An integrative analysis reveals coordinated reprogramming of the epigenome and the transcriptome in human skeletal muscle after training

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Abbreviations: β-HAD, 3-hydroxyacyl-CoA dehydrogenase; CpG, cytosine-guanine dinucleotide; CS, citrate synthase; DEG, differentially expressed gene; DMP, differentially methylated position; FDR, false discovery rate; LUMA, luminometric methylation assay; MDS, multidimensional scaling; MEF, myocyte enhancer factor; MRF, myogenic regulatory factor; PCA, principal component analysis

Introduction

Endurance exercise training is a strong physiological stimulus that leads to a multitude of health and functional improvements when performed regularly. The health benefits include prevention and/or treatment of a multitude of the most common diseases, e.g., cardiovascular disease, type II diabetes, and several forms of cancer.1 The health benefits following exercise training are elicited by gene expression changes in skeletal muscle, which are fundamental to the remodeling process.2 Epigenetic modifications through DNA methylation regulate gene transcription.3 DNA methylation has traditionally been considered to be a mitotically stable modification that could change only over long periods of time, e.g., in disease
development and during the general aging process. However, there is increasing evidence that more short-term environmental factors can influence DNA methylation. For example, dietary factors have the potency to alter the degree of DNA methylation in different tissues, including skeletal muscle, while exercise is less well studied. In one study, a single bout of endurance-type exercise was shown to affect methylation at a few promoter CpG sites. In the context of diabetes, exercise training has been shown to affect genome-wide methylation pattern in skeletal muscle, as well as in adipose tissue. Together, those data indicate that physiological stressors can indeed affect DNA methylation.

Adaptation in skeletal muscle depends on successive transient increases in mRNA encoding regulatory, metabolic, and structural proteins. However, the acute changes in gene expression are quite different from the more robust basal alterations that characterize a well-adapted muscle after a major lifestyle change, e.g., months of regular exercise.

In order to obtain a deeper understanding of the mechanisms underlying the massive functional and health benefits of regular exercise, we conducted a 3-month fully supervised human one-legged exercise training study. Training only one leg allowed for an intraindividual control, thereby excluding potential influence of diet, seasonal changes or unknown environmental factors, which are expected to affect both legs equally. We observed that the training intervention reshapes the epigenome and induces significant changes in DNA methylation, and the global fractions of decreased and increased methylation sites were similar. Importantly, changes in DNA methylation were enriched in regulatory enhancer regions. Functional categories related to muscle biology (e.g., regulation of cellular carbohydrate metabolism and structural remodeling) were overrepresented among differentially methylated sites. In addition, a coordinated transcriptional and epigenetic response was identified through network analysis.

Together, the findings from this tightly controlled human study strongly suggest that the regulation and maintenance of exercise training adaptation is to a large degree associated to epigenetic changes, especially in regulatory enhancer regions.

**Results**

**Endurance exercise training induces global alterations in DNA methylation**

Twenty-three young volunteers (Table 1), not regularly performing intense exercise, performed supervised one-legged knee-extension exercise training for 3 months (45 min, 4 sessions per week; Figure S1A), training only one randomized leg (trained leg), while the second leg (untrained leg) was used as an intraindividual control leg. Skeletal muscle biopsies from the vastus lateralis were taken from both legs at rest, before and after the training period (see Supplementary Methods S9 for details). Performance improvements and enzyme activity increases in the trained leg confirmed that the training response was highly significant ($P < 10^{-4}$, Fig. 1A-B and Figure S1B-C).

To address the effect of training on DNA methylation of specific sites across the human skeletal muscle genome we used the Illumina Infinium HumanMethylation450 arrays. Endurance training [after training (T2) vs. before training (T1)] induced significant (false discovery rate, FDR < 0.05) methylation changes at 4919 sites across the genome in the trained leg (Fig. 1C, Dataset S13). Of these, 839 sites had an absolute change of at least 5% in their mean methylation level (β-value) after training, with a maximum of 9%. The corresponding transcriptional analysis was performed using RNA sequencing, which identified 4076 differentially expressed genes (DEGs; Fig. 1D, Data set S13). Ontology and pathway analysis of DEGs showed that the transcriptional response robustly reflected key pathways involved in training adaptation and a trained muscle phenotype (Figure S2-S3 and Table S10). Clustering analysis identified training and gender as the major determinants of variability on autosomal DNA methylation and gene expression data (Fig. 1E). The clustering was independent of gene expression on sex chromosomes, as a corresponding grouping was obtained excluding genes on X and Y chromosomes (data not shown). Moreover, a complementary approach revealed that over 600 CpG sites correlated to the increase in citrate synthase activity, an objective measure of training response (Figure S4 and Dataset S14). This might imply that some of these sites could influence the degree of training response.

