAAAAG	60°C	412bp	NM_001003959.1
	rDnmt3b-2	ACCCACCAGCACCTCCAGACACTC			
rat INSIG1	rInsig1-1	TCTCCTCCGCCTGGTGGGTG	60°C	268bp	NM_022392.1
	rInsig1-2	CCCAGGCCACTTCGGGATCG			
rat IDI1	rIDI1-1	GTTGAAGTACGGCGCTCCGCA	60°C	418bp	NM_053539.1
	rIDI1-2	AGCGCTTCTGTGCTGCTCGTTT			
rat LUM	rLum1	GCCCGCTGGCCTTCCAACAT	55°C	350bp	NM_031050.1
	rLum2	GGGGATTGCCATCCAAGCGCA			
rat ID1	rIDB1	TGTCGGAGCAAAGCGTTGCCA	61.5°C	331bp	NM_012797.2
	rIDB2	GCTGGAACACATGCCGCCTCGG			
rat PRND	rPPD1	ACCGAAGCCCAGGTGGCTGA	60°C	373bp	NM_001102431.1
	rPPD2	CCAGCAGGCAGACCATCGCC			
rat ACTA1	rACTA1-1	GGCAATGAGCGCTTCCGTTGC	59°C	392bp	NM_019212.2
	rACTA1-2	AGACGCGGGTGCGCCTAGAAG			
rat MYH2	rMYH2-1	TGGAGGCCAGAGTGCGTGAAC	55°C	436bp	NM_001135157.1
	rMYH2-2	CACATGGGGACATGACCAAAGGC			
rat TXNIP	rTIP-1	AGTGGGTACACCCCCGCCTC	55°C	300bp	NM_001008767.1
	rTIP-2	TGGCTCTGTTGCCCCATGGC			
mm DNMT1	mDnmt1-1	GGTGGATGGCCGGGTCTACTGC	60°C	489bp	NM_001199431.1
D.111 472	mDnmt1-2	GGGCCCCCTTGTGAATAATCCT	CO 10	44.01	
mm DNM13a	mDnmt3a-1	CALCIAIGGGCIGCIGCGAAGAC	60°C	419bp	NM_007872.4
	mDnmt3a-2	GGCGGCCAGTACCCTCATAAAGT	6010		
mm DNM13b	mDnmt3b-1	IGICGGAGCAAAGCGIIGCCA	60°C	414bp	NM_001003961.4
0.14	mDnmt3b-2	IGGCAGCGCIGAGGGAGGCACAIA	CO ¹ O	5001	
mm Oct4	mOct4-1	ACCGCCCCAATGCCGTGAAGTT	60°C	593bp	NM_013633.3
10712	mOct4-2	IGGGGGCAGAGGAAAGGAIACAGC	5790	200	NIM 007202.2
mm ACTA2	mACTA2	AGICGCIGICAGGAACCCIGAGACG	570	296bp	NM_007392.3
			F7%2	22.44-	
MM IVIYH11	mMYH11-1	AGGAAALALLAAGGILAAGLA	5/-0	324bp	NM_013607.2
				220h	
mm SIVIZZ	mSN122a-1		57 C	3280p	NIVI_011526.5
	1112101229-2	GAATIGAGCCACCIGTICCATCIG			

 Table 2: Oligonucleotides used for COBRA analyses

	primer				
gene ID	name	primer sequence	annealing	product	Genbank
rat LINE1	rLINE1-s	TTTGGTGAGTTTGGGATA	56°C	163bp	AC229945
	rLINE1-as	CTCAAAAATACCCACCTAAC			(Pogribny et al. 06)
rat ER $lpha$	rERa-e1-s	TTTTGTTGTATTAGATTTAAGGGAA	59°C	272bp	NM_012689.1
	rERa-e1-as	ΑΑΑΑΑΑΑΑCCCCCAAACTATTAAC			(Monje et al. 07)
rat 14-3-3 σ	rSFN1-s	TTTTAGGAGTTATTTGGGTGTTGAT	60°C	229bp	XM_003750048
	rSFN1-as	ССССТАААСТАААТААААСТАСТССС			(Chen et al. 10)
rat PRND	rPP2-s	TTATTGGTAGTTTTTTGATGGGA	59°C	283bp	NM_001102431.1
	rPP2-as	ΑΑΑCCAAATAAAACCCAACAAA			
CMV	CMV-BSP-s	TGATTTTATGGGATTTTTTTATTT	52°C	269bp	X17403.1
	CMV-BSP-as	CAACTCTACTTATATAAACCTCCCA			

