The Effect of **ACACB** cis-Variants on Gene Expression and Metabolic Traits

Lijun Ma¹, Ashis K. Mondal¹, Mariana Murea¹, Neeraj K. Sharma¹, Anke Tönjes², Kurt A. Langberg¹, Swapan K. Das¹, Paul W. Franks³, Peter Kovacs², Peter A. Antinozzi¹, Michael Stumvoll², John S. Parks¹, Steven C. Elbein¹, Barry I. Freedman¹

¹ Wake Forest School of Medicine, Winston-Salem, North Carolina, United States of America, ² Interdisciplinary Center for Clinical Research, Leipzig University, Leipzig, Germany, ³ Clinical Research Center, Malmo General Hospital, Lund University, Malmo, Sweden

**Abstract**

**Background:** Acetyl Coenzyme A carboxylase (ACACB) is the rate-limiting enzyme in fatty acid oxidation, and continuous fatty acid oxidation in Acacb knock-out mice increases insulin sensitivity. Systematic human studies have not been performed to evaluate whether ACACB variants regulate gene expression and insulin sensitivity in skeletal muscle and adipose tissues. We sought to determine whether ACACB transcribed variants were associated with ACACB gene expression and insulin sensitivity in non-diabetic African American (AA) and European American (EA) adults.

**Methods:** ACACB transcribed single nucleotide polymorphisms (SNPs) were genotyped in 105 EAs and 46 AAs whose body mass index (BMI), lipid profiles and ACACB gene expression in subcutaneous adipose and skeletal muscle had been measured. Allelic expression imbalance (AEI) was assessed in lymphoblast cell lines from heterozygous subjects in an additional EA sample (n = 95). Selected SNPs were further examined for association with insulin sensitivity in a cohort of 417 EAs and 153 AAs.

**Results:** ACACB transcribed SNP rs2075260 (A/G) was associated with adipose ACACB messenger RNA expression in EAs and AAs (p = 3.8 x 10⁻⁵, dominant model in meta-analysis, Stouffer method), with the (A) allele representing lower gene expression in adipose and higher insulin sensitivity in EAs (p = 0.04). In EAs, adipose ACACB expression was negatively associated with age and sex-adjusted BMI (r = −0.35, p = 0.0002).

**Conclusions:** Common variants within the ACACB locus appear to regulate adipose gene expression in humans. Body fat (represented by BMI) may further regulate adipose ACACB gene expression in the EA population.

**Citation:** Ma L, Mondal AK, Murea M, Sharma NK, Tönjes A, et al. (2011) The Effect of ACACB cis-Variants on Gene Expression and Metabolic Traits. PLoS ONE 6(8): e23860. doi:10.1371/journal.pone.0023860

**Editor:** Maria Moran, Hospital Universitario 12 de Octubre, Spain

**Received** April 1, 2011; **Accepted** July 26, 2011; **Published** August 26, 2011

**Copyright:** © 2011 Ma et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

**Funding:** This study was supported by the Wake Forest School of Medicine Development Fund and Translational Science Institute Synergy Fund (SCE,BIF), National Institute of Diabetes and Digestive and Kidney Diseases (SCE), National Institutes of Health, and General Clinical Research Center grant 1UL1RR029889 from National Center for Research Resources, National Institutes of Health. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Competing Interests:** The authors have declared that no competing interests exist.

* E-mail: lima@wfubmc.edu

**Introduction**

Acetyl-CoA carboxylase (ACC1/ACACA and ACC2/ACACB) catalyze the synthesis of malonyl-CoA, the substrate for fatty acid synthesis and a regulator of fatty acid oxidation. Increased malonyl-CoA concentrations inhibit carnitine palmitoyltransferase-1 (CPT1) activity, decreasing the rate of fatty acid entry into mitochondria and subsequent fatty acid oxidation [1]. ACACB is the key regulator of the fatty acid oxidation pathway [2] and Acacb knock-out mice are reportedly protected from obesity and diabetes induced by high fat/high carbohydrate diets [3]. Continuous fatty acid oxidation in adipocytes of Acacb knock-out mice is one factor contributing to their high insulin sensitivity [4]. Human gene expression studies suggest that **ACACB** is abundantly expressed in both oxidative and lipogenic tissues [5,6].

When dietary intake exceeds the storage capacity of adipose tissue, excess lipid is delivered as ectopic fat to skeletal muscle, liver, pancreatic β-cells, and cardiac muscle. The resulting organ dysfunction, known as ‘lipotoxicity’, results in impaired insulin action and glucose homeostasis, and ultimately type 2 diabetes; however, the cellular mechanisms and intermediates are not fully understood [7–11]. **Acacb** knock-out mice [2,3,4] and rats treated with Acaca or Acacb anti-sense nucleotide inhibitors [12] suggested a potential therapeutic target for **ACACB** in insulin resistance, obesity, metabolic syndrome, and type 2 diabetes. Alternations in nutritional status may also regulate **ACACA** and **ACACB** expression [13,14].

**ACACB** single nucleotide polymorphism (SNP) rs4766587 [15,16] is associated with an increased risk of metabolic syndrome. Interestingly, a common SNP rs2268388 within **ACACB** is reproducibly associated with type 2 diabetes-related proteinuria.
and end-stage renal disease in non-African American (AA) populations [6,17]. Animal studies [3] suggest that the lack of strong association between ACACB variants and obesity/diabetes may be masked by the cross-regulation of ACACB expression and hormonal/nutritional status in insulin-sensitive tissues [18,19]. Gene-nutrient interactions may further influence the expression of functional variants in ACACB and their relationships with insulin sensitivity [16]. The ACACB cis regulatory SNPs rs2075259 and rs2075263 were significantly associated with ACACB messenger RNA levels in skeletal muscle (p = 3.0×10^{-8} and p = 5.2×10^{-7}, respectively) [20].

