Motor neurone disease (MND) is a progressive and terminal disease that results in degeneration of the motor neurones, or nerves, in the brain and spinal cord. Many people with MND will experience dysphagia (swallowing problems), which can make eating and drinking difficult. This can cause anxiety for people with MND and their carers/family, who may have concerns about choking on food and liquids. This information provides suggestions, medication options and practical tips for health and social care professionals working with people with MND.

Everyone with MND has a different experience of the disease. There is no standard rate of progression. Not all symptoms affect everyone, and it is unlikely that they will develop at the same time or in any specific order. Therefore, each person should have an individual assessment for treatment.

What is dysphagia?

Dysphagia is a disruption of the normal eating and swallowing process. It affects many people during the course of MND. People with MND may have swallowing problems caused by weak muscles in the bulbar region (face, mouth, tongue and throat). Those whose first symptoms affect this region (bulbar-onset) usually experience dysphagia at an earlier stage than those with other types of MND.

Find out more

See our information sheet for health and social care professionals:
• Information sheet P3 – Managing saliva problems in motor neurone disease

Information you can share

We have two relevant information sheets for people affected by MND:
• Information sheet 7A – Swallowing difficulties
• Information sheet 7B – Tube feeding

Download from our website at www.mndassociation.org/publications or see page 9 for details of how to order copies.

MyTube is a useful online resource containing short videos and information about tube feeding for people with MND. Search mytube.mymnd.org.uk/
**Referrals to other professionals**

As this symptom will get worse, it is crucial that referrals are made to the appropriate professionals as soon as a problem is identified. Referrals should be made by whichever professional first notices a problem. Specialists at MND care centres or networks, or through multidisciplinary teams (see below), can advise on dysphagia management and refer to appropriate services. Unregistered professionals have a responsibility to report concerns or deterioration immediately to their line manager.

The **specialist respiratory team** can determine whether the person with MND has respiratory involvement and risk of aspiration and/or choking, and can help the person with MND determine the course of action.

Referral to a **speech and language therapist (SLT)** should be a priority. They can assess the ability to eat and swallow and will put together a plan of action, which may include teaching swallowing techniques and altering the consistency of diet, for example by introducing a puréed diet. Often the SLT will indicate whether the person needs to see a dietitian and these two professionals will work closely together.

Weak bulbar muscles will also affect communication. The SLT can assess speech problems and whether any aids or equipment might be needed to help the person with MND communicate.

A **dietitian** can assess diet, nutritional and fluid intake and weight. Weight loss is often related to dysphagia, but it can also be affected by a person with MND not having the strength to feed themselves because of extreme fatigue or weakness in their upper limbs. Low mood or changes in bowel habit are other possible causes. A dietitian can suggest ways to optimise the person’s diet and nutritional intake (see page 3).

The person may be referred for an assessment for **enteral feeding** to determine whether alternative ways of feeding are needed.

A **physiotherapist** can advise on head supports and positioning and teach simple cough assist techniques to help with secretion management.

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**Signs of dysphagia:**

- coughing or choking when eating or drinking
- poor lip control
- reduced ability to cough
- problems swallowing medication
- several swallows needed for each mouthful of food
- weak chewing, which takes more effort
- eating and drinking is tiring
- food may not move to the back of the mouth because of a weak or absent swallow reflex, or because of an unco-ordinated swallow caused by weak pharyngeal muscles
- wet or muffled sounding voice after eating
- spasm of the bulbar muscle
- sialorrhoea – drooling due to impaired swallowing of saliva rather than excess production.

**Dysphagia can lead to:**

- dehydration and weight loss – because the person is unable to take in the food and fluids they need
- aspiration – where food or drink goes into the lungs – which can be life threatening
- recurrent chest infections – often caused by aspiration
- loss of strength
- tiredness
- constipation
- loose, dry and flaky skin, and poor skin integrity
- loss of wellbeing and reduced quality of life
- lengthy mealtimes, with food going cold
- feeling unable to take part in family mealtimes
- burden and stress for carer or family members, who may eat separately and prepare food that may not be eaten.

