Let’s Talk About...

Mental Health and deaf people
Shaping a better, stronger, supportive future

DeafVictoria
Mental Health and deaf people
Lets Talk About.. Mental Health and deaf people conference
Deakin University Burwood Victoria
Conference Report
16-17th November 2015
Written by Melissa Lowrie and Gary Kerridge
Deaf Victoria
Executive Summary

On 16th and 17th November 2015, Deaf Victoria hosted the Lets Talk About... Mental health and deaf people conference at Burwood Corporate Centre, Deakin University, Burwood. The purpose of the conference was to have a bottom up approach to a mental health discussion around what is currently working and not working in Australia’s mental health system. These discussions revealed how Deaf, hard of hearing and deafblind people are currently accessing services.

This conference would not have been possible without the generous support of our conference sponsors. Sponsors were:

- Mental Health Australia provided $20,000 in a conference grant to support this conference.
- Deakin University provided the use of the Burwood Corporate Centre and its world class Audio Visual technology free of charge.
- Vicdeaf/Auslan Connections provided heavily subsidized interpreting and captioning, as well as in kind support.
- Deaf Children Australia, Able Australia, Karli Health Centre, The Deaf Society of NSW and Beyond Blue all provided funds to assist with the plenary sessions. Deaf Victoria is most grateful for their support and looking forward to working with them further in the future to achieve the much needed outcomes.

The conference centered on the Deaf/Hard of hearing/deafblind consumers of mental health services and their experiences accessing the system. It was attended by consumers who are Deaf/hard of hearing/deafblind and a range of professionals in the mental health sector and deafness sector alike. Presentations and discussions at the conference allowed participants learn from each other and provided information that was central to devising a plan of action.

The conference covered a wide range of topics not limited to:

- Audism
- Domestic Violence
- Alcohol and Substance Abuse
- Mental Health in the Workplace
- Suicide and Self-Harm
- Deaf Indigenous Australians and Mental Health
- Deaf Youth
- Theoretical approaches

The topics were far reaching and covered Deaf, hard of hearing, deafblind, children of deaf adults and parents of deaf children.

This report will recommend the following actions, based on the evidence collected at this conference:

1. Develop a community based support model for deaf mental health
2. Develop training and support to increase knowledge of the mental health system around working with deaf people for communication support professionals.
3. Develop accessible resources for people who go online to seek help from the mental health system
4. Develop training and awareness programmed for mainstream mental health professionals and medical professionals
5. Develop programmes for deaf people that focus on prevention throught the development of positive mental health activities and strategies.
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This report will summarize discussions and presentations held at the conference and make recommendations for actions.

The Steering Committee

The conference was lead by a group of experts in their respective fields. The group also has strong links to the grassroots Deaf and hard of hearing community, organizations and services for Deaf and hard of hearing people, as well as the mental health sector in Victoria. The conference leaders included:

- Melissa Lowrie, Manager of Deaf Victoria leads the agency responsible for advocacy of Deaf and hard of hearing people in Victoria.
- Gary Kerridge, an expert in the disability field, is currently employed as the National Disability Coordination Officer at Deakin University.
- Dean Barton Smith, a leading expert in disability and mental health and current CEO of Victorian Mental Illness Awareness Council of Victoria.
- Karli Dettman, founder of Karli Health Centre and a qualified Deaf mental health counselor, specializing in Deaf counseling.
- Sandra Spadea, a deaf mental health worker, with a lived experience of the mental health system.

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Keynote Speaker

We were pleased to host Dr Brendan Montiero, Consultant Psychiatrist in the field of Mental Health and Deafness since 1987. Dr Montiero is from The St George Health Care Group in Manchester UK and presented to the conference as our keynote speaker on both days. On day one, Dr Montiero spoke on the concept of Audism and its relationship to the mental health of Deaf and hard of hearing people. On day two, Dr Montiero focused on the Mental Health system for Deaf and hard of hearing people in the UK, providing examples of best practice model could be used and adapted here in Australia. Dr Monteiro has a special interest in Forensic Aspects of Mental Health and Deafness and has been instrumental in setting up and directing Forensic Services for Deaf People in the United Kingdom.

Dr Montiero has published extensively in the field of Mental Health and Deafness including:

- At least 5 Peer Reviewed articles.
- Contributing a chapter on deafness and communication in a book “Neurological Boundaries of Reality” (Farrand Press 1994).
• Lecturing extensively on topics related to Mental Health and Deafness in the UK, Europe, USA, South Africa, Japan and Australia.

