Investigating the Biochemical Basis of Muscle Cell Dysfunction in Chronic Fatigue Syndrome

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April 2016

Thesis submitted for the degree of Doctor of Philosophy in the Institute of Cellular Medicine, Newcastle University
Abstract

Chronic fatigue syndrome/Myalgic Encephalomyelitis (CFS/ME) is a debilitating disorder of unknown aetiology and is characterised by severe disabling fatigue in the absence of an alternative diagnosis. Historically, there has been a tendency to draw psychological explanations for the origin of fatigue. However, this model is at odds with patient descriptions of their fatigue, with many citing difficulty in maintaining muscle activity due to perceived lack of energy and discomfort.

In vivo studies have revealed profound and sustained intracellular acidosis following a standardised exercise protocol, suggestive of underlying bio-energetic abnormality and pointing towards an over-utilisation of the lactate dehydrogenase pathway. Similarly, a recent in vitro pilot investigation reported aberrantly low intracellular pH in CFS/ME patient myoblast samples when compared to healthy controls. Remarkably, intracellular pH in CFS/ME myoblasts was normalised to control level following treatment with pyruvate dehydrogenase kinase (PDK) inhibitor dichloroacetate (DCA), suggesting bio-energetic dysfunction in CFS/ME may be modifiable and therefore treatable.

In this thesis, in vitro approaches were used to investigate possible mechanisms leading to muscle dysfunction and the fatigue phenotype exhibited in CFS/ME. Validation work was performed to assess the capacity of a novel pH responsive nanosensor system to measure intracellular pH in CFS/ME patient myoblast cells. The work was unable to reliably detect any acidosis in CFS/ME cells, or any difference between CFS/ME and control cells. In addition, DCA did not modify intracellular pH in either CFS/ME or control cells.

The fluorescent pH responsive dye 2’7’-bis (2-carboxyethyl)-5 (6) carboxyfluorescein (BCECF) was used to measure intracellular pH at rest, following electrical pulse stimulation (EPS) and after treatment with DCA in myoblast and differentiated myotube cells. Intracellular pH did not differ between CFS/ME patient and control cells at rest or post-EPS. In addition, treatment with DCA did not modify pH in either CFS/ME patient or control cells.

Glycolytic function was assessed via a combination of extracellular flux analysis (XF) and through the measurement of cellular L-lactate concentration. XF analysis revealed extracellular acidification rate (ECAR) measurements for all glycolytic
parameters to be comparable in CFS/ME patient muscle samples when compared to controls. Additionally, DCA did not alter ECAR in either group. L-lactate concentration was elevated at rest of post-EPS in CFS/ME cells compared to controls. DCA did not modify L-lactate concentration in either sample group.

Mitochondrial function was assessed via extracellular flux analysis. Bio-energetic function was investigated by manipulating glucose substrate availability in the assay medium. Basal oxygen consumption rate (OCR) was reduced in CFS/ME myoblasts under hypoglycaemic conditions compared to control cells, however this was not observed in CFS/ME myotubes. ATP-linked OCR was reduced in CFS/ME myoblasts under hyperglycaemic conditions compared to control cells but was not observed in CFS/ME myotube cells. There was no difference between CFS/ME and control cells for any of the other mitochondrial parameters tested.

A direct real-time electrochemical approach was used to monitor superoxide (O$_2^-$) generation in CFS/ME cells following ethanol stimulation and lactic acidification of the assay medium. O$_2^-$ generation was not elevated in CFS/ME cells compared to controls following ethanol stimulation or lactic acidification.

The in vitro muscle culture approaches reported in this thesis have enabled the investigation of the biochemical basis of muscle cell dysfunction in patients with CFS/ME. It is possible to conclude there to be no evidence of impaired muscle function in CFS/ME patients. Additionally, there was no impairment found in PDK enzyme function. Therefore, it can be determined that bioenergetic function is normal in CFS/ME patients and cannot be attributed to the excessive peripheral muscle fatigue phenotype frequently exhibited.
ii. Declaration

The work presented in this thesis was conducted within the Diagnostic and Therapeutic Technologies Department within the Institute of Cellular Medicine, Newcastle University between April 2013 and April 2016. All work reported is original unless acknowledged via reference.

No part of this work has been submitted for a degree, diploma or any other qualification at this University or any other institution.
iii. Acknowledgements

Firstly, I would like to thank my supervisors Professor Julia L Newton and Dr Philip Manning for their continuous support and guidance throughout the PhD process. I also offer my sincere thanks to the charity Action for ME who made this research journey possible.

I am incredibly thankful to Dr Aurora Gomez-Duran who provided technical training and outstanding support which was central to the successful completion of this thesis. Additionally, I would like to thank Dr Audrey Brown, a true expert in muscle cell culture who was always there to share her knowledge and experience.

To my friends Cara, Emily and Hannah thanks for keeping me sane with a cup of coffee and a chat when cells would not grow and experiments did not work, which was far too often. I am grateful to each of you.

To Mum and Dad and my brother Greg thank you for never doubting my capabilities, helping to put things in perspective and always being there with a positive outlook, you deserve a medal. Finally, thank you Lewis for your unwavering support and encouragement, even though at times you probably wanted to strangle me.
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<td>ADP</td>
<td>Adenosine diphosphate</td>
</tr>
<tr>
<td>AMPK</td>
<td>5' AMP-activated protein kinase</td>
</tr>
<tr>
<td>ANS</td>
<td>Autonomic nervous system</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>BCA</td>
<td>Bicinchoninic acid</td>
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<tr>
<td>BCECF-AM</td>
<td>(2',7'-Bis-(2-Carboxylethyl)-5-(and-6)-Carboxyfluorescein, Acetoxyethyl Ester</td>
</tr>
<tr>
<td>BMPO</td>
<td>5-tert- butoxycarbonyl-5 methyl-pyrrole-N-oxide</td>
</tr>
<tr>
<td>CBT</td>
<td>Cognitive behavioural therapy</td>
</tr>
<tr>
<td>CNS</td>
<td>Central nervous system</td>
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<tr>
<td>CFS</td>
<td>Chronic Fatigue Syndrome</td>
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<tr>
<td>CYP2E1</td>
<td>Cytochrome P450 2E1</td>
</tr>
<tr>
<td>DCA</td>
<td>Dichloroacetate</td>
</tr>
<tr>
<td>2-DG</td>
<td>2-deoxy-D-glucose</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s modified eagles medium</td>
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<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
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<td>DMPO</td>
<td>O$_2^-$ 5, 5 dimethyl01-pyrrole-N-oxides</td>
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<tr>
<td>DTSSP</td>
<td>3-3’-dithiobis (sulfosecnimidy1-propionate</td>
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<tr>
<td>ECAR</td>
<td>Extracellular acidification rate</td>
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<tr>
<td>EPS</td>
<td>Electrical pulse stimulation</td>
</tr>
<tr>
<td>ESR</td>
<td>Electron spin resonance spectroscopy</td>
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<tr>
<td>FADH</td>
<td>Flavin adenine dinucleotide phosphate</td>
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<tr>
<td>FCCP</td>
<td>carbonylcyanide p-trifluromethoxyphenylhydrazone</td>
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<tr>
<td>Abbreviation</td>
<td>Description</td>
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<tr>
<td>--------------</td>
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</tr>
<tr>
<td>FI</td>
<td>Fluorescent intensity</td>
</tr>
<tr>
<td>fMRI</td>
<td>functional magnetic resonance spectroscopy</td>
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<tr>
<td>GET</td>
<td>Graded exercise therapy</td>
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<tr>
<td>H₂O₂</td>
<td>Hydrogen peroxide</td>
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<tr>
<td>HF</td>
<td>High frequency</td>
</tr>
<tr>
<td>HR</td>
<td>Heart rate</td>
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<tr>
<td>IL</td>
<td>Interleukin</td>
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<tr>
<td>IOM</td>
<td>Institute of medicine</td>
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<tr>
<td>LDH</td>
<td>Lactate dehydrogenase</td>
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<tr>
<td>LF</td>
<td>Low frequency</td>
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<tr>
<td>MCT</td>
<td>Monocarboxylate transporters</td>
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<td>ME</td>
<td>Myalgic Encephalomyelitis</td>
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<td>MRF</td>
<td>Myogenic regulatory factor</td>
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<td>MRS</td>
<td>Magnetic resonance spectroscopy</td>
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<tr>
<td>MYOG</td>
<td>Myogenin</td>
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<tr>
<td>NADH</td>
<td>Nicotinamide adenine dinucleotide phosphate</td>
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<tr>
<td>NICE</td>
<td>National institute for health and care excellence</td>
</tr>
<tr>
<td>NK</td>
<td>Natural killer cell</td>
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<tr>
<td>O₂⁻</td>
<td>Superoxide</td>
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<tr>
<td>OCR</td>
<td>Oxygen consumption rate</td>
</tr>
<tr>
<td>-OH</td>
<td>hydroxyl radical</td>
</tr>
<tr>
<td>PEBBLE</td>
<td>Probes encapsulated by biologically localised embedding</td>
</tr>
<tr>
<td>PDC</td>
<td>Pyruvate dehydrogenase complex</td>
</tr>
<tr>
<td>PDK</td>
<td>Pyruvate dehydrogenase kinase</td>
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<tr>
<td>Abbreviation</td>
<td>Term</td>
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<tr>
<td>PEM</td>
<td>Post exertional malaise</td>
</tr>
<tr>
<td>PMA</td>
<td>Phorbol-12 myristate-13 acetate</td>
</tr>
<tr>
<td>PMRS</td>
<td>Phosphorus magnetic resonance spectroscopy</td>
</tr>
<tr>
<td>RNS</td>
<td>Reactive nitrogen species</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive oxygen species</td>
</tr>
<tr>
<td>SBP</td>
<td>Systolic blood pressure</td>
</tr>
<tr>
<td>SEID</td>
<td>Systemic Exertional Intolerance Disease</td>
</tr>
<tr>
<td>SOD</td>
<td>Superoxide dismutase</td>
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<tr>
<td>TCA</td>
<td>Tricarboxylic acid</td>
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<tr>
<td>XF</td>
<td>Extracellular flux</td>
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<tr>
<td>XOD</td>
<td>Xanthine oxidase</td>
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Chapter 1
General Introduction


1.1 Definition and History

Chronic Fatigue Syndrome (CFS/ME) also known as Myalgic Encephalomyelitis (ME), is a heterogeneous disorder of unknown aetiology [Bradley et al. 2013]. The condition is characterised by severe disabling fatigue in the absence of alternative diagnosis and is associated with a myriad of other symptoms including but not limited to post exertional malaise, sleep disturbance and cognitive dysfunction [Bradley et al. 2013; Jones et al. 2012; Perrin et al. 2007; Prins et al. 2007; Wessley et al. 1997]. One of the primary symptoms of CFS/ME is generalised abnormal muscle fatigue that occurs after relatively mild activity [Fukuda et al. 1994; Jones et al. 2010; Macintyre et al. 1992]. Fatigue can be defined as a progressive impairment in maximal force generating capacity that develops during muscular activity [Lancet (editorial) 1998].

The prevalence of CFS/ME in the adult population is estimated at around 0.2%-2.6% worldwide [Nacul et al. 2011; NICE. 2007; Prins. 2006; Wessley et al. 1997]. However, the Centre for Disease Control 1994 criteria (CDC4) [Fukuda et al. 1994] and the Canadian criteria [Carruthers et al. 2003] provide a strict definition and estimate prevalence to be 0.2%, lower than previously reported in a primary care setting whereby a less stringent criteria was applied [Wessley et al. 1997]. In the UK, the condition affects approximately 600,000 individuals, with a peak incidence in the 20-40 age group and a preponderance in females, at a ratio of 6:1 [NICE, 2007; Action for ME. 2012]. Moreover, CFS/ME has been reported in children as young as two years of age and would seem to affect all socioeconomic groups to a similar extent [Shepherd. 2006]. Therefore, CFS/ME is a major clinical problem, imposing substantial burden on the health of the UK population in addition to economic costs on society, mainly in the form of informal care and loss of employment. Thus, the development of treatments that recognise these impacts are vital [Perrin et al. 2011].
CFS/ME is historically a complex and poorly understood disorder [Lorusso et al. 2009] and has been the subject of a number of debates. One important challenge has been to define and delineate the illness, which has proven difficult [Hossenbaccus & White. 2013]. Furthermore, controversies surrounding the disorder largely date back to an editorial in the Lancet (1956) that introduced ME into medical language as a severe post-infectious illness involving symptoms that affected peripheral muscle and the brain. However, medical opinion remained sceptical and it was only during the 1980s that interest was renewed, one result being the redefinition and renaming of ME as CFS. In 1998, in an attempt to produce a degree of consensus as regards the diagnosis and management of the disorder, the chief medical officer appointed a working group to produce a report [Department of Health. 2002]. Consequently, CFS/ME was recognised as a genuine and disabling condition in the UK [Shepherd.2006].

Another subject of debate has been in the naming of the condition. For example, the condition was originally referred to as ME, which describes an unproven inflammatory process in the brain and spinal cord, a term that many health practitioners are reluctant to use. Alternatively, CFS is a term that makes no definitive assumptions regarding cause. Two major criticisms regarding the use of this term have been firstly that it fails to reflect symptomology and the severity of the illness. Secondly, it may be a convenient label that could be applied to anyone experiencing unexplained chronic fatigue [Hossenbaccus & White. 2013; Shepherd. 2006]. More recently, early in 2015 the Institute of Medicine (IOM) [IOM, 2015] issued a report that proposed a new case definition for CFS/ME, recommending renaming the illness Systemic Exertion Intolerance Disease (SEID). This new case definition requires a substantial reduction in ability to complete pre-illness activities, unrefreshing sleep, post-exertional malaise and either cognitive or orthostatic intolerance. In a recent study conducted by Jason et al. [2015] the new SEID criteria was reported to identify a group of patients comparable in size to the Fukuda criteria but a larger group than the Canadian criteria. Additionally, the name was reported to select more patients who had less impairment and fewer symptoms than a four item empiric criteria. Presently, there is considerable debate among scientist as regards which case definition to use for clinical and research purposes. As indicated in the IOM report [IOM, 2015] funding was limited so an inadequate number of studies have focused on the validity and reliability of the case definition [Jason et al. 2015]. Thus, further research utilising empirical methods is required to fully evaluate the criteria and develop a consensus among researchers, clinicians and the patient community [Jason et al. 2015].
Additionally, confusion also exists as to whether the nature of the condition is psychological or physiological which is exemplified by the way it is classified, for example the World Health Organisation have classified CFS under the international classification for diseases (ICD-10) as a neurological disease, with the classification suggesting a ‘fatigue syndrome’ should be classified as a neurasthenia in the mental and behavioural disorders chapter [Hossenbaccus & White. 2012; WHO.1992]. While the UK National Institute for Heath and Clinical Excellence emphasise the condition to be a genuine illness with physical symptoms which can be as disabling as multiple sclerosis, systemic lupus erythematosus, rheumatoid arthritis and other chronic conditions [Action for ME. 2012; NICE. 2007].

1.1.1 Clinical features and diagnosis

NICE recommend in their 2007 guidelines that health practitioners should consider the possibility of CFS/ME if the patient has fatigue with all of the following features; New or a specific onset (that is, it is not lifelong), persistent and/or recurrent, unexplained by other conditions, has resulted in a substantial reduction in activity levels, characterised by post-exertional malaise and/or fatigue (typically delayed, for example by at least 24-hours, with slow recovery over several days). In addition to one or more of the following symptoms; difficulty with sleeping (hypersomnia, unrefreshing sleep, a disturbed sleep-wake cycle), headaches, painful lymph nodes without pathological enlargement, sore throat, cognitive dysfunction (inability to concentrate, impaired short-term memory, difficulties with word finding), general malaise or ‘flu-like’ symptoms, nausea and/or dizziness and palpitations in the absence of identified cardiac pathology.

In this PhD thesis muscle samples were obtained from CFS/ME patient donors who met the Fukuda (1994) diagnostic criteria. The Fukuda criteria requires a medical professional to perform a full medical history, physical examination, mental status examination and laboratory tests to rule out other conditions that may require treatment. A patient is then classified as exhibiting CFS/ME by meeting the following criteria. Firstly, unexplained persistent or relapsing chronic fatigue is new or has a definite onset and is not caused by ongoing exertion and is not alleviated by rest, resulting in a substantial decrease in pre-illness activities e.g. occupational, educational, social and personal. Secondly, the individual must also exhibit 4 concurrent symptoms which include the following: A substantial impairment in
short term memory, sore throat, tender lymph nodes, muscle pain, joint pain without redness or swelling, headache that is new in type, severity or pattern, unrefreshing sleep and post-exertional malaise lasting more than 24 hours. To obtain a diagnosis the symptoms must have persisted or recurred during at least 6 months of illness and must not have occurred prior to the onset of chronic fatigue.

Interestingly, CFS/ME exhibits a number of similarities with fibromyalgia which is another condition characterised by severe fatigue and of unknown aetiology. However, to achieve a fibromyalgia diagnosis patients must have experienced widespread pain for up to 3 months to achieve a diagnosis. Although only 18% of fibromyalgia patients have also been diagnosed with CFS/ME it has been reported that around 80% of CFS/ME patients have a diagnosis of fibromyalgia. Demonstrating a clear link between the conditions [Courtois et al. 2015].

1.1.2 Treatment and prognosis

Currently, there is no recognised cure for CFS/ME, however, there are treatments recommended to reduce and manage symptoms [White et al. 2011]. For example, NICE recommend Cognitive Behavioural Therapy (CBT) and Graded Exercise Therapy (GET).

Cognitive Behavioural Therapy (CBT) is a psychological therapy model that is commonly used to treat a range of psychological and chronic pain conditions. CBT facilitates the identification of unhelpful, anxiety provoking thoughts and challenges these negative automatic thoughts and dysfunctional underlying assumptions [Price et al. 2008]. In relation to the treatment of CFS/ME, CBT combines a rehabilitative approach of a graded increase in activity with a psychological approach addressing thoughts and beliefs about CFS/ME that may hinder recovery [Price et al. 2008; White et al. 2011]. Effectively, a gradual increase in activity with a psychological approach addressing thoughts and beliefs, such as the link between increased physical activity and worsened physical symptoms can be tested.

Alternatively, GET is an approach that focuses on the basis of deconditioning and exercise tolerance theories of CFS/ME. Such theories assume that CFS/ME is perpetuated by reversible physiological changes of deconditioning and avoidance of activity. Consequently, these changes lead to deconditioning being maintained, which causes an increased perception of effort and leading to further inactivity. Thus, the aim of GET treatment is to gradually
return the patient to appropriate levels of physical activity, in a manner similar to sports training to reverse deconditioning and reduce associated fatigue and disability [White et al. 2011].

A number of systematic reviews support the use of CBT and GET in the management and improvement of symptoms in patients with mild to moderate CFS/ME. Nevertheless, a lot of this evidence is restricted to small trials [Edmonds et al. 2004; Malouff et al. 2008]. However, in the recent PACE study which is the largest CBT and GET study to date White et al. [2011] reported both CBT and GET to moderately improve CFS/ME outcomes when combined with specialist medical care.

Another systematic review conducted by Cairns et al. [2005] investigated patient prognosis in response to both interventions and no treatment. The authors concluded full recovery from untreated CFS/ME was rare, however symptoms may be improved through CBT and/or GET.

1.1.4 Charity involvement

Action for ME is the UKs leading charity for individuals with CFS/ME and their carers. The charity provides information and support, in addition to being at forefront of research promoting more effective treatment and better services since 1989. Action for ME works in partnership with other organisations in order to transform the lives of people with CFS/ME in the longer term [Action for ME. 2012]. This work was partially funded by Action for ME.

1.1.5 Cellular bio-energetic function overview

The cellular bio-energetic function of CFS/ME patient muscle samples will be discussed at length in this thesis, therefore the following section will provide an overview of the key bio-energetic processes occurring at the cellular level.

Mammalian cells require energy in the form of adenosine-5-triphosphate (ATP) to perform a wide variety of cellular processes. Cells rely upon oxidative phosphorylation and glycolysis to produce ATP [Calderon-Montano et al. 2011]. In terms of the overall contribution of each,
under normal conditions oxidative phosphorylation accounts for around 70% of total ATP yield [Zheng. 2012].

During aerobic glycolysis, pyruvate is able to enter the mitochondria to be oxidised to acetyl-coenzyme-A (Acetyl-CoA), which then combines with oxaloacetate to initiate the tricarboxylic acid cycle (TCA) cycle [Zheng. 2012]. This process enables the production of energy in the form of ATP, as well as the generation of nicotinamide adenine dinucleotide phosphate (NADH) and flavin adenine dinucleotide phosphate (FADH) [Harris et al. 2002]. In terms of oxidative phosphorylation, this oxygen-dependent process couples the oxidation of NADH and FADH generated during the TCA cycle with the phosphorylation of adenosine diphosphate (ADP) to form ATP [Cooper. 2000]. This is achieved through the movement of electrons originating from NADH and FADH2 oxidation through the electron transport chain, which is located in the inner mitochondrial membrane. Effectively, movement of electrons through the chain causes H+ to travel from the mitochondrial matrix into the intermembrane space and in the process producing an electrochemical gradient across the inner membrane mitochondrial space [Cooper. 2000]. The resultant electrochemical gradient promotes H+ movement into the mitochondrial matrix through ATP-synthase, enabling the regeneration of ATP from ADP and inorganic phosphate [Calderon-Montano et al. 2011]. In contrast during anaerobic glycolysis pyruvate is reduced to lactate by the cytoplasmic enzyme lactate dehydrogenase before lactate is excreted into the extracellular space through the monocarboxylate transporters (MCTs), the process producing only 2 ATP molecules [Zheng. 2012].

Pyruvate dehydrogenase complex (PDC) is 3-enzyme complex located in the mitochondrial matrix. It controls the conversion of pyruvate to Acetyl-CoA, functioning as the link step between glycolysis and oxidative metabolism. In terms of structure, PDC is completely nuclear encoded and consists of a variety of copies of 2 structures which are distinct but remain functionally interdependent enzymes (E1- E3) [Smolle et al. 2006]. PDC is a large complex which combines; pyruvate dehydrogenase (E1), dihydrolipoyl transacetylase (E2), E3 binding domain [E3BP] and dihydrolipoyl dehydrogenase (E3). Pyruvate dehydrogenase (E1) is the first enzyme component of PDC [Jha et al. 2012; Harris et al. 1997; Sanderson et al. 1996; Reed et al. 1990].

PDC is tightly regulated by the enzyme pyruvate dehydrogenase kinase (PDK). PDK is a kinase enzyme which operates by inhibiting PDC through phosphorylation with ATP. It is
regulated by the concentration of ATP and acetyl-coA. Four tissue specific isoforms of PDK have been described (1-4), in human skeletal muscle tissue isoforms 1 and 4 have been reported [Nellemann et al. 2013].

Elevated PDK4 expression has been reported in patients in a number of disease states including but not limited to diabetes [Nellemann et al. 2013; Kulkarni et al. 2012], pulmonary hypertension [Piao et al. 2013] and cancer [Sameen et al. 2011]. Interestingly, glycolytic inhibitors are the subject of intense research to explore the therapeutic potential of specific PDK inhibitors to treat diseases characterised by bio-energetic dysfunction [Jha et al. 2012]. Dichloroacetate (DCA) is a small molecule (150 Da) [Michelakis et al. 2008] which functions to activate PDH enzyme function through inhibition of PDK [Stacpoole, 1989], inducing a greater delivery of pyruvate to the mitochondria [Jha et al. 2012]. The molecule occupies the pyruvate-binding site in the N-terminal regulatory (R) domain of the PDK enzyme [Knoechel et al. 2006]. In terms of sensitivity, the isoform PDK2 has been reported as the most sensitive, PDK3 most resistant and PDK1 and PDK4 relatively sensitive [Baker et al. 2000; Bowker-Kinley, 1998]. Furthermore, the inhibitor has been reported to downregulate glycolysis in both in vitro and in vivo studies and has been reported to exhibit a substantial therapeutic benefit in many cancer types [Bonnet et al. 2007; Pan and Mak. 2007].

1.1.6 Bio-energetic muscle dysfunction

Although currently the aetiology of CFS/ME remains elusive, previous research has demonstrated inter-linked changes in muscle bio-energetic function in CFS/ME and has identified key therapeutic targets for the reversal of acidosis. For example, in vivo studies have demonstrated lowering of anaerobic threshold compared to age, sex and BMI matched controls. Phosphorus magnetic resonance (P MRS) approaches have demonstrated CFS/ME patients to exhibit profound and sustained acidosis when undertaken a standard level of activity (35% maximal voluntary contraction) [Jones et al. 2012]. Effectively, P MRS functions by measuring the chemical content of MR visible nuclei, which includes metabolically relevant phosphorus (31P). P MRS is particularly advantageous for assessing metabolism because chemical properties and environment of each nucleus determine the frequency at which it appears on the MR spectrum, giving rise to peaks corresponding not
only to specific metabolites but also the constituent nuclei of each metabolite [Befroy & Shulman. 2011].

Furthermore, pilot work performed by Boulton [2012] revealed aberrantly low intracellular pH in CFS/ME patient derived muscle following the utilisation of a novel pH nanosensor transfection approach, which enabled the fluorescent-based intracellular assessment of muscle cell pH in real time. The findings demonstrated significantly lower pH in CFS/ME derived muscle cells when compared to non-diseased control cells. Furthermore, dichloroacetate (DCA) a drug which up-regulates the function of PDH through inhibition of its inhibitory kinase PDK was reported to normalise pH. This observation suggested that muscle cell acidosis was in part a consequence of down-regulated PDC function and a concomitant increase in the metabolism of pyruvate to lactic acid. Importantly, this also suggested that intracellular acidosis was fully reversible and therefore peripheral muscle fatigue associated with CFS/ME may be treatable.

1.1.7 Myoblasts and differentiated myotubes

In this thesis, primary human muscle cells were used to investigate muscle bio-energetic function in CFS/ME. Muscle cultures were derived from satellite cells, which were isolated via muscle needle biopsy from the vastus lateralis of CFS/ME patients and control participants. The isolated cells first formed mononucleated myoblasts, which were then differentiated into multi-nucleated myotubes, which have been demonstrated to possess all the key characteristics associated with mature native skeletal muscle [Brown et al. 2015].

The fusion of myoblasts is pivotal to enable the formation of relatively mature multinucleated skeletal muscle myotubes. The myogenesis of skeletal muscle is a highly co-ordinated and complex process, which involves a wide range of intracellular signalling molecules [Brand-Saberi, 2005]. In cell culture, myoblast fusion can be induced by reducing the serum concentration of the media, which suggests differentiation is in part controlled by the signalling pathways of growth factors [Kitzman et al. 2001]. Human primary myoblasts and differentiated myotubes are displayed in culture in Figure 1.1.

In order for differentiation to occur cells must first leave the cell cycle and halt proliferation, to enable this requires the down-regulation of cell cycle activators such as cycling and Cdks
and the upregulation of key cell cycle inhibitors such as Rb and P2, which are negatively regulated by growth factors [Kitzmann et al. 2001]. Furthermore, myoblast differentiation is also regulated by the expression of muscle-specific helix-loop-helix transcription factors (myogenic regulators, MRFS) such as MYF-5 and MYOG. These transcription factors function as promoters of muscle specific genes that influence upon the fusion of mononucleated myoblasts into mature myotubes [Ridgeway et al. 2000]. For example, MYOG produces heterodimers with proteins of the E family and binds to the Ebox DNA sequence CANNTG [Tapscott and Weintraub, 1991] and is expressed during the later stages of the differentiation process, with evidence suggesting it is upregulated from day 4 differentiation and peaks at day 16. Alternatively, MYF5 is upregulated during the early differentiation stages and has been reported to be highest at day 1 in mammalian cell culture following treatment with differentiation medium. Additionally, expression has been shown to decrease in a time-dependent manner until day 16 differentiation [Kitzman et al. 2001].

![Figure 1.1: (a) Human primary myoblasts in culture and (b) following differentiation into myotubes](image)

### 1.1.8 Electrical pulse stimulation

Electrical pulse stimulation (EPS) was reported as an *in vitro* exercise model throughout this thesis, enabling the contraction of skeletal muscle cells as a strategy to examine muscle bioenergetic function in CFS/ME muscle samples. A number of studies have reported the use of EPS to induce muscle contraction in skeletal muscle cells. For example, Fujita et al. [2007]
reported EPS to accelerate *de novo* sarcomere assembly, in addition to the induction of $\text{Ca}^{2+}$ transients in C2C12 mouse skeletal muscle myotubes. Additionally, Nedachi et al. [2008] utilising the same experimental model reported EPS to induce activation of 5′ AMP-activated protein kinase (AMPK), increase glucose transport and enhance the release of chemokines such as IL-6. Interestingly, other papers have detailed the use of EPS to investigate human skeletal muscle myotube function [Lambernd et al. 2012; Nikolic et al. 2012], with these studies reporting the model to enhance sarcomere assembly, increase AMPK activation, glycolysis, glucose uptake as well as chemokine expression. More recently, Brown et al. [2015] reported EPS to provide an *in vitro* system to enable the pre-clinical testing of compounds that may influence upon muscle contraction and the metabolic changes associated with exercise. The authors utilised EPS to investigate muscle function in skeletal muscle myotube samples from patients with CFS/ME and controls. The authors reported alternating frequency (low/high) EPS to be successful in enabling muscle contraction and inducing metabolic changes associated with exercise. In healthy cells, EPS induced myotube contraction and increased AMPK activation whereas in the CFS/ME patient group AMPK activation and glucose uptake were impaired. The authors therefore concluded the exercise model to be an effective strategy to investigate metabolic and bio-chemical exercise associated dysfunction in cultured cells.
1.2 Literature Review

Historically, there has been a tendency to draw psychological explanations for the origin of fatigue in CFS/ME, however this model is at odds with patient perceptions of the nature of their condition, with many suggesting a ‘peripheral’ as opposed to ‘central’ basis [Jones et al. 2010]. Interestingly, patients frequently cite difficulty maintaining muscle activity due to perceived ‘lack of energy’ and/or ‘muscle pain’, which can often be serious enough to lead to the avoidance of exercise [Van Oosterwijck et al. 2010]. Thus, it is important to understand peripheral muscle dysfunction in CFS/ME further to inform potential effective treatments for fatigue.

The following sections will explore existing evidence for the role of physiological and biochemical abnormality in the pathophysiology of CFS/ME. Initially, providing an overview of the literature regarding autonomic and immune dysfunction before moving on to more specifically examine the role of peripheral muscle dysfunction as a pivotal cause of fatigue in CFS/ME

1.2.1 Autonomic dysfunction

The compromise of the vascular system and its regulation of the autonomic nervous system (ANS), is a consistent theme that spans throughout existing CFS/ME literature [La mancha et al. 1999; Newton et al. 2009; Rowe et al. 1998]. Effectively, such findings suggest that the abnormal regulation of the ANS plays a pivotal role in the pathogenesis and/or clinical expression of CFS/ME in all or sub-groups of patients [Hagglund et al. 2012; Newton et al. 2009]. The ANS functions as a major regulator of the cardiovascular system, modulating heart rate and blood pressure in the short-term to cope with everyday situations.

Parasympathetic (vagal) modulation functions to decrease heart rate and cardiac contractility, whereas activity of the sympathetic branch opposes these effects and regulates peripheral vasoconstriction. Balanced ANS function is based on strong parasympathetic and efficient but not overactive sympathetic modulation of the heart. Variability of these finely regulated mechanisms may therefore contribute to the expression of fatigue [Newton et al. 2009].

A number of studies have demonstrated the response of the ANS when standing to be abnormal in CFS/ME patients [Hollingsworth et al. 2010; La Manca et al.1999; Rowe et
al.1998]. For example, Hollingsworth et al. [2010] examined haemodynamic responses to immediate and prolonged standing in a large cohort of CFS/ME patients and matched controls. Results demonstrated left ventricular work index on standing to be significantly higher in the CFS/ME group, confirming that the hearts of the CFS/ME group appeared to be working harder in response to the stress of standing in comparison to controls. Similarly, Frith et al. [2012] reported autonomic dysfunction in CFS/ME patients, with results demonstrating systolic blood (SBP) pressure on standing to be significantly decreased with reductions in both sympathetic and parasympathetic components of SBP variability (P<0.0001). These findings were in agreement with previous studies that have demonstrated spectral indices of blood pressure variability (BPV) to be significantly lower in CFS/ME patients when compared to controls [Duprez et al.1998; Yoshiuchi et al. 2004].

In relation to the physiological mechanisms that may account for blood pressure variability, Frith et al. [2012] suggested it to be possible that CFS/ME patients suffer from pathological sympathetic activity over a period of time leading to autonomic effectors (heart and blood vessels) becoming resilient to physiological sympathetic stimulation.

Additionally, previous studies have also reported ANS dysfunction indicated by abnormalities in blood pressure measurements when performed over 24-hours [Newton et al. 2009; Yoshiuchi et al. 2004]. For example, in a study by Newton et al. [2009], CFS/ME and matched controls performed 24-hour ambulatory blood pressure assessment. Results demonstrated significantly lower SBP (P<0.0001), mean arterial blood pressure (P=0.0002), exaggerated diurnal variation (P=0.009) and a significant inverse relationship between increasing fatigue and diurnal variation of blood pressure (P<0.05). Thus, demonstrating lower BP and diurnal variation to occur in patients with CFS/ME. Effectively, the authors proposed three potential mechanisms to account for BP variation. Firstly, BP may be secondary to fatigue, i.e. reflecting a decrease in the amount of physical activity (PA) (reduction in exercise induced BP) performed by patients who view themselves as fatigued. However, an argument against this explanation is that, firstly the effect is principally expressed when physical activity is low and secondly, previous studies have demonstrated PA to only explain one part of BP and HR variability [Cavelaars et al. 2004 and 2002].