Dysphagia is also associated with impaired respiratory function.
An occupational therapist may advise on head supports, seating and positioning, food and drink preparation, and may recommend strategies and equipment to help with eating and drinking, such as adapted cutlery.

**Managing dysphagia**

When eating and swallowing become difficult the following may help:

**Food**
- Choose softer food which is easier to chew.
- Cut the food finely on the plate or mash it well with a fork so it requires less effort to chew.
- Add extra sauce or gravy, as moist food is easier to swallow.

As mealtimes become more difficult, people with dysphagia usually find that blended or puréed food is easier to swallow. It is often preferable to blend the different foods separately to retain the flavours, rather than blending the entire meal together.

In general, when someone has a swallowing problem, **they should avoid**:
- mixed textured foods and liquid with ‘bits’, eg minestrone soup
- foods that need a lot of chewing, eg fresh bread or some meats
- stringy food, eg celery or some green beans
- coarse, hard food, eg well done toast, or crumbly, dry food, eg biscuits or flaky pastry
- some vegetable skins which can be difficult to clear in the mouth, eg tomato or sweetcorn
- foods that become sticky in the mouth, eg bread or mashed potato.

However, it is best to seek advice from an SLT, as everyone’s swallowing ability is different.

**Drink**

People with MND sometimes cough when drinking. This may be due to weakness and lack of co-ordination of the muscles used to swallow. It may be a sign of aspiration (drink going down towards the lungs). Thicker drinks tend to move more slowly and are therefore less likely to cause aspiration. An SLT might recommend a thickening powder, available on prescription, which can be added to drinks.

If drinking is difficult, an individual may gradually reduce their fluid intake. Choosing moist foods with a high fluid content, eg soups, casseroles, stewed fruit, ice cream and custard, helps to prevent dehydration and associated symptoms.

**Assessment and management by the dietitian**

When someone has swallowing problems and can take in less food, the nutritional content of meals is particularly important. A dietitian can recommend a diet based around advice from the SLT. They will suggest different types of food and ways to fortify food to improve the nutritional content. They may recommend food supplements which are available on prescription.

As part of the management of a swallowing difficulty, alternative methods of feeding may also be discussed and considered (see page 5).

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**Information you can share**

Information sheet 7A – *Swallowing difficulties* contains information for people with MND and their carers on how to manage episodes of choking. You can download a copy of this sheet from [www.mndassociation.org/publications](http://www.mndassociation.org/publications) or call 0808 802 6262 to order.

**Tips for meal times**

These tips may help if you’re involved in the preparation and serving of food for someone with MND, at home or in a care setting.

- Serve several small meals rather than one large one: this can be less tiring for the person eating.
- Use garnishes and attractive colours to make food more appetising, and make sure they can see and smell the food.
- Allow plenty of time for the meal.
- Keep food warm and reheat if necessary. Using a keep-warm plate may help.
- Let the person concentrate on what they’re doing – allow them to swallow before asking questions.
- Leave a gap between courses if eating is slow and tiring.
- Sipping an iced drink or consuming anything cold or frozen (e.g. ice cream) before eating and between mouthfuls can stimulate a stronger swallowing reflex.
- A relaxed, quiet environment may help the person to feel less anxious.
- The person with MND may feel unable to eat with their family or friends because of dysphagia. They may instead join the family for social reasons and take a few spoonfuls, while eating most of their meal before or afterwards.
- Encourage good mouth hygiene before and after meals, including swilling the mouth, brushing teeth and/or using a suction unit to clear away food debris.

