A HOME EXERCISE BOOK

Physiotherapy management for Duchenne muscular dystrophy
Here at the Muscular Dystrophy Campaign we are very aware that there is an increasing need for guidance on physiotherapy and exercise for those with Duchenne muscular dystrophy.

This 2009 third edition of our popular guide, which was first published in 1993, passes on our understanding of the benefits of regular gentle exercise and physiotherapy for children with muscle disease. It is part of our ongoing commitment to provide free, expert information.

Written in a simple easy-to-follow format, with clear illustrations and input from experts in this specialist field, this resource offers up-to-date information and guidance for individuals, their families and professionals working with them.

Our thanks go to Lyn Jenkins, Children’s Physiotherapy Service Manager, who helped update the guide and advise on illustrations. Lyn works at the Children’s Centre at Frenchay Hospital in Bristol, a specialist clinic which provides physiotherapy assessment for a large number of children with Duchenne muscular dystrophy. Thanks also to the specialist physiotherapists who have given their time to reviewing previous editions of this publication.

This guide is a revision of the following Muscular Dystrophy Campaign publications:


Second edition of *Physiotherapy Management for Duchenne Muscular Dystrophy* 2001

And incorporates material from *Duchenne Muscular Dystrophy, A Team Approach to Management*, written by Helen Posselt, Australia

Should you require further information or support using this guide, or advice on other free resources available from the Muscular Dystrophy Campaign, please contact us:

0800 652 6352 (freephone)
info@muscular-dystrophy.org
www.muscular-dystrophy.org

September 2009
Muscular Dystrophy Campaign
Contents

1. Introduction .................................................. 2
2. Exercise ..................................................... 7
3. Good posture ............................................... 9
4. Mobility ...................................................... 13
5. Stretches ..................................................... 16
6. Back care for carers ....................................... 18
7. Children’s questions ..................................... 19
8. Contacts and resources .................................... 21
9. Glossary ..................................................... 23

Stretches and exercises  (enclosed sheets)
1. Stretch for the ankles
2. Stretch for the knees
3a, b, c. Stretch for the hips
4a, b. Iliotibial stretch
5a. Stretch for the elbows
5b. Stretch for the elbow and wrist
6a, b. Stretch for the wrist, elbow and fingers
7. Self-stretch for the calf
8a, b. Self-stretch for the knees
9. Deep breathing
10a. Postural drainage
10b. Postural drainage for older children
11. Assisted coughing
1. Introduction

This book is intended to support the practice of physiotherapy at home for children and young people with Duchenne muscular dystrophy.

What is physiotherapy?
Physiotherapy is the physical treatment and management of a disease or condition which enables people to reach their maximum physical potential. Physiotherapists help to ensure that their patients lead as fulfilling a life as possible by advising children, families, carers and school staff about how the condition affects physical development.

In Duchenne muscular dystrophy, the physiotherapist will help to:
- minimise the development of contractures and deformities through a programme of stretches and, where appropriate, exercises
- anticipate and minimise any secondary physical complications
- identify and prescribe aids and equipment (orthoses, callipers, wheelchairs and standing frames, for example)
- advise on moving and handling issues
- monitor respiratory function and advise on techniques to assist with breathing exercises and methods of clearing secretions.

Physiotherapy at home
The exercises recommended by your physiotherapist need to be done regularly if they are to be effective. This means developing a routine at home.

Any physiotherapy regime should be based on the:
- needs of your child
- advice of your physiotherapist
- needs of the family (a practical routine to suit your family’s lifestyle).
It is important that any routine is right for you and your child and, where possible, fits in with other family activities. If mornings are hectic, for example, it may be best to have a physiotherapy session during the evening, at bed or bath time. When the exercises are done is not important as long as they are part of an established routine.

Although the exercises should never be painful, stretching exercises may cause the muscle to feel different and your child will need to become accustomed to this. Some children will be able to do self-stretching exercises, as well as exercises with a carer, and a physiotherapist can help with this.