Another potential mechanism relates to dysfunction of the hypothalamic pituitary adrenal axis and finally BP deregulation may be causally linked to fatigue expression, for example
low BP may give rise to fatigue through either or both the central nervous system and peripheral hypo perfusion [Newton et al. 2009].

Alternatively, a number of studies have failed to confirm the presence of autonomic dysfunction during rest [Boneva et al. 2007; Wyller et al. 2008; Yamamoto et al. 2003], leading to the suggestion that CFS/ME may be a physiological condition of orthostasis. For example, Wyller et al. [2008] assessed autonomic function in an adolescent CFS/ME patient group during rest and a mild orthostatic challenge. Results demonstrated CFS/ME patients to exhibit greater variability in Heart Rate (HR) and HR variability during an orthostatic challenge in comparison to the control. However, these measures did not significantly differ between the CFS/ME and control group during rest.

Nevertheless, in contrast Wyller et al. [2011] reported abnormal autonomic dysfunction during rest and a mild orthostatic challenge in adolescent CFS/ME patients. HR and blood pressure were recorded continuously and none invasively during supine rest and lower body negative pressure of -20mmHg to stimulate mild orthostatic stress, indices of BP variability and baroreflex (α-gain) were computed from microvariate and bivariate spectra in the low frequency (LF) and high frequency (HF) band. Results demonstrated SBP in the HF range was lower in CFS/ME patients compared to controls as rest and during low body negative pressure (LBNP). Additionally, during LBNP compared to controls α-gain LF/α-gain HF increased more in CFS/ME patients. Thus, the authors concluded all results to be suggestive of a greater shift from parasympathetic to sympathetic baro-reflex control in CFS/ME. Furthermore, the authors postulated an increase in the sympathetic component of the baroreceptor feedback with even mild orthostatic stress in CFS/ME patients to indicate early sympathetic activation and potentially reflect diminished baro-reflex reserve for more severe stressors. These changes further suggest that the baro-reflex may have diminished ability to buffer a variety of internal and external influences on arterial pressure, but particularly those related to upright activity and ambulation. These findings agree with an earlier report regarding the combined effect of orthostatic stress and isometric exercise in CFS/ME patients [Wyller et al. 2008]. However, it is important to acknowledge that the Wyller [2011] study did exhibit crucial methodological limitations. Firstly, respiratory activity has been demonstrated to change during orthostatic challenge and therefore may have influenced cardiovascular variability [Cooke et al.1999] as this factor was not controlled during the
Secondly, blood and/or plasma volume were not measured, therefore hypovolemia could not be ruled out.

1.2.2 Central sensitisation

Central sensitisation is defined as an increase in the responsiveness of central neurons to input from unimodal and polymodal receptors [Meyer et al. 1995]. Central sensitisation involves a number of top-down and bottom-up mechanisms, which contribute to the hyper-responsiveness of the central nervous system to a variety of stimuli [Nijss et al. 2012]. It is important to note that an alteration in central pathways may impact upon peripheral muscle fatigue.

In terms of central impairment, the perception of fatigue during exercise is not always abnormal and serves an important function during significant physical exertion [Jones et al. 2012]. For example, in a study conducted by Amann and Dempsey [2008] peripheral muscle fatigue was induced in patients and consequently fatiguing muscle was reported to play a pivotal role in the determination of central motor drive and force output. Therefore, suggesting the presence of a feedback signal from peripheral muscle to the central nervous system, to ensure fatigue is confined to a certain level, preventing damage to the individual. However, it is plausible that peripheral fatigue experienced in CFS/ME is the direct result of excessive signal feedback, leading to a disproportionate perception of fatigue early in the fatiguing process associated with physical activity [Jones et al. 2010].

Interestingly, evidence exists to suggest the role of central impairment in CFS/ME patients. For example, Whiteside et al. [2004] reported CFS/ME patients to exhibit a dysfunction in nociceptive inhibition during exercise. This was evidenced by a decrease in pain threshold following exercise, which is abnormal as pain threshold usually increases during exercise due to a greater release of endogenous opioids and additional inhibitory mechanisms (descending inhibition). This exercise-induced abnormality was also reported in two additional studies [Meeus et al. 2010; Van Oosterwijck et al. 2010]. For example, Meeus and colleagues [2010] compared CFS/ME patients with chronic pain (n=26), healthy control participants (n=31) and chronic back pain patients (n=21). Participants all completed a submaximal aerobic exercise protocol, which was followed by venous blood sampling (nitric oxide) and algometry (hand,
arm, calf, lower back). Results demonstrated patients with CFS/ME to exhibit lower pain threshold compared to both the healthy control participants and chronic lower-back pain participants. Taken together the results demonstrated CFS/ME patients to have a lack of descending inhibition during exercise. The implication of a lack of endogenous inhibition has been suggested to account in part for the post-exertional malaise (PEM) experienced by CFS/ME patients.

Evidence also exists to suggest a role of generalised hyperalgesia in CFS/ME as outlined by Nijs et al. [2012]. It has been postulated that lower pain thresholds in symptomatic locations represents primary hyperalgesia due to sensitised nociceptors within injured peripheral muscle tissue. However, the authors also commented that when pain thresholds are detected in asymptomatic areas central sensitisation is at play. Two studies conducted by Vecchiet et al. [2003] investigated the effect of electrical stimulation of muscle tissue, skin and sub cutis in relation to pain threshold in CFS/ME patients and healthy control participants. Interestingly, in both studies CFS/ME participants reported there to be no significant difference between groups for electrical pain threshold of skin and sub cutis. Nevertheless, a much lower electrical pain threshold was observed in all sites of muscle tissue (trapezius, quadriceps and deltoid) for the CFS/ME group only, illustrative of hyperalgesia in CFS/ME.

1.2.3 Immune dysfunction

Increasing evidence suggests that CFS/ME is characterised by a profound complex imbalance in immune function [Broderick et al. 2010; Harvey. 2008]. In relation to immunological dysfunction, there is largely no universal agreement as regards the mechanisms due to the varying methodological approaches and study quality. However, evidence suggests natural killer cell function, oxidative stress, altered cytokine profiles/increased pro-inflammatory cytokines and movement towards a Th2 dominant profile to be possible aetiological mechanisms underlying CFS/ME.

In relation to NKC function Maher et al. [2005] observed a significant reduction in NKC associated perforin levels in CFS/ME plasma samples compared to healthy controls moreover the authors also reported reduced perforin content in cytotoxic T cells in CFS/ME subjects. Perforin is a cytolytic protein capable of non-specifically lysing a number of target cells, thus the authors suggested such a deficiency to demonstrate altered immune function in
patients with CFS/ME. Fletcher [2010] also reported diminished NKC function in CFS/ME subjects due to a markedly reduced CD26 density on lymphocyte surfaces and a reduced concentration of the enzyme in the plasma. Collectively, these findings demonstrate a loss of innate immune function and chronic immune stimulation with CFS/ME.

In relation to cytokine abnormalities existing research suggests the pro-inflammatory cytokines it play a pivotal role in CFS/ME [Fletcher.2009; Patarca. 2001; Gupta. 1998]. For example, Robson-Ansley et al [2004] reported recombinant IL-6 (rIL-6) administration to reduce exercise performance in trained endurance athletes. Moreover, the authors speculated these findings to be transferable to patients suffering from CFS/ME, as the condition is associated with reduced exercise tolerance and exaggerated symptoms of fatigue. They suggested an increase in IL-6 to result in impaired neural function; essentially IL-6 may activate the hypothalamic pituitary-adrenal axis by enhancing a natural rise in serum prolactin a marker of neuroendocrine 5-hydroxytrptamine (5-HT) activity. Thus, this rise may stimulate the 5-HT receptors that control prolactin release. Interestingly, the central fatigue hypothesis suggests that 5HT concentrations to relate to impaired CNS function during exercise [Blomstrand. 1995].

In contrast, Robinson et al. [2010] suggested F2 isoprostanes, a marker of oxidative stress to be a key molecular mediator in CFS/ME as no difference in plasma IL-6 or its receptors (sgp-130/SIL-6R) were observed at rest or during exercise in CFS/ME patients. However, F2 isoprostanes were consistently higher at rest, during exercise and 24 hours post exercise. Thus demonstrating a possible role of reactive O2 species in the pathology of CFS/ME, which has been reported by Jammes et al. [2005] and Kennedy [2005] who observed associations between levels of O2 stress and CFS/ME symptoms.

However, in a recent report Broderick et al. [2010] criticised previous studies that had examined the expression and function of individual cytokines in CFS/ME as individual cytokine levels often do not differ between CFS/ME patients and healthy controls. Therefore, the authors examined the co-expression of 16 cytokines in 40 female CFS/ME patients. Results illustrated a diminution of Th1 and Th17 function and a movement towards Th2 type immunity in addition to evidence of increased NK cell function. Similarly, several groups have reported a shift from a Th1 to Th2 cytokine profile in CFS patients. Skowera [2004] for example examined the frequency of type 1 and 2 regulatory CD4 and CD8T cells in 35 patients with CFS/ME. Results illustrated a bias towards a Th2 immune response.
Furthermore, Breu et al. [2011] reported the presence of a possible imbalance in Th1/Th2 response in CFS/ME characterised by significant increase in IL-10, IFNγ- and TNFα in a CFS patient cohort when compared to healthy controls. Such increases in IL-10 are suggestive of a persistent chronic infectious state and may be associated with a dampening of NK cell and CD8+ immune response.

Regarding the relevance of this shift in immune response in relation to CFS/ME, Torres-Harding [2008] reported an increased Th2 response to be associated with a higher reported sleep disruption in patients. Interestingly Th2 cytokines IL-4 and IL-10 have been associated with decreased sleep and an increased Th2 response has been associated with increased cortisol levels, which were reported to increase slow wave sleep and inhibit REM [Buckley & Schatzberg. 2005]

In a study conducted by Bradley et al. [2013] the authors suggested alterations in B-cell maturation to lead to an increased tendency towards autoimmunity and subtle humoral immune dysfunction in a CFS/ME patient cohort. A detailed characterisation of the proportions of different B-cell subsets in 33 patients who met the Fukuda criteria for CFS/ME and 24 age and gender matched controls. Results demonstrated CFS/ME patients to exhibit a greater number of naïve B-cells as a percentage of lymphocytes when compared to controls (6.3% verses 3.9% respectively, P=0.034), a greater number of transitional B-cells, 65% verses 47% in controls (P=0.003), increased numbers of transitional B-cells 1.8% verses 0.8% in controls (P=0.025) and reduced numbers of plasmoblasts (0.5% verses 0.9% respectively (P=0.013).

Remarkably, the finding that CFS/ME patients exhibited increased numbers of transitional B cells may suggest a defective negative selection checkpoint. B cell development in the bone marrow is an antigen independent and tightly regulated process and BCR expression enables negative selection of autoreactive B cells and their subsequent elimination by apoptosis (Keenan et al. 2008). Surviving B cells subsequently become transitional B cells (CD19+, CD38++, IgM+, IDG+) and travel via the blood into secondary lymphoid organs (e.g. spleen) where complete maturation occurs [Verma et al. 2007 ]. Generally, only 10-20% of immature B cells produced in the bone marrow reach the spleen, as a large number are depleted via the negative selection checkpoint because of the expression of defective autoreactive BCRs [Bradley et al. 2013]. Therefore, the authors postulated a defective negative selection checkpoint to lead to a subtle tendency towards autoimmunity in the CFS/ME cohort.
Interestingly, increased numbers of transitional B cells have also been reported in a number of other patient groups that exhibit humoral immunity, including patients with systemic lupus erythematosus [Cuss et al. 2006] X linked lymphoproliferated disease and patients recovering from hematopoietic transplantation [Lee et al. 2009]. Nevertheless, it is important to interpret the findings with caution, as it cannot be ascertained as to whether the changes exhibited are the cause of CFS/ME symptoms or simply the result of patient inactivity, sleep disturbance or raised stress. However, in a recent large placebo controlled clinical trial, symptomatic benefit was shown in 67% of CFS/ME patients after receiving two infusions of rituximab versus 13% of CFS/ME patients receiving placebo [Fluge et al. 2011]. Rituximab has many mechanisms in addition to depleting CD20<sup>+</sup> cells (B cells) down-regulating CD40L and CD80 on B cells, decreasing CD4 effector cells, reducing NK cell no and activation inducing macrophage maturation and reducing tumor necrosis factor (TNF) [Kessel et al. 2008]. Thus, indicating the involvement of B-cells in the pathology or perpetuation of CFS/ME or at least in a subset of patients.

When examining the evidence base, it is clear that CFS/ME is underlined by a complex cascade of immune abnormalities, resulting in a loss of innate immune function and chronic immune stimulation.

1.2.4 Oxidative stress

Reactive species and free radicals are molecules that due to their electronic instability (e.g. unpaired electron) promote oxidation reactions with other molecules such as proteins, lipids and DNA to become stabilised [Gomes et al. 2012]. Many reactive species are O<sub>2</sub> centred (e.g. H<sub>2</sub>O<sub>2</sub>) and are denominated reactive oxygen species (ROS). Some ROS are also free radicals e.g. superoxide anion (O<sub>2</sub><sup>−</sup>) and nitric oxide anion (NO) because they have an unpaired electron [Zadak. 2009]. One important source of ROS/free radical production is believed to be the result of the leakage of electrons in the mitochondrial respiratory chain, due to inadequate coupling of the electron transfer between complexes I and III [Vollard et al. 2005; Xu et al. 2009]. The movement of electrons through the electron transport complex (ETC) and subsequent sites of O<sub>2</sub><sup>−</sup> production are illustrated in Figure 1.2. It has been suggested that for every molecule of ATP generated by the mitochondria, one molecule of superoxide is generated [Jason et al. 2009]. O<sub>2</sub><sup>−</sup> then combines with nitric oxide to form
peroxynitrite which breaks down to release a hydroxyl radical that can induce genetic damage. Peroxynitrite (ONOO⁻) production also leads to an increase in the generation of both nitric oxide and superoxide, which act to produce more peroxynitrite, creating a self-sustaining cycle known as lipid peroxidation [Jason et al. 2009].

Interestingly, enhanced oxidative stress has been reported in CFS/ME patients. For example, studies have demonstrated that CFS/ME patients exhibit excessive production of ROS following physical exertion [Brkic et al. 2010; Jammes et al. 2012] as well as altered resting blood oxidant to antioxidant status [Brkic et al. 2010; Jammes et al. 2012; Maes et al. 2011]. Additionally, bio-chemical markers associated with oxidative stress play a pivotal role in skeletal muscle fatigue [Finsterer et al. 2012], which is often cited as a debilitating symptom experienced by CFS/ME patients [Van Oosterwijck et al. 2010]. Oxidative and nitrosative stress involves the enhanced production of ROS and reactive nitrogen species (RNS), in addition to other free radicals. These reactive species have the potential to disrupt cell membrane function through lipid peroxidation, as well as damage to functional proteins and DNA. This can ultimately lead to alterations in cell structure and disease initiating mutations [Bloomer et al. 2005].

![Diagram showing the complex I and II of mitochondria and the production of reactive oxygen species](image)

**Figure 1.2:** Complex I and II have both been accepted as major sites of electron leakage and $O_2^-$ production. If $O_2^-$ enters the cytosol it may further react with NO$^-$ and redox ions to form additional reactive oxygen species e.g. $H_2O_2$ and ONOO⁻.
Additionally, elevated ROS/RNS exhibit the capacity to profoundly impair mitochondrial function, this has been proposed to be due to the accumulation of oxidative modified mitochondrial proteins, lipids and DNA [Crane et al. 2013]. With the potential of these factors to induce electron transport chain dysfunction, an impairment in cellular bioenergetics and ultimately skeletal muscle fatigue. Furthermore, skeletal muscle is a post-mitotic tissue so is extremely susceptible to mitochondrial oxidative damage, due to being terminally differentiated and because of a slow cellular turn over and high metabolic rate [Crane et al. 2013].

Mitochondria are a major source of ROS generation in cells and are therefore highly susceptible to oxidative damage. For example, they exhibit reduced levels of antioxidants such as glutathione, in comparison to levels found in the cytosol [Fernandez-Checa. 1998]. This relative lack of protection enables mitochondrial DNA (mtDNA) to be damaged, leading to major changes in polypeptide synthesis. Finally, mt DNA repair enzyme activity is considerably lower than found in the nucleus [Wan et al. 2009]. The combination of the effects lead to a reduced electron transfer rate with subsequent reduction in the rate of ATP synthesis [Genova et al. 2004].

In relation to the impact of elevated oxidative stress on muscle dysfunction, muscle specific symptoms of fatigue have been reported to be proportional to the blood levels of a marker of ROS induced lipid peroxidation, named thiobarbituric acid reactive substances (TBARS) [Vecchiet et al. 2003]. TBARS occurs in the serum as a result of lipid peroxidation of low-density lipoproteins and oxygen mediated injury of myocyte membranes [Hulbert et al. 2005]. Lipid peroxidation of skeletal muscle fibres induces a loss of membrane excitability as a result of altered activation of $K^+$ channels [Jammes et al. 2012; Luin et al. 2011]. Additionally, muscle biopsies from healthy adults indicated that the intensity of membrane excitation was proportional to $K^+$ efflux measured in plasma [Marcos et al. 1995], with an increase in ROS generation acting to inhibit $Na^+\cdot K^+$ pump activity, thus reducing the $K^+$ outflow and muscle membrane excitability [Juel et al. 2006]. Furthermore, in another study [Fulle et al. 2003] CFS/ME patients were reported to exhibit a loss of $Na^+\cdot K^+$ and $Ca^{2+}$-ATPase pump regulation, in addition to alterations in the ryanodine channels within the sarcoplasmic reticulum membrane. This was related to increased fluidity of the sarcoplasmic membrane as a result of ROS induced formation of lipid hydroperoxides, which the authors
suggested to support the hypothesis that sarcolemma conduction system and some aspects of Ca^{2+} transport are negatively influenced in CFS/ME.

Furthermore, in another study [Jammes et al. 2012] blood oxidant status and muscle membrane excitability were measured pre and post exercise. The case control study compared CFS/ME patients (n=55) and healthy matched controls (n=40). However, within the CFS/ME cohort sub groups emerged, firstly those who had reported severe infection (e.g. Pneumonia, Septis, encephalomyelitis, H1N1 influenza) within 3-7 months preceding onset of CFS/ME symptoms. Secondly, those who had practiced sport to a high level (>6hrs per week) for 6 years prior to the onset of CFS/ME symptoms and with no history of severe infection. Thirdly, a combination group who had practiced sport to a higher level and had experienced an infection prior to the onset of symptoms. Finally, the last group of CFS/ME patients exhibited no relevant pre-illness history. Participants were required to complete a maximal incremental cycle based protocol to reach the point of maximal oxygen uptake. In relation to markers of oxidative stress, blood samples were obtained at pre, during and post exercise time points, enabling plasma concentration of TBARS and endogenous antioxidant (reduced ascorbic acid) to be measured. Additionally, action potential (M-wave) was evoked in the vastus lateralis to explore muscle excitability.

Results demonstrated that all CFS/ME patients to exhibit abnormal biochemical and electrophysiological measures indicated by elevated TBARS levels prior to exercise and an altered M-wave configuration during and after exercise. These findings are in agreement with other studies [Jammes et al. 2009 and 2005] which also reported an elevated blood oxidant status at rest, which was accentuated by exercise in addition to reduced muscle excitability. For example, one study described enhanced exercise induced oxidative stress, in CFS/ME patients indicated by early changes in TBARS and reduced ascorbic acid in response to incremental exercise. Additionally, CFS/ME patients also exhibited marked alterations in muscle membrane excitability; indicate by lengthened M-wave duration during the recovery period [Jammes et al. 2005].

However, a pivotal finding of the more recent study [Jammes et al. 2012] was that CFS/ME patients who had a history of infection exhibited significant accentuation of blood oxidant status at rest and muscle hypoexcitability at work, in addition to a significantly reduced potassium outflow in response to maximal exercise in this group. Therefore, CFS/ME patients with a history of previous severe infection exhibited greater biological and EMG
disorders in comparison to those who reported completing a high level of physical activity before the onset of fatigue related symptoms. The authors concluded that severe infection could act as a stressor responsible for the alteration in blood oxidant status, which may help to explain impaired exercise-induced K⁺ outflow and altered membrane excitability. However, it is important to acknowledge that the CFS/ME patients self-reported physical activity level for the 6 years prior to the onset of CFS/ME symptoms. Therefore, it is possible CFS/ME patients overestimated their physical activity habits prior to developing CFS/ME symptoms.

1.2.5 Mitochondrial dysfunction

There is evidence to suggest that mitochondrial dysfunction plays a key role in CFS/ME aetiology. Lowered ATP production, impaired oxidative phosphorylation and mitochondrial damage has been reported in patients with CFS/ME [Filler et al. 2014; Myhill et al. 2009]. Moreover, these CFS/ME patients share common skeletal muscle symptoms associated which diseases linked to mitochondrial dysfunction, for example muscle pain, fatigue and cramping [Morris & Maes. 2014; Fulle et al. 2007].

Additionally, there is accumulating evidence to suggest that abnormally high lactate levels and intracellular acidosis exhibited in patients with CFS/ME are the result of impaired mitochondrial function [Morris & Maes. 2014; VanNess et al. 2010]. For example, CFS/ME patients exhibit profound and sustained intracellular acidosis of the peripheral musculature following relatively low-level exercise, resulting in a decreased AT as a result of an overutilization of the lactate dehydrogenase pathway [Jones et al. 2010; Jones et al. 2012]. Upon the point of exhaustion CFS/ME, patients have been found to have intracellular ATP concentrations that are lower than those found in non-diseased controls. This would be indicative of disorders relating to oxidative metabolism. Moreover, in a recent review [Morris & Maes. 2014] the authors concluded the response to exercise in CFS/ME patients to be similar to that typically exhibited in individuals with mitochondrial disease. Additionally, there were also a number of similarities between symptoms of mitochondrial disease and the physio-somatic symptoms of CFS/ME. For example, muscle pain, cramps, weakness and myalgia’s [Morris & Maes. 2014; Fulle et al. 2007; Nijs et al. 2004].
Mitochondrial dysfunction in CFS/ME may be explained not only by elevated oxidative and nitrosative stress but also increased immune-inflammatory stress pathways [Morris & Maes. 2012]. For example, chronic low grade inflammation in CFS/ME has been demonstrated through increased levels of pro-inflammatory cytokines (IL-1, TNFα) and a movement towards a Th2 dependant immune response, in addition to inflammatory mediators including nuclear factor-κB (NF-κB) and elastase [Nijs et al. 2004]. Further, evidence of elevated oxidative and nitrosative stress is demonstrated by increased isoprostane levels, peroxides and protein carbonyl levels, indicating damage to lipids and mitochondrial protein as previously discussed [Jammes et al. 2012].

In relation to immune dysfunction a study was conducted to examine cytokine networks in 40 female CFS/ME patients and 40-case matched controls [Broderick et al. 2010]. The authors examined a total of 16 cytokines and results revealed a diminution of Th1 and Th17 function and a movement towards Th2 type immunity. Similarly, several other groups have reported a shift for a Th1 to Th2 cytokine profile in CFS/ME patients [Brenu et al. 2011; Fletcher et al. 2009. Skowera et al. 2004]. For example, Skowera et al [2004] examined the frequency of type 1 and 2 regulator CD4 and CD8T cells in 35 patients with CFS. Results illustrated a bias towards a Th2 immune response.

Elevated levels of the inflammatory mediator NF-κB [Maes et al. 2011] have also been reported in the blood samples of CFS/ME patients. NF-κB is a major upstream intracellular mechanism, which regulates inflammatory and oxidative stress mediators [Maes et al. 2007]. For example, it functions to trigger inducible nitric oxide synthetase (iNOS) expression, which promotes the production of nitric oxide (NO) by monocytes and macrophages [Paludan. 1998].

Maes and colleagues [2007] examined the production of NF-κB P50 in unstimulated 10ng/mL TNF-α and 50ng/mL phorbol 12-myristate 13-acetate (PMA) stimulated peripheral blood lymphocytes in 18 CFS/ME patients and 18 age-matched controls. Results demonstrated both unstimulated (10ng/mL) TNF-α (P=0.0009) and PMA (0.008) stimulated production of NF-κB to be significantly higher in CFS/ME patients compared to controls. Additionally, positive correlations were reported between the production of NF-κB and severity of illness in CFS/ME patients (as measured by fibro fatigue scale) and with symptoms, including muscular fatigue and tension. Interestingly, NF-κB signals the function of p53 is pivotal in the regulation of glycolysis and mitochondrial respiration as it reduces the
activity of the glycolytic pathway and stimulates mitochondrial O$_2$ consumption and aerobic respiration. However, when inhibited there is a shift to anaerobic glycolysis and reduced O$_2$ consumption [Morris and Maes. 2012].

Aside from immune dysfunction, several studies have reported mitochondrial dysfunction to be caused by abnormal levels of key mitochondrial enzymes. For example, in a paper by Smit and colleagues [2011] a significant reduction in citrate synthase in quadriceps biopsies from patients with CFS/ME was reported when compared to healthy control samples. Citrate synthase is an enzyme located in the mitochondrial matrix, which plays a critical role in the tricarboxylic cycle [Tymoczko. 2010]. Similarly, the McArdle group [1996] reported a decrease in citrate synthase in addition to succinate reductase and cytochrome-C oxidase (Complex IV), which are two of the four mitochondrial transmembrane enzyme complexes of the electron transport chain. However, in contrast to the McArdle group [1996], Smit and colleagues [2011] attributed the decrease in transmembrane enzymes to be the result of reduced physical activity levels frequently present in CFS/ME patients, as opposed to underlying mitochondrial dysfunction. Nevertheless, a paper by Edwards and colleagues [1993] reported there to be no significant difference in partial cytochrome-C oxidase in skeletal muscle biopsies between CFS/ME patients and healthy matched controls.

Evidence also exists to suggest that CFS/ME patients exhibit significantly reduced levels of Co-enzyme Q10, an important mitochondrial nutrient that functions as a co-factor for the production of ATP in the mitochondria and displays significant antioxidant activity [Jones et al. 2009]. In a study conducted by Maes and colleagues [2009], CFS/ME patients (n=58) displayed significantly lowered plasma co-enzyme Q10 concentration compared to healthy controls (n=22). Moreover, in CFS/ME patients there was a significant inverse relationship exhibited between plasma co-enzyme Q10 concentration and fatigue severity measured by means of the fibro fatigue scale.

Furthermore, CFS/ME patients may also exhibit alterations in L-carnitine and acyl carnitine homeostasis [Reuter et al. 2011]. L-carnitine is a ubiquitously occurring trimethylated amino acid that plays an important role in the transport of long chain fatty acids across the inner mitochondrial membrane, which is essential for energy production via fatty acid metabolism [Reuter et al. 2009]. Previous studies have reported a reduction in endogenous plasma L-carnitine and total carnitine levels in patients with CFS/ME [Karatsune et al.1998 and 1994]. Nevertheless, other studies have not always replicated these findings [Jones et al. 2005;
Majeed et al. 1995. Reuter and colleagues [2011] postulated this to be related to the use of varying methodological approaches, with some studies solely focused on free carnitine and total carnitine rather than the level of each individual acyl carnitine, which may be ‘cancelled out’ by normal levels of other acyl carnitines in CFS/ME patients. To overcome this weakness Reuter and colleagues [2011] utilised tandem mass spectrometry to quantify individual acyl carnitine levels in plasma samples to provide a more detailed carnitine profile. Results demonstrated significant alterations in C8:1, C12DC, C14, C16:1, C18, C18:1, C18:2 and C18:1-Oh acyl carnitines. What is more, significant correlations between acyl carnitine and clinical symptomology were observed.

1.2.6 Post-exertional malaise and immune function

CFS/ME patients report a changeable pattern to their symptoms and physical and cognitive capabilities, often with severe symptom exacerbation following physical exercise [Fukuda et al. 1994; Whiteside et al. 2004]. This is termed PEM, with approximately 95% of CFS/ME patients experiencing PEM [Prins et al. 2007]. As regards the cause of PEM, it has been suggested that exercise may exhibit the ability to amplify pre-existing immune abnormalities, in addition to oxidative and nitrosative stress [Twisk. 2015]. Immunological abnormalities have been reported following exercise in CFS/ME. For example, observations in CFS/ME symptom flare after moderate intensity exercise have been reported to be directly linked to the levels of Interleukin 1β (IL-1β), IL-12, IL-8, IL-10 and IL-13, 8 hours post-exercise [White et al. 2012]. Additionally, sustained increase in plasma TNFα in CFS/ME patients and not in healthy controls has been observed post-exercise [White et al. 2012]. Moderate intensity exercise has also been reported to induce a larger 48-hour post exercise area under the curve for IL-10 [Light et al. 2012].

However, a recent systematic review [Nijs et al. 2014] compared 23 case control studies regarding exercise-induced immunological changes in CFS/ME patients verses healthy control participants. The authors reported in comparison to healthy participants, CFS/ME patients exhibited a more exaggerated response in the complement system, indicated by C4a split product level, enhanced oxidative stress, combined with a delayed and reduced antioxidant response. Finally, the authors also reported there to be an apparent alteration in immune cell gene expression profile, which was evidenced by an increase in post-exercise...
IL-10 and toll like receptor 4 gene expression. Nonetheless, in contrast to previous work there was no reported change in circulating pro and/or anti-inflammatory cytokines. Effectively, the review confirmed CFS/ME patients to respond differently to an exercise-based stimulus, resulting in a more pronounced immune response.

1.2.7 Muscle bio-energetic dysfunction

1.2.7.1 Intracellular acidosis

Even a minimal decrease in muscle pH interferes with cross bridge binding and ATPase activity due to competitive binding and reduced enzyme function [Jones et al. 2009]. Decreased intracellular pH impairs oxidative enzyme activity and may adversely affect ryanodine receptor function [Bellinger et al. 2008]. Furthermore, recent studies confirm the presence of a peripheral bio-energetic abnormality in CFS/ME patients [Jones et al. 2010; VanNess et al. 2010; Jones et al. 2012].

In a cross-sectional study conducted by Jones et al [2010] novel P MRS techniques were utilised to investigate muscle acid handling following exercise in CFS/ME patients and the relationship with autonomic dysfunction. CFS/ME patients (n=16) and age and sex matched normal controls (n=8) performed an exercise protocol, which consisted of 3 minutes of plantar flexion at 35% load maximum voluntary contraction (MVC) at a rate of 0.5Hz, followed by 3 minutes recovery. After the period of exercise P MRS was utilised to investigate intramuscular acid handling. Results demonstrated, a significant suppression of proton efflux immediately post-exercise (P<0.05) in CFS/ME patients, in addition to a significantly (P<0.05) prolonged time taken to reach maximum proton efflux. In controls there was a strong inverse correlation between maximum proton efflux and nadir pH following exercise ($r^2=0.6$, P<0.01). However, in CFS/ME patients the significance of this relationship was lost ($r^2=0.003; P=ns$). Collectively, these findings demonstrated CFS/ME patients to exhibit abnormalities in the recovery of intramuscular pH following standardised exercise. Effectively, proton efflux is crucial for acidosis resolution, with the immediate post-exercise period associated with maximum proton efflux in healthy individuals; however this initial fast phase does not occur in CFS/ME patients [Jones et al. 2010]. Furthermore, the authors also acknowledged there to be a close relationship between the degree of intramuscular acidosis and proton efflux, demonstrating a closely regulated process, which
has been observed in healthy individuals [Kemp et al. 1997] in addition to the study control group. Nevertheless, they concluded this relationship to be lost in CFS/ME patients. However, the relatively small sample size of this study made it difficult to draw firm conclusions. Therefore, additional adequately powered studies are required to investigate the relationship further.

In contrast, Wong et al. [1992] reported no difference in intramuscular pH at rest, exhaustion and during early and late recovery, following a graded exercise test to exhaustion. Measurement of intramuscular pH of the gastrocnemius muscle was performed via 31P nuclear magnetic resonance (NMR) spectroscopy. However, the authors did report changes in PCr and pH to occur more rapidly at the onset of exercise in CFS/ME patients compared to controls, which was suggested to be indicative of accelerated glycolysis. Nevertheless, the authors postulated this finding to reflect a lower level of physical endurance due to inactivity in the CFS/ME patient cohort. Nonetheless, it is important to interpret this study with caution, as the CFS/ME patients were able to complete a maximal exercise test to exhaustion. Thus, suggesting the patients in this study were not severely physically incapacitated by the condition and therefore may not be representative of the wider CFS/ME population.

Alternatively, Jones et al. [2012] reported prolonged post-exercise recovery from acidosis. In this investigation CFS/ME, patients and age/sex matched healthy controls performed a similar exercise protocol (35% MVC plantar flexion for 180s, 390s recovery, repeated 3 times). In addition, participants were also required to perform a MVC assessment and a cycle based cardio-respiratory fitness test. Results revealed the ability to divide patients into two distinct groups; 8 (45%) demonstrated normal phosphocreatine (PCr) depletion in response to exercise at 35% MVC, with MVC strength values comparable to controls. In the second grouping, 10 CFS/ME patients exhibited low PCr depletion (generating abnormally low MVC values). Results demonstrated anaerobic threshold (AT), VO₂ and VO₂ peak to be significantly reduced in all CFS/ME patients compared to controls. Essentially, one implication of a reduced AT would be a reliance upon anaerobic as opposed to aerobic metabolism, with the predicted consequence of greater short-term acid generation within the muscle as a result of an over-utilisation of the lactate dehydrogenase pathway [Jones et al. 2010]. This was further confirmed by MRS demonstrating CFS/ME patients to exhibit markedly increased intramuscular acidosis compared to controls at a similar work rate.
following each 3-minute bout, with prolongation (almost 4-fold) in the time taken for pH to recover to baseline, replicating previous findings [Jones et al. 2010].