**Equipment that may help**

<table>
<thead>
<tr>
<th>Potential problem</th>
<th>Things to try</th>
<th>Who can help?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Posture</td>
<td>Sitting upright in a firm, high-backed chair</td>
<td>Occupational therapist</td>
</tr>
<tr>
<td></td>
<td>Keeping head erect with chin tucked in</td>
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<tr>
<td></td>
<td>Adequate support for arms</td>
<td></td>
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<tr>
<td>Weak neck muscles</td>
<td>A collar or chin support may help – although this may make swallowing more difficult</td>
<td>Orthotics, occupational therapist</td>
</tr>
<tr>
<td>Weakened grip and limited wrist movement</td>
<td>Specially designed plates, cups and cutlery</td>
<td>Occupational therapist</td>
</tr>
<tr>
<td></td>
<td>Attaching cutlery to splints</td>
<td></td>
</tr>
<tr>
<td>Difficulties in lifting food/drink to mouth</td>
<td>An adjustable cantilever table</td>
<td>Occupational therapist, speech and language therapist</td>
</tr>
<tr>
<td></td>
<td>Specially designed cups</td>
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<td></td>
<td>A bed tray (with small legs) on the table</td>
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<td></td>
<td>Mobile arm supports</td>
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<tr>
<td></td>
<td>A foot-operated feeding device</td>
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<td></td>
<td>Powered feeding devices</td>
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<td></td>
<td>Straws with a one-way valve</td>
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</table>

**Fear of choking**

Swallowing will get more difficult over time and there may be times when a person with dysphagia will have episodes of coughing and choking. An ineffective cough, resulting from weakened respiratory muscles, will make it more difficult to remove the irritant, whether it is food, drink or saliva. Read more about cough management in Information sheet P6 – *Evaluation and management of respiratory symptoms in MND* (see page 9 for how to order).

This can be very distressing for both the person with MND and those close to them, and it is common for people with MND to worry that they will die by choking. However, it is important to explain and reassure them that this is very rarely a direct cause of death in MND. Our information sheet 7A - *Swallowing difficulties* has guidance on managing choking. Download from [www.mndassociation.org/publications](http://www.mndassociation.org/publications) or see page 9 for details of how to order copies.
Other issues related to dysphagia

Laryngeal spasm
Uncontrolled/involuntary muscular contractions of the laryngeal (vocal) cords may be experienced due to an impaired swallow. This is often preceded by acid reflux. Laryngeal spasm can be extremely distressing as it can temporarily stop someone from breathing or speaking. The person affected should be encouraged to extend their neck backwards, take a slow breath in and a quick breath out. It can be accompanied by stridor: a loud, high-pitched respiratory sound.

A prescribing professional may consider lorazepam for laryngeal spasm.

Taking medication
Some people will have problems swallowing tablets, so speak to a pharmacist about alternative forms of any medications needed, for example syrups or patches. Many drugs can be taken via a gastrostomy tube.

Cognitive change
About half of all people with MND experience some degree of cognitive and/or behaviour change. This increases to 80% in the final stages of the disease. Additional support will be needed for people with dysphagia and cognitive impairment.
• Supervise the person’s eating more closely.
• People with more severe cognitive or behaviour changes may place too much food in their mouth at one time and cram food. Others may eat more food than they need, or they may have a preference for sweet food.
• Limit the amount of food on the plate at one time.
• Ask the carer to model eating at an appropriate pace.
• If food cravings are noticeable, question how much of a problem the behaviour is causing. It may be helpful to discuss with a dietitian.
• Those with poor swallowing may have trouble following medical advice to modify consistency or to thicken drinks.
• Refer to an SLT for assessment and advice about how to encourage safe eating, eg using the chin-tuck technique, counting to 10 when swallowing.
• Repeated reminders about swallow safety tips may be necessary.
• Ensure that mealtimes are protected from any distractions.

Alternative feeding methods
Alternative feeding methods should be discussed at an early stage, and revisited as oral intake becomes symptomatic. Introducing the option of a feeding tube early will give the patient time to make an informed decision.

Alternative feeding may be provided via a tube inserted through the nose into the stomach (nasogastric tube) or a stoma (hole) from the skin of the abdomen into the stomach, allowing a tube to be placed and feed passed through (gastrostomy).

Once the feeding system has been inserted, liquid feed may be delivered by:
• syringe
• gravity – a bag of liquid food is hung from a stand and allowed to drip through a tube
• pump method – a measured quantity is pumped into the tube, generally over some hours.

