MEA INFORMATION ON
Post-viral fatigue (PVF)
and Post-viral fatigue syndrome (PVFS)
following coronavirus infection

May 2020
INTRODUCTION

We are starting to receive reports about previously healthy people who have had (or probably had) coronavirus infection and have not been able to return to their normal level of health and energy levels in the weeks following the onset of symptoms. These reports are largely from people who have managed at home and not had a more serious infection that required hospital admission. Some reports are from health professionals. It seems likely that some of them are experiencing what is called post viral fatigue (PVF), or a post viral fatigue syndrome (PVFS).

We are also receiving reports from people with ME/CFS (myalgic encephalomyelitis/chronic fatigue syndrome) who have had this infection and now have a significant exacerbation of their ME/CFS symptoms - especially a further reduction in energy levels.

The ME Association has a lot of experience in dealing with people who develop prolonged and debilitating fatigue (sometimes with other symptoms as well) following a viral infection – as well as people with ME/CFS who relapse following another infection.

We are now expecting to see a number of new cases of ME/CFS that follow coronavirus infection fatigue. This is why we have produced some guidance on how we feel that convalescence and good basic management of post infection fatigue can lessen the chances of this turning into a more permanent and debilitating illness.

BACKGROUND

WHAT IS POST VIRAL FATIGUE (PVF) AND POST-VIRAL FATIGUE SYNDROME (PVFS)?

Some degree of post-viral fatigue (PVF) or debility is a fairly common occurrence after any type of viral infection. Fortunately, in most cases, this is short lived and there is a steady return to normal health over a period of a few weeks.

However, in some cases, a full return to normal health takes months rather than weeks. Additional symptoms may also develop, where the term post-viral fatigue syndrome (PVFS) may be a more appropriate diagnosis.

The situation with persisting fatigue following coronavirus infection appears to be rather more complicated than what happens with other viral illnesses. It could also be more serious - as fatigue and lack of energy are turning out to be a very characteristic symptom of coronavirus infection.

The precise explanation for what causes post-infection/viral fatigue remains uncertain. But one of the reasons why people have fatigue, loss of energy, muscular aches and pains, and generally feel unwell, when they have an acute infection is the production of chemicals called cytokines by the body’s immune system. These immune system chemicals form part of the front line attack on any viral infection. And it is interesting to note that in people who develop serious respiratory complications from coronavirus infection, this may be due to an overactive immune response involving what is being termed a ‘cytokine surge’.

When fatigue and flu-like symptoms persist for a longer period of time once the acute infection is over, as they do in ME/CFS, there is research evidence to indicate that what is a perfectly normal immune system response to the acute
Post-viral fatigue (PVF) and Post-viral fatigue syndrome (PVFS) following coronavirus infection

**BACKGROUND (cont.)**

infection has not returned to normal.

It is also possible that, as happens with ME/CFS, there is a problem with the way that energy production is taking place at a cellular level in structures called mitochondria.

We are still on a very steep learning curve when it comes to understanding how the coronavirus behaves. However, there is no evidence to indicate that it persists like hepatitis C infection or HIV. So the continuing fatigue does not appear to be due to a persisting viral infection. Consequently, people who have continuing fatigue, but no other coronavirus symptoms, are no longer infective to others in our current state of knowledge.

While most people with post-infection fatigue will improve and return to normal health, good management during the very early stage is an important factor in trying to help any natural recovery process take place. And patient evidence also indicates that good initial management of post-infection fatigue lessens the chances of this turning into an ME/CFS-like illness.

**WHAT ARE THE SYMPTOMS OF PVF AND PVFS?**

**ACTIVITY INDUCED FATIGUE**

As the name post-viral fatigue indicates, the main symptom is fatigue, or a failure to return to normal energy levels.

In practical terms this means no longer being able to carry out a normal range of physical activities in the home, at work or school, or anywhere else. In terms of severity this can range from a relatively mild impairment to a really quite severe functional impairment.

A useful comparison here is trying to operate a radio with a battery that no longer functions properly. The radio may work for short periods of time but then starts to fail quite quickly. After a rest it will work again - but only for another short period.

In a similar manner, people with PVF find they can carry out short periods of physical and/or mental activity but then have to stop and rest and ‘recharge their batteries’. They are no longer able to sustain normal levels of physical activity and find anything that involves using a large amount of energy in a short period of time, such as going for a run, is no longer possible.

**SLEEP DISTURBANCE**

Some form of sleep disturbance often accompanies PVF.