We hypothesized that: (a) cis SNPs may regulate ACACB expression in insulin sensitive tissues and by consequence affect insulin sensitivity, and (b) body fat, represented by body mass index (BMI), may regulate ACACB expression in insulin responsive tissues. Therefore, we investigated the association between transcribed SNPs in ACACB and subcutaneous adipose tissue and skeletal muscle ACACB gene expression, as well as insulin sensitivity in European Americans (EAs) and AAs. We also tested for relationships between BMI and ACACB messenger RNA expression in adipose and skeletal muscle tissues.

Results

Demographic characteristics of study subjects are listed in Table 1.

**ACACB transcribed SNPs are associated with adipose ACACB messenger RNA expression**

Eleven ACACB SNPs were genotyped in 105 EA and 46 AA subjects who were non-diabetic at the time of adipose and skeletal muscle biopsies. The linkage disequilibrium (LD) plots are shown in Figure 1. The LD pattern in EAs was similar to HapMap Caucasians, but differed from AAs and HapMap Yoruba Africans (Figure S1). The patterns of ACACB LD in AAs reflected those reported in HapMap for Yoruba Africans, with the exceptions of rs2075260 and rs2075263 for which the observed level was midway between HapMap Caucasians and Yoruba.

In subcutaneous adipose tissue, SNPs rs7135947, rs2075259, rs2075260, and rs2075263 were nominally associated with ACACB messenger RNA levels from the combined sample of EAs and AAs, after adjusting for age, sex, BMI, and race (p = 0.02-0.05; additive and dominant/recessive models). However, association was not replicated in EAs and no evidence of association was observed in the combined biopsy sample for any of the tested SNPs (Table 2).

**Allelic expression imbalance of ACACB transcribed SNPs in lymphoblast cell lines**

To identify a potential regulatory effect of ACACB transcribed SNPs, we applied an independent method to test AEI for six of the eight coding variants in lymphoblast cell lines from 95 Utah EAs under controlled cell culture conditions. In this group, 18 subjects were heterozygous for rs2075260 (A/G). The mean (±SD) normalized percentage of G allele expression in cDNA was 56.08 ± 2.97%, significantly higher than observed in the genomic DNA background (49.96 ± 0.34%; p = 0.048; see Figure 3). Other SNPs were either not significant or not consistent with adipose/skeletal muscle gene expression.

**Association of insulin sensitivity with ACACB cis transcribed SNPs in the metabolic sample**

SNP rs2075260, which showed the strongest association between adipose ACACB expression for EAs and AAs in the biopsy sample, together with rs2075259 and rs2075263, which

| Sample | Race | n (M/F) | Age (yr) | BMI (kg/m²) | TG (mg/dl) | TC (mg/dl) | HDL-C (mg/dl) | S1 (×10^{-8} min^{-1} lU/ml^{-1}) |
|--------|------|---------|----------|-------------|------------|-----------|-------------|----------------------------------|
| Biopsy Sample | EA | 105 (41/64) | 40.3 ± 11.2 | 28.9 ± 5.91 | 132 ± 123 | 183 ± 35 | 53.7 ± 18.3 | 3.70 ± 2.04 |
| | AA | 46 (29/17) | 43.5 ± 9.25 | 30.0 ± 6.33 | 109 ± 65 | 175 ± 40 | 51.4 ± 12.7 | 3.29 ± 1.85 |
| Metabolic Sample | EA (all) | 417 (163/254) | 38.3 ± 10.3 | 29.3 ± 6.1 | NA | NA | NA | 5.85 ± 5.97 |
| | EA(AAR) | 293 (112/181) | 37.6 ± 9.85 | 29.8 ± 6.0 | NA | NA | NA | 4.08 ± 3.0 |
| | EA(UUT) | 124 (51/73) | 40.1 ± 11.1 | 27.8 ± 5.9 | NA | NA | NA | 9.70 ± 8.75 |

Data expressed as mean ± SD. EA, European American; AA, African American; AR, Arkansas; UT, Utah; M, male; F, female; NA, not available; TG, triglyceride; TC, total cholesterol; HDL-C, HDL cholesterol; S1, insulin sensitivity.

doi:10.1371/journal.pone.0023860.t001
showed strong association with insulin sensitivity in 62 previously evaluated non-diabetic subjects (40 EA and 22 AA) (20), were further genotyped in the metabolic sample (416 EA and 153 AA). Rs7135947 was also genotyped in this sample due to significant association with adipose ACACB expression in AAs from the biopsy study sample. The association results with insulin sensitivity are shown in Table 3. Only rs2075260 was nominally associated with insulin sensitivity in EAs after adjusting for age, sex, BMI, sibship, and cohort (Utah or Arkansas) \( (p = 0.04, \text{ additive model}) \). Allele G, associated with higher ACACB expression in the adipose biopsy sample (Figure 2) and AEI (Figure 3), was also associated with lower insulin sensitivity in EAs after appropriate adjustment (Figure 4). No significant association of this SNP was seen in AAs (additive \( p = 0.71 \); Table 3).