There has been some concern that over-activity may cause more harm than good to a child or young person with muscular dystrophy. Exercise should be at a moderate level\(^1\) and not cause severe fatigue, however exercise with weights and eccentric exercise (where the muscles are having to work against the body load while slowly lengthening) should be discouraged. Any activity that a child does voluntarily and without becoming overtired should have a positive effect.

**Muscles**

Muscles allow us to move, stand, and perform the range of movements needed for daily living. Each muscle is made up of fibres (the type and amount of fibre varies depending on the sort of work the muscle does). The muscle fibres we use for standing are, for example, different to those in the muscles used to make fine, quick, finger movements.

Muscles are attached to bones via specially adapted parts of the muscle, called tendons. A muscle spans at least one joint and a movement occurs when it contracts or shortens. Muscles and their tendons are normally very flexible, allowing movement through lengthening and shortening. Usually when one muscle contracts or shortens the opposite muscle lengthens.

In Duchenne muscular dystrophy, muscle fibres break down and are replaced by fibrous and/or fatty tissue causing the muscle to gradually

---

\(^1\) A Moderate Level (borg scale 3-4) is feeling warmer, breathing faster but still able to hold a conversation.
weaken. The rate at which this happens can vary between children with the same condition. Some muscles will be affected earlier than others and if one muscle weakens sooner than another it can upset the normal balance of strength and cause contractures.

**Contractures**

When muscles are not used or become weak, they lose their stretchiness along with the associated tendons and ligaments (the tissue around the joints which connects bones, and controls the extent or range of movement). The joint becomes stiff and tight, usually more in one direction than the other. When a joint becomes fixed in one position, this is known as a contracture, and a deformity may occur.

As soon as possible after diagnosis, seek advice about physiotherapy and start treatment and management aimed at preventing contractures. Take action before there is any tightness or obvious deformity.

The most frequent contractures for children in the early stages of Duchenne muscular dystrophy occur at the ankles and hips. These are partly caused by the walking position which the child adopts – on the toes with feet apart – to maintain balance as the hip, knee and trunk muscles weaken. Children in the later stages of the condition spend more time sitting down which increases the tendency to develop hip, knee and ankle contractures.

**Adapting to changes**

Because a child with Duchenne muscular dystrophy will go through various stages of development, the physiotherapy you do at home should reflect these changing needs.

In *early developmental stages* the child will enjoy acquiring gross motor skills such as rolling, crawling and walking, and maybe running, jumping and hopping. All these activities provide good opportunities for learning and development but some will need to be adapted as the child loses strength and tires more easily. Over time, the child’s skill acquisition and strength will level out.
During this stage of the child’s development it is important to encourage activity that does not cause extreme fatigue. Exercise with weights and eccentric muscle work such as trampolining should not be encouraged. Parents or carers may want to consider introducing a physiotherapy and exercise programme that includes:

- regular stretches – self and/or manual stretches as well as passive stretching – for the muscle groups that are tightening (tendo-achilles, hamstrings and iliotibial band)
- swimming, hydrotherapy
- wearing orthoses (splints) at night to slow down contractures in the ankles
- insoles – may be helpful for flat feet, used in ordinary lightweight footwear
- general games and fitness encouraging cardiovascular fitness.

In the **later developmental stages**, there will be a progressive loss of function. Although the child will be walking for periods of time, he or she will also require a wheelchair for mobility, especially over long distances. There is a range of equipment that can be used to help, such as electric or manual wheelchairs, standing frames etc. Any equipment should be thoroughly assessed by a qualified professional before it is used. The upper limbs will also be weaker but daily function activities should be encouraged.

A physiotherapy programme at this later developmental stage may include:

- regular passive stretches for tendo-achilles, hamstrings, hip flexor and iliotibial band muscles; some self-stretches may also be recommended
- stretches to the upper limb muscles
- swimming, hydrotherapy
- wearing orthoses (splints) at night
- prone lying and other good positioning
- games to promote deep breathing including blowing bubbles and playing wind instruments.