Global cytosine methylation was also analyzed to look for globally unidirectional changes by luminometric methylation assay (LUMA), revealing no overall change with training, and no difference between the trained and the untrained leg (Figure S5A). Global CpG methylation was remarkably similar both within the same subject and between all subjects. This is consistent with the specific observations from the array data of approximately equal number of sites increasing and decreasing in methylation and no significant change in the sample average methylation after training. Analysis of global hydroxymethylation levels resulted in no overall change with training (Figure S5B).

As expected by a physiological environmental trigger on adult tissue, the observed effect size on DNA methylation was small in comparison to disease states such as cancer. Hence, to exclude

| Table 1. Baseline subject characteristics for males (n = 12), females (n = 11) and all subjects (n = 23). Data is presented as mean ± SEM. For details on the training program see Figure S1. |
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| **Age (yrs)** | **Height (cm)** | **BMI (kg/m²)** | **Peak VO₂ (ml·kg⁻¹·min⁻¹)** |
| Males | 27.5 ± 0.97 | 181 ± 2 | 24.8 ± 1.3 | 39.5 ± 1.1 |
| Females | 26.4 ± 1.31 | 169 ± 2 | 23.2 ± 0.9 | 38.4 ± 1.2 |
| All subjects | 27.0 ± 0.79 | 175 ± 2 | 24.0 ± 0.8 | 39.0 ± 0.8 |
Figure 1. Effects of endurance exercise training are mirrored by alternations in DNA methylation and gene expression. The significant (* \( P < 10^{-4} \)) effect of training in the trained leg is shown by the increase in a 15 min performance test (A) and the citrate synthase activity in muscle biopsies (B) in T2 (after training) vs. T1 (before training). For comparison, the untrained leg is also shown, where a smaller change is observed in performance and no change in CS activity. # indicates significant differences between the changes in the 2 legs. Additional physiological measurements are shown in Figure S1B-C. The physiological changes are mirrored by modifications in DNA methylation (C) and gene expression (D). For DNA methylation, the effect size is measured as the difference in M-values and points in black correspond to DMPs with FDR < 0.05. For gene expression, the \( \log_2 \) (Fold Change) is plotted against the average \( \log_2 \) (Counts Per Million) and red points correspond to genes with FDR < 0.05. Correlation between changes in DNA methylation and CS activity exists and results are shown in Figure S5. The clustering of the samples is shown in (E) using either DNA methylation (upper panels) or gene expression (lower panels). A segment connects to measurements from the same subject obtained before and after training. Samples are alternatively colored by group (T1 = blue, T2 = red) or by gender (M = green, F = magenta). For DNA methylation, Principal Component Analysis was employed using only autosomal DMPs, while for gene expression the top 1000 genes with largest biological variation were chosen and the biological coefficient of variation used to produce a multidimensional scaling plot.

Possible technical artifacts leading to false positive associations, we selected several genomic positions for bisulfite pyrosequencing validation. Overall, the methylation levels significantly correlated with the 450K array data (Figure S5C). Out of 7 selected sites, we observed the same direction of change for 6 positions (up/down/no change), thus providing support on the reliability of our data. The untrained leg was also included in the validation to verify that the observed changes were indeed an effect of the training itself and not due to other environmental effects. No significant change was detected in the untrained leg. For details, see Figure S5D-J and the corresponding text. Moreover, a Q-Q plot for the array results is showed in Figure S8.

Enrichment of training-induced DMPs in enhancers Annotation of the differentially methylated positions (DMPs) revealed a preferential localization outside of CpG Islands/Shelves/Shores. When considering the positioning of
the DMPs with respect to the genes, we could also detect an enrichment of the relative fraction of DMPs in gene bodies and intergenic regions and fewer DMPs in promoter regions (TSS200), as compared to the relative distribution of the probes present in the array (Fig. 2A–B). Given these observations, we asked whether the major source of the DMPs could originate from enhancers. Indeed, using the standard array annotation, DMPs were significantly enriched in enhancers (Fig. 2C). This finding was further corroborated using external sources of data and annotations (see Supplementary Methods S9). For example, when defining enhancers based on histone marks (either H3K4me1, H3K27ac, or H3K4me1 and H3K27ac jointly) and using completely independent data from human skeletal muscle tissue, we also confirmed that the changing sites were observed predominantly in enhancers (Fig. 2C). Additional evidence came from the definition of chromatin states by a chromatin segmentation algorithm and the corresponding public data available for human myoblast cultures. Here we observed an enrichment, relative to the array background, for strong and weak enhancers, but not for promoters (Fig. 2D). We also observed a slight preferential enrichment in regions of accessible chromatin, as defined by DNase Hypersensitive Sites (Figure S6). These observations have potential implications on the role of DNA methylation on regulating gene expression, a phenomenon likely mediated by altering the methylation status of enhancers and other regulatory elements.

