Fatigue Matters
A self-help guide for people living with rheumatoid arthritis
ABOUT US

The National Rheumatoid Arthritis Society (NRAS) is the only patient-led organisation in the UK specialising in rheumatoid arthritis (RA) and juvenile idiopathic arthritis (JIA). Due to its targeted focus on RA and JIA, NRAS provides truly expert and wide-ranging services to support, educate and campaign for, people living with these complex autoimmune conditions, as well as their families and the health professionals who treat them.

ASK US

Call our freephone helpline on 0800 298 7650, it is open from 9.30am to 4.30pm, Monday to Friday. Our trained helpline staff are there to answer your questions on all aspects of living with RA.

If you’d like to talk to someone else with RA, our staff can match you with one of our trained Volunteers, who will then call you back at a mutually convenient time to discuss whatever aspect of living with RA most concerns you. To be put in touch with a Volunteer who has RA, please call the helpline to make the arrangements.

Our website www.nras.org.uk has a wealth of information about all aspects of living with RA, treatments, the latest research and developments, as well as full details of other useful organisations that can help.

The NRAS online forum www.healthunlocked.com/nras is a safe space where you can get peer support and blog about your experiences.

If you don’t have access to the internet and want information sent to you in the post, please call us on 0845 458 3969

MEET US

Local NRAS groups meet regularly around the country. To find out if there is a group near you, visit www.nras.org.uk/groups, email groups@nras.org.uk or call 0845 458 3969

JOIN US

To find out how to join NRAS and support the work we do, thereby helping us to help even more people, please get in touch (see inside back cover of this booklet for full contact details).
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Introduction

Ailsa Bosworth MBE
Chief Executive,
National Rheumatoid Arthritis Society (NRAS)

In 2014 we conducted a UK-wide survey on fatigue which drew over 2000 completed responses. We believe this is the first time that so many people with RA have been surveyed about their experiences of chronic fatigue – including their views about levels of public awareness of the symptom, the effectiveness of existing management approaches, and the impact of chronic fatigue on their quality of life and work outcomes. The respondents ranked chronic fatigue as the least understood symptom of RA by the public. Nine out of ten respondents said they suffered with chronic fatigue, of which nearly two thirds said onset appeared before diagnosis. Equally troubling, the majority of respondents said they experienced chronic fatigue on three or more days during the previous week and that this lasted for several hours at a time. Over three-quarters of respondents said fatigue had a high or medium-level impact. Two-thirds of all respondents said their healthcare professional never or rarely asked them about their fatigue. The impact of fatigue on work was also very significant, with just under three-quarters of working-age people with RA who were not in work, saying that fatigue had contributed to their unemployment. These findings caused us to commit to developing this new publication devoted to the subject of fatigue.

We collaborated with King’s College Hospital to provide a work placement/internship for one of their Health Psychology students to work with us on the framework for this booklet in early 2017. I am extremely grateful to Dr Katie Druce for contributing her great insight and experience on the subject of fatigue to this booklet. Our Editor, Colin Richardson has done a brilliant job of editing all this work to create this informative and insightful resource, and thanks also go to Dr. Katie Hackett for peer reviewing the booklet. We know that healthcare professionals find it difficult to ‘treat’ fatigue as there is no magic pill or instant resolution to fatigue. As a consequence, and maybe because health professionals like to be able to provide solutions, it isn’t always discussed at clinic appointments. This means that patients often feel a bit isolated in their battle with this debilitating symptom. We therefore hope that health professionals will find this booklet a very useful resource to give to their patients. All at NRAS sincerely hope that you find this new and comprehensive booklet useful in gaining better control over the problems you may be experiencing with fatigue in your daily life.
I am delighted to provide the foreword for *Fatigue Matters*. People with rheumatic diseases, such as RA, tell us that fatigue is one of their most important and distressing symptoms. They also tell us that managing fatigue is problematic. Flares or fluctuations may occur out of the blue, for no obvious reason, and effective treatment options to manage fatigue are limited.

It is for these reasons that we often hear from people with RA that they try to manage their fatigue by themselves. However, they often feel like they are undertaking a constant battle of trial and error. The problem is that fatigue is complex. There is no one-size-fits-all experience of fatigue and many things may cause a symptom flare.

The management of fatigue is also difficult because it is invisible to others and therefore hard to relate to. As a researcher into fatigue I often hear people say that their healthcare providers, friends and family members don’t care about their fatigue. But I really believe the truth is that they don’t really understand it. Most of us know what it feels like to be tired, but few of us can understand the overwhelming feeling of physical and mental exhaustion reported by people with RA. Even as a researcher in this area, I am constantly surprised by the descriptions of fatigue that I hear: “It’s like walking through treacle”; “I feel like I’ve got a back-pack full of stones on my back”; “I feel absolutely fine and then suddenly – BAM! – it hits me and I feel like I’ve got a full suit of armour and chainmail on.”

That’s why one of the most important themes in this booklet is to talk about fatigue. Whether you are someone with RA, a healthcare provider, friend or family member, I urge you to talk more about fatigue. As someone with fatigue this is particularly important because people can’t support you to manage your fatigue unless they understand your experience of it. If you don’t yet feel ready to talk about your fatigue, there’s plenty of other help and advice in this booklet, including the importance of sleep, exercise and forward-planning.

In creating this booklet, NRAS has done a wonderful job of collating a number of resources, which I hope will help improve the management of fatigue. Whether you are someone with RA, a healthcare provider, or someone who knows a friend or family member with the condition, I hope that this booklet can help you.
What is fatigue?

Most people with RA experience fatigue, one of the most common and distressing symptoms of RA.

*I feel totally exhausted; like a tap suddenly opens and all my energy (what little I have) disappears in an instant. It can last hours, days, weeks or months.*

*I feel like I have limbs of lead, and even completing the smallest of tasks leaves me totally drained. It also leaves me feeling tired emotionally.*

*My fatigue makes me feel helpless, and unable to complete everyday tasks. One of the worst effects is how it affects my thinking. I feel unable to cope in social situations with strangers. It is a blight on every aspect of my life.*

All quotes from the 2014 NRAS survey: Invisible disease: Rheumatoid arthritis and chronic fatigue

Fatigue can have a severe impact on the quality of life of those affected. A survey conducted by NRAS (*Invisible Disease: Rheumatoid Arthritis and Chronic Fatigue*, 2014) found that 89 per cent of those surveyed experienced chronic fatigue, of whom getting on for half (40 per cent) said their fatigue was persistent and severe.

Fatigue can come on at any time of the day without warning. It can last from a few hours to several days at a time. For some it can persist for weeks or months. It tends to be worse during a flare.

Fatigue is so much more than ordinary tiredness. It is much more debilitating and all-encompassing than that. Fatigue is to tiredness as ‘flu is to the common cold.
Fatigue affects the great majority of people with RA.

Fatigue is a major but hidden symptom of RA.

Fatigue is an overwhelming feeling of exhaustion, both physical and mental.

Fatigue can come on at any time, without warning. It can last for a few hours or days but may persist for weeks or months.

Individual experiences of fatigue vary – it is not the same for everyone

These are some of the ways people experience it:

- Fatigue can be an overwhelming feeling of exhaustion, both physical and mental. You may feel completely drained of energy. You may just want to lie down and never get up again.

- Your limbs may feel heavy. Walking may feel like wading through treacle while wearing concrete boots. It may feel as though you have a heavy load on your back.

- You may find it hard to think or concentrate or perform the simplest of tasks. Some people talk of ‘brain fog’ descending out of a clear blue sky!

Fatigue is a ‘hidden symptom’ of RA

There is no agreed medical definition of fatigue and no easy way to measure it. People who experience it often find that others may not be very understanding of what it is nor of its effects. Unless you tell them, other people may not be aware that you have fatigue. The NRAS survey mentioned above (Invisible Disease, 2014) found that only 31 per cent – less than a third – of the general public identified fatigue as a symptom of rheumatoid arthritis (RA).

It can be frustrating having to explain all the time to people who don’t understand that you are not just tired. It is easy to feel that others think you are malingering or making it up. You may even start to believe it yourself. But when you do start to talk about it, you will find that people become more understanding and you will have to explain yourself less and less. And you will begin to find ways of coping with and tackling fatigue that work best for you.
What causes fatigue?