Based on the findings the authors went on to conclude that the profound intramuscular acidosis exhibited with repeat exercise was at least in part related to poor aerobic capacity. This in relation to the physiology of fatigue closely mirrors that observed in patients with the autoimmune disease primary biliary cirrhosis (PBC). PBC exhibits a comparable peripheral pattern and a similar level to fatigue to CFS/ME [Hollingsworth et al. 2010]. In a study by Hollingsworth and colleagues [2010], PBC patients exhibited profound and comparable intramuscular acidosis to the CFS/ME patients in the Jones et al. [2012] study following the same repeat exercise protocol. However, one pivotal difference between the conditions, which may contribute to the severity of fatigue in CFS/ME, is related to acid homeostasis. In contrast to CFS/ME patients, when PBC patients undergo repeat exercise the extent of acidosis within the muscle decreases with each repeated exercise bout. This may suggest a compensatory mechanism, which operates to resolve excess acidosis. One potential mechanism that may account for this is increased proton flux, in addition to the speed of onset of maximum proton excretion with repeat exercise [Hollingsworth et al. 2010; Jones et al. 2012]. This mechanism also plays a role in mitochondrial disease whereby increased proton efflux post-exercise helps to compensate for decreased aerobic capacity [Trenell. 2006]. Nevertheless, it would seem that in comparison to other conditions that exhibit reduced aerobic capacity and acidosis, CFS/ME patients are unable to compensate for an increased reliance upon anaerobic energy pathways during exercise [Jones et al. 2012].

While the production of protons as a by-product of anaerobic metabolism is a feature of normal metabolism, the body requires mechanisms to effectively manage protons as even small changes in pH dramatically alter enzyme kinetics, decrease muscle function and cause fatigue [Allen et al. 1995]. Thus, slow recovery from acidosis in CFS/ME may relate to ineffective exporting of protons from the recovering muscle. Protons are actively transported out of the muscle by 3 main groups of proton transporters [Juel et al. 2006]; Na+/H+ anti-porters, namely NHE1 [Street et al. 2001] sodium/bicarbonate co-transporters (NBCs) [Kristensen et al. 2004] and most predominantly monocarboxylate transporters (MCT), whereby in the latter group MCT-1 and MCT-4 isoforms seem to be of particular importance in human skeletal muscle [Wilson et al. 1998]. During rest, intramuscular pH is predominantly influenced by NHEs, with MCTs and NBCs playing a greater role during the
recovery from muscular contraction. These transporter systems are under autonomic regulation [Halenstrep & Prince. 1998]. It is therefore possible that the impaired function of acid transporters occurs in CFS/ME, which may be a consequence of autonomic dysfunction as previously discussed an abnormality found frequently in CFS/ME [ De Becker et al. 2000; Newton et al. 2007; Anderson et al. 2008]. Additionally, it is also possible that reduced vascular run off, (related to autonomic dysfunction) resulting in decreased vascular flow into and out of the muscle following exercise, which may have an effect on \( \text{O}_2 \) delivery, potentially limiting the function of the three enzyme complex pyrate dehydrogenase complex (PDC) [Anderson et al. 2008].

A sub group of patients (low PCr depletion) exhibited no excess acidosis, which appeared to be entirely the consequence of lower MVC values compared to normal PCr depletion controls. Interestingly, despite markedly lower MVC values the patients perceived themselves to be working maximally immediately following the MVC assessment. However, despite this perception the authors postulated that these findings related to a type of exercise avoidance behaviour. Kinesiophobia is defined as an excessive, irrational and debilitating fear of movement and activity resulting from a fear of vulnerability to painful injury or re-injury [Kori et al. 1990; Nijs et al. 2004] and has been reported to play a role in a variety of musculoskeletal disorders, including CFS/ME patients who experience widespread pain [Silver et al. 2002; Vlaeyen et al. 1995]. Therefore, the fear of the consequence of an action such as exercise may lead to avoidance behaviour in patients with CFS/ME patients [Jones et al. 2012] However, the study was limited, as it did not include a repeat assessment. Therefore, it is impossible to ascertain whether the groups were stable i.e. avoiding the first exercise session and consistently doing so.

Interestingly, previous studies have also reported the existence of sub-groups within CFS/ME patient cohorts in relation to glycolytic metabolism and intramuscular pH regulation [Lane et al. 1998a; Wong et al. 1997; Barnes et al. 1993]. However, unlike the studies conducted by Jones et al [2012 and 2010], the earlier studies have reported an absence in abnormal glycolytic metabolism in the majority of the CFS/ME patients. For example, Barnes et al [1993] explored intramuscular pH regulation in 46 CFS/ME patients via 31P MRS. Results demonstrated no consistent abnormalities in glycolysis or pH regulation at rest or following exercise when the group was taken as a whole. Nonetheless, 12 patients did exhibit abnormal PCr depletion following exercise, with 6 patients within this group displaying increased
intramuscular acidification in relation to PCr depletion and the other 6 demonstrating reduced acidification. This study illustrates the heterogeneity within the CFS/ME patient population and suggests sub-groups do exist in CFS/ME that display abnormal glycolytic metabolism and intramuscular pH. Similarly, in Lane’s studies [1998a and 1998b] CFS/ME patients completed a sub anaerobic threshold exercise protocol. Results revealed only a small sub-group (8%) of CFS/ME patients to have an increased blood lactate response to exercise and muscle biopsies revealed a relative increase in type 2 glycolytic fibres for this sub-group.

1.2.7.2 Acidosis as a consequence of impaired PDC function

It is possible that the previously demonstrated muscle cell acidosis in CFS/ME is the consequence of down-regulated PDC function and a concomitant increase in the metabolism of pyruvate to lactic acid (over-utilisation of lactate dehydrogenase (LDH) pathway). PDC is a 3 protein complex responsible for a series of reactions that convert pyruvate to acetyl coA during aerobic respiration. Principally, when the function of this complex is reduced, pyruvate that has been generated by glycolysis accumulates within the cells and is metabolised anaerobically to lactic acid. This accumulation causes a drop in pH and concurrent deterioration in muscle function [Forque et al. 2003].

The phenotype of fatigue exhibited by CFS/ME patients closely mirrors that seen in fatigue associated PBC. For example, Hollingsworth et al [2010] reported that PBC patients exhibited significant acidosis because of an over-utilisation of the lactate dehydrogenase pathway, following a low-level repeat exercise protocol [Hollingsworth et al. 2010]. Furthermore, the authors postulated that the increased dependence on anaerobic pathways of energy production resulted in the fatigue associated with PDC.

The idea that impaired PDC function leads to an over-utilisation of the lactate-dehydrogenase pathway is in agreement with other studies. For example, Murrough et al [2009] reported significantly higher levels of lactate in ventricular cerebrospinal fluid in CFS/ME patients in when compared to healthy controls. Similarly, Constantin-Teodosiu and colleagues [2009] in an experimental exercise model using rats, demonstrated that when PDC function was decreased via the use of PPAR modulating drugs which up-regulated PDK function, lactate accumulated intramuscularly which led to decreased muscle function.

Therefore, impaired energy generation in muscle, an increase in the lactate/pyruvate ration in CFS/ME patients and a propensity towards excess intra-muscular acidosis following limited
exercise suggests PDC dysfunction in the muscles of CFS/ME patients, which has implications in relation to the expression of fatigue. Furthermore, as previously discussed CFS/ME patients exhibit significant intramuscular abnormalities relating to both acid generation and clearance from tissue, which has been postulated to relate to a centrally-perceived “stop signal”, leading to a disproportionate perception of fatigue [Jones et al. 2009]. Figure 1.3 illustrates the effect of PDC downregulation, leading to an over-utilisation of the lactate dehydrogenase pathway.

Figure 1.3: Diagram outlining PDC function when down-regulated, leading to an over utilisation of the lactate dehydrogenase pathway.

1.2.8 Abnormal AMPK activation and glucose uptake

A recent study [Brown et al. 2015] reported striking biochemical differences in skeletal muscle cultures established from 10 CFS/ME patients and 7 age-matched controls. Samples were subjected to EPS for 24-hours and examined for exercise associated changes. Key differences emerged, in the basal state there was increased myogenin expression in CFS/ME samples but a decrease in IL-6 secretion during differentiation when compared to control samples. Following 16 hours EPS there was a significant increase (P<0.006) in AMP-activated protein kinase (AMPK) phosphorylation and glucose uptake (P<0.0001) in control samples when compared to unstimulated control cultures. Alternatively, CFS/ME samples demonstrated no increase in AMPK phosphorylation or glucose uptake. Nevertheless, glucose
uptake remained responsive to insulin, suggesting exercise related dysfunction. Furthermore, IL-6 secretion on response to EPS was significantly reduced (P<0.05 vs corresponding control) across all time points measured.

AMPK is a phlogenically-conserved fuel-sensing enzyme, consisting of a heterotrimeric complex composed of a catalytic α subunit and regulatory β and γ subunits [Richter et al. 2009]. During exercise under normal physiological conditions, AMPK is activated in the skeletal muscle of healthy humans, with exercise suggested to be the most powerful physiological activator of AMPK [Chen et al. 2003; Winder et al. 2000]. Upon activation AMPK sets into motion processes that increase ATP production such as glucose transport and fatty acid oxidation [Richter et al. 2009], while decreasing others that consume ATP, for example lipid and protein synthesis and cell growth and proliferation [Hardie et al. 2007; Kahn et al. 2005; Richter et al. 2009] Additionally, evidence also suggests AMPK to have a broader range of actions including mitochondrial bio-genesis [Winder et al. 2000; Jorgensen. 2005] and skeletal muscle angiogenesis [Ouchi et al. 2005]. Suggesting AMPK activation to play a key role in peripheral muscle function during exercise.

However, it is important to consider the role of physical activity on AMPK activation. For example, trained subjects have been reported to express higher levels of α1 AMPK in comparison to untrained individuals [Nielsen. 2002]. Furthermore, a 3-week endurance training intervention with young male participants resulted in increases in α1 and α2 AMPK protein expression in addition to ACC-β phosphorylation, which suggested basal activity of AMPK to be, increased [Frosig et al. 2004]. Therefore, it should be considered that the decreased AMPK activation reported in CFS/ME muscle samples after EPS may be the result of lowered physical activity levels of the CFS/ME sample donors, when compared to control donors. As regards study recruitment criteria, although participants were age matched it was not specified whether any measures had been taken to ensure donors were matched in terms of physical activity habits. Future work is required with patients and controls who are matched in terms of physical activity.

In addition to impaired activation of AMPK, the study also reported reduced IL-6 secretion in response to EPS. Interestingly, previous studies have reported IL-6 to activate AMPK in skeletal muscle by increasing the concentration of cAMP and secondly by increasing the AMP:ATP ratio [Kelly. 2009].
The inability of CFS/ME muscle cells to activate AMPK and glucose uptake in addition to reduced IL-6 secretion in response to EPS is suggestive of underlying peripheral muscle dysfunction in CFS/ME. However, further work is required to investigate the mechanisms that lead to impaired activation of AMPK in those with CFS/ME.

1.3 Conclusion

There is increasing evidence to suggest muscular bio-chemical abnormality to play a major role in CFS/ME associated fatigue. Patients have been reported to exhibit profound intramuscular dysfunction regarding acid generation and clearance, with a tendency towards an over-utilisation of the lactate dehydrogenase pathway following relatively low-level activity. However, the precise mechanisms underlying the dysfunction are yet to be fully elucidated but may relate to impaired function of PDC. Future work is required to examine PDC function in vivo, to determine the mechanisms responsible for muscle cell acidosis in and explore the capacity of drugs to normalise bio-energetic function and ultimately treat peripheral fatigue.

1.4 Aims

The aim of this thesis is to improve understanding of the mechanisms underlying peripheral muscle dysfunction and the associated perception of fatigue in patients with CFS/ME through the development of an in vitro muscle cell culture testing platform.

Preliminary work has demonstrated an aberrantly low intracellular pH in CFS/ME patients, which was subsequently normalised following treatment with PDK inhibitor DCA. Which was revealed when pH was measured using a novel pH sensing platform [Boulton. 2012]. Therefore, a key aim of this thesis is to validate and further develop this in vitro model as a pre-clinical testing system, to firstly determine the presence of acidosis in CFS/ME and secondly the extent in which pH is modulated by bio-energetic enzyme inhibition. The identification of therapeutic targets within the biological muscle system are warranted to enable the development of appropriate therapies to inform clinical trials in CFS/ME.

Specific chapter aims are stated below.
Chapter 2 will further validate and develop the fluorescent pH responsive nanosensor previously reported by Boulton [2012]. Additionally, PDK inhibitor DCA was added to the cells to assess the capacity of drugs to normalise intracellular acidosis *ex vivo*.

Chapter 3 will investigate O$_2$- generation in CFS/ME patient myoblasts *ex vivo*, performed following chemical stimulation with ethanol or incubation with lactic acid. Specific experimental hypotheses are (1) CFS/ME myoblasts exhibit elevated O$_2$- generation following ethanol stimulation, compared to controls. (2) O$_2$- elevated in CFS/ME patient samples compared to controls following incubation with lactic acid.

Chapter 4 will measure cytosolic pH via the application of the widely used free dye 2′7′-bis (2-carboxyethyl)-5 (6) carboxyfluorescein (BCECF) in CFS/ME myoblasts and differentiated myotubes at rest and following electrical pulse stimulation (EPS), performed *ex vivo* experimental work performed. The specific experimental hypothesis are (1) CFS/ME myoblasts and differentiated myotubes exhibit significantly lower intracellular pH at rest and following EPS, when compared to controls. (2) Treatment with PDK inhibitor DCA normalises intracellular pH in CFS/ME samples at rest and following EPS.

Chapter 5 will measure glycolytic function in CFS/ME patient myoblast and myotube samples via extracellular flux analysis, in addition to extracellular lactate quantification performed *ex vivo*. Which will be performed using extracellular flux glycolytic stress testing and L-lactate measurements via fluorometric assay. The specific experimental hypotheses proposed are (1) Increased glycolytic activity in CFS/ME patient myoblasts and myotubes when compared to controls. (2) Normalisation of glycolytic function comparable to controls following treatment with PDK inhibitor DCA.

Chapter 6 will investigate CFS/ME myoblast and myotube mitochondrial function *ex vivo* via XF analysis. Function will be assessed following the addition of key inhibitors of the mitochondrial electron transport chain. Further investigation into CFS/ME myoblast and myotube function will be performed by varying substrate availability (glucose) during the mitochondrial stress test. Specific experimental hypotheses are (1) CFS/ME myoblasts and myotubes exhibit impaired mitochondrial function. (2) CFS/ME samples exhibit a greater impairment in mitochondrial function compared to controls when substrate availability is reduced.

The conclusions of each chapter are summarised in Chapter 7.
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Chapter 2
Application of a Novel pH Sensing Nanosensor Platform to Measure Intracellular pH in CFS/ME Primary Myoblasts

2.1 Introduction

Previous research has demonstrated that CFS/ME patients exhibit profound and sustained intracellular acidosis with repeat exercise, compared to controls [Jones et al. 2012]. In addition to abnormality in the recovery of intramuscular pH following exercise [Jones et al. 2010], the authors suggested that these findings result from an over-utilisation of the lactate dehydrogenase energy-producing pathway, pointing towards a bio-energetic abnormality, which may offer the potential of therapeutic intervention. Furthermore, Boulton [2012] utilised a novel in vitro drug pre-testing platform to measure intracellular pH in CFS/ME muscle, to determine whether the PDK enzyme inhibitor DCA had the potential to modify intracellular pH. Results demonstrated that CFS/ME patient myoblasts exhibit an aberrantly low intracellular pH when compared to controls. Remarkably, this effect was completely normalised following treatment with PDK inhibitor DCA.

Briefly, PDK functions to inhibit PDC a 3-enzyme complex which is responsible for the conversion of pyruvate to acetyl-coA during aerobic respiration. However, when PDC functionality is decreased an increase in pyruvate generated by glycolysis accumulates in the cell and is metabolised to lactic acid, which due to ineffective oxidative phosphorylation can cause a decrease in pH and a concurrent reduction in muscle function. The compound DCA is a pyruvate analogue and functions to inhibit PDK, an enzyme which acts to inhibit PDC function therefore downregulating aerobic metabolism. DCA exhibits the capacity to reverse intracellular acidosis by inducing the active state of PDC and promoting the clearance of lactic acid by functioning to convert it back to pyruvate aerobically [Boulton, 2012; Forque, 2003].

The primary aims of this chapter were [1] to validate and develop the in vitro muscle pH sensing system reported by Boulton [2012] and further confirm the presence of abnormal pH in CFS/ME myoblasts. [2] To apply PDK inhibitors into the system in order to investigate the ability of drugs to modify intracellular acidosis.
2.1.1 Intracellular pH and Cell Function

Intracellular pH plays a pivotal role in nearly all aspects of cell function, including regulation of cell volume, vesicle trafficking, cellular metabolism, cell membrane polarity, growth and proliferation [Dennis et al. 2012; Loiselle et al. 2003; Busa et al. 1984]. The disruption of intracellular pH is associated with serious physiological consequences and an alteration in pH of as little as 0.1-0.2 pH units can induce metabolic depression [Gibbin et al. 2014]. Eukaryotic cells therefore exhibit mechanisms to prevent any significant fluctuations in pH, for example acute alterations as a consequence of metabolic reactions are neutralised via weak acids and bases located in the cytosol [Gibbin et al. 2014; Casey et al. 2010]. However, more significant long-term fluctuations involve the use of more permanent mechanisms such as transmembrane exchangers [Gibbin et al. 2014]. Cellular dysfunction is often associated with an abnormally low intra compartmental pH, which can function to denature proteins or activate enzymes, which are usually inactive at neutral pH [Han and Burgess, 2010]. Additionally, low intracellular pH may negatively influence upon other aspects of human physiological functioning particularly the nervous system and has been implicated in the pathophysiology of cancer [Izumi et al. 2003] and Alzheimer’s disease [Dennis et al. 2012]. Typically, cytosolic pH should be between 7.2-7.4, with even subtle alterations inducing cellular dysfunction [Dennis et al. 2012].

Given the central role, that intracellular pH plays in cell function it is not surprising that practically all cell types operate to regulate intracellular pH and ensure it remains within the optimal physiological range [Valli et al. 2005]. This steady state intracellular pH is determined by the balance between the rate of acid extruding processes and acid loading processes [Boron et al. 2004]. In terms of acid loading processes, this includes the uptake of HCO$_3$- and CO$_3$ by the Na$^+$-coupled members of the solute carrier family (SLCF) family, Na-H exchange, vacuolar H$^+$ pumps in the plasma membrane, in addition to lactate efflux. Alternatively, acid loading processes involve HCO$_3$- efflux, which is aided by the Cl$^-$-HCO$_3$ exchangers in the SLC4 family with help from the anion channels.

2.1.2 Acidosis and fatigue development

The role of pH and the precise physiological mechanisms that contribute to fatigue remain relatively elusive and the subject of intense debate and investigation [Lancha Junior et al.
2015]. However, evidence does exist to suggest several physiological mechanisms associated with acidosis contribute to the development of muscle fatigue. For example, competition between H\(^+\) and Ca\(^{2+}\) ions for the troponin binding site leads to an impairment in the ability of the contractile components to function effectively [Donaldson et al. 1978; Fabiato et al. 1978]. In addition, excessive H\(^+\) ions function to inhibit phosphorylcreatinine resynthesis [Sahlin et al. 1975]. In addition to inhibiting enzymes involved in the glycolytic pathway such as glycogen phosphorylase and phosphofructokinase, limiting the ability of the muscle to deal with energetic demand [Sutton et al. 1981]. *In vitro* muscle cell pH homeostasis is maintained by the flux of H\(^+\) out of the cell and into the extracellular environment, this is achieved through the monocarboxylate transporters (MCT), particularly MCT-1 and MCT-4 [Juel. 2008]. These transport proteins carry monocarboxylates across the cell membrane. Additionally H\(^+\) ions are also transported out of the cell via the sodium hydrogen exchanger and the sodium bicarbonate co-transporter [Lancha Junior et al. 2015].

### 2.1.2 Intracellular sensing techniques

Conventional intracellular sensing techniques have included the use of fluorescent dyes [Bkaily et al. 1999], optochemical sensors, NMR and surface enhanced Raman scattering (SERS) [Desai et al., 2014]. NMR can effectively discriminate between cytoplasmic and vacuolar pH. However, a major drawback associated with this technique is the inability to measure pH at the single cell level, as larger tissue sections are required [Sondergaard et al. 2014]. Alternatively, SERS may be used to measure the pH of single cells following the internalisation of metal nanoparticles, which is promising, however presently the resolution of SERS when performing live cell imaging does not compare to fluorescent-based measurement [Sondergaard et al. 2014]. In relation to the use of fluorescent dye, there are a diverse range of fluorophores commercially available, which provide a rapid response, which is measureable via confocal microscopy or spectrophotometry [Han and Burgess. 2009]. The use of fluorescent dye enables the real-time measurement of various analytes, including pH. The small size of the free dye molecules provide the benefit of high spatial resolution and allows information throughout the cell to be collected *en masse* [Coupland et al. 2009].

Nevertheless, the technique is not without limitation. For example, the direct contact between the dye and the intracellular environment may be associated with cytotoxicity, through the
induction of biochemical processes within the cell [Desai et al. 2014]. There is also the potential for non-specific protein binding within the cell leading to false positive results and sequestration [Srivastava. 1997]. Furthermore, simply retaining the dye within the cellular environment throughout the duration of the experiment is another difficult problem as many dyes are prone to leeching due to their low molecular weight [Coupland et al. 2009].

Alternatively, optochemical sensors (optodes), overcome some of the inherent weaknesses associated with the use of free dye (graphic representation of an optode displayed in Figure 2.1). For example, the tip of the optical fibre consists of a bio-compatible matrix protecting the cell from cytotoxicity associated with the dye. Nevertheless, there are crucial limitations associated with the use of the optochemical sensor. For example, the technique requires a fibre with a modified tip to be inserted directly into an individual cell [Buhlman et al. 1998; Vo-Dinh. 2003]. The large size of the fibre in comparison to a single cell causes damage to the cell membrane, which can trigger apoptosis. Further, the physical size of the probe bodies prevents more than two or three sensors being used per cell. This limits resolution of the intracellular sensing capabilities [Coupland et al. 2009].

Figure 2.1: Graphic representation of an ‘optode’ used for intracellular analysis
2.1.3 PEBBLE Nanosensors

The PEBBLE (Probes Encapsulated By Biologically Localised Embedding) nanosensor represents a major development within sensor-based technology, overcoming many of the inherent weaknesses associated with optochemical and free-dye sensing techniques. For example, due to their sub-micron diameter, inert matrix, signal intensity and ratiometric ability they can be used to accurately characterise intracellular compartments and allow real-time measurement in sub-cellular environments [Aylott et al. 2003]. Effectively, PEBBLE is a term, which can be used to describe a wide range of matrices and nano-fabrication techniques that have been used to miniaturise optical sensing technologies [Monson et al. 2014]. Their structure is spherical and they are around 30-500nm in size [Desai et al. 2014], taking up approximately 1ppb of the internal volume of a mammalian cell, resulting in minimal perturbation and ‘passive’ (non-destructive) analytical observations [Buck et al. 2004; Clarke et al. 1999]. The most common matrices used to fabricate the nanosensors are, polyacrylamide, sol gel silica and cross-linked methacrylate [Monson et al. 2014]. This chapter will focus upon the use of polyacrylamide nanosensors. See Figure 2.2 for a simplified illustration of nanosensor structure and function.

The nanosensor exhibits a wide range of benefits over conventional sensing techniques. For example, they consist of an inert matrix, which prevents any non-specific interaction between fluorophores and cellular components, which reduces potential cytotoxic effects exhibited by the dye. [Desai et al, 2014].

![Figure 2.2: Schematic representing a PEBBLE nanosensor](image-url)
A major limitation of traditional free fluorescent dyes is uneven dye loading within the cell. During the loading process, each cell will take up an arbitrary amount of fluorophore. This means that when performing single cell analysis variations in fluorescence between cells may be dependent on the concentration of fluorescent dye in each cell rather than the concentration of the analyte of interest. Nanosensors routinely contain two types of fluorophore; an indicator and a reference probe. The indicator functions to generate a signal, which is directly proportionate to the concentration of the analyte of interest within the cell. The reference dye does not respond to the analyte of interest and exhibits a constant signal at a different wavelength to the indicator dye. Although there may still be considerable variation in the number of nanosensors take up by individual cells, the analyte-dependent fluorescent signal can be ‘normalised’ with respect to the reference signal. Absolute fluorescent values may vary but the ratio of indicator to reference signal will be constant for a given concentration of analyte.

The combination of these dyes allows for accurate and ratiometric measurements to be performed. It is possible to add multiple pH sensitive dyes to the nanosensor matrix during production to allow for an extended range of pH measurements to be obtained across the physiological range [Chauhan et al. 2011; Sun et al. 2011]. The fluorophores used in this chapter were selected in reference to the Desai et al. [2013] protocol. This included the use of two pH sensitive dyes with identical emission spectra but with varying pKa values (5’6-carboxyfluorescein [FAM]) pKa 6.5, Oregon green pKa 4.8. In addition to a reference probe (5-(and-6)-carboxy-tetramethylrhodamine (TAMRA). Essentially, the optimal responsiveness of Oregon green is within the acidic range (3.5-5.5) whereas FAM is optimally stimulated in the neutral range (5.5-7.5). Therefore, at low pH levels Oregon green responds to pH, while FAM is largely inactive [Desai et al. 2013].
2.1.4 Nanosensor delivery

The cellular cytoplasm is separated from the external environment by a plasma membrane, in addition to lipid bilayer with membrane proteins, lipids and carbohydrates embedded. Before a nanosensor can effectively interpret sub-cellular pH it must be capable of first crossing the cell membrane, which can be challenging [Chou et al. 2011].

It is possible to aid intracellular delivery by bio-chemically engineering the synthetic nanosensor to aid passage through the plasma membrane into the intracellular environment. One method involves functionalising the nanosensor with specific ligands, which are capable of binding to the surface of the cell membrane, enabling receptor-ligand interaction and leading to receptor mediated endocytosis of the nanosensor [Chou et al. 2011]. Alternatively, cationic coating of the nanosensor is another internalisation strategy. Effectively, cationic molecules are able to interact with the negatively charged plasma membrane, to promote membrane permeability [Chou et al. 2011]. It has been hypothesised that increased permeability is due to the formation of nanoscale holes within the plasma membrane following interaction with cationic species [Al-Jamal et al. 2008; Al-Jamal et al. 2008]. A number of cationic coatings have been used in the transfection process and include; cationic liposomes, polypeptides and amine-containing polymers [Herrero et al. 2009; Chatterjee et al. 2008]. An example of a popular liposomal transfection reagent is lipofectamine. This reagent is able to complex with negatively charged, water-soluble quantum dots through simple electrostatic interaction during co-incubation in culture media. This delivery technique involves pinocytosis, a process which enables the cell to take in the surrounding extracellular environment as a result of vaginating and pinching off the plasma membrane into vesicles containing the extracellular fluid [Derfus et al. 2004]. A major benefit of liposomal delivery is that nanosensors can be delivered to a large number of cells simultaneously [Monson et al. 2014]. Figure 2.3 illustrates the various endocytic pathways into the cell, in addition to the possible transport routes the nanosensor could take through the endosomal network.
2.1.5 pH sensitive nanosensor application

Pilot work conducted by Boulton [2012] investigated the ability of the novel pH sensing nanosensor system incorporating to provide pH measurements in CFS/ME patient myoblasts. The nanosensors consisted of a polyacrylamide matrix and incorporated the free dyes Fluorescein isothiocyanate (FITC) and Alexa Fluor 568. The aim of the investigation was to use the nanosensors to confirm the presence of intracellular acidosis in CFS/ME patients, which has been previously reported via MRS techniques [Jones et al. 2010].

Boulton [2012] investigated the ability to successfully deliver the pH nanosensors to the intracellular environment of the myoblasts using the transfection reagent Lipofectamine 2000. Confocal microscope imaging was reported to confirm the nanosensor distribution throughout the intracellular environment (displayed in Figure 2.4) Nevertheless, it is important to interpret these images with caution, as higher quality imaging is required to confirm the sub-cellular location of the nanosensor or simply whether they were bound to the outer cell membrane.
Additionally, the nanosensors were also reported to quantify intracellular pH via fluorimetric measurement. CFS/ME myoblasts loaded with pH responsive nanosensors exhibited significantly lower intracellular pH compared to controls. The acidosis was reversed when the cells were treated with the PDK inhibitor DCA (See Figure 2.5). These findings suggested that CFS/ME related acidosis could be reversible and potentially treatable with drug intervention. Therefore, a primary aim of this chapter was to reproduce this pilot data to further confirm the presence of aberrantly low pH in CFS/ME myoblasts via the nanosensor system and the extent in which it could be manipulated following DCA treatment.

Figure 2.4 Preliminary data generated by Boulton [2012], depicting confocal imaging of discreet-z-slice in a control myoblast. A fair distribution of nanosensors throughout the cell was observed. FITC fluorescence A, Alexafluor568 fluorescence B, Bright field C and merged D.
2.1.6 Hypotheses

The experimental hypotheses were (1) CFS/ME myoblasts exhibit intrinsic intracellular acidosis compared to controls. This could represent a major advance in the understanding of mechanisms that underpin a major phenotypic characteristic of this condition. (2) A normalisation in CFS/ME intracellular pH to control levels could be achieved in response to the PDK inhibitor DCA. This could represent a novel treatment strategy in the management of symptoms associated with CFS/ME.

Figure 2.5 Preliminary data generated by Boulton [2012]. Myoblasts were treated with 16µM DCA. No change in pH between treated and untreated control myoblasts. DCA boosted pH beyond the baseline level of normal myoblasts. Untreated CFS/ME exhibited a significantly lower pH. Data presented ± SEM, N=4, *** denotes p<0.005
2.2 Materials and Methods

All diseased (CFS/ME) and non-diseased primary myoblasts were supplied by Dr Audrey Brown at Newcastle University.

All tissue culture reagents were purchased from Sigma-Aldrich (Poole, Dorset, UK) with the exception of Ham’s F10 myoblast growth media (Lonza, SLS, East Riding of Yorkshire, UK) and chick embryo extract (Seralab, West Sussex, UK). Tissue culture flasks, 24-well plates and serological pipettes were supplied by Greiner bio-one (Stonehouse, UK). Resazurin (7-Hydroxy-3H-Phenoxazin-3-one10-oxide) dye was purchased from Sigma-Aldrich.

pH responsive dyes and TAMRA were purchased from Life Technologies (Glasgow, UK).

Inhibitors of enzyme PDK (Potassium Dichloroacetate, Sodium Iodoacetate, Dehydroabietylamine) were purchased from Sigma Aldrich.

The delivery reagent Lipofectamine 2000 was purchased from Thermo Fisher Scientific (Paisley, UK).

2.2.1 Study participants

Muscle biopsy samples were obtained from patients with CFS/ME and healthy control participants. The CFS/ME and control individuals were gender and age matched, for example in this chapter the CFS/ME patient was a female aged 65 years and the control participant was a female aged 66 years. Recruitment was achieved through the NHS CFS clinical service within the Newcastle Upon Tyne Hospital Foundation Trust. The CFS/ME patients all met the Fukuda (1994) criteria for CFS/ME. All participants agreed to complete the study via formal written consent. Ethical approval for the study was provided by the Newcastle and North Tyneside joint ethics committee.

2.2.2 Cell culture and preparation

Muscle biopsies from both CFS/ME patients and control participants were obtained and isolated as previously described by Brown et al. [2015]. Briefly, samples were collected from
the vastus lateralis (Location illustrated in Figure 2.6) via fine needle biopsy. The precursor cells were then isolated via the Blau and Webster method [1981]. The biopsies were collected in medium designed for proliferation (Ham’s F10 supplemented with 20% (v/v) FBS, 2% chick embryo extract, 1% penicillin-streptomycin). The samples were then transferred into a petri dish and washed with PBS. Any adipose or connective tissue was removed with a scalpel before the samples were again washed in PBS. The samples were then added to a falcon tube which contained 5mL 0.05% trypsin-EDTA and spin-digested at 37°C for 15 minutes. The trypsin was then removed and 5mL of media added before centrifuging at 448 G for 5 minutes. The pellet containing the satellite cells was then re-suspended in medium and the spin dissociation process was repeated an additional 3 times. The pellets were then plated into a T25 culture flask with Ham’s F10 growth medium. The media was changed after 24 hours to remove any cellular debris. Cells were expanded via routine culture and proliferating myoblasts were passaged multiple times before performing experimental work.

Diseased (CFS/ME) and non-diseased primary human myoblasts were routinely cultured using Ham’s F-10 medium supplemented with 20% FBS 2% chick embryo extract, 500U/mL penicillin streptomycin and 1% amphotericin B. Cells were grown in a humidified environment 5% (v/v) CO₂ at 37.5°C throughout. Cells were grown to 80% confluence before passaging. Prior to experimentation growth media was removed and replaced with experimental medium which was unsupplemented Dulbecco’s modified eagles medium.

