Abstract

Essential elements of Myalgic Encephalomyelitis (ME) are muscle (weakness) and tenderness, cognitive deficits, neurological impairments, especially of cognitive, autonomic and sensory functions, but above all, post-exertional “malaise”: a prolonged increase of symptoms after a minor physical and mental exertion.

Chronic fatigue Syndrome (CFS) is defined as clinically evaluated, unexplained (persistent or relapsing) chronic fatigue, accompanied by at least four out of eight specific symptoms, e.g., sore throat, unrefreshing sleep, and headaches.

Since cognitive deficits and post-exertional “malaise” are not mandatory for the diagnosis CFS, only part of the CFS patient group meets the diagnostic criteria for ME.

So, post-exertional “malaise” is considered to be the distinctive feature of ME. However, “malaise” is an ambiguous and subjective notion. In order to assess post-exertional malaise objectively, several studies have employed widely used methods to quantify the deviant effects of exertion in ME (CFS). This review focuses on the long-lasting (negative) effects of exercise on the performance indicators of the physical exercise capacity, the cognitive deficits and the muscle power.

Keywords: Myalgic encephalomyelitis; Chronic fatigue syndrome; Exercise; Post-exertional malaise; Methods; Assessment

Introduction

ME and CFS are often qualified as enigmatic disorders [1], partly due to the fact that typical symptoms, e.g. fatigue and post-exertional “malaise”, are subjective notions. A substantial part of the confusion and controversy with regard to the diagnosis, the nature and the treatment of ME and CFS originates from these subjective notions. To end this debate and to move forward it is essential to use objective methods to assess the characteristic symptoms of ME and CFS objectively, as far as possible [2]. This review aims to discuss observations with regard to the impact of exercise on the physical exercise capacity, cognitive functioning and muscle power in ME (CFS).

Post-exertional “malaise” in this article is defined as ‘a pathological inability to produce sufficient energy on demand’ [3], resulting into a (delayed) increase of typical symptoms, e.g. weakness, muscular and/or joint pain, cognitive deficits, after a minor physical or mental exertion, with prolonged “recovery” times [4,5].

The Impact of Exertion on the Physical Exercise Capacity

A cardiopulmonary test (CPET) is considered an accurate method for assessing function [6]. The ventilatory or anaerobic threshold indicates the workload and oxygen consumption at which the anaerobic metabolism begins to predominate [7]. Therefore a negative effect of a CPET on this threshold and the accompanying oxygen uptake partly reflects the physical dimension of post-exertional malaise.

Figure 1: Exercise performance levels of ME/CFS patients compared to sedentary controls.

Although contradicted by some studies, e.g. [8] and [9], several studies (Table 1) have observed an extremely low exercise capacity when compared to sedentary healthy controls as illustrated in Figure 1. This is reflected by the low power output and oxygen uptake at the anaerobic threshold (W AT, respectively VO2 AT) or at the ventilatory threshold (W VT, respectively VO2 VT) and at exhaustion (Wmax and VO2max) at a CPET, despite sufficient effort, as implicated by the respiratory exchange ratio at exhaustion (RERmax). The average VO2max of 21.9 mL/min/kg observed at the first CPET by Keller et al.
for example is only 77.1% of the predicted VO2max for age/sex-matched sedentary controls [10].

The left and lower corner of the diamonds indicate the minimum performance levels and the right and upper corner of the diamonds reflect the maximum performance levels of sedentary controls (SC) and ME/CFS patients (P) observed in the exercise tests studies summarized in Table 1a) effort and oxygen uptake the ventilator y or anaerobic threshold, and b) effort and oxygen uptake at maximum effort.

Looking at the observations it seems likely that the exercise capacity of ME/CFS patients is profoundly compromised when compared to sedentary controls. This could explain easy muscle fatigability and profound "lack of energy" in ME/CFS.

<table>
<thead>
<tr>
<th>Study</th>
<th>ME/CFS patients</th>
<th>Sedentary controls</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>WR AT (Watt)</td>
</tr>
<tr>
<td>Snell et al. 2013 [6]</td>
<td>1</td>
<td>49.5 ± 20.4††††</td>
</tr>
<tr>
<td>Keller et al. 2014 [7]</td>
<td>2</td>
<td>51.4 ± 25.0</td>
</tr>
<tr>
<td>Vermeulen et al. 2010 [11] &amp; 2014 [12]</td>
<td>5</td>
<td>8.6 ± 24.2*</td>
</tr>
<tr>
<td>Vermeulen et al. 2014 [12]</td>
<td>78</td>
<td>10.9 ± 2.8*</td>
</tr>
<tr>
<td>De Becker et al. 2000 [13]</td>
<td>27</td>
<td>2.8 ± 1.5***</td>
</tr>
<tr>
<td>Inbar et al. 2001 [14]</td>
<td>5</td>
<td>13.5 ± 3.1***</td>
</tr>
<tr>
<td>Ickmans et al. 2013 [15]</td>
<td>0</td>
<td>114.2 ± 31.3***</td>
</tr>
<tr>
<td>Aerenhouts et al. 2014 [16]</td>
<td>2</td>
<td>117 ± 30.0***</td>
</tr>
</tbody>
</table>

Mean ± SD (rounded to 1 decimal). *: p>0.05; **: p<0.01; ***: p<0.001 between patients and controls.
^: ANOVA p= 0.009; ^^: 0.000^; ^^^: 0.093; ^^^^^: 0.019.
†: Effect size (Cohen’s d): 0.8; ††: 0.4; †††: 1.1; ††††: 0.5.
&: first of two CPETs with 24h rest in-between.
: females; : males.