The finding that endurance training especially influences enhancers is indubitably novel in the context of tissue adaptation to a physiological stimulus in humans. Therefore, we sought to identify the biological processes corresponding to these regulatory regions.

**Differential methylation is related to genes governing muscle related processes**

To assess the putative physiological relevance of the DMPs, we performed a functional annotation and ontology enrichment analysis, where we defined an association rule that considered gene regulatory domains (see Supplementary Methods S9). Indeed, we detected a clear enrichment of muscle ontology related processes for the genes in the vicinity of DMPs, demonstrating that the top enriched categories of molecular function, biological process and cellular component were linked to myogenesis as well as

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**Figure 2. DNA methylation changes are primarily localized in enhancers.** (A–C) For each annotation category, the relative fraction of positions located within each feature type is calculated for DMPs (red bars), non-DMPs (blue bars) and the entire position on the array (green bar). Panels A-B were obtained using Illumina annotation, while for panel C we used Illumina annotation (Enhancers) or publicly available ChIP-seq experiments on HSMM cells from which we derived H3K4me1 and H3K27ac peaks. The presence of H3K27ac mark denotes an active regulatory element, while H3K4me1+H3K27ac marks active enhancers and other distal elements. (D) The log2 fold enrichment for DMPs vs. the array was calculated for the relative fraction of probes falling in each category; data from chromatin segmentation algorithms in HSMM cells were used. For additional information, see Supplementary Methods S9. Significance codes: *P < 0.05; **P < 0.001; Fisher’s exact test.
Figure 3. For figure legend, see page 1562.
muscle structure, function and bioenergetics, in concordance with a trained muscle phenotype (Fig. 3). This prompted to a deeper characterization of DMPs, by extending bidirectionally the sequence of the position (±100 bp) and examining the enrichment of known transcriptional motifs. Concordantly, we observed a significant enrichment for binding motifs of the MRF (Myogenic Regulatory Factors), and MEF2 (Myocyte Enhancer Factor 2) classes in DMPs increasing in methylation after training (Fig. 3 and Table S11). Conversely, for DMPs that decreased their methylation level, an enrichment for the ETS family binding domain was observed (Fig. 3 and Table S11). The ETS transcription factors have been strongly implicated for example in angiogenesis, a key process in endurance exercise adaptation.

Specific differential methylation correlates with expression changes in an exercise-dependent manner

The finding that differential DNA methylation is observed at enhancers and regulatory regions with the associated enrichment of muscle ontology processes suggests a functional link between DNA methylation and RNA expression. To integrate those 2 data types, we initially performed correlation analysis, in order to highlight pairs of genes/methylation positions that showed dependence after training. Overall, when all probe/position pairs were considered, the Spearman correlation was centered on zero (Fig. 4A), revealing no sign of global trend. However, selection of the pairs resulting from a DMP and a DEG revealed that an effect of exercise training exists in specific genomic regions, indicated by peaks of both positive and negative correlation (Fig. 4A) that were preferentially observed according to the relation to gene regions. In particular, negative correlation was more prominent for probes in promoter/5’UTR/1st exon regions, while gene bodies had a stronger peak of positive correlation (Fig. 4B).