Physiotherapy advice and support continues throughout the wheelchair stage. Both good posture and a manual stretching programme are still important.
A physiotherapy programme at the **wheelchair stage** may include:

- regular stretches to minimise the development of contractures in hips, knees and ankles and ensure comfort in bed, ease in dressing and positioning in wheelchair
- use of ankle splints when sitting in wheelchair (instead of use of orthoses)
- chest clearance techniques and assisted cough
- stretches for the upper limbs to minimise contractures
- using a standing frame
- prone lying and other good positioning
- swimming, hydrotherapy
- use of orthoses.

Physiotherapy support may need to be adjusted following any surgery for scoliosis. In particular, the wheelchair may need adapting to accommodate the child’s improved posture.

**How to use this booklet**

This booklet provides guidelines and instructions on how you can use physiotherapy to help your child with Duchenne muscular dystrophy.

All the stretches and exercises are on separate sheets so they can be combined, on the advice of your physiotherapist, into an individual physiotherapy programme that suits your child’s needs. Any routine should be reviewed regularly as your child’s needs change.

The stretches, exercises and guidance included in this booklet are general, and additional ones may also be recommended.

**Please refer to the glossary on page 22 for help with any technical terms used in this guide.**
It is natural to worry about how much exercise your child should do and whether it’s possible to do too much or too little. Finding a balance can be difficult and very often your child is in the best position to say how much is right for him or her. Exercise should never be done to the point of extreme or severe fatigue, although it is unlikely that you could persuade your child to do this.

Research from the British Heart Foundation indicates that all children should exercise at a moderate level\(^2\) for at least one hour a day. A child with Duchenne muscular dystrophy should be able to exercise on a daily basis.

Walking is good exercise and should be encouraged whenever possible, although this can be difficult if your child falls frequently. Children with muscular dystrophy may tire more quickly but can walk for some distance and time if they are not rushed. They may struggle to walk uphill, downhill, and on uneven surfaces, such as sand or grass. Using a wheelchair for longer distances can alleviate tiredness, enabling your child to enjoy him or herself more on arrival.

It is important to not limit your child’s play by being too overprotective. Physical and appropriate sporting activities can be beneficial to a child with Duchenne muscular dystrophy, helping to maintain strength and increase self-confidence. Swimming, horse riding (depending on muscle strength), bicycling, tricycling (there are low geared tricycles available) and other general activities all help children to relax, enjoy themselves and socialise with their peer group.

Swimming is particularly good exercise at all ages, both for the muscles and the lungs, and children may be able to swim or take lessons at school. It is important that children with muscular dystrophy stay warm in the water, either by continually moving or perhaps by using a children’s pool, which is often warmer than a full size pool. The changing facilities also need to be warm, as well as accessible, to prevent the children becoming cold when they leave the water.

\(^2\) A Moderate Level (borg scale 3-4) is feeling warmer, breathing faster but still able to hold a conversation.
Children need to be as active as possible and not spend too much time sitting in front of the television, computer or using electronic games. They could play computer games while standing at a table, or lie on their tummy to help stretch the hips while reading or watching television. Encourage at least one activity a day that involves some exercise. It may be difficult in winter to find a suitable activity, but children can still help with household tasks such as tidying bedrooms or setting the table.

**Breathing exercises**

Breathing exercises become important as the child becomes less able to actively exercise.

When we breathe in (inspiration), muscles lift the rib cage up and out, making the chest larger. Air then rushes into the lungs to fill the extra space created. When we breathe out (expiration) the muscles relax and the air is pushed out by the elasticity of the lungs. We only use muscles of expiration when air is forced out, as in coughing.

As the respiratory muscles weaken in children and young people with Duchenne muscular dystrophy, this reduces the ability to inhale and exhale air forcefully. It becomes more difficult to cough and expel mucus from the lungs, which affects the amount of oxygen in the body and increases the likelihood of chest infections.

An incentive spirometer can help good expiratory (breathing out) technique. Inspiratory muscle training against resistance is not recommended. It may also be helpful to encourage your child to play a wind instrument or join a singing group. Younger children could blow bubbles.
Muscle weakness in key areas such as the spine and hips can affect the posture of a child with Duchenne muscular dystrophy.