If it is safe to do so, the person may continue to take some food and liquid by mouth, for pleasure or quality of life, and use the feeding tube to supplement nutrition. The quantity fed through the tube may increase as swallowing becomes more difficult.
Types of enteral feeding
The following may be used to ‘top up’ oral intake, to meet full nutrition and hydration needs and to administer medications:\textsuperscript{13}

- Nasogastric tube (NGT)
- Percutaneous Endoscopic Gastrostomy (PEG)
- Radiologically Inserted Gastrostomy (RIG)
- Per-oral Image-guided Gastrostomy (PIG)

With RIG, the feeding tube is inserted under x-ray guidance. PIG is a hybrid of PEG and RIG but at present is not widely available.

Nasogastric tube
This is a tube inserted through the nose, allowing food to be passed directly into the stomach. It is usually a short term option which may be appropriate for someone who is malnourished or dehydrated, while waiting for gastrostomy placement, or where gastrostomy placement is not possible.

This method may be used in people for whom other types of gastrostomy are not appropriate, or may be preferred by some. As with other types of gastrostomy, this method can improve quality of life, but it is often considered less comfortable than gastrostomy tube placement.\textsuperscript{14}

Gastrostomy placement
PEG is the preferred method of gastrostomy, when someone has good respiratory function, or PIG/RIG when there is significant compromise of respiratory function.

The optimal timing of gastrostomy is not entirely clear, but is likely to be when someone has lost around 5\% of their body weight (from measurement taken at diagnosis).\textsuperscript{15} Earlier placement of a gastrostomy tube is recommended, even if it isn’t used straight away, as when needed, it can improve/maintain quality of life.\textsuperscript{15}

Possible risks of a late gastrostomy include, continued weight loss, respiratory complications, dehydration, failed insertion, and a higher risk of mortality and procedural complications.\textsuperscript{16} Therefore, the conversation about alternative feeding should happen early, before a crisis is reached. The discussion may be triggered by these indicators:

- aspiration of food
- anxiety when eating
- fear of choking
- inadequate food or fluid intake
- length of time and effort taken over meals
- weight loss or dehydration
- fatigue from eating.

Making the decision
It is important to recognise that, while early placement of a feeding tube is recommended, insertion of gastrostomy is an elective procedure which people with MND delay due to personal perceptions and concerns. It may not be an option where support to manage the tube is not available locally.

Discussions about the need for a tube can be distressing, especially soon after diagnosis, and should be handled sensitively. Referral to psychological support services may be helpful.

The pros and cons of this option should be discussed to enable the individual to make an informed choice. Understanding the factors that influence decision making about artificial feeding can help professionals to support people with MND to decide whether gastrostomy is for them.\textsuperscript{17}
Any discussion the person with MND has about gastrostomy should cover:

- how alternative feeding can affect quality of life
- any possible side effects, including discomfort or pain
- how gastrostomy may be included in an Advance Decision to Refuse Treatment (ADRT).

Where possible the person should also have the opportunity to meet with the palliative care team, and other people with MND who are using gastrostomy.

Not everyone with MND will choose this type of intervention and their decision should be respected. However, as problems progress, they may change their mind.

**Careful assessment is also needed of:**

- the level of support required to manage living with this intervention. The person with MND or their carer will need to understand what is involved and how often feeds should be administered, to ensure they can manage physically. If care support is required, arrangements need to be made at an early stage
- the understanding by the person with MND of the need for the procedure, so they can make an informed choice about whether or not to have the tube
- any possible physical restrictions imposed by the time required for feeds or trips to the toilet at night if the feed runs through overnight.

**Maintenance and potential problems**

Disadvantages of gastrostomy include having to keep the placement site very clean, the potential for infection at this site, and for the tube to be displaced. Practicalities around the maintenance of the tube (e.g., cleaning and flushing through) and administration of feed should also be considered.

People with MND and their carers need adequate information and training both before and after feeding tube insertion to help them make the decision, but also to help with the transition from oral to gastrostomy feeding. They should know where to go and who to contact in case of any clinical complications.

Possible problems that may arise with use of the tube include bloating and changes in bowel habits. A dietitian or MND specialist should be consulted about these issues.

