Mitochondrial uncoupling proteins regulate angiotensin-converting enzyme expression: crosstalk between cellular and endocrine metabolic regulators suggested by RNA interference and genetic studies

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Uncoupling proteins (UCPs) regulate mitochondrial function, and thus cellular metabolism. Angiotensin-converting enzyme (ACE) is the central component of endocrine and local tissue renin–angiotensin systems (RAS), which also regulate diverse aspects of whole-body metabolism and mitochondrial function (partly through altering mitochondrial UCP expression). We show that ACE expression also appears to be regulated by mitochondrial UCPs. In genetic analysis of two unrelated populations (healthy young UK men and Scandinavian diabetic patients) serum ACE (sACE) activity was significantly higher amongst UCP3-55C (rather than T) and UCP2 I (rather than D) allele carriers. RNA interference against UCP2 in human umbilical vein endothelial cells reduced UCP2 mRNA sixfold ($P < 0.01$) whilst increasing ACE expression within a physiological range ($\leq 1.8$-fold at 48h; $P < 0.01$). Our findings suggest novel hypotheses. Firstly, cellular feedback regulation may occur between UCPs and ACE. Secondly, cellular UCP regulation of sACE suggests a novel means of crosstalk between (and mutual regulation of) cellular and endocrine metabolism. This might partly explain the reduced risk of developing diabetes and metabolic syndrome with RAS antagonists and offer insight into the origins of cardiovascular disease in which UCPs and ACE both play a role.

Keywords:
- ACE; association studies; endothelial cell; gene expression; genetics; uncoupling protein.

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Abbreviations
ACE, Angiotensin-converting enzyme; ACEI, Angiotensin-converting enzyme inhibitor; ADP, Adenosine diphosphate; Ang I, Angiotensin I; Ang II, Angiotensin II; ANCOVA, Analysis of covariance; ANOVA, Analysis of variance; ARB, Angiotensin II type 1 receptor blocker; AT1R, Angiotensin II type 1 receptor; AT2R, Angiotensin II type 2 receptor; ATP, Adenosine triphosphate; BAT, Brown adipose tissue; cDNA, Complementary deoxyribonucleic acid; Ct, Crossing threshold; D, Deletion; DNA, Deoxyribonucleic acid; H+, Hydrogen ion/proton; HMAP, Healthy Male Army Recruits Study; HUVECS, Human umbilical vein endothelial cells; GAPDH, glyceraldehyde 3-phosphate dehydrogenase; I, Insertion; LD, Linkage disequilibrium; mRNA, Messenger ribonucleic acid; O2, Molecular/diatomic oxygen; PCR, Polymerase chain reaction; RAS, Renin–angiotensin system; RNA, Ribonucleic acid; ROS, Reactive oxygen species; RT-PCR, Reverse transcription polymerase chain SACE, Serum angiotensin-converting enzyme; SD, Standard deviation; siRNA, Small interfering ribonucleic acid molecule; SNP, Single nucleotide polymorphism; T1DM, Type 1 Diabetes Mellitus Study; UCP, Uncoupling protein
Introduction

Both circulating and local (tissue/cellular) renin–angiotensin systems exist

As a pivotal component of the endocrine renin–angiotensin systems (RAS), angiotensin-converting enzyme (ACE) plays an important role in the regulation of the human circulation. Cleaved from an anchoring stalk on endothelial cells that line blood vessels, it is released into the circulation, acting upon the protein angiotensin I to yield eight amino-acid angiotensin II (Ang II). Ang II provokes release of the hormone aldosterone from the adrenal glands, which leads to salt and water retention by the kidney. It also causes constriction of small blood vessels in the arterial tree – actions which, together, serve to elevate blood pressure. In medical care, RAS antagonists, whether ACE inhibitors (ACEIs) or drugs antagonising the actions of Ang II at its type 1 receptor (AT1R blockers or ARBs), are widely used in the treatment of elevated blood pressure, or in the treatment of heart failure in which reduced vascular tone and reduced circulating volume may offer advantages.

But the RAS has functions far beyond this, being found in cells and tissues throughout the body. Here, Ang II action on cell surface AT1R and type 2 (AT2R) receptors affects the function of the cell that synthesised it (autocrine actions), or the function of nearby cells (paracrine actions). Ang II can also act as an intracellular signalling molecule (so-called intracrine action): some cells internalise Ang II made elsewhere, and others synthesise it de novo (reviewed in [1]). In this way, both circulating and tissue RAS can act independently or interact in the regulation of cell function.

ACE activity is influenced by naturally occurring variation in the ACE gene

Circulating and tissue ACE activity varies greatly between individuals, and common genetic variation in the ACE gene explains up to 40% of such differences. In particular, each of the two inherited ACE genes can exist in one of two forms. One form contains a small extra sequence of DNA (287 base pairs) and is known as the ‘insertion’ or ‘I’ variant (allele). If this fragment is missing, the gene variant is known as the deletion, or ‘D’, allele. In both the circulating/endocrine [2] and cellular [3] RAS, the ‘I’ allele is associated with lower ACE activity.