It was an enormous privilege to have him with us at this conference.

**Attendance and structure:**

160 people from all over Australia attended this conference. The balance of consumers and carers to professionals was approximately half. The first day of the conference was structured to focus on consumers and carers telling their stories and sharing their experiences accessing mental health services under a number of key themes. The second half of the day consisted of eight separate plenary session topics, which will be summarized throughout this report.

Day two focused on theoretical approaches to treatment and statistics with Deaf/hard of hearing/deafblind people and professional development for people who work with deaf/hard of hearing/deafblind people with mental health issues.

**Topics of Discussion**

**Audism:**

Dr Brendan Monteiro, Consultant Psychiatrist, for Deaf people in the UK, has an international reputation in this field and presented on Audism.

Dr Montiero defined the concept of Audism as an emotional superiority based on hearing status. This can be seen in three different ways:

Hearing people feeling/acting superior to deaf people because they can hear and treating them in a negative/oppressive attitude.

Deaf/hard of hearing people who use spoken language having a negative/oppressive attitude towards Deaf people who use signed language.

Dr Montiero demonstrated an example of audism in a mental health setting:

“As a psychiatrist in hearing services I will expect people to come in with an anxiety attack, to be seen for maybe half an hour, an hour, be diagnosed, a bit of on the surface counselling and some sort of care plan going forward, maximum two to three hours even if there is waiting time. From what I can see (from the deaf persons experience in the same setting found in “Anna’s Story” page 8) it was roughly 36 hours and not only that, she came in with an anxiety attack and the process made her even more anxious. So the anxiety was sort of magnified. Ludo Timmermans did some research in Belgium published in about 2004-05; the average inpatient stay for hearing parents in psychiatric hospitals was four months. The average inpatient stay for Deaf People in psychiatric hospitals was 17 years. So you can see the difference between that. I mean, this displays Audism to the Nth degree.” – Dr Montiero, 16th November 2015

Audism in a mental health setting is about recognizing that Deaf and hard of hearing’ patients have a hard time getting access to services and it needs to be corrected. In order to assess anyone with a mental illness, you need communication.

Communication is the tool in psychiatry/psychology that allows understanding and interaction between doctor and patient. Through it you assess the person, you diagnose the person, you develop treatment. There are certain facts about deafness that need to be accepted. The person will not be able to function without some sort of environmental aids, be it an interpreter, captioning or the correct environment to lip-read. This is not a correction of deficiency but as a conduit to better communication. Without it, you cannot talk to patients about what is going to happen to them, how any medications diagnosed will affect them, the side effects.

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1 "Deaf" refers to people who are culturally Deaf, engage with the Deaf community and use sign language. Hard of hearing people are people who have hearing loss but don’t identify with the Deaf community. “Deafblind” refers to people who are both deaf and blind.
Dr Montiero estimated that the average assessments of a person may take one hour, but with a Deaf person requiring an interpreter, it may take three hours, and also needs to factor in the cost of interpreting. The reality of this is that it takes time and costs more.

Dr Monterio noted that generally, current services are hearing focused. This contributes to audism because the general view is that hearing services are superior. Instead of the hearing services having to adjust for the needs of Deaf people, they expect Deaf people to adjust to meet the needs of the service.

Service providers often try to avoid the cost of interpreting. They try to fit an appointment that requires three hours into a one-hour appointment. Many Deaf people that go to hearing based mental health services have said that advocating for themselves to access the services was far more stressful than having the condition in the first place.

Dr Montiero described the UK model. The UK model that he described has three different services for Deaf and hard of hearing people. Awareness of these services is good and as a result general practitioners, when a Deaf person presents with a mental health issue, know exactly where to go for information and support.