**Refusing or withdrawing a feeding tube**

Someone with MND may make an Advance Decision to Refuse Treatment (ADRT) in which they refuse particular treatments. This could include instructions about initiating or withdrawing artificial nutrition and hydration by any means, including by feeding tube.

If someone decides against a gastrostomy or it is withdrawn, they may need symptom control for hunger and thirst. As the disease progresses, these desires usually decline naturally, but any discomfort may be minimised with the use of medication. Consult the specialist palliative care team.

**Cognitive change and decision making**

The potential for cognitive impairment highlights the importance of early planning. If someone experiences cognitive change, their ability to tolerate and understand the need for particular interventions may be affected. Advance care planning should include a person’s preferences in relation to alternative feeding.

The person should have the chance to discuss whether they would want to make an ADRT before they lose the ability to communicate or experience possible cognitive change. If this has not been done and the person is unable to make an informed decision, consult the person’s relatives and the specialist palliative care team if appropriate in your area.

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**Information you can share**

Information sheet 7B – *Tube feeding*. See page 9 for details of how to order copies, or download from [www.mndassociation.org/publications](http://www.mndassociation.org/publications)
References


8 British National Formulary. Feed thickeners and pre-thickened drinks. 2018; vol 75, p1478.


10 Crockford C et al. ALS-specific cognitive and behavior changes associated with advancing disease stage in ALS. Neurology. 2018; 91(15): e1370-e1380.


Further Reading

How the MND Association can support you and your team

We support health and social care professionals to provide the best possible care for people living with MND, their carers and families. We do this in a number of ways:

MND Connect

Our helpline offers help, information and support, and signposting to other services.

Telephone: 0808 802 6262

Email: mndconnect@mndassociation.org

Please contact MND Connect if you have any questions about the information in this publication.

MND Association website

Access information for health and social care professionals on our website at www.mndassociation.org/professionals
**Education**

Our education programme is designed to improve standards of care and quality of life for people living with and affected by MND. Opportunities include online modules and face-to-face training, such as conferences and masterclasses. Find out more at [www.mndassociation.org/education](http://www.mndassociation.org/education)

**Information resources**

We produce high quality information resources for health and social care professionals who work with people with MND. The range includes:

- Information sheet P3 – *Managing saliva problems in motor neurone disease*
- Information sheet P6 – *Evaluation and management of respiratory problems in MND*

We also have a wide range of resources for people living with and affected by MND, including:

- *Living with motor neurone disease* guide
- *Eating and drinking with motor neurone disease* – guide containing information, tips and easy swallow recipes
- Information sheet 7A – *Swallowing difficulties*
- Information sheet 7B – *Tube feeding*
- Information sheet 11B – *Mouth care*
- Information sheet 14A – *Advance Decision to Refuse Treatment (ADRT) and advance care planning*
- *End of life: a guide for people with motor neurone disease*

Downloads of all our information sheets and most of our publications are available from our website at [www.mndassociation.org/publications](http://www.mndassociation.org/publications) You can also order our publications directly from the MND Connect team.

**The MND Just In Case Kit**

The MND Just in Case Kit is a box designed to hold medication for the relief of choking, breathlessness or related anxiety. Its presence in the home provides tangible evidence for people with MND and carers that fears have been addressed and practical help is at hand. For the GP and district or community nurse it provides guidance on symptom management and medication and storage for the prescribed medications. A GP or other prescriber can request a kit for a person with MND from MND Connect. Call [0808 802 6262](tel:08088026262) or email [mndconnect@mndassociation.org](mailto:mndconnect@mndassociation.org) Find out more at [www.mndassociation.org/justincase](http://www.mndassociation.org/justincase)

**MND support grants and equipment loan**

Where statutory funding or provision has been explored and is not available, we may be able to provide a support grant or some equipment on loan.

Grants may be given to help with aspects of care or quality of life for people with MND, their carers, and younger members of the family.