RAS play an important role in regulating metabolism in health and disease

One important function of local and endocrine RAS is in the regulation of cellular and whole-body metabolism. This they do in numerous ways, influencing, for example, the storage and release of fatty acid fuels from fat cells (adipocytes) [4]; regulating islet cells in the pancreas, which are responsible for releasing the hormone insulin and thus regulating uptake and use of glucose [5]; and regulating the uptake and use of carbohydrate fuel by the liver [6]. But Ang II influences more than the uptake of metabolic substrates. It increases liver, skeletal muscle and whole-body oxygen consumption in rodents [7–9]. Conversely, ARBs and ACEI reduce oxygen consumption related to renal sodium transport [10]. Human data are supportive of such metabolic roles: the ACE I allele is associated not only with lower circulating and tissue ACE activity but also with successful physical performance in hypoxic environments [11–14] and with enhanced training-related falls in skeletal muscle oxygen consumption per unit of external work [15,16].

These metabolic roles of RAS appear to influence the development of disease in humans. Genetically determined high ACE activity (marked by the ACE D rather than I allele) is associated with the development of metabolic syndrome (hypertension, diabetes and abnormal blood lipid profile) [17], whilst reducing RAS activity (by the use ACEIs or ARBs) also reduces the risk of people developing diabetes [18], or of them suffering a myocardial infarction (heart attack), clinical signs or symptoms of heart failure, stroke or death from a cardiovascular cause [19].

Ang II has direct effects on mitochondria

The cellular metabolic effects of RAS may be mediated, in part, by direct action of Ang II on the mitochondrial respiratory chain (reviewed in [20]). Mitochondria are the intracellular organelles responsible for generating the body’s energy currency, adenosine triphosphate (ATP). The respiratory or electron transport chain of the inner mitochondrial membrane consists of an assembly of several discrete electron carriers, which are grouped into complexes. Three of these complexes (complexes I, III and IV) work as oxidation–reduction-driven proton pumps: electrons derived from diverse metabolic substrates combine with molecular oxygen to form water, and the energy released drives the translocation of protons (hydrogen ions, H+) from the mitochondrial matrix, across the otherwise impermeable inner membrane, and into the intermembrane space. This results in a chemiosmotic gradient (a mitochondrial membrane potential) across the inner membrane, which drives the flow of these protons back into the matrix through ATP synthase, which produces ATP from adenosine diphosphate (ADP) and inorganic phosphate. When the membrane potential is high (for instance at rest when no useful work is being performed and the demand for ATP is low), complexes I and III are also able to produce reactive oxygen species (ROS), where diatomic oxygen (O2) combines with a single electron only to form superoxide rather than being fully reduced to water. These ROS can cause substantial cell damage [21]. Exogenously administered Ang II traffics to mitochondria [22,23], where outer mitochondrial membranes may express AT1Rs [24]. Ang II will then stimulate production of ROS, NADPH oxidase-dependent superoxide and ADP-independent respiration – which reduces the activities of complexes I and III. Mitochondria may have the capacity to endogenously synthesise Ang II [25–29].

Uncoupling proteins can ‘short circuit’ the mitochondrial membrane and reduce the membrane potential

The ‘coupling’, which connects substrate energy with the derived ATP is, however, incomplete – protons can flow back into the matrix in a manner disconnected from ATP synthesis. This is in part controlled by nuclear-encoded, mitochondrial-
targeted uncoupling proteins (UCPs), of which five mammalian forms are recognised [30]. Of these, UCP4 and UCP5 are principally neuronally expressed [31]. The remaining three (UCP1–3) have close sequence homology. Expression of UCP2 is recognised in tissues including white adipose tissue, liver, and cardiac and skeletal muscle, whilst that of UCP1 is limited to brown adipose tissue (BAT), and of UCP3 largely to BAT and skeletal and, to a lesser extent, cardiac muscle [32]. By allowing protons to flow back across the inner mitochondrial membrane without coupled ATP synthesis, UCPs reduce the membrane potential and help protect the cell from the generation of excessive ROS from complexes I and III [33,34]. Thus, expression of UCP2 is increased as ROS levels rise, in a negative feedback regulatory system [35]. Indeed, it has been hypothesised that UCPs evolved in response to severe ROS production and inflammation. Linkage disequilibrium (LD) refers to the finding that some gene variants are not randomly distributed in a population: being in proximity, they tend to ‘travel together’. This is the case for this UCP2 variant (Fig. 1), which is in complete LD with a second promoter variant (−2723T>A). These also explain 71% of the variation in messenger RNA (mRNA) transcript ratio associated with the deletion/insertion (D/I) of 45 base pairs of DNA (in exon 8 of the gene) [41]. The UCP2-866A allele is associated with lower gene transcription (expression) in somatic non-β cells [42] but more effective UCP2 gene transcription in pancreatic β cells (those that make insulin). In humans, the UCP2-866A allele is thus associated with reduced insulin sensitivity [42] and glucose-stimulated insulin secretion [43], diabetes in obese subjects [42], oxidative stress in diabetic people [44] and future cardiovascular risk in diabetic and non-diabetic people [44]. The UCP2 I allele is associated with reduced UCP2 mRNA stability [41], and with elevated body mass index [45] and variably with basal metabolic rate [46].

