Title: Gene pathway development in human epicardial adipose tissue during early life

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#### Abstract

Studies in rodents and newborn humans demonstrate the influence of brown adipose tissue (BAT) in temperature control and energy balance and a critical role in the regulation of body weight. Here, we obtained samples of epicardial adipose tissue (EAT) from neonates, infants and children in order to evaluate changes in their transcriptional landscape by applying a systems biology approach. Surprisingly, these analyses revealed that the transition to infancy is a critical stage for changes in the morphology of EAT and is reflected in unique gene expression patterns of a significant proportion of thermogenic gene transcripts (~10%). Our results also indicated that the pattern of gene expression is a distinct developmental stage, even after the rebound in abundance of thermogenic genes in later childhood. Using weighted gene co-expression network analyses, we found precise anthropometric-specific correlations with changes in gene expression and the decline of thermogenic capacity within EAT. In addition, these results indicate a sequential order of transcriptional events affecting cellular pathways which could potentially explain the variation in the amount, or activity, of BAT in adulthood. Together, these results provide a novel resource to elucidate gene regulatory mechanisms underlying the progressive development of BAT during early life.

#### Introduction

The prevalence of childhood obesity has risen sharply during the past three decades, resulting in more than 42 million overweight children around the world (1). Animal and cell culture models have elucidated the pathogenic mechanisms of early life obesity and associated metabolic pathways (2, 3). However, to date, clinical trials based on these findings have not been translated into successful interventions (4). A potential reason for this is that the origin of metabolic diseases are rarely a consequence of an abnormality in a single gene but reflect perturbations in the complex web of intra-genomic regulatory activity, organ systems and their relationships to the environment (5, 6).

Elucidating the dynamic and functional changes which are occurring in adipose tissue at different stages during development is pivotal to understanding the detrimental consequences of childhood obesity. In early life, there is major transformation body energy usage as heat generating brown adipose tissue (BAT) cells are lost and are gradually replaced with white adipose tissue (WAT) (7). This process is especially important as the magnitude in reduction of thermogenic capacity is one of strongest phenotypic associations with metabolic related disorders (8). The impact of BAT on metabolic health primarily resides in its unique ability to rapidly produce heat and oxidise fat and glucose by the tissue-specific uncoupling protein (UCP)1 (9). Multilocular brown adipocytes, with cell numbers declining by age, were first identified in pericardial (presumably epicardial) fat in autopsies of infants and children (10). In adults, epicardial adipose tissue (EAT) consists of beige unilocular adipocytes which are smaller than white adipocytes and express abundant UCP1 mRNA and protein (11). Currently, no UCP1 immuno-histochemistry, gene expression profiles from microarray, or gene module, analyses have been described for any adipose tissue depot during early human development with congenital heart disease (CHD), although the molecular signatures have been described in a range of neonatal fat depots (12).

Following the rediscovery of BAT in human adults there has been an extensive search for UCP1expressing cells (13) including in EAT which has been shown to exhibit an intermediate (or 'beige') phenotype in adults (11). However, one of the most challenging aspects in unravelling the genomic changes that occur during early adipogenesis remains the practical difficulties in obtaining relevant samples from newborns and infants (14). To address this, through collaborations with cardiac surgeons, we collected EAT biopsies from pediatric patients. To analyse this data set, we used weighted gene co-expression network analyses to identify precise age-related specific changes in gene expression within EAT. Based on previous studies using animal models (15), we hypothesized that human EAT undergoes pronounced genotypic and phenotypic adaptations during the transition from the neonatal period through infancy and childhood.

#### RESULTS

### **Patient characteristics**

EAT samples were collected from children with CHD requiring cardiac surgery for clinical indications (Supplementary Table 1). For our analyses, we studied a total of 46 paediatric patients: 11 neonates (aged 6 to 24 days), 28 infants (aged 40 days to 12 months) and 7 children (aged 2 to 7 years). Physiological traits, such as weight and height, allowing calculation of Z scores for comparable parameters such as weight for age (WAZ) and height for age (HAZ), were recorded (Table 1; Supplemental Table 1). Growth Z scores were negative in the majority of participating children, reflecting the clinical status of the children with CHD approaching surgical intervention in our study cohort as although the intrauterine growth of the neonates was largely unaffected by their cardiac conditions, and early postnatal growth was appropriate in 14 out of 28 of the infants (defined as WAZ (-1.5< z-score)), there was greater variability in the WAZ and HAZ in children over 2 years reflecting, in part, the inherent difficulties of maintaining adequate ongoing nutrition.

# Histological and transcriptomic properties of EAT during the transition from the neonatal period to childhood

UCP1, the product of the primary BAT gene, was expressed in all EAT samples irrespective of age, whereas the histological appearance of UCP1 changed with age. In neonates, EAT was uniformly composed of multilocular, UCP1 positive adipocytes (Figure 1) whereas, in infants and children, it was predominantly composed of unilocular, UCP1 negative cells. Importantly, in 9/35 of those over 28 days of age, discrete islands of multilocular, UCP1 positive adipocytes were interspersed between white adipocytes.

Microarray gene expression analyses in EAT from infants (i.e. those aged 1-12 months) and children (aged 2-7 years) were similar (Supplemental Tables 2-5) but differed significantly from those of neonates (aged 0-24 days: FDR  $\leq$  0.05), with the difference being most marked between the neonatal and infant groups (Figure 2). Of the 1074 differentially expressed genes in EAT

during the transition to infancy (Supplemental Table 3), 103 transcripts were mitochondrial components involved in the activation of thermogenesis (p=5.8E<sup>-08</sup>), of which 83 were down regulated. Notably, the main marker of cell thermogenesis, *UCP1*, exhibited the largest reduction in gene expression with age (up to 3 log<sub>2</sub> fold-change; Figure 2C). However, there were only 127 differentially expressed genes between the EAT biopsies from neonates and children (Figure 2J), of which 107 genes were up or downregulated in the infant group (Figure 2A - Supplemental Table 4). Neither gene sets included *UCP1* (Figure 2G) or were significantly enriched with mitochondrial genes, indicating a possible repopulation of pro-thermogenic cells in this older age group.