**Adipose ACACB gene expression correlates with BMI in EA**

Adipose ACACB messenger RNA expression level was negatively associated with age and sex-adjusted BMI in EAs \( (r = -0.35, p = 0.0002; \text{ Figure 5a}) \), but not in AAs (Figure 5b). Significant differences in BMI or SI were not found between EAs and AAs in...
| No. | SNP ID     | allele | 12_ exp. | 11_ exp. | 11_ SE | 12_ SE | 22_ exp. | 22_ SE | p_add | p_dom | p_rec | Ethnicity | p_add_c | Tissue        |
|-----|------------|--------|----------|----------|--------|--------|----------|--------|-------|-------|------|-----------|---------|---------------|
| 1   | rs2878960  | C/T    | 0.34434  | 0.147    | 0.131  | 0.162  | 0.373    | 0.174  | 0.81  | 0.85  | 0.003 | 0.94     | 0.45    | adipose       |
| 2   | rs2878960  | C/T    | 0.680851 | 0.238    | 0.244  | 0.253  | 0.485    | 0.330  | 0.81  | 0.71  | 0.0007 | 0.98    | 0.79    | adipose       |
| 3   | rs7135947  | C/T    | 0.852381 | 0.052    | 0.056  | 0.099  | 0.557    | 0.552  | 0.79  | 0.84  | 0.0007 | 0.98    | 0.72    | adipose       |
| 4   | rs2878960  | C/T    | 0.978309 | 0.125    | 0.125  | 0.166  | 0.256    | 0.351  | 0.21  | 0.24  | 0.0009 | 0.98    | 0.49    | adipose       |
| 5   | rs2878960  | C/T    | 0.989362 | 0.152    | 0.152  | 0.211  | 0.167    | 1.010  | 0.003 | 0.90  | 0.003  | 0.95    | 0.50    | adipose       |
| 6   | rs11065772 | C/T    | 0.828395 | 0.131    | 0.131  | 0.162  | 0.373    | 0.212  | 0.81  | 0.85  | 0.003  | 0.94    | 0.45    | adipose       |
| 7   | rs2241220 | C/T    | 0.904634 | 0.125    | 0.125  | 0.166  | 0.256    | 0.351  | 0.21  | 0.24  | 0.0009 | 0.98    | 0.49    | adipose       |
| 8   | rs2241220 | C/T    | 0.969574 | 0.042    | 0.042  | 0.056  | 0.065    | 0.125  | 0.81  | 0.71  | 0.003  | 0.95    | 0.50    | adipose       |
| 9   | rs2075259  | A/G    | 0.828395 | 0.131    | 0.131  | 0.162  | 0.373    | 0.212  | 0.81  | 0.85  | 0.003  | 0.94    | 0.45    | adipose       |
| 10  | rs2075259  | A/G    | 0.978309 | 0.125    | 0.125  | 0.166  | 0.256    | 0.351  | 0.21  | 0.24  | 0.0009 | 0.98    | 0.49    | adipose       |
| 11  | rs7135947  | C/T    | 0.828395 | 0.131    | 0.131  | 0.162  | 0.373    | 0.212  | 0.81  | 0.85  | 0.003  | 0.94    | 0.45    | adipose       |
| 12  | rs2075259  | A/G    | 0.978309 | 0.125    | 0.125  | 0.166  | 0.256    | 0.351  | 0.21  | 0.24  | 0.0009 | 0.98    | 0.49    | adipose       |

**Table 2.** *ACACB* messenger expression vs. genotype in adipose and muscle tissues in EAs and AAs.

EA, European American; AA, African American. P values were adjusted by age, sex, BMI.

p_add, p value under dominant model; p_dom, p value under dominant model; p_rec, p value under recessive model; p_add_c, p value of combining EA and AA by additional adjustment of race.

11exp, 12exp, 22exp: *ACACB* relative expression levels in subjects with genotype 11, 12, and 22 respectively. Freq1: Frequency of allele 1; SE: standard error; N: number of subjects.

doi:10.1371/journal.pone.0023860.t002
the biopsy sample (Table 4). However, adipose ACACB expression was significantly lower in EAs than AAs in the biopsy sample (Table 4).

Association of BMI, triglycerides (TG), total cholesterol (TC), and HDL cholesterol (HDL-C) with ACACB cis transcribed SNPs in the metabolic sample

No significant associations were identified with BMI, TG, TC or HDL-C for the selected SNPs in the metabolic sample. SNP rs2075260 showed the lowest p-values for association with BMI in EAs and AAs (dominant p = 0.19 and 0.10, respectively; Table S1); however, significant association was not observed in the combined EA and AA sample. No association was identified with TG, TC, and HDL-C (data not shown).

Discussion

ACAGB is a key rate limiting enzyme in mitochondrial fatty acid oxidation. In this study, we found that the ACAGB transcribed SNP rs2075260 was associated with adipose tissue ACAGB gene expression in EAs and AAs and with insulin sensitivity in EAs; the allele associated with lower gene expression was also associated with higher insulin sensitivity. This is consistent with the mouse model, which demonstrated that inhibition of Acacb improved insulin sensitivity [3]. These data suggest that ACAGB cis-acting SNPs may regulate gene expression in humans, potentially altering insulin sensitivity independent of BMI. Although the association of rs2075260 with insulin sensitivity was not statistically significant in the AA sample, the effect of this allele was similar in magnitude and in the same direction as that observed in EAs. This result is similar to that seen in an independent German Sorb population sample whose LD pattern was similar to HapMap CEU with the association between rs2075263 (G/T) and homeostasis model assessment of insulin resistance (HOMA-IR; p = 0.22, beta = 0.06, n = 793) (personal communication: Stumvoll M, 2011) [21], as well as in Diabetes Genetics Initiative (DGI) for HOMA-IR (p = 0.03, beta = 0.11, n = 1393), where the meta-p value of these two cohorts reached 0.01 and 0.02 for Stouffer and Fisher’s methods respectively with minor allele C indicating lower insulin sensitivity, consistent with rs2075260 in Table 3 where minor allele G represented lower insulin sensitivity in the AR and UT Caucasian metabolic sample. Our inability to detect association at this locus in AAs may reflect low statistical power in this small sample. Concerns about multiple testing are relevant in genetic studies. The number of SNPs to account for in this study is impacted by the fact that rs2075259, rs2075260, and rs2075263 are in high genotypic concordance in Caucasians. Given this, we presented the raw p values without adjustment for multiple comparisons.

