Increased Flux through the Hexosamine Biosynthetic Pathway leads to the induction of Acetyl-CoA Carboxylase Gene Expression in the Heart

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Thesis presented in partial fulfilment of the requirements for the degree of Masters of Physiological Sciences at the University of Stellenbosch.

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Declaration

I, the undersigned, hereby declare that the work contained in this thesis is my own original work and has not previously in its entirety or in part been submitted at any university for a degree.

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Signature

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Date
Abstract

Gene expression of the cardiac isoform of acetyl-CoA carboxylase (ACCβ) is induced in a glucose-dependent manner. ACCβ produces malonyl-CoA, a potent inhibitor of mitochondrial fatty acid uptake. Previous studies show that increased flux through the hexosamine biosynthetic pathway (HBP) under hyperglycaemic conditions may contribute to the development of insulin resistance. In light of this, we hypothesised that increased HBP flux induces cardiac ACCβ gene expression thereby contributing to the onset of insulin resistance.

We tested our hypothesis by transiently transfecting cardiac-derived rat H9c2 myoblasts with a 1,317 bp human ACCβ promoter-luciferase construct (pPIIβ-1317) and an expression construct encoding the rate-limiting step of the HBP i.e. glutamine:fructose 6-phosphate amidotransferase (GFAT). Overexpression of GFAT increased ACCβ gene promoter activity by 75 ± 23% versus controls (n=6, p<0.001). When co-transfection experiments were repeated in the presence of varying concentrations of L-glutamine (0 mM, 4 mM, 8 mM), a substrate for the HBP, ACCβ promoter activity was dose-dependently increased. To further corroborate these findings, we employed two inhibitors of GFAT, i.e. 40 µM azaserine and 40 µM 6-diazo-5-oxo-L-norleucine were administered to transfected cells for a period of 24 hours. Here both azaserine and 6-diazo-5-oxonorleucine attenuated ACCβ gene promoter activity.

In agreement, co-transfections with two dominant negative GFAT constructs also diminished ACCβ gene promoter activity. We next inhibited two enzymes of the HBP acting downstream of GFAT, i.e. O-GlcNAc transferase and O-GlcNase using alloxan (0.1 mM, 1 mM and 2 mM) and streptozotocin (5 mM and 10 mM).
respectively, for a period of 24 hours. Addition of alloxan attenuated ACCβ gene promoter activity by 35.6 ± 1.9% (n=16, p<0.001) and streptozotocin increased activity by 32 ± 12% (n=12, p<0.001). We also investigated USF1 and USF2 as transcriptional regulatory candidates for HBP-induced ACCβ promoter regulation. Our data implicates USF2 as an important transcriptional regulator of HBP-induced ACCβ promoter regulation.

In summary, this study demonstrates that increased flux through the hexosamine biosynthetic pathway induces ACCβ gene promoter activity. We further propose that such an induction would reduce cardiac fatty acid oxidation, thereby leading to intracellular lipid accumulation due to a mismatch between sarcolemmal FA uptake and mitochondrial FA oxidation in the insulin resistant setting (i.e. hyperlipidaemia).
Opstomming

Geen uitdrukking van die kardiale isoform asetiel-KoA karboksilase (ACCβ) word in 'n glukose afhanklike wyse geënduseer. ACCβ produseer maloniel-KoA, 'n kragtige inhibeerder van mitochondriale vetsuuroopname. Vorige studies toon aan dat verhoogde fluks deur die heksosamien biosintestiese weg (HBW) onder hiperglukemiese toestande bydra tot die ontwikkeling van insulienweerstand. In die lig hiervan, word daar gehipotetiseer dat verhoogde HBP fluks kardiale ACCβ geenuitdrukking induseer en so bydra tot die ontstaan van insulienweerstand.

Ons hipotese is getoets deur die kardiale afkomstige rot H9c2 mioblaste met 'n 1.317 bp mens ACCβ-lusiferase promotor konstruk (pPII-1317) te transfekteer en 'n uitdrukking te konstrueer wat die tempo bepalende stap van HBP i.e. glutamien: fruktose-6-fosfaat amidotransferase (GFAT) kodeer. Ooruitdrukking van GFAT verhoog ACCβ geenpromotor aktiviteit deur 75 ± 23% teenoor kontrole (n=6, p<0.001). Die herhaling van ko-transfeksie eksperimente is herhaal in die teenwoordigheid van variëerbare L-glutamienkonsentrasies (0 mM, 4 mM, 8 mM), 'n substraat vir die HBP, ACCβ promotor aktiviteit is dosisafhanglik verhoog. Om die bevindinge verder te staaf, is twee inhibeerders van GFAT, i.e. 40 µM azaserien en 40 µM 6-diazo-5-oxo-L-norleusien aan transfeksie selle toegedien vir 'n tydperk van 24 uur. Beide azaserien en 6-diazo-5-oxo-L-norleusien verlaag ACCβ geenpromotor aktiwiteit.

In ooreenstemming met die bogenoemde het ko-transfeksies met twee dominante negatiewe GFAT konstrukte ook ACCβ geenpromoter aktiwiteit verminder. Die volgende stap is om twee ensieme van die HBP wat stroomaf van GFAT aktief is, vir
'n periode van 24 uur te inhibeer i.e. O-GlcNAc transferase en O-GlcNAcase deur alloxan (0.1 mM, 1 mM en 2 mM) and streptozotosien (5 mM en 10 mM) onderskeidelik vir 'n 24 uur periode te gebruik. Toevoeging van alloxan het die ACCβ geenpromotor aktiwiteit by 35.6 ± 1.9% (n=16, p<0.001) verlaag en streptozotosien aktiwiteit verhoog by 32 ± 12% (n=12, p<0.001). Ons het ook die USF1 en USF2 as transkripsie regulerings kandidate vir HBP-geïnduseerde ACCβ promotor regulering ondersoek. Ons data impliseer dat USF2 as 'n belangrike transkripsie reguleerder van HBP-geïndiseerde ACCβ promotor regulering is.

Samevattend het hierdie studie demonstreer dat verhoogde fluks deur die hexosamien biosintetiese weg ACCβ geenpromotor aktiwiteit induseer. Ons stel verder voor dat hierdie induksie die kardiale vetsuuroksidasie verlaag wat daartoe lei dat intrasellulêre lipied akkumulasie as gevolg van onparing tussen sarkolemma vetsuuroopname en mitochondriale vetsuuroksidasie in 'n insulin weerstandige situasie (i.e. hiperlipidaemia).
I dedicate this thesis to my twin sister, Lucille and to
my two nephews, Jesse and Michael.
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<tr>
<td>ACBP</td>
<td>Acyl-CoA binding protein</td>
</tr>
<tr>
<td>ACC</td>
<td>Acetyl-CoA carboxylase</td>
</tr>
<tr>
<td>acetyl-CoA</td>
<td>Acetyl-Coenzyme A</td>
</tr>
<tr>
<td>Acyl-CoA</td>
<td>Acyl-Coenzyme A</td>
</tr>
<tr>
<td>AMP</td>
<td>adenosine monophosphate</td>
</tr>
<tr>
<td>AMPK</td>
<td>AMP-activated protein kinase</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>BSA</td>
<td>Bovine serum albumin</td>
</tr>
<tr>
<td>CACT</td>
<td>Carnitine/acylcarnitine transferase</td>
</tr>
<tr>
<td>ChREBP</td>
<td>Carbohydrate response element-binding protein</td>
</tr>
<tr>
<td>CPT</td>
<td>Carnitine palmitoyl transferase</td>
</tr>
<tr>
<td>CTD</td>
<td>Carboxyl-terminal domain</td>
</tr>
<tr>
<td>dGFAT</td>
<td>Dominant negative GFAT</td>
</tr>
<tr>
<td>dH₂O</td>
<td>Distilled water</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s modified Eagle’s medium</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
</tr>
<tr>
<td>DNL</td>
<td><em>De novo</em> lipogenesis</td>
</tr>
<tr>
<td>DON</td>
<td>6-Diazo-5-oxo-L-norleucine</td>
</tr>
<tr>
<td>ECL</td>
<td>Enhanced chemiluminescence</td>
</tr>
<tr>
<td>EDTA</td>
<td>Ethylene diamine tetraacetic acid</td>
</tr>
<tr>
<td>eLF</td>
<td>Eukaryotic factor</td>
</tr>
<tr>
<td>ER</td>
<td>Estrogen receptor</td>
</tr>
<tr>
<td>F-6-P</td>
<td>Fructose-6-phosphate</td>
</tr>
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</table>
FABP  Fatty acid binding protein
FADH$_2$  Flavin adenine dinucleotide
FAS  Fatty acid synthase
FAT  Fatty acid transporter
FATP  Fatty acid transport protein
FCS  Fetal calf serum
FFA  Free fatty acids
GFAT  Glutamine: fructose 6-phosphate amidotransferase
Glc-6-P  Glucose-6-phosphate
GlcN-6-P  Glucosamine-6-phosphate
GlcNAc  N-acetylglucosamine
GLUT  Glucose transporter
GP  Glycogen phosphorylase
GS  Glycogen synthase
H$_2$O  Water
HBP  Hexosamine biosynthetic pathway
HCl  Hydrogen chloride
H-FABP  Heart-type fatty acid binding protein
HK  Hexokinase
HRP  Horse-radish peroxidase
IRS-1  Insulin receptor 1
kDa  kilodaltons
LCFAs  Long-chain fatty acids
LAR  Luciferase assay reagent
malonyl-CoA  Malonyl-Coenzyme A
MCD  Malonyl-Coenzyme A decarboxylase
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
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<tbody>
<tr>
<td>mg</td>
<td>Milligrams</td>
</tr>
<tr>
<td>ml</td>
<td>Millilitre</td>
</tr>
<tr>
<td>mM</td>
<td>Millimolar</td>
</tr>
<tr>
<td>mRNA</td>
<td>Messenger ribonucleic acid</td>
</tr>
<tr>
<td>n</td>
<td>Numbers</td>
</tr>
<tr>
<td>NADH</td>
<td>Nicotinamide adenine dinucleotide</td>
</tr>
<tr>
<td>nm</td>
<td>Nanometer</td>
</tr>
<tr>
<td>O-GlcNAc</td>
<td>O-linked β-N-acetylglucosamine</td>
</tr>
<tr>
<td>O-GlcNAcase</td>
<td>β-N-acetylglucosaminidase</td>
</tr>
<tr>
<td>OGT</td>
<td>O-linked β-N-acetylglucosaminyl transferase</td>
</tr>
<tr>
<td>PDH</td>
<td>Pyruvate dehydrogenase</td>
</tr>
<tr>
<td>PFK</td>
<td>Phosphofructokinase</td>
</tr>
<tr>
<td>PI-3 kinase</td>
<td>Phosphoinositide 3-kinase</td>
</tr>
<tr>
<td>PMSF</td>
<td>Phenylmethylsulfonyl flouoride</td>
</tr>
<tr>
<td>PPARs</td>
<td>Peroxisome proliferator-activated receptors</td>
</tr>
<tr>
<td>PPP</td>
<td>Pentose phosphate pathway</td>
</tr>
<tr>
<td>Pro</td>
<td>Proline</td>
</tr>
<tr>
<td>PUGNAc</td>
<td>O-(2-acetamido-2-deoxy-D-glucopyranosylidene) amino-N-phenylcarbamate</td>
</tr>
<tr>
<td>PVDF</td>
<td>Polyvinylidene difluoride</td>
</tr>
<tr>
<td>RIPA</td>
<td>Radio immuno precipitation assay</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
</tr>
<tr>
<td>RNA Pol</td>
<td>Ribonucleic acid polymerase</td>
</tr>
<tr>
<td>rpm</td>
<td>Revolutions per minute</td>
</tr>
<tr>
<td>RT</td>
<td>Room temperature</td>
</tr>
<tr>
<td>RT-PCR</td>
<td>Real-time polymerase chain reaction</td>
</tr>
<tr>
<td>Acronym</td>
<td>Full Form</td>
</tr>
<tr>
<td>---------</td>
<td>-----------</td>
</tr>
<tr>
<td>SBT</td>
<td>Strontium bismuth tantalate</td>
</tr>
<tr>
<td>SDS</td>
<td>Sodium dodecyl sulphate</td>
</tr>
<tr>
<td>Ser</td>
<td>Serine</td>
</tr>
<tr>
<td>STZ</td>
<td>Streptozotocin</td>
</tr>
<tr>
<td>TBS</td>
<td>Tris-buffered saline</td>
</tr>
<tr>
<td>Thr</td>
<td>Threonine</td>
</tr>
<tr>
<td>Tyr</td>
<td>Tyrosine</td>
</tr>
<tr>
<td>UDP</td>
<td>Uridine diphospho</td>
</tr>
<tr>
<td>UDP-GlcNAc</td>
<td>Uridine diphospho-(N)-acetylglucosamine</td>
</tr>
<tr>
<td>µl</td>
<td>Microlitre</td>
</tr>
<tr>
<td>USF</td>
<td>Upstream stimulatory factor</td>
</tr>
<tr>
<td>VLDLs</td>
<td>Very-low density lipoproteins</td>
</tr>
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Chapter 1

Introduction
1.1 Epidemiology

Obesity is defined as an excess of body fat which accumulates in adipose tissue, associated with increased fat cell size and number (68). Obesity has become more prevalent worldwide, resulting in an increase in related diseases such as type-2 diabetes. Although obesity is especially common in industrialised countries, its prevalence is also dramatically increasing in developing countries such as South Africa (4). The movement of populations from a rural type to a more “western-based” lifestyle with its increased availability of high-calorie foods is a key factor that has lead to a higher incidence of obesity. It is predicted that ~300 million people worldwide will be diagnosed with type-2 diabetes by 2025 (122, 126).