Unsupervised clustering and principal component analyses revealed that a majority of the samples from the neonatal age group form a distinct gene cluster, possibly highlighting the existence of different gene-to-gene interactions (Figure 3A, B-D). Intriguingly, both cluster analyses concluded that, although there were large changes in gene expression, the neonatal and infant groups each clustered closely together but away from the majority of the samples from children. Consequently, this identified a distinct transcriptome pattern undetected by standard statistical procedures. Therefore, the conjunction of statistical and cluster analysis indicates the potential existence of transcriptional changes as well as the replacement of thermogenic cells occurring during the first year of postnatal life. These findings remain consistent with the hypothesis that, in infancy and childhood, adipose tissue passes through discrete developmental stages characterized by increased white adiposity (16) and reduced thermogenic capacity (10).

# Construction and description of the neonatal and infant EAT functional gene networks

To understand the shift of gene function in EAT during the early transition to infancy, we performed weighted gene co-expression network analysis (WGCNA). This unsupervised and unbiased algorithm allocated co-expressed transcripts to a specific gene module (17). By applying this algorithm, we increased the likelihood of identifying the activation of discrete

biological pathways (17). It yielded 7 distinct clusters in neonates and 6 for infants (Figures 4A; 4B; Supplemental Tables 6 & 7), each having variable degrees of connectivity (*k*), most markedly in the neonatal networks, affecting their topography (Figure 4A and B). Further analysis of the neonatal network demonstrated that it was organized in a distinct hierarchy in which the top 30 interconnected genes were allocated to the same module (represented by the neonatal-turquoise module in Figure 4A). This was characterized by a strong intra-module connectivity (*k*/N<sub>average</sub>=108.3). By contrast, connectivity patterns of the 6 modules from the infant network were much weaker, as illustrated in the infant blue module (*k*IN<sub>average</sub>=64.5). Strikingly, 76 of the 362 neonatal-turquoise module genes were functional mitochondrial genes, including *UCP1* (p=2.06E<sup>-22</sup>; Figure 5A), emphasizing the importance of mitochondrial regulation during early development.

# Integration of EAT network co-expression analysis and the relationship with neonatal and infant physiological outcomes

Next, we assessed the physiological relevance of each module by examining the overall correlation of the module genes with anthropometric indicators from each patient involved in our study. In the entire neonatal network, only the red module was correlated with any of the physiological traits, specifically, being negatively correlated with BMI Z-score (Figure 4C). This neonatal module contained 78 genes and included those with the lowest connectivity and cluster membership values in the network ( $kME_{average}=0.69$ ). The neonatal-red module was an exception in this network as other modules showed a strong interconnectivity between all the gene members. These results lead us to conclude that the collective expression and interactions between genes may play a strict role in the specific developmental physiological responses (e.g. thermogenic activation) in the neonatal age group. In contrast, half of infant modules (3 out of 6) were correlated with four different physiological traits (Figure 4D). The two strongest correlations were negatively related to HAZ score for the infant red (p=0.003, R<sup>2</sup>=-0.54) and infant yellow (p=0.001; R<sup>2</sup>=-0.59) modules. In addition, the infant red module in this network was positively

related to child age (p=0.01;  $R^2$ =0.46) and negatively with WAZ score (p=0.02;  $R^2$ =-0.43). The infant green module correlated positively with WAZ (p=0.03;  $R^2$ =0.41) and HAZ (p=0.02;  $R^2$ =0.44) scores (Figure 4D). These results may demonstrate that the infant gene network captures the state-dependent interaction between genes and their expression in response to environmental factors, such as nutrition and physiological stress such as is reflected in a negative Z score. Additionally, the lower connectivity observed in the infant network may be indicative of a specialised function within each module, thus increasing the importance of the interactions between peripheral modular and hub genes as potential translators of environmental adaptations.

# Changes in the biological significance of the networks, capturing a transition from a programmed to an environmental responsive state

The specialization of cell function is an important factor in adipose tissue maturation during the transition from the neonatal period to infancy. In order to address this process, we examined each module for over-representation of relevant functional gene classifications (Figure 5; Supplemental Table 8). Closer examination of the relationship between functional enrichment and network analysis indicated striking differences (Figures 6A and 6B). For instance, in the neonatal group, on the neonatal-turquoise module, were allocated 32/37 significant functional gene classifications, whilst only two classifications were allocated to either the neonatal-red or neonatal-tan gene modules (Figure 5A). In contrast, for the infant group with its relatively lower intramodule connectivity coefficients for inter-gene connections, a more even assignment of gene functional classes was possible between each module (varying from 1-15 classifications per module; Figure 5B). We were unable to assign a gene function in 2/6 modules.

For the neonatal network, however, no functional category was over-represented in 4/7 modules. The disproportionate distribution of gene function allocation between the modules in neonates might be attributed to the strength of its network organization. The infant gene modules exhibited a more specialized enrichment indicative of an enhanced internal genetic re-organisation and

transition to a more committed cellular function (i.e. lipogenesis). We, therefore, evaluated the impact of the gene enrichment analysis on the internal organization of both networks and applied the betweenness centrality coefficient (Cb(n)). Remarkably, acetylation (Cb(n)=0.453), had the highest value enrichment list during the neonatal period whilst, in the infant network, there was a shift to protein phosphorylation (Cb(n)=0.503). The transition in the dominant roles between these different biochemical processes with age is demonstrated when these features are displayed as a functional enriched gene network (Figured 6A and B).