In principle, since we are analyzing longitudinal observations on the same subject before and after training, any detected pattern of correlation between expression and methylation could arise from the individual levels of methylation and gene expression being correlated at the subject level and not as a result of the experimental intervention. Indeed, a Q-Q plot (Fig. 4C) revealed that, after excluding DMP/DEG pairs, some residual correlation remained and this behavior is most likely explained by the presence of some baseline level of correlation between DNA methylation and gene expression, that is not explained by training, but instead by inter-individual factors (e.g., gender). However, when considering only the DMP/DEG pairs, we reduced the number of false positives or, in other words, we decreased the possibility that the dependence is not explained by the endurance training. This phenomenon is further exemplified by a starburst plot that illustrates the relationship between DNA methylation and expression changes (Fig. 4D). This integrative analysis identified 255 downregulated genes with significant increase in methylation, 203 upregulated genes with significant increase in methylation, 273 upregulated genes with significant decrease in methylation, and 70 downregulated genes with significant decrease in methylation. Those gene/position pairs correspond to interesting examples of changes shaped by the training-specific intervention and, therefore, provide the fraction of genes whose expression-methylation correlation is changed in different directions. Individual examples selected among the top-correlated genes are given in Figure 4D. We found several interesting genes in the context of muscle physiology and metabolism. Examples include the MIPEP gene that encodes for a mitochondrial peptidase, primarily involved in the maturation of proteins involved in oxidative phosphorylation, and GRK5, a GPCR kinase involved in multiple biological processes, proposed as a positive regulator of insulin sensitivity in mouse. In turn, some of the highly correlated gene/methylation probes were not differentially regulated after training, including many HLA genes and other genes whose expression could possibly be influenced by individual genotypes and not by the experimental intervention. This is exemplified by THNSL2 (in the bottom left panel of Fig. 4D), whose bimodal expression has been linked to a cis-eQTL in muscle tissue. For more examples, see Table S12.

Integrating transcriptional network analysis with DNA methylation identifies a coordinated training response

Having established that DNA methylation alterations are induced by training and are linked to corresponding genes that are also regulated by exercise training, we asked whether we could visualize epigenomic changes superimposed on a reconstructed transcriptional network. Hence, we reverse-engineered a mutual information-based transcriptional network (p < 10^-10) using count data from DEGs alone and, in a second independent step, we overlaid the DNA methylation data by summarizing the methylation changes of the linked DMPs (Fig. 5). This analysis uncovered 3 major network domains and some smaller modules (Figures S2-S3) that were grouped according to their relative fold change (T2 vs. T1) that corresponded to distinct ontologies. Domain A was overall downregulated (average log2FC = -0.31) and was populated with genes annotated for regulation of gene expression, DNA replication and cell cycle. Examples include MDM2, a E3 ubiquitin-protein ligase that mediates ubiquitination of p53, ZNF638, a transcription factor associated to PPAR expression, UACA involved in regulation of stress-induced apoptosis and CUL3, part of an E3 ubiquitin-protein ligase complex and previously shown to decrease in expression with endurance training. At the other extreme, domain C contained upregulated genes (average log2FC = 0.60) involved in morphological changes, including cell adhesion, blood vessel development and extracellular matrix organization. This domain included genes that have been shown to increase in expression.

Figure 3 (See previous page). Muscle related processes and factors are enriched in the up-methylated DMPs. GREAT analysis was performed to retrieved functional categories associated with DMPs increasing or decreasing methylation after training. Up to top 5 categories passing the threshold (see Supplementary Methods S9) are shown for GO Molecular Function, Biological Process and Cellular Component. We tested the presence of known enriched motif on a symmetrical 200 bp window around each DMP (P < 10^-15, consensus motif shown). Known profiles were clustered and a familial logo was drawn. For a corresponding ontology analysis of gene expression, see Figures S2-S3.
with training, such as several collagens (e.g., COL4A1 and COL4A2), the protease inhibitor A2M, the adhesive glycoprotein gene THBS4, and PDGFRB involved in blood vessel formation. Domain B contained a majority of upregulated genes, with an intermediate magnitude as compared to domain C (average log2FC = 0.25). This domain was clearly associated with cellular energetics, mainly oxidative phosphorylation. Many mitochondrial enzymes were found in this group, including CS, SDHA, HADHA, COX7A2, ATP5B and several NADH dehydrogenases.

Importantly, the independent integration of DNA methylation information on top of the transcriptional network identified a consistent pattern of inverse changes, possibly reflecting a
coordinate transcriptional plan (Fig. 5 and Data set S13). In fact, the upregulated domain C was associated with a significant decrease in DNA methylation, whereas the downregulated domain A had almost entirely corresponding positive changes in DNA methylation. Concordantly, domain B contained a comparable fraction of genes with corresponding positive or negative changes in DNA methylation. Interesting genes whose methylation changes were opposite to those of transcription included the collagens COL4A1 and COL4A2, and laminin LAMA4 that form the basement membrane around skeletal muscle cells. For oxidative metabolism, the enzymes MDH1 and a NADH dehydrogenase of the electron transport chain NDUFA8 represent interesting examples. In domain A, the myosin phosphatase PPPI12A and TRDN, likely involved in regulation of calcium release from the sarcoplasmic reticulum, both increase in methylation and decrease in expression.

**Discussion**

From this well-controlled, prospective and extensive human study we were able to create a map of coherent, consistent and biologically relevant DNA methylation changes in skeletal muscle tissue in response to a lifestyle intervention known to improve function and health. The significant changes in DNA methylation, that primarily occurred in enhancer regions, were to a large
extent associated with relevant changes in gene expression, even if no causal relationship could be definitely determined.