Table 1: Exercise performance levels of ME/CFS patients at a CPET in comparison with sedentary controls.

Although the enormous difference between the exercise performance levels of ME/CFS patients and those of sedentary controls seems to indicate otherwise, one could argue that this low exercise capacity is simply due to deconditioning. However, a phenomenon which is not observed in sedentary controls or patients with other diseases and seems to be unique to ME (CFS) is the profound negative effect of a CPET on the performance indicators 24 hours later at a second CPET.

Table 2 summarizes studies into the effect of a CPET (CPET1) on the performance at a second CPET (CPET 2) 24 hours in ME/CFS patients (Figure 2). To illustrate the size of the effect of CPET1 on the
performance levels at CPET2, in a recent study by Keller et al. [7] the ME/CFS patient group as a whole showed significant decreases at CPET2 in VO2max (13.8%), Wmax (12.5%), VO2 VT (15.8%), and W VT (21.3%), when compared to CPET1. VO2max decreased in most patients. Patients whose VO2max did not change instead exhibited a decrease in VO2 VT. So, all patients presented with clinically significant decreases in either VO2max and/or VO2 VT exceeding the well-established normative variation of ≤ 7%. Even if ME/CFS patients are able to perform at levels comparable to sedentary controls in the first CPET, as e.g. seen in a study by Snell et al. [6], the performance levels of ME/CFS patients at the second CPET are substantially decreased, while in sedentary controls most performance indicators are slightly improved in the second CPET. Data from the study by Keller et al. [7] and other studies [6,11,12] implicate that aerobic energy-producing systems fail to respond adequately to exercise stress in ME/CFS.

<table>
<thead>
<tr>
<th>Study</th>
<th>N</th>
<th>WR AT (Watt)</th>
<th>VO2 AT (mL/min/kg)</th>
<th>WR VT (Watt)</th>
<th>VO2 VT (mL/min/kg)</th>
<th>Wrtmax (Watt)</th>
<th>VO2max (mL/min/kg)</th>
<th>WR AT (Watt)</th>
<th>VO2 AT (mL/min/kg)</th>
<th>WR VT (Watt)</th>
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<th>Wrtmax (Watt)</th>
<th>VO2max (mL/min/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Snell et al. 2013 [6]</td>
<td>5</td>
<td>1</td>
<td>49.5 ± 20.4</td>
<td>12.7 ± 2.9</td>
<td>109.6 ± 28.9</td>
<td>21.5 ± 4.1</td>
<td>22.2 ± 16.5</td>
<td>(-7.0%, *)</td>
<td>11.4 ± 2.9</td>
<td>(-10.8%, *)</td>
<td>101.6 ± 30.7</td>
<td>(-7.2%, *)</td>
<td>20.4 ± 4.5</td>
</tr>
<tr>
<td>Keller et al. 2014 [7]</td>
<td>2</td>
<td>2</td>
<td>51.4 ± 25.0</td>
<td>12.2 ± 3.7</td>
<td>122.7 ± 28.8</td>
<td>21.9 ± 4.75</td>
<td>41.4 ± 28.8</td>
<td>(-21.3%</td>
<td>9.9 ± 2.89</td>
<td>(-15.8%</td>
<td>003</td>
<td>105.7 ± 33.57</td>
<td>(-12.5%</td>
</tr>
<tr>
<td>Vermeulen et al. 2010 [11]</td>
<td>1</td>
<td>5</td>
<td>58.6 ± 24.2</td>
<td>12.6 ± 3.0</td>
<td>132 ± 30.0</td>
<td>22.3 ± 5.7</td>
<td>54.5 ± 20.9</td>
<td>(-7.0%)</td>
<td>11.9 ± 2.9</td>
<td>(-7.0%)</td>
<td>125.0 ± 35.0</td>
<td>(-5.3%)</td>
<td>20.9 ± 5.5</td>
</tr>
<tr>
<td>VanNess et al. 2006/2007 [17,18]</td>
<td>6</td>
<td>1</td>
<td>15.0 ± 4.9</td>
<td>26.2 ± 4.9</td>
<td>11.0 ± 3.4</td>
<td>(-26.7%, *)</td>
<td>20.5 ± 1.8</td>
<td>(-21.8%</td>
<td>012)</td>
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</table>

Table 2: Exercise performance levels of ME/CFS patients at repeated CPETS.