The causes of fatigue in RA are complex and poorly understood.

There is no one reason why most people with RA experience fatigue at one time or another. Rather, there are several overlapping, interwoven and fluctuating factors at play, which, as we shall see later on, have to be tackled on several fronts.

Research indicates that it is principally the effects of having RA that can contribute to fatigue – notably, pain, lack of sleep, low mood, disease activity, disability, inactivity, and social isolation.

Pain

There is a strong link between levels of pain and levels of fatigue. Pain can contribute to fatigue directly or indirectly. However, while effective pain control may reduce fatigue in many people with RA, it is not the solution for everyone.

Pain can be very tiring. Being in constant pain is extremely wearing and if you have to modify your movements because of pain and/or disability, you will likely find yourself expending more energy than usual.
Pain can also contribute to fatigue indirectly by disturbing sleep and/or by causing low mood or depression.

- The causes of fatigue in RA are poorly understood.
- There is no one cause of fatigue in RA. Instead there are many overlapping and interwoven factors that lie behind it.
- Some of the main symptoms of RA, if not properly treated, can contribute to fatigue. Pain is most strongly linked with fatigue.
- At the same time, fatigue may make coping with the other symptoms of RA more difficult.

Poor sleep

Inadequate or disturbed sleep makes fatigue worse. Sleep helps the body to recover and repair and to restore energy levels. If you do not sleep properly or for long enough, you will have less energy to deal with the day ahead; you won’t feel rested, your joints may feel stiffer and more painful than usual. Lack of sleep may also depress your mood.

Therefore, establishing good sleeping habits (sometimes referred to by health professionals as ‘sleep hygiene’) can help and include:

- establishing fixed times for going to bed and waking up
- creating a relaxing bedtime routine
- only going to bed when you feel tired

- maintaining a comfortable sleeping environment that’s not too hot, cold, noisy or bright
- not napping during the day
- avoiding caffeine, nicotine and alcohol late and avoiding eating a heavy meal late at night.

Talk to your GP or specialist nurse if you continue to have poor sleep as they will be able to help. See also the NRAS leaflet on **Sleep Hygiene**.

Low mood

Low mood is also strongly linked with fatigue. Excessive tiredness can be a symptom of depression. And having fatigue may itself cause you to be depressed. In our survey of people with RA (Invisible disease, 2014), almost all those surveyed (90 per cent) said fatigue had caused them to feel down or depressed during the previous week, of whom over a third said they felt very down or depressed.

Depression can also make everything seem worse than it is. If you feel
depressed, your fatigue may feel more than usually severe and you may feel that there is nothing that can be done about it. This is entirely understandable and you shouldn’t blame yourself for feeling like that.

Disease activity

There is a link between disease activity and fatigue in RA, but the exact nature of the link isn’t clear and isn’t the same for everyone. For some people, controlling their disease to the point that they are considered to be in remission leads to a big improvement in the severity and duration of their fatigue. It may be that being in remission helps to minimise the other symptoms of RA, which may, in turn, lead to a reduction in fatigue. But for others, remission may have little or no impact on fatigue. In either case, minimising disease activity remains the central aim of the treatment of RA, whatever its impact on reducing fatigue, because of its overall beneficial effect on health in the long-term.

Other factors

Your level of physical functioning can affect your fatigue. RA can lead to disability, which, in turn, can affect your fatigue. Disability can cause physical activity to be harder and you may find that your activities become restricted. You may find that the changes you have to make to compensate for these restrictions mean that even the most basic tasks require more effort, leading to an increase in fatigue. You may also find that when you decrease your levels of physical activity your muscles become weaker, which means that physical tasks require more effort to complete.

Social isolation may be linked to fatigue. When you have fatigue, ordinary social activities, such as going to the cinema or visiting family or friends, may be beyond you. You may be
too exhausted to contemplate leaving the house. At the same time, the unpredictability of fatigue can make it difficult to plan ahead. You may find yourself cancelling arrangements at the last minute. And because constantly having to explain yourself can be very tiring, your friends may not understand and may not ask you out so often in future. All that means you may have less support than you otherwise might; and you will probably find your mood becoming depressed, too.

**Eating well is important.** A poor diet can make you feel run-down and tired.

Some medications can make you feel drowsy. A review of your medications can help identify those that are making you feel sleepy and it may well be possible to find substitutes.

It can be difficult to adjust to having fatigue. You may try to push through it and continue with your normal level of activity. However, **overdoing it is likely to worsen your fatigue.**

As we will discuss later on, learning to rest and pace yourself will help you cope better with your fatigue.
Are there any treatments for fatigue in RA?

It is important to be clear that there are no licensed treatments specifically for fatigue.

There is no cure for fatigue. But scientists are working hard to find out more about what causes fatigue in RA. They hope to find effective treatments and even a cure.

It is equally important to say that there are plenty of things that can be done to reduce fatigue and to make it easier to manage.

- There is, at present, no cure for fatigue.
- But there are plenty of things that can be done to reduce your fatigue and to help you cope better.
- Emotional support, a medication review, effective pain control, exercise, getting a good night’s sleep, and planning your day to conserve energy can all help.

Emotional support

There is much evidence that talking about fatigue is good for you. First of all, by talking to others about your fatigue, you can start to get the help and support you need. Secondly, ‘talking therapies’ such as cognitive behavioural therapy (CBT), have been shown to be very effective in tackling fatigue by combating depression and low mood and by helping you develop better coping strategies. Anxiety and stress are also commonly experienced by people with RA – often, hand in hand with depression. Feeling stressed and anxious is very wearing and can worsen your fatigue and your ability to manage it. Any ‘talking therapy’ aimed at tackling depression will also look at ways of combating anxiety and stress.

A medication review

Some medications can cause drowsiness, which can make fatigue worse; some may even list fatigue as a known side effect. Ask your clinician, pharmacist or healthcare provider for a review of all the medications you take – not just those you take for your RA – which should identify any that might be contributing to fatigue. It may be possible to find a substitute for any medication that is making you feel sleepy.

Pain control

Effective pain control can greatly reduce fatigue for many people, though it may not be the answer for everyone. In either case, keeping pain under control is a key element of the
treatment of RA. Even if it has no direct effect on your fatigue, minimising pain will improve other symptoms of RA (such as poor sleep) and will make it easier to cope.

It is important to note that increased or high levels of pain may indicate that your disease is not being well-controlled. If you think this may be the case, seek a review with your rheumatology team by contacting the rheumatology nurse helpline at your hospital.

**Exercise**

It may seem to go against common sense to suggest that a way to deal with fatigue is to take some exercise, but it really can help. Most people with RA are affected by a condition known as *rheumatoid cachexia*, which causes muscle wastage and an increase in body fat. When you lose muscle, it takes more energy to accomplish even the simplest of physical tasks. And an increase in weight will also mean you expend more effort on everyday activities, as well as not being good for your general health. But muscle wasting and an increase in weight will not automatically be reversed when your RA is under control. Exercise will help you recover muscle mass, thereby building up your strength, and decrease body fat, which will bring your weight down. Strengthening your muscles and losing weight will both help to counter fatigue. Exercise can also lift your mood and contribute towards a good night’s sleep, both of which will help alleviate fatigue.

**A good night’s sleep**

There’s little evidence that tackling poor sleep has a direct effect on fatigue. But sleeping well will make you feel better, which will help you cope better. You will likely have more energy
if you have slept well than if you have slept badly. And poor sleep may indicate that other things are wrong, so it should not be ignored.

**Sleep, page 9**

Planning ahead and conserving your energy

Although keeping active and taking exercise is an important part of any programme for tackling fatigue, it is also important not to overdo it. Fatigue limits your available energy. Planning ahead can help you make the most of the day. Knowing when to rest and when to be active, when to ask for help and when to delegate tasks to others are all about good self-management and part of any energy conservation programme. Later in this booklet, we have a wealth of tips and ideas for getting the best out of your available energy.

**Planning Ahead, page 18**

At the same time, people who are in employment may find it much harder to organise their time and incorporate regular rest periods into their day. However, there are things you can do, as we explain later on.