The main findings of this study were that 3 months of endurance training in healthy human volunteers induced significant methylation changes at almost 5000 sites across the genome and significant differential expression of approximately 4000 genes. The genes associated with the DMPs that increased and decreased in methylation, respectively, with training, represent distinct ontologies. DMPs that increased in methylation were mainly associated to structural remodeling of the muscle and glucose metabolism, while the DMPs with decreased methylation were associated to inflammatory/immunological processes and transcriptional regulation. This suggests that the changes in methylation seen with training were not a random effect across the genome but rather a controlled process that likely contributes to skeletal muscle adaptation to endurance training.

CpG methylation is subjected to spatial (tissue- or cell-specific methylation) or temporal variability (age-dependent, disease-associated, or environmental-mediated differential methylation). In order to eliminate confounding factors, we formulated a paired study design, where 2 samples from the same individual were contrasted before and after the training intervention. We verified, for selected sites, that the observed effect, despite small, was specific for the trained leg, corroborating the hypothesis that the measured changes in DNA methylation represent explicit molecular manifestations of exercise training. Unlike certain diseases, e.g., cancer, environmentally induced changes in DNA methylation are typically small, and susceptible to large individual variation. By employing an intraindividual control leg for validation of methylation and physiological changes, we minimize any influence of diet or other environmental factors that may otherwise be a challenge in environmental epigenomics.

Here we successfully report the identification of significant and ontologically coherent changes in DNA methylation in a human genome-wide study of the effect of endurance exercise in muscle tissue, which was made possible by the combination of a well-controlled, prospective study design with a comprehensive, integrative bioinformatic analysis.

A well-known source of heterogeneity in DNA methylation studies is represented by the cell composition of the analyzed sample. The epigenome varies between cell types and correcting for cellular composition has been shown to reduce confounding due to cell type bias in blood. Nevertheless, for complex solid tissue samples, the correction for cell composition is not straightforward, as it would require the isolations and profiling of individual cell types. An increased vascularization following endurance exercise results in a higher endothelial cell content per skeletal muscle fiber. In line with this, we confirmed that some exclusive and non-exclusive endothelial markers were differentially expressed after training (data not shown). Hence, we cannot completely rule out the possibility that a minor fraction of the observed changes in DNA methylation and/or gene expression could reflect changes in cellular fractions.

Correlation of the changes in DNA methylation to the changes in gene expression showed that the majority of significant methylation/expression pairs were found in the groups representing either increases in expression with a concomitant decrease in methylation or vice versa. The fraction of genes showing both significant decrease in methylation and upregulation was 7.5% of the DEGs or 2.3% of all genes detected in muscle tissue with at least one measured DNA methylation position. Correspondingly, 7.0% of the DEGs or 2.1% of all genes showed both significant increase in methylation and downregulation. This could reflect the classical view on promoter methylation with a reciprocal relationship between methylation and expression. The pairs showing a concordant change were skewed in the positive direction. In fact, 5.6% of the DEGs or 1.7% of all genes were significantly changing in the positive direction, whereas only 1.9% of the DEGs or 0.6% of all genes had significantly negative changes. In total, we show that DNA methylation changes are associated to gene expression changes in roughly 20% of unique genes that significantly changed with training.

One previous study has investigated genome-wide alterations in DNA methylation after long-term training using MeDIP-ChIP and related those to gene expression changes by microarrays. Various methods are now available for determining CpG methylation status, each potentially exhibiting advantages and limitations and differing in the ability to detect differential methylation. We employed a method (450K arrays) that quantifies methylation levels at specific loci and does not require correction for CpG bias, coupled to RNA-seq, in order to uncover the connection between the epigenetic and transcriptional responses, and therefore obtain a deeper understanding of the impact of regular exercise. The methodological differences render a direct comparison more difficult, however, both studies show reduced methylation of some genes involved in muscle physiology, for example RUNX1 and COL4A1. In contrast, for the gene THADA also highlighted in Nitert et al. we observe an increased methylation of 2 CpG sites. Our sites are, however, located in the gene body, while changes in Nitert et al. occur in promoter regions, something that likely produces different effects on transcription.