The non-synonymous SNP rs2075260 (G/A) encodes an amino acid substitution Val2141Ile [NM_001093]. We are unable to predict ACACB protein structural changes caused by this amino acid substitution (http://snpeffect.vib.be/snp_main.php?id=34162381). We scanned this SNP using MatInspector (http://www.genomatix.de) and TFSEARCH (http://www.cbrc.jp/research/db/TFSEARCH.html) and have not identified potential anchoring loci for a transcription factor binding site at rs2075260. Therefore, we are unable to determine whether this SNP is a functional variant. However, rs2075259 (G/A), in high genotypic concordance with rs2075260, forms activating protein (AP1)/v-maf musculoaponeurotic fibrosarcoma oncogene homolog (MAF) anchoring sites for allele G (matrix similarity = 0.96-0.98), while allele A abolished these potential binding sites. As we demonstrated, the G allele of rs2075259 is associated with lower ACACB gene expression in adipose tissue (Table 2). AP1/MAF may act as a suppressor, or impact other repressors, of gene expression [22,23]. This is supported by the strong negative correlation between messenger level of ACACB and MAF/AP1S2 (AP1 subunit 2) in our adipose gene global expression study in non-diabetic subjects (unpublished data).
No SNP was associated with BMI in the metabolic sample (Table S1). However, adipose ACACB expression was negatively associated with age and sex-adjusted BMI in EAs (Figure 5a). This correlation cannot be explained by the known regulatory function of ACACB on body weight and insulin sensitivity in the mouse model [2,3,4]. We suspect that BMI may regulate adipose ACACB expression in EAs. Although this cross-sectional study cannot provide direct evidence, this hypothesis is supported by our observation that HepG2 cell ACACB expression was down-regulated after treatment with 1 mM palmitate [13]. It is conceivable that nutritional stress (e.g. diet-induced obesity and free fatty acid exposure) regulates ACACB expression. Down regulation of Acacb was observed in visceral fat tissue in rats fed a high-fat diet, while Juniperus chinensis extract significantly reduced this effect [14]. AMPK (AMP activated protein kinase) may be the link between ACACB gene expression and nutritional status [24]. ACACB activity [25] can be directly inhibited by AMPK. This may constitute a physiologic link between higher BMI and lower ACACB expression and supports the metabolic adaptation model proposed by Weyer et al based on energy expenditure, fat oxidation, and body weight regulation in Pima Indians [26]. The lack of association between adipose ACACB gene expression and BMI in AAs may be due to the smaller sample size or imprecision of BMI as a measure of adiposity. Alternatively, AAs may be less sensitive to nutritional stress in terms of regulating adipose ACACB expression. Adipose ACACB expression levels are significantly higher in AAs than EAs (Table 4). However, BMI and insulin sensitivity were nearly equivalent in EAs and AAs in the biopsy study sample (Table 4). This finding may reflect racial differences in ACACB expression in response to similar body fat. Different genetic or environmental factors may affect ACACB expression between EAs and AAs. ACACB variants have not been shown to be associated with BMI or diabetes using GWAS. However, functional variants (e.g. transcribed SNPs) may still regulate gene expression and impact insulin sensitivity. Our biopsy sample was modest and generated

Table 3. Association between selected transcribed SNPs and insulin sensitivity (SI) in the metabolic sample.

| SNP       | Allele (1/2) | Race | S_i vs.Genotype/n | genetic power* | P_add | P_dom | P_rec | P_add_comb |
|-----------|--------------|------|-------------------|----------------|--------|--------|--------|------------|
| rs7135947 | C/T          | EA   | 6.37±0.56         | 6.79±0.37      | 7.16±0.79 | 0.66   | 0.72   | 0.76       |
|           |              | AA   | 3.28±0.27         | 4.58±0.48      | 2.55±0.36 | 0.92   | 0.10   | 0.36       |
|           |              | n=416| 81                | 217            | 118    | 0.99   | 0.71   |
|           |              | n=153| 73                | 64             | 16     | 0.60   |        |
| rs2075259 | A/G          | EA   | 6.39±0.72         | 6.69±0.46      | 6.94±0.45 | 0.83   | 1.00   | 0.33       |
|           |              | AA   | 3.23±0.42         | 3.65±0.32      | 3.93±0.44 | 0.60   | 0.74   | 0.52       |
|           |              | n=411| 17                | 148            | 246    | 0.99   | 0.75   |
|           |              | n=149| 14                | 67             | 71     | 0.58   |        |
| rs2075260 | A/G          | EA   | 7.05±0.44         | 6.50±0.47      | 5.87±0.51 | 0.04   | 0.87   | 0.046      |
|           |              | AA   | 3.81±0.38         | 3.71±0.36      | 3.48±0.38 | 0.71   | 0.25   | 0.98       |
|           |              | n=415| 257               | 140            | 18     | 0.99   | 0.07   |
|           |              | n=152| 79                | 61             | 12     | 0.59   |        |
| rs2075263 | C/T          | EA   | 5.62±0.66         | 6.59±0.47      | 6.99±0.43 | 0.58   | 0.57   | 0.94       |
|           |              | AA   | 3.90±0.58         | 3.75±0.36      | 3.73±0.36 | 0.41   | 0.61   | 0.20       |
|           |              | n=416| 12                | 134            | 270    | 0.99   | 0.68   |
|           |              | n=152| 7                 | 59             | 86     | 0.59   |        |

EA European American; AA African American.

Data are presented as least squares mean±SEM, after controlling for age, sex, and BMI.

Location (Arkansas/Utah) is additionally adjusted for in EA.
P_add reflects p-value for the additive model; P_dom reflects p-value for the dominant model; P_rec reflects p-value for recessive model.
P_add_comb is additionally adjusted for race when EA and AA are combined for analysis.

*Genetic power was estimated to detect 10% of the variation in insulin sensitivity and other metabolic traits in the metabolic sample (assuming a type 1 error rate = 0.0005).
doi:10.1371/journal.pone.0023860.t003
approximately 86% power in EAs and 30% power in AAs to detect 20% of the variation in gene expression levels (assuming a type 1 error rate = 0.0005). Larger populations will be necessary to replicate these results. The lack of association of ACACB variants with BMI/diabetes may also reflect the masking effect of gene*gene and gene*environment interactions. A recent gene*environment interaction study of ACACB variants and metabolism suggested that uncovered causative variants may have been overlooked in published GWAS [16]. Acacb knock-out mice were only protected from obesity and diabetes when fed high fat/high-carbohydrate diets; when fed normal chow, significant differences in body weight were not observed [3].