During the actual infection, and shortly afterwards, this may involve sleeping for much longer than usual - as well as wanting to sleep for periods during the day. This is known as hypersomnia and is a natural part of the way in which human body responds to and recovers from an infective illness.

**COGNITIVE DYSFUNCTION**

Just as energy is required for physical activity, it is also required for mental activity. So in a similar way to easily running out of energy when doing a physical task, some people cannot cope with longer periods of mental activity and start to lose concentration and their ability to process and retrieve information. Short-term (working) memory may also be affected.

**PVFS AND POSSIBLE PROGRESSION TO ME/CFS**

When the symptoms discussed above persist, and are accompanied by other flu like symptoms, it may be more appropriate to diagnose a post-viral fatigue syndrome (PVFS).

If the symptoms persist for more than two to three months, and continue to cause a significant degree of functional impairment that is preventing a return to education, employment or other key activities, a diagnosis of ME/CFS should also be considered.
SYMPTOMS WHICH CAN FORM PART OF A PVFS AND ARE ALSO SUGGESTIVE OF A DIAGNOSIS OF ME/CFS INCLUDE:

- Alcohol intolerance
- Feeling wobbly or unsteady
- Headaches of new onset
- Difficulty in sustaining an upright posture for long periods – known as orthostatic intolerance
- Muscular aches and pains
- Sore throats and tender glands
- Problems with temperature control
- Post exertional malaise/symptom exacerbation
- The MEA has an information leaflet providing more guidance on the early and accurate diagnosis of ME/CFS HERE.

CONVALESCENCE

The most important aspect of management of post infection fatigue involves good old fashioned convalescence. In practice, this means taking things easy, and having plenty of rest and relaxation. This should be combined with gentle activity within your limitations, having a good night’s sleep, eating a healthy diet, avoiding stressful situations, and not returning to work, school or domestic duties until you are well again.

Some people find that adding in approaches like meditation or yoga can help with relaxation. Others do not.

Overall progress can be very variable with some people making a steady improvement back to normal health. Others follow a less predictable pattern with good days and bad days.

Doing too much on a good day will often lead to an exacerbation of fatigue and any other symptoms the following day. This characteristic delay in symptom exacerbation is known as post-exertional malaise (PEM). As already noted, where this persists for more than two to three months it should certainly raise the possibility of a diagnosis of ME/CFS.

ACTIVITY MANAGEMENT

Resting and relaxing are both essential components of any convalescence. As energy is required for both physical and mental activity, it’s important to make sure that you are not exceeding what you feel comfortable doing, stopping an activity when you are starting to feel more fatigued, and not carrying on with a task when you are not able to do so.

Finding the right balance on an individual basis between activity and rest is a process called pacing and many people find it helpful to alternate small flexible amounts of physical and mental activities with a longer period of rest/relaxation in between.

Physical activities could include things like household tasks and going for a short walk – when you feel ready to do so. Mental activities could include social activities with people you can still be in contact with, reading, listening to the radio, watching some TV, or doing a small amount of computer work.

Do try to limit screen time on computers, phones and TV, and avoid using electronic devices to catch up on your normal work!

Having established what is a safe baseline of physical and mental activity that is not exacerbating symptoms, the next step is to gradually try and increase the amount you are doing – whilst sticking to the rule of not exceeding limitations and living within what is called your ‘energy envelope’.

Activity management needs to be carried out in a very flexible manner. So any increase in activity levels need to be gradual and within your physical and mental limitations.
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MANAGEMENT OF PVF AND PVFS

Any kind of vigorous, sporting or prolonged physical activity should be given a complete miss until you have fully recovered and remained well for a period of time.

There is clearly quite a lot of trial and error involved in getting the balance right between activity and rest and great care needs to be taken to make sure that you are only doing what you are comfortable with. You cannot exercise your way out of PVF or PVFS!

MENTAL WELL BEING

Looking after your mental and emotional health at this time is another important part of management. In particular, this should include avoiding stressful situations wherever possible.

Not surprisingly, having PVF or PVFS, along with all the restrictions that this imposes on what you are able to do, can make people feel fed up, frustrated and sometimes even anxious or depressed.

If you are feeling anxious, or your mood is obviously low, and you are feeling depressed (with tearfulness, loss of interest in activities, loss of self-esteem etc), you must speak to your GP about how this should be managed.