	primer			
gene ID	name	sequence	annealing	product
rat ACTA1	ACTA1-ps-1	TTTGGTTTTTAGTATTATGAAGATTAAGGT	56°C	205bp
	ACTA1-ps-2	ACTCCTACTTAATAATCCACATCTACTA		
	ACTA1-seq	TTTAGTATTATGAAGATTAAGGTG		
	sequence	GATGAYGYGT TTGGTATGGG YGGAGATTAG GGGYGGGGGG		
	analysed	AATTYGTAGG [+2554 to +2700 from transcriptional start]		
rat MYH2	MYH2-ps-1	TAGATTTTGAAGGTGAAAGAAGATTAGA	56°C	178bp
	MYH2-ps-2	ACATTAAAAACCCCTAATATAAATCAC		
	MYH2-seq	AGATAGAGGATATGGTTATGATG		
		ATTTATTTTT AYGAGTTYGT TGTGTTGTAT AATTTTAAAG		
	sequence	AGYGTTAYGT AGTTTGGATG ATTTAYGT [+2379 to +2502		
	analysed	from transcriptional start]		
rat PRND	PRND-ps-1	GGGATTTATTAAGAAGGTTGTTTTGAAG	56°C	221bp
	PRND-ps-2	AAACCCAACAAACAAACCA		
	PRND-seq	GTGATTAAGGAGGTGTT		
		GGTGATTYGT TGYGTTAAYG TTATTTAGGY GGTTAATTAG		
	sequence	GTTGAGTTTT TT Y GGGAGAA [+311 to +405 from		
	analysed	transcriptional start]		
rat TXNIP1	TXNIP-ps-1	GGTTTAATTATGGTGATGTTTAAGAAGATT	56°C	205bp
	TXNIP-ps-2	ATCTATTTACACTACTAAAACCCTTACAT		
	TXNIP-seq	ATGGTAGAGGGGAGAA		
	sequence	GGTGGTYGGT YGGGTGATAG TGGAAGTGTG TGAAGTTATT		
	analysed	YGAGTTAAAG TYGTTAGGAT TTTGGTTTGY G [+210 to +357]		
	sequence	YGGAATGTTAYGTTTTTTYGGAYGA [PM00547638] [+308 to		
rat INSIG1	analysed	+343 from transcriptional start]	56°C	163bp
	sequence	YGGGTAGGYGYGTTTTAGYGTAGYGYGA [PM00543697]		
rat ID1	analysed	[+174 to +202 from transcriptional start]	56°C	208bp
	sequence	TTYGTAYGTTTAGTTGTTTTGYGGT [PM00504119] [-65 to -23		
rat IDI1	analysed	from transcriptional start]	56°C	112bp

 Table 3: Oligonucleotides used for pyrosequencing analyses and gene sequences analysed

Table 4: Top 20 gene expression changes in response to folate deficiency. Fold change of gene expression between A7r5 cells grown in in folate sufficient (100ng/ml) and folate deficient (2.5ng/ml) medium

name	gene description	fold change	Genbank
	expression increased in 100ng/ml vs 2.5ng/ml folate		
Txnip	thioredoxin interacting protein	8.97	U30789
ltgb8	integrin subunit beta 8, transcript variant X1	8.61	XM_006240697.3
Lum	lumican	8.34	NM_031050
Insig1	insulin induced gene 1	8.10	NM_022392
ldi1	isopentenyl-diphosphate delta isomerase	6.55	NM_053539
Ednra	endothelin receptor type A	6.04	BF414702
Ptn	pleiotrophin	5.99	NM_017066
ltgb8	integrin subunit beta 8, transcript variant X1	5.97	XM_006240697.3
Scd2	stearoyl-Coenzyme A desaturase 2	5.84	NM_031841
Cdc20	cell division cycle 20 homolog	5.44	U05341
ldi1	isopentenyl-diphosphate delta isomerase	4.61	BI290053
LOC684841	histone H3.2-like	4.48	XM_006222398.2
Slc29a1	solute carrier family 29 (nucleoside transporters), member 1	4.46	NM_031684
Cenpw	centromere protein W	4.41	NM_001246319.1
Kif20a	kinesin family member 20A	4.38	BE111697
Cdca3	cell division cycle associated 3	4.34	BF417638
Nrp1	neuropilin 1	4.21	AF016296
Arrdc4	arrestin domain containing 4	4.18	XM_006229367.3
Hmgcs1	3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1	3.93	NM_017268
Kif11	kinesin family member 11	3.93	BE116384
	expression increased in 2.5ng/ml vs 100ng/ml folate		
Prnd	prion protein dublet	24.95	BF551984
Myh2	myosin, heavy polypeptide 2, skeletal muscle, adult	11.52	BI277586
ld1	inhibitor of DNA binding 1	9.79	M86708
Acta1	actin, alpha 1, skeletal muscle	9.11	NM_019212
F3	coagulation factor III	9.10	NM_013057
Tnnt1	troponin T1, skeletal, slow	8.92	AF399874
Plcxd2	phosphatidylinositol-specific phospholipase C	7.81	NM_001134481.1
Myh3	myosin, heavy polypeptide 3, skeletal muscle, embryonic	7.38	NM_012604
ld2	inhibitor of DNA binding 2	6.96	BC086391.1
Scg2	secretogranin II	6.73	NM_022669
Vsnl1	visinin-like 1	6.51	AI227991
Rnf181	ring finger protein 181	6.31	AI008549
Tnc	tenascin C	6.31	AI176034
lsg12(b)	putative ISG12(b) protein	6.13	AA819034
Prss35	protease, serine, 35	5.88	AA866443
Myl2	myosin, light polypeptide 2, regulatory, cardiac, slow	5.38	BF419995
Trib3	tribbles homolog 3	5.37	AB020967
Рср4	Purkinje cell protein 4	5.10	NM_013002
Ppp1r1a	protein phosphatase 1, regulatory (inhibitor) subunit 1A	5.00	NM_022676
Plekhb2	pleckstrin homology domain containing, family B member 2	4.99	AW254369

