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Disclaimer Statement: The information in this guide is not medical advice. Talk to your ALS care team before making any decisions about your health or treatment. Together, you and your care team can find a treatment plan that works for you.

Last Reviewed: October 15, 2021
Good nutrition and hydration are very important for people living with ALS. People living with ALS burn more calories because of the increased energy used in breathing, moving and muscle twitching. You may have challenges with low energy, maintaining weight and decreasing muscle mass. Proactively addressing your challenges can result in greater energy, strength and independence.

If you are having trouble eating, drinking, or taking medicine on your own, you may not be able to get all of the nutrition your body needs. This can lead to weight loss, infections, and other health problems down the road. This guide walks you and/or your caregivers through proactive strategies to maintain weight.
Maintain weight

You may benefit from a proactive nutritional approach to any of these symptoms:

- **Having difficulty chewing and swallowing** (coughing or choking)
- **Losing weight** without meaning to despite eating a "normal" amount of food
- **Eating less food** than you usually would or having a poor appetite
- **Taking longer** than expected to eat a meal because of arm or hand weakness or chewing and swallowing problems
- **Having problems from not getting enough water** (or dehydration) This can include constipation or thick secretions

Even if you’re not having these symptoms, it’s never too early to be proactive with ensuring a healthy diet and adequate hydration.

If you ignore weight loss it may be hard to gain weight back. Severe weight loss contributes to muscle loss, decrease in independence and energy.
Calories

You are burning more calories at rest when compared to people who are not living with ALS

This makes maintaining weight more difficult, so people with ALS need to increase their calories to maintain weight. The challenge is to increase the number of calories without significantly increasing the amount of food.

A few ways to increase calories would be:

- **Eat small frequent meals during the day every 2 to 3 hours.**
- **Healthy fats have a lot of calories without much volume.** Foods like nuts, nut butters, seeds, avocados, olives, and oils (olive or canola) are high in healthy calories.
- **Avoid diet foods and choose the full-fat version for more calories.**
- **Add high protein foods such as cheese, eggs or powdered milk to casseroles and soups to increase calories and protein.**
- **Drizzle olive oil over foods such as vegetables, meat and soup.**
- **Look for nutritional supplements with the word “plus”, rather than the “high-protein” varieties.** These supplements contain more calories and protein.

Talking with a registered dietitian or nutrition specialist can help guide you in selecting the right foods and supplements.
Eating

Chewing and swallowing difficulties (dysphagia) may make mealtimes more challenging or tiring for you. Concentrating and eating more slowly may cause a sense of isolation due to others having already finished eating. As a result, you may not finish your meal leading to a decrease in caloric intake. Some people listen to the radio or podcasts to lessen their isolation.

Signs of swallowing difficulty include:

- Choking or coughing while eating
- Increased saliva or excessive drooling
- Sensation of food getting stuck in your mouth, throat, or chest
- Needing more time to finish a meal
- Frustration during meals
- Avoiding certain foods
- Loss of appetite
- Increased nasal secretions

Dietary Supplements
ALSUntangled reviews alternative and off label treatments, with the goal of helping people with ALS make more informed decisions about them.

alsuntangled.com
Try to determine what types of foods and liquids are the easiest to tolerate. You might want to consider a soft diet of foods that are easy to chew and swallow.

How to prepare soft foods?

- Cut food into small pieces that are 1/2 inch or smaller because they are easier to swallow.
- Use chicken broth, beef broth, vegetable broth, gravy or sauces to cook or moisten meats and vegetables.
- Cook vegetables until they are soft enough to mash with a fork.
- Use a food processor to grind or purée foods to make them easier to chew and swallow.

Drinking and dehydration

Adequate fluid intake is essential for avoiding health problems.

What are some of the signs of dehydration?