Referrals for support grants or equipment loan need to be made by an occupational therapist, speech and language therapist or other relevant health or social care professional. For enquiries about MND support grants or equipment loan, call MND Connect on [0808 802 6262](tel:08088026262), email [mndconnect@mndassociation.org](mailto:mndconnect@mndassociation.org) or visit [www.mndassociation.org/getting-support](http://www.mndassociation.org/getting-support)

**Communication aids service**

This service helps people affected by MND and health and social care professionals with queries about communication aids. The service provides limited financial support for communication aids or some items on loan, if unavailable or delayed through health and social care services. Our aim is to improve provision and information on a local and national level, through collaboration with professionals. Call [0808 802 6262](tel:08088026262) or email [communicationaids@mndassociation.org](mailto:communicationaids@mndassociation.org)
Research into MND
We fund and promote research that leads to new understanding and treatment and brings us closer to a cure. Contact the Research Development team on 01604 611880 or email research@mndassociation.org Alternatively, visit www.mndassociation.org/research for more information.
For the latest research news, visit our research blog at www.mndresearch.blog
Our peer-to-peer research and care community blog (ReCCoB) has a number of contributors who report on MND-related workshops and events at www.reccob.wordpress.com

International Symposium on ALS/MND
Each year we organise the world's largest clinical and biomedical research conference on MND. It is the premier event in the MND research calendar for discussion on the latest advances in research and clinical management. Find out more at www.mndassociation.org/symposium

MND Association membership
Join us now and help fight MND. Membership costs £12 for individuals. It is free for people with MND and for carers, spouses or partners of people living with MND. Call 01604 611855 or email membership@mndassociation.org

Local support
Regional care development advisers
We have a network of regional care development advisers (RCDAs) covering England, Wales and Northern Ireland. RCDAs have specialist knowledge of the care and management of MND. They work closely with local statutory services and community care providers to ensure effective support for people affected by MND. RCDAs provide education for health and social care professionals in MND, and are champions at influencing care services in their respective areas.

MND care centres and networks
We help fund and develop care centres and networks across England, Wales, and Northern Ireland, which offer specialist clinical expertise from diagnosis onwards.

Branches and groups
We have volunteer-led branches and groups nationwide providing local support and practical help to people with MND and their carers.

Association visitors (AVs)
Association visitors are volunteers who provide one-to-one local support to people affected by MND.

We welcome your views
Your feedback is really important to us, as it helps improve our information for the benefit of people living with MND and those who care for them. If you would like to provide feedback on any of our publications, you can access an online form at:
https://www.smartsurvey.co.uk/s/professionalinformation
You can request a paper version of the form or provide direct feedback by email at infofeedback@mndassociation.org
Or write to: Information feedback, MND Association, PO Box 246, Northampton NN1 2PR
If you would like to help us by reviewing future versions of this or other resources, please email us at infofeedback@mndassociation.org
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Kiri Elliott, Policy Officer (Professional Development), The British Dietetic Association

About us

The MND Association was founded in 1979 by a group of volunteers with experience of living with or caring for someone with MND. Since then, we have grown significantly, with an ever-increasing community of volunteers, supporters and staff, all sharing the same goal – to support people with MND and everyone who cares for them, both now and in the future.

We are the only national charity in England, Wales and Northern Ireland focused on MND care, research and campaigning.

Our mission

We improve care and support for people with MND, their families and carers.

We fund and promote research that leads to new understanding and treatments, and brings us closer to a cure for MND.

We campaign and raise awareness so the needs of people with MND and everyone who cares for them are recognised and addressed by wider society.

About MND

• MND is a fatal, rapidly progressing disease that affects the brain and spinal cord.

• It attacks the nerves that control movement so muscles no longer work. MND does not usually affect the senses such as sight, sound and feeling.

• It can leave people locked in a failing body, unable to move, talk and eventually breathe.

• It affects people from all communities.

• Some people may experience changes in thinking and behaviour, with a proportion experiencing a rare form of dementia.

• MND kills a third of people within a year and more than half within two years of diagnosis.

• A person's lifetime risk of developing MND is up to 1 in 300.

• Six people per day are diagnosed with MND in the UK.

• MND kills six people per day in the UK.

• It has no cure.

MND Association

PO Box 246, Northampton NN1 2PR
Tel: 01604 250505
Email: enquiries@mndassociation.org
www.mndassociation.org

@mndeducation
@mndassociation