The Impact of Exertion on Cognitive Functioning

Although contradicted by some studies, several studies observed mild to substantial deficits in the speed of processing of simple and complex information [19] and in cognitive tasks which require working memory over a longer period of time (endurance) [19,20], (concurren) processing of information [21,22] and conflict-monitoring (interference control) [23]. Cognitive deficits, especially impairments related to attention, memory and reaction times, only seem to manifest themselves if adequate tests and measured are employed [24]. This relevant observation is substantiated by a meta review of 79 cognitive tests applied in studies [19]. All tests for attention, including attention span and working memory, showed significant deficits in ME/CFS and all five tests aimed at reaction times disclose impairment for responses to both simple and complex (choice) stimuli. However, specific word list learning and recall tests seem to be more sensitive than others to the memory deficits reported by ME/CFS patients.

Although research into the effects of physical exercise on cognitive functioning is limited, studies seem to implicate that exertion has a (prolonged) negative impact on cognitive performance, e.g. focused and sustained attention [25], simple reaction time and choice reaction times [26], cognitive speed and disinhibition, psychomotor speed and vigilance [27]. This negative effect seems to oppose the positive effect of exercise on cognitive performance observed in sedentary controls.
A recent study [33] found that head-up tilt (HUT) induced a sequence. Various studies have observed reduced muscle power (strength and endurance) in ME/CFS. Neu et al. [34] e.g. found reduced hand grip strength during tonic trials. They also observed that patients exhibited higher fatigability during phasic trials, in which a subject must grip the handle as strongly as possible and relax immediately during successive trials, with a significant decline of the maximal grip strength at the end of the ten phasic trials. Siemionow et al. [35] observed that the hand grip strength of patients was significantly lower than that of sedentary controls and that the maximum voluntary contraction force in the ME/CFS group dropped significantly to 83% of the strength at baseline after 50 contractions with 10s and 50 contractions with 5s rest between contractions. Lawrie et al. [36] observed that patients were able to sustain a 10 kg handgrip contraction less shorter than healthy controls, and Fulcher and White [37] found that the maximum twitch interpolated isometric contraction force of the quadriceps muscle of the dominant leg was significantly lower patients compared to sedentary controls.

Next to reduced muscle power (endurance), recovery of muscle exercise seems to be prolonged or inadequate. In a study by Paul et al. [38] ME/CFS patients and healthy sedentary controls were subjected to an exercise comprising 18 maximum voluntary contractions (MVCs) of the quadriceps (10s contraction, 10s rest). The MVC forces in the controls were consistently higher than those of the patients, with a decline in force during the exercise in both groups. Recovery was prolonged in the ME/CFS patient group, with a significant fall in the MVCs in the recovery phase (200 minutes) and at 24 hours post-exercise (73 ± 9% of the baseline MVC force levels). Recently Meeus et al. [39] found that patients exhibited reduced MVC levels for handgrip strength and insufficient, delayed recovery in the 45 minutes after intensive muscle exercise (18 maximum contractions lasting 5 seconds with 5 seconds rest in-between), when compared to MS patients and sedentary controls.

**Discussion**

Muscle (weakness), cognitive deficits, neurological impairments, and in particular post-exertional “malaise” are mandatory elements of ME and optional features for the diagnosis CFS: (persistent or relapsing) chronic fatigue. There is controversy about the diagnosis, the nature of the symptoms and potentially effective therapies.

Virtually all research into ME and CFS up to day is based upon subjective measures and questionnaires. In order to resolve the debate and to develop effective therapies for ME and CFS, it seems essential to make a distinction between people with muscle (weakness), cognitive deficits and post-exertional “malaise”, designated as ME, and patients with “chronic fatigue” but without this characteristic symptoms, labeled as CFS, and to assess ambiguous symptoms, e.g. muscle weakness and post-exertional “malaise”, employing objective methods. The relevance of a distinction between ME and CFS has been contested by some authors [3,40,41]. Although one could ask oneself whether the new diagnostic criteria for ME [3] are too restrictive or not, but numerous studies have pointed out that the need for a more concise definition of ME and emphasized the disparity between patients with distinctive symptoms, e.g. post-exertional “malaise” and patient without these symptoms [42].

Objective methods for assessing characteristic symptoms and pattern recognition analysis methods for diagnostic criteria should be employed to develop more effective strategies for future research into ME and CFS and for clinical practice.

Although various aspects of post-exertional malaise (an amplification of symptoms after a minor exertion) can be assessed objectively, e.g. the effects of exercise and orthostatic stress on physical functioning, cognitive performance, and muscle power, due to their subjective nature, other aspects of post-exertional malaise, e.g. a (prolonged) increase of pain and “fatigue” after exertion, cannot be quantified. However, looking at various findings [1] exercise-induced exacerbation of these symptoms can plausibly be explained, e.g. by an long-lasting increase of metabolite-responsive pain receptors and a impaired stress system response to exertion [43].

**Conclusion**

Post-exertional malaise, a long-lasting increase of typical symptoms after what is considered to be a minor exertion, is a distinctive
phenomenon of ME (CFS). Several studies have demonstrated that the potentially profound impact of exertion and orthostatic stress on characteristic symptoms of ME (CFS), e.g. physical functioning, cognitive symptoms and muscle weakness, can be quantified using objective tests (Figure 3). To end the debate about the diagnosis, the nature of the symptoms and the effect of therapies, objective methods and measures should be employed.

References