Two independent groups reported that Acacb knock-out mice have similar body weights as wild type mice [27,28]. Molecular explanations for the phenotypic differences observed between Olson’s model of Acacb deletion and the original model (prone to resistant to obesity, diabetes and insulin resistance) published by Abu-Elheiga et al. [2] are not clear. Although the Acacb biotin-binding site was deleted in both models, different targeting strategies were employed. The targeting strategy employed in the original study replaced only the exon containing the biotin binding motif [2]. RNA splicing across the targeting cassette might leave the mRNA in frame, resulting in a mutated but otherwise intact protein lacking a catalytic domain. If produced, such a protein might potentially have “dominant negative” activity toward ACACA [27]. This may be supported by the effects of soraphen, an inhibitor of both ACACA and ACACB, improving peripheral insulin sensitivity in mice fed high-fat diets [29]. It suggests that lower ACACB expression alone may be insufficient to improve insulin sensitivity. As in the present study, association between rs2075260 and S1 was weaker compared to the association with gene expression. It is noteworthy that this result was based on transcript messenger level only, as protein level and enzyme activity data are not available yet. Limitations of this study included the small sample of AA subjects, lack of longitudinal follow-up data and lack of dietary intervention. These factors limit interpretation of gene expression effects. We also lack quantifiable life-style data to use as covariates in our database. Free fatty acid (FFA) levels fluctuate even with fasting. Healthy non-diabetic, non-drug or tobacco using subjects should be recruited on balanced diets for three days, with blood samples drawn after an overnight fast. We are not confident that our blood samples meet these criteria. Therefore, we elected not to measure FFAs in our sample. We were unable to test the effects of specific ACACB inhibitors on human insulin sensitivity. Finally, due to the large size of the ACACB gene (Ensembl transcript ID ENST00000338492: 55 exons spanning 150 kb), where 142 SNPs were available in HapMap database (NCBI builder 36), we prioritized the effect of transcribed SNPs on gene expression. The most significant SNP for adipose ACACB expression in the biopsy sample (rs2075260) was in high genotypic concordance with rs2075259 and rs2075263, which rank among the top SNPs in skeletal muscle eQTL mapping [20]. However, rs2075260 was not associated with skeletal muscle ACACB expression in the biopsy sample. This observation is not uncommon in terms of eQTLs in different cell types [30]. This could also result from different transcript forms presented across different cell types or variation in participant characteristics in the two study samples. In addition, the ACACB probe (Agilent expression array) applied in the eQTL mapping [20] was located at the 3’ UTR of the ACACB gene, while the expression primer set (for RT-PCR) used in the current study was designed across exon 13 and 14, in an attempt to cover all of the reported transcripts. Transcribed SNPs are a logical source for testing AEI as a complementary prioritization method for gene expression based on genotype. We chose lymphoblast cell lines for AEI studies instead of adipose tissue due to mixed sources of adipose RNA, as well as the restricted amount of tissue RNA available. It can be argued that lymphoblast cell lines are not the most relevant cell types to evaluate insulin action. However, our prior study revealed that lymphoblast cell lines provided confirmatory evidence for eQTLs in insulin-response tissues [31]. Our data suggest that the common transcribed SNP rs2075260 (A/G) in the coding region of the ACACB gene is associated with adipose tissue ACACB messenger RNA expression in EAs and AAs. The G allele, representing higher levels of gene expression, is also associated with lower insulin sensitivity in EAs. Body fat, represented by BMI, may serve as a “negative feed-back” signal down-regulating ACACB expression in adipose tissue, possibly as a mechanism of metabolic adaptation. Longitudinal intervention studies will assist in interpreting whether high ACACB expression is a risk factor for obesity and type 2 diabetes. Gene*gene interaction studies in larger cohorts will be helpful to identify any undetected diabetes or obesity genes that interact with ACACB. Analysis of alternative splicing of ACACB in adipose tissue and skeletal muscle may be helpful to address tissue-specific transcripts and regulatory SNPs involved in gene expression.

Materials and Methods

Subjects and phenotypes

The “biopsy sample” consisted of 105 EAs and 46 AAs from Little Rock, Arkansas who lacked diabetes based on ADA diagnostic criteria 2010 [32]. All participants underwent a screening visit during which height, weight, fasting blood lipids, and blood insulin and glucose concentrations (fasting, 30, 60 and 120 minutes after a standard 75-g oral glucose load) were measured, and adipose and skeletal muscle biopsies were performed using a Bergstrom needle under local (lidocaine) anesthesia at University of Arkansas for Medical Sciences (UAMS), Little Rock, Arkansas. Biopsy samples were immediately rinsed in normal saline, cut, and snap frozen in liquid nitrogen. Premenopausal women were studied in the follicular phase of the menstrual cycle.

The “metabolic sample” consisted of 440 non-diabetic EAs (417 with measures of insulin sensitivity) and 163 non-diabetic AAs (153 with measures of insulin sensitivity). AAs were recruited in Arkansas, whereas 293 EAs were recruited from Arkansas and 124 were siblings from 62 nuclear families of Northern European descent ascertained in Utah. Subjects in the previously described “biopsy sample” were a subset of the Arkansas “metabolic sample”. An insulin-modified (0.04 U/kg), frequently sampled...
intravenous glucose tolerance test (FSIGT) was performed, as reported [33]. Insulin sensitivity ($S_I$) was calculated from the FSIGT using either the MinMod (Utah sample) or MinMod Millenium (Arkansas Sample) programs [34,35]. These programs use the same algorithms and provide nearly identical estimates of $S_I$.

Figure 5. Adipose ACACB expression vs. adjusted BMI. Figure 5a Adipose ACACB expression vs. adjusted BMI in EA. Figure 5b Adipose ACACB expression vs. adjusted BMI in AA.

doi:10.1371/journal.pone.0023860.g005
Subjects provided written, informed consent under protocols approved by either the Institutional Review Board of the University of Utah Health Sciences Center or UAMS. Information from these subjects was de-identified and samples transferred to the Wake Forest University School of Medicine (WFUSM). This study was approved by the WFUSM Institutional Review Board.