UCPs alter metabolism through mechanisms in addition to altering mitochondrial membrane potential

This said, UCP2 may have metabolic roles other than, or beyond those related to, proton gradient and ROS generation (well reviewed in detail in [37]). Indeed, it appears to increase fatty acid oxidation [38], whilst long-chain fatty acid intake is also known to induce UCP2 expression [39]. Further, UCP2 appears to play roles in the regulation of food intake [14,15], although the mechanisms by which such actions occur remain poorly understood [40]. One means by which to explore the physiological roles of UCPs is through the study of genetic variants that are associated with differences in their expression. Helpfully, even a common genetic variation in the UCP3/2 locus influences UCP expression.

Functional variation exists in the genes that encode the UCPs

A common variant exists in the UCP2 gene’s promoter region (the area that initiates gene transcription), comprising the presence of an adenine base (rather than the more frequent guanidine: UCP2-866G>A, rs659366). This variant lies in a region containing binding sites for transcription factors (factors that stimulate gene transcription) influenced by hypoxia and inflammation. Linkage disequilibrium (LD) refers to the finding that some gene variants are not randomly distributed in a population: being in proximity, they tend to ‘travel together’. This is the case for this UCP2 variant (Fig. 1), which is in complete LD with a second promoter variant (−2723T>A). These also explain 71% of the variation in messenger RNA (mRNA) transcript ratio associated with the deletion/insertion (D/I) of 45 base pairs of DNA (in exon 8 of the gene) [41]. The UCP2-866A allele is associated with lower gene transcription (expression) in somatic non-β cells [42] but more effective UCP2 gene transcription in pancreatic β cells (those that make insulin). In humans, the UCP2-866A allele is thus associated with reduced insulin sensitivity [42] and glucose-stimulated insulin secretion [43], diabetes in obese subjects [42], oxidative stress in diabetic people [44] and future cardiovascular risk in diabetic and non-diabetic people [44]. The UCP2 I allele is associated with reduced UCP2 mRNA stability [41], and with elevated body mass index [45] and variably with basal metabolic rate [46].

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Diagrammatic representation of the human UCP3 and UCP2 gene locus (chromosome 11q13) with the relative positions and linkage disequilibria (LD) of the three common genetic variants in this study: the UCP3-55C>T promoter variant, UCP2-866G>A promoter variant and the UCP2 D/I variant. UCP2 is approximately 7 kb downstream from UCP3. They share 72% sequence homology. The exonic structures are shown (rectangles) as well as the transcription start points (angled arrows), translation start codons (ATG) and translation stop codons (TGA). UCP3 has two translation stop codons, yielding either a short (TGA0) or long (TGA1) transcript. The short transcript does not code for a functioning protein. There are three ATG codons at positions 888, 924 and 948 within an open reading frame in exon 2 of UCP2, which could initiate translation of a putative short protein (depicted as ATGS). However, translation only starts at the ATG codon in exon 3 to yield UCP2 protein [47]. LD (or D') is the non-random association of alleles or genetic markers in a population and is measured between 0 and 1. When two alleles are completely randomly associated, D' is zero. When two alleles are completely linked or associated, then D' = 1. In the figure, D' is shown between the three gene variants for the HMAR (§) and T1DM (†) studies. In both studies, there was significant LD between the UCP2-866G>A and UCP2 D/I variants (D’ = 0.38 – 0.54; P = 0.0001) and negative LD (−0.31 to −0.26; P = 0.0005) between the UCP3-55C>T and UCP2 D/I variants, but no LD between the UCP3-55C>T and UCP2-866G>A variants. The common UCP3-55C allele therefore showed significant allelic association with the rare UCP2 I allele, and the latter with the rare UCP2-866A allele.
Meanwhile, a common variant in the regulatory 'promoter' region of the *UCP3* gene (the presence of a thymidine base rather than the more frequent cytosine: -55C>T, rs1800849) is in negative LD with the *UCP2* D/I polymorphism [48], the T allele being associated with obesity in a recessive manner [48] and with higher circulating low-density lipoproteins cholesterol levels [49]. For clarity, such LD is demonstrated in Fig. 1, which relates to our study populations described later. We have recently shown the combined presence of both the *UCP3*-55C allele and the *UCP2*-866A allele (the 'UCP3-55C/UCP2-866A haplotype') to be associated with gains in muscle performance following endurance training [50] in a similar pattern to that previously associated with the ACE I allele (lower ACE activity) [51].

ACE may regulate UCP expression. But do UCPs regulate ACE expression?