Next, we explored the function of the hubs genes in each module (or the gene with largest number of connections to other genes, known in graph theory as 'degree') and its location in the networks. The genes with the highest degrees in the neonatal network were located on neonatal-turquoise module and included leucine zipper-EF-hand containing transmembrane protein 1 (*LETM1*; *k*IN=119.2). Further interrogation by the propinquity (i.e. nearness index) of gene connections to *LETM1*, determined that this list included genes such as solute carrier family 25 member 1 (*SLC25A1*; *k*IN=102.6) and cytokine induced apoptosis inhibitor 1 (*CIAPIN1*; *k*IN=104.6), which are key mitochondrial regulatory transcripts whose critical functions expand from organelle maintenance to respiration, influencing the activation of thermogenesis (18-20). In addition, *UCP1* (*k*ME=0.88) was allocated to the turquoise module in this age group, indicating the importance of the genetic regulation of thermogenesis immediately after birth (Figure 6A).

A similar analysis of the infant network revealed that the gene hub in this network is adaptorrelated protein complex 2 (*AP2S1*; *k*IN=68.5) located in the infant blue module. A brief inspection of the genes nearby to *AP2S1* in the blue module identified that these are mainly RNA transcripts of proteins performing similar functions (e.g. *GANAB*; *k*IN=63.4 and *COPE*; *k*IN=63.2). The phosphorylated proteins products of these genes have crucial roles in all phases of the protein translation and degradation process, including cargo selection and formation of vesicles (21, 22).

#### The activation of thermogenesis is closely linked to the child's ability to develop normally

We evaluated changes in the infant green module including associated thermogenic and other mitochondrial genes as well as UCP1. Although we cannot exclude that possibility hypoxia affected gene expression patterns and profiles, in our cohort (Table 1) we did not have sufficient the statistical power to examine this possibility. Critically, the majority of infants in the study were underperforming in growth (Table 1). Therefore, the positive relationships of the infant green module to WAZ and WHZ (Figure 4D) suggest that infants with a higher Z-score show less pronounced declines in UCP1, as well as other pro-thermogenic genes. Inhibition of cell thermogenesis and adipocyte differentiation occurs during periods of extreme food deprivation (23), indicating that these responses can be considered to be a normal response to environmental stress. Indeed, our analysis indicated that thermogenic genes are progressively replaced by other active genes associated with lipogenesis and tissue remodelling on the infant red module (24), such as PCOLCE2, ANG and ADIRF. However, the over activation of these transcripts may have long term adverse health consequences, as these genes are also associated with the progression of cardiovascular diseases and with insulin resistance (24). Our results indicate that gene expression patterns are not uniquely linked by biochemical pathways or cellular compartment location, but that other, yet unidentified, factors play an important role in the formation of gene modules.

Gene expression patterns are also influenced by a diverse range of factors including transcriptional binding-sites, transcription factors and mRNA pro-degradative elements (25). Thus, the infant red module contained a number of important genes, such as *SETD7* (*k*ME=0.89), *TWIST2* (*k*ME=0.89) and *EIF4EBP2* (*k*ME=0.8), which are transcriptional factors playing central roles in the transcriptional activation of genes involved in angiogenesis and insulin regulation (26-28). These results are in broad agreement with previous research on the physiological functions of EAT and its role in the development of cardiovascular disease (29). A widely accepted assumption is that mature EAT acts both as a store of free fatty acids for the myocardium and also displays some thermogenic capacity (11). However, it can also be subjected to a wide range

of pathological circumstances varying from lipodystrophy, poor cardiac contractility and obesity, with abnormal secretion of inflammatory and fibrotic factors and white adipocyte hypertrophy, leading to deterioration in cardiovascular function (30, 31).

#### Discussion

Here, we demonstrate, for the first time, how the use of contemporary systems biology approaches, e.g. through the construction of gene networks, can be used to elucidate the transcriptional function of EAT. This computational methodology, through the development of open source packages (e.g. WGCNA, DAVID and Cytoscape), represents a complementary method in the understanding of adipose tissue biology. Our results demonstrate that this approach can be used to subdivide genes into pathways on the basis of the regulation of their expression. By segregating genes into functional groups, we demonstrate that changes in the topology intra-modular networks can be very useful for identifying biologically important genes within these pathways. One key example is the neonatal network which was characterised by high connectivity hub genes regulating cellular activities, particularly those associated with cell differentiation and mitochondrial function. Therefore, one of the most striking observations from this study was the intricate co-ordination of multiple processes implicated in the control of thermogenesis in EAT of neonates, and the way in which these gene-to-gene interactions shift towards lipogenesis through infancy. Although our data in older children will need to be validated in larger studies, there appears to be a distinct thermogenic pathway, which is in state of latency in older children. These findings raise the possibility that this pathway may be amenable to recruitment recruited in studies of the prevention of childhood obesity and insulin resistance (32). Our analysis highlights the complexity and time-dependent nature of these adaptations and it is apparent that simply targeting single genes is likely to be of limited benefit in reactivating thermogenesis through UCP1 in humans.

Another finding observed in the infant network is the identification of a cluster of genes (the infant red module) which are potent stimulators of vascular remodelling and metabolic dysfunction (24, 26-28). This module revealed striking correlations, both positive and negative, between patient age and negative Z scores for growth, and may ultimately link EAT dysfunction to environmental factors which may promote cardiovascular diseases. However, as we were unable to define a functional gene category, it remains to be determined whether this is only a statistically significant

relationship or whether a common mechanistic association linking these genes exists. Interestingly, our analysis suggests that the EAT transcriptome in children is sensitive to longer term reduction of cardiac performance as shown in adults with advanced coronary artery disease (31).

Despite limitations inherent in the sample size in our cohort, participants clustered into relatively homogenous age groups allowing elucidation of distinct gene networks reflecting distinct biological entities. Our data, therefore, supports the interpretation of the existence of time-dependent trajectories in postnatal developmental pathways in adipose tissue. These results may reflect the unique endocrine environment during the transition from birth to early childhood (7, 10, 16, 19), which induces a complex series of events in the cells constituting EAT, including cell differentiation, proliferation and the change in primary function between heat production and energy dissipation to energy storage.