**Work Matters, page 32**

Disease management

As has already been said, some people whose disease is in remission find that their fatigue is also greatly reduced. But this isn’t the case for others. However, keeping your disease under control remains the central aim of the treatment of RA, whatever its direct impact on your fatigue. It limits damage to your joints and helps to minimise symptoms such as pain, inflammation, swelling and stiffness, which may, in turn, help reduce the severity of your fatigue.

**Diet**

There is no evidence that diet can directly affect fatigue. However, eating a balanced diet, with regular meals, is generally beneficial. As with getting a good night’s sleep, eating healthily will make you feel better, which will make it easier to cope with your fatigue.

Both over-eating and under-eating can have an indirect impact on your fatigue. Eating too much and becoming overweight puts more pressure on your joints and can make carrying out everyday tasks more tiring. Eating too little (by skipping meals, for example) and becoming underweight will likely reduce your energy. Being hungry is tiring. Having low blood sugar from not eating enough can make you feel tired and muddle-headed.

Keeping up your fluid intake is also important. Dehydration is tiring. Make sure you drink enough – regular cups of tea and glasses of water, for example – especially during the day.

People with RA are also prone to anaemia (iron deficiency), which, among other things, causes tiredness and lack of energy. Anaemia is treated with iron supplements and by increasing iron-rich foods in the diet – such foods include dark-green leafy vegetables, pulses and beans, nuts and seeds, eggs, meat, fish and tofu.
What can you do to tackle fatigue?

Talk about it.

Fatigue is a hidden symptom of RA, little understood and too infrequently talked about. The first step in tackling fatigue, therefore, is to talk about it – to your healthcare team and to those close to you. That way you can explain how fatigue affects you and begin to get the help, support and treatment you need.

Of course, it’s not always easy to talk about how you feel. It can be embarrassing, or it may be difficult to find the words to explain your fatigue – particularly if talking about your feelings is something you don’t do naturally and have never done. You may not want to talk to someone close to you about how you feel about living with RA because you don’t want to upset them. It can sometimes seem easier to say nothing. This is where the NRAS peer-support service can help by putting you in touch with a trained volunteer with RA who understands fatigue and its impact. Sometimes talking to a stranger is easier than talking to a close family member or friend. Call our freephone helpline on 0800 298 7650 to access the peer-support service.

You may also find viewing the NRAS Behind the Smile videos helpful, the videos show how difficult it can be to tell people how you really feel. These powerfully emotive and informative videos encourage you to admit that you’re struggling and show the benefits of opening up. You can view them on our website at www.nras.org.uk/behind-the-smile
If you can find a way to open up about the things that you find difficult, you will almost certainly find it helps to make things less difficult. And in any case, those close to you will be able to see when things are hard for you. In the case of fatigue, they will see you struggling to get things done but may not understand why; and then you’re more likely to upset them by not talking.

**Talk to your healthcare team** about your fatigue so that they are aware of how it affects you. Try not to play down how you feel, even if you’re having a good day. Tell them how you feel on the bad days. Don’t be afraid to ask questions or raise any concerns you have. They are there to help.

If you are finding it difficult to talk to your healthcare team, think about taking someone with you to your next appointment for moral support. And write down in advance any questions you particularly want to ask and take the list with you.

You may find it useful to keep track of your fatigue, by means of a diary or an app on your mobile phone or tablet. You can then show the results to your consultant, GP or specialist nurse at your next appointment to give them a clear picture of how fatigue affects you over time. NRAS offers several ways to monitor your fatigue and there is more on keeping track, including a sample fatigue diary, later on in this booklet.

**Keeping track, page 24**

**Talk to your family, partners, friends and employer(s),** to explain how fatigue affects you and what you need from them when it strikes. Some people find it helpful to have a way of describing their fatigue that they can use whenever telling others about it. For example, some say it’s like having the plug pulled or the battery on their phone running down and having nowhere to charge it up again. Others talk of their energy levels as a fixed daily quantity – say, six chocolate bars a day; if they use up four bars on one task, then they only have two left to accomplish everything else they have to do that day. And when the last bar is gone, that’s it until tomorrow.

**Family life & relationships, page 29**

Another way of explaining how fatigue affects you is the ‘spoon theory’. A spoon represents a person’s ability to
tackle a task. People who don’t have fatigue have an unlimited number of spoons. They don’t even have to think about using them. They get up in the morning, get ready for work, say, and leave the house without undue effort. But if you have fatigue, you only have a limited supply of spoons. Getting up in the morning can itself be an effort that requires a spoon. And by the time you have got ready, had breakfast, taken your medication and left the house, you have few spoons left – you know you are going to have to plan the rest of your day very carefully if you are not to run out of spoons altogether. You could ‘borrow’ against tomorrow’s spoons, but that means you will be able to accomplish less the next day as you’re starting with a ‘spoon’ deficit. Imagine that each imaginary spoon is a spoonful of energy.

As well as describing your fatigue to those close to you, be prepared to ask them for help. Explain that there’s only so much you can do and ask them to take on some of your jobs, whether at work or at home. Don’t be afraid to delegate. Equally, don’t take on any additional tasks that you know you won’t be able to accomplish that day. Explain that you already have enough to do and that someone else will have to do it or that it will have to wait until another day. There’s more on talking to those close to you later in this booklet.

If you are in paid employment, it may not be easy to work your day around your fatigue. But it’s not impossible, and you may also be able to arrange a more flexible way of working. Later on, we outline the possibilities.

Talk to other people with RA, to find out how they manage their fatigue. As we said earlier, the NRAS Helpline (0800 298 7650) can arrange for a trained Volunteer with RA to call you to chat about how they cope and what works for them.

The Helpline team can also talk through possible ways to manage fatigue and how to explain it to others. Fatigue is a common Helpline enquiry and staff are used to talking about it in an impartial and confidential way.

NRAS has many groups around the UK. Going along to a group is a great way of getting out and meeting other people with RA who will share with you what they’ve found to be helpful in managing their fatigue. You can find out if there is an NRAS group near where you live by calling 0845 4583969 or email groups@nras.org.uk or visit www.nras.org.uk/groups

The NRAS online forum, HealthUnlocked, also offers the opportunity to talk to other people with RA. You can visit the forum at www.healthunlocked.com/nras

NRAS has developed a range of free (to users) self-management workshops designed to bring people with RA together to share their experiences and to help them better manage their RA. The workshops are only available if the commissioning groups, health boards or hospital trusts in your area commission them (ie provide the funding to put them on). To date they have been funded in only a very few areas as the reality is that the NHS is so financially strapped that we are really struggling to get these programmes funded in more areas. We know that courses can
be life changing and our experience is that patients and clinicians are interested in them and those who have run them or attended them understand their value to people with RA and the long-term benefits to the health system. For more information, please email enquiries@nras.org.uk or call 0845 458 3969.

Talk to a counsellor, life coach, therapist or psychologist. As we have already said, almost everyone with RA-related fatigue suffers from low mood or depression at some point. At the same time, depression can make your fatigue worse. Being depressed is nothing to be ashamed of. Recognising that you may be depressed is the first step to getting help and support, which in turn may help reduce your fatigue.

The evidence shows that ‘talking therapies’, such as cognitive behavioural therapy (CBT), delivered by a trained counsellor or psychologist, are very effective in combating depression and reducing fatigue. If you experience depression, ask your GP, specialist nurse or consultant about the available therapies in your locality and ask for a referral.

Pace yourself

Fatigue limits your available energy. Understanding how your fatigue affects you will help you plan your day and pace yourself, so that you can make the most of your energy supply.

Tracking your fatigue with a diary or with an app on your mobile phone or tablet is a good way to start recognising the patterns: to see when fatigue is likely to affect you and how long it is likely to last; to recognise how a busy day or a bad night’s sleep can make things worse; to get a feeling for when are the best times to rest and the best times to be active.

You can use an ordinary paper diary and fill it in as often as you can and in as much detail as you like. We have designed a simplified diary, using letters to mark busy times, rest periods and episodes of overwhelming fatigue. There is a sample of the diary, with suggestions as to how best to use it, later in this booklet. There is a digital version on our website that you can fill in online and then print off. Or you can print blank versions and fill them in by hand. And we have paper copies available on request – to get a copy call us on 0845 458 3969.