Some of the metabolic effects associated with ACE and Ang II (refer to the previous discussion) are now thought to be mediated through regulation of UCP expression: Ang II upregulates expression of UCP1 [52], UCP2 [53] and UCP3 [54]. However, an unexpected (and completely novel) finding made by our group suggested an inverse relationship: that UCP activity might also regulate ACE expression. We had previously reported a correlation between lower serum ACE (sACE) activity and decreased risk of hypoglycaemia in Danish diabetic adult, going on to show a similar association of the ACE I allele (i.e. genetically determined low sACE) with a lower risk of severe hypoglycaemia [55]. Low ACE activity is favourable for performance and hypoglycaemia awareness when substrate availability is limited, for example, during hypoglycaemia. As part of our ongoing work, we recently genotyped these subjects for the *UCP3*-55C>T polymorphism, finding an association between *UCP3* genotype and sACE activity. Such data suggest a role for UCPs in the regulation of ACE activity, rather than the (previously recognised) converse observation.

We sought to confirm the validity of the finding by performing more detailed genetic analysis in this sample of 210 type 1 diabetic study patients (≥18 years old; 117 women; mean age 45.5 ± 13.7 years; none taking RAS antagonists): the T1DM study. We then sought replication of our findings in a second completely independent dataset, comprising 250 Caucasian British healthy male recruits (mean age 19.4 ± 2.2 years): the HMAR study.

Finally, as sACE is derived from endothelial cells, we performed *in vitro* experimentation to confirm the co-expression of both ACE and *UCP2* mRNA in human umbilical vein endothelial cells (HUVECs) before determining whether changes in UCP expression (by RNA interference) could directly alter endothelial ACE expression.

Results

**Gene variants in the adjacent UCP2 and UCP3 loci are tightly associated**

Genotype distributions (Table 1) were in Hardy–Weinberg equilibrium (meaning that they are consistent with stability over generations) and similar to those previously reported in Caucasians [41,45,48]. There was significant LD between the *UCP2*-866G>A and *UCP2* D/I variants (D 0.38–0.54; P = 0.0003; Fig. 1) and negative LD (−0.31 to −0.26; P = 0.0005) between the *UCP3*-55C>T and *UCP2* D/I variants, as previously described in a South Indian cohort [48], but no LD between the *UCP3*-55C>T and *UCP2*-866G>A variants. The *UCP3*-55C allele therefore showed allelic association with the rare *UCP2* I allele, and the latter with the *UCP2*-866A allele. There was no association between any genotype and common descriptive population variables.

**sACE activity is strongly associated with UCP2/3 genotype**

sACE activity was normally distributed in both studies and weakly correlated with age (HMAR r = −0.19; P = 0.02, T1DM r = 0.12; P = 0.04) but no other population variables. Age-adjusted sACE was therefore used in subsequent analyses. As expected, sACE activity was higher in T1DM subjects, with levels similar to those of previously published data [56]. The wide distribution of ACE activities in the T1DM group may partly reflect differences in diabetic phenotype, such as the presence or absence of nephropathy [56].

In both studies, sACE activity differed by *UCP* genotype. For *UCP2*, the I allele was associated with significantly higher sACE activity in both the HMAR study (25.8 ± 8.3 vs. 29.2 ± 10.3 vs. 29.9 ± 7.3 nmol his-leu/ml/min for DD vs. DI vs. II, linear trend P = 0.04; I allele vs. DD homozygotes P = 0.02) and T1DM study (49.9 ± 15.5 vs. 50.5 ± 17.6 vs. 60.4 ± 24.2 nmol his-leu/ml/min; P = 0.04 analysis of variance [ANOVA]). Considering *UCP3*, the ACE activity for CC versus

| Gene variant | *UCP3*-55C>T | *UCP2*-866G>A | *UCP2* D/I |
|-------------|-------------|---------------|------------|
| HMAR        | 128/71/13   | 82/108/22     | 104/92/16  |
|             | 0.229 (0.189–0.269) | 0.358 (0.313–0.404) | 0.292 (0.249–0.336) |
| T1DM        | 116/78/16   | 84/98/28      | 116/76/18  |
|             | 0.262 (0.220–0.304) | 0.367 (0.321–0.413) | 0.267 (0.224–0.309) |
| P HMAR vs. T1DM | 0.53         | 0.97          | 0.74       |

There was no significant difference in genotype distribution between the two studies.
of this variation accounted for by the statistical interaction between UCP2-866G>A and UCP2D/I genotypes ($P = 0.001$).

The expected sACE activity phenotypic means according to estimated haplotypes (groupings of the different gene variants, found in combination) for study subjects in the HMAR and T1DM studies are depicted in Fig. 3. Uncommon haplotype groups with a frequency <0.05 in both studies were excluded from subsequent analysis (i.e. the UCP3-55T/UCP2-866A/UCP2I and the UCP3-55T/UCP2-866G/UCP2D haplotypes). There was insufficient statistical power to detect a haplotypic effect in predicting sACE activity in the HMAR study ($P = 0.24$), perhaps contributed to by the limited number in such haplotype groups and compounded by the different distribution on ACE activity observed in this group when compared with T1DM subjects (see section on Discussion). However, there was a significant haplotype effect in predicting sACE activity in the T1DM subjects ($P = 0.01$). The lowest sACE activity was seen amongst individuals with the UCP3/2 haplotype CAD in both studies. A single nucleotide polymorphism (SNP) change from the UCP3/2 CAD to CAI haplotype led to a significant mean increase in sACE activity of 10.0 nmol his-leu/ml/min in the T1DM study ($P = 0.008$). The UCP2 I allele has been associated with decreased mRNA stability, with expression dependent on variation in the UCP2G>A polymorphism [41]. Thus, genotypes associated with putative lower UCP2 expression were associated with higher sACE activity.