Most patients who are obese develop insulin resistance which is characterised as an impairment of insulin to mediate glucose uptake and metabolism by muscle and adipose tissue. Cardiovascular disease is the primary cause of mortality in obese individuals and in patients diagnosed with type-2 diabetes mellitus (60, 68, 112, 127). Although the heart plays a small role in the development of insulin resistance throughout the whole body, cardiovascular complications are the main causes of death in insulin resistant obese and type-2 diabetes patients (23). Diabetes mellitus is a cluster of metabolic perturbations characterized by high blood glucose levels or hyperglycaemia, which result from defects in insulin secretion, or action, or both. Diabetes mellitus, can easily be identified with high glucose levels found in urine and excessive muscle loss (121).
South Africa has a unique, heterogeneous population originating from a diverse range of ethnic backgrounds. The ~ 46 million individuals that reside in South Africa largely consist of African (79%), Caucasoid (9.6%), mixed ancestry (“coloured”) (8.9%) and Indian backgrounds (2.5%). More recently, migrants from other African countries have also settled in South Africa. This diverse cultural and ethnic diversity makes it difficult to elucidate trends with respect to changes in diet and health behaviours of the entire population. There are also limited studies performed to assess the prevalence of metabolic syndrome (a precursor to obesity and diabetes) in South Africa. However, recent data show that mortality rates from ischaemic heart disease among whites, coloureds and Indians were found to be more than 2x the rate for blacks, while stroke death rates among blacks and coloureds were double compared to whites (Table 1).

There are many sociological implications why the prevalence of cardiovascular diseases has become more common. For example, for the black community, the highest incidences of obesity are observed in black women (4). This may, in part, depend on the cultural background in the black population where obesity is often regarded as a reflection of health and wealth (4).
Table 1: Age-standardised mortality rates per 100,000 population, South Africa (112).

<table>
<thead>
<tr>
<th></th>
<th>Ischaemic heart disease</th>
<th>Stroke</th>
<th>Hypertensive heart disease</th>
<th>Diabetes</th>
</tr>
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<tbody>
<tr>
<td>Black</td>
<td>70</td>
<td>143</td>
<td>88</td>
<td>56</td>
</tr>
<tr>
<td>White</td>
<td>230</td>
<td>72</td>
<td>10</td>
<td>22</td>
</tr>
<tr>
<td>Coloured</td>
<td>171</td>
<td>139</td>
<td>37</td>
<td>59</td>
</tr>
<tr>
<td>Indian</td>
<td>392</td>
<td>392</td>
<td>29</td>
<td>103</td>
</tr>
<tr>
<td>South Africa</td>
<td>123</td>
<td>124</td>
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Other associated lifestyle risk factors may also influence the epidemiology of cardiovascular diseases. Risk factors such as physical inactivity, increased smoking, hypertension and hypercholesterolemia may play an important role in the increase in cardiovascular disease and type-2 diabetes (112, 131). For example, although measures were adopted to reduce smoking (higher retail prices and a smoking ban) a recent South African report found that the prevalence of young smokers (14 years) increased by 30% (112). The survey was performed from 1998 to 2003 among a population of 15,124 school children in South Africa. Furthermore, the prevalence of sedentary behaviour has increased in recent years (112). Together these data highlight the increased burden of diabetes and cardiovascular diseases faced by developing nations such as South Africa.
In light of this our laboratory has begun to investigate the basic mechanisms underlying the development of diabetes and cardiovascular diseases. In particular, we are focusing on the role of altered metabolism in the pathogenesis of type-2 diabetes and heart diseases. For the next part of this Introduction, I will now review some basics aspects of the heart’s metabolism and thereafter focus on my particular interest, i.e. the hexosamine biosynthetic pathway.
1.2 Metabolism of the heart

The heart pumps blood for the complete lifespan of an individual. This constant workload demands a high capacity for energy production in the form of adenosine triphosphate (ATP), produced by the mitochondria of the heart. In terms of its fuel substrate preferences, the heart is like an omnivore, utilising fatty acids, glucose, lactate and ketone bodies and is able to switch between either one depending on substrate availability (23). The normal adult mammalian heart prefers fatty acids as a fuel source, i.e. obtaining ~70% of its energy from fatty acid oxidation with the remainder provided by glucose and lactate (23). In the foetal heart carbohydrates are major substrates.

The heart utilises ketone bodies as a fuel source during fasting conditions. Metabolism of ketones generates NADH2 and FADH2 which can be used to generate energy by the electron transport chain (36). It has been shown that ketone bodies are able to suppress cardiac fatty acid oxidation in diabetes (46).

Studies have shown that in conditions of increased exercise, hypoxia, and anoxia excess pyruvate from glycolysis is converted into lactate (37). Brooks et al. (2000) introduced the hypothesis of a “lactate shuttle” (9). This hypothesis proposes that lactate formation and its distribution throughout the body can act as a mechanism to coordinate metabolism in different tissues. It is known that during intense exercise lactate flux can exceed glucose flux (9, 37).
1.2.1. Fatty acid metabolism

Fatty acids have four major physiological roles in metabolism. Firstly, it forms building blocks in the formation of phospholipids and glycolipids; secondly its involved in the modification of proteins that are targeted to the cell membrane; thirdly it can serve as hormones and intracellular messengers; and finally fatty acids are utilised as fuel substances (113). Fatty acids are stored as triacylglycerol in adipose tissue until it is needed, i.e. then to be broken down by lipolysis (113).

Long-chain fatty acids (LCFAs) enter the circulation in two forms, either in a complex with albumin or esterified in a lipid core of very-low density lipoproteins (VLDLs) and chylomicrons (23, 109, 118). Free fatty acids (FFA) are released into the bloodstream by adipose tissue and taken up by non-adipose tissue via sarcolemmal transporters. It was first believed that LCFAs were transported across the sarcolemma into cardiomyocytes by passive diffusion, but it is now accepted that most of LCFAs are taken up by membrane transporters (40, 72, 73, 92). Two such fatty acid transporters are fatty acid translocase, a rat homologue of human CD36 (FAT/CD36) and fatty acid binding protein (FABP) (8, 82). There are also two isoforms of the fatty acid transport protein (FATP) family, i.e. FATP1 and FATP6 present in cardiomyocytes (110). Both exhibit acyl-CoA synthetase activity. FATP6 is found exclusively and in higher abundance in the heart. FATP has been found to colocalise with FAT/CD36 and both these LCFA transport proteins act in concert with each other.
Once inside the cardiac myocyte, LCFAs bind to a cytoplasmic heart-type fatty acid binding protein (H-FABP). This protein transports non-esterified LCFAs towards a site where they are converted and activated by acyl-Coenzyme A (acyl-CoA) synthetase to form fatty acyl-CoA. Acyl-CoA binding protein (ACBP) then binds to these acyl-CoAs and can either incorporate acyl-CoA into intracellular lipid pools or shuttle it to the mitochondria to be metabolised.

Fatty acyl-CoA is transported into the mitochondria by the action of three proteins which function as a complex. At the outer membrane is carnitine palmitoyl transferase (CPT1) which catalyses the formation of acylcarnitine (23, 29). Connected to CPT1 is carnitine/acylcarnitine transferase (CACT) which transports acylcarnitine into the mitochondria. The final enzyme (CPT-II) is found on the inner mitochondrial membrane and releases acyl-CoA into the mitochondrial matrix (23). The process of mitochondrial LCFA uptake is regulated by CPT1 which is the rate-limiting enzyme for this process (Figure 1).

After uptake is complete the acyl-CoAs are oxidised by β-oxidation producing acetyl-CoA as a by-product. Acetyl-CoA from β-oxidation enters the citric acid cycle to be degraded along with acetyl-CoA from glucose oxidation (23, 34, 103). The result is the generation of FADH$_2$ and NADH that enter the mitochondrial respiratory chain. In oxidative phosphorylation, ATP synthesis is coupled to the flow of electrons from NADH or FADH$_2$ to oxygen by a proton gradient across the inner mitochondrial membrane.
Free fatty acids  
\[ \text{FAT/CD36} \]  
\[ \text{Sarcolemma} \]  
\[ \text{LCFA} \]  
\[ \text{Acyl- CoA synthetase} \]  
\[ \text{CPT1} \]  
\[ \text{Mitochondrion} \]  
\[ \text{Acyl-CoA} \]  
\[ \text{Acetyl-CoA} \]  
\[ \text{Malonyl-CoA} \]  
\[ \text{Fatty acid β-oxidation spiral} \]  
\[ \text{Citric acid cycle} \]  
\[ \text{FADH}_2 \text{ and NADH} \]  

**Figure 1:** Diagram of fatty acid metabolism.  
A proton gradient is created by pumping protons out of the mitochondrial matrix into the inter-mitochondrial membrane space. Thus a proton gradient is formed which creates a membrane potential. The protons then flow back into the mitochondrial matrix through ATP synthase which drives ATP production (113).

Even though LCFAs are utilised for energy production they also play a role in the regulation of genes involved in their own metabolic pathway. LCFAs induce genes that increase fatty acid oxidation by activating a family of ligand-activated nuclear receptors called the peroxisome proliferator-activated receptors (PPARs) (114, 117). There are three isoforms of PPARs, i.e. PPARα, β/δ and γ (114, 117). PPARα and β/δ are the main isoforms expressed in cardiomyocytes and activation of these genes results in an increased expression of regulators of fatty acid oxidation and fatty acid uptake, i.e. FAT/CD36 and CPT1 (35).

Fatty acid oxidation is regulated depending on fatty acid availability, its uptake by mitochondria and by its breakdown. Mitochondrial uptake of long-chain fatty acyl units is controlled by CPT1. A key molecule responsible for the regulation of CPT1 is malonyl-CoA (66, 68, 89, 100, 111, 130). Malonyl-CoA is a potent inhibitor of CPT1 and is produced from acetyl-CoA by an enzyme known as acetyl-CoA carboxylase (ACC) (66, 68, 89, 100, 111, 130). Another enzyme, malonyl-CoA decarboxylase (MCD) degrades malonyl-CoA into acetyl-CoA (28, 111). ACC has two isoforms, ACCα and ACCβ, that have different physiological roles based on their distinct subcellular distributions (44).
ACCα is a cytosolic enzyme (molecular mass of 265 kDa) that supplies malonyl-CoA to fatty acid synthase (FAS) and is committed to de novo lipogenesis (DNL) in many tissues via subsequent nutritional and hormonal regulation (3, 39, 44, 61, 97). In contrast, ACCβ (molecular mass of 280 kDa) is anchored to the mitochondrial surface via a unique N-terminal domain that includes 20 hydrophobic amino acids (1, 2, 39, 44). ACCβ is responsible for malonyl-CoA production in the heart. ACCβ overexpression increases malonyl-CoA production resulting in a decrease in fatty acid uptake (28, 66, 89, 100, 111, 130). AMP-activated protein kinase (AMPK) plays an important role in the regulation of CPT1 by phosphorylating and inhibiting ACCβ, resulting in an increase in fatty acid oxidation. Phosphorylation of ACCβ by AMPK is well documented (27, 54). It has been suggested that MCD is also phosphorylated by AMPK (27, 101). Therefore, AMPK plays a distinct and important role in regulating both malonyl-CoA levels and fatty acid oxidation in the heart.