Whilst we have demonstrated variations in cellular heterogeneity intrinsic to EAT in the three agegroups, representative histological examinations (e.g. Figure 1) demonstrate that the tissue samples taken span all cell types, minimising any influence on expression analyses. As the cells composing the EAT respond to both functional and environmental challenges, they appear to apply different transcriptional regulatory mechanisms. These lead to a reduction, or more controlled thermogenic activity in some cells, whilst others simply become energy stores (19). Such heterogeneity in genetic responses may underlie the separation of genes into modules as well as the differences observed in the topology of each network. However, as we noted in our cohort, this transition needs a precise temporal coordination of gene expression which in turn is dependent on the nutritional environment.

In summary, the structure and architecture of EAT differs between the neonate, infant and child with profound regulatory effects on *UCP1* (Figure 6C). EAT retains discrete islands of UCP1

positive cells persisting into childhood. These thermogenic cells exhibit significant differences in their transcriptional architecture which could alter gene activation at different age ranges in early life. In the neonate, EAT has limited flexibility and responsiveness. With age and maturation, the depot changes to become more responsive to environmental stimuli. Furthermore, we expect that the intensity of these changes is the product of both cellular programmable events and external environmental factors. The range of processes in EAT which we have identified as age and growth-dependent, including acetylation, protein phosphorylation and mitochondrial function, have the potential to regulate adipose tissue function. These unique, developmentally-regulated gene interactions and metabolic-related pathways are potential targets for intervention strategies including diet and/or pharmaceuticals designed to promote BAT function.

#### Methods

# **Tissue collection**

With regulatory approval and informed consent, as detailed below, samples of adipose tissue overlying the right ventricle were collected between April 2012 and October 2013. Due to the variable aggregation and amount of EAT overlying the right ventricle, samples were collected from the region that was readily and safely accessible to sampling. EAT samples were taken immediately prior to the child undergoing cardio-pulmonary bypass to eliminate any potential effects of intraoperative therapeutic cooling on EAT. All children were cared for by their families prior to surgery at thermoneutrality, notably no child's residence in this central England location was equipped with air conditioning (the environmental temperature being 10.1°C). External environmental temperatures prior to surgery undertaken for children with CHD were not different between age groups.

Measures of height and weight were undertaken using standardized, validated equipment in regular clinical use. Data on participant's age, weight, height, medications and other medical conditions were recorded prospectively. Anthropometric assessments were performed using the Emergency Nutrition Assessment Tool and the World Health Organization Anthro (v 3.2.2).

# Immunohistochemistry

Formalin fixed tissue samples were rehydrated, loaded into Histosette II (Simport, Quebec, Canada) cassettes and processed by ethanol dehydration followed by xylene rinse. The samples were then embedded in paraffin wax by the Shandon Excelsior<sup>™</sup> tissue wax processor (Thermo Scientific, Waltham, MA, USA). Five micrometre tissue sections were cut from the samples using a sledge microtome (Anglia Scientific, Cambridge, UK) and mounted on to Superfrost<sup>™</sup> Plus slides (Menzel-Gläser Inc, Braunchweig, Germany). At least two slides per participant alongside a negative control were labelled with a random identifier and loaded into the Lecia BondMax<sup>™</sup> IHC slide processor (Leica Microsystem, Wetzlar, Germany) and run on an automated software

programme (Vision Biosystems Bond version 3.4A) using bond polymer refine detection reagents (Leica Microsystem) using 1:500 dilution of pre-optimised primary rabbit polyclonal antibody to UCP1 (ab10983, Abcam, Cambridge, UK) for the antibody positive samples.

### RNA isolation, quantification, and quality control

For RNA isolation, 100-1000 mg of EAT biopsies were dissected for analysis into 2ml of TRI<sup>®</sup> reagent (Sigma Chemical Co. Poole, UK). Total RNA was extracted using RNeasy Plus<sup>®</sup> kit (Qiagen, West Sussex, UK) according to the manufacturer's instructions. Total RNA quantity was measured with a NanoDrop ND-1000 Spectrophotometer (Thermo Scientific). Optical density ratios at 260/280 were all >1.9. Total RNA quality was assayed by Agilent BioAnalyzer RNA 6000 Nano Kit (Agilent Technologies, Santa Clara, CA, USA). Quality control required intact, distinct ribosomal peaks (RIN >7).

#### Transcriptome profiling with Affymetrix GeneChip

Sample mRNA was labelled and hybridized onto Human Genome U133A plus 2 arrays according to manufacturer's recommendations using the GeneChip® 3' IVT Express kit (Affymetrix, High Wycombe, UK). Detection used a GeneChip Scanner 3000 7G (Affymetrix).

# Expression array analysis

Normalization and network analyses were performed, unless otherwise stated, using free and open source packages, from the R project (<u>http://cran.r-project.org/</u>). The open source Bioconductor community (<u>http://www.bioconductor.org/</u>), provides specific resources for microarray analysis including the Human Genome U133A plus 2 arrays (Affymetrix). We used the function "gcrma" embedded in the "affy" package for pre-processing data stages, including background correction, normalization and probe match verification. For statistical analysis of gene expression, we used the "limma" library, which enabled us to perform empirical Bayesian statistical modelling (33) between all the three age groups. For all other statistical analysis, unless otherwise stated, we applied the false discovery rate (or FDR) (34) approach and

considered q≤0.05 as a significant result. The R-package "gplots" was used for fold change, heatmap, and volcano expression plots.

#### Accession numbers

All original microarray data were deposited in the NCBI's Gene Expression Omnibus (accession GSE82155).