NUTRITION

Some people will lose a significant amount of weight during a serious acute infection. This type of weight loss may need to be built up again if you are not overweight before being ill.

Try to stick to a healthy balanced diet with plenty of fresh fruit and vegetables, possibly with small frequent meals if you find this easier.

Make sure your fluid intake is adequate. It’s probably better to avoid alcohol, or cut right down and abstain, especially if alcohol is making you feel worse.

If your appetite is still poor, or you need to put on some weight again, ask for help from a dietitian on foods that are worth trying.

SLEEP

As already noted, having an increased sleep requirement (hypersomnia) is an important part of the body’s natural healing process during, and sometimes following, an acute infection.

So whilst going bed and staying there day and night is not the solution, making sure that you have plenty of good quality sleep, possibly including a daytime nap if you need one, is a key part of good management.

There are a number of simple self-help strategies that can improve sleep quality:

- Relax before going to bed by having a warm bath or listening to a relaxation CD.
- Avoid caffeine-containing drinks – coffee, cola, tea – in the evening. Caffeine interferes with the natural process of going to sleep. Try a warm milky drink instead.
- Avoid heavy meals and alcohol shortly before going to bed. Alcohol may help you to get off to sleep but it can disrupt sleep during the night.
- Try to establish a routine for getting up and going to bed at roughly the same time each day that also accommodates any increased sleep requirements.
- Keep the bedroom as a quiet, peaceful and relaxing place for going to sleep – not too hot or too cold and not a place for watching TV or doing computer work.
- If you don’t have increased sleep requirements, or the need to sleep for excessive periods has reduced, try to
have eight or nine hours solid uninterrupted sleep each night
The MEA has an information leaflet covering all aspects of sleep management in more detail [HERE](#).

**WORK AND EDUCATION**
The simple message here is to switch off from anything to do with work and do not go back to work until you are fit to do so.
This is especially important if your job involves a significant amount of hard physical activity or high levels of mental stress.
Do keep in touch with your employer or education provider - as there may be a number of options to explore once you start to improve and reach a point where a return to work or school/college looks possible.
This could involve doing some work from home, or having a part time or flexible return to normal activities at school/college or duties in the workplace.

**FINANCES**
If you are employed, and entitled to state or other sickness benefits, you will need to discuss sick pay, including what is called statutory sick pay, with your employer and obtain a fit note (aka a sick note) from your GP, or other health professional who can confirm that you are unable to return to work. A fit note will inform your employer that the health professional has confirmed that you are either ‘not fit for work’ or ‘may be fit for work’.

Information from the Citizens Advice Bureau on Statutory Sick Pay
There are other state benefits from the DWP, including Universal Credit, that you may be entitled to during the coronavirus epidemic and lockdown. The MEA has produced a guide to benefits that might help you to navigate the benefit system and other guides to specific types of benefit that may be needed in the longer term if your health does not improve.

**DRUG TREATMENTS**
Finally, it’s worth noting that there are no drug treatments or supplements for treating PVF or PVFS. And there is no value in taking products like iron supplements (which are unnecessary and can even be harmful) and high doses of vitamin C or any other vitamins. However, if you are not having a healthy balanced diet it would be worth taking a multivitamin supplement.
As many people are no longer going outside in the sunshine, or only doing so for short periods, the risk of vitamin D deficiency is increasing. Vitamin D is made in the body in response to sunshine and is essential for bone and muscle health - so taking a daily 10 microgram vitamin D supplement is a sensible precaution.
Do talk to your GP or pharmacist if you require help with a specific symptom such as muscle aches and pains.
The MEA has information leaflets providing detailed information on the drug and non drug management of orthostatic intolerance, pain and sleep disturbance.

WHEN TO CHECK WITH YOUR GP
Although a face-to-face consultation with your GP may be difficult to arrange, most GPs are happy to speak to their patients on the phone or by a video link at the moment.
If your post-infectious fatigue is continuing beyond four weeks, and is not improving, it’s still worth speaking to your GP to keep him/her appraised of the situation - even though there is no specific drug treatment. You may also need help from a GP when it comes to sick pay and benefit applications.
The diagnosis of PVF and PVFS is made on your symptoms. There is no diagnostic blood test. However, if you are having symptoms that are not consistent with post-infection fatigue, blood tests will be arranged to check for other conditions that can cause persisting fatigue (such as hypothyroidism/low thyroid function).