There are also two free apps that can help you keep track of your RA and its symptoms, including fatigue. They work on iPhones or iPads or on any mobile phone or tablet that runs the Android operating system. You can download them by visiting our website (search for ‘app’).
Know your DAS is a free app that is aimed at tracking your DAS28 so that you can see how well your RA is being controlled. DAS28 is a measure of disease activity. The lower the score, the better your RA is being controlled. A DAS28 score of 2.5 or lower means that your disease is in remission, which is the main goal of RA treatment. As well as tracking your DAS28, this app has a simple diary feature that lets you keep an eye on all the main symptoms of RA, including fatigue. The DAS app can also help you remember when to take your medications as prescribed.

RheumaBuddy is a free app for people with RA and JIA. It helps people get a better understanding of what influences the good or the bad days and with this understanding, increase the number of good days.

The app is suitable for people over the age of 16 years to use independently. The pain body-map allows you to pin point and note specific areas of pain. You can also log sleep, exercise and hours of work or school. You can also keep a daily record of the severity of your symptoms, including fatigue, and it is possible to chat with other patients and learn from their experiences.

The app helps you to gain greater control of your RA or JIA. By getting a better overview and discovering patterns you can figure out what you can do to influence the disease in a more positive direction. The app enables you to share the findings with your rheumatologist.

RheumaBuddy is a perfect tool to document the development in your disease since your last visit.

As well as helping you to see when you should plan to rest and when you can expect to have more energy, diaries and apps are useful tools for educating others – friends, family, healthcare professionals, employers and work colleagues – about the ways in which fatigue affects you.

Keeping a diary that describes how all your symptoms (including fatigue) affect you is also very useful if you need to apply for either of the two disability-related benefits: Personal Independence Payment (PIP) and Employment and Support Allowance (ESA). This is because you will have to undergo an assessment of how your disability affects your ability to carry out a variety of tasks. A diary will help give a good picture over a period of weeks or months. There is more on this later on.

Benefits & fatigue, page 37
Whether you use a diary or not, it is helpful to plan ahead and pace yourself.

- If you have a busy day tomorrow, try to have a less busy day today, with regular breaks, and plan to have plenty of rest the day after tomorrow. At the same time, make sure that you plan to take some rest breaks during your busy day.

- You may find that you have more energy at certain times of the day than at others, so try to structure your day accordingly.

- Re-evaluate your ‘to do’ list. Do you need to do everything you had planned for the days ahead? Are there things that someone else could do? More than likely, there are things you can put off or strike off the list altogether.

Some people make use of a ‘traffic-light’ system when planning ahead. They assign tasks in their diary or on their calendar a colour – red, amber or green, according to how tiring they are; red activities being those that take the most out of you, and green the least. Then they aim to ensure they have no more than one red-light activity on any given day and, say, no more than two amber-light activities.

People who use this system say that once they get used to thinking about activities in this way, it really helps them to plan ahead and conserve their energy. That way, they find they get more done.

- Break down the tasks you feel you can’t avoid into manageable chunks, taking time out to rest in between. Try to spread big tasks out over the day or even over several days. Little
and often may be the best way to approach tasks rather than trying to do everything all in one go

- See if there are ways in which you could do things more efficiently or in a less strenuous way. If you’re preparing a meal, for example, try to keep everything you need together, close by, so that you don’t have to keep going off to fetch things.

There are a number of gadgets that can help around the home – such as devices to help with opening jars with minimal effort. You may also want to consider getting a perching stool that you can rest on while cooking or using the bathroom sink. You’ll still be at more or less the same height as when you’re standing up but you will be supported, which will help you conserve your energy.

You may not need to use such aids every day, but they can be helpful to have on hand for less good days.

The Disabled Living Foundation is a charity that provides impartial advice, information and training on equipment for independent living. Visit their website at www.dlf.org.uk or call their Helpline on 0300 999 0004

- Be kind to yourself. Accept that there may be times when you will be able to do a lot less. Even when your energy levels are high, be careful not to overdo it.

Of course, this is all very well if you have complete control over your time. However, if you are in paid employment or you have regular commitments – say, due to the demands of a young family – it may not be so easy. Later on, we look at what you can do at work to better accommodate your fatigue – flexible working hours may be possible, for example.
If you are not in paid work but have regular demands on your time – child care, for example, or other caring responsibilities – you may need to ask others to help so that you can build some rest periods into your day. Can someone else collect the kids from school today or take the toddler to the nursery in the morning? Can someone do the shopping for you or do a spot of ironing?

**Exercise**

Taking even a moderate amount of exercise helps to strengthen muscles and build fitness, which will help to make it easier and less tiring to carry out everyday tasks. Exercise can also help you get a good night’s sleep.

Exercise doesn’t have to involve going to the gym and doing a strenuous workout. It can be as simple as going out for a brisk walk or weeding a flowerbed.

You could take a cycle ride, go for a swim or take part in an exercise class.

Joining an exercise class or taking part in any other kind of organised activity – a walking group, a dance class or a cycle club for example – is also a good way of meeting people and getting out and about.

Take it very gently to begin with. Even if you were used to exercising hard before being diagnosed with RA, it’s probably best not to go for the burn every day. Build up gradually. In time, you will find yourself able to do more and more. There will still be days when you just won’t be able to do much more than rest. That’s fine. But when the energy returns, beware of trying to make up for lost time; don’t overdo it.

As you exercise more regularly, you will become aware of your limitations as well as your capabilities.
If you do want to join a gym or an exercise class, talk to your healthcare team first. You should be able to talk to a physiotherapist who can give you advice on the kind of exercise that would be suitable for you. They may also be able to point you towards suitable classes where you live.

Some leisure centres run classes tailored for people with long-term conditions and/or reduced mobility. It may be worth contacting the centres near to you to see what they have to offer. You could also do an internet search to see what, if any, classes or activities are run locally. Or visit your local library for information.

Some local NRAS groups also run exercise groups or may know of local suitable classes. Contact your local NRAS group. They may have details of exercise classes nearby. They may also be able to put you in touch with other people with RA who live locally and who already participate in regular exercise classes or activities. To find out if there is a patient group near you visit www.nras.org.uk/groups, phone 0845 458 3969 or email groups@nras.org.uk

You can also find some useful exercise videos on our website at www.nras.org.uk/exercise-videos

Sleep and diet

Eating well, making sure you stay hydrated (by drinking plenty of fluids during the day) and sleeping well will help you to manage your fatigue better. If you are well-rested and well-nourished, you will feel better able to cope with daily tasks.
## Fatigue diary

This is a sample diary. Paper copies are available to download from the NRAS website and there is also an interactive version on our website that you can fill in and print out. Visit [www.nras.org.uk/diary](http://www.nras.org.uk/diary) See also pages 18 to 20 of this booklet.

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<td>11 November 2018 Sunday</td>
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### KEY

- **B** Busy: Busy time; doing a lot, few breaks
- **M** Moderately active: You’re active but not madly busy
- **R** Rest: Sitting or lying quietly, reading, watching TV, but not sleeping
- **S** Asleep: Sleeping
- **C** Crash: When fatigue stops you doing anything
# Fatigue Diary

**Start of Week: Monday, 5 November 2018**

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**MMB B R R C C C C S**

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**NRAS HELPLINE**

0800 298 7650

**YOUR NHS NUMBER**

12345CDR6789

**YOUR HOSPITAL NUMBER**

00F4G77J99910

**HOSPITAL PHONE NUMBER**

020 8800 1199

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Your name
Working with your healthcare team

I feel as if even the medical profession think I am making up my fatigue. So this just makes me feel worse about it!

Invisible disease: Rheumatoid arthritis and chronic fatigue (NRAS survey 2014)

Many people with RA feel that healthcare professionals don’t take fatigue seriously enough. In fact some people have told us that they’ve been told by their health professional that fatigue is nothing to do with their RA! Wrong. Part of the problem is that there are no approved treatments for fatigue in RA so doctors and rheumatology consultants may feel they have little or nothing to offer. Instead, their focus may be on controlling your disease. For many people, being in remission will minimise all the symptoms of RA, including fatigue. However, for many others, this is not the case.

Pain and fatigue can persist even when you are in remission. If that is the case for you, let your healthcare team know. Being in remission is not just about getting your DAS28 score as low as possible; it is also about how you feel. So, if you’re not feeling well, despite all your tests showing your disease is under control, don’t pretend otherwise.