ACCβ is expressed abundantly in heart, skeletal muscle, and liver (1, 61, 87, 133). ACCβ transcripts contain two species of 5′-UTRs, which contain either the sequence of exon 1a or of exon 1b via the alternative usage of two promoters, i.e. promoter 1 and promoter 2 (P1 and P2) (87). Exon 1a and exon 1b are located ~15 kilobases apart in human genome but are both connected to exon 2 in mRNA after splicing (87). However, they both use the same ATG start codon for translation, which is found in exon 2 and therefore the two transcripts encode for the same protein (39, 67, 87). A differential regulation of ACCβ gene expression originates from alternative usage of promoters, such as P1 and P2 in different tissues. P1 is the sole promoter.
found in the heart and skeletal muscle of rats, although both P1 and P2 are active in human skeletal muscle (87).

In human skeletal muscle and P2 is regulated by myogenic regulatory factors (MRFs) (67, 87). MRFs, including Myf5, MyoD, myogenin, and MRF4, are basic helix-loop-helix transcription factors involved in myogenic differentiation. These factors all recognize the same consensus sequence, i.e. E-box (CANNTG) (87). Myogenin and MRF4 play a major role in the expression of muscle genes in fully differentiated myotubes, while Myf5 and MyoD have been shown to establish the myogenic lineage during embryogenesis (87, 98, 99, 102, 108). It is important to note that myogenic regulatory factor-binding sites found in the human ACCβ P2 are not conserved in rat P2. This would contribute to this difference in P2 usage between human and rat skeletal muscle (87).

The level of ACCβ is higher in the heart than in skeletal muscle. It is currently not known which promoter controls ACCβ expression in the heart. Cardiomyocyte-specific transcription factors, such as Csx/Nkx2.5, GATA4, MEF2, and eHand have been implicated in cardiac development and cardiac gene expression. Unlike in skeletal muscle MRFs have not been shown to be involved in this regulation (58, 63, 84, 87, 88).

The nucleotide sequence of the cDNA of the human liver ACCβ carboxylase has an open reading frame of 7,449 nucleotides that encode 2,483 amino acids. The nucleotide sequences and the predicted amino acid sequences
from the cDNA of ACCβ, has ~60 and 80% in similarity to that of ACCα, respectively. Ser77 and Ser79 have been found to be critical for the phosphorylation and of rat ACCα (Ser78 and Ser80 of human ACCα) (1, 87). These amino acids are conserved in ACCβ and are represented as Ser219 and Ser221, respectively. Another phosphorylation site, Ser1200, in rat ACCα (Ser1201 of human ACCα) has been found to be absent in ACCβ.

Most of the homology between the amino acid sequences of the human ACC isoforms is found downstream of residues Ser78 and Ser81 in human ACCα and their equivalent residues in ACCβ, i.e. Ser219 and Ser221 (1). It has been suggested that the first 218 amino acids at the N terminus of ACCβ represents a unique peptide that could be responsible for the variation between the two carboxylases (1). Despite the similarities between these two isoforms, studies with rat liver ACCα and ACCβ showed that the two isoforms do not cross-react immunochemically (1, 128). It was shown that when the amino acid sequences of the human ACCα and ACCβ are aligned, an extra 142 amino acids can be found in ACCβ (i.e. 426 bp in ACCβ cDNA) (1, 87). It is believed that the extra 142 amino acids are involved in controlling the localisation of ACCβ in the cell, i.e. to the mitochondrion.
1.2.2. Glucose metabolism

The uptake of extracellular glucose by myocytes is mediated by glucose transporters (GLUTs). There are two glucose transporters that can be found in the heart, i.e. GLUT1 and GLUT4 that are located not only in the sarcolemma but also in intracellular storage compartments (6, 23, 48-50, 62, 77, 78, 80, 105). GLUT1 is the foetal isoform and can be found in less abundance than the adult, insulin-stimulated glucose transporter (GLUT4). After glucose has entered the cardiomyocytes it is rapidly phosphorylated by hexokinase into glucose-6-phosphate.

Once converted to glucose-6-phosphate, glucose can be metabolised in six different ways (Figure 2). Firstly, glycogen synthesis can take place and glucose-6-phosphate can be converted by glycogen synthase (GS) into glycogen for storage. This process is reversible and when required glycogen phosphorylase (GP) can convert glycogen back into glucose-6-phosphate (113). Some of the glucose-6-phosphate can enter the pentose phosphate pathway (PPP) where it has been proposed that xylulose-5-phosphate activates a specific isoforms of protein phosphatase 2A which, in turn, dephosphorylates the transcription factor carbohydrate response element-binding protein (ChREBP) (53, 129). In the liver ChREBP translocates from the cytosol to the nucleus where it regulates the expression of glycolytic and lipogenic enzymes (57).
Most of the glucose-6-phosphate enters glycolysis. After the first step of glycolysis, phosphoglucone isomerase converts glucose-6-phosphate into fructose-6-phosphate (Figure 2). At this point most of the fructose-6-phosphate continues down the glycolytic pathway where it is converted to pyruvate after a long series of steps. However, a small percentage of the fructose-6-phosphate is diverted to another pathway responsible for nutrient sensing, i.e. the hexosamine biosynthetic pathway (77, 95) (to be discussed in more detail later).

**Figure 2: Diagram representing the different pathways of glucose metabolism (reproduced from (53)).**
In the glycolytic pathway, fructose-6-phosphate is converted into fructose-1,6-bisphosphate by phosphofructokinase which is the rate-limiting enzyme of glycolysis (113). The production of pyruvate marks the end of the glycolytic pathway. In the absence of oxygen pyruvate can be reversibly converted to lactate. Under aerobic conditions pyruvate is transported into the mitochondria by pyruvate dehydrogenase (PDH), the rate limiting enzyme of glucose oxidation, where it undergoes oxidative decarboxylation into acetyl-CoA (113). Acetyl-CoA from glucose metabolism, together with acetyl-CoA from fatty acid oxidation, enters the citric acid cycle (Krebs cycle) where it is oxidised to carbon dioxide, NADH and FADH$_2$. The NADH and FADH$_2$ produced are then used in oxidative phosphorylation to produce ATP after donating their electrons to oxygen (113).

Another pathway that utilizes glucose is the polyol pathway. The polyol pathway consists of two steps in which glucose is converted to sorbitol and then converted into fructose (70). During this process NADPH is converted to NADP$^+$ (70). The polyol pathway mainly functions to remove excess glucose from glycolysis and then return it to the glycolytic pathway again (70).

The glyoxylate pathway is found mainly in plants and yeast (69). This pathway converts acetyl-CoA into oxaloacetate by bypassing the steps in the citric acid cycle. It can therefore use fats for the synthesis of carbohydrates (69).

The last pathway that utilises carbohydrates is the biosynthesis of oligosaccharides and glycoproteins which are then expressed on the surface of cell membranes (20).
1.2.3 The Randle cycle

The Randle cycle (named after Philip Randle, its first proposer), which has been used to explain the reciprocal relationship between fatty acid oxidation and glucose oxidation, has long been implicated as a potential mechanism for hyperglycaemia and type-2 diabetes mellitus (106). The Randle cycle states that increased fatty acid oxidation causes a decrease in glucose oxidation. Thus in the setting of excess FFA and glucose supply (insulin resistant state), this is thought to lead to lower glucose uptake and eventually hyperglycaemia (106). Here, acetyl-CoA and NADH derived from fatty acid oxidation can suppress pyruvate oxidation by inhibiting pyruvate dehydrogenase (33, 85). Increased fatty acid oxidation has also shown to result in the inhibition of phosphofructokinase and prevent glycolysis. This would increase glucose flux through other glucose pathways and result in glucose accumulation (85).
1.2.4. Hexosamine biosynthetic pathway

Since the focus of my thesis is on the hexosamine biosynthetic pathway, I will now discuss this in more detail. The hexosamine biosynthetic pathway (HBP) is a relatively small branch glucose utilising pathway. Only ~3% of the total glucose utilised in the cell enters the HBP (77, 95). The pathway is catalysed by the rate-limiting enzyme glutamine: fructose 6-phosphate amidotransferase (GFAT). During this first step, fructose-6-phosphate and glutamine is converted to glucosamine-6-phosphate and glutamate (Figure 3). Thereafter, through a series of steps glucosamine-6-phosphate is converted to glucosamine-1-phosphate. After the addition of uridine, it is converted into uridine diphospho-\(N\)-acetylglucosamine (UDP-GlcNAc) and CMP-sialic acid. UDP-GlcNAc is the end product of the HBP pathway and also functions as an inhibitor of GFAT (12).

![Figure 3: Description of the hexosamine biosynthetic pathway.](image)

UDP-GlcNAc functions as the substrate for O-linked β-N-acetylglucosamine transferase (OGT). OGT catalyses the reversible modification of various proteins and transcription factors by cleaving UDP from GlcNAc and transferring GlcNAc in O-linkage to serine/threonine residues on proteins (12, 64, 71). O-GlcNAc modification has two novel mechanisms of action. Firstly, O-GlcNAc is not elongated into a more complex structure (115). Secondly, it has a nucleoplasmic distribution instead of being localised to the cell surface like other glycoproteins (45, 52). O-GlcNAc modification has been implicated in modulating different mechanisms that include (i) regulating protein phosphorylation and function; (ii) altering protein degradation; (iii) altering the localisation of proteins; (iv) modulating protein-protein interactions and (v) mediating transcription (132).

The sites of O-GlcNAc modification are often identical or adjacent to known phosphorylation sites, suggesting that “O-GlcNAcation” plays a role in regulation of a wide range of pathways (12, 125). It has been shown that O-GlcNAc regulation can modify proteins in competition with phosphorylation (21). In some instances O-GlcNAc and phosphorylation can exist on separate and distinct subsets of a protein. For example, c-Myc and RNA polymerase both contain threonine or tyrosine sites that can be phosphorylated or glycosylated, but although, there has been no proof that the two modifications can exist on one protein, there is a possibility that this dual-protein modification can occur. In particular, RNA polymerase II exists in two distinct forms, i.e. RNA Pol IIA and RNA POL IIO (21, 132). RNA polymerase II contains a highly conserved carboxyl-terminal domain (CTD) consisting of 52 tandem repeats of the consensus sequence Tyr-Ser-Pro-Thr-Ser-Pro-Ser (21, 24, 26, 132). The CTD of the IIO isoforms is found to be phosphorylated on
the serine and threonine residues. In contrast, the IIA isoform is non-phosphorylated and exhibits extensive O-GlcNAc modification. The existence of both O-GlcNAc and phosphorylation site implies a precise regulation of protein activity (21).

The modifications of proteins by OGT with O-GlcNAc are also closely regulated. Another enzyme responsible for the regulation of O-GlcNAc isation is β-N-acetylglucosaminidase (O-GlcNAcase). Although OGT is responsible for binding O-GlcNAc to serine/threonine residues of proteins, O-GlcNAcase functions to remove O-GlcNAc. O-GlcNAc modification is thus regulated in the same manner as phosphorylation (Figure 3).

O-GlcNAc modification targets numerous proteins, including transcription factors (21, 132). For example, Sp1, an important transcription factor in the regulation of several target genes, has been shown to have multiple O-GlcNAc residues (12, 21, 132). O-GlcNAc has been shown to alter protein degradation by two different mechanisms, i.e. (i) by altering the targeting of proteins to the proteasome or (ii) by altering the activity of the proteasome. O-GlcNAc modifies eukaryotic factor (eIF) 2α-p67, Sp1 and estrogen receptor (ER)-β prolonging the half-life of these proteins (132). Insulin has been reported to increase O-glycosylation and nuclear content of Sp1 (74).

Incubation with high glucose or increasing flux through HBP by overexpressing GFAT increased the expression of upstream stimulatory factor 1 and 2 (USF1 and 2), although these transcription factors are apparently not O-GlcNAc modified (12, 123).
The gene encoding OGT (O-linked β-N-acetilglucosamine transferase) is essential for embryonic and stem cell development in mammals (42), making it difficult to produce a transgenic knockout model to investigate HBP regulation. Hanover et al. (2005) examined the role of OGT using an ogt-1 deletion strain of *Caenorhabditis elegans* (42). This strain exhibited no obvious developmental phenotype that was found in homozygous animals and could be used successfully as a model for nutrient-driven insulin resistance. One of the main findings of this model was that homozygous (rat/mouse) lacking ogt-1 had increased levels of glucose and glycogen, accompanied by a decrease in fat stores (42). This would imply that the HBP was involved in the regulation glycogen synthesis and fatty acid oxidation.