- Dark colored urine
- Dry itchy skin
- Confusion
- Dizziness or Lightheadedness
- Headaches
- Flushing/Fever
- Increased fatigue
- Decreased urine output
Drinking thin liquids might be difficult causing you not to drink enough fluids. If thin liquids like water cause coughing, drink thicker liquids such as:

- Milkshakes, smoothies, or drinkable yogurt
- Puréed soups
- Carbonated fluids
- Thicker fruit juice with pulp
- Use thickening powders or gels, such as Resource Thicken Up Clear, Thick-It, or Simply Thick, to thicken fluids without altering the taste
- Other thickening options for thin liquids, which also help increase calorie intake, include: baby rice cereal, mashed potato flakes, puréed baby foods

You might also be dehydrated because you want to avoid increased trips to the restroom, which can require assistance. Talk with your ALS care team about different bathroom aids and strategies.

Ensure proper fluid intake with a goal of six to ten 8-ounce cups of liquid a day.

Alcohol and caffeinated beverages can be dehydrating (although can be consumed with adequate hydration).

Consider the use of a sports drink with electrolytes or other flavored beverages to increase fluid intake.

Certain foods such as Jello, sherbet, and fruit will also help to increase fluid intake.
## Aids for eating and drinking

<table>
<thead>
<tr>
<th>Aids</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Built-up handles</td>
<td>Use foam tubing or wash-cloths on lightweight eating utensils for decreased grip</td>
</tr>
<tr>
<td>Universal cuff</td>
<td></td>
</tr>
<tr>
<td>Large handled cup for hot and cold liquids</td>
<td>This allows four fingers to fit through the handle to increase stability.</td>
</tr>
<tr>
<td>Offset spoon or fork that can be angled right or left</td>
<td></td>
</tr>
<tr>
<td>Clear plastic, clip-on plate-guard</td>
<td></td>
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<tr>
<td>A non-slip disc; or Dycem, a gel pad used to stabilize plates and cups on the table</td>
<td></td>
</tr>
<tr>
<td>Long rigid or flexible straws</td>
<td></td>
</tr>
<tr>
<td>Straw holder clip</td>
<td></td>
</tr>
<tr>
<td>Small pliers</td>
<td>to open water bottles and jar openers</td>
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<tr>
<td>Sports drink container with a straw</td>
<td></td>
</tr>
<tr>
<td>Nosey cup</td>
<td>The special cut out provides space for the nose, allowing you to tilt the cup for drinking without bending the neck or tilting the head</td>
</tr>
<tr>
<td>Flexi-Mug and cupholder</td>
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Choking
As face, throat and jaw muscles weaken, the risk of food and/or liquid going down the wrong way increases.

You can reduce choking episodes in several ways:

- **Limit talking and other distractions while eating.**
- **Limit foods with different textures like chicken noodle soup or dry cereal with milk.**
- **Serve smaller, but more frequent meals, avoiding dry, crumbly or large chunky foods.**
- **Take smaller bites of food and little sips of liquid. Swallow several times to clear each bite.**
- **Keep mucus and saliva thin by ensuring adequate fluid intake.** (Try papaya, pineapple or purple grape juice. These juices each contain enzymes which helps to breakdown protein in mucus. You can also add lemon juice to water.)
- **Ask to talk with a speech pathologist or swallowing specialist for specific strategies that might make swallowing easier for you.**
- **Consider getting a feeding tube to reduce your risk of choking.**

Food going down “the wrong way” means breathing food or liquid into the lungs (aspiration) instead of swallowing down the esophagus into the stomach.

Aspiration can cause respiratory infections or a frightening choking spell and is a leading cause of pneumonia, a potential life-threatening event for people living with ALS.
Feeding tubes

How does a feeding tube work?
If you cannot maintain weight by eating, or swallowing becomes too exhausting, time-consuming or dangerous, a feeding tube may be considered. A feeding tube is a type of medical device that can help you get the nutrition you need. You will need to have a procedure for a doctor to place the tube inside your stomach. The other end will be on the outside of your skin but is easy to cover up under clothes.

Feeding tubes can lessen the stress you feel when you are unable to eat enough to maintain your weight. It may be a much better way to maintain nutrition and hydration than trying to eat or drink everything by mouth.

A feeding tube allows you to eat what you are able by mouth and supplement with adequate calories through the tube. Vitamins and medications can also be easily given through the tube when crushed.

