





# A Quick Guide to CF in High School

CF is a genetic disease that causes thick, sticky mucus in the body, mainly affecting the lungs and digestive system. There is no cure for CF and daily treatment is needed to keep healthy. With a little bit of planning and good communication, CF can be successfully managed at school. A Student Medical Plan detailing any CF care required at school is recommended.

## **Symptoms**

CF affects various organs and symptoms will vary from one person to another. It is important to know what's normal for the student in your care so you can discuss any change in symptoms with them and their parents.

The most common respiratory symptom is a cough, which is not contagious to others. People with CF need to cough to clear the mucus from their lungs. Coughing should not be discouraged, however students may choose to discreetly leave the classroom to avoid embarrassment.

Many children with CF will also have digestive symptoms, including difficulty gaining weight, abdominal pain, bloating, constipation or diarrhoea as well as strong smelling wind.

### **Nutrition**

A healthy diet is different for everyone. Recess and lunch for a student with CF might contain more high fat, high calorie foods because of digestive issues as well as the extra energy their body uses to breathe, cough and fight infection in the lungs.

A healthy weight is important for maintaining healthy lungs



# **Digestive Enzymes**

Most people with CF require enzymes before they eat, to help digest food as their pancreas doesn't work properly. Enzymes are not harmful to other students.

It is important to develop a plan with the student and their parents to ensure enzymes are taken each day. This should be managed independently in high school. If enzymes are skipped, the student can experience abdominal pain and diarrhoea.

## **Dehydration**

People with CF are at a greater risk of dehydration due to increased salt loss in sweat. Dehydration can make mucus in the airways even stickier and more difficult to clear.

It is important to know the signs of dehydration such as lethargy, irritability and salt crystals on the skin.







You can help prevent and manage dehydration by:

- · Allowing easy access to water at all times
- · Encouraging students to drink during sport
- Allowing extra salt or electrolytes to be taken as needed

## **Infection Control**

Due to thick sticky mucus in the lungs, colds, flus and other infectious diseases can be more serious for people with CF. Good school cleaning practices, encouraging good hand hygiene and cough and sneeze etiquette in the classroom are the best ways to minimise the spread of germs. Students who are unwell should not attend school until their symptoms have cleared.

#### **Environmental Risks**

There are also infection risks in the environment for people with CF, such as still or stagnant water, damp soil, hay and mulch. Bacteria found in these environments can cause damage to the lungs of a person with CF. Discuss any precautions with the student and their parents.

#### **Cross Infection**

People with CF should avoid close or prolonged contact with each other to avoid sharing infections. It is recommended that they maintain a distance of 4m apart, particularly in small, enclosed spaces. If you have more than one individual with CF attending your school, including staff and parents, please ensure strategies are put in place to keep everyone safe. Please consider the confidentiality of individuals when having these discussions. People with CF should not be in the same classroom and should avoid close contact at whole school events, incursions/excursions and on buses.

These recommendations may also apply to other people in the school community with chronic lung conditions or impaired immunity, however, does not include people from the same household with CF for example, siblings.

#### Other Considerations

Children with CF have the same capacity to learn and you should have the same expectations of their behaviour as other students in your class. They can participate in all school activities including camps and excursions, however some considerations may need to be discussed including environmental risks and ensuring treatment can continue while they are away.

Each person with CF will have their own unique challenges and may have additional issues that need to be managed at school including other CF related conditions or more complex care needs. These additional challenges can all be successfully managed with good communication and some extra support where needed.

CF treatment has come a long way, and is continuing to advance, enabling people with CF to live longer, healthier lives. Learning how to manage CF at school is an important part of helping children with CF develop independence in a supportive environment.

## More Information

For more information about CF, we have free eLearning modules and other resources for educators, available on our CFSmart website.