Table 5: TOP-GO pathway analysis of transcriptome changes in response to folate deficiency. elimKS here gives the p-values from topGO specific Kolmogorov-Smirnov Test with elimination of genes of significant GO-terms in parent terms.

GO.ID	Term	Annotated	Significant	Expected	elimKS
GO:0007076	mitotic chromosome condensation	17	16	5.02	3.40E-07
GO:0006695	cholesterol biosynthetic process	65	38	19.18	1.30E-06
GO:0051301	cell division	621	249	183.24	1.40E-06
GO:0000381	regulation of alternative mRNA splicing	80	43	23.61	2.20E-06
GO:0070507	regulation of microtubule cytoskeleton	288	104	84.98	6.10E-06
GO:0007059	chromosome segregation	412	188	121.57	1.10E-05
GO:0034508	centromere complex assembly	31	19	9.15	1.70E-05
GO:0031055	chromatin remodelling at centromere	14	12	4.13	1.70E-05
GO:0045214	sarcomere organization	87	45	25.67	1.90E-05
GO:0071897	DNA biosynthetic process	215	98	63.44	2.00E-05







С





D



[-]RA [-]puro



[+]RA [-]puro



×**

[+]RA [+]puro







D



Ε



Kolb et al.; Fig. 4













ng/ml folate



F

F

DFB





fold change from CON [-]RA

120





MYH11





confluent



В





D

























thioredoxin interacting protein [TXNIP]

В

insulin induced gene 1 [INSIG1]



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D





Ε



F

a-actin skm [ACTA1]



CON

E FCS

DÆ

DF

*

DNMT3a



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В



D









D

Dnmt1 - ID 19991



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F

С

Dnmt3b - ID 14143







В







120

4









[-] F gene list [p<0.01; fold change >2] genes activated by folate deficiency

HF-1-alpha trans	cription factor network, Homo sapient, 20ef2bB1-6193-13e5-8ac5-06603eb7f303
PAR1-mediated th	rombin signaling events, Homo sapieris, be5084c0-6194-11e5-8ac5-06603eb7f303
UPA receptor medi	ated events_Nomp sagies_46994cde=6194-11e5-8ac5-06603eb7f303
Validated targets of	d C-MYC transcriptional repression_Homo sapiens_6bbdafa6-6196-11e5-8ac5-06603eb7f303
Plasma membrane	extrogen secretor signaling. Homo sapiens_dcc37895-6194-11e5-8ac5-06603eb7f303
Betal integrin cell	surface interactions, Nome supiens, 2fd0bc63-618d-11e5-8ac5-06603eb7f303
North-mediated in	CS/19 Y network_Homo sapiens_8ee56389-6194-11e5-8ac5-06603eb7f103
Arts mattering re-	Homo sapiens_7a5b8f09-618a-11e5-8ac5-06603eb7f303
Rule of Calciments	-dependent NFAT signaling in lymphocytes, Homo sapiens, 61bdd46d-6195-11e5-8ac5-06603eb7f3
SIFI authenty, Hor	mo sapiens_7327884f-6195-11e5-8ac5-06603eb7f303



А

cardiomyopathy	
myopathy.	
cardiomyopathy_dilated	
blood	
muscular_dystrophy	
leukemia	
anemia	
colorectal_cancer	
neuropathy	







[+] F gene list [p<0.01; fold change >2] genes silenced by folate deficiency



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