The association of UCP2/3 genotype with sACE activity might in part be mediated through UCP’s regulation of ACE expression

In theory, differences in UCP expression may drive systemic (whole body) physiological effects (such as changes in insulin secretion, or in inflammatory or immune responses), which might change sACE activity in ways yet unknown. Alternatively, changes in cellular UCP expression might more directly influence ACE expression in those same cells. We sought to clarify the latter hypothesis through in vitro experimentation. Specifically, we sought to determine whether cellular ACE gene expression (ACE mRNA levels) would alter when UCP expression (UCP mRNA activity) was reduced by introducing targeted ‘small interfering RNA’ molecules (siRNA). We performed this experiment in HUVECs, given that circulating ACE is derived from such vascular endothelial cells.

Firstly, the mRNA co-expression of $ACE$, $UCP2$, $UCP3$ and $GAPDH$ (a housekeeping gene) in HUVECs was confirmed using a two-step reverse transcription polymerase chain reaction amplification (RT-PCR). Both $UCP2$ and $ACE$ mRNA were found to be expressed abundantly in cultured HUVECs, whilst expression of $UCP3$ was confirmed at a lower level (Fig. 4).

Subsequent quantitative PCR confirmed that $UCP2$ mRNA expression was reduced at all time points (between threefold and sixfold, $P < 0.0001$, Fig. 5) in endothelial cells transfected with two different siRNAs against $UCP2$ (siRNA1 and siRNA2), when compared with scrambled siRNA (a negative control, not specifically targeted at UCP2). In keeping with data from our genetic study, this effect was associated with a significant increase in ACE expression, ACE mRNA rising at 30 and 48 h (1.2- to 1.4-fold and approximately 1.8-fold, respectively, $P < 0.001$).
Genetic variation in the UCP3/2 locus is associated with sACE activity

Ours is the first study to demonstrate an association of sACE with a genetic variant outside the ACE gene itself [57–59]. We showed that the UCP3-55C (rather than T) and UCP2 I (rather than D) alleles were associated with higher age-adjusted sACE activity, the interaction between the three genotypes accounting for 4.1% and 5%, respectively, of the inter-individual variation in age-adjusted sACE activity in two independent genotyping studies—one of healthy young UK Caucasian men and one of Scandinavian middle-aged diabetic people. The lowest sACE was associated with the UCP3-55C/UCP2-866G/UCP2D (UCP3/2 CAD) haplotype in both studies, an SNP change from UCP3/2 CAD to CAI increasing sACE by 10.0 nmol his-leu/ml/min in the T1DM. We are unable to confidently attribute the observed effects to a precise allele, given that the UCP3-55C allele showed allelic association with the rare UCP2 I allele, and the latter with the UCP2-866A allele. Even the haplotypes described may simply mark functional variation elsewhere in the cluster. The use of exome sequencing in substantially expanded cohorts is thus suggested.

The replication of the association of UCP genotype with sACE activity in two independent and diverse groups strengthens confidence in the validity of the findings. It is worthy to note that ACE activity was substantially higher in the T1DM group than in HMAR group—a finding unlikely related to the use of two different methodologies of ACE assay. Indeed, an established literature shows ACE activity (when assayed using the same commercial Sigma assay we used) to be elevated in diabetic people to levels commensurate with those we identified and with similar levels in controls to those we report [56].

Figure 3. A: The UCP3-55C>T, UCP2-866G>A and UCP2D/I haplotype frequencies are shown in descending order of frequency for both the HMAR and T1DM studies. The most common haplotype is the UCP3-55C/UCP2-866G/UCP2D (CGD) allele. Subsequent analysis was confined to common haplotypes with a frequency greater than 0.05. B: Predicted UCP3-55C>T, UCP2-866G>A and UCP2D/I haplotype effects on age-adjusted serum ACE (sACE) activity in the HMAR and T1DM study subjects. Data are mean ± SD for each single allele effect. Haplotypes with a frequency less than 0.05 are not represented in the figure because of the very small group size and resulting wide standard deviation for sACE activity. §There was a significant haplotype effect in predicting sACE activity overall in the T1DM subjects (P = 0.01), but not in the HMAR study. *The bracket shows the comparison in the T1DM study between those individuals with the CAD haplotype who had the lowest mean sACE activity and those with the CAI haplotype who had the highest mean sACE activity. A SNP change from the UCP3/2 CAD to CAI haplotype led to a significant mean increase in sACE activity of 10.0 nmol his-leu/ml/min in the T1DM study (P = 0.008). Individuals with the CAD haplotype in the HMAR study also had the lowest sACE activity but are not shown owing to the low sample size.