Weakened hip extensor muscles cause lordosis (pronounced inward dip of the lower spine when standing). Weakness of the trunk muscles can cause scoliosis (curvature of the spine). When one side of the body is stronger than the other, this can cause an asymmetrical posture.

Your child may adopt unusual postures – in sitting, standing and lying – to compensate for muscle weakness, limited mobility and contractures. It is important to correct these postures because, if left, they can cause further problems, particularly in the spine. Good seating at all times helps to maintain good posture.

**Sitting**

The feet should be at a 90° angle to the legs when the child is sitting down. The seat of the chair should be firm and, ideally, not too wide. The back of the chair also needs to be firm and either upright or slightly slanting backwards (10°). The seat should be as deep as the thigh is long, so that the child is encouraged to use the back of the chair and not slump. The armrests need to be at the right height and not too far apart so that the elbows can be supported without causing hunched shoulders or leaning.
Positioning

The way a child moves and the positions adopted – to write, eat or rest, for example – are a direct response to losing muscle strength and having contractures. The child will naturally find the easiest and least tiring option, without thinking about it. Sometimes muscle strength and/or the stiffness of a contracture may be different on each side of the body. When this happens, an asymmetry or imbalance occurs which can cause scoliosis.

Passive stretching and night splints can delay the onset of contractures but it is important to know which positions to encourage and which to discourage, without nagging.

Prone lying

The prone lying position (face downwards) is good for resting. It can also help prevent contractures developing in the hips and knees. Prone lying can be combined with activities such as reading or watching television.

The child lies face down on a floor, couch or similar firm surface. Place a small pillow or wedge just below the hips (which should be level and the pelvis down) to encourage hip extension. The weight of the lower leg will straighten out the knees but it is important that the feet are free.

Discourage asymmetrical positions as these reinforce development of contractures and scoliosis.
Standing

Standing helps bone density and posture as well as assisting in the management of contractures. It should be encouraged, during the day, for short periods (i.e. half an hour) or longer blocks of time (two or three hours if possible, but you need not be prescriptive).

When an older child or young adult finds it difficult to stand unsupported, but callipers are unsuitable, it may be helpful to use a standing frame, swivel walker or tilt table. They reduce the muscular effort required to stand upright and provide total body support, enabling the hip flexor, knee flexor and calf muscles to be fully stretched. Using a standing frame every day can delay the onset of scoliosis as well as aiding digestion and circulation. Children who have callipers often use these for standing.

Night splints

As the name suggests, these are designed to be worn at night and are usually only for the ankles. They help slow down contractures by keeping the joint in the best position for the child. Night splints are made from a variety of materials, including polypropylene. The splint starts at the toes and finishes just below the knee. They must be comfortable and fit properly, as poorly fitting splints are unlikely to be worn and may put the child off using splints.

Research has shown that using night splints in conjunction with passive stretching is the most effective way of delaying the development of contractures. Night splints are never, however, a substitute for passive stretching and should only be used in combination with stretching once there is an obvious feeling of tightness.

Day ankle splints are very rarely worn by walking children as they can adversely affect mobility, make it more difficult to walk and increase the risk of falling. However, children who use wheelchairs can and should wear day ankle splints to prevent contractures.
Sleep systems

When the child has difficulty moving or turning in bed it is important that he/she is positioned carefully to prevent the likelihood of muscle shortening (contractures). There are a range of sleep/positioning systems, which can help prevent pressure areas from developing and keep the child in a good position for longer. A sleep system can also reduce the number of times parents need to turn the child, thereby improving quality of life for all family members. Your physiotherapist can conduct a sleep posture assessment to determine whether a sleep system is appropriate for your child’s needs.
4. Mobility

‘What happens when walking becomes difficult?’ is a question asked by all parents and an area of understandable concern. This chapter includes information on alternative choices and ways to help your child. Do not be afraid to also discuss this issue with your doctor and/or physiotherapist.