![Figure 2.6 Schematic of the lower limb anatomy as depicted by Dixit et al. [2007]](image-url)
2.2.3. Intracellular optical sensing

2.2.3.1 pH responsive nanosensor production

PEBBLE (Probes Encapsulated by Biologically Localised Embedding) Nanosensors were synthesised during a multi-stage process developed by Chauhan et al. [2011]. The method involved dissolving acrylamide (7.6mM) and N,N Methylene bisacrylamide (1.3mM) in 1.5mL of deionised water by sonication to create a monomer solution. Dextran linked reference fluorophore TAMRA-D and indicator dyes FAM-D and Oregon green-D were added to the monomer solution at 10mg/mL and covered to avoid exposure to light as they are photosensitive. A surfactant mixture was prepared consisting of Dioctyl sulphasuccinate (3.6mM) and Brij 30 (8.5mM) and deoxygenated. The surfactant was added to 42mL of deoxygenated hexane by perfusion with argon gas to form a micro emulsion. The monomer/fluorophore solution was then added to the surfactant mixture and left for 10 minutes. The polymerisation process was then initiated by the addition of 30µL of ammonium persuphate (10% w/v) and 15µL of -N,N,N’,N’tetramethylethylethylenediamine. Following 2 hours of polymerisation hexane was removed by rotary evaporation in the absence of oxygen and the remaining nanosensors washed 5 times in absolute ethanol, which was removed by rotary evaporation during the final wash. The remaining sensors were stored in an airtight container at 4°C.

2.2.3.2 pH sensitive nanosensor calibration

The pH nanosensor calibration was performed following the Desai et al. [2013] protocol. Buffer stock solutions were prepared before completing the calibration. A 0.2M stock of sodium phosphate dibasic was prepared by adding 7.098g of sodium dibasic to 250mL of deionised water into a volumetric flask before sonicating. A 0.1M stock solution of citric acid monohydrate was also prepared by adding 5.524g of citric acid monohydrate to 250mL deionised water and then sonicated. Buffer solutions between pH 4-8 and the volumes of sodium diphosphate and citric acid monohydrate were added to 50mL Falcon tubes as detailed in Table 1.1. A pH meter was then used to confirm the pH of each solution.
Nanosensor suspensions were prepared by dissolving nanosensors in buffer solutions over a range of pH values (4-8) in each buffer solution to achieve a final concentration of 10mg/mL. The nanosensors contained 3 dye probes OG-D, FAM-D and Tamra-D (reference dye). The nanosensor/buffer solution was then added (200 µL) in triplicate to a 96-well plate, in addition to a control triplicate containing only buffer solution to ensure that background fluorescence did not interfere or overlap with the response obtained from the dye nanosensors. Fluorescent intensity (FI) measurements were performed as detailed in section

<table>
<thead>
<tr>
<th>pH</th>
<th>Sodium phosphate dibasic (0.2M)</th>
<th>Citric acid monohydrate (0.1M)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>7.72</td>
<td>12.28</td>
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<tr>
<td>5</td>
<td>10.28</td>
<td>9.72</td>
</tr>
<tr>
<td>6</td>
<td>12.84</td>
<td>7.16</td>
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<tr>
<td>7</td>
<td>17.44</td>
<td>2.56</td>
</tr>
<tr>
<td>8</td>
<td>19.53</td>
<td>0.47</td>
</tr>
</tbody>
</table>

Table 1.1. The volumes required of the stock concentrations of sodium phosphate dibasic (0.2M) and citric acid monohydrate (0.1M) required to make buffer solutions between pH 4-8.
2.2.3.2 Fluorometric measurement strategy

Fluorescence measurements were obtained using a Tecan Infinite 200 fluorimeter (Tecan, Mannedorf, Switzerland). Data was attained using the companion Magellan software. The measurement settings for the active dyes OG-D and FAM-D were (λ<sub>ex</sub> = 490nm, λ<sub>em</sub> = 525nm) and for reference dye TAMRA-D (λ<sub>ex</sub>=555, λ<sub>em</sub>=580). The software was set to perform multiple reads per well and a gain value of 50 was selected remained constant across all experiments.

2.2.4 Resazurin cytotoxicity assay

Prior to all cytotoxicity assays CFS/ME and control myoblasts were routinely cultured and seeded into 96-well testing plates at a density of 1x10<sup>4</sup> cells per well. Following seeding cells were incubated in a humidified environment at 37.5°C and 5% CO<sub>2</sub>. Myoblast viability was measured by adding resazurin dye (concentration 0.03%) to each well. Resorufin produced as a result of resazurin bio-reduction was measured using a Tecan-I fluorometric plate reader at λ<sub>ex</sub>560nm / λ<sub>em</sub>590nm. Essentially, resazurin is a none fluorescent dye, however in the presence of viable cells it is reduced by the mitochondria into highly fluorescent forms of the dye namely resorufin and dihydroresorufin.

2.2.4.1 Dye free nanosensor toxicity

To assess the biocompatibility of the polyacrylamide nanosensor matrix, nanosensors containing no fluorophores (blanks) were added to the wells of the testing plate at a 10mg/mL concentration. Resazurin dye was added at a concentration of (0.03% w/v) and fluorometric analysis were performed as described in section 2.2.4.

2.2.4.2 pH sensitive nanosensor cytotoxicity

Cell viability was also investigated following incubation with nanosensors containing pH responsive dyes OG-D and FAM. Briefly, nanosensors were added to the wells of the testing
plate at a 10mg/mL concentration in addition to resazurin dye (concentration 0.03% w/v). The plate was then read over the following 3,4,5,6 hours as detailed in section 2.2.4.

2.2.4.3 Free dye FAM and Oregon Green cytotoxicity

The nanosensor has been reported to provide a protective barrier between the cell and the potential cytotoxic effects of the dyes which are encapsulated during the fabrication process [Desai et al. 2013]. Therefore, the cytotoxicity of the pH responsive dyes FAM-D and OG-D was assessed as these dyes were incorporated into the nanosensors described in this chapter. Concentrations (5-20µg/mL) of FAM-D and OG-D were added to the cell culture plate. Viability was investigated following the addition of resazurin (concentration 0.03% w/v). The plate was monitored over the following 3 hours as described in section 2.2.4.

2.2.4.4 Lipofectamine 2000 cytotoxicity

In this chapter lipofectamine 2000 was transfection reagent used to deliver nanosensors into the intracellular environment. The cytotoxicity of the reagent was assessed at varying concentrations (0.003-0.3%) which were added to the wells of the testing plate, followed by 0.03% resazurin and incubated for 3 and 6 hours. Fluorescent intensity (FI) was then measured as detailed in section 2.2.4.

2.2.4.5 DCA cytotoxicity

The impact of PDK inhibitor DCA upon intracellular pH was investigated in this chapter. It was important to assess the cytotoxicity of the drug before applying it to the cell system. DCA was added to 96-well testing plates at varying concentrations (1µM-1mM). Resazurin (concentration 0.03% w/v) and plates were incubated for 3 and 6 hours. The plate was read as described in section 2.2.4.
2.2.4.6 pH measurement of DCA in a cell-free system

The impact of PDK inhibitor DCA upon myoblast intracellular pH was investigated in this chapter. Prior to applying the drug to a cell system it was essential to firstly determine the inherent pH of the drug in a cell-free system. To achieve this nanosensors at a concentration of 10mg/mL were added to the wells of a 96-well testing plate and then treated with varying concentrations of DCA (0-40µM). FI measurements were performed as described in section 2.2.3.2.

2.2.5 pH sensitive nanosensor internalisation with varying lipofectamine concentration

Primary myoblasts were routinely culture and seeded at a density of 1x10⁴ cells per well, in triplicate into 96-well testing plates. Following 12-hours incubation in a humidified environment at 37°C and 5% CO₂ day, nanosensors at 10mg/mL were dissolved in experimental media containing lipofectamine 2000 at varying concentrations (0.003-3%). The nanosensor/lipofectamine mixture was allowed to complex for 30 minutes at 37.5°C. The culture media was then removed from the testing plates and the nanosensor/lipofectamine mixture was added to the wells and left for 3 hours at 37.5°C and 5% CO₂ to allow for nanosensor uptake. The plate was washed twice with PBS before replacing with experimental media and incubated for a further 30 minutes at 37.5. Intracellular pH was then measured over the next 6 hours. FI measurements were performed as described in section 2.2.3.2.

2.2.6 pH sensitive nanosensor internalisation and DCA treatment

Primary myoblasts were routinely cultured and seeded into 96-well testing plates at a density of 1x10⁴ cells per well. Following attachment, a nanosensor (10mg/mL) and lipofectamine (0.03%) complex was allowed to complex for 30-minutes before being to the well and incubated at 37.5°C and 5% CO₂ for 3 hours to enable nanosensor uptake. The plate was washed twice with PBS before replacing with experimental media, which was then dosed with various concentrations of DCA (0-20µM). Intracellular pH was measured over the following 6 hours. FI measurements performed as described in section 2.2.3.2.
2.2.7 Data analysis

The statistical model used to analyse the data was Minitab 17 statistical software. Inferential statistics were performed to investigate intracellular pH in CFS/ME and control skeletal muscle samples. An independent sample T-test was used to examine intracellular pH differences between different samples (CFS/ME vs control). A paired sample T-test was utilised to investigate within group differences, for example pH at baseline verses DCA treatment. The descriptive statistics used were mean ± SD.
2.3 Results

2.3.1 pH Sensitive nanosensor calibration curve in a cell-free system

pH responsive nanosensors at 10mg/mL were initially analysed in the absence of cells over a range of pH values. Fluorescent intensity (FI) ratio measurements for OG-D and FAM in relation to reference dye TAMRA-D were obtained. FI ratio values were shown to be linearly dependent on pH \((y=6.0624, R^2=0.9885, N=3)\). The linear regression of the calibration curve was then utilised to determine intracellular pH for nanosensor-doped myoblast samples. The data is presented in Figure 2.7.

![Figure 2.7 Calibration of pH responsive nanosensors containing FAM-D and OG-D. The Fluorescent ratio was shown to be linearly dependent upon pH \((y=6.0624x)\). The data is presented as mean ± SD, n=5, \(R^2=0.9885\).](image-url)
2.3.2 pH responsive nanosensor optimisation

2.3.2.1 Myoblast viability following exposure to dye-free (blank) nanosensors

CFS/ME and control myoblast viability was assessed following the addition of dye-free nanosensors and was measured via the resazurin viability assay. Figure 2.8 demonstrated myoblast viability over time following incubation with 10mg/mL dye-free nanosensors. After 24-hours CFS/ME and control myoblast viability was significantly reduced (% of untreated control) when compared to all other time points (P≤0.0005). There was no significant difference between any other time points.

![Viability graph](image)

Figure 2.8 CFS/ME and Control myoblast viability (% of untreated control) following incubation with 10mg/mL of dye free nanosensors. Data presented as mean of 3 replicate measurements ± SD. Significant difference in viability between 24 hours and all other time points denoted by **** (P≤0.0005), n=2
2.3.2.2 Myoblast viability following exposure to Oregon green and FAM

Figure 2.9 demonstrated CFS/ME myoblast viability following treatment with varying concentrations (5-20µg/mL) of pH responsive free dyes OG-D and FAM-D, simultaneously. Viability was decreased to less than 70% (of untreated control values) across all concentrations after 1 hour.

CFS/ME cell viability was significantly decreased (P≤0.0005) between 1 and 2 hours for a free dye concentration of 20µg/mL. Similarly, there was a significant decrease (P≤0.0005) between 2 and 3 hours, resulting in a final % viability of less than 40%. The higher the dye concentration and the longer the exposure period the higher the cell death. Cell death was only seen to be concentration dependent beyond 2h incubation.

![Graph showing cell viability (%) over time](image)

Figure 2.9 Control myoblast viability (%) after incubation with varying concentrations (5-20µg/mL) of pH sensitive Oregon green and FAM. Data presented as mean ± SD, n=3. Significant difference in viability between 1 and 2 hours at 20µg/mL, and similarly between 2 and 3 hours as denoted by **** (P≤0.0005)
2.3.2.3 Myoblast viability following treatment with Lipofectamine 2000

Following 3-hours incubation with varying concentrations of lipofectamine myoblast viability is shown in figure 3.1. During this time period CFS/ME myoblasts exhibited significantly reduced (P<0.005) viability at 3% verses 0.003% lipofectamine concentration, a decreased viability of 33%. Similarly, control myoblast viability was also significantly reduced at 3% versus 0.003% lipofectamine concentration (P<0.05) which resulting reduction in viability by 44%.

Figure 3.0 CFS/ME myoblast viability (%) after incubation in varying concentrations (5-20µg/mL) of pH sensitive Oregon green and FAM. Data presented as mean of triplicate measurements from n=2 donors ± SD. Significant difference in viability between 1 and 2 hours at 20µg/mL, and similarly between 2 and 3 hours as denoted by **** (P≤0.0005)
2.3.2.4 Myoblast viability following incubation with DCA

Myoblasts were treated with varying concentrations (1µM-1mM) of the PDK inhibitor DCA and viability was assessed via the resazurin assay. Figure demonstrated viability (%) of untreated control) after 3-hours incubation and Figure 3.3 following 6-hours. Results demonstrated good viability at both 3 and 6 hours incubation, across all concentrations and in both CFS/ME and control samples (>88%).
Figure 3.2 CFS/ME (n=2) and control (n=2) myoblast viability following 3-hours incubation with 1µM-1mM DCA. Viability (% untreated control) was assessed following the addition of 0.03% resazurin. Myoblast viability remained high across all concentrations (>90%) in both CFS/ME and control sample groups. Data presented as mean ±SD.

Figure 3.3 CFS/ME (n=2) and control (n=2) myoblast viability following 6-hours incubation with 1µM-1mM DCA. Viability (% untreated control) was assessed following the addition of 0.03% resazurin. Myoblast viability remained high across all concentrations (>87%) in both CFS/ME and control sample groups. Data presented as mean ±SD.
2.3.2.4 pH measurement of DCA in a cell-free nanosensor system

Figure 3.4 demonstrated the calculated pH values for varying concentrations of DCA (0-40µM) measured by pH responsive nanosensors at a concentration of 10mg/ml. All concentrations of DCA maintained neutral pH during course of experiment.

![Graph showing calculated pH for varying concentrations of DCA (0µM-40µM) measured by nanosensors at 10mg/mL concentration. Data presented as mean of 3 replicate measures ±SD.](image)

2.3.3 Nanosensor application

2.3.3.1 Myoblast pH with varied lipofectamine 2000 concentration

Figure 3.5 demonstrated CFS/ME and control myoblast intracellular pH measurements with varied lipofectamine 2000 concentration (0.003-3%). Control myoblast intracellular pH was significantly lower at all lipofectamine 2000 concentrations (P<0.005) compared to CFS/ME samples.
2.3.3.2 Myoblast pH with varied DCA treatment

Intracellular pH was measured using nanosensors at a 10mg/mL concentration. Following nanosensor internalisation, cells were treated with varying concentrations of DCA (10-20µM) and incubated for 3-hours. pH was significantly lower in control compared to CFS/ME myoblasts at baseline (0µM DCA) and with all DCA concentrations. Displayed in Figure 3.6.
Figure 3.6 CFS/ME & control myoblast intracellular pH at baseline (0µM) and following treatment with DCA (10-20µM). pH measured with nanosensors at 10mg/mL concentration. Significantly lower intracellular pH for control compared to CFS/ME myoblasts at baseline and following treatment with all concentrations of DCA. P<0.005 denoted by ***, Data presented as mean of triplicate measurements ±SD, n=2 donors
2.4 Discussion

Existing literature has demonstrated the role of impaired bio-energetic function in CFS/ME patients. For example, MRS studies have reported patients to have a delayed recovery from intramuscular acidosis following a standardised exercise protocol [Jones et al. 2010], as well as profound and sustained intracellular acidosis with repeat exercise [Jones et al. 2012]. Furthermore, Boulton [2012] reported the presence of inherently low intracellular pH in CFS/ME myoblasts compared to controls when measuring using the novel intracellular nanosensor system. Additionally, the author also reported DCA to completely normalise intracellular pH in the CFS/ME patient samples. Thus, suggesting bio-energetic dysfunction in CFS/ME may be modifiable and therefore treatable.

In this chapter the intracellular pH of CFS/ME and control myoblasts was determined using pH responsive fluorescent nanosensors. Specific aims were [1] Repeat pilot work conducted by Boulton [2012] to confirm the presence of abnormal pH in CFS/ME myoblasts measured using pH responsive nanosensors. [2] Apply PDK inhibitors to the cells to investigate the ability of drugs to treat intracellular acidosis and potentially open a new route to therapeutic intervention.

The work carried out in this chapter was unable to reliably detect any acidosis in CFS/ME cells or any difference in intracellular pH between CFS/ME and control cells. Secondly, DCA did not modify intracellular pH in either CFS/ME or control myoblasts contrasting previously reported data. The following sections will contextualise the findings and suggest possible reasons for the conflicting results.

2.4.1 pH nanosensor optimisation

Optimisation work was required before the nanosensor experimental platform could be utilised to interpret intracellular pH in CFS/ME patient samples. This included varying dye ratios in order to optimise signal strength and investigating innate cellular toxicity associated with the nanosensors.

The nanosensors were successfully calibrated over a diverse pH range (4-8), however this was could only be achieved following chemical modification of the pH sensitive dyes FAM.
and Oregon green. During the initial calibration process there was a large degree of fluctuation in the signal strength provided by the pH responsive dyes. Additionally, it was observed during the washing steps of the fabrication process that there was a substantial amount of dye leaking from the sensors with each wash. It was felt that the variation in signal strength could be linked to the apparent dye leeching observed during fabrication. Therefore, as suggested by Desai et al [2013] dextran linked fluorophores were incorporated into the fabrication process. Effectively, dextran is an inert molecule, which is large enough in size to entrap the fluorophores in the nanosensor matrix to prevent the low molecular weight dye leaking from the sensor.

The nanosensor has been reported to exhibit a wide range of benefits over the use of conventional free-dye based sensing techniques. For example, the polyacrylamide shell has been described as biologically compatible, preventing non-specific interaction between fluorophores and cellular components, therefore minimising cytotoxicity [Desai et al. 2014]. Thus, a key aim of the optimisation work was to investigate whether blank (dye-free) nanosensors exhibited a cytotoxic effect on the CFS/ME and control myoblasts. Interestingly, the nanosensors were found to be bio-compatible with short term incubation (≤6 hours), however after 24-hours viability in both CFS/ME and control samples was significantly reduced (<40%) when compared to other time points (See Figure 2.8 and 2.9). In contrast to this finding Lee et al. [2010] reported blank polyacrylamide nanosensors to exhibit a non-toxic effect (>90% viability) in a glioblastoma cell line following 20-hours incubation. However, the nanosensor concentrations used (1-4mg/mL) were considerably lower than used in this chapter (10mg/mL). The higher sensor concentration was chosen to give a more clearly defined intracellular signal.

It is not clear why prolonged exposure caused substantial cell death in the present investigation. However, one possible explanation could relate to chemical residue left on the nanosensor surface following the manufacturing process. Following synthesis the nanosensors were washed in several changes of ethanol in order to remove plasticisers used during the fabrication process. Although the sensors were washed 7 times in ethanol as previously recommended [Desai et al. 2013], it is possible that residual contaminants may have remained on the surface and it may have taken prolonged exposure to these contaminated probes before significant cell death occurred.
Another potential factor that could be attributed to cell death with prolonged exposure could relate to contamination within the culture. When working with a primary cell culture the need for sterility is very important and the introduction of any nonsterile product to cell culture greatly increases the risk of contamination [Burns et al. 2011]. Nevertheless, contamination can usually be avoided through the use of conventional sterilisation techniques. However, the ability to sterilise the nanosensors via typical sterilisation techniques such as autoclaving and UV irradiation is not possible. For example, UV irradiation may photo bleach the fluorophores making them less responsive to changes in pH [Benjaminson et al. 2011]. Additionally, sterilisation via autoclaving is not possible as excessive heat exhibits the capacity to limit fluorophore function and disrupt the polyacrylamide structure of the nanosensor shell. Therefore, the inability to sterilise the nanosensors is an inherent weakness in the ability to use the platform to measure pH in cell cultures where sterility is a necessity.

Additional toxicity work was performed to investigate the impact of exposure to pH responsive dyes Oregon green and FAM. A key benefit of the nanosensor has been reported to be its ability of the polyacrylamide matrix to protect the cell from possible fluorophore cytotoxic effects. Following exposure to FAM and Oregon green both CFS/ME and control myoblasts exhibited a significant reduction in viability (<70% compared to untreated controls) after only 1 hour. Similarly, viability was also significantly reduced between 1 and 2 hours and again between 2 and 3 hours at the higher dye concentration (20µg/mL) and in both samples (See Figure 3.0). Therefore, it would appear that the higher the concentration of the dyes and the longer the exposure, the greater the cytotoxic effect. Interestingly, while blank nanosensors demonstrated biocompatibility during early measurements, it would appear the free dyes Oregon green and FAM are acutely cytotoxic (i.e. within 1 hour of exposure to the cells).

Toxicity work was also performed on the transfection reagent Lipofectamine 2000 (Displayed in Figure 3.1). It has been reported to exhibit negligible cytotoxicity in a wide variety of cell-lines [Cui et al. 2012]. Results demonstrated good viability (>70% compared to untreated controls) at the lower concentrations (0.003, 0.03%) however after 3-hours incubation at the highest concentration (3%) there was a significant reduction in viability (<40% cells still viable) in both CFS/ME and control samples. Therefore, concentrations below 3% were used in the nanosensor internalisation experiments.
The PDK inhibitor DCA was added to cells that were loaded with nanosensors to test the ability of DCA to ‘normalise’ CFS/ME myoblast intracellular pH. The cytotoxic profile of the inhibitor was investigated and it was associated with good viability (>89%) when measured over a 24-hour period (See Figure 3.2 and 3.3). This is in agreement with the data presented in a review by Papandreou et al. [2011], who reported clinically relevant concentrations of DCA (<1mM) to exhibit no direct cytotoxic effect in vitro.

Finally, it was also important to determine the innate acidity/alkalinity of the inhibitor and the potential effect it may have in relation to intracellular pH. pH was measured via the nanosensors and DCA maintained a neutral pH during the course of the experiment. Therefore, DCA was considered acceptable to use within the nanosensor experimental system.

2.4.2 pH Nanosensor application

The discovery that control myoblasts exhibited significantly lower intracellular pH compared to CFS/ME patient samples, contrasts previous findings [Jones et al. 2010, 2012; Boulton, 2012], which measured intracellular pH via MRS and using a pH responsive nanosensor platform, similar to the one reported in this chapter. However, the design incorporated only one pH responsive fluorophore (FITC), with a limited pH range (Pka 6.5). In contrast, in this chapter a dual-fluorophore approach to extend the dynamic pH range, with fluorophores exhibiting the same emission spectra but different PKa values, increasing the pH sensing range (3.5-7).

Importantly, unlike the aforementioned studies the data generated in this chapter demonstrated both CFS/ME and control myoblasts to exhibit intracellular pH readings outside the cytosolic physiological range of around 7.2 [Casey et al. 2010; Han and Burgess, 2009; Llopis et al., 1998]. For example, intracellular pH was <4.5 for both patient and control samples following incubation with varying concentrations of transfection reagent lipofectamine 2000 [Displayed in Figure 3.5]. Similarly, following treatment with 0-20µM DCA intracellular pH was <5.7 for both samples [See Figure 3.6].

When interpreting these findings it is important to consider the heterogeneity of the intracellular environment and the difficulty in delivering nanosensors to specific sub-cellular
compartments. For example, mitochondrial pH has been reported as high as 8, compared to 4.7 in lysosomes, which are significantly different to the value of 7.2 reported in the cytoplasm [Panariti et al. 2012].

Despite many studies being conducted in the intracellular drug-delivery field, the ability of a nanoparticle to reach a specific sub-cellular compartment remains a substantial barrier, largely due to the complex and dynamic nature of the intracellular environment [Kristl et al. 2013; Ruenraroensak, 2010]. In the present investigation the sub-cellular component of interest was the cytoplasm, this is because with a greater glycolytic rate within the cell there is a concomitant increase in the release of protons in the cytoplasm [Robergs. 2001]. An increased rate of glycolysis is associated with a concomitant greater release in protons in the cytoplasm as a direct result of glycolysis increases greater glycolytic rate there is a concomitant increase in proton release in the cytoplasm as a direct result of glycolysis and ATP hydrolysis.

To understand why nanosensors provided a measurement outside the cytosolic range it is first important to consider how nanosized particles are trafficked within the cell. Intracellular trafficking processes are still not fully understood, however various internalisation pathways have been recognised [Zhang et al. 2012]. In the present study, cationic mediated delivery (Lipofectamine 2000) was utilised. This transfection approach has been reported to rely primarily on clathrin-dependent mediated pathways through the transmembrane [Cui et al. 2014]. Once endocytosed it has been suggested that nanosensors are encapsulated within transport vesicles which move though the endosomal pathway. It has been reported that nanoparticles taken up by clathrin-dependent endocytosis are typically marked for lysosomal degradation [Zhang, 2012; Bareford and Swan, 2007], which would explain the low pH values reported in the present study.

Although unpublished pilot work conducted by Boulton [2012] reported intracellular pH measurements within the cytosolic pH range, presently no published work has demonstrated the ability to deliver nanosensor to the cytosolic compartment. Interestingly, in agreement with the present study other authors have reported the nanosensor platform to provide pH measurements consistently outside the cytosolic range in a variety of cell lines. This finding has been associated with the endosomal and lysosomal localisation of the nanosensor once internalised [Sondergard et al. 2014; Benjaminsen et al. 2011; Coupland et al. 2009; Burns et al. 2006]. For example, in a study conducted by Benjaminsen et al. [2011] the authors
investigated the use of a polyacrylamide nanosensor, which covalently incorporated 2 pH responsive dyes (fluorescein and Oregon green) and reference probe rhodamine, to measure intracellular pH in a HepG2 cell line. The authors tracked pH using live cell imaging confocal microscopy and reported localisation of nanosensors to the endosome during early measurement time-points (1 hour: 5.1±0.6, 2 hours: 4.9±0.6) and exclusively the lysosome after 24-hours (4.5±0.4). Likewise, Coupland et al. [2009] reported pH values of 4.88 and 5.10 and combined with cell imaging confirmed endosomal localisation after 2-3 hours of uptake of a Tat conjugated polyacrylamide nanosensor in a CHO-K1 cell line. Furthermore, Burns et al. [2006] used an alternative nanosensor (matrix, silica gel), however similarly reported low pH values (5.11-6.6) an hour after uptake in a RBL-2H3 cell line. It is important to note that mildly acidic intracompartmental pH in the endosome and a significantly more acidic lysosomal pH is not an indication of cellular dysfunction. In terms of cell biology, the endosome plays a pivotal role in receptor recycling and the degradation of foreign material. Additionally, the lysosome is recognised as the principle site of intracellular digestion and is filled with hydrolytic enzymes that are used for controlled digestion of macromolecules involved in cellular metabolism [Han and Burgess. 2009; Alberts et al. 2002].

It is also important to note that the resting intracellular pH measurements reported by Boulton [2012] were lower (6.3-6.5 CFS/ME and 6.5-7.0 controls) than previously reported (7-7.1) by Jones et al. [2012] via MRS. Interestingly, even after exercise CFS/ME patients in the Jones et al. [2012] study exhibited an intracellular pH measurement of >6.6 which was not as acidic as the resting values reported by Boulton [2012]. It is therefore possible that the aberrantly low pH values reported in the present study and by Boulton [2012] were the result of nanosensor localisation within the acidic endosomal and lysosomal network. However, neither studies used live-cell imaging technology so it is difficult to draw firm conclusions.

An additional aim of this chapter was to investigate the capacity of the PDK enzyme inhibitor DCA to alter intracellular pH. Boulton [2012] reported DCA treatment to induce pH changes in CFS/ME myoblast samples, effectively normalising the highly acidic intracellular environment in CFS/ME cells to a level comparable to control samples, alternatively DCA did not exhibit any effect on control cells. The authors postulated the observed effect to be related to underlying abnormality in normal pyruvate metabolism. This could be either through reduced function PDH within multi-enzyme complex of PDC or due to over activity of PDK, leading to an over-reliance upon the lactate dehydrogenase energy-producing
pathway and localised acidosis. The data presented in this chapter showed that DCA did not alter intracellular pH in either CFS/ME or control myoblast samples (See Figure 3.4). However, the ability of this compound to modify cytosolic pH cannot be clearly defined as results would also suggest that the nanosensors were located in the endosomal and lysosomal intracellular compartments.

Aberrantly low pH values recorded in this and other work using intracellular pH responsive nanosensors would suggest that they are being sequestered by subcellular organelles. It is possible to release sensors from lysosomal entrapment but this is a complex task and requires chemical surface modification of the nanosensors. Benjaminson et al. [2013] suggested polyamines such as poly (ethylene imine) (PEI) which is positively charged to potentially enable the nanosensors to leave the lysosome via the ‘proton sponge effect’. This hypothesis suggests unprotonated amines of PEI to exhibit the ability to absorb protons while they are pumped into the lysosome. Subsequently, more protons are pumped into the lysosome, inducing an increased influx of Cl⁻ ions and water of the resulting osmotic swelling causes the lysosomal membranes to rupture and the contents are released into the cytoplasmic environment [Benjaminson et al. 2013; Nel et al. 2009; Behr et al. 1997]. Such an approach though potentially useful, was beyond the scope of the work presented in this chapter.

2.4.3 Limitations

The aim of this chapter was to repeat preliminary work conducted by Boulton [2012]. Therefore, care was taken to ensure the fabrication and internalisation methods performed were in line with pilot work. However, evidence suggested that nanosensor transfection via lipofectamine mediated delivery to be ineffective in delivering the nanosensor to the cytosolic compartment. This issue may have been rectified by incorporating a chemical surface modification step within the nanosensor fabrication process, to enable the nanosensors to break through the endosomal/lysosomal membrane and into the cytosolic environment.

Another potential limitation is that confocal microscopy was not used to confirm nanosensor internalisation in the present investigation. Previous studies using this imaging approach confirmed that lipofectamine was effective at internalising nanosensors into alveolar cell [Henderson et al. 2009]. It was felt that recorded pH values discussed in this chapter were so
far from physiological acceptable values that there were obvious limitations to the nanosensor approach that rendered imaging of little value.

In preliminary work conducted by Boulton [2012] images were provided to suggest the successful internalisation of the pH nanosensors. However, the image quality was limited and it was difficult to determine whether the nanosensors were internalised or bound to the outer membrane of the cell. Furthermore, it was impossible to determine the sub-cellular compartmental localisation of the nanosensors. Therefore in the present investigation the aforementioned imaging technique was not performed. To overcome this weakness a more sophisticated imaging technique could have been utilised. For example, Desai et al. [2014] reported fluorescence co-localisation via Wide field microscopy to enable the identification of the nanosensor sub-cellular location. This technique involves the genetic labelling of the nanosensor with fluorescent protein constructs.

2.4.4 Conclusion

The use of fluorescent-based pH responsive nanosensors revealed significantly lower pH in control myoblasts compared to those isolated from CFS/ME patient samples. This finding contrasted pilot work that reported CFS/ME samples to exhibit an aberrantly low intracellular pH, which could be returned to normal control levels following treatment with DCA. However, the pH values reported in this chapter are in line with other studies that utilised a similar experimental protocol and confirmed the endosomal/lysosomal localisation of the nanosensors via live-cell confocal imaging [Sondergard et al. 2012; Benjaminson et al. 2011; Coupland. 2009]. Thus, it would seem that without chemical modification the nanosensors are unable to freely locate within the cell cytosol. Future work could centre on the investigation of different nanosensor surface modification strategies to promote cytosolic delivery and prevent lysosomal entrapment. Additionally, fluorescent co-localisation microscopy techniques should be used to investigate nanosensor intracellular trafficking.
2.5 References


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Chapter 3

Direct, Real-time Electrochemical Detection of Superoxide Generation in CFS/ME Patient Myoblasts

3.1 Introduction

The purpose of this chapter was to utilise a direct, real-time electrochemical sensing technique to investigate the potential role of elevated O$_2^-$ production in the generation of the CFS/ME fatigue phenotype. For example, a series of in vivo investigations confirmed the elevation of blood oxidant status both pre and post-exercise, in a variety of CFS/ME patient cohorts [Jammes et al. 2012, 2009 and 2005]. Additionally, bio-markers associated with oxidative stress have been reported to play a pivotal role in skeletal muscle fatigue [Finsterer et al. 2012], which is regularly described as a debilitating symptom experience by CFS/ME patients [Van Oosterwijck et al. 2010]. Overproduction of ROS/RNS has been linked to impaired mitochondrial function. This may relate to the accumulation of modified mitochondrial proteins, lipid and DNA, which may interfere with the electron transport chain leading to cellular bio-energetic dysfunction. Presently, there have been no in vivo studies that have attempted to measure cellular superoxide generation directly and in real-time in CFS/ME patient muscle samples. This chapter will use an electrochemical superoxide sensor and myoblasts derived from CFS/ME patients.