When talking to your healthcare team, be honest, not brave. You may be so used to telling friends and family you’re OK, answering every ‘How are you?’ with ‘I’m fine’, that it’s hard to be honest with healthcare professionals about your pain or other concerns. But your team needs to know the real picture so that they can give you the help you need. However good your health team are they are not mind readers so be open with them.

You might find it helpful to take a look at our emotive and informative Behind the Smile videos on our website at www.nras.org.uk/behind-the-smile

If you still feel that your doctors or nurses aren’t taking your fatigue
seriously, consider keeping a fatigue diary (see pages 18-20 and 24). Take it to your next appointment. You may also want to take someone with you who can also testify to the ways in which fatigue affects you.

Don’t be afraid to ask for help if fatigue is making life difficult

Ask for a review of all your medications to ensure that none of your medications are worsening your fatigue

Emotional and psychological support can make a difference to how you experience fatigue. Your doctor should be able to make a suitable referral

Members of your healthcare team – the specialist nurse, physiotherapist and occupational therapist – may be able to help you better manage daily living.

One of the things your healthcare team can do that may have a direct impact on your fatigue is to carry out a review of your medication to identify any drugs you are taking (including any medications you are taking for conditions other than RA) that may be making you feel drowsy. If they haven’t offered you a review, don’t be afraid to ask for one or speak with your pharmacist about a medicine use review (MUR). It may be possible to substitute other medications for those that are causing you to feel tired.

Your healthcare team can also refer you to other healthcare professionals who can help you to tackle fatigue:

- **A physiotherapist** can help you devise an exercise programme that is suitable for you. They may also be able to refer you to local exercise classes or activities

- **A counsellor or psychologist** can help you to tackle low mood, depression and/or anxiety. It can be difficult to admit that you need such help and difficult to ask for it. But your GP, consultant or specialist rheumatology nurse will understand and be sympathetic; you can always take someone with you to an appointment for moral support.

At the same time, it’s better that you ask rather than wait to be asked. Research suggests that more than three-quarters of people with RA are not asked about social and emotional issues by their healthcare team.

There are a range of ‘talking therapies’ available on the NHS, including cognitive behavioural therapy (CBT) and Mindfulness. There are courses you can follow online or you can see a therapist in person, on a one-to-one basis or as part of a group. These therapies have been shown to be very effective in helping people deal with their fatigue. Another useful resource is the RefRAme RA campaign www.refRAmeRA.co.uk

There are generally quite long waiting times for these services on the NHS and it may be some time before you can get an appointment. But it is worth the wait.
• **A dietitian** can help you adopt a healthy diet and maintain a healthy weight

• **An occupational therapist**, also known as an **OT**, can help you improve or maintain your ability to do everyday activities. He or she works with you to see what you’re finding difficult and to find out ways to reduce the problems. Among other things, this might be by:

  - **Changing how you do things** – helping you to use your time and energy effectively or finding ways to save energy. You might break activities down into smaller parts or do less at any one time

  - **Helping you to reduce the impact of RA on your daily activities**. This could mean teaching you about looking after your joints during everyday activities or about ways to manage pain and fatigue

  - **Offering you support in making changes**, for example with individual counselling or group programmes

  - **Adapting your home or workplace to make activities easier** – the right piece of equipment or structural alteration can make all the difference. Occupational Therapists can also liaise with employers to ensure that appropriate adaptations are provided through statutory services, where possible. There’s more on workplace adjustments later on.

The National Institute for Health and Care Excellence (NICE) offers guidance to NHS healthcare professionals on the best practice for treating a wide range of conditions, including **rheumatoid arthritis**. NICE guidance applies to England and Wales and is generally followed in Northern Ireland. In Scotland, the equivalent body is the Scottish Medicines Consortium (SMC).

NRAS contributes at a national level to the NICE (England and Wales) and Scottish Medicines Consortium Health Technology Appraisals for the introduction of new drugs, as well as the standards and guidelines produced by NICE and the Scottish Intercollegiate Guidelines Network (SIGN), and whilst they refer to symptom management they do not go into detail about managing fatigue, which is one of the reasons we have produced this booklet.

The NICE website is at [www.nice.org.uk](http://www.nice.org.uk)
The SMC website is at [www.scottishmedicines.org.uk](http://www.scottishmedicines.org.uk)
NICE does offer advice to healthcare professionals on managing fatigue, but it is not specifically tailored to people with RA – you can find it here: [https://cks.nice.org.uk/tirednessfatigue-in-adults#!scenario](https://cks.nice.org.uk/tirednessfatigue-in-adults#!scenario)
Family life and relationships

One of the aspects of fatigue that many people with RA find especially difficult to deal with is its impact on their relationships with family and friends.

“I have two wonderful grandchildren who I cannot play with when I’m fatigued. It’s horrible.”

“It’s difficult to explain to my toddler that I’d rather sit on the sofa than go to the park with her.”

In our survey, *Invisible Disease*, almost half (48 per cent) of those who responded said they felt guilty about the impact of their fatigue on other people. Nearly three-quarters (70 per cent) of respondents reported that fatigue had prevented them from taking part in activities with their family or partner; and over a half (54 per cent) reported that fatigue negatively affected their sex life.

In our survey in 2013 on emotions, relationships and sexuality, 63% said they had been less able physically to have sex in the way they would like to.

In our ‘emotions’ survey referred to above, a third (33 per cent) of respondents rarely or never talked to those close to them about how they feel.

- Fatigue can affect your closest relationships and negatively affect your sex life.
- Fatigue may disrupt or limit your involvement in family and social activities.
- Finding ways to talk to those close to you about how fatigue affects you and the support you need can make a big difference.
- Psychological support and counselling can be a great help in tackling these difficulties.

It is never easy to talk about difficult feelings such as guilt or about intimate matters such as sex. You may find yourself avoiding any discussions that
you find awkward. And that’s entirely understandable. But your husband, wife, partner or best friend may prefer you to open up. They may wonder why you don’t want to go out, for example, and they may think it’s something to do with them rather than being the result of fatigue. In return, you may find yourself being angry and irritable or withdrawn and silent, which may make things worse. But if you can find a way to start talking about what you are finding difficult, it can make a big difference to your close relationships.

It can be difficult explaining to young children how you feel and you may find yourself making up stories that you think they will find easier to understand. But children often notice more than you think they do and they may not entirely believe them if you say you can’t take them out to play because you’ve got a headache. They may instead think that things are worse than they are and become worried and upset.

There is no one simple way to limit the negative impact of your fatigue on your close relationships. One way to start is by talking to other people with RA to find out how they manage. Our Helpline (0800 298 7650) may be able to arrange for a trained Volunteer with RA who has faced similar difficulties to call you for a chat. You can also talk things through with Helpline staff. Or you may want to join one of our NRAS groups or chat to people through our online forum, HealthUnlocked – there are more details on page 17. It’s sometimes easier to talk to people you are not emotionally involved with.

Alternatively, you may prefer to talk to a healthcare professional, such as your GP or specialist rheumatology nurse.
They can also refer you to professional sources of support, such as counselling.

It is also a good idea to make sure that friends and family are aware that fatigue may at times make it difficult for you to meet social commitments. If you find yourself cancelling arrangements at the last minute, your friends and/or family may be less likely to include you in future activities. Explain to those close to you that you often have to say ‘no’ because you are often laid low by fatigue, which can come without warning. However, having to say ‘no’ frequently does not mean that you never want to be asked to join in. They will be sure to understand. There may be more days than you would like when you have to stay at home, but there will still be plenty of occasions when you are up to going out. Just because you can’t go out sometimes, it doesn’t mean you don’t ever want to go out.

If you keep any kind of fatigue diary, you may want to show those close to you. It may help you to explain to them how fatigue affects you.

If you are concerned about the impact of fatigue or any other aspect of living with RA on your sexual relationships, you (and your partner, if that’s what you both want) could talk to a sex and relationship counsellor. Your GP or nurse may be able to make a referral. Or you could contact the national charity, Relate, who offer help with all aspects of relationships. Visit their website at www.relate.org.uk or call 0300 100 1234

NRAS also publishes a very useful booklet, *Emotions, relationships and sexuality*, which draws on the results of our 2013 sex and relationships survey and is full of helpful advice and information. You may also want to look at our *Family Matters* survey (2012), which looks at how the families of people living with RA are affected. Both publications can be downloaded from our website or you can phone NRAS on 0845 458 3969 and ask to be sent a copy.
Work matters...