If you have any of the following symptoms, which are not part of PVF or PVFS, you must speak to your GP:

- Continuing fever or periodic spikes of fever
- Persisting cough, shortness of breath or respiratory symptoms
- Weight loss
- Feeling depressed

- If symptoms are still present after two to three months, and are still having a significant impact on daily living, consideration will have to be given as to whether a PVFS has progressed into ME/CFS. As with PVF and PVFS, this is a diagnosis based on symptoms. There is no diagnostic blood test for ME/CFS.

The MEA has an information leaflet providing detailed guidance on the early and accurate diagnosis of ME/CFS [HERE].

Several small epidemiological studies have looked at the development of fatigue syndromes following an acute infection.

An Australian research group followed up 253 people with glandular fever (Epstein Barr virus infection) Q fever (Coxiella burnetii) and Ross River Fever (epidemic polyarthritis) in a rural part of Australia at regular intervals over a period of 12 months. Approximately 12% developed a post-infection syndrome involving fatigue, cognitive dysfunction, musculoskeletal pain and mood disturbance. The predictability of developing a post-infectious fatigue syndrome was largely determined by the severity of the triggering infection and not by any demographic or psychological factors, or the type of infection that triggered the symptoms.

Reference [HERE].

During an outbreak of SARS (Severe Acute Respiratory Syndrome - another coronavirus infection) in Toronto during 2002 and 2003, 273 people were diagnosed with the infection, of whom 44 died. After the outbreak ended, researchers at the University of Toronto, studied 22 people, mostly health workers, who had on-going health problems that prevented them from returning to work.

They concluded that there was a chronic post SARS infection syndrome characterised by persistent fatigue, non-restorative sleep, daytime fatigue, pain and weakness in muscles all over their body, and depression.

Reference [HERE].

As to how research into post infective fatigue and ME/CFS might be carried out after an acute coronavirus infection is over, this was discussed at a recent meeting of the ME Biobank Steering Group - as there are epidemiologists at the London School of Hygiene and Tropical Medicine, some of whom are looking at Cv19 - and at the last CMRC (CFS/ME Research Collaborative) Board meeting.

One possible route here would be to make use of the baseline information on people who already have clinical data and blood samples stored at the UK Biobank but did not have PVF, PVFS or ME/CFS at the time of enrolment, and then go on to develop prolonged fatigue, or a fatigue syndrome following Cv19 infection.

We also need to look at how the immune system is responding, especially the levels of various cytokines, in people are not recovering and have post-infectious fatigue. The MEA Ramsay Research Fund would be very happy to consider any research proposals to investigate immune system status in coronavirus infection.

FURTHER INFORMATION

Article on coronavirus fatigue that appeared in the New Scientist

MEA website information on coronavirus infection and ME/CFS
ME ASSOCIATION WORKING ARRANGEMENTS

We issued a statement to provide information on all aspects of our work and the services we provide. Almost all key activities are continuing as normal – the main change being that we have now closed Head Office in Buckingham and office staff are now working from home.

The main impact here is that we will not be able to send out any paper literature, purple books, or merchandise from the office for the foreseeable future. ME Connect – the MEA telephone helpline – remains operational, seven days a week, for information and support.

ME Association statement:
The ME Association and Coronavirus: New working arrangements

OTHER INFORMATION

This information and guidance is based on patient evidence collected over many years on the symptoms and management of PVF and PVFS. The author has personal experience of a PVFS following chickenpox infection, which was caught from a patient. No form of clinical trial research has been carried out and published into the management of PVF and PVFS – so this information and guidance is largely based on clinician and patient evidence.

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ME CONNECT
We’re here to help

Do you need to talk?
CALL
0844 576 5326
10am-12noon
2pm-4pm, 7pm-9pm
every day of the year

THE ME ASSOCIATION:

- Provides information on M.E. and campaigns on issues such as research, the NICE guideline, NHS service provision and care
- Provides support through our ME Connect helpline, ME Essential members magazine and our website and social media
- Funds biomedical research – including the UK ME/CFS Biobank which is managed by an expert team at the London School of Hygiene and Tropical Medicine – through the Ramsay Research Fund
- Is a member of the Forward ME Group of charities and patient representatives that is chaired by the Countess of Mar, and the CFS/ME Research Collaborative, chaired by Professor Stephen Holgate, which aims to raise the profile of M.E. and attract greater research investment

CORONAVIRUS AND ME/CFS

We have a new MEA information leaflet covering Coronavirus and ME/CFS which is updated weekly. This is available as a free download on our website.