FACT SHEET

COMMUNICATING WITH PRIMARY AGED CHILDREN ABOUT GENETIC CANCER RISK

Most parents wonder about how and when they should talk to their child about cancer genetic risk. There may be many reasons why it is appropriate to begin to help your child understand more about genetic cancer risk. Reasons include; your child may be required to have a genetic test, your child may be recommended to have procedures to screen, treat or reduce the risk of cancer, your child may be witnessing you or other family members having procedures or surgery to manage cancer risk or treat cancer.

Why is it important to talk to your child?

Conversations with your child are important. These conversations can help your child to understand the family situation and to make sense of what they are experiencing. When your child does not understand what is happening in the family they can worry, feel alone and/or fearing the worst, which is very common in late primary and early teens, and misunderstand the situation. They may feel personally responsible and worry about you and your health and safety. Helping your child to understand cancer genetic risk and what it means for them and your family will:

- help your child to know that it is okay to talk about it.
- allow your child to ask questions and get the correct information.
- help them come to you (or others) when they are worried or feeling overwhelmed.
- build a shared understanding that can strengthen relationships.

This resource will help you to prepare for conversations with your child about genetic cancer risk so you can help them to understand what it means. It steps you through:

- how your child might understand health and illness
- thinking about how you feel about your genetic cancer risk
- thinking about what your child notices and experiences
- preparing to talk with your child

How do children understand health and illness?

Thinking further about what children understand about health and illness can help you in thinking about how to approach a discussion with your child.

Children hold complex concepts of health and illness and these concepts change as they develop. Much of how children understand these concepts comes from their personal experiences of illness and learnings about health promotion. Younger children's concepts are often different to older children and adults who use more abstract experiences of health to inform their understanding.

Children view the general concept of being "healthy" as being able to participate in activities, maintaining their normal routine, not taking medications or feeling sick and being happy.

As children less commonly experience serious or chronic disease, most children's personal experiences of illness come from their experiences of contagious diseases such as colds and viral illnesses. This forms a 'causal and effect' concept of illness e.g. that one event leads to another.

Children's experience of health promotion also informs their understanding of illness. Children understand the importance of behaviours such maintaining personal hygiene, eating well and not being around passive smoke and how these exposures relate to illness. Because of this they link healthy behaviours to the concept of health.

In essence children's understanding is complex but the conclusions they draw may appear simplistic. This can be of relevance when children come across a new illness scenario. Researchers have found that children may often 'fill in the gaps' and find ways for their experience to fit within their known illness construct. Examples of this might be that children report that their illness is a punishment for something they have done or that things like their natural environment such as 'cold air' or 'pollution' may have caused their illness when the reason is not able to be identified.

It may be helpful to think about how you can describe the impact of genetic risk information in ways that can fit into the constructs your children hold.

For example, if you are considering genetic testing for your child it may be explained in ways such as;

"We have lots of genes in our body, these make up who we are and what we look like, for example what colour eyes we have. Genes are not something we can change, we get them when we are born. They are not like germs that give us a cold. Sometimes we get genes that mean we have to do extra things to help us stay healthy. We can do a test on our genes, to see if we need to have extra check ups or speak with other doctors, to make sure we stay as healthy as possible."



COMMUNICATING WITH PRIMARY AGED CHILDREN ABOUT GENETIC CANCER RISK cont...

How do you feel about your genetic cancer risk?

For primary school aged children parents are often the most important people in their life and they look to their parents for guidance in understanding the world around them. Because of this, children of this age are often very sensitive and attuned to their parents' emotions and behaviours (even when parents try to hide them). How parents behave and the emotions they experience often inform how children feel and understand new information and situations.

Thinking about how you feel about cancer genetic risk and how you feel about your child's situation is important in how your child may understand and adjust to any information you share.

Asking yourself the following questions can assist in helping you understanding how you feel about genetic cancer risk and the idea of sharing information with your child;

- What do you feel when you think about your cancer risks?
- What do you feel when you think about your children being at risk of a genetic cancer risk?
- What does your child's other parent about cancer genetic risk?
- What do you feel when you think about having conversations with your child/children about genetic cancer risk?
- What do you understand about how the genetic cancer risk may affect your child?

TIP -Whilst it is normal to feel sad, anxious, guilty or worried it may be helpful to talk with a health professional and/or peers or friends to work through these emotions and feelings before talking to your children.

Understanding what your child notices and experiences is also important, the questions below may help you in thinking about this further;

- What may they have noticed or experienced that has changed since I learnt of the genetic cancer risk?
- what have you noticed about their reactions?
- what do you think they might be feeling?
- what might they understand or not understand?
- What do you think might worry your child the most?

Practical tips in preparing for conversations

Children best understand information over time. As your child grows, their need for information will also change, so be prepared to have multiple conversations.

When and where to start the conversation/s?

Timing and location is important and it may be helpful for you to identify what a good time may be;

- is your child have a difficult time at school or with social groups? (would delaying a short time be possible?)
- what time of day is your child best able to focus and concentrate on what you are saying?
- Where do you have the best conversations with your child, when they are in the bath, at the breakfast or dinner table, at bed time, in the car?
- who else will be around when you plan to share the information? (if you have more than one child you may need to consider if different approaches or times may be best)
- Who else may be able to help support you in sharing information and/or keeping an eye on how your child may respond e.g. other relatives, friends or school teachers.

Tips for talking to your child?

- Think about what words or language they will understand
- Stop and pause after each new bit of information. Sometimes the
 conversation may be very short don't worry if this happens they
 small conversations are just as important as big ones, they all add
 up and help children learn at their pace.
- Children will be often worried about if it will impact on your ability to take care of their needs and if on their health they may only understand concepts such as will I get better and when.
- Give your child time to think and to ask questions (the questions might not come straight away, your child may need thinking time).
- Know what to do if you don't know the answer. E.g. tell them that you will find out, or even find out the information together and follow through.
- Encourage your child to ask questions or raise concerns whenever they want and make sure when they do, answer them or make time to answer them as soon as possible so they feel valued and trust that you aren't avoiding the topic. Often children may ask questions when you least expect them to!
- Watch your child's behaviour or for signs such unexplained tummy aches, wanting to stay home from school as this may be a sign you need to check in with them about whether there is anything worrying them.
- Set up a support network for your child so that your child also can seek answers from a person that you both trust (e.g. a family member, a family friend or a health professional).