The network reconstruction resulted in domains classified mainly as structural, metabolic and regulatory and suggested a coordinated pattern of change in DNA methylation and gene expression. Examples of structural genes include COL4A1, COL4A2 and LAMA4. These genes have also been identified as important for differences in responsiveness to endurance training, where methylation status could be part of the mechanism behind variable training response. Among the metabolic genes, MDH1 catalyzes the reversible oxidation of malate to oxaloacetate, utilizing the NAD/NADH cofactor system in the citric acid cycle and NDUFA8 plays an important role in transferring electrons from NADH to the respiratory chain. Regulatory genes downregulated with training include MDM2, targeting p53 for proteasomal degradation, and PPP1R12A, a subunit of myosin phosphatase that is also capable of inhibiting HIF1AN-dependent suppression of HIF1A activity. The resulting inhibition of HIF1A could be an advantage for endurance adaptation as HIF1A has a negative effect on oxidative metabolism.

Recently it was shown that enhancers have dynamic methylation and that their methylation levels are more closely associated
with gene expression alterations than promoter methylation in cancer. In the present study, methylation predominantly changed in enhancer regions with enrichment for binding motifs for different transcription factors suggesting that enhancer methylation may be highly relevant also in exercise biology. Regions with sites increasing in methylation were enriched for myogenic regulatory factors or MRFs and myocyte enhancer factors, or MEFs. The MRFs include Myf5, MyoD, myogenin, and Mrf4. These are basic helix-loop-helix transcription factors of the myogenic lineage that control the determination and differentiation of skeletal muscle cells in the embryo. Of special interest in the biology of endurance training may be that MRFs, through binding to the PGC-1α core promoter, can regulate this well-studied co-factor for mitochondrial biogenesis. Animal studies indicate that MyoD promotes slow-to-fast fiber transformation while myogenin could possibly work as an ultimate transcription factor binding directly to promoters of mitochondrial enzymes, thereby promoting a more oxidative profile of muscle tissue.

The myocyte enhancer factor 2 (MEF2) transcription factor links extracellular signals (partly through regulation of class IIa histone deacetylases, HDACs) to the activation of genetic programs involved in cell differentiation, proliferation, morphogenesis, survival and apoptosis in many different cell types. KO experiments suggest that some of the MEF2 isoforms promote slow fiber phenotype during development. That endurance training led to an increased methylation in enhancer regions containing motifs for the MRFs and MEFs is somewhat counterintuitive since it should lead to the repression of the action of the above discussed transcription factors. Possible explanations include either: a) a dynamic regulation of enhancer accessibility and activity, involving active methylation as an inactivation mechanism at the time (i.e., 3 months) when morphological changes have already taken place and a negative feedback should be provided; b) a confounding effect due to increased representation of different cell types after training, which would result in the detection of a change in methylation because of the underlying differences in methylomes between muscle and non-muscle cell types and c) possible incompletely understood phenomena such as the differential methylation that inhibits transcription of an inhibitory RNA molecule, as it was shown for p15. However, these factors regulate a large number of genes and there are several that decrease with training in this study, including CDCH15, MYH3, TNNT2, RYRI and SH3GLB1, which would be expected from an increased enhancer methylation. Also, the expression of MEF2A itself decreased with training.

Regions with sites decreasing in methylation with endurance training were enriched for the ETS family of transcription factors, a large group with a well-conserved binding motif. A reduction in methylation with training in enhancer regions containing this motif potentially allows for further activation of their response genes. There are several members of this group that are known to regulate exercise-responsive genes, thus proposing an epigenetic regulation with endurance training. GABPA, also known as nuclear respiratory factor 2, controls expression of several genes involved in mitochondrial respiration in human skeletal muscle and is itself regulated by the transcriptional coactivator PGC1α. Another interesting example is ETS1, that targets genes important for endothelial migration e.g., matrix metalloproteinases, angiopoietin 2 and the vascular endothelial growth factor receptor 2 (KDR), which increase in expression in this study. ELF1 targets TIE1 and TIE2, also important factors in angiogenesis known to increase with endurance training in human skeletal muscle. The ETS family of transcription factors have been shown to interact with RUNX1 which has been identified as one potential key transcription factor in the regulation of the endurance training induced transcriptome. The mechanisms responsible for inducing these training specific effects are yet to be described. However, altered metabolism is known to induce epigenetic changes and metabolites from the citric acid cycle such as α-ketoglutarate are used as substrates for several of the enzymes known to catalyze methylation reactions.

In conclusion, this study demonstrates that the transcriptional alterations in skeletal muscle in response to a long-term endurance exercise intervention are coupled to DNA methylation changes. We suggest that the training-induced coordinated epigenetic reprogramming mainly targets enhancer regions, thus contributing to differences in individual response to lifestyle interventions. We provide a valuable and novel perspective on the fields of human physiology and environmental epigenomics, showing that a physiological health-enhancing stimulus can induce highly consistent modifications in DNA methylation that are associated to gene expression changes concordant with observed phenotypic adaptations.