Studies in adipocytes suggest that glucose-induced insulin resistance is caused by impaired translocation of insulin-responsive glucose transporters to the cell membrane (such as GLUT4) and that an increase in glucose flux through the HBP plays a major role in the development of insulin resistance (10-12, 79, 96). There are several observations to suggest that the HBP increases the development of insulin resistance (6, 13, 18, 47, 48, 55, 77, 81, 105, 120). Pre-exposure to glucosamine inhibits basal and insulin-stimulated glucose transport and decreases insulin-stimulated glycogen synthesis in rat muscles without affecting insulin receptor signaling (6, 13, 18, 47, 48, 55, 77, 81, 105, 120). Increasing HBP flux has been shown to alter glucose uptake due to increased O-GlcNAc modification of proteins involved in the regulation of the insulin-signaling cascade, i.e. IRS-1, PI-3 kinase and Akt (31, 90, 132).
A much lower concentration of glucosamine than glucose is required to elicit insulin resistance. The main difference between these two substrates is that glucose is utilised by several pathways where glucosamine is utilised only by HBP. Glutamine or a mixture of amino acids is also an important requirement for the development of glucose-induced insulin resistance of glucose transport in adipocytes (55). When transamidases are added to adipocytes treated with a mixture of amino acids the effect is reversed (55).

Several studies have shown that by increasing the concentration of extracellular glucose and glucosamine, or by increasing glucose uptake by overexpressing glucose transporters (GLUTs) results in insulin resistance (6, 48-50, 62, 77, 78, 80, 94, 105). For example, it was shown that by blocking GFAT with pharmacological agents inhibited glucose-mediated insulin resistance (77). Moreover, other studies found that GFAT overexpression mimicked the effect of treating cells or rats with elevated glucose/glucosamine (18, 22, 25, 80). Also by increasing O-GlcNAc levels in mice and in cell culture genetically or by pharmaceutical intervention, resulted in insulin resistance (5, 14, 42, 81, 90, 120). Together these studies therefore support a strong link between increased HBP flux and the development of insulin resistance/type-2 diabetes.
1.3. Hypothesis

Since the expression of ACCβ is elevated under hyperglycaemic conditions (1-3, 7, 32, 33, 39, 66, 75, 76, 100), we hypothesized that increased HBP flux induces cardiac ACCβ gene expression (Figure 4).

We further propose that since this would result in greater malonyl-CoA production, less fatty acids will be transported into the mitochondria to be oxidised. In the insulin-resistant setting (hyperlipidaemia), failure to oxidise fatty acids is predicted to result in an accumulation of intracellular long chain fatty acids. This, in turn, could result in deleterious effects, for e.g. cell death and contractile dysfunction (68) (Figure 4).
Figure 4: Description of hypothesis.
1.4. Aims

1. **Transfection** - we will transfect rat cardiac-derived H9c2 myoblasts with a human ACCβ promoter-luciferase construct and a GFAT overexpression construct. The myoblasts will be treated with various inhibitors to test whether the HBP regulates the ACCβ promoter.

2. **Western blotting ***(in vitro)*** – Protein extracted from transfected myoblasts will be analysed by Western blotting for the expression of GFAT and O-GlcNAc.

3. **Western blotting ***(in vivo)*** – Proteins will be extracted from cardiac tissue from a transgenic mouse model of obesity-induced type-2 diabetes (db/db mouse) and analysed for altered expression of GFAT and O-GlcNAc.
Chapter 2

Methods
2.1. Transfections

2.1.1 Background to principles of the technique

Transfection, i.e. experimental exogenous transfer of DNA into a target cell is a useful method to exploit in order to measure gene promoter activity (Figure 5). Two components are required for a successful transfection. Firstly, a transfection reagent is required that will bind to plasmid DNA to be transferred into the cytosol of cells. For example, for this study Fugene 6 transfection reagent (Roche, Penzberg, Germany) was employed. The second requirement is a plasmid DNA that will be transfected together with the promoter-luciferase construct. The former is constitutively expressed and used to normalise transfection results according to cell number and transfection efficiency.

The gene promoter of interest is bound to a firefly luciferase gene, allowing promoter activity to be measured by the amount of luciferase protein synthesized by the cell. The normalising construct employed for this thesis was pRL-CMV (Promega, Fitchburg, WI, USA). The luciferin protein expressed by the pRL-CMV construct is isolated from *Renilla reniformis*. After transfection, luciferin protein is extracted by cell lysis. Thereafter a substrate called luciferase assay reagent II (LAR II) is added to activate the luciferin protein, resulting in light emission. The latter can be measured using a luminometer. Since the luciferin protein produced by the normalising agent and the promoter construct are different, each can be measured separately from the same sample. Thus, a Dual-Luciferase Reporter Assay Kit
(Promega, Fitchburg, WI, USA) was used where two substrates were added to the same sample, i.e. Luciferase Assay Reagent II (LAR II) (measuring promoter activity) and “Stop and Glo” (neutralizes LAR II substrate and activates the *Renilla* luciferin).

Figure 5: Outline of principles governing gene promoter activity measurements using luciferase assay.

Transfections were performed as a 5-day experiment (Figure 6). On the first day cells were seeded on 12-well plates. On day 2 cells were transfected, while media of myoblasts was replaced on day 3. This ensured that myoblasts were supplied with sufficient nutrients and also to remove excess transfection reagent. At this stage inhibitors/drugs that were being tested were added (to be discussed later). On day 4, after 24 hours treatment, cells were lysed and the lysate stored at -80°C. The samples were rapidly thawed on day 5 to further enhance cell lysis. Samples were subsequently plated on a 96-well luminometer plate and promoter activity measured (Figure 6).
Day 1: Seed H9c2 myoblasts in 12-well culture plates (seeding density ~35,000 cells/well)

Day 2: Performed transfection

Day 3: Change media on cells ± experimental manipulation

Day 4: Extract cell lysates using Dual-Luciferase Reporter Assay Reagent Kit and store -80°C

Day 5: Perform a luciferase assay using Dual-Luciferase Assay Reporter Kit and luminometer

Figure 6: Outline of transfection schedule.
2.1.2. Cell culture

H9c2 rat cardiac-derived myoblasts were chosen for experiments because they are a precursor cell line to cardiomyocytes (19, 38, 91). Precursor cells are also easier to differentiate than terminally differentiated myotubes. H9c2 myoblasts were cultured in T75 culture flasks with Dulbecco’s modified Eagle’s medium (DMEM) (Highveld, South Africa) with 10% GibCo foetal calf serum (Invitrogen, Carlsbad, CA, USA) and 4 mM GibCo L-glutamine (Invitrogen, Carlsbad, CA, USA). Cells were not allowed to grow to a confluency greater than 80-90% and were cultured for a maximum of 8 passages before growing new cells. We used passages 9-15 for transfection experiments. In our initial optimizing experiments passages 9-15 were used and it was decided to continue using them in order to ensure consistency between results.

Myoblasts were grown as described and plated at 35,000 cells per well on 12-well culture plates (Greiner, Kremsmünster, Austria) in 1 ml of completed DMEM with 10% foetal calf serum and 4 mM L-glutamine. The cells were incubated for 24 hours at 5% CO₂, 20% O₂ and 95% humidity at 37°C prior to transfection.
2.1.3. Promoter-luciferase and DNA constructs used for transfection experiments

pGL3-Control (Promega, Madison, WI, USA) was used in all transfection experiments to normalise results according to cell number and transfection efficiency. pGL3-Control is a plasmid constitutively expressing luciferase from an SV40 promoter. pGL3-Basic is a plasmid lacking a promoter and therefore expresses only baseline levels of luciferase. The latter was used to normalise the total amount of DNA used per transfection to ensure comparable transfection efficiency between experiments. The total amount of DNA transfected for each experiment was 0.75 µg, and pGL3-Basic was used to make up the remaining DNA needed. H9c2 myoblasts were transiently transfected with a 1,317 bp human ACCβ promoter-luciferase reporter construct (pPIIβ-1,317) previously described (Makaula et al., 2006) (75). 0.25 µg of pPIIβ-1,317 was transfected ± 0.25 µg of a human pcDNA3-GFAT expression vector (123). Two dominant negative constructs, i.e. pcDNA3-GFAT577 and pcDNA3-GFAT667 were also employed in this study (123). Both dominant negative constructs were separately transfected with pPIIβ-1,317 and GFAT. There is a great amount of sequence homology between the rat and human isoforms of GFAT (91%) and ACCβ (90%) and therefore it is unlikely that this would represent a problem when expressing human constructs in a rat cardiac-derived cell line.
Description of plasmid constructs:

1. pcDNA3-GFAT (see Figure 7) contains a full-length human GFAT cDNA generated by RT-PCR and cloned into the expression vector pcDNA3.1 (see Figure 8) (Invitrogen, Inchinnan, Scotland). The PCR product was verified by sequencing and shows identity to human GFAT (also known as GFAT1 = glutamine:fructose-6-phosphate transaminase 1, GenBank accession number M90516). This construct was kindly donated to us by Dr. Cora Weigert (University of Tübingen, Germany).

![Diagram of pcDNA3-GFAT vector](attachment:image.png)

**Figure 7: Sketch of human GFAT gene cloned into pcDNA3.1 vector (123).** (from Weigert et al., 2003) (pRL-CMV: Vector, GFAT: glutamine:fructose-6-phosphate amidotransferase, EcoR1 and BamH1 are restriction splice sites).
2. pcDNA3-GFAT/577 contains human GFAT1 cloned into the pcDNA3.1 vector but with histidine 577 mutated to alanine, resulting in the complete loss of GFAT enzyme activity (Weigert et al., 2003).

3. pcDNA3-GFAT/677 contains human GFAT1 cloned into the pcDNA3.1 vector but with lysine 667 mutated to alanine, leading to complete loss of GFAT enzyme activity (Weigert et al., 2003).

![Diagram of pcDNA3 vector](from brochure supplied by Invitrogen, Carlsbad, CA, USA).

*There is an ATG upstream of the Xba I site.*
4. pPIIβ-1,317 is a full-length human ACCβ promoter reporter luciferase construct that contains 4 E-boxes (CANNTG) (Figure 9) (Makaula et al, 2006).


ACCβ promoter region = 1,317 bp

Four “E-boxes” identified (E1-E4): important regulatory elements for transcription factors such as upstream stimulatory factor (USF)

5. TransLucent USF Reporter Vector (USF-L) contains promoter recognition sites for both upstream stimulatory factor 1 (USF1) and upstream stimulatory factor 2 (USF2) cloned into a pTransLucent Vector (catalog number LROO86, Panomics, Redwood City, USA) (Figure 10).
2.1.4. Preparation of plasmid DNA

Each expression vector was amplified in *Escherichia coli* cultures (JM109 competent cells, Promega, Madison, WI, USA) and extracted using the Qiagen® Plasmid Purification Maxi Kit (Qiagen, Invitrogen, Carlsbad, CA, USA). Purified DNA was quantified using a spectrophotometer (wavelengths of 260 nm and 280 nm) and its quality checked by restriction enzyme analysis. The DNA was electrophoresed on a 1% agarose gel to check for the quality of the DNA.
2.1.5. Transfection procedures

On Day 2 of transfection experiments the cells were transfected with the DNA of interest (Figure 6). Transfections were performed in triplicate for each experiment and repeated to generate the necessary numbers for statistical analysis. First, a stock solution of pGL3-Control DNA (pRL-CMV) was made in media concentration of 10 ng/ml (see step 1 of Figure 11). The media contained DMEM and 4 mM L-glutamine. The stock solution was aliquoted into separate microfuge tubes to a final volume of 165 µl for every transfection experiment (consisting of three replicates for each experiment) (see step 2 of Figure 11). DNA was aliquoted into its respective microfuge tubes with pGL3-Basic making up the total DNA mass to 0.75 µg (step 3 of Figure 11).

A second stock solution was then prepared with an equal volume of media containing Fugene 6 Transfection Reagent (Roche, Penzberg, Germany). Here, a 2:1 ratio of Fugene 6: DNA (with DMEM and 4 mM L-glutamine) was used. 165 µl of the Fugene 6 solution was then added to each of the microfuge tubes containing DNA and incubated at room temperature for 15 minutes (steps 4, 5 of Figure 11).

Meanwhile, 0.9 ml of fresh medium (containing DMEM, 10% FCS and 4 mM L-glutamine) was added to the H9c2 cells before the transfection. The final volume of the DNA/Fugene 6 cocktail therefore equalled 330 µl in each microfuge tube for each transfection experiment.
Step 1:  
10 ng PRL-CMV  
+  
DMEM + 4 mM L-glutamine

Step 2:  
Aliquot 165 µl in each 1.5 ml microfuge tubes

Step 3:  
Aliquot DNA into tubes up to a total of 0.75 µg DNA

Example:
Tube 1:  
= 0.75 µg of pGL3-Basic

Tube 2:  
= 0.25 µg pPIβ-1317 + 0.5 µg pGL3-Basic

Step 4:  
Aliquot 165 µl in each 1.5 ml microfuge tube with DNA

1.5 µl Fugene 6 transfection reagent (n = 3)  
+  
DMEM + 4 mM L-glutamine

Step 5:  
Incubate for 15 minutes at room temperature

Step 6:  
Aliquot 100 µl DNA:Fugene cocktail into each of three wells representing a triplicate

Example:
Tube 1:  
=  
(Fugene:DNA = 2:1)

Figure 11: Outline of day 2 transfection procedure.  
(n: number of samples, DMEM: Dulbecco’s modified Eagle’s medium, pGL3-Basic: vector, PRL-CMV: Renilla luciferase construct).