Discussion

Genetic variation in the UCP3/2 locus is associated with sACE activity

Ours is the first study to demonstrate an association of sACE with a genetic variant outside the ACE gene itself [57–59]. We showed that the UCP3-55C (rather than T) and UCP2 I (rather than D) alleles were associated with higher age-adjusted sACE activity, the interaction between the three genotypes accounting for 4.1% and 5%, respectively, of the inter-individual variation in age-adjusted sACE activity in two independent genotyping studies—one of healthy young UK Caucasian men and one of Scandinavian middle-aged diabetic people. The lowest sACE was associated with the UCP3-55C/UCP2-866G/UCP2D (UCP3/2 CAD) haplotype in both studies, an SNP change from UCP3/2 CAD to CAI increasing sACE by 10.0 nmol his-leu/ml/min in the T1DM. We are unable to confidently attribute the observed effects to a precise allele, given that the UCP3-55C allele showed allelic association with the rare UCP2 I allele, and the latter with the UCP2-866A allele. Even the haplotypes described may simply mark functional variation elsewhere in the cluster. The use of exome sequencing in substantially expanded cohorts is thus suggested.

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The use of two different methodologies relates to the different geographical locations of sample acquisition and analysis. Both techniques are considered reliable and are well validated. A small systematic bias, related to the assay used, would not in any way affect the findings or conclusions of our study.

**UCP2 RNA interference increases ACE mRNA expression**

Most importantly, however, our data suggest that this association may partly result from UCP2's modulation of ACE expression. We confirmed the expression of UCP2 reported in endothelial cells of diverse origin [60–63]. Those of the vasculature are central to the pathophysiology of atherosclerosis. Indeed, UCP2 genotype is associated with the development of vascular disease [44], whilst ACE inhibition appears protective [64].

In HUVECs, a threefold to sixfold reduction in UCP2 mRNA (induced by transfection with two separate and different siRNA against UCP2) was reproducibly associated with an increase in ACE mRNA expression (1.2–1.8-fold at 30 h, and a near doubling at 48 h, \( P < 0.001 \)) – the scale of altered ACE expression (1.2–1.8-fold) by UCP2 RNA interference being similar to that observed in the physiological context [65–68]. We cannot discount an additional influence on ACE stalk cleavage, however, and would advocate further investigation of this possibility.

**Is there negative feedback between UCP2 and ACE at the cellular level?**

UCPs regulate mitochondrial metabolism. Mitochondrial regulation by RAS [7,9,20,25,69] may partly be mediated through modulation of UCP expression: Ang II upregulates expression of tissue UCP1 in brown fat [52] and UCP2 expression in brain [70], pancreatic beta-cells [53,71], and hepatic [72], renal [73] and cardiac [74] tissue. ACE inhibition reduces UCP2 expression in adipose and cardiac tissue [68,75], whilst adipose
UCP3 expression is reduced in AT1R knockout rodents [54]. However, Ang II’s influence on UCP expression may be both tissue and context dependent, given the existence of seemingly conflicting data: in other studies, Ang II action at the AT1R reduces cardiac UCP2 and UCP3 expression [76]; adipose UCP1 expression is elevated in AT1R gene knockout mice [77] and those treated with AT1R antagonists [78]; whilst ACE inhibition may have no effect on cardiac/skeletal muscle UCP3 upregulation in response to thyroid hormones [79] or may actually increase UCP2 expression in retinal tissue [80].

Meanwhile, RAS influences whole-body metabolism in other ways (e.g. through altered adipocyte function [4] and hepatic metabolism [6]). Given this, our data suggest several hypotheses. Firstly, ACE-UCPs feedback control may exist: this would function by Ang II altering mitochondrial function through changes in UCP expression, the latter feeding back to change Ang II synthesis through regulation of ACE expression. Secondly, UCPs might be considered to regulate mitochondrial metabolism through changes in coupling, but also through independent mitochondrial effects of Ang II. Such hypotheses would be supported were UCP expression is shown to regulate that of ACE in cells beyond those derived from the endothelium.

The regulation of endocrine sACE by cellular UCP may represent a novel interface between local cellular metabolism and endocrine metabolic regulation