Specifically, O$_2^-$ generation was measured in response to ethanol stimulation, which has recognised as a key chemical stimulant of O$_2^-$ in other cell lines (e.g. hepatocytes). Ethanol is metabolised by the enzyme cytochrome P450 2E1 (CYP2E1), which has been recognised as a key generator of oxidative stress (Gonzalez. 2007). While predominantly located in the liver, genes encoding CYP2E1 have been reported to be clearly expressed in human skeletal muscle [Molina-Ortiz et al. 2013], suggesting skeletal muscle to play a role in xenobiotic metabolism. Ethanol has also been reported to stimulate the mitochondria specific CYP2E1 enzyme [Bansal et al. 2010]. Given the presence of CYP2E1 in human skeletal muscle and as myoblasts contain a high volume of mitochondria, the use of ethanol as a chemical stimulant to assess the capacity of CFS/ME myoblasts to generate O$_2^-$ was justified.
O$_2^-$ generation was also measured in response to lactic acidification, to assess the impact of extracellular acidification on oxidative stress. As previously described CFS/ME patients have been reported to exhibit significant intracellular acidosis when measured via MRS [Jones et al. 2012; 2010], in addition an *in vitro* study reported aberrantly low intracellular pH in CFS/ME compared to control myoblasts [Boulton, 2012].

### 3.1.1 Free-radical overview

Oxygen-derived free radicals are produced as a result of a one electron reduction in molecular oxygen resulting in an extremely unstable configuration. Radicals rapidly react with other radicals or molecules to achieve stability (Displayed in Figure 3.7). Oxygen is a di-radical possessing two unpaired electrons. When oxygen gains an additional electron O$_2^-$ is produced which is the precursor to other ROS and RNS. For example, the reaction of O$_2^-$ with nitic oxide produces a highly aggressive oxidant, peroxynitrite (ONOO-) which acts to greatly limit nitric oxide availability. Furthermore, in aqueous solutions O$_2^-$ may also steadily dismutate to hydrogen peroxide (H$_2$O$_2$), which is catalysed by superoxide dismutase (SOD) in the body. Finally, in the presence of trace metals, including copper and iron, O$_2^-$ and H$_2$O$_2$ can produce hydroxyl radical ·OH which rapidly reacts with a wide variety of biological molecules. The hydroxyl radical has a particularly high affinity for lipid and is associated with lipid membrane degeneration via lipid peroxidation [Brandes and Janiszewski. 2005].

In terms of the function of ROS, under normal physiological conditions the cellular concentration of ROS is maintained at a constant level with this balance modulated by processes that both produce and eliminate free radicals. Effectively, the source of ROS can be separated into two categories. Firstly processes that release ROS through normal physiological processes as a waste product such as mitochondrial oxidative phosphorylation. Secondly, processes that generate ROS purposefully, in response to xenobiotics, cytokines and bacteria invasion, either through molecular synthesis of breakdown and as part of a signal transduction pathway, or functioning as part of cell defence mechanism [Finkel et al.2011; Zhang et al. 2016].

In terms of the impact of ROS on skeletal muscle cells, ROS have been reported to induce and promote mitochondriogensis a key factor in muscle differentiation. This has been
reported to occur via peroxisome proliferator-activated-receptor-gamma-coactivator-1α (PGC-1α) activated signal transduction pathway. However, in excess ROS may target mt DNA effectively shutting down myogenic differentiation [Rochard and Sejtili, 2000]. The occurrence of each of these processes is determined by the level and duration of ROS targeting muscle cells, the antioxidant status of the cell (SOD content) and the DNA repair capability of the cell. The differentiating stage of the muscle cell (satellite cell, differentiating myoblast or myotube) also is capable of redirecting the cell through different signalling pathway and further modifying cellular response to limit damage [Barbieri et al. 2012]. Therefore, it would appear that while ROS generation can promote muscle cell differentiation and repair, in excess it exhibits the capacity to be highly detrimental to muscle cell growth and development [Barbieri and Sejtili, 2012].

![Stable Molecule](image)

**Figure 3.7** Free radicals are atoms that contain an unpaired electron in their outermost shell, resulting in a highly unstable configuration and radicals rapidly reacting with other molecules and radicals to achieve stable configuration.

### 3.1.2 \( \text{O}_2^\cdot \) Detection techniques

#### 3.1.2.1 Spectroscopic \( \text{O}_2^\cdot \) measurement

Spectroscopic techniques have been used to measure \( \text{O}_2^\cdot \) flux in real-time [Tarpey and Fridovich. 2001]. One method being the reduction of cytochrome c, which has been used to measure the rate of \( \text{O}_2^\cdot \) generation from enzymes, tissue extracts and whole cells [Tarpey and Fridovich. 2001]. In terms of the reaction, it occurs at a constant of \( 1 \times 5 \times 10^5 \text{mol/L}^{-1} \), pH8 and at room temperature, absorbance is then measured spectrophotometrically at \( \lambda \text{550nm} \)
[Land and Swallow. 1971; Ballou et al. 1969]. During the reaction, ferricytochrome c is reduced to ferrocytochrome c by receiving an electron from O$_2^\cdot{}^-$. When ferricytochrome c is reduced its spectrophotometric absorbance is altered in a specific manner, with absorbance at 550nm increased, whereas at $\lambda$540nm and $\lambda$560nm absorption remains unchanged and serve as isobestic points [Dikalov et al. 2007].

The reduction of cytochrome c is considered by many researchers to be the ‘gold standard’ technique for the detection of O$_2^\cdot{}^-$. However, while the technique is effective in measuring O$_2^\cdot{}-$ when it is present in large amounts e.g. during respiratory burst in neutrophils and in isolated enzyme reactions, it may not be as effective at detecting O$_2^\cdot{}-$ in tissues that produced lower levels such as endothelial and muscle cells [Dikalov et al. 2007]. Tarpey and Fridovich [2001] outlined further precautions to consider when using the technique to measure O$_2^\cdot{}-$.

For example, tissue extracts contain compounds such as ascorbate and glutathione that can reduce cytochrome c, in addition to reductases that enzymatically catalyse cytochrome c reduction. What is more, reduced cytochrome c can be reoxidised by cytochrome oxidases, peroxidases and oxidants, which include H$_2$O$_2$ and ONOO$^-$. This reoxidation by diminishing apparent rates of cytochrome c reduction, which can lead to the underestimation of O$_2^\cdot{}-$ formation.

Nitroblue tetrazolium (NBT) is another technique commonly applied in spectroscopic O$_2^\cdot{}-$ detection and is based upon the reduction of NBT dye in the presence of O$_2^\cdot{}^-$. [Brandes and Janiszewski. 2005]. A limitation associated with this technique is that the reduction of NBT is not isolated to O$_2^\cdot{}^-$. Other substances including cellular reductases can donate an electron to NBT, forming the NBT radical. The major problem is that under aerobic conditions the NBT radical is capable of reacting with environmental O$_2$, which acts to generate O$_2^\cdot{}-$ artificially [Warwar et al. 2011; Tarpey and Fridovich. 2001].

The NBT radical intermediate can react with molecular oxygen under aerobic conditions which acts to generate O$_2^\cdot{}-$ artificially, which further reduces NBT and a false positive result [Warwar et al. 2011].

### 3.1.2.2 Electron spin resonance spectroscopy

Electron spin resonance spectroscopy (ESR) also referred to as electron paramagnetic resonance is a technique that detects paramagnetic species with one or more unpaired
Electrons [Hogg. 2010]. ESR exhibits a number of benefits when compared to other superoxide detection techniques because of its unique ability to detect either short or long lived free radicals with specificity and sensitivity [Tarpey and Fridovich. 2001]. ESR is capable of taking advantage of the paramagnetic state of O$_2^-$, however the low steady-state concentration, short lifetime as well as the rotational angular momentum of the relatively small diatomic molecule limits the ability to direct detection, requiring the need for spin-trapping [Warwar et al. 2011]. Spin traps are molecules that react with O$_2^-$ to produce a relatively stable paramagnetic species [Warwar et al. 2011; Khan et al. 2003]. The spin-trapping technique utilises a nitrone or nitroso compound to react with a free radical to produce a nitroxide (spin adduct), which exhibits substantially greater stability than the parent free radical [Ledoux et al. 2004; Gornicki et al. 2001]. Traditionally, the most common spin trap used for superoxide detection has been O$_2$-5,5 dimethyl-1-pyrroline-N-Oxides (DMPO). However, DMPO-OOOH (DMPO/O$_2^-$) adduct is incredibly unstable and decays rapidly with a half-life of approximately 60 seconds at pH7. Due to this limitation, more recently other spin-traps have been synthesised including ester group 5-tert-butoxycarbonyl-5-methyl-pyrrolline-N-oxide (BMPO). BMPO exhibits an increased rate constant for spin trapping O$_2^-$ compared to DMPO and BMPO-OOH exhibits a greater half-life than DMP-OOH [Rana et al. 2010].

Nevertheless, even with improvements in spin-trapping capability, due the low-levels of O$_2^-$ in vivo makes detection via ESR techniques incredibly difficult. Furthermore, ESR has been primarily used to measure O$_2^-$ in cell lysates and purified proteins and lacks the ability to detect O$_2^-$ in whole cell systems [Rana et al. 2010; Tarpey and Fridovich et al. 2001]

3.1.2.3 Amperometric extracellular O$_2^-$ monitoring

Electrochemical techniques for the detection of free radicals exhibit a key benefit over other free radical detection methods by enabling the direct, real-time measurement of free radicals within a biological system, with negligible disruption to the sample throughout experimentation [McNeil and Manning. 2002]. Amperometric O$_2^-$ detection devices are a type of electrochemical sensor, which continuously measure current resulting from redox reactions occurring at the electrode surface [Grieshaber et al. 2008]. The electrodes that are commonly used are produced from conductive materials such as gold, platinum and graphite
In the 1970s it was realised that electrode surfaces following functionalisation were capable of interacting with proteins, to provide stable and direct electrochemistry that was not disturbed by artefacts [McNeil and Manning, 2002; Armstrong et al, 1990]. Cooper et al. [1993] first described the use of a gold electrode with immobilised cytochrome c on the surface. The covalent attachment of cytochrome c on the modified surface was achieved through a carbodiimide initiated condensation reaction, with cytochrome c forming an integral part of the amperometric $O_2^-$ sensor.

Manning et al. [1998] described a simplified protocol for cytochrome $c$ immobilisation at a gold electrode surface in comparison to previously described methods [Cooper et al. 1993]. In this method the $O_2^-$ electrode was functionalised via a two-step process (See Figure 3.8)

Firstly, the link molecule 3-3'-dithiobis (sultosuccinimidyl-propionate) (DTSSP) was attached to the gold surface of the electrode, which was possible as DTSSP possesses a disulphide group to enable covalent attachment to gold surface. Once attached the molecule exhibits two carboxyl groups which undergo amide linkage with cytochrome $c$. The electrode was then poised at a working potential of +100mV verses a silver/silver chloride reference electrode [McNeil and Manning, 2002]. In a study conducted by Tammeveski et al. [1998] the authors reported the use of a gold surface electrode functionalised with cytochrome $c$ to be effective in detecting $O_2^-$ generated through the enzymatic breakdown to xanthine to uric acid by xanthine oxidase. Furthermore, Manning et al. [1998] utilised the cytochrome $c$ functionalised electrode to measure in vitro extracellular flux of $O_2^-$ from phorbol-12 myristate-13 acetate (PMA) stimulated astrocytes. This demonstrated the ability of the technique to measure $O_2^-$ production...
directly and in real-time from a cellular system. More recent studies have also reported cytochrome c functionalised electrodes to be effective in measuring O$_2^\cdot$ generated in live cells as well as isolated mitochondria [Aitken et al. 2007; Henderson et al. 2009]. Other functionalisation methods have been reported, for example, Shleev et al. [2006] described an O$_2^\cdot$ electrode that utilised azurin rather than cytochrome c. However, this method exhibited a crucial limitation in that the stability period of the electrode was reduced when compared to that described by Manning et al. [1998].

Recent studies have also reported cytochrome c functionalised electrodes to be effective in measuring O$_2^\cdot$ generation in various in vitro biological systems [Aitken et al. 2007; Henderson et al. 2009; Manning et al. 2001]. Other functionalisation methods have been reported, for example, Shleev et al. [2006] described an O$_2^\cdot$ electrode that utilised azurin rather than cytochrome c. However, this method exhibited a crucial limitation in that the stability period of the electrode was reduced when compared to that described by Manning et al. [1998]. Despite a more prolonged functionalisation process.

### 3.1.3 Hypotheses

The aim of the chapter was to examine O$_2^\cdot$ generation in CFS/ME patient myoblasts, performed following chemical stimulation with ethanol or incubation with lactic acid.

Specific experimental hypothesis were:

1. CFS/ME myoblasts exhibit elevated O$_2^\cdot$ generation following ethanol stimulation, compared to controls.

2. O$_2^\cdot$ elevated in CFS/ME patient samples compared to controls following incubation with lactic acid.
3.2 Materials and Methods

CFS/ME and control myoblast samples were provided by Dr Audrey Brown from the diabetes research group, Newcastle University. In this chapter the myoblast samples were derived from a female CFS/ME patient aged 65 years and a female control participant aged 66 years.

Tissue culture reagents were supplied by Sigma-Aldrich (Poole, Dorset), apart from Ham’s F10 growth medium (Lonza, SLS, East Riding, Yorkshire) and chick embryo extract (Sera Lab, West Sussex, UK).

Plastic consumables such as tissue culture flasks, serological pipettes, 24-well plates, falcon tubes and cell scrapers were supplied by Greiner Bio-one (Stonehouse, UK).

Superoxide dismutase (SOD), Xanthine, Xanthine oxidase (XOD), cytochrome c and lactic acid were purchased from Sigma Aldrich. DTSSP was supplied by Pierce (Chester, UK). Ethanol was purchased from Fisher (Cramlington, UK).

3.2.1 Cell culture and preparation

CFS/ME and control muscle samples were collected and processed as described in chapter 2. Myoblast samples were cultured in Ham’s F10 growth medium, which the addition of 20% FBS, 2% chick embryo extract, 500U/mL penicillin streptomycin and 1% amphotericin B. Cells were grown to 80% confluence prior to experimentation in a T75 flask. Cells were maintained in a humidified incubator at 5% (v/v) CO$_2$ at 37°C.

3.2.1.1 Direct real-time electrochemistry

3.2.1.2 Preparation of superoxide specific electrode

The electrode was prepared as detailed by Manning et al. [1998]. A 2 mm O.D. solid gold electrode (BioAnalytical Systems, Cambridgeshire, UK) was polished with a 0.2μm aluminium oxide slurry bound to a micro-cleaning cloth (BioAnalytical Systems, Cambridgeshire, UK). The electrode surface was then cleaned by sonicating in 100% ethanol for 5 minutes, before rinsing with deionised water. Subsequently, the electrode was incubated
with 50mM of the thiol linker DTSSP and incubated for 5 minutes at room temperature before rinsing with deionised water. The electrode was then treated with a 2mM of cytochrome c prepared in PBS and incubated over night at 4°C. Prior to experimentation, the electrode was once again rinsed with deionised water to remove any unbound protein.

3.2.1.3 Electrode calibration

An enzyme-based reaction (Xanthine/ XOD) was used to calibrate the electrode. See Figure 3.9.

Throughout the calibration the electrode was poised at +100mV and used in conjunction with an Ag/AgCl electrode which acted as a combined reference and counter electrode. The working electrode and the Ag/AgCl reference electrode were both placed in a 24-well plate containing 1.5mM xanthine in PBS. Following the achievement of a stable baseline varying concentrations of XOD (0,0.25, 0.5, 0.75, 1 U/mL) were added to the reaction volume and the resultant change in current was observed. To ensure XOD was equally and rapidly distributed throughout the well a magnetic based stirring system was applied. To test the specificity of the electrode to \( \mathrm{O}_2^- \) the scavenger superoxide dismutase (SOD) radical was added to the well at a concentration of 500U/mL to quench any responses at the electrode surface. Measurements were obtained using a 2-channel potentiostat and data was acquired using UiEchem software (Buxton, UK).
Following routine culture cells were seeded into a clear, 24-well plate at a density of $1 \times 10^5$ cells per well and placed in an incubator overnight to attach. Cells were maintained in a humidified environment at 5% CO$_2$ at a temperature of 37.5°C.

During experimentation the working and reference electrode were connected to the potentiostat and suspended just above the cell later. A steady baseline current was achieved before adding 50μL of PBS (control), or varying concentrations of ethanol (2-5%). The chemicals were used to induce the generation of O$_2^-$ radicals by the myoblasts. Current was produced at a rate that was directly proportional to the rate of O$_2^-$ production by the cell. The specificity of the electrode response to O$_2^-$ was assessed by using the superoxide specific scavenger, SOD (500 U/mL) was added to scavenge superoxide after a clear maximal current value had been obtained following the addition of the chemical stimulant.

### 3.2.1.4 Cellular superoxide measurement following ethanol stimulation of myoblasts

CFS/ME and control myoblasts were seeded into 24-well testing plates at a density of $1 \times 10^5$ cells per well in triplicate and were allowed to attach overnight. The following day, ethanol was added to the wells of the testing plate at the concentration 2, 3, 4 and 5%. To confirm the presence of superoxide the enzyme superoxide dismutase (SOD) was subsequently added at a concentration of 500U/mL to scavenge any superoxide radicals present.

### 3.2.1.5 Cellular superoxide measurement following acidification of myoblasts

CFS/ME and control primary myoblasts were seeded into 24-well testing plates at a density of $1 \times 10^5$ per well in triplicate. Following overnight attachment, cell culture media was acidified by adding 2mM lactic acid directly to each well, which corresponded to pH 5.36 (determined prior to the experiment using a pH meter (Hanna, Bedfordshire, UK)). Cells were incubated in a humidified environment at 5% CO$_2$ and at a temperature of 37.5°C for 30 minutes, 5 hours and 24 hours.

Following achievement of a maximal response (indicated by a stable plateau in current values) 500U/mL of SOD was added to the well to scavenge any superoxide radicals.
3.2.2 Cell viability in response to lactic acidification

The role of lactic acidification as a potential stimulator of $\text{O}_2^-$ generation was investigated in this chapter. Therefore, it was necessary to assess the cytotoxicity of lactic acid following 24-hours incubation. Briefly, myoblasts were seeded at a density of $1 \times 10^4$ cells per well and allowed to attach overnight. The following day, growth medium was removed, cells were washed once with PBS and replaced with DMEM and varying concentration of lactic acid (0.5mM-2mM). The plate was then incubated for 24-hours.

3.2.3 Data analysis

Minitab statistical software was the statistical model used to examine the data. The descriptive statistics used were mean ± SD. Inferential statistics were used to investigate CFS/ME and control $\text{O}_2^-$ generation in response to ethanol stimulation and lactic acidification. Specifically, the statistical tests used were paired samples T-test and independent T-test. Significance was accepted at the P<0.05 level.
3.3 Results

3.3.1 Direct real-time electrochemistry

3.3.1.1 Xanthine/XOD calibration curve

Figure 4.0 demonstrated that current recorded at the electrode was directly proportional to the concentration of XOD added to the system (in the presence of excess xanthine) \( (R^2=0.9582) \).

![Superoxide calibration curve with varying concentrations of xanthine oxidase (0.25-1.0μM) in the presence of constant xanthine (1.5mM) produced a linear increase in current \( (R^2=0.9582, n=4 \pm SD) \)](image)

3.3.1.2 Cellular superoxide response to ethanol in myoblasts

Figure 4.1 (a) showed there to be a slight linear relationship \( (R^2 = 0.7007) \) between increased ethanol concentration and a greater recorded current in CFS/ME myoblasts. Similarly, Figure
4.1 (b) also demonstrated control myoblasts to exhibit a small linear increase ($R^2=0.6814$) in current with a rise in ethanol concentration. Additionally, responses were quenched with (500U/mL) SOD which was added directly to the well, confirming the presence of $O_2^-$. There was no significant difference in current for CFS/ME compared to control myoblasts at any of the ethanol concentrations tested when each sample. Statistical analyses were performed by comparing triplicate measures for CFS/ME and control samples for each of the ethanol concentrations.

![Figure 4.1 (a): Superoxide generation in CFS/ME myoblast cells following stimulation with varying concentrations (2-5%) of ethanol. There was a small linear relationship between ethanol concentration and recorded current ($R^2=0.7007$). The response was inhibited with 500U/mL of SOD. Data presented as Mean ± S.D of 3 replicate measures, n=2.](image-url)
3.3.1.3 Superoxide generation following acidification in myoblasts

Figure 4.2 showed acidification of the medium to pH 5.36 (2mM lactic acid) over-time to induce no significant difference in superoxide generation (nA) in either CFS/ME or control myoblasts at any of the time points.

Figure 4.2: Lactic acidification (2mM, pH 5.36) over time (30 minutes-24 hours) did not induce any meaningful change in superoxide generation over-time in CFS/ME or control myoblasts samples. Data presented as Mean ± S.D of 3 replicate measures, n=2.

Figure 4.1 (b): Superoxide generation in CFS/ME myoblasts cells following stimulation with varying concentrations (2-5%) of ethanol. There was a small linear relationship between ethanol concentration and recorded current ($R^2=0.6814$). The response was inhibited with 500U/mL of SOD. Data presented as Mean ± S.D of 3 replicate measures, n=2.
3.3.2 Cytotoxicity with lactic acidification

Figure 4.3 demonstrated both CFS/ME and control myoblast cell viability measured with 0.03% resazurin following 24-hour incubation with varying concentrations of lactic acid (0.5mM-2mM). CFS/ME cells exhibited significantly greater viability when compared to control cells following 2mM acidification (pH 5.36 (P<0.0005), 1.5mM (pH 6.11 (P<0.0005) and 1mM (pH 6.61 (P<0.005). There was no significant difference following 0.5mM (pH 7.09) acidification.

![Graph showing viability over lactic acid concentration](image)

Figure 4.3: Lactic acidification of CFS/ME and control myoblasts over 24-hours resulted in significantly lower viability in control compared to CFS/ME cells following treatment with 1mM, 1.5mM and 2mM Lactic acid (pH 6.61, 6.11 and 5.36 respectively). There was no significant difference between CFS/ME and control cells following 0.5mM acidification (pH 7.09). Data presented as mean ±SD of 3 replicate measures, n=2. **** denotes P≤0.0005, *** denotes P≤0.005 as determined by independent t-test.
3.4 Discussion

Previous studies have confirmed CFS/ME patients to exhibit increased oxidative stress and decrease antioxidant status at rest, which was related to the extent of symptomology. In these studies blood oxidant markers such as TBARS and ascorbic acid were measured [Jammes et al. 2005; Fulle et al. 2000; Keenoy et al. 2001; Richards et al. 2000]. More recently, in vivo exercise based investigations have reported accentuated oxidative stress and altered muscle membrane excitability, which the authors suggested to explain muscle pain and PEM experienced by individuals with CFS/ME [Jammes et al. 2012; 2009 and 2005]. Elevated ROS has been reported to impact upon skeletal muscle function by inducing lipid peroxidation, which may lead to a loss of membrane excitability as a consequence of altered activation of K+ channels within skeletal muscle [Luin et al. 2011].

In this chapter, a direct, real-time approach to monitoring \( \text{O}_2^- \) generation in CFS/ME patient muscle samples was used. The electrochemical sensing technique was first reported by McNeil et al. [1992] and later developed by Manning et al. [1998]. Briefly, the method involved the covalent attachment of cytochrome \( c \) to the surface of a gold working electrode through surface modification with DTSSP. The generation of \( \text{O}_2^- \) by Xanthine/XOD resulted in the one electron reduction of cytochrome \( c^3 \) to cytochrome \( c^2 \), with the reduced protein then reoxidised at the electrode surface (poised at 100mV against Ag/AgCl reference electrode). The current rates recorded were found to be directly comparable to \( \text{O}_2^- \) production by XOD [Manning and McNeil. 2011]. The electrochemical technique has been reported to successfully detect \( \text{O}_2^- \) in a number of in vitro studies including, isolated mitochondria, cultured glial cells, human glioblastoma and B16 mouse melanoma [Henderson et al. 2009; Valverde et al. 1996; Manning et al. 2001; Manning et al. 1998].

In this chapter \( \text{O}_2^- \) generation was measured in CFS/ME myoblast samples following ethanol stimulation or lactic acidification. The work performed in this chapter did not confirm the presence of elevated \( \text{O}_2^- \) generation in CFS/ME muscle samples. Firstly, stimulation with ethanol was associated with a small linear increase in \( \text{O}_2^- \) generation in both CFS/ME and control samples, however no significant difference in \( \text{O}_2^- \) generation between sample groups was found (Figure 4.1 a and b). Secondly, following lactic acidification of the assay medium, \( \text{O}_2^- \) generation did not significantly differ between CFS/ME and control muscle samples (displayed in Figure 4.2). However, cytotoxicity work did reveal CFS/ME myoblasts viability to be significantly higher than controls following incubation with lactic acid (See Figure
4.3), suggesting CFS/ME myoblasts to be better able to buffer the effects of extracellular acidification.

### 3.4.1 Ethanol stimulation

In this chapter $O_2^-$ generation was not elevated in CFS/ME muscle samples following ethanol stimulation. It is difficult to contextualise this finding as presently no other studies have used electrochemical sensing approaches to directly measure $O_2^-$ in the muscle of CFS/ME patients.

However, this finding contrasts previous *in vivo* studies that have alternatively reported CFS/ME patients to exhibit enhanced oxidative stress at rest, which was accentuated during exercise [Jammes et al. 2012, 2009 and 2005]. Nevertheless, it is important to acknowledge the differing methodological approaches used. Free-radicals exist for a limited period *in vivo* and cannot be measured directly [Halliwell and Whiteman, 2004]. Therefore, previous studies have measured blood-based markers associated with oxidative stress. These have included TBARS and endogenous antioxidant such as RAA and erythrocyte reduced glutathione [Jammes et al. 2012, 2009]. These markers give an indication of the degree of systemic oxidative stress. Unlike, the aforementioned *in vivo* studies a key benefit of the electrochemical sensing approach reported in this chapter is its ability to directly measure $O_2^-$ at the cellular level. Nevertheless, it is important to recognise that *in vitro* effects may not be representative of the wider biological system as a whole. For example, *in vitro* cell systems lack the *in vivo* microenvironment, communication and direct contact with other cells and bioactive substances [Aas et al. 2013].

It is also important to note that previous studies measured blood oxidant markers at rest and following exercise, representing a normal biological state [Jammes et al. 2012, 2009]. However, in this chapter myoblast cells were chemically manipulated to induce oxidative stress. Ethanol is metabolised by the CYP2E1 enzyme, with this process recognised as a key generator of $O_2^-$ [Bansal et al. 2010; Gonzalez, 2007; Jimenez-Lopez and Cederbaum, 2005]. While the expression of CYP2E1 has been reported in the skeletal muscle of adults and children [Molina-Ortiz et al. 2013]. It is accepted that the enzyme is predominantly located in the liver and the levels found in the muscle are substantially lower [Jimenez-Lopez et al. 2005].
and Cederbaum. 2005]. Therefore, ethanol stimulation may not have been the most effective way to promote oxidative stress in skeletal muscle cells.

3.4.2 Extracellular acidification

This chapter also investigated O$_2^-$ generation in response to lactic acidification, to assess whether extracellular acidification impacted upon oxidative stress. As previously described, several in vivo studies have revealed excessive intramuscular acidosis in CFS/ME patients, suggestive of bio-energetic dysfunction and an over utilisation of the lactate dehydrogenase pathway. In addition, intracellular pH has also been reported to be significantly lower in CFS/ME myoblasts compared to controls [Boulton. 2012].

Nevertheless, in this chapter lactic acidification of the assay medium did not lead to increased O$_2^-$ generation in CFS/ME myoblasts when compared to healthy controls. Contrary, to previous investigations that have reported elevated blood-oxidant status in CFS/ME patients and concurrent muscle dysfunction [Jammes et al. 2012, 2009 and 2005]. Thus, extracellular acidification does not appear to induce oxidative stress.

However, it is important to note that chemical manipulation to induce extracellular acidification may not be representative of intracellular acidosis that has been reported in CFS/ME patients following exercise, which may alternatively be related to bio-energetic dysfunction [Jones et al. 2012 and 2010]. Therefore, in this chapter it would have been advantageous to simulate exercise in vitro through the EPS of muscle samples, to investigate the impact of muscle contraction on O$_2^-$ generation in CFS/ME. However, it was not possible to use EPS in conjunction with the electrochemical sensor as the device function to electrically stimulate muscle samples to contract. As the electrode measures current in the well it would have been difficult to decipher between artefacts due to EPS and changes in the current caused by O$_2^-$ generation.

Although, lactic acidification work did not impact upon O$_2^-$ generation, a cytotoxicity assay revealed CFS/ME myoblasts to exhibit greater viability than control cells following treatment with lactic acid. This was an interesting finding as it suggests the CFS/ME patient cells to tolerate substantial acidification, although it is difficult to determine the mechanism behind this occurrence. However, as previously described several studies have revealed profound
intramuscular acidosis in CFS/ME patients [Jones et al. 2012; 2010], it is possible as a consequence that cells have developed an improved buffering capacity to cope with excessive acidification.

3.4.3 Limitations

It is important to acknowledge a number of limitations associated with this technique. For example, the concentration of ROS at the site of production is very high. However, as $O_2^-$ was measured in the extracellular medium and was required to diffused out of the cell it is possible that the amount of $O_2^-$ measured by the electrode was not reflective of the intracellular environment. Additionally, the presence of trace metals in the extracellular medium may potentially function to decrease cellular $O_2^-$ generation. For example, copper has been reported to serve an essential co-factor for several oxidative stress related enzymes such as superoxide dismutase which quenches cellular $O_2^-$ generation [Tchounwou et al. 2012].

An enzyme based reaction (Xanthine/XOD) was performed to calibrate the $O_2^-$ electrode, this process was carried out in a stirred system to ensure equal diffusion of the enzyme and substrate throughout the well. However, when $O_2^-$ was measured in vitro, cells were attached to the surface at the bottom of the well. It was therefore not possible to use a magnetic stirrer without disturbing the cell layer. This was a limiting factor as following the addition of the chemical stimulants (i.e. ethanol/lactic acid) it was not possible to achieve equal diffusion across the well. It could have been possible to incorporate stirrers from above, however as the $O_2^-$ electrode also was positioned above the cell layer such equipment risked disturbing the electrode.

Additionally, although each well was seeded at the same density ($1 \times 10^5$ cells), following attachment cells did not provide even coverage across the well. Consequently, it was impossible to position the probe in an area of exactly the same coverage during each measurement, bringing the reliability of the technique into question.
3.4.4 Conclusion

The direct real-time electrochemical sensing approach used in this chapter enabled the measurement of $O_2^-$ generation in CFS patient myoblast cells. Contrary, to previous *in vivo* studies that have reported evidence of elevated oxidative stress in CFS/ME patients this study alternatively revealed CFS/ME patient myoblasts to exhibit $O_2^-$ generation comparable to healthy control cells following chemical stimulation. Therefore, oxidative stress does not appear to contribute towards the muscle dysfunction and fatigue phenotype associated with CFS/ME.
3.5 References

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Chapter 4

Application of 2’7’-bis (2-carboxyethyl)-5 (6) Carboxyfluorescein to Measure Intracellular pH in CFS/ME Myoblasts and Myotubes

4.1 Introduction

Chapter 2 investigated intracellular pH in CFS/ME myoblasts using a novel pH sensing nanosensor platform. However, the nanosensor system was unable to reliably determine the cytosolic pH. It was therefore determined that the technique required further development, which was beyond the scope of this thesis. The aim of this chapter was to investigate variations in cytosolic pH using the well characterised, pH responsive fluorophore 2’7’-bis (2-carboxyethyl)-5 (6) carboxyfluorescein (BCECF) in CFS/ME myoblasts and differentiated myotubes at rest and following electrical pulse stimulation (EPS).

Evidence exists to suggest CFS/ME patients exhibit profound acidosis following low-level exercise, when measured via MRS [Jones et al. 2010]. Additionally, pilot work using the fluorescent nanosensor system (detailed in Chapter 2) to measure intracellular pH showed that CFS/ME myoblasts exhibited a significantly lower pH at rest compared to non-diseased control cells. Further, this innate acidosis could be normalised following addition of the PDK inhibitor, DCA [Boulton. 2012]. This represented the first in vitro analysis of pH in myoblasts derived from CFS/ME patients and suggested that these cells were intrinsically acidic which could have a significant influence on the CFS/ME phenotype. Importantly, this acidosis could be reversed by PDK inhibition. This observation, if correct, could open up novel treatment strategies for CFS/ME patients. However, in the absence of subsequent reliable nanosensor data it was decided free fluorescent dye would be used in an attempt to measure intracellular myoblast pH and further examine these initial observations.

This chapter developed also upon Chapter 2 by investigating intracellular pH in myoblasts and differentiated myotubes. Multinucleated myotubes represent the best alternative to intact human skeletal muscle and have been reported to exhibit the morphological, metabolic and biochemical properties of intact adult muscle cells [Gaster et al. 2001; Henry et al. 1995; Nikolic et al. 2012].
4.1.1 Intracellular sensing

Techniques that can accurately and precisely measure intracellular pH under physiological conditions are highly advantageous as intracellular pH plays a pivotal role in cell function, with organelle metabolic processes directly affected by H$^+$ ion concentration [Han and Burgess. 2010; Orkan et al. 2002]. Even slight deviations in cytosolic pH (typically 7.2-7.4) are associated with a reduction in cell function, growth and division, which is observed in diseases such as cancer and Alzheimer’s disease [Izumi et al. 2003; Davies et al. 1993].