Materials and Methods

Human physiological measurements

The study, labeled EpiTrain (“Epigenetics in Training”), was approved by the Ethics Committee of Karolinska Institutet and conformed to the Declaration of Helsinki. Twenty-three young, sedentary volunteers (Table 1) trained only one randomized leg during 3 months, and the other leg was used as an untrained intradividual control leg. Two one-legged knee-extension performance tests were conducted before and after the training period. Skeletal muscle biopsies from vastus lateralis were taken before and 24 h after the last training session from both legs. The post-training performance tests were conducted 3–6 d after the biopsies. Enzyme activity assays included citrate synthase (CS) and β-HAD activity. For details, see Supplementary Methods S9.

DNA methylation methods

The total amount of DNA methylation (at CCGG sites) in the genome, before and after training in both legs, was analyzed with LUMA (Luminometric Methylation Assay). Genome-wide DNA methylation profiling was generated with the Infinium HumanMethylation450 BeadChip array on bisulfite-treated DNA from biopsies collected before (T1) and after training (T2, trained leg), for 17 subjects in total. Bisulfite pyrosequencing was adopted for technical validation. We analyzed global hydroxymethylation with a colorimetric antibody-based
method from Epigentek according to the manufacturer’s specifications. More details are given in Supplementary Methods S9.

**Bioinformatics analysis of DNA methylation data**

We employed a pre-processing and normalization pipeline as reported previously. Color-bias adjustment, quantile normalization, probe type bias adjustment and batch correction were performed. For further details see Supplementary Methods S9 and Marabita et al., where the current dataset was named “Data set B.” Differentially methylated positions (DMPs) were defined using limma on M values, including the group (T2 vs. T1) and the subject as covariates, in order to account for the paired design. DMPs were selected if FDR < 0.05. For sample clustering, principal component analysis was performed using only data from the DMPs. Standard Illumina annotation was used to annotate methylation data. Additional sources included NIH Roadmap Epigenomic (http://www.roadmapepigenomics.org) data for skeletal muscle and ChromHMM chromatin segmentation tracks for HSMM cell line in Encode (http://encode-project.org). For each annotation category the relative fraction of positions located within each feature type was calculated for DMPs, non-DMPs and the entire array. GREAT (http://great.stanford.edu) was used to discover functional categories associated with DMPs. The enrichment for known transcriptional motifs was tested with HOMER (http://homer.salk.edu/homer/) and motifs were clustered using STAMP (http://www.benoslab.pitt.edu/stamp/index.php). Full details are given in Supplementary Methods S9.

**RNA sequencing**

Total RNA was used to prepare libraries, which were sequenced as paired-end, 2 × 100 bp on an Illumina HiSeq2000 and generated an average of 21 million paired-end reads per sample.

Briefly, we performed quality control, trimming, mapping, PCR duplicate removal and gene count summarization. EdgeR was used to normalize the data and extract differentially expressed genes (DEGs). We included the group (T2 vs. T1), the library preparation and the individual as covariates. DEG were selected if FDR < 0.05. Multidimensional scaling plots were used for sample clustering. Gene ontology analysis was done taking length bias into account, as implemented in goseq and enriched KEGG pathways were visualized with pathview. Full details are given in Supplementary Methods S9. The baseline muscle transcriptome has been described elsewhere, where a subset of the samples before training (n = 12) has been analyzed.

**Integration between DNA methylation and transcriptomics**

A list of genes and the corresponding measured DNA methylation positions was obtained and correlation was calculated for each pair. The distribution of Spearman rho statistics between DNA methylation and gene expression was calculated either including all pairs of genes/methylation positions or only pairs formed by a DMP and a DEG. Transcriptional networks were reconstructed from gene expression data applying the Mutual Information (MI) method developed in ARACNE. The resulting network components were visualized and analyzed with Cytoscape. For each independent network domain, gene ontology was tested using the BiNGO tool (http://www.psb.ugent.be/cbd/papers/BiNGO/Home.html). For additional information see Supplementary Methods S9.

**Data availability**

Data are available on GEO under the accession numbers GSE60655 (DNA methylation), GSE58608 and GSE60590 (gene expression).