Laboratory measurements

Insulin levels were measured using an immunochemilumino-

metric assay (Molecular Light Technology, Wales, UK) and plasma glucose by a glucose oxidase assay. Standard clinical assays (lips, glucose) were performed at LabCorp (Burlington, NC).

Genotyping

Eight common transcribed ACACB SNPs were genotyped in the biopsy sample (rs2078960, rs4766516, rs11065772, rs2300455, rs7135947, rs2241220, rs3742023, rs2075260), all with minor allele frequencies >0.05 in Caucasians and Yoruba Africans based on HapMap data (http://hapmap.ncbi.nlm.nih.gov/). Additionally, rs2268388 (significantly associated with diabetic nephropathy) [6, 17] and rs2075259/rs2075263 (two top SNPs for ACACB expression quantitative trait loci [eQTL] in skeletal muscle) [20] were genotyped. Genotyping of the biopsy sample was performed on a PSQ 96 Pyrosequencer (Biotage, Uppsala, Sweden). Additional genotyping in metabolic sample participants was performed by pyrosequencing or the ABI TaqMan assay (Applied Biosystems, Foster City, CA). Genotype distributions for all variants met Hardy-Weinberg expectations (p>0.01). The overall genotype call rate was above 98%. Sixty-nine duplicated QC samples were randomly distributed across genotyping plates of the metabolic sample to assure 100% reproducibility. For the biopsy sample, in addition to the fact that pyrosequencing is a highly reliable genotyping method which visualizes and quantifies both alleles of target SNPs, AEI requires genotyping of both genomic and cDNAs, which virtually serve as duplicates to assure accuracy.

Linkage disequilibrium plots (D’ and r2) were generated using the HaploView program (http://www.broadinstitute.org/haploview).

Gene expression

Total RNA was isolated from subcutaneous adipose tissue using the RNAeasy Lipid Tissue Mini Kit (QIAGEN, Valencia, CA) and from skeletal muscle using the Ultraspec RNA kit (Biotex Laboratories, Houston, TX). The quantity and quality of isolated RNA were determined by ultraviolet spectrophotometry and electrophoresis, respectively, using the Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA), and 1 μg was reverse transcribed using random hexamer primers with Qiagen reverse transcribed reagents (QIAGEN, Valencia, CA). All RNA samples from a single study population and tissue were reverse transcribed using the same kit on the same day. The standard curves were generated using pooled RNA from assayed samples. Primers were designed to capture most known splice variants where the amplicon spanned an intron. ACACB expression was measured by real time PCR (SybyrGreen) on an ABI 7500-Fast Real time-PCR system (Applied Biosystems, Foster City, CA) using 18S ribosomal RNA as a normalization standard. Primer sequences were as follows: 18S forward: ATCAACTTTCGATGG TAG-TCG, 18S reverse: TTCCTTGGATGTGGTAGCCG, ACACB-Forward: GGGCTCTCCGTGCTCCTTACA, ACACB – Reverse: CGTCTCTCTCTTGGGCAAACACAG [36].

Allelic expression imbalance (AEI)

Transformed lymphocytes were cultured from an independent sample of 95 unrelated HapMap Utah Caucasians (EAs) as reported [30]. Total RNA was extracted using the RNAeasy mini kit (QIAGEN, Valancia, CA), quantity and quality was assessed on an Agilent 2100 Bioanalyzer (Agilent Technologies, Inc, Santa Clara, CA). Unequal expression of ACACB alleles was sought as evidence for cis acting regulatory variants by comparing peak heights in individuals heterozygous for the synonymous coding SNPs rs2878960, rs4766516, rs2300455, rs7135947, rs2241220, and rs2075260 [37]. Rs11065772 and rs3742023 failed in assay design. Briefly, total RNA was reverse transcribed using random hexamers. Allelic specific quantitation of both cDNA and genomic DNA samples was determined using the same assay for pyrosequencing on a PSQ 96 Pyrosequencer (Biotage, Uppsala, Sweden) with peak height quantified using Allele Quantification software (Biotage, Uppsala, Sweden).

Statistical analysis

S1 was estimated from the insulin and glucose data using either the MinMod (Utah sample) or MinMod Millenium (Arkansas sample) programs. Gene expression levels were normalized to 18S RNA and the ratio was used in all calculations. Statistical analyses were performed using the SAS 9.1 software of the SAS Institute (Cary, NC). Correlations between ACACB expression and BMI or S1 were assessed using a general linear model after controlling potential confounders (age, race, sex, BMI). To approximate normality, the logarithm of ACACB expression was used in all analyses. Allelic specific expression was assessed by comparing the percentage of normalized genomic and cDNA expression on observed alleles using the method of Fogarty et al [38]. For the biopsy samples, a generalized linear model (GENMOD) was used to assess the association between genotype and ACACB gene

| Table 4. Comparison of ACACB expression, S1, and BMI between EAs and AAs in the biopsy. Sample. |
|-----------------------------------------------|
| EA | AA | P value | Adjustment |
|---|---|---|---|
| Adipose ACACB expression | 1.68±0.10 | 2.33±0.15 | 0.0004 | age, sex, BMI |
| Adipose ACACB expression | 1.76±0.10 | 2.37±0.17 | 0.002 | age, sex, BMI |
| SM ACACB expression | 0.83±0.03 | 0.93±0.06 | 0.15 | age, sex |
| SM ACACB expression | 0.83±0.03 | 0.93±0.06 | 0.15 | age, sex, BMI |
| SM ACACB expression | 0.80±0.04 | 0.90±0.06 | 0.18 | age, sex, BMI |
| S1 (× 10^-4 min^-1 [μU/ml]^-1) | 3.65±0.19 | 3.60±0.30 | 0.88 | age, gender, BMI |
| BMI (kg/m^2) | 28.1±0.54 | 29.6±0.86 | 0.15 | age, gender |

Data are least squares mean ±SE. The controlled covariates are listed in the adjustment column.