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The DNA/Fugene 6 solution was added to the H9c2 myoblasts (100 µl per well to make a final volume of 1 ml per well). The solutions were applied in triplicate (step 6 of Figure 11). The plates were gently rocked and then incubated at 37°C for 24 hours.

After 24 hours the media on the cells was changed. During this media change various drugs of interest were added to cells. We employed the following agents: 0.1 mM, 1 mM and 2 mM alloxan (Sigma-Aldrich, St. Louis, Missouri), 40 µM and 80 µM azaserine (Sigma-Aldrich, St. Louis, Missouri), 40 µM and 80 µM 6-Diazo-5-oxo-L-norleucine (DON) (Sigma-Aldrich, St. Louis, Missouri) and 5 mM and 10 mM streptozotocin (STZ) (Sigma-Aldrich, St. Louis, Missouri). Twenty-four hours later cells were lysed, protein extracted and expression of luciferase measured using the Glomax luminometer (Promega, Fitchburg, WI, USA).

The protocol and reagents used were as stipulated in the manual of the Dual-Luciferase Reporter Assay Kit (Promega, Fitchburg, WI, USA). First the cells were washed with phosphate buffer saline and then 200 µl of Passive Lysis Buffer (Promega, Fitchburg, WI, USA) was added to each well. The 12-well plate was then incubated on a shaker at room temperature for 15 minutes. The lysis buffer and cells from each well were transferred to microfuge tubes. A separate microfuge tube was used for each well of the 12-well plate. The microfuge tubes carrying the cells and lysis buffer were stored at -80°C (Figure 12). On Day 5 the lysate was thawed in water and vortexed. The microfuge tubes were centrifuged at 12,000 rpm at 4°C for 2 minutes with a ALC multispeed refrigerated centrifuge PK 121R (Intergrated Services, New
Jersey, USA). 10 µl of each sample was aliquoted into a separate well on a 96-well luminometer plate (Amersham, Buckinghamshire, UK).

**Step 1:** Remove media and wash of cells with phosphate buffer saline

**Step 2:** Add 200 µl of Passive Lysis Buffer from Dual-Luciferase Reporter Assay Kit in each well of 12-well culture plate

**Step 3:** Leave plate on a shaker for 15 minutes at room temperature

**Step 4:** Extract lysates and place each into separate microfuge tubes and store at -80°C

**Example:**

```
1 2 3 4

Store at -80°C
```

Figure 12: Outline of day 4 lysate extraction.
Two reagents had to be prepared. Reagent 1 contained Luciferase Assay Reagent II (LAR II) and reagent 2 Stop and Glo reagent (Figure 13). Both reagents were provided with the Dual-Luciferase Reporter Assay Kit (Promega, Fitchburg, WI, USA). The luminometer plate, together with the two reagents, was placed in the luminometer. The luminometer was set up to add 50 µl of LAR II, delay for 2 seconds and then take a reading for 8 seconds. This gave a measurement for human ACCβ promoter (pPIIβ-1,317) activity. The luminometer then added “Stop and Glo” to the same well and delayed for 2 seconds before measuring the light released (for another 8 seconds). This yielded the measurement for the Renilla construct (pRL-CMV) of the same sample. The process was repeated for each well, providing two readings for each sample (Figure 13).
**Figure 13: Performing the luciferase assay on day 5.**

**Step 1:** Thaw samples in a waterbath at room temperature

**Step 2:** Vortex samples and then centrifuge at 12,000 rpm at 4°C for 2 minutes

**Step 3:** Aliquot 10 µl of each sample onto a 96-well luminometer plate

**Step 4:** Place in luminometer with reagent 1 and reagent 2

**Step 5:** For each sample the luminometer will add 50 µl of reagent 1, take a reading, add 50 µl reagent 2 and then take a second reading.

**Reagent 1:** Luciferase Assay Reagent (measures activity of firefly luciferase gene, i.e. promoter of interest)

**Reagent 2:** Stop and Glo Reagent (measures activity of Renilla luciferase gene, i.e. pRL-CMV which is the normalising control)

Normalize for transfection efficiency and against the vector and perform statistical analysis
2.1.6. Statistical analyses of transfection results

Luminometer firefly readings for each experiment were normalised against its respective Renilla luminometer reading by dividing the firefly reading by its renilla reading. Each experiment was performed in triplicate each time a transfection was carried out. Each replicate of an experiment was added together and then divided by 3 to give a mean value for each experiment. This provided a mean value for each experiment that was normalised according to transfection efficiency and cell number. To eliminate any vector effects (i.e. in which the construct were cloned) on the experiment, each of the mean values were divided by the vector’s mean value.

The transfection experiments were repeated numerous times to provide a larger number of replicates for each experiment. Measurements from all the same experiments were combined and Graphpad Instat version 3.01 (GraphPad Software Inc., San Diego CA) used to perform statistical analyses of these values. First the mean and standard error of the mean and standard deviation was calculated. A 95% confidence interval was also calculated. Replicates of an experiment that were outside the 95% confidence interval or were not within two standard errors of the mean were discarded. The remaining replicates were used to plot a graph. Student-Newman statistical test was used to see if there were in significant differences between controls and the experiments in each set of experiments. Values $p< 0.05$ were taken as being significant.
2.2. Protein and RNA extraction from transfected cells

On day 4 of transfection experiments, protein and RNA were extracted from H9c2 myoblasts using an Allprep RNA/Protein extraction kit (Qiagen, Invitrogen, Carlsbad, CA, USA). This was done in duplicate so that a luciferase assay could be performed to confirm that transfection experiments were indeed successful. Protein extracted from these cells was used in Western blotting experiments, while the RNA was stored for future real-time PCR experiments.

2.3. Western blotting

2.3.1. Sample preparation and quantification of protein from cells

The protein extracted using the Allprep RNA/Protein extraction Kit was quantified using the Qubit fluorometer (Invitrogen, Carlsbad, CA, USA). Protein was prepared with an equal volume of sample buffer and boiled for 5 minutes, centrifuged at 14,000 rpm for 5 seconds before loading onto a SDS-polyacrylamide gel for electrophoresis.
2.3.2. Tissue biopsies acquired from db/db transgenic mouse model of type-2 diabetes

Previously collected heart tissues from 20-week-old male and female db/db mouse and matching controls (db/+), were used. This mouse has a point mutation in the leptin receptor gene and gains weight with age (15). They also develop insulin resistance and hyperglycaemia, eventually progressing to type-2 diabetes. The symptoms of type 2 diabetes are similar to that observed in the human condition making this a good model to explore my hypothesis in an in vivo setting. We used heterozygous (db/) instead of homozygous (+/) as controls because of a lack of animal supply.

2.3.3. Sample preparation and quantification of protein from heart tissues

Protein was extracted from male and female db/db mouse and heterozygous mouse heart muscle tissue biopsies and stored in RIPA buffer (refer to Appendix 1 for extraction method). Total protein was quantified using the Bradford method for protein determination (see Appendix 2 for Bradford method). 50 µg of sample was then added to an equal volume of sample buffer and boiled for 5 minutes. The samples were then electrophoresed on a SDS-polyacrylamide gel.
2.3.4. Gel analysis

Protein extracts were electrophoresed on a 10% SDS-polyacrylamide gel with a 4% stacking gel at 200 V for 1 hour. 10 µl of a protein marker (BioRAD precision Plus protein standard cat. # 161-0374, Bio-RAD Laboratories, California, USA) was used in lane 1 of gels. The amount of protein loaded per well was 50 µg for each sample. The protein was then transferred to a PVDF membrane using a semi-dry transfer apparatus (Bio-RAD Laboratories, California, USA) for 1 hour (See Appendix 3 for electrotransfer procedure).

The membranes were used to probe for three proteins of interest:
1) O-GlcNAc was detected using an O-GlcNAc Western Blot Detection Kit (Pierce, catalogue number 24565, Rockford, Illinois) (Refer to Appendix 4 for complete protocol).

2) GFAT (Refer to Appendix 5 for the complete protocol). The GFAT antibody was kindly donated by Dr. Cora Weigert (University of Tübingen, Germany).

3) β-actin (Cell Signaling, Danvers, MA, USA) was also employed to check for equal protein loading (Refer to Appendix 6 for protocol).
2.3.5. Analysis of Western blot results

Results from western blots were analysed by densitometry. For the in vivo studies db/db protein extracts were probed with O-GlcNAc or GFAT antibody, reprobed with β-actin (Cell Signaling, Danvers, MA, USA) and then densitometry was performed. The densitometry was performed by scanning the blots with an HP Scanjet 3500c scanner (Hewlett Packard, Palo Alto CA) and densitometry was performed on the scanned pictures using Un-Scan-It Gel version 5.1 (Silk Software, Orem, UT). The densitometry value for a sample was normalised against its β-actin densitometry value. Then all the normalised densitometry values from db/db were pooled together (likewise for db/+). The combined protein expression for db/+ was compared to that of db/db using appropriate statistical analyses to evaluate if there was a significant change in protein expression.
Chapter 3

Results
3.1. Transfections

Before investigating our hypothesis, my initial experiments were focused on optimising the transfection procedure for H9c2 myoblasts. Here myoblasts were transfected with different concentrations of the ACCβ promoter-luciferase reporter construct (pPIIβ-1317/+65) ± different concentrations of GFAT expression vector ± different concentrations of dominant negative GFAT/577 or GFAT/677 constructs. After considerable effort, I eventually was in a position to select optimal DNA concentrations for use in transfection experiments (Figure 14). For these initial experiments a total of 1 µg of DNA per experiment was employed. The results obtained showed inconsistency. The transfection procedure was then re-optimised. Thus, for the experiments which followed we used 0.75 µg of DNA per experiment; however the Fugene 6 to DNA ratio remained at 2:1.

![Figure 14: Results of pilot co-transfection experiments performed with ACCβ promoter, GFAT and dominant negative constructs (GFAT/577 and GFAT/667).](image)

(* p<0.001 vs. ACCβ [n=9], ** p<0.05 vs. ACCβ [n=9], *** p<0.01 vs. ACCβ [n=9], # p<0.01 vs. ACCβ [n=9], ## p<0.001 vs. ACCβ+ 0.4 µg GFAT [n=5]. Vector: pRL-CMV, ACCβ: pPIIβ-1317/+65, GFAT: glutamine:fructose-6-phosphate amidotransferase).
After establishing and optimising the transfection procedure I next proceeded to investigate my hypothesis. To ensure that results were statistically relevant experiments were repeated several times (at least an n=6).

I began by transiently transfecting H9c2 myoblasts with the human ACCβ promoter-luciferase reporter construct (pPIIβ-1317/+65) ± a GFAT expression vector ± dominant negative GFAT/577 or GFAT/677 constructs. The results showed that GFAT overexpression induced ACCβ gene promoter activity by 75 ± 23% (p<0.001, n=6). Co-transfection with either GFAT/577 or GFAT/667 constructs resulted in marked attenuation of this induction (Figure 15).

![Figure 15: GFAT overexpression induces ACCβ gene promoter activity.](http://scholar.sun.ac.za)

* + GFAT/577
* + GFAT/667

0 1 2 3 4 5 6 7
Fold induction

Vector ACCβ GFAT ACCβ+GFAT

Figure 15: GFAT overexpression induces ACCβ gene promoter activity.

(∗ p<0.001 vs. ACCβ [n=9], ∗∗ p<0.001 vs. ACCβ+GFAT [n=6], # p<0.001 vs. ACCβ+GFAT [n=6]. Vector: pRL-CMV, ACCβ: pPIIβ-1317/+65, GFAT: glutamine:fructose-6-phosphate amidotransferase).
To assess the extent of GFAT regulation on ACCβ gene promoter activation, H9c2 myoblasts were treated with varying doses of L-glutamine (0 mM, 4 mM and 8 mM) a substrate for the hexosamine biosynthetic pathway (HBP). Here, glutamine induced ACCβ promoter activity in a dose-dependent manner (Figure 16). There was no significant change in ACCβ promoter activity in the absence of GFAT (Figure 16).