There are several signs that suggest your child could be finding walking more difficult:

- He/she may seem quieter or less inclined to join in with activities.
- Teachers might report that your child has been left behind.
- The number of falls may increase.

Falls are more likely to occur when the child is outside, negotiating uneven ground, slopes and kerbs. Falls are not necessarily caused by tripping; muscle weakness may make the child drop. If the trunk muscles are weak, for example, it can be difficult to make the necessary postural adjustments and the child can over-balance. Falling can also occur more frequently when the child is tired.

It may be apparent at your child’s regular physiotherapy assessment that muscle weakness and possible contractures in the hips, knees or ankles are reaching a critical point. A sudden growth spurt, a bad fall or another illness may also precipitate loss of independent walking.

**Callipers (KAFOS)**

Some children can carry on walking independently for up to two years or longer by using orthoses called Knee Ankle Foot Orthoses (KAFOs). This treatment was developed at Hammersmith Hospital, London, based on work done in Chicago, USA. The young person uses a wheelchair for long distances – as a car is used for speed and convenience – but walks short distances at home. Because children with Duchenne muscular dystrophy have muscle weakness in the arms, shoulders and trunk, they cannot use walking aids such as crutches.
A KAFO is an ‘ischial weight bearing knee ankle foot orthosis’, which makes use of the lordosis with which the child has become accustomed. It extends from the toes to the hip and the child sits on, or is supported by, the lip at the top of the thigh piece. KAFOs are made of polypropylene and have a hinge which allows the knee to bend when sitting. No special shoes are required and they can be worn beneath trousers, so are fairly unobtrusive.

The decision to use orthoses must be made by both the family and child, based on information and advice from a doctor and physiotherapist. There are many things, unique to each child and family, which need to be considered and discussed, such as the number and age of other children in the family, lifestyle, personality, type of school and attitude of teaching staff etc. KAFOs will not be suitable for everyone and your child may choose to use a standing frame instead.

**How are they fitted?**
To fit the orthoses, the foot must be at a right angle to the leg. To do this, an orthopaedic surgeon usually releases the tight tendo-achilles (heel cord). This is a small operation, which generally causes very little pain. Your child’s consultant or physiotherapist can explain what choices are available locally, or at the Muscle Centre you attend.

**When to do it**
Orthoses do not need to be fitted before the child has either stopped walking independently or is falling more frequently. It must be done, however, before the child has been dependent on a wheelchair for more than two or three months, and is best done when the child has only just lost the ability to walk independently. It is important to discuss the use of KAFOs well in advance of losing the ability to walk so that you know who to contact, what to do and when.
Who is suitable?
- The child with enough strength in the hips and trunk to balance.
- The child who wants, and will accept, them.
- The family who can manage.

What are the advantages?
- Potential to increase independence at home and school.
- Being the same eye level as friends.
- A delay in the development of scoliosis and contractures in the hips and knees.
- Easier transfer between chair and car, etc.
- Easier toileting.

When the young person is unable to use the orthoses for walking they can still be used for standing, and help to make transfers easier between chair, car and bed etc.

Wheelchairs
The majority of children with Duchenne muscular dystrophy will need a wheelchair for transportation and independent mobility, long before they have lost the ability to walk. Often two wheelchairs – one manual, one electric – are required, to suit life at school and home. Choosing the right wheelchair is absolutely essential for the child’s well being. Technical decisions about seat size, seat cushion, back support, headrest, position of power controls, tilt-inspace facility etc, must be made with great care. Experts such as your physiotherapist and/or occupational therapist, the physiotherapist at the local muscle centre/specialist clinic, local wheelchair services, Disabled Living Centre, Whizz-Kidz assessor should be consulted.

It is essential that any wheelchair, once in use, is routinely checked and monitored so that it continues to meet the child’s requirements. As the child grows and their physical condition changes, the wheelchair will need to be modified to offer additional support and maintain independence.
5. Stretches

It is very common in muscular dystrophy for muscles and tendons to tighten. Some muscles will be affected earlier than others. The first muscle to tighten is usually the calf muscle/achilles tendon at the ankle, but the muscles around the hips, knees, elbows and fingers can also be affected.