SM: skeletal muscle; EA: European American; AA: African American.

doi:10.1371/journal.pone.0023860.t004
expression after adjusting for age, sex, and BMI, although age and sex were not obvious confounders (Table S2). For analysis under an additive model, homozygotes for the allele (1/1), heterozygotes (1/2), and homozygotes for the allele (2/2) were coded to a continuous variable (0, 1, and 2). The dominant model was defined as contrasting genotypic groups 1/1 vs. 1/2+2/2. The generalized estimating equations (GEE) procedure was used to account for sibships in the metabolic sample in addition to age, sex, BMI, and cohort, since Utah subjects were family-based. We analyzed each SNP using the same genetic model for all analyses, irrespective of the study sample. A meta-analysis was performed using Stouffer [39] and Fisher’s method [40]. P-values <0.05 were considered to represent a nominal level of statistical significance.

Supporting Information

Figure S1  Linkage Disequilibrium (LD) plot of studied ACACB SNPs in HapMap. S1a: Linkage Disequilibrium (LD) plot of studied ACACB SNPs in HapMap Caucasians. S1b. Linkage Disequilibrium (LD) plot of studied ACACB SNPs Yoruba Africans. (TIF)

Figure S2  Adjusted ACACB expression by sum of eQTL-increasing alleles in adipose. Adjusted ACACB expression levels were controlled for age, sex, BMI, and ethnicity. eQTL-increasing alleles: G for rs2075260; C for 7135947. P values were obtained under general linear model (GLM). EA: European American; AA: African American; All: EA and AA combined.

(TIF)

Table S1  Association between selected transcribed SNPs and BMI in the metabolic sample. Data are least squares mean ± SE, after controlling age and sex. Cohort location (Arkansas/Utah) is additionally adjusted for in EA. (XLS)

Table S2  Summary of Type III testing for age and sex as individual coefficients of ACACB expression in the biopsy sample. EA: European American; AA: African American; DF: degree of freedom. (XLS)

Acknowledgments

The authors are grateful to DGI investigators for generously making their data publicly available, and thank Dr. Clifton Bogardus for his critical reading of this manuscript. This work is dedicated to the memory of the late Dr. Steven C. Elbein.

Author Contributions

Conceived and designed the experiments: SCE BIF LM PAA MM. Performed the experiments: LM AKM NKS KAL. Analyzed the data: LM PWF BT. Contributed reagents/materials/analysis tools: BIF PWF PK SKD JSP PAA. Supervised the project: MM. Performed the experiments: LM AKM NKS KAL. Analyzed the data: LM PWF BT. Contributed reagents/materials/analysis tools: BIF PWF PK SKD JSP PAA. Wrote the paper: LM. Contributed data: AT PK MS. In-depth review and editing: BIF PWF PK SKD JSP PAA. Supervised the study: BIF SCE.