As discussed in Chapter 2 a number of pH intracellular sensing techniques exist, including the use of optochemical sensors, fluorescent dye, NMR, SERS and PEBBLE nanosensors. This chapter utilised the fluorophore 2’7’-bis (2-carboxyethyl)-5 (6) carboxyfluorescein (BCECF) to investigate intracellular pH. BCECF was introduced in 1982 [Rink et al. 1982] for the measurement of cytoplasmic pH and has since been widely used for pH measurement in mammalian cells, living tissues and individual organelles [Han and Burgess. 2010]. In terms of intracellular delivery, chemical modification of BCECF can occur to incorporate acetoxymethyl ester (AM), which is synthesised from carboxyfluorescein by the addition of two extra carboxylate groups via short alkyl chains. BCECF-AM is therefore a no charge, non-fluorescent form of BCECF. In terms of function, the presence of AM ester enables BCECF to rapidly diffuse through the cell membrane into the cell, once inside the cell hydrolysis of the acetyl ester linkage by enzymatic cleavage regenerates the less permeable and fluorescent original compound BCECF [Gdovin et al. 2010] (Displayed in Figure 4.4) which is capable of generating a fluorescent signal in relation to intracellular pH [Gdovin et al. 2010; Han and Burgess. 2010; Orkan et al. 2002].

![Figure 4.4: The synthesis and hydrolysis of AM and acetate ester as illustrated by Han and Burgess [2010].](Image)

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BCECF remains one of the most widely used fluorophores for intracellular pH measurement. Its popularity is due to it exhibiting a number of benefits when compared to other dyes, which were detailed in a review conducted by Han and Burgess [2010], regarding the use of fluorescent indicators for intracellular pH measurement. For example, the dye is retained well inside the cell as at physiological pH it exhibits 4-5 negative charges. BCECF also exhibits a pKₐ of 7.0 which makes it well suited for the measurement of cytosolic pH which is typically in the 7.2-7.4 range. Additionally, BCECF AM esters are membrane permeable enabling non-destructive dye loading into the cell. Furthermore, non-fluorescent BCECF-AM esters are converted into fluorescent BCECF acid in a highly efficient manner, so much so it may be used for cell viability assays [Perez-Sala et al. 1995]. Finally, BCECF is a ratiometric fluorophore, which enables fluorescence to be measure at two excitation wavelengths. For example, when excited at 440nm (isosbestic point) the dye is pH insensitive and when excited at 495nm the dye is pH sensitive [Gdovin et al. 2010]. Effectively, ratiometric dyes correct for a number of shortcomings inherently associated with the use of free-dye, including variable dye loading, dye leakage and detector sensitivity [Grant & Acosta.1997].

Although BCECF remains the most widely used intracellular pH indicator, other popular fluorophores for pH measurement include fluorescein and fluorescein derivatives (carboxyfluorescein). However, without additional modification these dyes are able to leak through the cell membrane allowing for erroneous pH measurement [Han and Burgess. 2010]. For example, at 37°C intracellular concentrations of 5- and 6-carboxyfluorescein have been reported to be decreased 30-40% after the first 10 min following washing [Rink et al. 1982]. Alternatively, although BCECF-AM has been reported to leach from the cell, the rate is around 10% 20 minutes after washing, which is a lot lower than other commercially available dyes. Therefore, it would seem that in terms of approaches to measure intracellular pH, BCECF-AM combined with fluorescent spectroscopy is a popular choice, exhibiting high sensitivity, good spatial and temporal observation of pH change as well as being relatively operationally simple [Han and Burgess et al. 2010].
4.1.2 Electrical pulse stimulation

In this chapter electrical pulse stimulation (EPS) was applied to enable the contraction of skeletal muscle cells in vitro as a strategy to investigate the effect of physical activity on intracellular pH in CFS/ME patient muscle samples. The EPS protocol that was used has previously been described by Brown et al. [2015] and EPS was performed using the C-pace EP cell culture pacer (Ion Optix, Dublin), Displayed in figure 4.5.

![EPS device](image)

Figure 4.5: EPS was performed using the C-pace EP cell pacer (Ion Optix, Dublin)

A number of studies have reported the use of EPS to induce muscle contraction in skeletal muscle cells. For example, Fujita et al. [2007] reported EPS to accelerate de novo sarcomere assembly, in addition to the induction of Ca\(^{2+}\) transients in C2C12 mouse skeletal muscle myotubes. Additionally, Nedachi et al. [2008] utilising the same experimental model reported EPS to induce AMPK activation, increase glucose transport and enhance the release of chemokines such as IL-6. Interestingly, other papers have detailed the use of EPS to investigate human skeletal muscle myotube function [Lambernd et al. 2012; Nikolic et al. 2012], with these studies reporting the model to enhance sarcomere assembly, increase AMPK activation, glycolysis and glucose uptake as well as chemokine expression. Interestingly, Brown et al. [2015] demonstrated EPS to simulate exercise in cultured CFS/ME and control myotubes. For example, the authors reported alternating frequency (low/high)
EPS to be successful in enabling muscle contraction and inducing metabolic changes associated with exercise. In healthy cells, EPS induced myotube contraction and increased AMPK activation whereas in the CFS/ME patient group AMPK activation and glucose uptake were impaired. The authors therefore concluded the *in vitro* use of EPS be effective strategy to investigate metabolic and bio-chemical exercise associated dysfunction in cultured cells.

The cytotoxic effects of EPS on skeletal muscle cells have also been investigated. For example, Nikolic et al. [2012] reported acute high frequency and chronic low frequency EPS to exhibit no significant alteration in lactate dehydrogenase [LDH] concentration, when measured in the experimental medium following 24 and 48-hours stimulation. Furthermore, Brown et al. [2015] also reported there to be no significant increase in the release of LDH into the assay medium following 24-hours EPS, in CFS/ME myotubes compared to controls. Therefore, suggesting both acute and chronic EPS of myotubes to not exhibit any notable toxic effect on skeletal muscle cells.

4.1.3 Hypotheses

The aim of this chapter was to investigate the intracellular pH of CFS/ME myoblasts and differentiated myotubes at rest and following (EPS) to mimic exercise. Intracellular pH was measured via fluorescent spectrophotometry following treatment with fluorophore BCECF-AM. The PDK inhibitor DCA was utilised to investigate whether PDC function impacts upon intracellular pH

The experimental hypotheses were:

(1) CFS/ME myoblasts and differentiated myotubes exhibit significantly lower intracellular pH at rest and following EPS, when compared to controls.

(2) Treatment with PDK inhibitor DCA normalises intracellular pH in CFS/ME samples at rest and following EPS.
4.2 Materials and Methods

CFS/ME and control myoblasts were supplied by Dr Audrey Brown from the diabetes research group at Newcastle University. Myoblasts were obtained from a total of 5 (4 females, 1 male) CFS/ME patients which consisted of 4 females and 1 male aged 43± 12.73 years. A total of 4 myoblast samples were obtained from control participants (3 females, 1 male) who were aged 50.14± 12.81 years.

All tissue culture reagents apart from Ham’s F10 growth medium (Lonza, SLS, East Riding, Yorkshire) and chick embryo extract (Sera Lab, West Sussex, UK) were purchased from Sigma-Aldrich (Poole, Dorset).

Plastic tissue culture flasks, serological pipettes, 6-well plates, falcon tubes and cell scrapers were purchased from Greiner Bio-one (Stonehouse, UK).

pH responsive dye BCECF-AM was purchased from Abcam (Cambridge, UK). Intracellular pH calibration kit was purchased from Thermo Fisher Scientific (Loughborough, UK). PDK inhibitor potassium DCA was purchased from Sigma-Aldrich.

4.2.1 Cell culture and preparation

As described in Chapter 2, CFS/ME and control muscle cell samples were collected and isolated as detailed in the study by Brown et al (2015). The myoblast samples were routinely cultured in Ham’s F10 medium, supplemented with 20% FBS, 2% chick embryo extract, 500 U/mL penicillin-streptomycin and 1% amphotericin B. Myoblasts were grown to 80% confluence in a T75 flask before being trypsinised and seeded into the vessel of choice. Cells were grown and maintained in a humidified incubator at 5% (v/v) CO₂ at 37° C.

Myoblasts were seeded into 6-well testing plates at a density of 2x 10⁵ cells per well and left overnight to attach. BCECF-AM pH calibration or experimentation was performed the following day. Growth media was removed and replaced with Minimum Essential Medium (MEM), which was supplemented with 2% (V/V) FBS and 1% Penicillin-streptomycin. MEM was replaced every 48-hours over a 7 day period.
4.2.2 Intracellular pH calibration

To enable the accurate determination of intracellular pH in cells treated with BCECF-AM an *In situ* pH calibration was performed for both myoblast and myotube samples. The calibration was conducted using reagents provided by a commercially available pH calibration kit (Thermo Fisher Scientific, Loughborough).

Myoblasts were seeded into 96-well testing plates at a density of 3x 10^4 cells per well and at 2x 10^5 cells per well in 6-well plates to promote differentiation into myotubes. Immediately prior to the calibration, cells were washed once with PBS. To ensure equilibration of extracellular and intracellular pH, cell-loading solution was prepared by dissolving ionophores nigericin and valinomycin in DMSO at a stock concentration of 100μM before adding them to each pH buffer solution (4-7) at final concentration in the well of 10μM at 37°C. Following 5-minutes incubation BCECF-AM was added to the wells at a final concentration of 2μM and further incubated at 37°C for 30 minutes, before FI measures were performed.

FI measurements were then acquired using the Tecan-I 200 fluorimeter and data was provided by using the compatible Magellan software. Fluorescence was determined by measuring excitation at 440nm (non-pH dependent) and 490nm (pH dependent), with emission detected at 535nm. FI data was then analysed to provide FI ratios, this was achieved by dividing the FI at λ490 by FI at λ440. A standard curve was then produced to display the relationship between pH and FI ratio. A polynomial second-order curve was then applied to determine the pH of cells treated with BCECF-AM.

4.2.3 Myoblast viability following treatment with BCECF-AM

It was necessary to investigate the biocompatibility of BCECF-AM before using the fluorophore to measure intracellular pH in skeletal muscle samples. To assess the level of cytotoxicity, varying concentrations of the pH responsive dye (0.5-2μM) were incubated with CFS/ME and control myoblasts. Briefly, myoblasts were seeded into 96-well plates at a density of 1x10^4 cells per well. BCECF-AM was then added to the plate and incubated for 3,6 and 24-hours. Viability was measured following the addition of resazurin (0.03%) and fluorescence was interpreted at λex=560nm, λem= 590nm, via the Tecan- I 200 fluorimeter.
4.2.4 BCECF-AM intracellular pH measurement

Intracellular pH measurements were acquired for myoblast and myotube samples following the internalisation of fluorophore BCECF-AM.

Prior to experimentation culture medium was removed and testing plates were washed once with PBS. Assay medium (DMEM) in addition to BCECF-AM was then added to the wells at a final concentration of 2µM. The plate was then incubated for 35 minutes at 37°C. The assay media was aspirated and the plate was washed once before adding fresh assay medium or assay medium supplemented with DCA (40µM).

Fluorescence was measured using the Tecan 2000 fluorimeter and measurement setting were $\lambda_{\text{ex}} 440\text{nm}$ (non-pH dependent) and $\lambda_{\text{ex}} 490\text{nm}$ (pH dependent) / $\lambda_{\text{em}} 535\text{nm}$.

4.2.5 Electrical pulse stimulation

EPS was performed to simulate exercise in CFS/ME and control myotube samples. The ability of EPS to model exercise in vitro has previously been confirmed, with a number of studies utilising the technique to investigate cellular metabolic function [Orfanos et al. 2016; Brown et al. 2015].

The EPS protocol that was used has previously been described by Brown et al. [2015]. EPS was performed using the C-pace EP cell culture pacer (Ion Optix, Dublin). Myoblasts were plated into 35mm culture dishes at a density of $2 \times 10^5$. The cells were differentiated for 7 days before stimulation was initiated for 24-hours at alternating frequencies, with one hour low frequency (5 volts, 24ms, 2Hz ) followed by one hour high frequency (5 volts, 24ms, 0.2Hz).

4.2.6 Data analysis

The statistical model used to analyse the data was Minitab 17 statistical software. Inferential statistics were carried out to examine intracellular pH in CFS/ME and control skeletal muscle
samples. The statistical tests used were the paired samples T-test and independent T-test and statistical significant was accepted at P<0.05. The descriptive statistics used were mean ± SD.
4.3 Results

4.3.1 BCECF-AM pH In Situ Calibration

Cultured primary myoblasts and differentiated myotubes were treated with pH responsive free-dye BCECF-AM (2µM). To ensure intracellular and extracellular pH equilibration cells were incubated with varying pH buffer solutions (4-7), which were doped with ionophores valinomycin and nigericin (10µM). FI measurements were obtained and ratios were calculated (490/440nm) for all samples. A second-order polynomial curve fitting was applied to generate the calibration curve. The polynomial fitting was successful in determining the buffer pH within the pH range tested in both myoblast (y=0.3758x^2 - 3.4706x + 9.4155, R^2=0.999, n=3) and myotube (y=0.4119x^2 - 3.8228x + 10.10^2, R^2=0.995, n=3) cell samples. Data presented for myoblast samples in Figure 4.6 and myotube samples in Figure 4.7.

![Figure 4.6: In situ calibration curve conducted in myoblast samples following incubation with BCECF-AM. A polynomial second-order fitting enabled the successful determination of pH within the physiological range from the FI ratio measures provided (y=0.3758x^2 - 3.4706x + 9.4155, R^2=0.999). The data is presented at mean ±SD (n=3).](image-url)
4.3.2 Myoblast viability following exposure to BCECF-AM

Figure 4.8 a, b and c demonstrated CFS/ME and control myoblast viability following incubation with BCECF-AM for 3, 6 and 25-hours respectively. Viability remained high for both samples across all time-points and dye concentrations (>70%). No significant differences in viability were exhibited (% untreated control) between samples, across time-points or as a result of varied BCECF-AM concentration.

![Graph showing myoblast viability (%)](image)

Figure 4.8 (a): CFS/ME & control myoblast viability after 3-hours incubation with BCECF-AM at 0.5-2μM concentrations. No significant differences in viability was observed between samples or with varying concentrations of BCECF-AM. Data presented as mean ±SD, n=4)
Figure 4.8 (b): CFS/ME and control myoblast viability after 6-hours incubation with 0.5-2μM BCECF-AM concentrations. No significant differences in viability was observed between samples or with varying concentrations of BCECF-AM. (Data presented as mean ± SD, n=4)

Figure 4.8 (c): CFS/ME and control myoblast viability after 24-hours treatment with 0.5-2μM concentrations of BCECF-AM. No significant differences in viability was observed between samples or with varying concentrations of BCECF-AM. (Data presented as mean ± SD, n=4)
4.3.3 Myoblast intracellular pH determination with DCA treatment

Figure 4.9 demonstrated intracellular pH measurements for CFS/ME (n=5) and control (n=4) myoblast samples following the internalisation of free-dye BCECF-AM (2μM). Intracellular pH did not significantly differ between the CFS/ME and control group samples at baseline or following DCA treatment across all time-points. In terms of within group effects, DCA did not induce any significant alteration in pH in both samples across any of the time-points.

Figure 4.9: CFS/ME & control myoblast intracellular pH measurements following BCECF-AM internalisation. No significant differences in pH were observed between CFS/ME & control samples at baseline or following incubation with DCA, across any of the time-points. Additionally, no significant within group differences in pH for both samples were exhibited following treatment with DCA (40μM). Data presented as mean ± SD, n=9

Figure 5.0: (a) myoblasts in culture, (b) Myotubes 7-days differentiation
4.3.4 Myotube intracellular pH measurement at rest and post-EPS

The intracellular pH of CFS/ME (n=4) and control (n=4) myotubes was investigated at rest, following 24-hours EPS and in conjunction with DCA treatment. In resting samples, the intracellular pH of both CFS/ME and control samples did not significantly differ at baseline or following incubation with DCA (40μM). Moreover, there was no significant difference in the intracellular pH measurements within sample groups at baseline or following DCA treatment, displayed in Figure 5.1 a.

In terms of EPS, myotubes were stimulated for 24-hours at alternating frequencies. When compared to resting pH measurements there was no significant alteration in pH immediately post-exercise or throughout the recovery period for CFS/ME (see Figure 5.1 a and b) and control samples (see Figure 5.1 a and c) and DCA did not exhibit any effect on pH in either sample at any of the time points measured (CFS/ME samples see Figure 5.1a and 5.1b, Controls see figure 5.1 a and c). Furthermore, post EPS there was no significant difference in pH following DCA treatment compared to baseline for CFS/ME (see Figure 5.1c) and control samples (see Figure 5.1 c) immediately post-exercise and throughout recovery. Finally, when comparing CFS/ME to control samples there was no difference in the pH exhibited immediately post-EPS or throughout recovery at baseline and with DCA treatment (see Figure 5.1b and 5.1c).
Figure 5.1a: CFS/ME (n=4) & control (n=4) resting myotube pH at baseline and following treatment with DCA (40µM). No significant difference in pH between groups at baseline or following DCA treatment. DCA did not exhibit any within group differences in pH when compared to baseline for both samples. Data presented as mean ±SD, n=8.

Figure 5.1b: CFS/ME myotube intracellular pH measurements post-EPS stimulation at baseline and following treatment with DCA (40µM). DCA did not induce any significant alteration in pH at any point during the recovery period. Data presented as mean ± SD, n=4.
Figure 5.1 c: Control myotube intracellular pH measurements post-EPS stimulation at baseline and following treatment with DCA (40µM). DCA did not induce any significant alteration in pH at any point during the recovery period. Data presented at mean ± SD, n=4
4.4 Discussion

Previous research has described intracellular pH abnormalities in the muscle of patients with CFS/ME. In a MRS study conducted by Jones et al. [2012] patients were reported to exhibit profound and sustained acidosis, following a relatively low-level repeat exercise protocol. The authors postulated the findings to be evidence of underlying bio-energetic abnormality, due to an over-utilisation of the lactate dehydrogenase energy-producing pathway.

Intracellular pH dynamics were investigated in more detail in pilot work performed by Boulton [2012]. As described in Chapter 2, a novel pH sensing nanosensor system was utilised to investigate intracellular pH in patient myoblast samples. The primary finding was the presence of aberrantly low intracellular pH in CFS/ME myoblasts compared to controls. Secondly, intracellular pH was normalised to a level comparable to control samples following treatment with the PDK inhibitor DCA. These findings suggested bio-energetic abnormality in CFS/ME patient muscle as a consequence of impaired PDC function.

A pivotal aim of Chapter 2 in this thesis was to repeat the preliminary work of Boulton [2012], as well as further validating and developing the pH nanosensor system. However, it was not possible to reproduce the findings and the intracellular pH of CFS/ME myoblasts was not found to be significantly lower than controls. Additionally, DCA did not exhibit any significant effect on intracellular pH in either CFS/ME or control myoblast samples. Moreover, the pH measurements obtained for both myoblast samples were outside the cytosolic pH range and were suggestive of endosomal/lysosomal intracellular pH localisation. Work presented here therefore suggested that rather than being free to passively register pH changes in the cytoplasm, nanosensors may have been sequestered into subcellular organelles. Interestingly, this finding is in agreement with others who have also reported a low intracellular pH measurement following the internalisation of polyacrylamide nanosensors albeit in alternative cell-lines [Benjaminson et al. 2011; Coupland et al. 2009].

A primary aim of this chapter was to therefore utilise an alternative intracellular sensing technique to investigate the pH dynamics in CFS/ME skeletal muscle samples. In addition, as Jones et al. [2012] reported CFS/ME patients to exhibit acidosis following low-level exercise, EPS was used to stimulate muscle cells in an attempt to mimic physical activity in an in vitro setting. Specific aims were:
a) To measure the intracellular pH of CFS/ME myoblasts at rest and differentiated myotubes at rest and following electrical pulse stimulation (EPS).

b) To investigate the ability of PDK enzyme inhibitor DCA to impact upon intracellular pH at rest and post-EPS in myoblasts and differentiated myotube samples.

The results of this chapter showed that there was no significant difference in intracellular pH levels in CFS/ME myoblasts compared to controls (Figure 5.0a). Similarly, pH did not significantly differ between CFS/ME and control myotubes at rest and post-EPS (Figure 5.0b and 5.0c). Moreover, treatment with DCA did not significantly alter intracellular pH at rest in myoblast samples or at rest and following EPS in myotube samples (Figure 4.9 and 5.0a). These findings contrast the experimental hypotheses detailed in Section [4.1.3].

4.4.1 Optimisation

The toxicity profile of BCECF-AM was investigated before the dye was utilised to determine intracellular pH. Encouragingly, the pH responsive dye was well tolerated in the myoblast samples and cell viability remained high ( >70%), even following 24-hours incubation (Figure 4.8c). BCECF-AM is a derivative of fluorescein which has also been reported as a low-toxic compound [Richoz et al. 2013]. Alternatively, in high doses the dye purportedly exhibits a toxic effect in cell culture and in vivo [Alford et al. 2009]. BCECF-AM was deemed suitable to apply to the in vitro assessment of myoblast and myotube intracellular pH.

In terms of additional optimisation work, two in situ calibration curves were produced for both myoblast and myotube cell samples. BCECF-AM was successfully calibrated over the physiological pH range ( Figure 4.6 and 4.7).

4.4.2 Intracellular pH determination

CFS/ME resting muscle samples did not exhibit lower intracellular pH when compared to control samples which contrasts previous work [Boulton. 2012], which reported aberrantly
low pH in CFS/ME myoblasts, which was subsequently normalised to control level following treatment with DCA.

A possible reason for the discrepancy may relate to the varying sample sizes utilised. In the present study, n=5 CFS/ME myoblasts and n=4 control samples were investigated, compared to n=1 CFS/ME and n=1 control in the Boulton [2012] study. It is important to realise that failing to test multiple CFS/ME patient samples is methodological weakness as the sample tested may not be representative of the wider CFS/ME patient population. Furthermore, distinct sub-groups within patient population have been reported in terms of intracellular acidification following exercise. For example, Barnes et al. [1993] measured intramuscular pH via 31pMRS in 46 CFS/ME patients. Results demonstrated no significant abnormality in pH when the group were taken as a whole following exercise, however 6 patients did exhibit intramuscular acidification. Considering the heterogeneity of the CFS/ME patient population it is clear that the study conducted by Boulton [2012] was statistically underpowered due to low sample size, effectively low statistical power negatively impacts the likelihood of a nominally statistical significant finding reflects a true effect [Button et al. 2013]. Thus, any conclusions drawn from the data must be interpreted with caution.

An additional aim of the present chapter was to investigate the impact of physical activity on intracellular pH in an in vitro muscle culture system. This was achieved through the application of EPS and myotubes were stimulated for 24-hours at alternating high and low frequencies to produce skeletal muscle contraction and to induce metabolic changes associated with physical activity. Results demonstrated EPS to exhibit no significant effect on either CFS/ME or control myotube intracellular pH when each group when post-EPS measures were compared to rest [Figure 5.0a, b and c]. In addition, no difference in pH was observed between CFS/ME and control myotubes immediately post-EPS or during the recovery period [Figure 5.0b and c]. The findings contrast with earlier in vivo studies, which demonstrated when exercising to a comparable level to control participants CFS/ME patients exhibited significant intramuscular acidosis and a 4-fold increase in the time taken to recover from acidosis [Jones et al. 2010; 2012]. Nevertheless, in agreement the work presented here other studies have reported limited evidence of reduced intramuscular pH in patients [Barnes et al.1993; Wong et al. 1992; Lane et al. 1998; 1998b]. For example, Wong et al. [1992 ] reported there to be no difference in pH at rest or the recovery phase following graded exercise to exhaustion and measured by NMR. Similarly, Barnes et al. [1993] found no
Evidence of abnormality at rest or following exercise when CFS/ME patients (n=46) were taken as a whole. However, 6 patients were reported to exhibit increased acidification in response to exercise. Likewise, in studies conducted by Lane [1998a;1998b], following sub-anerobic threshold exercise CFS/ME only 8% of the group were reported to exhibit an elevated blood lactate response. All the studies compared CFS/ME patients to control participants and measured intramuscular pH via NMR or MRS. Importantly, the contrasting findings illustrate the heterogeneity of the CFS/ME patient population and the differing response to exercise.

It is important to note that contrasting patient bio-energetic responses to exercise exhibited with in vivo studies may be due to the level of patient engagement, which is a key limitation of that mode of investigation. For example, Jones et al. [2012] reported CFS/ME patients to fall into 2 distinct categories in relation to Phosphocreatine (PCr) depletion in response to exercise. The first group demonstrated normal bio-energetic dysfunction, exhibiting PCr depletion to a comparable level to controls when exercising at the same level of MVC. Conversely, the second group exhibited low-level PCr depletion and no exercise induced acidosis as a consequence. The authors postulated this effect to be evidence of some form of exercise avoidance behaviour. In contrast, the in vitro exercise model utilised in the present study eliminates the need for patient compliance in the exercise protocol and enables the bio-energetic function of all patient muscle samples to be examined equally following the same EPS strategy.

Nevertheless, it is important to acknowledge limitations associated with in vitro bio-energetic assessment for example, culture conditions are not homeostatic as there is a continuous depletion of nutrients and the generation of waste products within the culture medium. Additionally, oxygen supply is not always sufficient with dissolved oxygen typically consumed during the first few hours or medium replacement resulting in a diffusion limited supply which can induce anaerobic culture conditions (glycolysis) [Hartung and Daston, 2009; Coecke et al. 2006] Furthermore, while myotubes have been reported to closely resemble mature human skeletal muscle in terms of morphological, biochemical and metabolic characteristics [Olsson et al. 2015; Aas et al. 2013; Berggren et al. 2007]. However, when compared to intact skeletal muscle, myotube contractile force during EPS has been reported as lower. In addition, fused tetani is achieved at a reduced stimulating frequency and kinetic parameters are slower [Madden et al. 2015; Olsson et al. 2015].
Further, it is difficult to directly compare effects exhibited \textit{in vivo} when the whole organism is exercising as a system compared to the localised effects occurring \textit{in vitro} with cultured muscle samples. Similarly, Aas et al. [2013] suggested \textit{in vitro} myotube systems to lack the \textit{in vivo} microenvironment and communication with other cells, as well as the associated direct contact with other bioactive substances. Additionally, Brown et al. [2015] suggested monolayer cultures to be a good model to investigate exercise \textit{in vitro}. However, it was recognised that the 2D nature of the culture environment was a potential limitation, as it did not enable the complete alignment of myotubes, which can impair intracellular signalling. To overcome this issue the authors suggested the development of a 3D cell culture exercise model. Interestingly, 3D cultures have been reported in primary human skeletal muscle [Martin et al. 2013] and differentiated C212 myoblasts [Player et al. 2014; Sharples et al. 2012], with these models reporting the 3D structure to closely mirror the structural, functional and myogenic characteristics associated with native skeletal muscle. Although, presently no studies have investigated the application of EPS in 3D muscle cultures, which Brown et al. [2015] suggested to be an important consideration for future studies.

However, a number of studies have validated the use of EPS and have demonstrated the model to improve lipid oxidation, increase glycolysis, glucose metabolism, AMPK activation, glucose transport and IL-6 release [Brown et al. 2015; Nikolic et al. 2012; Nedachi et al. 2008], which are well known effects exhibited with \textit{in vivo} exercise. Therefore, confirming the application of \textit{in vitro} muscle culture platforms combined with EPS to be an advantageous experimental platform to investigate metabolic disease and dysfunction.

Data generated in this chapter demonstrated intracellular pH to be substantially lower in CFS/ME and control myoblast samples compared to resting myotube samples. Interestingly, this finding is in line with findings from other researchers [Wagatsuma et al., 2013; Barberi et al. 2011; Leary et al., 1998; Moyes et al. 1997]. For example, Wagatsuma et al. [2013] reported there to be a gradual shift in the dominant energy producing pathways utilised due to myogenic differentiation. Additionally, Leary et al. [1998] investigated metabolic rates during various stages of myogenic differentiation. The authors reported that in proliferating myoblasts approximately 30% of the ATP utilised by the cells was generated via oxidative phosphorylation. Alternatively, in terminally differentiated myotubes oxidative phosphorylation was reported to account for approximately 60% of ATP production and thus the primary source of metabolic energy. It therefore appears that throughout the
differentiation process there is a steady increase in the movement from glycolysis to oxidative phosphorylation as the primary energy source. Moreover, from a molecular perspective mitochondrial enzyme activity (e.g. citrate synthase, cytochrome-c-reductase, succinate dehydrogenase and cytochrome C oxidase) has been reported to dramatically increase during myogenic differentiation. Additionally, respiratory chain complex content have also been reported to be increased in myotubes compared to myoblasts [Barberi et al. 2011; Moyes et al. 1997].

4.4.3 Limitations

Differentiated myotubes have been reported to closely resemble mature human skeletal muscle, in relation to morphological, biochemical and metabolic characteristics [Berggren et al. 2007]. However, as recognised by Brown et al. [2015] monolayer culture systems do not enable complete alignment of myotubes. To overcome this issue, it would be beneficial for future studies to develop novel 3D in vitro culture platforms which incorporate EPS stimulation, to enable in vitro studies to more accurately replicate in vivo investigations.

4.4.4 Conclusion

The in vitro muscle culture system developed in this chapter enabled the determination of intracellular pH in CFS/ME skeletal muscle samples. In contrast to Boulton [2012] the pH-sensing technique applied in this chapter revealed there to be no significant difference in muscle cell pH for CFS/ME patients compared to controls. Similarly, in terms of differentiated myotubes, both sample groups displayed comparable intracellular pH measurements following EPS, which was applied to simulate exercise in vitro. Finally, the PDK inhibitor DCA did not exhibit any significant effect on intracellular pH in any of the muscle samples tested.

This work has highlighted the importance of using significant patient numbers when investigating a heterogeneous population. It has also highlighted the difficulties of comparing in vitro and in vivo data. Taking these limitations into consideration it is still possible to
conclude that, based on current findings, there is no evidence for the role of impaired PDK enzyme function in the peripheral muscle of patients with CFS/ME.
### 4.5 References


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Chapter 5

An Exploration of Glycolytic Function in CFS/ME Primary Myoblasts and Myotubes via Extracellular Flux Analysis and L-lactate Quantification

5.1 Introduction

The purpose of this chapter was to investigate glycolytic function in the skeletal muscle of patients with CFS/ME via extracellular flux (XF) analysis and through the measurement of cellular L-lactate concentration. Previous in vivo studies have reported CFS/ME patients to exhibit excessive acidosis following relatively low level physical activity, which was postulated to be the result of an over-utilisation of the lactate dehydrogenase energy-producing pathway [Jones et al. 2010; Jones et al. 2012]. Additionally, a pilot in vitro study revealed CFS/ME skeletal muscle samples to exhibit aberrantly low intracellular pH in culture, which was normalised following treatment with DCA, further supporting the presence of heightened anaerobic glycolysis in CFS/ME [Boulton. 2012].

5.1.1 Overview of cellular glycolysis

Glycolysis occurs in the cytoplasm and functions as an intracellular bio-chemical energy producing pathway, which acts to convert a single molecule of glucose into two molecules of pyruvate, with a parallel generation of two molecules of ATP (Teslaa & Teitell. 2014). The molecule ATP is the primary source of cellular energy and acts to capture and transfer free energy within biological systems (Pelltier et al. 2014). ATP is generated by both glycolysis and oxidative phosphorylation (mitochondrial respiration). It is important to note that oxidative phosphorylation and glycolysis are tightly coupled and operate as a molecular interconversion system. Under normal conditions, cells are able to consume energy supplied primarily by oxidative phosphorylation, however in hypoxic situations glycolysis is dramatically increased to maintain the energetic balance [Zheng. 2012]. During anaerobic conditions pyruvate is reduced to lactic acid in the cytosol by the enzyme lactate dehydrogenase, in aqueous solutions lactic acid almost completely dissociates into lactate and H⁺. Effectively, acidosis occurs if there is an impairment in oxidative phosphorylation capacity, leading to a net gain in H⁺ [Winer et al. 2014; Zheng, 2012]. Protons are pumped
out of the cell via various mechanisms to maintain intracellular pH [Winer et al. 2014; Casey et al. 2010] and it is the efflux of protons into the extracellular space or medium that results in extracellular acidification (Hochachka et al. 1983; Lane et al. 2009) which can be readily measured via a number of techniques. Overview of the glycolytic pathway demonstrated in figure 5.2

![Figure 5.2: Schematic illustration of cellular glycolysis, adapted from Winer et al [2011].](image)

### 5.1.2 Techniques to measure cellular glycolytic flux

Cellular glycolytic function can be measured via glucose uptake and lactate excretion. Effectively, glucose is transported into the cell via glucose transporters Glut 1-4. Alternatively, lactate excretion occurs through monocarboxylic transporters 1-4. A number of techniques are available to quantify extracellular lactate and glucose in the cell culture media (Teslaa & Teitell. 2014).
One method is via the entrapment of radioactive isotopes, this generally involves the uptake of glucose analogues to measure glycolytic flux. Typically, cells are seeded into culture dishes and then treated with radiolabelled deoxyglucose, which is then phosphorylated in the cytosol of the cell trapping the radioactivity, which is deemed proportional to glycolytic rate of the cell (Bittner et al. 2010). However, this technique does exhibit limitations as noted by Bittner et al (2010). For example, glucose analogues are treated differently by hexokinase than glucose and also may be acutely toxic to the cell even in low doses (Kurtoglu et al. 2007). Additionally, the accumulation of glucose is not only determined by metabolism but is also controlled by the ability of glucose to be transported via the transporters. Furthermore, isotopic measurement provides low spatio-temporal resolution, which makes it impossible to quantify the contribution of individual cells or the detection of rapid phenomena. Finally, Bittner et al. (2010) concluded the technique to be relatively insensitive and exhibit a need for radioactive isotope manipulation which makes them inadequate for the purposes of high throughput analysis.