What are the benefits of using a feeding tube?
Getting a feeding tube early can improve your quality of life and help you live longer.

Using a feeding tube can help you:

- Lower your risk of choking or pneumonia (a serious type of lung infection)
- Have more energy throughout the day
- Participate in activities you enjoy, like spending time with family and friends
- Worry less about whether you’re eating and drinking enough
- Maintain your strength and keep you from losing too much weight
- Make sure you’re getting the adequate your body needs

Considering a feeding tube

ALS is a progressive disease. This means your ALS symptoms will get worse over time.

If you choose not to use a feeding tube, it may become harder for you to swallow safely and you may be at an increased risk of choking.
Eating and drinking with a feeding tube
Some people can eat and drink in addition to using a feeding tube. Other people get all their nutrition through the feeding tube. This may also change over time as your ALS symptoms progress.

How will the doctor insert my feeding tube?
You will need to have a procedure to get a feeding tube. This procedure will make a few small cuts to place the feeding tube inside your stomach.

There are two different types of procedures. The difference between them is how the doctor looks inside your stomach:

**Percutaneous Radiological Insertion of Gastrostomy (RIG):** A special doctor called a radiologist will take an X-ray of your stomach. They carefully guide instruments through tiny incisions in the skin to make an opening in the stomach. You may have a Nasogastric Tube placed temporarily the day before the procedures. This is a flexible tube of rubber or plastic that is passed through the nose, down through the esophagus, and into the stomach to ensure accurate feeding tube placement.

**Percutaneous Endoscopic Gastrostomy (PEG):** A gastroenterologist, or a doctor who specializes in the digestive tract, will use a camera called an endoscope to take pictures of your stomach. The doctor will place the endoscope down your throat so they can see inside your stomach.

Most ALS doctors recommend the RIG procedure. Talk with your ALS care team about which type of procedure is best for you. There are risks associated with the feeding tube procedure—like infections, breathing problems and reactions to medicines you get during surgery.

Your ALS care team can go over all the risks with you and provide you with more details about the procedure.

You can use My ALS Decision Tool™ to learn more about using a feeding tube. Then, talk with your ALS care team. Together, you and your care team can find a treatment plan that works for you.

ALSDecisions.org/Nutrition
How do I care for a feeding tube?
While you are in the hospital after the procedure, your ALS care team and hospital staff will show you how to take care of your feeding tube. Your ALS care team will give you instructions to follow when you go home.

For a few weeks after the procedure, the feeding tube requires special attention to prevent infection. After the site is healed, daily cleansing with soap and water is all that is necessary.

It is recommended that the feeding tube is exchanged every 6-9 months.

Don’t worry—after the first procedure to place the tube in your stomach, it’s much easier to replace with a new one. You will not need to stay overnight in the hospital.
Learn more

The Les Turner ALS Foundation exists to guide you to answers, support you and your loved ones and advance scientific research. To learn more about living with ALS visit, lesturnerals.org/resources.

My ALS Decision Tool™
If you have ALS, you will need to make some important decisions about your health care. As your disease progresses, your ALS care team may recommend different care options. You can use this tool to learn about some common ALS treatments, answer a few questions to help you think through what is most important to you and get ready to talk with your ALS care team about your options. To learn more, visit: alsdecisions.org.

ALS Learning Series
Our online ALS Learning Series aims to empower the ALS community through the latest information and insights. Educational webinars and interactive Q&A’s covering a diverse array of topics, from nutrition to respiratory care, are offered monthly featuring members of the Foundation’s Support Services team, our Lois Insolia ALS Clinic at Northwestern Medicine and other national ALS experts. To learn more about ALS care and research, visit: alslearningseries.org.

My ALS Communication Passport to Quality Care
My ALS Communication Passport to Quality Care was created to make your life easier. You will be able to share health information and care preferences with caregivers. You have a lot of information to keep track of, and this tool will help you do that. To find out more, visit: lesturnerals.org/passport.

Support Groups
We facilitate support groups to provide people living with ALS and their caregivers and family the opportunity to share their experiences, give encouragement and help each other navigate their journey with ALS. To find out more, visit: lesturnerals.org/support-groups.