Contractures can make some movements and activities more difficult. Regular daily stretches help maintain muscle length and keep joints mobile.

Your physiotherapist can create an individual programme for your child from the exercises at the back of this book. The descriptions and diagrams are only reminders and anybody undertaking passive stretching must receive professional guidance and instruction.

There are three different types of stretches: **passive**, **active assisted** and **self-stretches**.

**Passive stretches**

Passive stretches are an essential aspect of physiotherapy management and an essential aspect of any programme, at all stages of the condition. It is never too soon to introduce passive stretching.

As the name suggests, the child does not actively take part in the stretching process. Passive stretches are done by a parent, carer or therapist. Slow and firm passive stretching will not harm the joint or muscle and can be done every day. Tight and/or shortened muscle tissue is stretched by moving the joint as far as possible and maintaining the position for at least ten seconds (your physiotherapist may recommend longer, depending on your child’s needs). Done properly and effectively, passive stretching is not painful but your child will experience a sensation of pulling and be aware of gentle but sustained pressure being applied. Some children put up a mild protest at passive stretching but this is usually overcome once their confidence has been gained and a routine established.
It doesn’t matter what time of day you do the passive stretching but most people find it helps to establish a routine that fits in with the other demands of family life. Make it a special time for the child, when the day’s events can be shared and emotional ties strengthened.

It may help to do the stretching:
- after a bath
- with tapes or stories, singing, story telling
- after massaging the muscles to be stretched

Position the child so that he or she is well supported and comfortable, and the joints not being moved are stabilised. The child must relax completely and not make any active movement or resist the stretch. If the stretching is done too quickly, the child is more likely to resist and become frightened.

Discuss and agree the duration of the stretch. Start the stretch gently and gradually increase to a maximum intensity, without pain. Overstretching should be avoided.

**Active assisted stretches**

Active assisted stretches are done by the parents with the child assisting the movement. When a joint becomes contracted, the tight tissue prevents the opposite muscle group from working properly. Active assisted stretches can stretch the tightened muscle and work the opposing muscle group at the same time.

Active assisted stretches are particularly useful for the ankle. While you stretch the Achilles tendon, for example, your child pulls up his or her toes. The harder you work together, the more effective the stretch will be. This form of stretching helps the time pass more quickly and makes the stretches less boring for your child.

**Self-stretches**

Self-stretches, as the name suggests, are stretches that the child is taught to do him or herself. These are most effective in children who are still walking and are particularly useful for the ankles, knees and hips.
Parents and carers can look after themselves by following some basic general rules:

- always bend the knees and keep the back straight when bending down to pick something up
- always be sure that you can manage the weight of whatever you are picking up
- always hold whatever you are carrying as close to your body as possible
- avoid carrying heavy loads up and down stairs

One of the most common ways to incur a back injury is to ‘lift and twist’ (when lifting a child out of a bath or car, for example). Always plan the best way to lift something or someone, do not rush, and use available equipment to help.

Physiotherapists can advise you on safe ways to move your child and you can also consult the manual handling advisors and risk assessors at your child’s school. If hoisting is recommended, your physiotherapist can refer your child to a social services occupational therapist (OT). Most doctors’ surgeries also have leaflets on how to protect the back when lifting.

For further support, information and general advice on looking after your back, contact Back Care – the charity for healthy backs – on 0845 130 2704 or at www.backcare.org.uk
7. Children’s questions

Q. What is the point of stretching exercises?

A. Stretching exercises are perhaps the most important exercises for all ages and keep your joints comfortable and moving. Athletes do these exercises to keep their muscles in shape.

Anybody who sits for a long time in the same position will get stiff and uncomfortable. Physios and doctors call this stiffness a ‘contracture’. You may not have a problem bending your knee, but sitting down a lot can make it difficult to straighten up.

Joints that don’t have a full range of movement can get achy and it’s sometimes difficult to find a comfy position in bed. So best to just get on and do those stretching exercises!