Work is important not just for financial reasons but also because it is generally beneficial to health and wellbeing.

On days when I work, I can do nothing else; I am completely drained. No one knows this but me. Even thinking about telling anyone makes me cry.

_Invisible disease: Rheumatoid arthritis and chronic fatigue (NRAS survey 2014)_

Work is as important to people with RA as it is to anyone else: three-quarters of people with RA are diagnosed when of working age. But RA and its symptoms, especially fatigue, can complicate matters.

Fatigue may disrupt your working life. It may make it hard to get a job in the first place and to retain a job when you have one. It can hold you back from promotion. You may find yourself worrying that absences or poor performance due to fatigue could cost you your job.

In our _Invisible Disease_ survey of people with RA, almost three-quarters (71 per cent) of those of working age who were unemployed said that fatigue was the reason they did not work. And almost a quarter (24 per cent) of all those surveyed said that they had had to change jobs because of fatigue.

The same survey also reported that almost a quarter (23 per cent) of respondents in employment had taken more than 10 days off in the preceding 12 months due to fatigue; for those diagnosed for less than a year, the proportion rose to more than a third (34 per cent). But almost a half (49 per cent) of respondents said their line manager had not discussed or offered any changes to help them better manage their chronic fatigue in the workplace.
Fatigue may disrupt your working life

Take all the breaks you are entitled to at work – get away from the workplace or at least work-station, for lunch if you can. And make sure you get plenty of rest and support outside working hours

Talk to your employer and/or work colleagues about the support you need

Consider taking this booklet to work, to share with your manager and work-mates

You have a right to ask for and your request to be considered for flexible working hours, whether you are disabled or not

The law offers protection from discrimination at work because of your disability

Unions, professional bodies and other organisations may be able to support you in any discussions with or challenges to your employer. You are not on your own.

But while fatigue can complicate matters, there is much that can be done to improve your working life. To begin with, here are some tips for helping you cope better with the working week:

Pace your activities for work as you would at home. If some elements of your work are more physically demanding, see if you can spread this out across the day/week, rather than doing them all in one session

Re-examine the way that you work on a regular basis to see if there’s an easier way for you to do the job. For example, are you sometimes performing tasks while standing when you could be sitting?

If you drive to work, find out whether there’s a car-share scheme in your area. That way, you can try and rest while someone else drives

Fresh air helps. Take a 10-minute walk during your lunch break to get the circulation going and to get away from the working environment. It will also give your brain a break

Take your lunch break away from your work station if possible – a change is as good as a rest

Talk about the impact of fatigue with family and friends; how it affects you after a day’s work and the support you might need

Keep meals during the week simple and nutritious. Get your partner and family to help you

When you cook, perhaps at the weekend, make extra and freeze it for use during the working week

If you live with someone else, ask them to cook a couple of nights a week

Invest in a slow cooker to cook things overnight or during the day so that you come home to it all ready for you

Keep any housework during the week to a minimum

Get help with shopping or, better still, do it online and get it delivered
• Get to bed early on the nights (or days, if you do night shifts) before you have to go to work

• Help the family understand that you may need a lie-in at the weekend.

But it’s not all up to you. Your colleagues should also play their part. Start by talking to them about your RA. It may be that the reason why you have not been offered any support at work is that your employers are not aware that you need any support. It is a good idea to start that conversation, perhaps with your line manager or with the HR department – or, if you work in a small business, with your employer. You should explain how fatigue affects you. You may want to show them your fatigue diary and/or give them a copy of this booklet.

You also have a legal right to request **flexible working**. That means that your start and finish times can be flexible and/or that you can work at home on some days. Your employer must respond to your request in a ‘reasonable manner’ – this would include weighing up the advantages and disadvantages of your application; meeting with you in person to discuss your requirements; and offering you a right to appeal against their decision. If your request is not handled in a reasonable manner, you can take your employer to an industrial tribunal. Your healthcare team may be able to refer you to an OT or you can seek a consultation with a private OT (who will charge you for their time).

The rules on flexible working are the same in England, Scotland and Wales. They are slightly different in Northern Ireland. For more information, visit [www.gov.uk](http://www.gov.uk), [www.acas.org.uk](http://www.acas.org.uk) or, for people in Northern Ireland, [www.nidirect.gov.uk](http://www.nidirect.gov.uk)

Apart from offering flexible working, employers can help you by **making adjustments** in other ways. Such adjustments might include obtaining specialist equipment to help you do your job with less effort; changing the workplace layout to minimise the distances you have to move about during the working day; offering a car parking space and time off for medical assessments and treatment. By law, employers are required to make ‘reasonable adjustments’ for employees with a chronic health condition or disability.

**An occupational therapist (OT)** can assess your workplace and the way in which you carry out your job in order to make recommendations about the adjustments that you require. An OT can also talk to your employer on your behalf about the adjustments, including flexible working, that would help you. Your healthcare team may be able to refer you to an OT or you can seek a consultation with a private OT (who will charge you for their time).

There is also a government-funded organisation called **Fit for Work** that offers support and advice from a team of OTs to employees with a chronic health condition or disability. Employees can access the service themselves. Employers and GPs can also refer employees or patients to the scheme to help you return to work if you’ve been on sick leave for four weeks or more.
If you live in England or Wales, visit www.fitforwork.org or call the advice line on 0800 032 6235

In Scotland, the scheme is no longer taking referrals, but you can get advice from their website at fitforworkscotland.scot or their free helpline on 0800 019 2211 (Monday to Friday, 9am to 5pm). Working Health Services Scotland is an NHS service that aims to help employees of small-to-medium enterprises remain in or return to work quickly after a period of absence.

For more information, visit the Healthy Working Lives website at www.healthyworkinglives.com or call 0800 019 2211 and select option 3.

The Fit for Work scheme does not apply in Northern Ireland. To see what support is available there, visit www.nidirect.gov.uk or talk to your GP, employer or trade union.

The Access to Work scheme can advise disabled employees about how to get the support they need at work. They may also be able to offer grants to cover some of the cost of making any reasonable adjustments you require. You can access the scheme online at www.gov.uk/access-to-work or by talking to a Disability Employment Adviser (DEA) at your local JobCentre or Jobs and Benefits Office. A DEA can also advise you about returning to work if you have recently become disabled or if you have lost your job because of your disability. There’s also information at www.gov.uk and www.nidirect.gov.uk
In Northern Ireland, The Disability Employment Service (DES) provides support to employers to help them recruit people with health issues or disabilities. It also helps employers to retain employees who become sick or disabled in their jobs. Visit www.nibusinessinfo.co.uk or www.nidirect.gov.uk

The law protects people with a disability or long-term health condition from discrimination at work. This also applies to people applying for work. For more information if you live in England, Scotland or Wales, contact The Equality and Human Rights Commission (EHRC) by visiting www.equalityhumanrights.com or calling 0808 800 0082. In Northern Ireland, contact The Equality Commission for Northern Ireland at www.equalityni.org or on 028 90 500 600

If you are a member of a trade union or professional body, contact them for advice about your rights at work. They can also support you in any dispute with your employer, including providing a representative to accompany you to any meetings about the dispute. If you are not a member of a union – and not all workplaces recognise unions – contact the Trades Union Congress (TUC) at www.tuc.org.uk or on 020 7636 4030, or if you live in Northern Ireland you should contact the Irish Congress of Trade Unions Northern Ireland Committee at www.ictuni.org or on 028 90 247 940

NRAS cares very much about the impact that RA can have on work (or that JIA has on education) and we have done a lot of work in this area. We carried out surveys on work in 2007, 2010 and 2017 and we are working on provision of new resources for working people with RA/JIA.

NRAS is an active member of the Fit for Work Coalition, which brings together healthcare professionals, policy makers, employers, and patients to improve the productivity of Britain’s workforce, by empowering and supporting people, including those with long-term conditions, to remain in or return to work and be economically active. NRAS is also a member of the Expert Advisory Group to the Work and Health Unit, which spans the Department of Health and the Department of Work and Pensions.