Dr Montiero provided an example of why the right advice and support is critical:

“In 1987 when I started as a consultant psychiatrist, I would receive a letter from a GP reading something like: This Deaf man has assaulted another person. He might be mentally retarded, schizophrenic or anything, please see and advise.” Literally. What he had done is put the top of a tin can on another guy’s head. They were having a fight after a few drinks. He managed to be closer to the can otherwise he might have been the receiver. So this case definitely needed specialist counseling from a service that could provide professionals with deaf awareness and Sign Language skills. So the attitude changed mainly because of exposure to services.”- Dr Montiero, 16th November 2015

This is an example of best practice in a mental health service that is inclusive of Deaf and hard of hearing people. Some examples of best practice are, but not limited to:

- The use of interpreters, experienced in mental health settings
- Development of community based support to provide individual support and advice to mainstream services
- Development of training and awareness around designing and implementing Deaf mental health support.
Anna's Story: An Admission to Hospital with the help of a friend:

One night I had a severe anxiety attack. Having had a history of mental illness for a very long time, I knew I had to go to the hospital. I drove to the hospital and presented myself to the nurse there. I asked for an Auslan interpreter. I told her that I have severe anxiety and was having a breakdown. I told her I am having suicidal thoughts. We were writing back and forth on paper and she told me to sit and wait.

I sat there and waited in the waiting room for 30-45 minutes. My anxiety got worse and worse. I was on FaceTime signing to my friend. At 11pm, I was then taken to a hospital bed and waited until the interpreter arrived. She arrived at 12.15pm. I was amazed at how quickly the interpreter got there- it was only because my friend Andrew is an advocate and made some calls through his networks to make it happen. Everything went smoothly after that. By 2.30am there was a plan in place for the morning- we agreed that -zat 10am, the doctors would visit me with an interpreter to discuss whether I could go home or have to stay in a psych ward. I was satisfied with this and went to sleep.

At 10am the next morning, the nurse came in and told me that there was no interpreter available. So I had to wait even longer so they could source an interpreter for me to see the doctor. After waiting hours and hours, at 3pm they admitted me straight into the psych ward with no discussion or anything. I panicked because I had not yet seen a doctor with an interpreter and this was not what I agreed to. I wanted to go home with special support. I texted Andrew for help.

I was still in emergency at that time and had been there since 11 o’clock the night before. 3 o’clock the next day I am still in emergency. I started to panic really badly and started machine gun texting Andrew. In this state, I can text 100 words per five seconds. I was very, very distressed. I couldn’t calm myself down or face the doctor. In the end, Andrew sent me a text and asked me to show the text to the doctor. In the text, he had explained a few things; that I am deaf and needed an interpreter, and not to try and diagnose me with anything and I might not understand what was happening. He offered to help them find an interpreter for me. Andrew also asked that the hospital let me access Wi-Fi so that I could then at least get FaceTime on my iPad to communicate and was able to get an interpreter through that if I needed to. Dealing with Andrew, the hospital was really great. Hospitals are normally not allowed to text. But in this case, they did text Andrew so that Andrew could give them the advice to help them get an interpreter.

By 7.30pm, I had an interpreter. I was scared because I have never been to a psych ward. Up until then I thought I had just suffered with depression. It was a Wednesday and the doctor told me that I should be ok to go home on Friday. However they wanted to see me again the next day and would book an interpreter for 10am. I asked the nurse later that day if they had booked an interpreter yet, and the nurse said, “I need to check the website. It’s a $370 booking fee. I will do it in the morning at 8.00 o’clock when they open”. I started to panic again- I knew that two hours notice to book an interpreter was not enough time.

The next morning, I was waiting in my bed at 10am and the three doctors came towards me- with no interpreter. I was just so tired from the day before. I started to get really upset when they tried communicating with me via pen and paper and I wouldn’t respond. They thought I was really angry, but I really wasn’t. I was just so tired.

I had texted Andrew again, and once again, Andrew was texting the doctor asked them not to go any further and he will come into the hospital at 1 o’clock and advise them of some way to work with her so he could help get interpreters in quickly. The doctors again to their credit, were really good and they listened. After Andrew arrived and was able to secure an interpreter, I asked him to sit in on my counseling session. When the doctors asked a question that I found difficult, Andrew was able to explain things, particularly related to deafness, deaf culture and interpreting access. The doctors really listened.

For the first time in a 14 year struggle, I was finally diagnosed with the right mental health issue and able to access the right treatment.

The system, it failed me. I had to use my friend Andrew and his networks to get access to what the hospitals should have known and provided in the first place. If it weren’t for his help, I would have been in the psych ward for weeks. This just demonstrates that something needs to change.
Amandas Story- My struggle with mental health issues and raising a family with autism and deafness.