Q. If my muscles are weak why can’t I use weights to make them stronger?

A. Using weights won’t make your muscles stronger because your muscles are different from other peoples. Overworking the muscle by lifting heavy weights could damage it. But there are other active exercises which you can do that are good for you.

Q. Why does my physio want me to go swimming or have hydrotherapy when I just think it’s a pain getting undressed and dressed again?

A. Hydrotherapy is probably the best way to exercise if you have a muscle condition. The warm water supports your body and relaxes your muscles, which makes stretching exercises much easier. The best thing about swimming or hydro though is being able to move much more freely than you usually can and it’s really good fun!
Q. **What about my hands and arms?**

A. Hands and arms need to be stretched too. Playing computer games is fun and your fingers need to be flexible in order to be an expert! Ask your physio what you can do (yes – it’s more stretches!).

Q. **What about my feet?**

A. Looking good is important too. If contractures of the feet and ankle develop, this might make it difficult to wear the sort of trainers or shoes you prefer. Wearing splints will keep your feet in the right position so that you can look good (and yes, there are stretches that will help too!). The position of the upper leg is very important when looking at your foot posture. If your legs flop apart, your feet turn onto their sides (try it!).

Q. **What if it gets hard for me to walk?**

A. There is nothing wrong with using a wheelchair for some of the day to save energy for playtime. Keeping your legs stretched with the help of your physio or carer will help keep you walking for as long as you can. Even when it gets really hard, there are other ways to take those important steps, such as using callipers (KAFOs).
8. Contacts and resources

Contacts

Muscular Dystrophy Campaign
61 Southwark Street
London SE1 0HL
0800 652 6352 (freephone)
info@muscular-dystrophy.org
www.muscular-dystrophy.org

Duchenne Family Support Group
78 York Street
London W1H 1DP
0870 241 1857
0800 121 4518 (helpline)
www.dfsg.org.uk

Action Duchenne
41 West Street
London, E11 4LJ
020 8556 9955
www.actionduchenne.org

Resources

Inclusive Education for Children with Muscular Dystrophy and other Neuromuscular Conditions (second edition 2007)
Muscular Dystrophy Campaign
ISBN 0-903561-08-5

Wheelchair Provision for Children and Adults with Muscular Dystrophy and other Neuromuscular Conditions – Best Practice Guidelines
Muscular Dystrophy Campaign
ISBN 0-903561-13-1

Duchenne Muscular Dystrophy – A Team Approach To Management
book and two videos (one video is An Overview of Physiotherapy Management). Produced by MontroseACCESS.
information@montroseaccess.org.au
Booklets for children and young people
(available from Muscular Dystrophy Campaign)

Everybody’s Different, Nobody’s Perfect is aimed at four to ten-year-olds with a neuromuscular condition. 
DMD On the Ball is aimed at teenage boys aged 10 to 14 who have Duchenne muscular dystrophy. 
Hey, I’m Here Too is aimed at siblings of children with Duchenne muscular dystrophy.

Factsheets (all written by clinicians and health specialists on behalf of the Muscular Dystrophy Campaign)
- Duchenne muscular dystrophy
- Duchenne muscular dystrophy – the older child
- Exercise and orthoses guidelines
- Making breathing easier
- Steroids and Duchenne muscular dystrophy

All Muscular Dystrophy Campaign literature is free and downloadable from our website.

For further support:
- call our freephone Information Line on 0800 652 6352
- email info@muscular-dystrophy.org or
- visit www.muscular-dystrophy
9. Glossary

**Glossary of physiotherapy terms**

**Abduction** – movement of a limb away from the body

**Adduction** – movement of a limb towards or across the body

**AFO** – ankle foot orthosis. In Duchenne muscular dystrophy this usually means night splints worn in bed to prevent the foot from pointing down