UCP2 negatively regulates both pancreatic β cell glucose-stimulated insulin secretion [81] and central neuronal glucose control of peripheral glucose utilisation [82]. Fatty acids induce both the expression and activity of UCP2 in pancreatic β cells [83,84], and superoxide activation of UCP2 is dependent on the presence of fatty acids [35]. Skeletal muscle UCP2 and UCP3 mRNA expression is induced by a high fat diet [85], and plasma free fatty acid levels are positively correlated with cardiac UCP2 and UCP3 protein levels [66]. Starvation leads to the depletion of glycogen stores and an increased reliance on energy transduction from fatty acid oxidation. A similar period of starvation reduces sACE activity [86] and also increases UCP expression in skeletal muscle [65] – possibly as a protective mechanism against oxidative stress [35]. Furthermore, lower ACE activity in HUVECs promotes a survival advantage during starvation [87]. Perhaps of greater relevance, RAS antagonists prevent diabetes and improve insulin sensitivity (reviewed in [18]), whilst the RAS appears to play a role in the pathogenesis of the ‘metabolic syndrome’ (hypertension/dyslipidaemia) [57,88,89]. Indeed, increased RAS activity is common in obesity and is considered a possible link with associated diabetes and hypertension [89]. In this regard, elevated RAS activity in adipocytes may play a particular role [90], raising the prospect that the recognised association between UCP2/3 genotype and obesity/diabetes [41,42,45,46,49,91,92] might be partly adipose-RAS mediated. This does not discount, of course, a role for elevated RAS activity of endothelial (and UCP regulated) origin also playing a role. However, we would advocate further experiments to explore the influence of adipocyte UCP expression on that of ACE.

Similarly, UCP3-55C/UCP2-866A haplotype is associated with training-related gains in muscle performance [50], and it is tempting to speculate that some of this effect might be ACE mediated, given the existence of similar associations with ACE genotype [51].

Future directions

Whilst confident in the validity of the observations we have made, the implications remain speculative. Further studies are also required to explore the means by which UCP might alter ACE expression: is it direct or mediated, for instance, via altered ROS activity? Inhibition of UCP2 or UCP3 could reduce ROS and thereby increase ACE activity. Might UCPs additionally alter sACE activity through ACE stalk cleavage and thus increase release into the circulation? In such regards, real-time PCR analysis of the expression of other RAS components might be performed in HUVECs, transfected in the manner that we have described. RAS activity can prove protective to blood vessels (especially, perhaps, through activation of AT1R rather than AT2R) – and UCPs role in regulating such vasoprotective effects is worthy of exploration (given that reduced ROS activity would also be vasoprotective). Recent data also suggest a role for mitochondrial-associated membrane RAS in regulating mitochondrial function – and the study of ACE activity in such regions in response to altered UCP expression might also be suggested. Exome sequencing may help identify the precise functional UCP2/3 variant affecting sACE activity. Experimentation should also be extended to other cell types.

Materials and methods

Genetic studies

The association of UCP3/2 genotype with sACE activity was explored in two distinct cohorts of European Caucasians: young healthy British men and adult Danish patients with type 1 diabetes. Ethics committee approval was obtained, as was written informed consent from all participants, with study protocols adhering to the Declaration of Helsinki (Defence Medical Services Clinical Research Committee at the Army Training Regiment, Basingbourn, UK, and Hillerød Hospital, Hillerød, Denmark).

Subjects

The HMAR study sample consisted of 250 consecutive healthy male Caucasian British Army recruits (mean age 19.4 ± 2.2 years), as previously described [93], 212 of whom had complete genotypic data.

The T1DM patients study comprised 262 consecutive Danish adults (≥18 years old; 117 women; mean age 45.5 ± 13.7 years) with a >2 year history of T1DM, as previously described [94]. Analysis was restricted to the 210 with complete genotype data who were both ACEI and ARB naïve.

Genomic DNA was extracted from 10 ml EDTA venous blood. Serum was aspirated from centrifuged citrated blood and stored at −20°C for subsequent blinded analysis of sACE.
Measurement of sACE activity

For the HMAR study, sACE was assayed with a modified fluorometric method using carbobenzoxy-phenyl-alanyl-histidyld-leucine (Z-phe-his-leu) as a substrate [95]. Interassay and intraassay coefficients of variation were 13% and 11%, respectively. For the 1DM study, sACE was measured using a commercial kinetic assay (Sigma Diagnostics, St Louis, MO, USA) based on the hydrolysis of the synthetic tripeptide N-[3-(2-furyl)acryloyl]-l-phenylalanylglycylglycine to furylacryloylphenylalanine and glycylglycine. ACE activity was determined by comparing sample reaction rate with that derived from an ACE reference. The use of two different methodologies relates to the different geographical locations of sample acquisition and analysis. The observed difference in mean sACE activity between two studies does not seem to be due to differences in the assays employed.