#### Gene co-expression network construction and analysis

The core of network construction analysis was performed using the WGCNA (Weighted Gene Co-Expression Network) (17) R package for weighted correlation network analysis as previously described (17, 35, 36), using methods developed at the David Geffen School of Medicine (University of California, Los Angeles, USA).

The analysis was restricted to differentially expressed genes, removing those with low expression (i.e.  $\log_2$  fold change  $\ge 0.5$  or  $\le -0.5$ ; (uncorrected) p value  $\le 0.01$ , and an average gene expression of >4.5). Each probe set was treated as a unique gene. This method yielded the two gene networks subsequently used in our analysis. We identified gene transcripts outliers by applying a canonical Euclidean distance based network  $(A(uv)=1-||S(u)-S(v)||^{2}/maxDiss)$ , transformed to a standardized connectivity index  $(Z.k_u=(k_u-mean(k))/(sqrt(var(k)))$ .

To distinguish between gene repression and activation, thereby retaining biologically relevant information, we implemented signed co-expressed network algorithms. To maintain the scale free topology criterion for this analysis, we choose a  $\beta$  (soft threshold) value of 16, where we noted that R<sup>2</sup> reached a saturation point. Then, as described on the WGCNA analytical algorithm,  $\beta$  served as a power function for the Pearson's correlation formula ( $a(ij) = |0.5 + 0.5 * cor(x(i), x(j))|\beta$ )), to transform the similarity matrix (-1;1) into an adjacency matrix (0;1). The value of  $a_{ij}$  indicates the specific relationship or connection strength between an individual pair of genes. The

generated adjacency matrix was then applied as a measure of node dissimilarity on which the topological overlap matrix has been built.

We used the dynamic branch cutting method (0.25%) to define modules as branches of the hierarchical clustering tree. To allocate the 1041 selected genes into modules, we used "moduleManualhybrid" from the WGCNA R package (17). By calculating the module, and clinical characteristics, eigenvectors we could evaluate similarity or proximity of a specific gene to physiological trait(s); as well as deriving module membership eigenvectors (*k*ME) on interesting modules using Pearson correlation. Within-module connectivity (*k*IN) for each probe set was determined by summing the connectivity of that probe set with other individual gene members in the cluster of interest (17).

#### **Functional annotation charts**

The over-expression functional annotation analysis was performed individually for each gene module list, and analysed using DAVID (Database for Annotation, Visualization and Integrated Discovery <u>http://david.niaid.nih.gov/david/ease.htm</u>). DAVID combines results from various data mining tools including gene ontology (GO); Kyoto Encyclopedia of Genes and Genomes (KEGG); and SwissProt to attribute term and fold enrichment scores for each gene list filtered by statistical significance (i.e. FDR; q≤0.05)(37).

# **Network visualization**

For network visualization, WGCNA R gene expression profiles were imported into Cytoscape (open source v.3.0) (38). Each WGCNA file was subjected to statistical filters for fold changes in gene expression, *k*IN values and gene ontology enrichment analysis. Network figures shown here use the spring-loaded organic layout from Cytoscape to aid visualization.

# Statistics

For statistical analysis of gene expression, we used the "limma" library, which enabled us to perform empirical Bayesian statistical modelling (33) between all the three age groups. For all other statistical analysis, unless otherwise stated, we applied the false discovery rate (or FDR) (34) approach and considered q≤0.05 as a significant result. The R-package "gplots" was used for fold change, heatmap, and volcano expression plots.

#### Study Approval

The collection of epicardial adipose tissue samples in all participants in this study was conducted with ethical approval from the U.K. National Research Ethics Committee (NRES) (NRES East Midlands: Nottingham 2 REC Ref: 12/EM/0214). Additionally, institutional approval was granted by the University Hospitals of Leicester NHS Trust. Informed written consent was received from parents/guardians of participating children as well as from the young person themselves if they were deemed to possess the capacity to give informed consent. Tissue samples were transferred to Academic Child Health, University of Nottingham, a facility approved under the Human Tissue Act (HTA) UK 2004. All procedures were conducted within the HTA regulatory framework.

**Author contributions**: All authors contributed to the research and reviewed the manuscript. SO, HPF, MES & HB designed the study; AAL, GP, SO & VW obtained informed written consent, collected the samples and phenotypical data; SO performed Histological analyses; HPF, MC & SM performed the microarray analyses; HPF, SO, & MC undertook bioinformatic analyses. HPF, SO & HB prepared the table, figures and supplementary material. HPF, SO, MES and HB led the biological interpretation of results. SO, HPF, HB & MES wrote the manuscript. All analysis and writing group authors extensively discussed the analysis, results, interpretation and presentation of results.

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#### References

1. World Health Organisation. Fact sheet N°311. Web Site. http://www.hoint/mediacentre/factsheets/fs311/en/. Updated January 25, 2015. Accessed May 19, 2016..

2. Regazzetti C, et al. Hypoxia decreases insulin signaling pathways in adipocytes. *Diabetes*. 2009;58(1):95-103.

3. Sharkey D, et al. Maternal nutrient restriction during pregnancy differentially alters the unfolded protein response in adipose and renal tissue of obese juvenile offspring. *FASEB J*. 2009;23(5):1314-1324.

4. Han JC, Lawlor DA, Kimm SY. Childhood obesity. *Lancet.* 2010;375(9727):1737-1748.

5. Claussnitzer M, et al. FTO Obesity Variant Circuitry and Adipocyte Browning in Humans. *N Engl J Med.* 2015;373(10):895-907.

6. Lidell ME, et al. Evidence for two types of brown adipose tissue in humans. *Nat Med*. 2013;19(5):631-634.

Symonds ME. Brown adipose tissue growth and development. *Scientifica* (Cairo). 2013;
 Article ID 305763 doi: 10.1155/2013/305763.

Virtanen KA, et al. Functional brown adipose tissue in healthy adults. N Engl J Med.
 2009;360(15):1518-25.

