Adulthood is seen as a time of growth and independence. Parents take great pleasure in watching their children take control of their own lives and begin to plan for a new stage in their own future. Acquired Brain Injury (ABI) can turn this all upside down. An adult child may lose their ability to be functionally independent and/or they may experience changes in their personality and behaviour. Parents can be called upon to step into a role of carer and/or supervisor, giving up their own plans and future. It is not surprising that in this time of upheaval and change that feelings and tensions run high. This fact sheet illustrates some possible reactions that parents may experience, how relationships and roles may change, practical consequences of ABI and useful strategies.

My Adult Child has an ABI

Introduction

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Possible Reactions

In coming to terms with an adult child’s ABI a parent can experience a range of emotions. These could include:

- Shock and disbelief – at the immediacy and severity of the injury and its consequences.
- Anger – at the circumstances of the injury, at how it has impacted on your life and how it will change your child’s life.
- Grief and loss – at the loss of a ‘normal’ future, the loss of hopes and dreams (both your child’s and your own) and the loss of your child as you imagined them in the future.
- Guilt – for the feeling that you could have “done something better” and for being angry, resentful or frustrated at the changes.
- Loneliness and isolation – as your life takes different directions with different priorities from that of your friends.
- Recurrent grief – grief reoccurs as your child remains in your life but as an altered person. You are periodically reminded of personality and characteristics that have been lost while learning to relate to a different person.
Relationships and Role Changes

An ABI can affect relationships and change family roles.

Often the greatest change is the need for parents to take on a carer role with an adult child who requires basic personal care and assistance with daily living tasks.

This can be like parenting a young child again and the relationship is no longer an adult relationship.

This can be especially difficult if the person’s ABI has also resulted in challenging behavioural problems.

Changes in relationships with an adult child can have many flow on effects, including:

- Worry about relationships with your other children e.g. being overprotective of your other children.
- Diminished relationships with friends and extended family as caring duties take time and energy and your interests shift to those of a carer.
- Strain in your relationship as a couple as you struggle to cope, have less time for each other or maybe have differing views about how care should be provided.
- Acting as a mediator/support in your child and their partners relationship as it changes.
- Changes in previously valued roles e.g. giving up work to be a carer, loss of retirement plans.
- Providing care to grandchildren.
- Providing support to your child’s partner or spouse as they come to terms with changes.
- Educating and supporting other family members and friends about brain injury.
- Allowing external carers or lifestyle support workers to come into the home making lives less private.

Practical Consequences of ABI

There may be many practical consequences of the brain injury. You may find that you need to be involved in provision of:

Accommodation

- Providing accommodation or sourcing alternative accommodation.

Care

- Providing or sourcing ‘hands on’ assistance e.g. feeding, bathing, supervision etc.
- Asking other family members or community services for help.
- Providing transport to appointments and community activities.

Maintaining connections

- Helping maintain pre-existing relationships.
- Providing social support and social outings.

Finances

- Assisting in managing finances and making decisions including applying for Centrelink benefits.
- Assisting in managing the effects of the loss of income when the injured person was the primary income earner e.g. mortgage repayments, school-fees.

Behaviour

- Managing behaviour changes resulting from the brain injury (professional assistance may be required in some cases).
Useful Strategies

Some strategies to help coping are:

- Becoming involved in the hospitalisation and rehabilitation stages.
- Learning as much as possible about the brain injury and its impacts on physical, mental and emotional function.
- Thinking about family members strengths and weaknesses—where can they help and where will they need help.
- Identifying areas where assistance would be beneficial and asking for help.
- Talking about concerns with other family members/friends.
- Keeping daily routines as normal as possible.
- Being open to involvement in support groups and counselling.
- Spending quality time with other family members to maintain relationships.
- Developing and maintaining a good network of friends and activities to avoid becoming isolated.
- Encouraging the person with ABI to gain as much independence as possible by supporting them to learn new skills and to be as independent as possible.
- Pay attention to and recognise the things (personality traits, behaviour traits and interests) that haven’t changed.
- Pay attention to and recognise the things the person with the ABI can do.
- Using supports such as regular respite when needed.
- Take the time to rest, rejuvenate, and care for yourself.

Further Information can also be obtained from the Synapse website http://synapse.org.au/information-services.aspx?category=Carers+%26%26+family+members

Resources


Notes:

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For Review Sept 2018: ABIOS Social Worker