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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**Author Contributions**

The study was designed and conducted by M.E.L. and C.J.S. H.R. and T.J.E. participated in design and conceptual discussions. F.M., D.G.C. designed and performed bioinformatic analysis of gene expression, DNA methylation and their integration, network reconstruction and analysis. D.G.C. and J.T. supervised the bioinformatic analysis and gave conceptual advices. M.E.L. performed molecular, physiological and statistical analyses. M.E. L. and F.M. drafted the manuscript. All authors participated in interpretation of data, manuscript preparation and have read and approved the final manuscript.

**Supplemental Material**

Supplemental data for this article can be accessed on the publisher’s website.
45. Mootha VK, Handschin C, Arlow D, Xie X, St Pierre J, Shiag S, Yang W, Ahshuler D, Puigserver P, Patterson N, et al. Erralpha and Gabpa/b specify PGC-1alpha-dependent oxidative phosphorylation gene expression that is altered in diabetic muscle. Proc Natl Acad Sci USA 2004; 101:6570-5; PMID:15100410; http://dx.doi.org/10.1073/pnas.0401401101

46. Oettgen P. Regulation of vascular inflammation and remodeling by ETS factors. Circul Res 2006; 99:1159-66; PMID:17122446; http://dx.doi.org/10.1161/01.RES.0000251056.85990.db

47. Timmons JA, Sandberg CJ. Oligonucleotide microarray expression profiling: human skeletal muscle phenotype and aerobic exercise training. IUBMB Life 2006; 58:15-24; PMID:16540428; http://dx.doi.org/10.1080/15216540500507390

48. Hollenhorst PC, Shah AA, Hopkins C, Graves BJ. Genome-wide analyses reveal properties of redundant and specific promoter occupancy within the ETS gene family. Genes Dev 2007; 21:1882-94; PMID:17652178; http://dx.doi.org/10.1101/gad.1561707

49. Lu C, Thompson CB. Metabolic Regulation of Epigenetics. Cell Metab 2012; 16:9-17; PMID:22768835; http://dx.doi.org/10.1016/j.cmet.2012.06.001

50. Bibikova M, Barnes B, Tsan C, Ho V, Klorze B, Le JM, Delano D, Zhang L, Schroth GP, Gunderson KL, et al. High density DNA methylation array with single CpG site resolution. Genomics 2011; 98:288-95; PMID:21839163; http://dx.doi.org/10.1016/j.ygeno.2011.07.007

51. Marabita F, Almgren M, Lindholm ME, Ruhrmann S, Fagerström-Billai F, Jagodic M, Sandberg CJ, Ekström TJ, Teschendorff AE, Tegnér J, et al. An evaluation of analysis pipelines for DNA methylation profiling using the Illumina HumanMethylation450 BeadChip platform. Epigenetics 2013; 8:333-46; PMID:23422812; http://dx.doi.org/10.4161/epi.24008

52. Teschendorff AE, Marabita F, Lechner M, Bartlett T, Tegnér J, Gomez-Cabrero D, Beck S. A beta-mixture quantile normalization method for correcting probe design bias in Illumina Infinium 450 k DNA methylation data. Bioinformatics (Oxford, England) 2013; 29:189-96; PMID:23175756; http://dx.doi.org/10.1093/bioinformatics/btt180

53. Smyth GK. Linear Models and Empirical Bayes Methods for Assessing Differential Expression in Microarray Experiments. Stat Appl Genet Mol Biol 2004; 3; PMID:16646809; http://dx.doi.org/10.2202/1544-6155.1027

54. Robinson MD, McCarthy DJ, Smyth GK. edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics (Oxford, England) 2010; 26:139-40; PMID:19910308; http://dx.doi.org/10.1093/bioinformatics/btp616

55. Young MD, Wakefield MJ, Smyth GK, Obhler A. Gene ontology analysis for RNA-seq: accounting for selection bias. Genome Biol 2010; 11:R14; PMID:20312535; http://dx.doi.org/10.1186/gb-2010-11-2-r14

56. Luo W, Brouwer C. Pathview: an R/Bioconductor package for pathway-based data integration and visualization. Bioinformatics (Oxford, England) 2013; 29:1830-1; PMID:23470750; http://dx.doi.org/10.1093/bioinformatics/btt285

57. Lindholm ME, Huo M, Solnesnø BW, Kjelleqvist S, Lundeberg J, Sundberg CJ. The human skeletal muscle transcriptome: sex differences, alternative splicing, and tissue homogeneity assessed with RNA sequencing. FASEB J 2014.

58. Basso K, Margolin AA, Stolovitzky G, Klein U, Dalla-Favera R, Califano A. Reverse engineering of regulatory networks in human B cells. Nat Genet 2005; 37:382-90; PMID:15778709; http://dx.doi.org/10.1038/ng1532