9. Nedergaard J, Cannon B. The browning of white adipose tissue: some burning issues. *Cell Metab.* 2014;20(3):396-407.

10. Heaton JM. The distribution of brown adipose tissue in the human. *J Anat.* 1972;112(Pt 1):35-9.

11. Sacks HS, et al. Uncoupling protein-1 and related messenger ribonucleic acids in human epicardial and other adipose tissues: epicardial fat functioning as brown fat. *J Clin Endocrinol Metab.* 2009;94(9):3611-3615.

12. Shinoda K, et al. Genetic and functional characterization of clonally derived adult human brown adipocytes. *Nat Med.* 2015;21(4):389-394.

13. Cypess AM, et al. Identification and importance of brown adipose tissue in adult humans. *N Engl J Med.* 2009;360(15):1509-1517.

14. Stiers J, Aguayo C, Siatta A, Presson AP, Perez R, DiGeronimo R. Potential and Actual Neonatal Organ and Tissue Donation After Circulatory Determination of Death. *JAMA Pediatr.* 2015;169(7):639-645.

15. Symonds ME, Pope M, Budge H. The Ontogeny of Brown Adipose Tissue. *Annu Rev Nutr.* 2015;35:295-320.

Spalding KL, et al. Dynamics of fat cell turnover in humans. *Nature*. 2008;453(7196):783 7.

17. Zhang B, Horvath S. A general framework for weighted gene co-expression network analysis. *Stat. Appl. Genet. Mol. Biol.*. 2005;4:Article17.

18. Fedorenko A, Lishko PV, Kirichok Y. Mechanism of fatty-acid-dependent UCP1 uncoupling in brown fat mitochondria. *Cell.* 2012;151(2):400-13.

19. Rockstroh D, et al. Direct evidence of brown adipocytes in different fat depots in children. *PloS one*. 2015;10(2):e0117841.

20. Rosell M, et al. Brown and white adipose tissues: intrinsic differences in gene expression and response to cold exposure in mice. *Am J Physiol Endocrinol Metab.* 2014;306(8):E945-E964.

21. Ge L, Baskaran S, Schekman R, Hurley JH. The protein-vesicle network of autophagy. *Curr Opin Cell Biol.* 2014;29:18-24.

22. Le Roy C, Wrana JL. Clathrin- and non-clathrin-mediated endocytic regulation of cell signalling. *Nat Rev Mol Cell Biol.* 2005;6(2):112-126.

23. Fazeli PK, et al. FGF21 and the late adaptive response to starvation in humans. *J Clin Invest.* 2015;125(12):4601-4611.

24. Sun K, Kusminski CM, Scherer PE. Adipose tissue remodeling and obesity. *J Clin Invest.* 2011;121(6):2094-2101.

25. Wan Y, Kertesz M, Spitale RC, Segal E, Chang HY. Understanding the transcriptome through RNA structure. *Nat Rev Genet*. 2011;12(9):641-655.

26. Lee YS, et al. Twist2, a novel ADD1/SREBP1c interacting protein, represses the transcriptional activity of ADD1/SREBP1c. *Nucleic Acids Res.* 2003;31(24):7165-7174.

27. Martens JH, Verlaan M, Kalkhoven E, Zantema A. Cascade of distinct histone modifications during collagenase gene activation. *Mol Cell Biol.* 2003;23(5):1808-1816.

28. Pause A, et al. Insulin-dependent stimulation of protein synthesis by phosphorylation of a regulator of 5'-cap function. *Nature*. 1994;371(6500):762-767.

29. Sacks HS, Fain JN. Human epicardial adipose tissue: a review. Am Heart J. 2007;153(6):907-917.

30. Mazurek T, et al. Human epicardial adipose tissue is a source of inflammatory mediators. *Circulation*. 2003;108(20):2460-2466.

31. McAninch EA, et al. Epicardial adipose tissue has a unique transcriptome modified in severe coronary artery disease. *Obesity (Silver Spring)*. 2015;23(6):1267-1278.

32. Harms M, Seale P. Brown and beige fat: development, function and therapeutic potential. *Nat Med.* 2013;19(10):1252-1263.

33. Wettenhall JM, Smyth GK. limmaGUI: a graphical user interface for linear modeling of microarray data. *Bioinformatics*. 2004;20(18):3705-3706.

34. Benjamini Y, Hochberg Y. Controlling the False Discovery Rate - a Practical and Powerful Approach to Multiple Testing. *J Roy Stat Soc B Met.* 1995;57(1):289-300.

35. Horvath S, et al. Analysis of oncogenic signaling networks in glioblastoma identifies ASPM as a molecular target. *Proc Natl Acad Sci USA*. 2006;103(46):17402-17407.

36. Oldham MC, Horvath S, Geschwind DH. Conservation and evolution of gene coexpression networks in human and chimpanzee brains. *Proc Natl Acad Sci USA*. 2006;103(47):17973-17978.

37. Huang da W, Sherman BT, Lempicki RA. Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. *Nat Protoc*. 2009;4(1):44-57.

38. Su G, Morris JH, Demchak B, Bader GD. Biological network exploration with Cytoscape 3. *Curr Protoc Bioinformatics*. 2014;47:8 13 1-24.

# Figures



# Figure 1: Representative immunohistochemical detection of uncoupling protein 1 in a histological section of epicardial adipose tissue from a neonate.

A) Light microscopy (scale bar: 400µm) of UCP1 immune-stained epicardial adipose tissue typically observed in a neonatal sample demonstrating an area of positively stained brown adipocytes. B) Light microscopy (scale bar: 25µm) of UCP1 immune-stained epicardial adipose tissue demonstrating the histological differences between BAT and WAT during the neonatal period.