NRAS campaigns for and on behalf of better support in the workplace for people with long-term fluctuating conditions. This includes raising awareness of the invisible symptoms of RA and JIA such as pain and fatigue with the government agencies who carry out assessments for benefits such as PIP and ESA.
Does fatigue affect my eligibility for benefits?

There are a number of benefits you may be able to claim; but there are two where your fatigue may play a key role in determining your eligibility.

**Personal Independence Payment (PIP)** (the replacement for Disability Living Allowance, or DLA) and **Employment and Support Allowance (ESA)** are benefits for people with a disability or chronic health condition, such as RA. PIP can be claimed by people in or out of work; ESA is for people with limited capability for work due to disability.

PIP and ESA are for people of working age, after retirement, anyone who needs additional help because of illness or disability can claim Attendance Allowance.

To be eligible for either PiP or ESA, it is not enough that you have a diagnosis of RA. You have to show that your RA limits your ability to work and/or carry out a range of everyday tasks and activities.

- If you are applying for Personal Independence Payment (PIP) or Employment and Support Allowance (ESA) you will have to undergo a medical assessment.
- These assessments are not good at taking into account hidden and fluctuating symptoms such as fatigue. Be sure to explain clearly how fatigue affects you. A diary (see pages 18-20 and 24-25) that tracks your fatigue, as well as other symptoms, is useful evidence to submit with your claim.
- The healthcare professional in charge of your care will be asked about how RA affects you. Make sure they are up-to-date with how all your symptoms, including fatigue, are affecting your ability to carry out daily tasks.

When you first apply for either benefit, you will be asked to **complete a detailed questionnaire about your health**, which focuses on your ability to carry out a number of everyday tasks. It is important that you include fatigue among your symptoms and show
clearly how it limits your ability to manage day-to-day activities. If fatigue affects your thinking – the 'brain fog' that people speak of – be sure to mention that, too.

The questionnaire does not include a specific question about fatigue. However, there are opportunities on these questionnaires to give examples of how fatigue affects you. The assessment form for PIP, for example, asks about your ability to mix with other people (question 11). There, you can explain how fatigue can make mixing (ie meeting people socially at home or elsewhere) difficult or, on occasion, impossible. Question 14 asks about your ability to move around without severe discomfort. Again, this is a place to mention fatigue. You can explain that it is not only pain and stiffness that can make moving about extremely difficult; fatigue can also have that effect.

As well as requiring you to submit a detailed assessment form, the assessors for PIP and ESA will ask you for the name of the person in charge of your treatment (your GP or consultant rheumatologist, for example) so that they can be contacted for information about your health. It is a good idea to talk to that person before submitting your claim form to make sure that they are fully up-to-date with all your symptoms, including fatigue (it may have been a while since your last appointment, for instance). They may also be able to give you some advice about what to say on the form.

If you keep any kind of fatigue diary, it is a good idea to include that as part of the evidence to accompany your benefit claim. You may also have a diary that records all your symptoms (not just fatigue) – include that as well.

Once you have submitted your claim, you will be asked to attend a medical assessment. This usually takes place at an assessment centre. However, if you are unable to travel, it may be possible to arrange for the assessment to take place in your home. They will need to see a letter from your doctor that explains that you cannot travel before they will agree to a home assessment.

The in-person medical assessment tends to focus on your physical capabilities on the day. Such assessments are not good at assessing the impact of hidden and fluctuating symptoms such as fatigue. For example, it may be that you are having a good day when the assessment takes place but on the following day your fatigue is so severe that you cannot function at all. That’s why a diary and as much other evidence about how you are affected over a period of time (such as an app on your mobile phone or tablet) is very helpful and should be taken along to the assessment.

If you are turned down for either benefit or if you think the level of benefit awarded is too low or, in the case of ESA, if you are judged to have some capability for work when you don’t think you have any, you can ask for a 'mandatory reconsideration'
of the decision. You can ask to be sent copies of all the evidence used in reaching their decision. In return, you should state in clear terms why you are asking them to reconsider: for example, you could say that you believe they have underestimated the extent of your disability and its effect on your ability to carry out everyday tasks. This would be another opportunity to emphasise the impact that fatigue has on you. If the reconsideration does not lead to a change in the original decision, you can then lodge an appeal. It is important to note that many people do win on appeal, so it well worth pursuing.

For detailed information on PIP and how to apply for it, see our booklet *How to claim Personal Independence Payment (PIP).* For further information on ESA and a range of other benefits, see our publication *Benefits and rheumatoid arthritis.* There is also information on our website and our Helpline can help answer any questions you might have.

The website [www.gov.uk](http://www.gov.uk) has up-to-date information about all benefits and how to apply for them. Your local *Citizens Advice* office can also offer advice and support – visit [www.citizensadvice.org.uk](http://www.citizensadvice.org.uk) for contact details. If you live in Northern Ireland, visit [www.nidirect.gov.uk](http://www.nidirect.gov.uk) for information about benefits.
Glossary

Anaemia

is a condition where the number of red blood cells in your blood is at a low level. The red blood cells carry oxygen around the body and if they are depleted your muscles and other body tissues won’t get enough oxygen. One of the main symptoms of anaemia is tiredness. The main causes include iron deficiency, vitamin B12 deficiency and several diseases, including leukaemia and sickle-cell disease. Anaemia can be diagnosed by means of a blood test. If it is caused by iron or vitamin B12 deficiency, it can be treated with dietary supplements and by increasing the foods in your diet that are rich in iron and vitamin B12.

Autoimmune

describes a condition where the body’s immune system attacks part of the body itself. The immune system is our defence against infection. It is an incredibly complex system that, for most of us, for most of the time, works well. But sometimes, something goes wrong and the immune system mistakes some parts of the body for a damaging invader (a virus, for example) and attacks it. In such cases, a person develops an autoimmune condition, the ‘auto’ part of the word here meaning ‘self’. RA is an autoimmune disease, in which the immune system attacks the lining of the joints. Other autoimmune diseases include multiple sclerosis, in which the protective coating of nerves (myelin) in the brain and spinal cord is progressively attacked and weakened by the immune system, and type 1 diabetes, where the pancreas is attacked. One of the main damaging mechanisms of autoimmune diseases is inflammation.

DAS28

is a measure of disease activity (see below) in RA. It combines the results of four measures: the number of joints (out of 28 examined) that are swollen; the number of joints (out of the same 28) that are tender; the levels of certain markers of inflammation in your blood; and your own assessment of your overall health. A score greater than 5.1 indicates that your disease is active; less than 3.2 indicates low disease activity; and lower than 2.6 means your disease is in remission.
Depression is a state of low mood. It may be caused by a difficult life event, such as bereavement or illness. It may also be a result of a long series of events, including a difficult childhood. People with a family history of depression have an increased likelihood of developing it themselves. Some medications can cause depression. Depression can be relatively mild and of short duration; it can be severe and long-lasting. It can be associated with feelings of guilt, worthlessness, anxiety (as if something dreadful is about to happen) and a lack of interest in any activities. People with depression tend to become withdrawn and inactive. They may have suicidal thoughts. They may also be irritable, sometimes angry, and can experience insomnia and a lack of sex drive. Too many people seek help for depression very late, partly because of not recognising the early symptoms of depression. Yet depression is not something to be ashamed of; it is a treatable condition that many people experience at some time in their lives. Treatments include ‘talking therapies’ such as cognitive behavioural therapy (CBT) or psychiatric support. Antidepressant medication may also be offered.

Disease activity refers to the severity of your disease. The more active it is, the more damage it is likely to cause to your joints. One of the principal aims of treatment of RA is to keep disease activity as low as possible. Disease activity is measured by the DAS28 tests.

Flare (or flare-up) is when RA disease activity is very high and symptoms are severe. During a flare, you will tend to experience increased joint pain, swelling, fatigue and stiffness. You may also feel increasingly low in mood. Fatigue may be an ‘early warning system’ for flares, a sign that your disease is becoming more active. Many flares can be self-managed (see our website or refer to the Living Better with RA booklet for help or contact our Helpline), but if they are occurring frequently, you should contact your healthcare team as soon as possible.

