Appendix A8: Information and resources for clinicians after a diagnostic assessment

The following are some approaches to providing support and intervention for an individual and/or their parents or caregivers after a diagnosis of FASD has been made. They are relevant for children, adolescents and adults. These approaches are also important to address patient needs even when the diagnostic assessment is inconclusive or FASD has been excluded.

1. Explain the diagnosis:
   Using a non-judgemental approach that recognises the range of emotions that might be experienced by individuals, parents or caregivers when a diagnosis of FASD is given, explain that making a diagnosis can:
   - Improve understanding of FASD.
   - Improve understanding of the individual’s difficulties while also identifying their strengths and help parents and carers adjust their expectations and provide support accordingly.
   - Provide opportunities for individuals, parents and caregivers to express and/or process a possible range of emotions.
   - Facilitate early intervention to improve a child’s development.
   - Identify individuals and/or their family members who are in need of assistance e.g. referral to alcohol and other drug services.

2. Provide individuals, parents and caregivers with:
   - The reports of assessments from health professionals.
   - The outcomes of the assessments, e.g. diagnoses; provisional diagnoses; need for further assessment.
   - The details and implications of a FASD diagnosis (or non-diagnosis).
   - Some ‘plain English’ information about FASD and contact details for NOFASD Australia (Printable information on pages 63 and 64), and/or contact details for RFFADA.
   - A contact number for a clinician who can respond to any questions that arise following diagnosis about the assessment and/or management plan.

3. Develop a management plan with individuals and/or their parents and caregivers so they:
   - Can identify their priorities and goals for inclusion in the management plan.
   - Are aware of therapy options and family support mechanisms available as appropriate interventions put in place.
   - Are empowered during future assessments, management and support.
   - Are aware of accessible parent, caregiver, family and personal networks in their community.
   - Are aware of support organisations.
Are aware of the need for referrals and further medical review and of potential waiting times for services.

Receive a copy of the management plan.

4. Consider support and interventions:

Build therapeutic interventions around the individual’s:

- Strengths, interests and positive attitudes.
- Willingness to participate in family, school or institutional activities and routines.
- Engagement with their family, peers and/or caregivers.

Key approaches include:

- Educating individuals, parents and caregivers about FASD and related impairments.
- Improving parent, caregiver and teacher understanding of interactions with the child, adolescent or adult living with FASD.
- Ensuring appropriate educational support and accommodations are implemented.
- Targeting therapy programs towards supporting the individual’s key functional difficulties.
- Medication (when indicated and appropriate).
- Advocating for the individual e.g. in education, child protection or justice systems.

Challenges to address may include:

- Challenges of daily life – e.g. caregiver fatigue, the need for routine and repetition for many individuals living with FASD, emotional or behavioural problems including aggression.
- Family’s need to access multiple health services, potentially with limited communication between different service providers.
- Service providers with limited knowledge about FASD.
- Need for individuals, parents and caregivers living with FASD to educate teachers, health and other professionals about FASD.
- Lack of recognition of a FASD diagnosis as a disability, providing a hurdle to obtaining funding for educational and other assistance.
- Lack of recognition of co-existing mental, developmental or physical health conditions.

Eight Magic Keys

- These eight strategies underpin successful strategic interventions for students with FASD and are one example of an approach that can be taken.
- They are simple, functional strategies to use with young people with FASD and can be used by caregivers, teachers and health professionals.
- They were developed for use by the FASD Centre for Excellence, Substance Abuse and Mental Health Services Administration [http://come-over.to/FAS/brochures/EightMagicKeysBroch.pdf](http://come-over.to/FAS/brochures/EightMagicKeysBroch.pdf).
1. Concrete Terms

- Children living with FASD do well when parents/carers and educators talk in concrete terms. Refrain from using words with double meanings, idioms etc. The social emotional understanding of children living with FASD is often below their chronological age, therefore it helps to ‘think younger’ when providing assistance, giving instructions etc. It is also important not to make deficit judgements.

2. Consistency

- Due to the difficulty that children with FASD experience in generalising learning from one situation to another, they do best in an environment with few changes. This includes consistency in language and routines. Educators and parents/carers should coordinate with each other to use the same words and/or gestures for key phrases. Communication books are effective ways of sharing what’s happening and advising on language use and behaviours in classrooms and homes.

3. Repetition

- Children with FASD have chronic short term memory problems. They forget things they want to remember, as well as information that has to be learned and retained for a period of time. In order for them to commit something to long-term memory, it often needs to be repetitively retaught.

4. Routine

- Stable routines and consistent visual cues that do not change from day to day make it easier for children with FASD to know what to expect next, and decrease their anxiety, enabling them to learn.

5. Simplicity

- Remember to keep input short and sweet. Children with FASD are easily over-stimulated, leading to ‘shutdown’, at which point they can take in no more information. Breakdown tasks and always communicate the task in the positive: “we walk inside” instead of “don’t run”.

6. Specific Language

- Say exactly what you mean. Remember that children with FASD have difficulty with abstractions, generalisations and ‘filling in the blanks’ when given an instruction. Tell them step-by-step what to do. This will help them develop appropriate habit-forming patterns. Keep instructions concise and broken into achievable chunks.