References

1. Walk SJ, Abu-Elheiga L (2009) Fatty acid metabolism: target for metabolic syndrome. J Lipid Res 50(Suppl): S138–43.
2. Abu-Elheiga L, Matzuk MM, Aba-Hashema KA, Walk SJ (2001) Continuous fatty acid oxidation and reduced fat storage in mice lacking acetyl-CoA carboxylase 2. Science 291: 2613–2616.
3. Abu-Elheiga L, Oh W, Kordari P, Walk SJ (2003) Acetyl-CoA carboxylase 2 mutant mice are protected against obesity and diabetes induced by high-fat/high-carbohydrate diet. Proc Natl Acad Sci U S A 100: 10207–10212.
4. Oh W, Abu-Elheiga L, Kordari P, Gu Z, Shainkova T, et al. (2005) Glucose and fat metabolism in adipose tissue of acetyl-CoA carboxylase 2 knockout mice. Proc Natl Acad Sci U S A 102: 1384–1389.
5. Castle JC, Hara Y, Raymond CK, Garrett-Engele P, Ohwaki K, et al. (2009) ACAC2 is expressed at high levels in human white adipose and has an isoform with a novel N-terminus [corrected]. PLoS One 4: e4949.
6. Maeda S, Kobayashi MA, Araki S, Babbaz Ho, Freedman BL, et al. (2010) A single nucleotide polymorphism within the acetyl-coenzyme A carboxylase beta gene is associated with proteumina in patients with type 2 diabetes. PLoS Genet 6: e1000882.
7. Poutou V, Roberton RP (2008) Ghrelinolipotoxicity: fuel excess and beta-cell dysfunction. Endocr Rev 29: 351–366.
8. Boden G (1997) Role of fatty acids in the pathogenesis of insulin resistance and NIDDM. Diabetes 46: 3–10.
9. Kelley DE, Goodpaster B, Wing RR, Simoneau JA (1999) Skeletal muscle fatty acid metabolism in association with insulin resistance, obesity, and weight loss. Am J Physiol 277(Pt 1): E1139–1141.
10. SrivathLindeos A, Velhkaara S, Hakkinen AM, Goto T, Westerbacka J, et al. (2002) Fat accumulation in the liver is associated with defects in insulin suppression of glucose production and serum free fatty acids independent of obesity in normal men. J Clin Endocrinol Metab 87: 3023–3030.
11. Unger RH (2002) Lipotoxic diseases. Annu Rev Med 53: 319–336.
12. Savage DB, Choi CS, Samuel VT, Liu ZX, Zhang D, et al. (2006) Reversal of fatty acid metabolism in association with insulin resistance, obesity, and weight loss. J Lipid Res 50(Suppl): S138–43.
13. Das SK, Mondal AK, Elbein SC (2010) Distinct gene expression profiles characterize cellular responses to palmitate and oleate. J Lipid Res 51: 2121–2131.
14. Kim SJ, Jung JY, Kim HW, Park T (2008) Anti-obesity effects of Juniperus chinensis extract are associated with increased AMP-activated protein kinase expression and phosphorylation in the visceral adipose tissue of rats. Biol Pharm Bull 31: 1413–1421.
15. Szabo de Edelenyi F, Goumidi L, Bertrais S, Phillips C, Macmanus R, et al. (2008) Prediction of the metabolic syndrome status based on dietary and genetic parameters, using Random Forest. Genes Nutr 3: 173–176.
16. Phillips CM, Goumidi L, Bertrais S, Field MR, Copples LA, et al. (2010) ACC2 gene polymorphisms, metabolic syndrome, and gene-nutrient interactions with dietary fat. J Lipid Res 51: 3500–3507.
17. Tang SC, Leung VT, Chan LY, Wong SS, Chu DW, et al. (2010) The acetyl-coenzyme A carboxylase beta (ACACB) gene is associated with nephropathy in Chinese patients with type 2 diabetes. Nephrol Dial Transplant 25: 3931–3934.
18. Kim KH (1997) Regulation of mammalian acetyl-coenzyme A carboxylase. Annu Rev Nutr 17: 77–95.
19. Field DJ, Born E, Murthy S, Mathur SN (2002) Polymutated fatty acids decrease the expression of sterol regulatory element-binding protein-1 in CaCo-2 cells: effect on fatty acid synthesis and triacylglycerol transport. Biochem J 368(Pt 3): 853–864.
20. Cox NJ (2011) personal communication.
21. Tönjes A, Koriat M, Schénitz D, Dietrich K, Bolteker Y, et al. (2009) Genetic variation in GPR133 is associated with height: genome wide association study in the self-contained population of Sorbs. Hum Mol Genet 18: 4662–4666.
22. Balasubramanian S, Eckert RL (2007) Curcumin suppresses API transcription factor-dependent differentiation and activates apoptosis in human epidermal keratinocytes. J Biol Chem 282: 6707–6715.
23. Hansen A, Henderson S, Lagos D, Nitsenko L, Coutere E, et al. (2010) KSHV-encoded miRNAs target MAF to induce endothelial cell reprogramming. Genes Dev 24: 195–205.
24. Huypens P (2007) Adipokines regulate systemic insulin sensitivity in accordance to existing energy reserves. Med Hypotheses 69: 161–163.
25. Cho YS, Lee JT, Shin D, Kim HT, Jung HY, et al. (2010) Molecular mechanism for the regulation of human ACC2 through phosphorylation by AMPK. Biochem Biophys Res Commun 391: 187–192.
26. Weyer C, Pratley RE, Saffel DE, Bogardus C, Ravussin E, et al. (2000) Energy expenditure, fat oxidation, and body weight regulation: a study of metabolic adaptation to long-term weight change. J Clin Endocrinol Metab 85: 1087–1094.
27. Olson DP, Pulimood U, Cline GW, Shulman GI, Lewis BB (2010) Gene knockout of Acc2 has little effect on body weight, fat mass, or food intake. Proc Natl Acad Sci U S A 107: 7586–7603.
28. Hoehn KL, Turner N, Swarbrick MM, Wilks D, Preston E, et al. (2010) Acute or chronic upregulation of mitochondrial fatty acid oxidation has no net effect on whole-body energy expenditure or adiposity. Cell Metab 11: 79–76.
29. Schrurs M, van Dijk TH, Gerdling A, Hovinga R, Reinoudt DJ, et al. (2009) Soraphen, an inhibitor of the acetyl-CoA carboxylase system, improves peripheral insulin sensitivity in mice fed a high-fat diet. Diabetes Obes Metab 11: 967–991.
30. Dimas A, Deutsch S, Stranger BE, Montgomery SB, Borel C, et al. (2009) common regulatory variation impacts gene expression in a cell type-dependent manner. Science 325: 1246–1250.
31. Sharma NK, Langberg KA, Mondal AK, Elbein SC, Das SK (2011) Type 2 diabetes (T2D) associated polymorphisms regulate expression of adjacent transcripts in transformed lymphocytes, adipose, and muscle from Caucasian and African-American subjects. J Clin Endocrin Metab First published ahead of print November 17, 2010.
32. American Diabetes Association (2010) Diagnosis and classification of diabetes mellitus. Diabetes Care 33: S5–S11.
33. Elbein SC, Chu WS, Das SK, Yao-Borengasser A, Hausteit SJ, et al. (2007) Transcription factor 7-like 2 polymorphisms and type 2 diabetes, glucose homeostasis traits and gene expression in US participants of European and African descent. Diabetologia 50: 1621–1630.
34. Pacini G, Bergman RN (1986) MINMOD: a computer program to calculate insulin sensitivity and pancreatic responsivity from the frequently sampled intravenous glucose tolerance test. Comput Methods Programs Biomed 23: 113–122.
35. Boston RC, Stefanovski D, Moare Pj, Sumner AE, Watanabe RM, et al. (2003) MINMOD Millennium: a computer program to calculate glucose effectiveness and insulin sensitivity from the frequently sampled intravenous glucose tolerance test. Diabetes Technol Ther 5: 1003–1015.
36. Morse DI, Carroll D, Weberg L, Bergstrom MC, Ranger-Moore J, et al. (2005) Determining suitable internal standards for mRNA quantification of increasing cancer progression in human breast cells by real-time reverse transcriptase polymerase chain reaction. Anal Biochem 342: 69–77.
37. Wang H, Elbein SC (2007) Detection of allelic imbalance in gene expression using pyrosequencing. Methods Mol Biol 373: 157–176.
38. Fogarty MP, Xiao R, Prokunina-Olsson L, Scott LJ, Mohlke KL (2010) Allelic expression imbalance at high-density lipoprotein cholesterol locus MMAB-MVK. Hum Mol Genet 19: 1921–1929.
39. Stouffer SA, Suchman EA, DaVinney LC, Star SA, Williams RM (1949) How the volumes were produced. In The American Soldier, Volume I: Adjustment to Army Life PrincetonNJ: Princeton University Press. 45 p.
40. Fisher RA (1948) "Questions and answers #14". The American Statistician 2(5): 30–31.