Glucose and lactate may also be measured through the use of commercially available kits, which function to quantify the amount of the substrate in the cell culture media. Measurement is made via calorimetric of fluorometric detection via standard laboratory spectrophotometry (Teslaa & Teitell. 2014).

Cellular glycolytic function may also be assessed via extracellular flux analysis (Seahorse Bioscience). The XF analyser was introduced in 2006/2007 and was developed to address the need for a high-throughput system for the determination of mitochondrial dysfunction (Horan et al. 2012). In terms of measurement of cellular acidification, the XF analyser builds upon probe technology developed during the late 1980’s and early 1990’s which resulted in the production of the microphysiometer, which functioned to measure proton excretion in culture (Parce et al. 1989; Owicki et al. 1992). However, the instrument overcomes many of the weaknesses associated with the microphysiometer and exhibits increased sensitivity, improved throughput and is capable of measuring in real-time the uptake and excretion of metabolic end products, with the additional option of compound injection (Ferrick et al. 2008; Wu et al. 2007). The flux analyser operates on a 24 or 96-well microplate format and a key benefit is that the assay is based on a single plate which containing multiple samples which can be analysed with high resolution, therefore providing sensitivity and high-throughput capability (Horan et al. 2012). The system consists of a novel fluorescent sensor
probe, which fits over the culture plate and is submerged in the wells of the plate [Horan et al. 2011] (Figure 5.3). Proton extrusion from the cells into the surrounding medium causes rapid and measurable alterations in pH, which is reported as extracellular acidification rate (ECAR), and is measured in milli-pH units per minute (mpH) (Wu et al. 2007). The sensor probe also contains 4 delivery ports, which enable key reagents to be transferred into the culture media to manipulate cellular metabolism.

The XF analyser enables glycolytic stress testing by the sequential delivery of glucose, oligomycin and 2-Deoxy-D-glucose (2-DG). Glucose is supplied to feed glycolysis and the difference between ECAR prior and after injection of glucose is a measure of glycolytic rate. Oligomycin acts to inhibit ATP synthase in the electron transport chain (ETC) which causes a reduction in the ATP/ADP ratio, thus stimulating glycolysis. The difference between ECAR before and after oligomycin addition is equal to the glycolytic capacity. The difference between the glycolytic capacity and glycolysis defines glycolytic reserve capacity. 2-DG glucose acts to inhibit glycolysis through competitive binding with glucose hexokinase and therefore provides baseline ECAR measurement. ECAR after the addition of 2-DG represents non-glycolytic acidification (Teslaa & Tietell, 2014; Wu et al. 2007). Glycolytic flux in response to the injection of key glycolytic stress test reagents is displayed in Figure 5.4.
5.1.3 Hypotheses

The aim of this chapter was to thoroughly investigate glycolytic function in CFS/ME skeletal muscle cells. Specifically, by utilising XF stress testing to reveal key glycolytic parameters, in addition to quantifying L-lactate concentration at rest and following exercise simulation. In each experiment cells were also treated with PDK inhibitor DCA to explore the capacity of the compound to alter cellular glycolytic function.

The specific experimental hypotheses proposed were (1) Increased glycolytic activity in CFS/ME patient myoblasts and myotubes when compared to controls. Evidenced by elevated ECAR for each glycolytic parameter obtained via XF analysis. (2) Elevated cellular L-lactate concentration in CFS/ME muscle samples compared to controls. (3) Normalisation of glycolytic function and L-lactate concentration comparable to controls following treatment with PDK inhibitor DCA.

Figure 5.4: Schematic illustration of glycolytic flux following the addition of stress test reagents to provide the parameters glycolysis, glycolytic capacity and non-glycolytic acidification.
5.2 Materials and Methods

CFS/ME and non-diseased primary myoblasts and myotubes were supplied by Dr Audrey Brown at Newcastle University. A total of 5 (4 females, 1 male) CFS/ME patient myoblast samples were used for experimentation with a donor age of 43.83 ± 12.73. Control samples were obtained from 3 females participants and 1 male aged 50.14 ± 12.81.

All tissue culture reagents for myoblast and myotube routine culture were purchased from Sigma-Aldrich (Poole, Dorset, UK) with the exception of Ham’s F10 myoblast growth media (Lonza, SLS, East Riding of Yorkshire, UK) and chick embryo extract (Seralab, West Sussex, UK).

Tissue culture flasks, 6 well culture plates and serological pipettes were supplied by Greiner bio-one (Stonehouse, UK).

XF-96 (V3) polystyrene cell culture plates, sensor cartridge and cartridge calibration buffer were supplied by Seahorse Bioscience (North Billerica). Glycolysis stress test reagents glucose, oligomycin and 2DG were supplied by Sigma-Aldrich, in addition to Dulbeccos modified eagles medium DMEM which was used as experimental assay medium and L glutamine.

L-lactate assay kit (calometric/fluorometric) was purchased from Abcam (Cambridge, UK).

PDK enzyme inhibitor Potassium Dichloroacetate was purchased from Sigma-Aldrich.

Pierce™ Bicinchoninic acid assay (BCA) protein quantification kit was purchased from Thermo Fisher scientific (Cramlington, UK) and complete Lysis-M buffer from Roche diagnostics (Risch-Rotkreuz, Switzerland).

5.2.1 Study participants

Muscle biopsy samples were obtained from patients with CFS/ME and healthy control participants. The CFS/ME and control individuals were gender and age matched and both groups included both male and female volunteers. Recruitment was achieved through the NHS CFS clinical service within the Newcastle Upon Tyne Hospital Foundation Trust. The
CFS/ME patients all met the Fukuda (1994) criteria for CFS/ME. All participants agreed to complete the study via formal written consent. Ethical approval for the study was provided by the Newcastle and North Tyneside joint ethics committee.

5.2.2 Cell culture and preparation

As previously described in chapter 2, CFS/ME and control muscle cell samples were collected and isolated as detailed in the Brown et al. (2015) study. The myoblast samples were routinely cultured in Ham’s F10 medium, supplemented with 20% FBS, 2% chick embryo extract, 500 U/ML Penicillin-streptomycin and 1% amphotericin B. Myoblasts were grown to 80% confluence in a T75 before being trypsinised and seeded into the vessel of choice. Cells were grown and maintained in a humidified incubator at 5% (v/v) CO₂ at 37°C.

For XF glycolysis stress testing myoblasts were seeded at a density of 3x10⁴ cells per well into XF-96 culture plates and allowed to attach overnight, assays were performed the following day. For L-lactate experimental work myoblasts were seeded at a density of 2 x 10⁵ cells per well and tested the following day.

Differentiation was induced by replacing Ham’s F10 medium with MEM supplemented with 2% (V/V) FBS and 1% Penicillin-streptomycin. Cells were allowed to differentiate over 7 days and experimentation was performed on day 7 differentiation. Media was replaced every two days and all myotubes were tested at passage 6-7.

5.2.3 Electrical pulse stimulation

Electrical pulse stimulation (EPS) was carried out using the C-pace EP cell culture pacer (Ion Optix, Dublin). Following 7 days differentiation EPS stimulation was initiated for 24-hours at alternating frequencies, with one hour low frequency (volts, 24ms, 2Hz) followed by one hour high frequency (5 volts, 24ms, 0.2Hz).
5.2.4.1 L-lactate assay

The commercially available L-lactate kit (Abcam) was used to assess L-lactate concentration in myoblast and myotube samples. Additionally, L-lactate was also measured in stimulated myotubes following EPS.

5.2.4.2 Standard curve and sample preparation

A standard curve was prepared using the L-lactate standard concentrations supplied by the manufacturers. Briefly, standards were diluted in assay buffer to achieve a final L-lactate concentration range of between 0-100pmol/well, which were then seeded into a 96-well plate in duplicate. Fluorescent measurements were performed at 535/587nm. The linear regression of the curve was then utilised to determine the L-lactate concentration in cell samples.

In terms of sample preparation cellular supernatant was diluted 10,000-30,000x in assay buffer as directed by the manufactures. A series of dilutions were performed until the samples readings were within the range of the standard curve.

5.2.4.3 Assay procedure

Prior to running the assay growth medium was removed from the myoblast and resting myotube samples and replaced with DMEM for untreated cells and DMEM supplemented with 40µM DCA for cells receiving treatment. Media was replaced in EPS myotubes after 23.5 hours stimulation. DMEM was removed 30-minutes later in all samples and replaced with lactate assay buffer. The cells were then scraped from the culture vessel and transferred into eppendorfs for sonication (30-seconds) and centrifugation (2000rpm, 5-minutes, 2°C). Supernatant was removed (50µl) and added to the well of a 96-well plate in addition to 50µl of reaction mix. The plate was incubated for 30 minutes, at room temperature and protected from light, before fluorometric measurement.
5.2.4.4 L-lactate fluorometric measurement setup

Fluorescent intensity measurements were performed using the TECAN I 200 fluorimeter and data was obtained using the compatible Magellan software. In terms of set up, the L-lactate kit recommended the following measurement settings, $\lambda_{ex}$ = 535nm, $\lambda_{em}$ = 587nm, additionally gain was set to 70 throughout all experiments and multiple reads (4x4) per well option was selected.

5.2.5 Extracellular flux analysis

5.2.5.1 Assay preparation

The day prior to running the assay myoblasts were seeded into XF-96 culture plates, myotubes were seeded into plates 7 days prior to the assay. The day before testing the XF sensor cartridge was hydrated by adding 100µl of calibrant (Seahorse Bioscience) to the utility plate. The cartridge was then incubated overnight at 37.5°C without CO₂.

On the day of the assay, growth media was removed from culture plates and replaced with DMEM supplemented with 2mM L-glutamine and pH adjusted to 7.35 +/- 0.05 and warmed to 37.5°C. Cells were washed twice with experimental media before the addition of DCA (40µM). The culture plate was then placed in an incubator at 37°C and no CO2 for 60 minutes before initiating the assay to allow for media temperature and pH to reach equilibrium.

5.2.5.2 Compound injection

All glycolytic stress test stock reagents stored at -20°C were defrosted and warmed to 37°C prior to loading the sensor cartridge. Each port of the sensor cartridge was loaded with 25µl of the required compound. Glucose was added to port A for a final well concentration of 10mM. Oligomycin was added to port B for a final well concentration of 3µM and 2-deoxyglucose was added to port C for a final concentration within the well of 100mM.
5.2.5.3 Glycolytic stress test

A glycolysis stress test assay was created on the XF controller, the template incorporate a calibration step for the sensor cartridge carried out without the cell culture plate. Once calibrated the utility plate was removed and replaced with the culture plate and the assay was initiated. The default template involves mix-wait-measure with timings 3min-0min-3 min, with 3 basal rate measurements performed prior to the first injection, followed by 3 rate measurements after each injection.

5.2.5.4 Normalisation

Normalisation for total protein concentration in the well was performed after each glycolytic stress test via the BCA assay. Following the stress test media was removed from the culture plate and cells were washed once with PBS, before the addition of lysis buffer (Thermo fischer) to each well. The wells were then scraped and the contents of the well removed and placed into a new 96-well plate. A 50:1 solution of BCA reagent A to BCA reagent B (Thermo fischer) was added to the wells as per manufacturer’s instructions. The plate was agitated for 30s on a plate shaker and then incubated at 37°C for 30 minutes and protected from light. Absorbance was measured at 562nm on the Tecan Infinite 200.

5.2.6 Data analysis

Results are presented as mean ± standard deviation. Data was analysed via independent and paired T-test. Statistical analysis was performed using Minitab 17 statistical software (Coventry, UK)
5.3 Results

5.3.1.1 L-Lactate standard curve

L-lactate standards were diluted in assay buffer to achieve a final concentration range of 0-100pmol/well. An increase in mean FI exhibited a linear relationship with L-lactate concentration ($y=99.613x$, $R^2=0.973$, $n=5$). The linear regression of this standard curve was utilised to determine the L-lactate concentration in the myoblast and myotube samples. (Figure 5.5).

![Figure 5.5: Standard curve conducted with varying L-lactate standard concentrations. Mean FI was linearly dependent upon the L-lactate concentration ($y=99.613x$, $R^2=0.973$, $n=5$). The data is presented as mean± SD, n=5](image.png)

5.3.1.2 Myoblast L-lactate measurement

L-lactate concentration was measured in CFS/ME (n=5) and control (n=4) myoblast samples. At baseline (0μM DCA) CFS/ME and control myoblasts, samples did not exhibit significantly different L-lactate concentrations. Similarly, DCA treatment (40μm) did not induce any significant effect on either group. Displayed in Figure 5.6.
5.3.1.3 Myotube L-lactate measurement at rest and post-EPS

L-lactate concentration was measured in CFS/ME (n=4) and control myotubes (n=3) at rest and following 24-hours EPS. When CFS/ME myotubes were compared to controls at rest and following EPS there was no significant difference in L-lactate concentration at baseline (0µM) or following DCA (40µM) treatment. Similarly, EPS stimulation exhibited no significant alteration in L-lactate concentration when compared to resting levels in both samples at baseline and following DCA treatment. See figure 5.7.
5.3.2 Extracellular Flux Analysis

5.3.2.1 Myoblast glycolytic stress testing with DCA treatment

Extracellular flux analysis provided a glycolytic profile for CFS/ME and control myoblasts. The sequential injection of inhibitors glucose, oligomycin and 2-DG enabled the measurement of glycolysis and glycolytic capacity and allowed the calculation of glycolytic reserve capacity.

Figure 5.8 demonstrated glycolysis measurements in CFS/ME (n=5) and control (n=4) myoblasts. ECAR was not significantly different in CFS/ME and control myoblasts at baseline (0µM), however was significantly lower in control compared to CFS/ME samples following 40µM DCA treatment. No within group (0µM vs 40µM DCA) effects as a consequence of DCA treatment were observed for either sample.

Figure 5.7: L-lactate concentration in CFS/ME and control myotubes at rest and following 24-hours EPS. L-lactate concentration did not significantly differ for CFS/ME myotubes compared to controls at rest or following EPS at both 0µM and 40µM DCA conditions. The data is presented as mean ± SD, n=7.
The glycolytic capacity of the myoblast sample's was demonstrated in Figure 5.9. ECAR at baseline (0μM) and following DCA treatment (40μM) was not significantly different between CFS/ME and control myoblasts. In terms of within-group effects, DCA treatment exhibited no significant alteration in ECAR for either the CFS/ME or control sample groups.

Figure 5.8: Glycolysis ECAR in CFS/ME and control myoblasts following treatment with DCA (40μM). No significant difference in ECAR at baseline (0μM) between samples, however ECAR significantly reduced in control compared to CFS/ME samples following DCA treatment. No within-group effects as a result of DCA treatment were observed. Data presented as mean ±SD, n=8, ** denoted P<0.05
Figure 5.9: Glycolytic capacity ECAR in CFS/ME and control myoblast samples. No significant difference was exhibited at baseline or following DCA treatment for the CFS/ME verses control myoblast samples. No within group effects were observed for either group following DCA treatment. Data presented as mean ±SD, n=8

Figure 6.0 demonstrated myoblast glycolytic reserve capacity. CFS/ME and control myoblast ECAR did not significantly differ at baseline (0µM) or following treatment with DCA (40 µM). In terms of within-group effects there was no significant alteration in ECAR from baseline following DCA treatment in either group.

Figure 6.0: Glycolytic reserve capacity ECAR in CFS/ME and control myoblast samples. No significant difference was exhibited at baseline and following DCA treatment for the CFS/ME verses control myoblast samples. No within group effects were observed for either group following DCA treatment. Data presented as mean ±SD, n=8
5.3.2.2 Myotube glycolytic stress testing with DCA treatment

Extracellular flux analysis was also performed to assess the glycolytic profile of differentiated myotube samples.

Glycolysis ECAR was not significantly different for CFS/ME (n=3) samples compared to controls (n=3) at baseline (0µM) or following incubation with DCA (40µM). Similarly, DCA treatment did not induce any significant alteration in ECAR in comparison to baseline measures in either group. Displayed in Figure 6.1.

![Figure 6.1: Glycolysis ECAR in myotube samples. No significant difference was exhibited at baseline and following DCA treatment for the CFS/ME versus control myotube samples. No within group effects were observed for either group following DCA treatment. Data presented as mean ±SD, n=8](image)

Glycolytic capacity ECAR is displayed in Figure 6.2. There was no significant difference in ECAR exhibited at 0µM or following 40µM DCA treatment in CFS/ME compared to control myotubes. Additionally, there was no significant alteration in ECAR following DCA treatment compared to baseline (0µM) in either sample group.
Glycolytic reserve capacity is displayed in Figure 6.3. ECAR was significantly lower (P<0.05) in control myotubes at baseline compared to CFS/ME samples. However, following treatment with 40µM DCA this significant relationship was lost. No other significant effects were observed.

Figure 6.2 : Glycolytic capacity ECAR in myotube samples. No significant difference in ECAR at 10µM and 40µM in CFS/ME compared to control myotubes. No significant alteration in ECAR as consequence of DCA treatment in either group samples. Data presented as mean ±SD, n=8

Figure 6.3: Glycolytic reserve capacity ECAR in CFS/ME and control myotube samples. Significantly lower ECAR at baseline in control samples. No other significant effects observed. Data presented as mean ± SD, ** denoted p<0.05
5.4 Discussion

CFS/ME patients frequently report the perception of generalised muscle fatigue, which is exacerbated following relatively low-level physical activity [Jones et al. 2010; Fukuda et al. 1994; MacIntyre et al. 1992]. *In vivo* studies have supported this assertion and have demonstrated the presence of profound intramuscular acidosis, a slowed time to recovery from acidosis post-exercise and a reduced anaerobic threshold in patients [Jones et al. 2010; 2012]. The authors postulated the findings to be related to an over-utilisation of the lactate dehydrogenase energy-producing pathway, even at relatively low exercise intensities. An *in vitro* pilot study also reported CFS/ME patient muscle samples to exhibit an aberrantly low intracellular pH compared to control samples. Nevertheless, this effect was completely normalised following treatment with PDK inhibitor DCA [Boulton. 2012].

The aim of chapter 2 in this thesis was to validate and further develop the *in vitro* muscle pH sensing system reported by Boulton [2012] and to confirm the presence of abnormal pH in CFS/ME myoblasts. However, the technique exhibited a number of weakness and it was not possible to accurately measure cytosolic pH. Therefore, in chapter 4 an alternative intracellular pH sensing technique was performed. Contrary to previous findings, CFS/ME muscle samples exhibited an intracellular pH comparable to controls at rest and following exercise simulation.

The aim of this chapter was to build upon chapter 2 and 4 and investigate cellular glycolytic function in more detail. Glycolytic function was assessed in myoblast and myotube samples via XF analysis, in addition to cellular L-lactate concentration, which was measured in resting samples and following exercise simulation. The results of this chapter demonstrated, firstly in terms of L-lactate concentration, levels were comparable for CFS/ME and control muscle samples at rest and following EPS, which contrasts the experimental hypotheses detailed in section 5.1.3. Additionally, contrary to the experimental hypothesis myoblast XF analysis revealed ECAR measurements for all glycolytic parameters to be comparable for both sample groups. However, glycolysis was reduced in control samples following DCA treatment. Finally, in contrast to the experimental hypothesis glycolytic reserve capacity was lower in control myotubes.
5.4.1 L-lactate concentration

In this chapter, cellular glycolytic function was investigated in vitro. This is the first study to measure L-lactate concentration in CFS/ME patient skeletal muscle samples at rest and following EPS. Lactate concentration is typically interpreted as a marker of anaerobic metabolism [Rogatzi et al. 2015]. Under anaerobic conditions, pyruvate is converted to lactic acid by the enzyme lactate dehydrogenase, which then dissociates to produce lactate and H⁺. If there is an impairment in oxidative pathways during lactate production, there is a net increase in hydrogen ions, resulting in a reduction in pH and the end point of acidosis [Phypers and Pierce. 2006].

The discovery that L-lactate concentration was not elevated in CFS/ME skeletal muscle samples at rest and following 24-hours EPS (Figure 5.6) is in agreement with the findings reported in chapter 4 of this thesis. Chapter 4 as previously described investigated intracellular pH as an alternative marker of anaerobic glycolysis and reported no evidence of elevated intracellular acidosis in CFS/ME muscle samples. Likewise, as discussed in earlier chapters several in vivo studies have measured intracellular pH dynamics using MRS at rest and post-exercise. These studies have reported a lack of evidence to suggest all CFS/ME patients exhibit excessive intracellular acidosis as a consequence of enhanced glycolytic function [Wong et al. 1992]. Additionally, Barnes et al. [1993], proposed the presence of a sub-group within the CFS/ME patient cohort which exhibit intramuscular acidification, however suggested this finding to not be uniform across the whole patient population.

In contrast, Boulton [2012] reported a decreased intracellular pH in CFS/ME myoblasts, which could be interpreted as an over-utilisation of lactate dehydrogenase energy-producing pathway. The reason for the discrepancy may relate to the previously described weaknesses associated with this study.

5.4.2 Glycolytic parameters

Skeletal muscle glycolytic function was assessed via XF analysis, with the technique enabling the determination of glycolytic parameters through the measurement of ECAR in the surrounding culture medium. Quite simply ECAR is a measurement of the excretion of lactic acid per unit of time following its conversion from pyruvate [Wu et al. 2007].
Glycolysis was the first parameter obtained and was measured following the injection of glucose, which was added to promote glycolytic metabolism [Das. 2013]. Glycolysis was not found to be elevated in CFS/ME skeletal muscle samples and DCA treatment did not alter ECAR (Figure 5.7 and 6.0). In contrast, control myoblast ECAR was reduced following the addition of DCA (Figure 5.7). It is difficult to ascertain the reason why ECAR was not similarly reduced in CFS/ME samples, however the finding does suggest control myoblasts to be more sensitive to DCA.

Glycolytic capacity was measured following the injection of oligomycin, which functioned to inhibit mitochondrial ATP production and shift energy production to glycolysis, with increases in ECAR revealing the maximal glycolytic capacity of the cells. CFS/ME muscle samples did not exhibit enhanced glycolytic capacity compared to controls and DCA treatment did not meaningfully alter ECAR in either group (Figure 5.8 and 6.1).

Finally, glycolytic reserve capacity was determined as the difference between glycolytic capacity (maximal in the presence of glucose + glucose) and glycolysis in the presence of glucose [Das, 2013]. Cells with a reduced reserve capacity have been reported to be more dependent upon glycolysis, with a higher reserve capacity indicative of greater tolerance to metabolic stress [Zheng et al. 2012]. Unexpectedly, in the present study control myotubes exhibited a reduced reserve capacity, however following treatment with DCA this effect was normalised (Figure 6.2). Suggesting, CFS/ME muscle samples to have an enhanced tolerance to metabolic stress.

Crucially, the findings suggest CFS/ME skeletal muscle samples to display normal cellular glycolytic function when under metabolic stress, which is in agreement with the previous section, which reported normal L-lactate concentration at rest and following 24-hours of EPS in CFS/ME muscle samples. Similarly, chapter 4 also reported no evidence of enhanced intracellular acidosis at rest and following EPS. However, the results do contrast previously described in vivo studies that measured intracellular pH following exercise [Jones et al. 2012; Jones et al. 2010]. Nevertheless, it is important to acknowledge this is the first in vitro study to directly measure glycolytic function in CFS/ME patient muscle samples, thus it is not possible to directly compare the findings to previous research.

If glycolytic function is not enhanced in CFS/ME, it is possible that other pathophysiological mechanisms may contribute to the muscle fatigue phenotype frequently reported. For
example, Allen et al [1995] suggested low pH and elevate intracellular lactate to impact upon the contractile apparatus of the muscle to a limited degree. In addition, Bogdanis et al. [2012], conducted isolated animal muscle and single fibre experiments and reported fatigue to be in part caused by increased inorganic phosphate leading to a reduction in force production and K\(^+\) accumulation inside the T-tubules. With this effect negatively influencing upon action potential propagation.

Alternatively, in a CFS/ME specific study, Brown et al. [2015] reported several bio-chemical differences in CFS/ME muscle samples following EPS. These were impaired AMPK activation, impaired stimulation of glucose uptake and diminished release of IL-6. Interestingly, AMPK is a key regulator of cellular energy homeostasis, through increased AMP/ATP and ADP/ATP ratios [Jenkins et al. 2013]. It functions both acutely and chronically to restore and maintain cellular ATP levels. In the first instance AMPK stimulates glucose transport and fatty acid oxidation in skeletal muscle [Smith et al. 2005; Balon et al. 2001]. Chronically, AMPK functions to up-regulate proteins involved in substrate availability and oxidation capacity [Winder et al, 2000; Zheng et al. 2001]. Importantly, it acts to maximise mitochondrial function by promoting mitochondrial biogenesis [Jenkins. 2013; Hardie. 2011]. Given the role AMPK in promoting mitochondrial oxidative phosphorylation, reduced activation may contribute to mitochondrial dysfunction, which may contribute to skeletal muscle fatigue in CFS/ME patients

5.4.3 Conclusion

Glycolytic function in CFS/ME patient skeletal muscle samples was assessed by L-lactate measurement and XF analysis. Contrary to previous reports [Jones et al. 2012; 2010] there was no evidence of increased glycolytic function in CFS/ME samples in the basal state, following 24-hours EPS, or when metabolically stressed. Peripheral muscle fatigue experienced by CFS/ME does not appear to be the result of an impaired bio-energetic function and a concurrent over-utilisation of the lactate dehydrogenase energy-producing pathway as previously considered.
5.5 References


Chapter 6

An Investigation of Mitochondrial Function in CFS/ME Patient Myoblast and Myotube samples via Extracellular Flux Analysis

6.1 Introduction

The work reported in this chapter centred on assessing mitochondrial function in myoblasts and myotubes isolated from CFS/ME patient samples. Evidence suggests mitochondrial dysfunction may play a pivotal role in CFS/ME aetiology. Lowered ATP production, impaired oxidative phosphorylation and mitochondrial damage have been reported in patients with CFS/ME [Filler et al. 2014; Myhill et al. 2014]. Moreover, CFS/ME patients share common skeletal muscle symptoms associated with diseases linked to mitochondrial dysfunction, for example muscle pain, fatigue and cramping [Morris & Maes. 2014; Fulle et al. 2007]. However, presently no in vitro studies have been conducted to investigate mitochondrial function in CFS/ME muscle samples via extracellular flux analysis, therefore experimentation using this novel technique to measure cellular mitochondrial function under stress is required.

6.1.1 Oxidative phosphorylation overview

In chapter 5 the first pathway of breaking down glucose into pyruvate was discussed. Glycolysis occurring in the cytosol produces only a small amount of ATP. In contrast the oxidation of pyruvate in the mitochondria results in approximately 15 times more ATP produced compared to glycolysis [Huttemann et al. 2007]. The metabolic reactions involved in oxidative phosphorylation are energy-transducing processes, whereby the oxidation-reduction reactions are essential for the synthesis of ATP. During these reactions electrons are removed by the oxidation of fuel molecules and are transferred to the electron carrier coenzymes nicotinamide adenine dinucleotide (NAD+) in addition to Flavin adenine dinucleotide (FAD) converting them to their reduced form (NADH and FADH₂). The electrons are then transferred to from NADH to O₂ through a series of protein complexes that form the electron transport chain [Hutteman et al. 2007]. The electron transport chain (ETC) is the site of oxidative phosphorylation and comprises the NADH-dehydrogenase (complex I), succinate dehydrogenase (complex II), ubiquinone, bc1 complex (III), cytochrome C and
cytochrome C oxidase (Complex IV). Effectively, electrons derived from NADH are carried through the ETC and enter the chain via complex I. Complex II passes electrons to the ubiquinone/ubiquinone pool and these electrons are transferred through the chain to molecular oxygen. The movement of electrons causes an exergonic reaction that is utilised to pump protons out of the mitochondrial matrix and in doing so generating a membrane potential across the intermembrane space in addition to electrochemical energy in the form of a proton-motive force in the intermembrane mitochondrial space. This process drives the synthesis of ATP as protons (H\(^+\)) are able to move passively to the mitochondrial matrix via a pore which is associated with ATP-synthase (complex V), thus driving ATP synthesis [Huttemann et al. 2007; Yong-Ling et al. 2008] (Displayed in Figure 6.3).

![Image of the electron transport chain]

Figure 6.3: Created by Yong-ling et al [2008]. The electron transport chain with 4 membrane bound complexes (I-IV) Electrons pass through the complexes with assistance from electron carrier Ubiquinone (Q) and cytochromes. The movement of electrons is associated with proton (H\(^+\)) pumping from the mitochondrial matrix to the intermembrane space. This creates a proton motive force, which drives ATP synthesis.

### 6.1.2 Evidence of mitochondrial dysfunction in CFS/ME

Increasing evidence exists to suggest mitochondrial dysfunction plays an underlying role in CFS/ME pathophysiology [Filler et al. 2014; Morris and Maes. 2014]. For example, Booth et al. [2012] investigated mitochondrial function in a large cohort of CFS/ME patients. The authors conducted an ATP profile test, which incorporated a variety of biochemical tests to yield 6 numerical factors that described the availability of ATP, and the efficiency of oxidative phosphorylation in mitochondria located in neutrophils. Additionally, all patients were reported to exhibit measurable mitochondrial dysfunction, which correlated with severity of illness.
Although the mitochondrial investigations were performed in neutrophils the authors suggested the findings to be transferable to other cells in the body.

A variety of studies have investigated mitochondrial function in CFS/ME patient muscle biopsies. Interestingly, a number of these studies have reported the presence of structural abnormalities, which have included mitochondrial degeneration, atrophy of Type II fibres as well as fusion and branching of the cristae [Morris and Maes 2014; Behan et al. 1991]. Further, evidence also exists to demonstrate the presence of key skeletal muscle bio-energetic abnormalities including a reduction in the rate of ATP production and re-synthesis following exercise, when measured via 31 PMRS [Lane et al. 1998; 2003]. Additionally, studies utilising the same methodological approach have also reported an impairment in the bio-energetic function of cardiac muscle [Hollingsworth et al. 2010].

Nevertheless, the alterations in mitochondrial bioenergetics do not appear to be uniform across the CFS/ME patient population, with Barnes et al. [1993] reporting evidence of impaired mitochondrial function in only a sub-group of patients. Alternatively, McCully et al. [2003] reported no evidence of mitochondrial bioenergetic abnormality despite utilising the same methodology. Similarly, Vermeulen et al. [2010] reported CFS/ME patients to exhibit a normal oxidative phosphorylation capacity in the muscle. In this study, patients and control participants completed a controlled repeat exercise intervention to exhaustion. Results demonstrated CFS/ME patients in response to exercise to exhibit a reduced anaerobic threshold, despite an abnormal oxidative phosphorylation capacity. This finding lead the authors to suggest the impairment in anaerobic threshold to be linked to impaired oxygen transport capacity rather than mitochondrial dysfunction.

6.1.3 Mitochondrial respiration measurement techniques

The synthesis of ATP is a key function of the mitochondria and is typically determined experimentally in an in-direct manner through the measurement of mitochondrial O₂ consumption or respiration rate [Perry et al. 2013]. Traditional approaches to the investigation of mitochondrial respiration have predominantly utilised amperometric techniques. Perhaps the most common amperometric approach being the application of the Clarke-type electrodes. Perry et al. [2013] and Gnaiger et al. [2008] described the Clarke electrode as consisting of either a gold or platinum cathode and a Ag/AgCl anode, which are separated via a KCl solution.
Voltage can be applied to both half-cells, which kept apart from the experimental medium via an O\textsubscript{2} permeant material (polyvinylidene difluoride). O\textsubscript{2} diffuses through the membrane and is reduced by electrons at the cathode, generating hydrogen peroxide. This causes the oxidation of the Ag/AgCl anode, which acts to generate an electrical current, which is representative of the partial pressure in addition to the concentration of O\textsubscript{2} in the experimental medium. The changes in the concentration of O\textsubscript{2} in the medium inversely relate to respiratory rate of the cell and allow for the quantification of O\textsubscript{2} consumption. However, classical Clark-type O\textsubscript{2} electrodes are associated with a number of limitations including, the potential of the electrode to consume O\textsubscript{2}, the sensitivity to mass exchange as a result of stirring requirements that may prove damaging to the cell and the need for a large assay volume, thus the electrode is position at quite a distance from the cell monolayer [Diepart et al. 2010].

Two systems have been developed to overcome the weaknesses associated with the use of traditional O\textsubscript{2} electrodes. The systems enable subtle changes in mitochondrial respiration to be determined. These systems include the high-resolution oxygraph- 2K (O\textsubscript{2}K, Oroboros instruments, Austria) and the sensitive high-throughput extracellular flux analyser (Seahorse bioscience). The O\textsubscript{2}K was produced during the 1990’s and has since been extensively used in the field of mitochondrial respiratory analysis. In terms of set-up, the system combines two separate 2-mL chambers, which contain polar graphic O\textsubscript{2} sensors, which enable the real-time measurement of O\textsubscript{2} concentration in (nanomoles/millilitre) and O\textsubscript{2} consumption (picomoles/second/millilitre) within each chamber. The system works by measuring both concentration and consumption of O\textsubscript{2} while injecting key substrates directly to the cells in suspension in the testing chamber, to stimulate various components of the electron transport chain [ Gnaiger et al. 1995; Horan et al. 2012]. The O\textsubscript{2}K exhibits a number of benefits over the conventional use of Clark-type electrodes. For example, one key advantage is that O\textsubscript{2} concentration in the measurement chambers can remain elevated throughout the duration of the assay, or until functional stability of the sample is reduced. This can enable extended substrate-uncoupler-inhibition-titration protocols to be completed, which is incredibly advantageous as such protocols allow for the evaluation of mitochondrial coupling and respiratory control measurements to be performed and as each substrate can be manually injected into the testing chamber each component of the electron transport chain can be investigated independently [Gnaiger, 2009; Horan et al. 2012]. However, the O\textsubscript{2}K does exhibit some crucial limitations. For example, there is a high level of operator input required during experimentation, as there is a need for constant monitoring of the assay and adjustment of concentrations of the injectable
reagents to obtain an optimal signal intensity, thus a fairly labour intensive procedure. Additionally, the system is not capable of high-throughput analysis as only two samples can be tested at one time, with each experiment lasting approximately 1 hour, meaning only 16 samples can be tested in an 8-hour period [Horan et al. 2012].