Figure 16: Glutamine induces ACCβ promoter activity in a dose-dependent manner. (* p<0.05 vs. ACCβ + GFAT 0 mM L-Glutamine [n=7], ** p<0.01 vs. ACCβ+GFAT + 0 mM L-Glutamine [n=7]. Vector: pRL-CMV, ACCβ: pPIIβ-1317/+65, GFAT: glutamine:fructose-6-phosphate amidotransferase).
Since GFAT-induced ACCβ promoter activity was inhibited by dominant negative constructs, I next performed additional confirmatory experiments by pharmacological inhibition of GFAT. Two inhibitors of GFAT were used, i.e. 6-Diazo-5-oxo-L-norleucine (DON) and azaserine (Figure 17).

![Diagram of metabolic pathways involving GFAT inhibition]

**Figure 17: Description of inhibitors of HBP and where they operate.**
Glc-6-P: glucose-6-phosphate, F-6-P: fructose-6-phosphate, GlcN-6-P: N-glucosamine-6-phosphate, UDP-GlcNAc: uridine diphospho-N-acetylglucosamine, OGT: O-linked β-N-acetylglucosamine transferase, GFAT: glutamine:fructose-6-phosphate amidotransferase, O-GlcNAc: O-linked N-acetylglucosamine
Two concentrations of DON were used (40 µM and 80 µM) to determine the effectiveness of DON inhibition (Figure 18). It was suggested in the literature that 40 µM of DON would be a physiologically relevant dose to achieve the required effect. Since the cell line used in this study was different to published data the dose was also doubled to 80 µM in case the 40 µM dose failed to inhibit GFAT.

Figure 18: Results of co-transfection experiments performed with ACCβ, GFAT and different concentrations of inhibitors of HBP.

(* p<0.05 vs. ACCβ [n=9], ** p<0.05 vs. ACCβ+GFAT [n=10], *** p<0.01 vs. ACCβ+GFAT [n=9], # p<0.01 vs. ACCβ+GFAT+STZ [n=7]. Vector: pRL-CMV, ACCβ: pPIIβ-1317/+65, GFAT: glutamine:fructose-6-phosphate amidotransferase, DON: 6-Diazo-5-oxo-L-norleucine, STZ: Streptozotocin).
Since both the 40 µM and 80 µM concentrations worked equally well (Figure 19) we continued by using the 40 µM dose as it was more cost effective.

**Figure 19: Pharmacological inhibition of GFAT abolishes ACCβ promoter induction.**

\[* p<0.001 vs. ACCβ [n=35], ** p<0.001 vs. ACCβ+GFAT [n=25]. Vector: pRL-CMV, ACCβ: pPIIβ-1317/+65, GFAT: glutamine:fructose-6-phosphate amidotransferase, DON: 6-Diazo-5-oxo-L-norleucine].

Two concentrations of azaserine were tested (40 µM and 80 µM) (Figure 19). Again, as before we employed the 40 µM dose because of cost effectiveness. Azaserine markedly blunted GFAT-induced ACCβ promoter activity (Figure 20). Therefore both the dominant negative and pharmaceutical data substantially strengthen our hypothesis that the HBP regulates ACCβ promoter activity.
Additional pharmaceutical agents were next employed to further examine the regulation of ACCβ promoter activity by downstream targets of the HBP. Here H9c2 myoblasts were treated with various concentrations of alloxan (0.1 mM, 1 mM and 2 mM), an inhibitor of O-linked β-N-acetylglucosaminyl transferase (OGT) (Figure 17). The results show that increasing concentrations of alloxan result in a corresponding decrease of ACCβ promoter activity in response to GFAT overexpression (Figure 21). Moreover, at the 2 mM alloxan concentration there was a significant reduction in ACCβ promoter activity (Figure 21).
A second pharmacologic inhibitor, i.e. streptozocin (STZ), an inhibitor of β-N-acetylglucosaminidase (O-GlcNAcase) was used to investigate downstream targets of the HBP on the regulation of ACCβ promoter activity (Figure 17). Thus I expected to observe an increase in ACCβ promoter activity in this instance. Two concentrations of STZ were used (5 mM and 10 mM) in preliminary testing, and we thereafter employed the 5 mM dose (Figure 19). Interestingly, these results show that STZ further enhanced GFAT-induced ACCβ promoter activity (Figure 22).
These data therefore show that the human ACCβ gene promoter is regulated by these two downstream HBP regulators, i.e. OGT and O-GlcNAcase. To further strengthen these findings I next performed experiments using both an inhibitor of GFAT (i.e. DON) and an inhibitor of O-GlcNAcase (i.e. STZ) (Figure 17). In agreement with our earlier findings, DON attenuated STZ-mediated upregulation of ACCβ promoter activity (Figure 23).

Figure 22: Pharmacological inhibition of O-GlcNAcase further increases ACCβ promoter activity.
(* p<0.001 vs. ACCβ [n=35], ** p<0.001 vs. ACCβ+GFAT [n=12]. Vector: pRL-CMV, ACCβ: pPIIβ-1317/+65, GFAT: glutamine:fructose-6-phosphate amidotransferase, STZ: Streptozotocin).

These data therefore show that the human ACCβ gene promoter is regulated by these two downstream HBP regulators, i.e. OGT and O-GlcNAcase. To further strengthen these findings I next performed experiments using both an inhibitor of GFAT (i.e. DON) and an inhibitor of O-GlcNAcase (i.e. STZ) (Figure 17). In agreement with our earlier findings, DON attenuated STZ-mediated upregulation of ACCβ promoter activity (Figure 23).
To gain further insight into transcriptional mechanisms underlying GFAT-mediated induction of ACCβ promoter activity, we employed a candidate transcription factor approach. Since our laboratory previously implicated upstream stimulatory factors (USFs) in ACCβ promoter regulation (Makaula et al., 2006) I next tested USF1 and USF2 as transcriptional regulators in the GFAT-mediated induction of ACCβ gene promoter. Here, USF2 induced ACCβ promoter activity by 44 ± 23% (p<0.001, n=6). However, when co-transfected with GFAT, USF2 did not further increase ACCβ promoter activity (Figure 24).

Figure 23: Pharmacological inhibition of inhibition of GFAT together with inhibition of O-GlcNAcase attenuates ACCβ promoter induction.

Figure 24: Pharmacologic inhibition of inhibition of GFAT together with inhibition of O-GlcNAcase attenuates ACCβ promoter induction. (* p<0.001 vs. ACCβ [n=33], ** p<0.001 vs. ACCβ+GFAT [n=10], *** p<0.001 vs. ACCβ+GFAT [n=12]. # p<0.05 vs. ACCβ+GFAT [n=7], # p<0.001 vs. ACCβ+GFAT+STZ [n=7]. Vector: pRL-CMV, ACCβ: pPIIβ-1317/+65, GFAT: glutamine:fructose-6-phosphate amidotransferase).
We further tested USF-mediated regulation of the ACCβ promoter by co-transfecting a GFAT expression construct together with a USF luciferase reporter construct (USF-L = Upstream Stimulatory Factor TransLucent™ Reporter Vector) that contains multiple promoter binding sites for USFs. Here, we found a marked induction of the USF-L reporter construct by 80 ± 12% (p<0.001, n=5) compared to the vector (Figure 25).

Figure 24: USF2 overexpression induces ACCβ promoter activity.
(∗ p<0.001 vs. ACCβ [n=6], ** p<0.001 vs. ACCβ [n=6]. Vector: pRL-CMV, ACCβ: pPIIβ-1317/+65, GFAT: glutamine:fructose-6-phosphate amidotransferase, USF1: upstream stimulatory factor 1, USF2: upstream stimulatory factor 2.)
Figure 25: GFAT overexpression enhances USF transcriptional activation.
3.2. Western blotting

3.2.1. In vitro experiments

To assess the degree of O-GlcNAcation after transfection, we performed Western blotting using our well-established protocol. We employed an Allprep RNA/Protein extraction Kit (Qiagen, Invitrogen, Carlsbad, CA, USA) to simultaneously isolate and purify RNA and protein. Protein extracts were probed using an O-GlcNAc Western blot Detection Kit (Pierce, catalogue number 24565, Rockford, Illinois). The membrane was stripped and reprobed with β-actin antibody. Our preliminary results suggest little difference in the level of O-GlcNAc modified proteins when transfected with ACCβ ± GFAT (n=3). However, there is a decreasing trend when cells were treated with 40 µM DON (Figure 26). Additional samples need to be tested to further assess this.

![Figure 26: Degree of O-GlcNAcation after GFAT transfection into H9c2.](image)

(The densitometry readings were normalised for protein loading using β-actin. Vector: pRL-CMV [n=1], ACCβ: pPllβ-1317/+65 [n=3], GFAT: glutamine:fructose-6-phosphate amidotransferase [n=3], DON: 6-Diazo-5-oxo-L-norleucine [n=3]).
3.2.2. In vivo experiments

We investigated our hypothesis in a mouse model of type 2 diabetes. Total protein was purified from previously collected heart tissue from heterozygous (db/+ ) and homozygous (db/db) female mice and probed for degree of O-GlcNAcation. Due to animal limitations, we could only test 3 female db/db and 3 matched db/+ controls. Our preliminary blotting data suggest that O-GlcNAc modification of proteins are decreased in the female db/db mouse (Figure 27).

![Figure 27: Degree of O-GlcNAcation in db/db female mouse heart.](http://scholar.sun.ac.za)

(The densitometry readings were normalised for protein load using β-actin. * p<0.0008 vs. female db/+ [n=3]).
To test whether alterations in O-GlcNAc modification are due to a change in GFAT expression, we also assessed GFAT peptide levels in db/db versus db/+ female mouse hearts. Our preliminary results suggest a decreasing trend in GFAT protein expression (Figure 28). However, additional sample numbers are required to confirm whether these trends are indeed true.

![Figure 28: GFAT peptide levels in female diabetic mouse hearts.](image)
(The densitometry readings were normalised for protein load using β-actin).
To elucidate whether these observations were gender dependent, we also analysed male db/+ and db/db mouse heart tissue using the GFAT antibody (due to time constraints we could not perform O-GlcNAc analysis for male heart tissues). Again, we were limited by sample numbers (n=3 for male db/db and db/+). Our early results suggest no differences in GFAT expression between male db/+ and db/db mice (Figure 29). However, further studies are needed to confirm our preliminary data and whether gender-dependent differences do indeed exist.

**Figure 29: GFAT peptide levels in male diabetic mouse hearts.**
(The densitometry readings were normalised for protein load using β-actin [n=3]).
Chapter 4
Discussion
Discussion

For this study we tested the hypothesis that increased flux through the hexosamine biosynthetic pathway (HBP) induces cardiac ACCβ gene expression. A secondary hypothesis resulting from this is that HBP-mediated induction would reduce mitochondrial fatty acid uptake/oxidation and result in an accumulation of intracellular lipid in the insulin resistant setting (hyperlipidaemia). This lipid accumulation will trigger signaling pathways resulting in cell death, insulin resistance and contractile dysfunction. The secondary hypothesis now forms part of my ongoing studies. The major novel finding of this study is that the human ACCβ gene promoter is inducible by increased flux via the HBP. Moreover we find that upstream stimulatory factors play a crucial role in transcriptional regulation of this process.

Increased flux through the hexosamine biosynthetic pathway has been linked to an increase in glucose and glucosamine availability. An increase in HBP flux results in higher UDP-GlcNAc production and a general increase in O-GlcNAc modification of a large number of proteins and transcription factors. O-GlcNAc has been shown to operate in opposition to phosphorylation, glycosylating proteins that are dephosphorylated and vice versa (21, 59, 65). Studies have proposed that the hexosamine biosynthetic pathway functions as a cellular nutrient sensor and numerous papers have showed a correlation between increased flux through the HBP and insulin resistance (10-12, 41, 79, 96). For example, Marshall et al. (1991) first proposed a role for glucose flux via the HBP in insulin resistance from a series of experiments performed in isolated rat adipocytes (77). Other studies found a correlation between the
HBP and insulin resistance in rodents treated with glucose/glucosamine (18, 22, 25, 51). Recently, increased O-GlcNAc modification has been attributed to altered glucose uptake resulting in insulin resistance (79, 80, 96, 132). A study investigating overexpression of OGT in adipose and muscle of mice resulting in increased O-GlcNAcation showed insulin resistance and hyperleptinaemia (81, 132). These studies would suggest that increased HBP flux and increased O-GlcNAcation can impact on fatty acid and glucose pathways. In this study we tried to confirm this \textit{in vivo} by investigating GFAT expression and O-GlcNAc modification in the mouse heart. Although, we were able to see some differences in our results, the sample size was too small to draw any meaningful conclusions.

Several experiments completed for this study supports our primary hypothesis.

1. We found that GFAT overexpression increased \textit{ACC}β promoter activity by \textasciitilde 75\% in cardiac myoblasts. This would suggest that increased HBP flux results in O-GlcNAc modification of transcriptional targets thereby inducing \textit{ACC}β promoter activity.