Inflammation is one of the main mechanisms by which the body’s immune system deals with infection. If, for example, you get a splinter in your finger, you will soon notice redness and swelling where the skin has been pierced. That is typical of an inflammatory response. A sore throat is another example of inflammation, usually caused by the immune system responding to an infection such as the common cold. In RA, the immune system causes inflammation in the lining of the joints, which can cause damage to the joints themselves. One of the goals of treatment of RA is to minimise inflammation.
Some scientists think that some of the chemicals released into the bloodstream during inflammation may be one of the causes of fatigue, which is seen in many autoimmune diseases, including RA.

**Juvenile Idiopathic Arthritis (JIA)**

refers to inflammation in a child’s joints that isn’t caused by any other condition:

- **Juvenile**: the child was aged 16 or under when the problem started
- **Idiopathic**: the cause is unknown
- **Arthritis**: There is inflammation inside the joint, which can be seen through swelling, warmth and reduced movement.

There are different types of JIA, some of which can be short term, while others continue into adulthood.

**Remission**

derives from a word meaning slackening or weakening. In medical terms it refers to a period when a person’s disease has greatly weakened or abated. RA is said to be in remission when a person’s DAS28 score is lower than 2.6. It means that disease activity is minimal. For most people that also means that their symptoms are minimal, too. However, for some people some symptoms – pain and fatigue, for example – may persist even when they are in clinical remission. If that is the case for you, let your healthcare team know as there may be things that can be done to make you feel better.

**Rheumatoid Arthritis (RA)**

is an autoimmune condition in which your immune system attacks the lining of your joints (the synovial lining). This causes inflammation, which leads to symptoms such as pain and stiffness. RA is a symmetrical arthritis, meaning that it usually affects both sides of the body in a similar pattern, although this is not always the case. It tends to affect the small joints of the hands and feet first – often the knuckle joints in the fingers. It is described as a polyarthritis, meaning that many joints can be inflamed. RA is a systemic disease – it doesn’t just affect joints. RA can affect a person’s whole system, including organs such as the lungs, heart and eyes. Around 400,000 people in the UK are living with RA, more than two-thirds of whom are women.
Rheumatoid cachexia affects most people with RA. ‘Cachexia’ means weakness and wasting of the body due to chronic illness. In RA, it causes muscle wasting and an increase in body fat. The condition is not automatically reversed when people are in remission. Andrew Lemmey, Professor of Clinical Exercise Physiology, School of Sport, Health and Exercise Sciences at Bangor University, reported in the Winter 2015 issue of NRAS Magazine that ‘we regularly find that two-thirds of our stable RA patients are significantly muscle wasted and 80 per cent are obese’. However, he goes on to say that exercise can reverse muscle loss and weight gain. In a study he and his colleagues conducted, people experienced a doubling of their muscle strength and a significant loss of body fat following a 24-week exercise programme.
Organisations that can help

NRAS
www.nras.org.uk

NHS Choices
www.nhs.uk

Tackling fatigue

Disabled Living Foundation  (page 21)
A national charity that provides impartial advice, information and training on equipment for independent living. Visit their website at www.dlf.org.uk or call their Helpline on 0300 999 0004

Family and relationships

Relate  (page 31)
Help with all aspects of relationships.
www.relate.org.uk
0300 100 1234

Work

Access to work  (page 35)
This government-funded scheme may be able to provide a grant to provide special equipment, adaptations or support worker services to help you do things like answer the phone or go to meetings and/or to help you get to and from work.
www.gov.uk/access-to-work
www.nidirect.gov.uk

Arbitration  (page 36)
The Advisory, Conciliation and Arbitration Service (ACAS) provides information, advice, training, conciliation and other services for employers and employees to help prevent or resolve workplace problems.
www.acas.org.uk
0300 123 1100
Disability discrimination
For advice on your rights at work and on what action you can take if you feel you are being treated unfairly, contact:

The Equality and Human Rights Commission (England, Scotland or Wales)
www.equalityhumanrights.com
0808 800 0082

The Equality Commission for Northern Ireland
www.equalityni.org
028 90 500 600

Fit for Work (page 34)
Fit for Work offers free, expert and impartial advice to anyone looking for help with issues around health and work.

England and Wales
www.fitforwork.org
Advice line on 0800 032 6235 (English) or 0800 032 6233 (Cymraeg/Welsh)

Scotland
fitforworkscotland.scot
0800 019 2211 (Monday to Friday, 9am to 5pm)
www.healthyworkinglives.com
0800 019 2211 (select option 3)

Northern Ireland
Fit for Work does not apply in Northern Ireland – to see what support is available, visit NI Direct:
www.nidirect.gov.uk

Trade Unions (page 36)
Trades Union Congress (TUC)
www.tuc.org.uk
020 7636 4030

The Irish Congress of Trade Unions Northern Ireland Committee
www.ictuni.org
028 90 247 940

Benefits
For information and advice about what benefits are available to you and for help with making an application, contact:

GOV.UK (page 39)
www.gov.uk

Citizens Advice
www.citizensadvice.org.uk
Other NRAS publications

New2RA

*New2RA* is for people who are newly or relatively newly diagnosed and contains the information you need to help you make sense of and understand what RA is and what you have been told by your rheumatology team.

Living Better with RA

*Living Better with RA* is a self-help guide for people with established disease, including Juvenile Idiopathic Arthritis.

Medicines in Rheumatoid Arthritis

We believe it is essential that people living with RA understand why certain medicines are used, when they are used and how they work to manage the condition.

Invisible Disease: Rheumatoid Arthritis and Chronic Fatigue

NRAS in collaboration with 2020health, a leading health think tank, today announces the results from *Invisible Disease: RA and Chronic Fatigue 2014*, a survey of 1,954 people with Rheumatoid Arthritis (RA), to assess the impact of chronic fatigue on their capability for work, emotional and mental health as well as the overall experience of chronic fatigue caused by their RA.

For all other NRAS publications, see [www.nras.org.uk/publications](http://www.nras.org.uk/publications)

To obtain copies of any of our other booklets, or more copies of this one, please call 0845 458 3969 or email enquiries@nras.org.uk. Copies can also be downloaded from our website at [www.nras.org.uk/publications](http://www.nras.org.uk/publications)
The impact of RA on Emotions, Relationships and Sexuality

This booklet tackles emotions, relationships and sexuality, and how these very personal and intimate issues are affected by being diagnosed with and living with rheumatoid arthritis.

Work Matters

A UK wide survey of adults with Rheumatoid Arthritis and Juvenile Idiopathic Arthritis on the impact of their disease on work

I Want to Work

In this booklet you will find up-to-date and accurate advice and information, to make sure you can find the help you need to stay in your job.

When an Employee has Rheumatoid Arthritis

If you’re an employer of someone who has rheumatoid arthritis, we hope you find this booklet useful.

RA & Benefits

This guide will make it easier for you to understand how to find out more about benefits that may be relevant for you and your family.

RA & Personal Independence Payment

A self-help guide to claiming Personal Independence Payment for adults with rheumatoid arthritis.
Your phone numbers and contacts

Your hospital number
Your consultant’s name
Telephone number for consultant’s secretary
Your rheumatology specialist nurse
Pharmacist number
Your GP’s name
GP surgery telephone number
GP out-of-hours number
Medications homecare delivery number
NRAS freephone number

0800 298 7650

It might be helpful to you to record your medication details here:

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References

For a full list of all the references used in compiling this booklet (such as articles in medical journals), please call NRAS on 01628 823524.

For general enquiries, call: 0845 458 3969/01628 823524 or email enquiries@nras.org.uk

NRAS – Changing Minds, Changing Services, Changing Lives for those living with RA and JIA in the UK
Join us

You can become a Member of NRAS for as little as £20 a year.*

You will be helping our work and helping us to provide help and support to all affected by RA. In addition, you will become part of a vibrant and supportive community of people affected by RA. You will also receive a variety of Member benefits, including:

- the NRAS Members’ magazine three times a year
- monthly email updates
- an invitation to attend our FREE meetings and events

Join us today by calling
**0845 458 3969**

or by visiting our website at
**www.nras.org.uk/join**

There is an online form you can fill in or you can download a printable application form to complete and send in by post. If you are reading the hard copy of this booklet, please tear off the membership form attached to the bookmark, complete it and send it to us.

*if you pay by direct debit.
Fatigue Matters
A self-help guide for people living with rheumatoid arthritis

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