Neonatal period to infancy

Infancy to childhood

в				C				D	Log <sub>2</sub> Fold	-4	
Gene Symbol	Log₂ fold change	Q-value	Rank	Gene Symbol	Log₂ fold change	Q- value	Rank	Gene Symbol	Log <sub>2</sub> fold change	Q-value	Ra
PCOLCE2	2.45	3.8E-06	1	UCP1	-3.28	2.7E-02	756	ADH1B	2.78	2.12E-04	
C19 orf 80	-3.12	1.1E-04	2	SCD	-3.27	9.7E-03	318	RARRES1	2.71	5.74E-04	
SORBS1	2.25	1.1E-04	3	KCNK3	-3.26	7.4E-03	273	S11B	2.67	7.90E-04	
C6	2.65	1.7E-04	4	SPP1	-3.13	4.7E-03	179	C6	2.65	1.74E-04	
TMEM132C	2.13	1.7E-04	5	C19 orf 80	-3.12	1.1E-04	2	PPP1R1A	2.51	8.34E-04	
E				F				G			
Gene Symbol	Log₂ fold change	Q-value	Rank	Gene Symbol	Log₂ fold change	Q- value	Rank	Gene Symbol	Log₂ fold change	Q-value	R
CXCL14	3.35	6.0E-08	1	UCP1	-4.33	9.2E-02	281	AZGP1	3.72	9.1E-04	
FMOD	1.98	1.3E-07	2	DLK1	-4.20	1.8E-03	5	TF	3.54	1.1E-02	
GIPR	-2.63	3.7E-07	3	KCNK3	-3.58	8.3E-02	249	CXCL14	3.35	6.0E-04	
LRIG1	1.73	7.3E-07	4	CATSPERD	-2.97	2.6E-02	66	FGF2	3.30	8.3E-03	
DLK1	-4.20	8.8E-07	5	SPP1	-2.96	1.1E-01	406	S100B	3.24	1.0E-02	
	۱ • -	en de la composition de la com					J	52 - · · · · · · · · · · · · · · · · · ·		e Stational Stational Stationae	
	-log <sub>10</sub> p-value 2 3						anyan-d <sup>or</sup> doj-	R			
					•			0			
	-J	-2	-1 0	1 2	3			4 2	0	2 4	4
			Log <sub>2</sub> fold	l change					Log <sub>2</sub> fold change		

# Figure 2: Differential gene expression at distinct developmental periods in epicardial adipose tissue (EAT).

A) Venn diagram summarising the number of differentially regulated genes between neonatal and infant EAT (red) versus that between infants and children (green). B) Top 5 differentially expressed genes during the transition from the neonatal period to infancy (eBayes moderated tstatistics) and F) from the neonatal period to childhood. C) Genes with the largest decline in expression between neonatal period and infancy and G) between neonates and children (grey background: genes with more than 2.5 log<sub>2</sub> fold change but with false discovery rate (FDR) q>0.05. D) Top 5 genes with the greatest increases in gene expression between the neonatal period and infancy and H) between the neonatal period to childhood samples. I & J) Volcano plots showing changes in EAT gene expression between I) neonatal period and infancy and J) infancy and childhood (cutoffs: log-fc=1, log-p=1.3, q<0.05: FDR correction: eBayes moderated tstatistics).



Figure 3: Comparison of gene expression analysis in epicardial adipose tissue (EAT) between age groups.

A) Heatmap and unsupervised hierarchical clustering dendrogram for the top 100 differentially expressed gene transcript comparisons identified on microarray analyses of neonate (blue, n=11), infant (red, n=28), and children (green, n=11) EAT samples (average linkage, Euclidean distance metric; eBayes moderated t-statistics (q<0.05)). Gene expression was transformed to a Z-score, blue indicates an increase and red a decrease in gene expression by age. Principal Component Analysis (PCA) of EAT gene expression patterns for B) three age groups and individual interactions between C) neonatal and infant EAT and D) neonatal and children EAT samples. Every participant is represented by a sphere and is color-coded to indicate the specific age group to which they belong. The PCA was performed using probes (n=2454) with a p value <0.05 (without false discovery rate correction).



Figure 4: Summary of the strength of intra modular connectivity for co-expressed genes within neonatal and infant EAT and the relationships between the gene modules and clinical characteristics for each age group.

Pearson correlations were generated for the 1041 genes co-expressed in neonates (n=11) and infants (n=28) and compared. All pairs of genes were transformed into network connection strengths (denoted by colour intensity: yellow to red). Each cell represents an individual gene for either A) neonates or B) infants. Each cell shows the correlation coefficient (and p-value after Bonferroni correction) between the module eigenvector (rows) to clinical characteristic (columns) for C) neonates and D) infants. Significance (p<0.05) is highlighted in red. Coloured rectangles identify gene modules.