2. \textit{ACC}β gene promoter activity was induced in a dose-responsive manner to increasing glutamine concentrations (glutamine being an HBP substrate).

It is important to note that the media used in culturing cells utilises 4 mM L-glutamine for standard growing conditions. Cells that were cultured at 0 mM L-
glutamine grew slower than those at 4 mM and 8 mM L-glutamine. We therefore had to ensure that the L-glutamine concentration of our growing media maintained the same concentration for all experiments to ensure that results were reproducible. We are considering using varying doses of glucosamine (another substrate for the HBP) in future experiments.

3. Cotransfection of myoblasts with two dominant negative constructs (competitive inhibitors of GFAT) attenuated dGFAT-mediated ACC\(\beta\) promoter induction.

4. Two specific pharmaceutical inhibitors of GFAT, i.e. azaserine and 6-Diazo-5-oxo-L-norleucine attenuated GFAT-induced upregulation of ACC\(\beta\).

5. To determine whether downstream regulators were induced in this process we employed appropriate pharmaceutical inhibitors of the HBP.

The first downstream target of the HBP we chose to investigate was OGT, an enzyme that converts UDP-GlcNAc to O-GlcNAc and then binds it to serine and threonine residues of proteins, lipids and transcription factors (12, 64, 71). Alloxan (OGT inhibitor) administration reduced GFAT-induced ACC\(\beta\) promoter activity. Of note, O-GlcNAcation is a very important regulatory step in developmental processes (42). Previous studies found that complete inhibition of OGT is lethal (42, 43, 86, 104). However for these experiments performed no obvious signs of large scale cell death were observed on examination under the light microscope at 10,000 X magnification. I propose that this could be because the concentration we used was low (0.1 mM - 2 mM) and
probably did not completely inhibit OGT activity. Another possibility may be that we were investigating the HBP in rat cardiac-derived myoblasts, and hence the aforementioned concern would only be applicable to tissue and whole organ development.

We also tested a pharmaceutical inhibitor of O-GlcNAcase, i.e. streptozotocin (132). Here we found that streptozotocin treatment enhanced ACCβ promoter activity. Of note, streptozotocin is an inhibitor that is widely used to investigate type 1 diabetes (95), known to destroy pancreatic β-cells (93). However, in our model we did not observe any cell death. This was confirmed by visually comparing cell viability between wells that contained seeded cells treated ± streptozotocin between media changes on day 3 and day 4 of the transfection procedure (Figure 2 of Methods section). I propose that this is probably due to the cell type (H9c2 myoblasts) used in this study. It has also been shown that an increase in O-GlcNAcation is linked to insulin resistance (5, 14, 42, 81, 90, 120). This would explain why streptozotocin is lethal to pancreatic β-cells and why it is so effective in creating a type 1 diabetic model, i.e. the type 1 diabetes is created by destroying the insulin-producing cells after they fail to maintain β-cell compensation (16, 93). Cardiomyocytes and H9c2 myoblasts (precursor cells to cardiomyocytes) do not produce insulin or function as β-cells and thus would not be affected in this way.

To further strengthen our hypothesis we combined DON and streptozotocin since these inhibitors have opposite effects on the HBP. We found that DON decreased ACCβ promoter activity while streptozotocin increased it. Since
DON acts higher up in the HBP pathway DON inhibition probably outweighed streptozotocin inhibition, resulting in a decreased ACCβ promoter activity.

The data from transfection experiments strongly supported the hypothesis that increased flux via the HBP induces ACCβ promoter activity. However, increased HBP flux effects are expected to be mediated via O-GlcNAcated transcriptional regulation. It is likely that a number/cluster of transcription factors form a complex with each other to regulate the ACCβ promoter in response to high HBP flux. In light of this we followed a candidate transcription factor approach. Here we focused on upstream stimulatory factors since it was previously shown to upregulate the ACCβ gene promoter (75). Promoter studies in cardiomyocytes have suggested that USF1 can mediate ACCβ promoter activity in a glucose-dependent manner, but implicates the pentose phosphate pathway (75). USF1 has also been shown to induce TGF-β, which has been shown to be upregulated by the HBP (124).

These experiments were performed toward the end of my thesis and thus the results are of a preliminary nature. However, my data revealed that USF2, and not USF1, regulates ACCβ promoter activity via the HBP. When the ACCβ promoter was cotransfected with GFAT together with USF2, this did not show a further increase versus USF2 alone. This could be because a “ceiling level” of overexpression of the USF2 construct was reached. We also cotransfected GFAT with a USF reporter-promoter construct (USF-L) containing consensus binding sites for USFs. Interestingly, the USF-L construct was induced to a much greater extent then with USF2 alone. These data therefore suggest that USF1-USF2 heterodimers may induce ACCβ to a
greater extent. In agreement, other studies have shown that USF1 and USF2 can form either homo- or heterodimers (107, 119).

USF1 has also been associated with the metabolic syndrome and type 2-diabetes (83). Due to time constraints I could not perform a conventional transfection with USF1 and USF2 together. Unfortunately, due to a limitation of animal numbers and time constraints our Western blotting results can only be considered as preliminary. Preliminary results of Western blotting showed that O-GlcNAcation was increased when the cells were cotransfected with GFAT and ACCβ reporter-promoter construct but that O-GlcNAcation decreased when the myoblasts were treated with DON. In our preliminary mouse studies we saw a reduction of GFAT protein in the female db/db from control (db/+), but saw no significant difference in the male mice. We also found a decrease in O-GlcNAcation of proteins in female db/db from control (db/+). Further experiments will have to be carried out before any assumptions can be made as to the relevance of this data. Although, we can speculate that an inability of the male diabetic heart to reduce CPT1 inhibition could explain why female hearts show a relatively higher degree of protection compared to male hearts (30, 56, 116). Should this be found to be the case, we may assume that hormones such as estrogen or testosterone may be involved in the regulation of the HBP.
Conclusion

Our data reveal a novel finding, i.e. that increased flux through the hexosamine biosynthetic pathway activates USFs thereby inducing $\text{ACC}\beta$ gene promoter activity in cardiac-derived myoblasts (Figure 30). To date no data, as far as I am aware, has been published in this regard. Further studies are underway to test our secondary hypothesis, i.e. that HBP-induced $\text{ACC}\beta$ induction reduces fatty acid oxidation, thereby leading to intracellular lipid accumulation due to a mismatch between sarcolemmal FA uptake and mitochondrial FA oxidation in the insulin resistant setting (Figure 30). Subsequently, we propose that intramyocardial lipid accumulation triggers signaling pathways resulting in cell death, insulin resistance and contractile dysfunction.

Figure 30: Schematic diagram of completed studies and planned future investigations.
Limitations

A limitation of the transfection technique is the time it takes to optimise the transfection protocol. This limited the initial progress of this project.

The effectiveness of an inhibitor of any pathway is determined by its specificity for its target. The inhibitors we employed for our experiments are well documented in the literature (17, 132). We tried to use as many inhibitors for the same target site as we could to strengthen our results. However, we are aware these inhibitors could also inhibit/enhance other intracellular signaling pathways. However, we also employed two dominant negative GFAT constructs which considerably strengthened our data.

Due to time constraints and limitation on animal numbers we were unable to investigate the differences in regulation of the HBP between male and female db/db mice. This means that our Western blotting results are only preliminary and more experiments would have to be performed to confirm this result.
**Future Studies**

- We are currently performing real-time PCR on RNA extracted from transfected myoblasts to determine whether the increase in ACCβ promoter activity results in a concomitant increase in endogenous RNA expression.

- We will also be performing additional USF transfection experiments and DNA binding assays (ELISAs and CHiP) to confirm the role of USF1 and USF2 in the transcriptional regulation of ACCβ.

- We would also like to confirm our L-glutamine transfection experiments by administering different doses of glucosamine to transfected myoblasts. Since it is more potent than L-glutamine, we expect to see a greater effect on ACCβ promoter activity.

- We are also currently investigating an additional inhibitor, i.e. O-(2-acetamido-2-deoxy-D-glucopyranosylidene) amino-N-phenylcarbamate (PUGNAc) (132), proposed to be more potent than streptozotocin.

- We also intend to probe heart tissues from various type-2 diabetic rodent models (db/db, Zucker and diet-induced rat model).

- To investigate our secondary hypothesis we will measure fatty acid accumulation with oil red staining and measure malonyl-CoA levels (Figure 30).
References


13. **Buse M, Robinson KA., Gettys TW., McMahon EG., and Gluve EA.**


14. **Buse M, Robinson KA., Marshall BA., Hresko RC., and Mueckler MM.**


17. **Champattanachai V, Marchase RB., and Chatham J.** Glucosamine protects neonatal cardiomyocytes from ischemia-reperfusion injury via

18. **Chen H, Ing BL., Robinson KA., Feagin AC., Buse MG., and Quon MJ.**


21. **Comer F, and Hart GW.** O-GlcNAc and the control of gene expression.


22. **Cooksey R, Hebert LF Jr., Zhu JH., Wofford P., Garvey WT., and McClain DA.** Mechanism of hexosamine-induced insulin resistance in transgenic mice overexpressing glutamine:fructose-6-phosphate...


33. **Garland P, and Randle PJ.** Control of pyruvate dehydrogenase in the perfused rat heart by the intracellular concentration of acetyl-coenzyme A. *Biochem J* 91: 6c, 1964.


83. **Miyake K, So WY., Poon EW., and Lam VK.** The linkage and association of the gene encoding upstream stimulatory factor 1 with type 2


94. **Robinson K, Sens DA., and Buse MG.** Pre-exposure to glucosamine induces insulin resistance of glucose transport and glycogen synthesis in


96. **Rossetti L.** Perspective: hexosamines and nutrient sensing.


99. **Sabourin L, and Rudnicki MA.** The molecular regulation of myogenesis.


111. Stanely W, Morgan EE., Huang H., McElfresh TA., Strek JP., Okere IC., Chandler MP., Cheng J., Dyck JRB., and Lopaschuk GD. Malonyl-CoA decarboxylase inhibition suppresses fatty acid oxidation and reduces lactate
production during demand-induced ischemia. *Am J Physiol Heart Circ Physiol*


smoke on biological processes important in atherogenesis. *BMC Cardiovasc


132. Zachara N, and Hart GW. Cell signaling, the essential role of O-


133. Zhang S, and Kim, KH. Acetyl-CoA carboxylase is essential for

nutrient-induced insulin secretion. *Biochem Biophys Res Commun* 229: 701-

705, 1996.
Chapter 5

Appendix
Appendix 1: Preparation of modified RIPA buffer

Preparation notes:

Label eppendorf tubes (±98).
Be sure to keep all inhibitors on ice and to add them last as indicated in the protocol.
Many of the reagents have been made up before use.

For 100ml modified RIPA buffer:

1. Prepare 50 mM Tris-HCl (a buffering agent that prevents protein denaturation):
   - Add 790 mg Tris to 75 ml distilled H₂0. Add 900 mg NaCl and stir.
   - pH to 7.4 using HCl.

2. Add 10 ml of 10% NP-40 to the solution. (Made up previously)

3. Add 2.5 ml of 10% Na-deoxycholate (an ionic detergent to extract protein, protect from light) and stir. (Made up previously in H₂0)

4. Add 1 ml 100 mM EDTA pH 7.4 (Acts as a calcium chelator). (Made up previously)
5. Now add the following protease inhibitors:

   - Phenylmethylsulfonyl fluoride (PMSF) – 200 mM stock made up in isopropanol and stored at room temperature (extremely short half life in aqueous solutions)
   - Leupeptin (1 mg/ml in H₂O)
   - SBT1 (5 mg/ml in H₂O)
   - Benzamidine (1 M)

(All of the above are stored in the freezer with the exception of PMSF which is available as a stock)

6. Add the following protease phosphatase inhibitors:

   - Activated Na₃VO₄ – 200 mM stock made up in H₂O (stored in freezer along with the protease inhibitors)
   - NaF (200 mM stock) – (Made up previously)

(Note: Do not add phosphatase inhibitors if lysates are to be used for phosphatase assays)

7. Add 1 ml of Triton X-100.

8. Make up to 100 ml with distilled H₂O using a graduated cylinder.

9. Aliquot 1 ml of RIPA buffer into each eppendorf tube before finally storing at -20°C
Final concentrations:

- Tris-HCl: 50 mM
- NP-40: 1%
- Na-deoxycholate: 0.25%
- EDTA: 1 mM
- PMSF: 1 mM
- Leupeptin: 1 µg/ml
- SBT1: 4 µg/ml
- Benzamidine: 1 mM
- Na$_3$VO$_4$: 1 mM
- NaF: 1 mM
Appendix 2: Bradford protein quantification and sample preparation

Bradford reagent (5x concentrated):

Dilute 500 mg of Coomassie Brilliant blue G in 250 ml 95% ethanol.
Add 500 ml of phosphoric acid before mixing thoroughly.
Make up to one liter with distilled H\textsubscript{2}O (dH\textsubscript{2}O).
Filter and store at 4°C.