**Assisted coughing** – a technique used to help clear secretions on the chest

**Asymmetry** – when one side of the body is different to the other

**Atrophy** – decrease in muscle size and strength

**Callipers** – see KAFO

**Cardiac** – affecting the heart

**Contracture** – when a joint cannot move through full range because the muscles are tight

**Distal** – part of the limb the furthest away from the body (e.g. the hand or foot)

**Dorsiflexion** – pulling the foot up to a right angle

**Extension** – a straightening movement

**Flexion** – a bending movement

**Hydrotherapy** – water based exercise, usually done in a warm pool under the supervision of a physiotherapist

**Hypertrophy** – increase in muscle size, usually in the calf muscles (in Duchenne muscular dystrophy this is usually because the muscle contains a lot of fat)

**Gastrocnemius** – the calf muscle

**Gluteal muscles** – the buttock muscles, used to stand upright and climb stairs

**Gower’s sign/Gower’s manoeuvre** – a way of getting up from the floor into a standing position by pushing the hands on the legs

**Gross motor skills** – movements such as crawling, running or jumping which use the large muscles of the body

**Hamstrings** – the muscles at the back of the knee which help it to bend and also stabilise the pelvis

**Ilio-tibial band (ITB)** – the fibre on the outside of the thigh running from the hip to knee, which has muscles attached to it

**KAFO** – a shortened name for a knee ankle foot orthosis which extends from the toes to the hip
**Lordosis** – the extended position of the lower back, visible as an inward dip at the base of the spine, typical in children with Duchenne muscular dystrophy

**Muscle biopsy** – removing a small piece of muscle for examination

**Night splints** – made of polypropylene and worn at night to prevent contractures; usually start at the toe and finish just below the knee

**Orthoses** – another term for splints, callipers or anything worn externally to support the limb

**Passive stretching** – a technique used to stretch tight muscles by moving the joint as far as possible and maintaining that position

**Patella** – kneecap

**Physiotherapy** – the physical treatment or management of a disease or condition through a specially designed programme of exercising, stretching, positioning etc

**Plantarflexion (of the foot)** – pointing the foot down

**Postural drainage** – using physiotherapy to clear phlegm from the chest

**Prognosis** – the expected course and outcome of a condition or disorder

**Pronation (of the forearm)** – turned palm down

**Prone** – lying face down

**Proximal** – part of a limb nearest to the body (e.g. the shoulders or pelvis)

**Quadriceps** – the muscles at the front of the thigh that straighten the knee

**Resistance** – using weight or manual pressure to strengthen the muscles

**Rotation** – a movement which turns a part of the body

**Scapula** – shoulder blade

**Scoliosis** – sideways curvature of the spine

**Spinal jacket** – a corset or brace made of polypropylene or leather worn to keep the spine straight

**Standing frame** – equipment that holds the child in a standing position with minimal effort from the child

**Steroids** – drugs used primarily to treat muscle inflammation but also able to slow down deterioration of the muscle

**Supination (of the forearm)** – turned palm up

**Supine** – lying face up

**Tendon** – the fibrous part of the muscle which is fixed to the bone

**Tendo-achilles (TA)** – the large tendon at the back of the heel

**Tilt table** – equipment that can take a child from a standing position to a lying position without any effort from the child

**Walking frame** – equipment, usually on wheels, designed to give the child stability while walking
Get involved

Physiotherapy Network

The Muscular Dystrophy Campaign set up this network to bring together physiotherapists who work with those living with muscle disease, and their carers.

The network enables those who work in specialist care centres and the community to share information and best practice. It also works in partnership with our National OT Network. It is free to join. Members receive updates on training opportunities for professionals as well as details of information and study days and social events for parents.

To join the Physiotherapy Network, please contact us on: 0800 652 6352 (freephone), email info@muscular-dystrophy.org or visit www.muscular-dystrophy.org

My Online World

My Online World is a fabulous new website aimed at children aged 18 and under. It features specially designed games, jokes, riddles, news, links to videos, information about muscular dystrophy and four fabulous chatrooms. We’ve split the site in two so children under 13 see different content to those over 13.

You can design your own character and check out other people’s crazy styles. If you want to share your ideas with the world you can blog or keep it short by tagging the thought tree. The site was funded by Schroders.

Check out www.myonlineworld.org.uk