Gene module: Turquoise			B Gene module:Blue		
Term	Fold Enrichment	Gene Count	Term	Fold Enrichment	Gene
membrane	1.37	140	phosphoprotein	1.36	1
GO:0005739~mitochondrion	3.45	76	acetvlation	2.39	1
cetvlation	1.67	72	GO:0000166~nucleotide binding	1.75	
mitochondrion	4.33	59	GO:0005783~endoplasmic reticulum	2.56	
GO:0031090~organelle membrane	2.66	59	GO:0031090~organelle membrane	2.12	
transport	1.94	53	endoplasmic reticulum	3.56	
GO:0044429~mitochondrial part	4.31	52	GO:0031982~vesicle	2.60	
GO:0005783~endoplasmic reticulum	2.36	46	GO:0046907~intracellular transport	2.75	
GO:0031967~organelle envelope	3.26	41	topological domain: I umenal	3.48	1
30:0031975~envelope	3.25	41	GO:0044432~endonlasmic reticulum part	3.35	i i
andonlasmic reticulum	3 34	30	GO:0044092~pegative regulation of molecular function	3.53	i i
GO:0005740~mitochondrial envelope	4.48	39	GO:0051082~unfolded protein binding	7.44	
CO:0031966-mitochondrial mombrano	4.20	25	GO:0001002 - unioided protein binding	7.00	
ransit pontido	9.00	21	CO:0042770-mienant granule	7.00	
ransit peptide	3.90	20	CO.0046770-pigment granule	11.00	
ransit peptide.witochondrion	3.90	30	GO:0005755~ER-Golgi Intermediate compartment	11.02	
GO:00000742 mitochandrial incomponent	2.07	28	Constructule Pressure		
30:0003743~mitochondriai inner membrane	4.19	20	Gene module:brown	Fold Fasishment	0
GO:0019866~organelle inner membrane	3.90	20	Term	Fold Enrichment	Gene
GO:0008610~IIpid biosynthetic process	4.21	25	GO:0000739~mitochondrion	4.83	
GO:0031980~mitochondriai iumen	4.35	20	mitochondrion	0.49	3
30:0006769~mitochondrial matrix	4.35	20	transit peptide	8.51	2
mitochondrion inner membrane	5.06	16	GO:0044429~mitochondrial part	5.43	2
GO:0007005~mitochondrion organization	5.51	14	transit peptide:Mitochondrion	8.26	2
GO:0006839~mitochondrial transport	8.66	11	GO:0005740~mitochondrial envelope	4.50	1
somerase	5.60	11	GO:0031966~mitochondrial membrane	4.44	1
hsa03320:PPAR signaling pathway	6.09	10	GO:0008610~lipid biosynthetic process	5.71	1
lipid synthesis	6.23	10	GO:0005759~mitochondrial matrix	6.52	1
repeat:Solcar 1	10.11	9	GO:0031980~mitochondrial lumen	6.52	1
repeat:Solcar 2	10.11	9	GO:0007005~mitochondrion organization	11.14	1
IPR018108:Mitochondrial substrate/solute carrier	9.05	9	lipid synthesis	13.77	٤ ا
mitochondrion outer membrane	7.85	9	GO:0005783~endoplasmic reticulum	6.66	
repeat:Solcar 3	9.52	8	Cone module: Orean		
Gene module:Red			Term	Fold Enrichment	Gene
Term	Fold Enrichment	Gene Count	GO:0005739~mitochondrion	5.32	1
phosphoprotein	1.73	47	GO:0044429~mitochondrial part	8.18	1
cetylation	3.04	30	mitochondrion	6.85	1
			GO:0005740~mitochondrial envelope	10.17	1
Gene module:Tan			GO:0031967~organelle envelope	6.87	1
Term	Fold Enrichment	Gene Count	GO:0031975~envelope	6.85	i i
cetvlation	2.97	11	GO:0031966~mitochondrial membrane	9.27	i i
3O:0005739~mitochondrion	4.28	8	GO:0005743~mitochondrial inner membrane	7.96	i i
	4.20		mitochondrion inner membrane	12.92	i i
			Gene module:Yellow-green		
			Term	Fold Enrichment	Gene
Fold Enrichment			phosphoprotein	1 36	Jene 1
1 13			prosproprotein	1.00	1
Gene Count					
1 140					

# Figure 5: Biological processes and cellular components enriched in the EAT in each age network.

Hypergeometric enrichment lists, using the DAVID database, of biological processes or cellular components (q<0.05) found in each module of A) neonatal and B) infant gene networks using the 1041 probe sets allocated to each gene network. The enrichment analysis was performed using each gene network module probes separately. Normalized enrichment score, and prediction of fraction of genes affected in each term shown (additional details, p-values and full gene lists: Supplementary Material, Table 8).



- Thermogenic genes independent of environmental factors
  Greater gene interdependence with
- restricted gene-gene interactions
- Enhanced expression of acetylation (cell differentiation) genes

- Thermogenic genes related to markers of growth

- Greater diversity in gene regulation with flexible mature gene networks

- Enhanced expression protein

phosphorylation genes committed to lipogenesis

# Figure 6: Cytoscape pathway analysis of the change in primary networks within epicardial adipose tissue (EAT) with age and schematic summary of changes in adipose tissue composition, structure and primary regulatory pathways.

Comparative sub-network visualization for hub gene modular topology in A) neonates and B) infants. Complete networks for each age group are shown in the smaller diagram on each figure. Layouts are based on "organic distribution" and gene node shading depicts gene expression fold change. Red and green denote positive and negative gene expression respectively. Relative numbers of direct gene interactions are indicated by node size and edge colour (from positive-red to negative-green lines) describing the extents (strength, intensity, or capacity) of interactions between two genes (weighted correlation). C) Summary of effects on transcriptional architecture which accompany age from the neonatal period into childhood.

	Age Groups			
	Newborns	Infants	Children	
Number of participants	11	28	7	
Males	8	17	4	
Age in days (mean ± SD) (range)	10 ± 2 (3-24)	207 ± 18 (40-360)	1575 ± 267 (796-2612)	
Caucasian	8	23	7	
WAZ (mean ± SD)	-0.40 ± 0.41	-1.60 ± 0.29	-1.84 ± 0.96	
HAZ (mean ± SD)	-0.41 ± 0.56	-0.37 ± 0.36	-1.63 ± 1.16	
BMI z-score (mean ± SD)	-0.24 ± 0.36	-0.63 ± 0.20	-0.83 ± 0.30	
Baseline oxygen saturation <92% in air (n)	9	8	0	
Regular beta adrenergic antagonist therapy (n)	1	3	1	
External environmental temperature (mean ± SD)	12.4 ± 3.3	9.3 ± 4.2	9.2 ± 6.9	

# Table 1. Participant clinical and anthropometric characteristics

Demographic characteristics for children presenting for surgery for congenital heart disease in different age groups. WAZ: weight for age z-score; HAZ: height for age z-score; BMI: body mass index. Mean external environmental temperatures for the month preceding surgery. Anthropometric assessments were undertaked using the Emergency Nutrition Assessment Tool and the World Health Organization (v3.2.2, January 2011).