Bradford working solution:

Dilute stock in a 1:5 ratio with dH\textsubscript{2}O.
Filter using 2 filter papers (at the same time).
Solution should be a light brown color.

Bradford method:

- Thaw 1 mg/ml BSA stock solution.
- Thaw protein samples if in -80°C freezer. Keep on ice at all times.
- Make up a working solution of 100 µl BSA:400 µl dH\textsubscript{2}O. Vortex mixture.
- Mark 7 microfuge tubes for the standards as well as tubes for the samples to be tested.
• Now add BSA and water to marker microfuge tubes as follows:

<table>
<thead>
<tr>
<th></th>
<th>BSA (µl)</th>
<th>Water (µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blank</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>2 µl protein</td>
<td>10</td>
<td>90</td>
</tr>
<tr>
<td>4 µl protein</td>
<td>20</td>
<td>80</td>
</tr>
<tr>
<td>8 µl protein</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>12 µl protein</td>
<td>60</td>
<td>40</td>
</tr>
<tr>
<td>16 µl protein</td>
<td>80</td>
<td>20</td>
</tr>
<tr>
<td>20 µl protein</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>Each sample</td>
<td>0</td>
<td>95 H₂O</td>
</tr>
</tbody>
</table>

• Briefly vortex all the tubes.

• Now add 900 µl of Bradford reagent to each microfuge tube. Vortex again.

• Let the solutions stand for ~5 minutes (switch on the spectrophotometer in the meantime).

• Read absorbencies, twice each, at 595 nm.
  o If sample values fall outside the range of the highest standard then dilute with RIPA buffer.

• Make use of Excel to make a linear plot of absorbencies and then calculate the amount of each sample to be added to aliquots.
Sample preparation:

- Begin by setting beaker of water to boil.
  - Remember to keep protein samples on ice at this point.
- Make up a stock solution containing 850 µl of sample buffer and 150 µl of mercaptoethanol.
- Vortex the solution.
- Calculate the number of sample sets needed, each containing one representative of each protein sample.
- Add sample buffer to each aliquot (do so under the fume hood to avoid exposure to harmful fumes).
- Add a volume of sample buffer equal to 1/3 of the final volume.
- Now add the amount of sample calculated previously to each respective microfuge tube.
- Punch small pin size holes in each tube then place in boiling water to stand for a period of 5 minutes.
- Spin tubes for a moment (~5 seconds) using the tabletop centrifuge.
- Samples can now be stored at -80°C.
Use of samples:

In the case that samples have been stored in the -80°C freezer:

- Start by bringing a beaker of water to the boil.
- Remove samples from the freezer.
- Make sure that small pin size holes have been punched in the top of each tube.
- Place in boiling water for a period of 5 minutes.
- Spin down momentarily (20 seconds) on the tabletop centrifuge (take care not to over centrifuge, especially if samples have been obtained from tissue.
- Samples can now be used for Western blotting.
Appendix 3: Electrotransfer of proteins (Semi-dry apparatus)

Solutions for PVDF membrane

Anode buffer 1:
- 0.3 M Tris-base, pH 10.4
- 20% methanol

Anode buffer 2:
- 25 mM Tris-base, pH 10.4
- 20% Methanol

Cathode buffer:
- 25 mM Tris-base
- 40 mM ε-aminohexanoic acid
- pH to 9.4
- Add 20% methanol

Need 8 Whatman 3MM papers and 1 PVDF membrane per gel cut to the required size.

1. Soak PVDF membrane in methanol for 15 seconds. Put the membrane in slowly, allowing it to soak by osmosis and prevent air bubbles from being trapped.

2. Soak 3 Whatman papers in anode buffer 1.


5. On the anode plate of the semi-dry apparatus put the 3 papers from step 2 (anode 1), roll with a glass tube.

6. Add 1 paper from step 3 (anode 2) and roll, making sure to remove bubbles.

7. Add the PVDF membrane and roll gently.

8. Take the gel with the proteins that have been electrophoresed and soak it in anode buffer 2. After soaking the gel, place it on top of the PVDF membrane.

9. Roll the gel with a wet tube making sure remove any bubbles.

10. Add the 4 Whatman papers from step 4 (cathode) on top of the gel and roll it with the glass tube.

11. Close the apparatus with the lid and set it to transfer for 1 hour at 15 volts and a current of 0.5 A.

12. After the electrotransfer is complete fix the protein to the PVDF membrane by placing it in methanol for a few seconds.
Appendix 4: O-GlcNAc Western blotting and detection

Note: The following description is reproduced as supplied by the O-GlcNAc Western blot detection kit (Pierce, catalogue number 24565, Rockford, Illinois).

Material Preparation:
(All reagents are supplied in the O-GlcNAc Western blot Detection Kit)

Wash Buffer Reconstitute one BupH™ Phosphate Buffer Saline (PBS) pack with 500 ml of ultrapure water and add 2.5 ml of 10% Tween®-20 for a final concentration of ~0.05%.

Blocking Buffer Dilute the 10x “Dilution buffer” (also functions as blocking buffer) to 1x in Wash buffer (e.g. 1 ml of 10x Dilution Buffer to 9 ml of Wash buffer).

Goat Anti-Mouse IgM -HRP Conjugate Prepare a 1 mg/ml stock solution by reconstituting the antibody in 75 µl of ultrapure water. Aliquot 6 µl the secondary antibody into separate micofuge tubes and add an equal volume of glycerol per aliquot. Store the aliquots at -20°C. By adding glycerol the final concentration of each aliquot changes to 0.5 mg/ml.
Procedure:

1. Remove PVDF membrane from the transfer apparatus and block nonspecific sites on the membrane by completely immersing the membrane with the blocking membrane (approximately 50 ml) at room temperature (RT) overnight at 4°C.

2. Prepare a 1:5,000 dilution of the anti-\(\beta\)-GlcNAc monoclonal antibody in blocking buffer.

3. Remove blocking buffer and add ~ 30 ml of the diluted antibody to the membrane. Incubate the blot overnight at 4°C. Alternatively, we used 2 µl of primary antibody in 10 ml of blocking buffer in a 50 ml Falcon tube and left the membrane to incubate overnight at 4°C.

4. Wash the membrane six times by completely immersing in wash buffer (approximately 50 ml) and shaking for 5 minutes.

5. Prepare a 10 µg/ml secondary antibody solution by diluting the 0.5 mg/ml stock solution 1:100 in blocking buffer. Prepare approximately 30 ml of a 1:5,000 dilution of the 10 µg/ml goat anti-mouse IgM-HRP in blocking buffer. For 30 ml of blocking buffer we added 12 µl of secondary antibody.

6. Remove wash buffer, add the diluted secondary antibody and incubate for 1 hour at room temperature with gentle shaking.
7. Wash the membrane 6 times by completely immersing the membrane in wash buffer (approximately 50 ml) and shaking for 5 minutes.

8. Prepare SuperSignal® West Dura Working Solution by mixing 5 ml of the Stable Peroxide Solution and 5 ml of the Luminol/Enhancer solution. Apply working solution to the membrane and incubate for 5 minutes at room temperature. Use 0.125 ml substrate solution per cm² of membrane.

9. Remove membrane from working solution and place in a plastic membrane protector. (We used two pieces of acetate).

10. Place the protected membrane in a film cassette with the protein side facing up.

11. The membrane was exposed to Hyperfilm (Amersham, Buckinghamshire, UK) in a dark room for up to 7 seconds and then placed in developer for ± 30 seconds. The film was washed and fixed.
Appendix 5: GFAT Western blotting and detection

Buffers used:

10x Tris Buffer Saline (TBS): (5 litres)

121 g Tris
40 g NaCl
pH solution to 7.6

TBS-T Wash Buffer Add 10 ml of 10x TBS to 90 ml of ultrapure or distilled water and then add 1 ml of Tween-20.

Blocking Buffer Add 5 mg of milk powder (Elite skim milk powder) to 100 ml of TBS-T.

Procedure:

1. Remove PVDF (Millipore, Billerica, USA) membrane from the transfer apparatus and wash the membrane 3 times with TBS-T for 5 minutes.

2. Block nonspecific sites on the membrane by completely immersing the membrane with the blocking membrane (approximately 50 ml) at room temperature (RT) overnight at 4°C.

3. Prepare a 1:3,000 dilution of the GFAT antibody in blocking buffer.
4. Incubate the blot overnight at 4°C. We used 17 µl of primary antibody in 5 ml of blocking buffer in a 50 ml Falcon tube and left the membrane to incubate overnight at 4°C.

5. Wash the membrane six times by completely immersing in TBS-T wash buffer (approximately 50 ml) and agitating for 5 minutes.

6. Prepare approximately 1:2,000 dilution of goat anti-rabbit-horse radish peroxidase in blocking buffer. For 5 ml of blocking buffer we added 5 µl of secondary antibody.

7. Remove wash buffer, add the diluted secondary antibody and incubate for 1 hour at room temperature with gentle shaking.

8. Wash the membrane six times by completely immersing the membrane in TBS-T wash buffer (approximately 50 ml) and shaking for 5 minutes.

9. Wash the membrane for 15 minutes with TBS.

10. Add 0.5 ml of ECL Reagent 1 to the membrane and then add ECL reagent 2 to the membrane. Make sure to spread the ECL reagents over the whole membrane. Incubate for 1 minute at room temperature while continuously washing the ECL reagent mixture over the membrane.
11. Remove membrane from ECL reagent and place in a plastic membrane protector (we used two pieces of acetate).

12. Place the protected membrane in a film cassette with the protein side facing up.

13. The membrane was exposed to Hyperfilm (Amersham, Buckinghamshire, UK) in a dark room for up to ± 7 minutes and then placed in developer for ± 30 seconds. The film was washed and fixed.
Appendix 6: β-Actin Western blotting and detection

Buffers used:

TBS-T Wash Buffer  Add 10 ml of 10x TBS to 90 ml of ultrapure or distilled water and then add 1 ml of Tween-20.

Blocking Buffer  Add 5 mg of milk powder (Elite skim milk powder) to 100 ml of TBS-T.

Procedure:

1. Remove PVDF (Millipore, Billerica, USA) membrane from the transfer apparatus and wash the membrane 3 times with TBS-T for 5 minutes.

2. Block nonspecific sites on the membrane by completely immersing the membrane with the blocking membrane (approximately 50 ml) at room temperature (RT) overnight at 4°C.

3. Prepare a 1:1,000 dilution of the anti-β-Actin monoclonal antibody (Cell Signaling, Danvers, MA, USA) in blocking buffer.

4. Incubate the blot overnight at 4°C. We used 5 µl of primary antibody in 5 ml of blocking buffer in a 50 ml Falcon tube and left the membrane to incubate overnight at 4°C.
5. Wash the membrane five times by completely immersing in TBS-T wash buffer (approximately 50 ml) and agitating for 5 minutes.

6. Prepare approximately 1:2,000 dilution of goat anti-rabbit-horse radish peroxidase in blocking buffer. For 5 ml of blocking buffer we added 5 µl of secondary antibody.

7. Remove wash buffer, add the diluted secondary antibody and incubate for 1 hour at room temperature with gentle shaking.

8. Wash the membrane 3 times by completely immersing the membrane in TBS-T wash buffer (approximately 50 ml) and agitating for 5 minutes.

9. Wash the membrane for 15 minutes with TBS.

10. Add 0.5 ml of ECL reagent 1 to the membrane and then add ECL reagent 2 to the membrane. Make sure to spread the ECL reagents over the whole membrane. Incubate for 1 minute at room temperature while continuously washing the ECL reagent mixture over the membrane.

11. Remove membrane from ECL reagent and place in a plastic membrane protector (we used two pieces of acetate).
12. Place the protected membrane in a film cassette with the protein side facing up.

13. The membrane was exposed to Hyperfilm (Amersham, Buckinghamshire, UK) in a dark room for up to ± 7 minutes and then placed in developer for ± 30 seconds. The film was washed and fixed.
Appendix 7: Stripping protocol

1. Make up the following solution:
   
   100 mM β-Mercaptoethanol: 700 ml/100 ml
   2% SDS: 20 ml/100 ml (from 10% stock)
   62.5 mM Tris-HCl pH 6.7: 0.757 g/100 ml
   
   Make up to 100 ml with H₂O

2. Incubate membrane in solution at 60°C for 30-60 min with occasional agitation.

3. Wash the membrane in TBS-Tween before blocking in 5% milk.