7. Structure

- Structure is the ‘glue’ that enables a child with FASD to make sense of the world. If this glue is taken away things fall apart. A child with FASD achieves and is successful because his or her world provides appropriate structure as a permanent foundation for learning.

8. Supervision

- Due to their cognitive challenges, children with FASD bring a naivety to daily life situations. They need constant supervision, as with much younger children, to develop habit patterns of appropriate behaviour and ensure safety and wellbeing at all times.
Fetal Alcohol Spectrum Disorder (FASD) is a condition that may be diagnosed in a person who, before they were born, was exposed to alcohol. The alcohol in any alcoholic drink (beer, wine or spirits) is rapidly absorbed into the mother’s blood stream and crosses the placenta to the unborn child to change otherwise healthy development. FASD is characterised by damage to the developing brain, leading to abnormalities in how the brain works. This can show up in several different ways, such as problems with learning, memory, language, judgement, decision-making and planning, movement or sensation. Some, but not all individuals can also have facial features that are characteristic of FASD.

Alcohol can cause harm to the unborn child at any time during pregnancy (including before pregnancy is confirmed) and the level of harm depends on the pattern of the mother’s alcohol use - the percentage of alcohol in drinks, the number of drinks, and over what time the alcohol drinks were consumed. Binge drinking for example, means a high level of alcohol is consumed in a shorter period of time.

In addition to the alcohol exposure, the vulnerability of a pregnancy and an unborn child may also be affected by other factors like genetics, family alcohol use across generations, the father’s alcohol use prior to conception, the mother’s age and general health (for example, nutrition, tobacco use) and other environmental factors like stress (exposure to violence, living with poverty, factors at work).

FASD is not always obvious at birth and might not be noticed until the child doesn’t reach developmental milestones or behaviour and learning difficulties become a worry once the child starts school. FASD can also be first diagnosed in adolescence or adulthood. Different professionals might need to be involved to assess the areas of the child’s life where help is most needed.

A person who was exposed to alcohol before they were born might now be any age. A proper diagnosis, appropriate services and support can help any person living with FASD to prevent behaviour from worsening, encourage attendance and participation at school, and help sustain work and build understanding, social relationships and friendships. Parents, families and communities need to be involved in this individual’s life and work together.

FASD lasts a lifetime but with the right help and caring, a good quality of life is possible. Care at home is incredibly important but can be challenging. Parents and carers need to care for themselves and be offered support too. NOFASD Australia can help. Please contact us on 1800 860 613.

With grateful acknowledgement to NOFASD Australia, a non-government national organisation registered as an incorporated association in South Australia under the Associations Incorporation Act 1985.
**Post Diagnosis – Support**

| NOFASD Australia is a strong and effective voice for people living with FASD and offers information, resources and ongoing support to individuals and families via telephone, email, online or by post. NOFASD Australia has a wide network of parents and carers in most locations across Australia and we can connect you with other experiences parents, and people who understand what you are going through. FASD lasts a lifetime but a better quality of life is always possible. Our knowledge and experience in supporting individuals, parents and families before and after diagnosis can help you. We work with people to share information, resources and offer professional support to service providers who might already be supporting your family or we can help connect you with these people in your community. | **CONTACT DETAILS**
Telephone: 1800 860 613
Email: enquiries@nofasd.org.au
Website: [http://www.nofasd.org.au](http://www.nofasd.org.au)
Online contact: [http://www.nofasd.org.au/contact-us](http://www.nofasd.org.au/contact-us) |
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NOFASD Australia raises public awareness of FASD through community education for individuals, parents/carers or groups and we deliver training to service providers who support families.

Parents, carers and their supporters can join the NOFASD Network and receive our monthly e-newsletter. If you do not have email, we can post out copies of the newsletter each month. NOFASD Australia has a Facebook page on which we post daily news and items of interest for individuals, parents and families from Australia and around the world.

The information you provide is private and confidential, we will always seek your written consent to share any personal information for any purpose and we respect your right to choose anonymity.

NOFASD Australia is non-government national organisation registered as an incorporated association in South Australia under the Associations Incorporation Act 1985 and has held Health Promotion Charity status since 2007.
### Australian FASD websites and resources

<table>
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<tr>
<th>Organisation</th>
<th>Website</th>
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<tbody>
<tr>
<td>FASD Hub Australia</td>
<td><a href="https://www.fasdhub.org.au/">https://www.fasdhub.org.au/</a></td>
</tr>
<tr>
<td>Australian Indigenous Alcohol and Other Drugs Knowledge Centre</td>
<td><a href="https://aodknowledgecentre.ecu.edu.au/learn/health-impacts/fasd/">https://aodknowledgecentre.ecu.edu.au/learn/health-impacts/fasd/</a></td>
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</tbody>
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### International websites

<table>
<thead>
<tr>
<th>Country</th>
<th>Support Group</th>
<th>Research/Other</th>
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Australian Parenting Information and Programs

- The Australian Parenting website [http://raisingchildren.net.au/](http://raisingchildren.net.au/)

Foster Carer Associations

- Foster Carers Association NT [http://fostercarersnt.org.au](http://fostercarersnt.org.au)
- Connecting Foster Carers – South Australia [http://cfc-sa.org.au](http://cfc-sa.org.au)