The extracellular flux analyser (Seahorse Bioscience) overcomes a number of the weaknesses associated with Clark-type electrodes and the O₂K. For example, the XF analyser exhibits high-throughput unlike the Clark-type electrode and O₂K, as the platform is available in a 96-well format, enabling the measurement of mitochondrial bioenergetics in up to 96 samples per plate [Salabei et al. 2014; Ferrick et al. 2008]. Thus, it is possible to test up to 96 samples in one experiment. Additionally, the system is fully automated so requires minimal operator input so proves less labour intensive than the O₂K. Additionally, the XF analyser is capable of measuring O₂ levels in very small volumes of media, just above the cell monolayer, which was not possible with Clark-type electrodes [Wang et al. 2013].

As previously discussed in chapter 5 the XF analyser has now become a mainstream technique to measure cellular bioenergetics (Salabei et al. 2014). Through the use of the previously described optical sensors it is capable of measuring both ECAR and oxygen consumption (OCR) simultaneously. OCR is measured in picomoles/minute and is a determination of the rate of mitochondrial oxidative phosphorylation [Horan et al. 2012]. In terms of function, the sensor probe contains specific fluorophores sensitive to O₂, the piston like sensor is injected into the wells of the plate periodically to form a transient microchamber slightly above cell monolayer (200 microns). To measure OCR fibre optic bundles are injected into the sensor probe to provide excitation and emission light for the fluorophore [Perry et al. 2013]. O₂ levels are measured over a couple of minutes and provide an indication of the extent of cellular O₂ consumption [Perry et al. 2013].

Reagents that inhibit components of the electron transport chain can be added to the sensor cartridge to perform a mitochondrial ‘stress testing’ technique and the inhibitors allow for the determination of 6 mitochondrial parameters. These are, basal OCR, proton leak OCR, maximal OCR, reserve capacity OCR and non-mitochondrial OCR. The parameters are obtained following the sequential delivery of oligomycin (to inhibit ATP synthase), carbonylcynaide p-trifluromethoxyphenylhydrazone (FCCP, to uncouple the mitochondrial inner membrane and allow for uninhibited electron flow though the ETC) and a combination of antimycin a and rotenone (to inhibit complex I and III) [Dranka et al. 2011], which is
displayed in Figure 6.4. The difference between basal OCR and oligomycin-insensitive OCR determines the amount of O$_2$ consumption that is ATP-linked [Dranka et al. 2011; Jekabsons et al. 2004]. The FCCP injection allows for protons to move uninhibited across the mitochondrial inner membrane, leading to maximal electron flow through the ETC, which results in an increase in O$_2$ consumption and enables a determination of maximal OCR [Drank et al. 2011; Jekabsons et al. 2004; Nicholls et al. 2002]. The difference between FCCP OCR and basal OCR provides a measure of cellular respiratory reserve capacity. Finally, a combination of inhibitors antimycin A and rotenone act to completely halt electron movement by acting upon complex I and III, preventing electron movement through complex IV and therefore providing a measure of O$_2$ consumption from non-mitochondrial sources [Dranka et al. 2011; Jekabsons et al. 2004].

![Figure 6.4: Mitochondrial respiration OCR during XF stress test. Mitochondrial parameters depicted in relation to substrate injection. Image provided by Seahorse Bioscience.](image)

However, as with the techniques previously described the XF analyser also exhibits some limitations. Firstly, the injectable compounds are expensive to purchase and require optimisation prior to use. Additionally, the fluorescent plates, which are costly, can only be used in one assay [Horan et al. 2012]. Finally, it is possible that the injectable compounds may
interact with the fluorescent probe or plastic plate generating inaccurate results [Sauerbeck et al. 2011].

6.1.4 Assessment of mitochondrial function

In this chapter CFS/ME skeletal muscle mitochondrial bioenergetics were investigated in vitro by manipulating the glucose substrate availability. The concentration of glucose in the assay media was altered to maximise oxidative phosphorylation. Interestingly, muscle cells have been reported to be highly glycolytic when grown and differentiated in high glucose conditions relative to muscle tissue in vivo, which complicates the accurate study of mitochondrial function [Aguer et al. 2011]. For example, myoblasts are routinely cultured in high glucose media, which has been reported to diminish mitochondrial function. Additionally, cancer cell lines have been reported to be highly glycolytic when grown in high glucose media, an occurrence referred to as the ‘Crabtree effect’, whereby excess glucose functions to impair oxidative phosphorylation [Marroquin et al. 2007; Shulga et al. 2010; Rossignol et al. 2004]. Furthermore, this effect is not limited to cancer cell-lines and has also been reported in primary human myotubes [Aguer et al. 2011], embryonic tissues and proliferative thymocytes [Aguer et al. 2011; Ibsen et al. 1961].

To combat the ‘Crabtree’ effect a number of studies have substituted glucose for galactose, with a goal to promote the utilisation of mitochondrial oxidative phosphorylation and enable a thorough investigation of mitochondrial function/dysfunction [Aguer et al. 2011; Elkalaf et al. 2013; Marroquin et al. [2007]. However, Elkalaf et al. [2013] reported galactose supplementation of the culture medium to impair parameters of mitochondrial function (maximal respiration and spare respiratory capacity) in C2C12 myoblast and myotube cell lines.

6.1.5 Hypotheses

The aim of this chapter was to examine CFS/ME myoblast and myotube mitochondrial function via XF analysis. Function was assessed following the addition of key inhibitors of the mitochondrial electron transport chain. Further investigation into CFS/ME myoblast and
myotube function was performed by varying substrate availability (glucose) during the mitochondrial stress test.

The specific experimental hypotheses were:

(1) CFS/ME myoblasts and myotubes exhibit impaired mitochondrial function indicated by reduced OCR measurements for all mitochondrial parameters.

(2) CFS/ME samples exhibit lower OCR for mitochondrial parameters compared to controls when substrate availability is reduced.
6.2 Materials and Methods

CFS/ME and control myoblasts were provided by Dr Audrey Brown at Newcastle University. Tissue culture reagents including penicillin streptomycin, amphotericin B, Fetal bovine serum, trypsin EDTA, phosphate buffered saline and modified Eagles medium were supplied by Sigma-Aldrich (Poole, Dorset, UK). Ham’s F10 growth media was supplied by Lonza (SLS, East riding, Yorkshire). Chick embryo extract was supplied by Sera lab (West Sussex, UK).

Plastic consumables including serological pipettes, 96-well plates, graduated pipette tips, cell scrapers, falcon tubes and tissue culture flasks were supplied by Greiner Bio (Stonehouse, UK).

XF flux 96-well plates, sensor cartridge and calibration buffer was supplied by Seahorse Bioscience (North Billerica). Assay medium (DMEM), L-glutamine, glucose and mitochondrial stress test reagents oligomycin, FCCP, antimycin and rotenone were all purchased from Sigma-Aldrich.

Pierce™ BCA protein assay kit was supplied by Thermo scientific (MA, USA). Complete Lysis-M was purchased from Roche diagnostics (Switzerland).

6.2.1 Study participants

Muscle biopsy samples were obtained from the vastus lateralis of CFS/ME patients and healthy control participants (as previously outlined in chapter 2). Recruitment was conducted through the Newcastle Upon Tyne Hospital foundation trust. All CFS/ME patients met the Fukuda criteria for CFS/ME. All participants supplied formal written consent and ethical approval was granted from the Newcastle and North Tyneside ethics committee. A total of 4 CFS/ME patient samples were used for experimentation (3 females, 1 male) with an average donor age of 46.12± 10.1 years. The control sample group consisted of 3 females and 1 male with a donor average donor age of 46.13 ±7 years.
6.2.2 Cell culture and preparation

CFS/ME and control myoblast samples were collected as detailed by Brown et al (2015). Myoblasts were grown in Ham’s F10 medium, supplemented with 20% FBS, 2% chick embryo extract, 500U/mL Penicillin streptomycin and 1% amphoteric B. The cells were stored in a humidified CO₂ incubator (5% v/v) at 37°C.

For the mitochondrial stress test myoblasts were seeded at a density of 3 x 10⁴ cells per well into XF-96 plates. This seeding density was determined during optimisation work. Myoblast experimentation was performed the following day. Differentiation was performed by replacing growth media with MEM supplemented with 2% FBS (v/v) and 1% penicillin-streptomycin. The cells were allowed to differentiate over a 7-day period; media was replaced every two days. Experiments were performed on day 7 differentiation. All assays were performed on myoblast and myotubes at passage 6-7.

6.2.3 Extracellular flux analysis

6.2.3.1 Assay preparation

Prior to experimentation myoblasts were seeded into XF-96 plates and allowed to attach overnight or differentiated into myotubes for 7-days. The sensor cartridge was hydrated overnight in 100µl calibrant per probe. The cartridge was incubated in a CO₂ free incubator at 37°C.

On the day of the assay the experimental medium was prepared by supplementing DMEM with pyruvate (1mM) and L-glutamine (2mM), before adding the desired concentration of glucose. The glucose concentrations tested were 2.5mM (hypoglycaemic) 5mM (normal) and 10mM (hyperglycaemic). The media was then pH adjusted to 7.35 +/- 0.05 and warmed to 37.5°C. The media of the cell culture plate was removed, the plate washed and DMEM added. The culture plate was then placed in a CO₂ free incubator for one hour to enable an equilibration in the pH and temperature of the experimental media.
6.2.3.2 Compound preparation

Prior to the loading the sensor cartridge, the mitochondrial stress test stock reagents stored at -20°C were defrosted. The ports of the sensor cartridge were loaded at a volume of 25µl. Port A was loaded with oligomycin at a final well concentration of 3µM, port B was loaded with FCCP at a final well concentration of 1µM and finally port C was loaded with 1.2µM antimycin A and rotenone.

6.2.3.3 Optimisation

A series of optimisation assays were required before completing experimental work. Firstly, cell seeding density was determined for both myoblast and myotube cell samples. The XF-96 manufacturers (Seahorse Bioscience) recommend an optimal seeding density to exhibit an OCR of 20-160pmol/min during the final basal measure during mitochondrial stress testing. Therefore, seeding densities between 1x10⁴ and 5x10⁴ were investigated to determine the optimal seeding density. In both myoblast and myotube samples 3x10⁴ was found to exhibit an OCR within the recommended range and also provided a good signal and consistent response throughout the experiment.

The concentration of stress test inhibitors was also optimised. This involved titrating concentrations of each inhibitor and choosing the concentration which exhibited the greatest and most consistent response. The inhibitor concentrations that provided the best response in both myoblast and myotube samples were 3µM oligomycin, 1µM FCCP and 1.2µM rotenone and antimycin-A.

6.2.3.4 Mitochondrial stress test

The mitochondrial stress test assay template was created on the XF controller. The assay incorporated a pre-calibration step followed by a calibration of the sensor cartridge. Once complete the utility plate was removed and replaced with the cell culture plate. The default template mix-wait-measure with the timings 3min,0 min, 3min was utilised in addition to three basal measurements performed before injection 1 and 3 rate measurements after each injection.
6.2.3.5 Normalisation

The OCR for each well was normalised for total protein concentration per well after each mitochondrial stress test. This normalisation process was justified as a number of studies have reported this normalisation strategy to be capable of successfully revealing mitochondrial bioenergetic differences between different cell samples [Salabei et al. 2014; Dott et al. 2014].

This was performed via BCA assay and briefly involved removing assay medium from the culture plate and adding 25μl of lysis buffer to each well. The contents of the well was then removed and placed into a clean 96-well plate. BCA reagent A and B were added in a 50:1 solution to each well as per manufacturer’s instructions. The plate was agitated for 30s on a plate shaker and then incubated at 37°C for 30 minutes while protected from light. Absorbance was measured at 562nm on the Tecan Infinite 200.

6.2.4 Data analysis

Results are presented as mean ± standard deviation. Data was analysed via independent and paired T-test. Statistical analysis was performed using Minitab 17 statistical software (Coventry, UK).
6.3 Results

6.3.1 Mitochondrial stress testing with varied glucose

Cellular mitochondrial function was assessed by varying the availability of the substrate glucose in the experimental assay media. Extracellular flux analysis provided a measurement of mitochondrial function following treatment with key mitochondrial ETC inhibitors; oligomycin, FCCP and a mixture of rotenone and antimycin A.

6.3.2 Basal respiration

Figure 6.5 demonstrated CFS/ME (n=4) and control (n=4) myoblast basal OCR in response to varied glucose media concentration. At the 2.5mM glucose concentration OCR was significantly lower (P<0.05) in the CFS/ME sample group compared to controls. No other significant alterations in OCR were observed as a consequence of varied glucose concentration.

![Figure 6.5: Basal OCR in myoblasts with varied glucose media concentration. CFS/ME samples exhibited significantly lower OCR compared to controls at 2.5mM glucose concentration. Data presented as mean ± SD, n=8, ** denoted P<0.05](image-url)
Figure 6.6 demonstrated myotube basal respiration OCR following the same glucose substrate treatment regime. No significant differences in OCR were exhibited with varying glucose concentration when CFS/ME (n=4) myotubes were compared to controls (n=4). Additionally, no significant within-group differences were observed.

6.3.3 ATP-linked respiration

Myoblast ATP-linked respiration is demonstrated in Figure 6.7. CFS/ME (n=4) myoblasts exhibited significantly lower (P<0.05) OCR following incubation in 10mM glucose media when compared to controls (n=4). No other significant relationships were observed.
Myotube ATP-linked respiration is illustrated in Figure 6.8. There was no significant difference between CFS/ME and control samples associated with any of the glucose media concentrations. Similarly, there were no significant within-group differences as a consequence of varied glucose concentration.

Figure 6.7: ATP-linked OCR in myoblast samples following incubation with varied concentrations of glucose media. At 10mM glucose, media concentration OCR was significantly lower in the CFS/ME sample group compared to controls. No significant within-group differences were found as a consequence of varied glucose concentration. Data presented as mean ±SD, n=6, ** denoted P<0.05

* denotes P<0.05
6.3.4 Maximal respiration

Myoblast maximal respiration is displayed in Figure 6.9. No significant difference in OCR was exhibited between CFS/ME and control sample groups at any of the glucose media concentrations. No significant within-group differences were demonstrated as a consequence of varied glucose media concentration. Data presented as mean ±SD, n=6.

Figure 6.8: ATP-linked OCR in myotube samples following incubation with varied concentrations of glucose media. No significant difference in OCR for CFS/ME verses control myotubes at any glucose media concentration. No significant within-group differences were found as a consequence of varied glucose concentration. Data presented as mean ±SD, n=6.

Figure 6.9: Myoblast maximal respiration with varied concentrations of glucose media. No significant difference in OCR for CFS/ME verses controls across glucose media concentrations. No significant within-group differences were found as a consequence of varied glucose concentration. Data presented as mean ±SD, n=6.
Myotube maximal respiration is demonstrated in Figure 7.0. OCR did not significantly differ between CFS/ME and control myotubes at any of the glucose media concentrations tested. Similarly, varied glucose media concentration did not result in any significant within-group differences.

![Figure 7.0: Myotube maximal respiration with varied concentrations of glucose media. No significant difference in OCR for CFS/ME versus controls across glucose media concentrations. No significant within-group differences were found as a consequence of varied glucose concentration. Data presented as mean ±SD, n=6](image)

### 6.3.5 Spare respiratory capacity

Myoblast spare respiratory capacity is demonstrated in Figure 7.1. Spare capacity (%) in CFS/ME and control myoblasts did not significantly differ at any of the glucose media concentrations tested. Additionally, no within-group differences were observed across the concentration range tested.

Myotube spare respiratory capacity is demonstrated in Figure 7.2. As described in myoblast samples CFS/ME and control myotube spare respiratory capacity (%) did not differ significantly at any of the glucose media concentrations tested. Moreover, no within-group differences were observed at the media glucose concentrations investigated.
Figure 7.1: Myoblast spare respiratory capacity. OCR was not significantly different in CFS/ME compared to control myoblasts at any of the glucose media concentrations investigated. No within-group differences between any of the glucose concentrations was observed.

Figure 7.2: Myotube spare respiratory capacity. OCR was not significantly different in CFS/ME compared to control myoblasts at any of the glucose media concentrations investigated. No within-group differences between any of the glucose concentrations was observed.
6.4 Discussion

Previous research suggested that the peripheral muscle fatigue phenotype expressed in CFS/ME may be related to impaired bio-energetic function. This energetic abnormality subsequently leading to intracellular acidosis in response to exercise [Jones et al. 2012; 2010]. Likewise, in vitro assessment of muscle intracellular pH revealed CFS/ME myoblasts to exhibit an aberrantly low pH, which was later normalised following treatment with enzyme inhibitor DCA. Thus, taken together preliminary work hypothesised impaired function of the 3-enzyme complex PDC as the pathophysiological mechanisms underlying muscle fatigue in CFS/ME.

However, the data generated in previous chapters has not supported the assertion of impaired PDC function. This was evidenced following treatment with pH responsive dye BCECF-AM, which revealed CFS/ME skeletal muscle samples to exhibit an intracellular pH comparable to controls at rest and following EPS. Additionally, glycolytic capacity was investigated via extracellular flux analysis, revealing no abnormality in key glycolytic parameters when compared to controls. Finally, the compound DCA failed to modulate pH and ECAR in all of the in vitro assays performed.

The present chapter aimed to investigate the potential role of an alternative pathophysiological mechanism in CFS/ME that could be associated with muscle dysfunction and fatigue. Interestingly, a number of the muscle symptoms experienced by CFS/ME patients such as muscle pain, fatigue and cramping are also reported in disease states associated with mitochondrial dysfunction [Fulle et al. 2007; Morris and Maes. 2014]. Additionally, impaired oxidative phosphorylation, ATP production and mitochondrial damage have been reported in patients with CFS/ME (Filler et al. 2014; Myhill, 2008]. Nonetheless, the present chapter is the first to assess mitochondrial function in CFS/ME muscle samples in vitro via XF analysis.

The primary findings were firstly, basal respiration was lower in CFS/ME myoblast samples compared to controls when incubated in media supplemented with 2.5mM glucose media, which was in agreement with the experimental hypotheses. However, no difference was observed in basal OCR across all glucose media concentrations for CFS/ME compared to control myotube samples. Therefore, contrasting the experimental hypotheses.
Secondly, ATP-linked respiration was reduced in CFS/ME myoblasts compared to controls following treatment with 10mM glucose, in agreement with the experimental hypothesis. However, this effect was not observed in myotube samples.

Finally, contrary to the experimental hypotheses no difference in maximal respiratory capacity or spare respiratory capacity was exhibited between CFS/ME and control myoblast and myotube samples with any of the glucose media concentrations tested, therefore rejecting the experimental hypotheses.

6.4.1 Mitochondrial parameters

In the present chapter skeletal muscle samples were exposed to varying glucose substrate concentrations (2.5-10mM). It is important to note that the 2.5mM glucose concentration was representative of hypoglycaemia as typically normal serum glucose levels are maintained within 4mM-6mM (approximately 72-108mg/dL). However, in the presence of low nutrient availability levels can drop to around 2.5mM (45mg/dL) with tissue levels reportedly lower [Zhuang et al. 2014]. The utilisation of low glucose media to assess mitochondrial function was justified by previous studies that have reported muscle cells to be highly glycolytic in the presence of excess glucose availability, thus impairing mitochondrial function. A glycolytic phenotype with hyperglycaemia was first noted in cancer cell lines and is referred to as the ‘Crabtree effect’. However, this effect has since been reported in a variety of other cell lines including skeletal muscle cells [Elkalaf et al. 2013; Marroquin et al. 2007; Shulga et al. 2010; Rossignol et al. 2004].

The first mitochondrial parameter obtained from XF testing was basal respiration. This measure has been described as an OCR, which reflects coupled mitochondrial respiration and the uncoupled consumption of O₂ to form reactive oxygen species at mitochondrial and non-mitochondrial enzymatic sites [Hartman et al. 2014]. In terms of XF analysis basal respiration was measured prior to the injection of any mitochondrial inhibitors. In this study basal OCR was reduced in CFS/ME myoblasts compared to control samples following incubation with the lowest glucose media concentration only, which is displayed in Figure 6.6, however CFS/ME myotube basal OCR was comparable to controls at all glucose media concentrations, displayed in Figure 6.7.
It is important to note that when cells are exposed to low glucose medium ATP is exclusively produced by oxidative phosphorylation [Marroquin et al. 2007]. Therefore, in the present study by reducing substrate availability the muscle cells were experimentally forced to utilise oxidative phosphorylation as the primary energy-producing pathway. At first glance, a lowered basal respiration in CFS/ME myoblasts would appear to be suggestive of underlying mitochondrial dysfunction when the cells are experimentally manipulated to produce ATP solely via oxidative phosphorylation. However, this interpretation is questionable, as low substrate availability did not impact upon basal OCR in CFS/ME myotube samples.

This discrepancy could be related to the shift in the dominant energy-producing pathway during myogenic differentiation, initially with naïve muscle cells primarily utilising glycolysis before steadily moving towards predominantly oxidative phosphorylation [Wagatsuma et al. 2013]. For example, it has been reported that in proliferating myoblasts approximately only 30% of ATP is produced via oxidative phosphorylation, compared to 60% in differentiated myotubes [Leary et al. 1998]. Additionally, from a molecular perspective myoblasts have been reported to exhibit a reduction in mitochondrial enzyme activity and a lowered respiratory chain complex content when compared to myotubes [Barberi et al. 2011; Moyes et al. 1997]. While it is evident that myoblasts and myotubes exhibit distinct differences in mitochondrial capacity it is difficult to ascertain why control myoblasts did not exhibit a comparable OCR when incubated in low glucose media.

The additional mitochondrial parameter ATP-linked respiration was also measured in the present study. Effectively, the difference between cellular basal OCR and the oligomycin insensitive OCR yields the amount of O$_2$ that is linked to ATP-production [Jekabsons. 2004]. XF analysis revealed CFS/ME myoblasts to exhibit a lower OCR, translating to a reduced cellular oxidative ATP production but only under hyperglycaemic conditions (Figure 6.8). However, no differences in OCR were observed in myotube samples (Figure 6.9). Possible reasons for this finding may relate to the previously described glycolytic phenotype acquired when cells are cultured in high glucose medium [Elkalaf et al. 2013]. For example, as previously described studies have demonstrated myoblasts during early stage differentiation to rely predominantly on lactate production from glucose so meet their cellular demands [Elkalaf et al. 2013; Leary et al. 1998]. Again, it is difficult to ascertain why control myoblasts did not exhibit a comparable OCR with hyperglycaemia but suggests CFS/ME myoblasts to exhibit a
more glycolytic phenotype prior to differentiation; however, this effect appears to be lost once terminally differentiated.

Maximal respiration was also determined in the present study. No differences were observed between CFS/ME and control muscle samples (Figure 7.0 and 7.1). Maximal respiration OCR was determined following the injection of FCCP. This inhibitor functions to allow the uninhibited movement of protons across the mitochondrial inner membrane, acting to collapse the mitochondrial membrane capacity. Oxygen consumption is then dramatically increased enabling the determination of maximal OCR [Dranka et al. 2011]. Interestingly, a decrease in maximal respiration has been reported to be a pivotal indicator in determining potential cellular mitochondrial dysfunction. For example, Elkalaf et al. [2013] reported uncoupling rates to provide a measurement of the maximum capability of the ETC and substrate oxidation that the cells are capable of when under assay conditions. Therefore, comparable OCR of CFS/ME and control samples would suggest CFS/ME skeletal muscle cells do not exhibit mitochondrial dysfunction.

Similarly, spare respiratory capacity has been suggested to be an important diagnostic measure of mitochondrial dysfunction. It was calculated by dividing the OCR response following FCCP injection by the basal OCR. Effectively, the measure provides an indication of cellular stress and provides information regarding the capability of the ETC and substrate supply to respond to increased energy demand, with cells with a higher capacity better able to respond to stress [Brand and Nicholls. 2011]. In the present study, spare respiratory capacity was comparable in CFS/ME and control myoblast samples (Figure 7.2).

Taken together the findings suggest CFS/ME myoblasts do exhibit a reduced oxidative phosphorylation capacity when compared to controls. However, once terminally differentiated into myotubes this effect appears to be lost and suggests that CFS/ME patients exhibit normal mitochondrial function. It is important to remember that muscle is formed by the fusion of many muscle pre-cursor cells (myoblasts) to form multinucleated myotubes which then mature into myofibres [Berendse et al, 2003]. Unlike myoblasts, myotubes have been suggested to exhibit morphological, metabolic and biochemical properties akin to intact skeletal muscle fibres [Olsson et al. 2015] and in terms of experimentation represent the best alternative to intact human skeletal muscle [Nikolic et al.2012].
It is difficult to directly compare the results of the present investigation with previous research, as this is the first study to utilise an in vitro testing platform to experimentally induce mitochondrial stress in CFS/ME muscle samples. Presently, the majority of studies have investigated skeletal muscle mitochondrial function in vivo typically via MRS approaches. Lane et al. [1998] utilised the aforementioned technique and in contrast to the present study reported a reduction in ATP-production and re-synthesis in CFS/ME patients. Nevertheless, in agreement with the present study others have reported limited evidence of mitochondrial dysfunction in the CFS/ME patient cohort. For example, Vermeulen et al. [2010] reported normal oxidative phosphorylation capacity in muscle following a repeat exercise protocol to exhaustion. Alternatively, Barnes et al. [1993] reported mitochondrial dysfunction in some but not the entire patient cohort, suggestive of a sub-group of CFS/ME patients that exhibited mitochondrial dysfunction. Possible reasons for the discrepancies between the present study and others that have alternatively reported mitochondrial dysfunction in CFS/ME may relate to the use of skeletal muscle cells. Although myotubes have purported to exhibit morphological, metabolic and biochemical properties similar to intact adult skeletal muscle fibres [Olsson et al. 2015], it is important to note that they also exhibit key differences, including the expression of immature muscle proteins [Larkin et al. 2006], alterations in the abundance and distribution of glucose transporters and a slightly more glycolytic phenotype [Baker et al. 2003; Sarabia et al. 1992].

6.4.2 Limitations

CFS/ME patients frequently report a changeable pattern to their symptoms and physical capabilities, often with severe exacerbation following physical exercise [Whiteside et al. 2004; Fukuda et al. 1994]. Therefore, in terms of experimentation the ability to assess mitochondrial parameters post-exercise is warranted and has been conducted in previous in vitro studies [Lane et al, 2003; Vermeulen et al. 2010; Hollingsworth et al. 2010]. To overcome this obstacle EPS was applied in previous chapters to simulate physical activity in vitro. However, in this study it was not possible to perform EPS as the process required cells to be seeded and differentiated in 35mm dishes which were compatible with the c-pace EP cell culture pacer (Ion Optix, Dublin). Whereas for the XF analysis it was necessary to seed cells into XF-96 culture plates. It would be advantageous to develop an EPS device, which would be compatible with the XF-96 plate to enable mitochondrial bioenergetics to be interpreted following exercise simulation.
6.4.3 Conclusion

This chapter investigated CFS/ME skeletal muscle mitochondrial function via XF analysis and is presently the first study to use this methodology to provide a direct measurement of cellular mitochondrial bioenergetics in the muscle of CFS/ME patients. Ultimately, CFS/ME skeletal muscle cells exhibited normal mitochondrial function, which was evidenced by OCR values comparable to controls for all myotube mitochondrial parameters. This finding is in agreement with some studies that have reported normal oxidative phosphorylation capacity in the muscle of CFS/ME patients [Vermeulen et al. 2010] and contrasts others who have alternatively reported impaired ATP production and resynthesis in a CFS/ME patient cohort [Lane et al, 1998].
6.5 References


Lane, RJ; Barrett, MC; Taylor, DJ; Kemp, GJ; Lodi, R. (1998). Heterogeneity in chronic fatigue syndrome: evidence from magnetic resonance spectroscopy of muscle. *Neuromuscul Disord.* 8, 204-209


Chapter 7
General Conclusions and Future Directions

7.1 Conclusions

The overriding aim of this PhD thesis was to provide a new insight into the biochemical basis of muscle cell dysfunction in patients with CFS/ME through the use of novel *in vitro* technologies.

One of the primary symptoms of CFS/ME is generalised abnormal muscle fatigue that occurs following relatively mild physical activity. Additionally, patients frequently report an inability to maintain muscle activity due to a perceived ‘lack of energy’ and ‘muscle pain’, which can be severe enough to lead the patient to avoid physical activity completely. Previous *in vivo* studies have revealed CFS/ME patients to exhibit profound and sustained intramuscular acidosis following a standardised exercise protocol [Jones et al. 2012; 2010]. Whereas others have reported either no evidence of enhanced acidosis in CFS/ME patients post-exercise [Wong et al. 1992] or in only some of the CFS/ME patient cohort [Barnes et al. 1993], demonstrating the heterogeneity of the CFS/ME patient population.

An *in vitro* pilot study reported aberrantly low intracellular pH in CFS/ME patient myoblasts, which was normalised following treatment with PDK inhibitor DCA [Boulton. 2012]. In this study intracellular pH was measured via a novel fluorescent pH responsive nanosensor system. This study provided preliminary evidence to suggest a role of bio-energetic dysfunction in CFS/ME because of an over utilisation of the lactate dehydrogenase pathway. Therefore, a key aim of this PhD thesis was to build upon pilot work to further develop and validate the pH responsive nanosensor system to confirm the presence of acidosis in CFS/ME patient muscle cells and also the capacity to modulate bio-energetic function and inform the treatment of peripheral muscle fatigue.

Chapter 2 aimed to develop and further develop the fluorescent pH responsive nanosensors system, however the work carried out in this chapter was unable to reliably detect intracellular acidosis in CFS/ME patient myoblast cells. Additionally, DCA did not modulate intracellular pH in either CFS/ME or control cells.
In chapter 4 the fluorophore BCECF-AM was used as an alternative strategy to measure intracellular pH, to confirm the presence of acidosis in CFS/ME patient muscle and investigate the ability of DCA to modulate normalise pH. Additionally, EPS was used to simulate exercise *in vitro*. In contrast to chapter 2, BCECF enabled the reliable determination of intracellular pH. However, contrary to pilot data, no evidence of elevated intracellular acidosis at rest or following EPS was found in CFS/ME patient muscle samples. Rather intracellular pH measurements were comparable to control cells. Additionally, DCA did not modulate intracellular pH in either CFS/ME patient or control cells. This work highlighted the importance of adequate patient numbers when investigating a heterogeneous patient population. Furthermore, it also illustrated the difficulty in directly comparing *in vitro* and *in vivo* data.

Chapter 5 further developed upon chapter 4 by using alternative techniques to measure cellular glycolytic function, these were XF analysis and the measurement of L-lactate. In agreement with chapter 4, no evidence of increased glycolytic function was found in CFS/ME patient muscle cells when compared to controls. This was demonstrated by comparable measurements in the basal state, post-EPS and following glycolytic stress testing. Similarly, DCA did not alter glycolysis in either patient or control cells.

In chapter 6 muscle mitochondrial function was assessed via XF analysis. In this chapter, glucose substrate availability was manipulated to induce mitochondrial stress. CFS/ME patient muscle cells exhibited normal mitochondrial function, which was evidenced by OCR values comparable to control cells for all mitochondrial parameters.

The possible role of enhanced oxidative stress in CFS/ME was investigated in chapter 3. A direct real-time electrochemical technique was used to measure $O_2^-$ generation in CFS/ME patient myoblast cells. To stimulate $O_2^-$ generation cells were treated with ethanol or experienced lactic acidification. The data revealed there to be no evidence of enhanced oxidative stress in CFS/ME patient muscle samples, rather levels of $O_2^-$ generated were comparable to control cells.

Taken together the *in vitro* muscle culture approaches reported in this thesis have enabled the investigation of the biochemical basis of muscle cell dysfunction in patients with CFS/ME. It is possible to conclude there to be no evidence of impaired muscle function in CFS/ME patients. Additionally, there was no impairment found in PDK enzyme function. Therefore, it
is possible to determine that bioenergetic function is normal in CFS/ME patients and does not explain the excessive peripheral muscle fatigue phenotype exhibited in patients with CFS.

7.2 Future Work

In this thesis, there was no evidence of biochemical abnormality at the skeletal muscle level of patients with CFS/ME. However, investigating the muscle in isolation failed to take into account systemic biochemical factors that may influence muscle function in vivo. It would be advantageous for future studies to investigate bioenergetic function in isolated peripheral blood mononuclear cells. Specifically, this could be achieved with XF analysis to enable glycolytic and mitochondrial stress testing. Also in line with previous in vivo investigations, blood sampling could be obtained from CFS/ME patients in the resting state and also following a standardised exercise protocol. Such strategies would help to bridge the gap between in vivo and in vitro investigations and promote improved understanding of the impact of systemic factors on muscle function.
7.3 References