Genotyping

UCP3-55C>T, UCP2-866G>A and UCP2D/I genotypes were determined by PCR amplification using published primers and conditions [41,45,48]. Products were resolved on a 7.5% polyacrylamide gel and stained with ethidium bromide prior to analysis. The samples were resolved on a 1.5% agarose gel against a 1 kb DNA ladder. ACE (Hs01106052_m1) and GAPDH (Hs99999905_m1) gene expression levels were measured by qPCR. The qPCR efficiencies were 99% and 99% for ACE and GAPDH, respectively. Three housekeeping genes were used: UBC (Hs00226760), β-actin (Hs99999903_m1) and GAPDH (Hs99999905_m1). We did not analyse for changes in expression of an unrelated gene to check for unintentional global translational repression by siRNA treatment. However, we did confirm that the expression of the three housekeeping genes, UBC, β-actin and GAPDH, did not show a significant difference upon siRNA treatment of UCP2, by comparing each of the housekeeping genes with each other; that is, comparing the UCP2 siRNA versus no treatment cells, there was no significant difference in GAPDH mRNA levels using UBC and GAPDH as the housekeeping control, and similarly with other pairwise comparisons. Each reaction was repeated three times in parallel and for each condition. Crossing thresholds (Ct) were obtained using relative quantification application on the SDS2-1 software (Applied Biosystems, Life Technologies Ltd, Paisley, UK). Any value that was not consistent with the rest of the triplicate was suppressed before analysis. As a positive control using GAPDH siRNA, GAPDH mRNA was down-regulated by a factor of 5.48 (P=0.001). The Ct values included in statistical analyses are shown in Table 2.

RNA extraction, reverse transcription and RT-PCR

Cellular RNA was isolated using the RNeasy® Mini Kit (Qiagen, Crawley, UK) spin protocol. The RNA was stored at –80°C and quantified using a nanodrop spectrophotometer (ND-8000, Labtech, East Sussex, UK) prior to reverse transcription. Resultant complementary DNA samples were stored at –20°C.

Human ACE, UCP2, UCP3 and GAPDH (housekeeping gene) target sequences were amplified using forward and reverse primers (Table 2) by PCR to confirm co-expression. Amplification products were separated by electrophoresis on a 2% agarose gel against a 1 kb DNA ladder.

RNA was extracted at 24, 30 and 48 h after HUVECs were transfected with two different siRNAs against UCP2 (siRNA1 and siRNA2). Quantitative RT-PCR was performed on ABI prism 7900HT. Assays Hs00174179_m1, Hs00163349_m1 and Hs0106032_m1 were used in combination to TaqMan gene expression master mix to amplify ACE, UCP2 and UCP3 mRNA, respectively. Three housekeeping genes were used: UBC (Hs0082473_m1), β-actin (Hs99999903_m1) and GAPDH (Hs99999905_m1). We did not analyse for changes in expression of an unrelated gene to check for unintentional global translational repression by siRNA treatment. However, we did confirm that the expression of the three housekeeping genes, UBC, β-actin and GAPDH, did not show a significant difference upon siRNA treatment of UCP2, by comparing each of the housekeeping genes with each other; that is, comparing the UCP2 siRNA versus no treatment cells, there was no significant difference in GAPDH mRNA levels using UBC as the housekeeping control, and similarly with other pairwise comparisons. Each reaction was repeated three times in parallel and for each condition. Crossing thresholds (Ct) were obtained using relative quantification application on the SDS2-1 software (Applied Biosystems, Life Technologies Ltd, Paisley, UK). Any value that was not consistent with the rest of the triplicate was suppressed before analysis. As a positive control using GAPDH siRNA, GAPDH mRNA was down-regulated by a factor of 5.48 (P=0.001). The Ct values included in statistical analyses are shown in Table 2.

Table 2. Forward and reverse primers used during non-quantitative RT-PCR of human ACE, UCP3 and UCP2 and the housekeeping gene GAPDH and the resulting amplicon size

| Gene | cDNA accession no. | Primers | Amplicon size, bp |
|------|--------------------|---------|------------------|
| hGAPDH | J04038 | F: GCCGAAGGTTAACGGCTGAG | 228 |
| hUCP2 | U84763/AF050113 | R: CCTGGAAGATGGTGATGGGAT | 235 |
| hUCP2 | NM_003355/AF019409 | F: GCCGTGGAAGGATGGTGATGGG | 296 |
| hACE | J04144 | R: ACCAATGACCGAGAAG | 207 |
analysis were the means of triplicates. Experiments were repeated between three and five times.

Statistical analysis

Allele frequencies were estimated by gene counting. A \chi^2 test was used to compare the observed numbers of each genotype with those expected in a population in Hardy–Weinberg equilibrium. LD between sites in pairwise combination was estimated \cite{96}. One-way analysis of covariance (ANCOVA) tested for confounders. Univariate and multivariate analyses were used to measure significance of association. Differences in descriptive population variables and sACE were compared between genotype groups and allele groups using two-way ANOVA with log-transformed values as appropriate. Genotype association with sACE activity in the HMAR study was compared between genotype groups (including those defined by the presence/absence of a specific allele) using one-way ANOVA, two-tailed unpaired \( t \)-tests, linear trend analysis and one-way ANCOVA with sex as a covariate, using either raw or standardized values of \( t \)-tests, linear trend analysis and one-way ANCOVA with sex as a covariate, using either raw or log-transformed values as appropriate. Genotype association with sACE activity in the HMAR study was compared between genotype groups and allele groups using two-way ANOVA with measured values on one factor (time). All data were analysed using SPSS (SPSS Inc., IBM Corporation, USA) and Microsoft® Office Excel® 2007 (Microsoft Corporation 2006). Data are presented as means ± standard deviation unless otherwise stated. \( P \) values of <0.05 were considered statistically significant.

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