I was born in Canberra in the 1970s and I am the only person in the family of 5 who is deaf. I was diagnosed profoundly deaf at 18 months old. I was enrolled in a small school at the age of three and stayed there for four years. I was happy there and was taught with both speech and sign language. Suddenly one day I was taken out of that school and put in a large mainstream school. Signing was banned and I had to use my speech and what little hearing I had to communicate. I became an angry child, wild and out of control. Schooling and growing up as an oral child was enormously difficult.

At 14 years of age, I was moved to Sydney and enrolled at a school with a deaf unit. Seeing the deaf students changed me. I no longer thought I would die. I used to cry myself to sleep because I was so afraid of dying as I thought all deaf people die young and I was the only one. I had dreadful social skills and it took me a while to adjust to fit in with my peers.

I endured a lot of trauma in my life that contributed to my downward spiral. I became a drug addict and had a drinking problem for many years. I was gang raped as a young woman, and tried to take my life three times. I was sexually molested by my great grandfather when I was 9 till I was 11. I was angry and very depressed for so long and I had forgotten how to live.

I also fell pregnant at a young age, and gave birth to a deaf little girl, Sophie. I had no support and was unable to raise her. The grief of not raising my child tore me apart for many years and it became unbearable when I hit rock bottom. Today she is a beautiful young adult and I am her mother, I have reconnected with her and we are close. I am grateful for that. My daughter has ADD and suffers from depression too.

I struggled with depression since high school because of the bullying I endured everyday. I had things such as drawings on the blackboard with “Go back where you come from”; locked up in lockers because I had a shrill scream, and the bullies loved hearing me screaming. I got into endless physical fights and never really understood why. I hated the fact that I was deaf. My mother wouldn’t talk about it, she wanted me to be ‘hearing’ and would tell me over and over that the Deaf community and Deaf people who sign are bad influence and stupid. She said I was ‘too good’ for them but I struggled with trying to be ‘hearing’ and to be good enough for her.

After school, I moved out of home and enrolled at TAFE, where I found my freedom to immerse myself into the Deaf world. It changed me, and since then, I had found my identity as a Deaf person.

My relationship with my mother became toxic and dysfunctional. My mother is a wonderful person and many people loves her, she is very kind and caring. However the damage between my mother and I became too hard to be repaired. I could not continue to live feeling responsible for my mother’s depression and sadness. I could not continue to feel that I was not good enough for her. I made the choice to cut the invisible string between my mother and I. It was the hardest thing I had to do.

I am glad to say I am ok now, I have been positive and living my life happy as I am for 2 years now. I learnt one thing – accepting yourself, as you are, deaf and all then it is easier to grow up as a deaf person.
Domestic Violence

Debra Swann and Kate Frost, two Deaf women survivors of domestic violence presented on the topic of domestic violence, their experiences getting help and the barriers for Deaf people.

Deaf victims of domestic violence often face unique circumstances:

- Information can travel quickly within a Deaf/deafblind community, compromising confidentiality and the victim’s safety.
- Law enforcement and shelters are often not skilled at communicating with Deaf, deafblind, or hard of hearing individuals and often don’t have interpreters.
- The perpetrator may take away their communication devices.
- The perpetrator may give false information to the victim to make them believe they have fewer options.
- The victim may be isolated from family, friends, services, resources and options.
- Ringing a hotline is very difficult. The hotline counsellors often want to ring you back, and won’t email or SMS you due to the risk of the perpetrator finding the email or text.
- AVO’s have to list a person’s address. However if the victim is Deaf, the perpetrator could attack when they are sleeping, because they know the victim cannot hear.

Some examples of what victims face and the tactics perpetrators use to abuse the Deaf:

- Intimidation through gestures, facial expressions, or exaggerated signs, floor stomping and pounding on the table or door
- Signing very close to a victim’s face when angry
- Criticizing the victim’s sign language skills or communication style
- Not informing the victim when people try to call on the phone or try to catch their attention
- Excluding the victim from important conversations
- Leaving the victim out in social situations with hearing people
- Talking negatively about the Deaf community
- Wrongly interpreting to manipulate the situation if the police are called
- Not allowing their children to use sign language to talk with the victim
- Not allowing their children to be proud of Deaf culture
- Criticizing the victim’s speech and English skills

Swann and Frost had three specific recommendations for tackling domestic violence for Deaf people:

- Develop a website on domestic violence specifically for Deaf people based on the “Deaf Hope International” website in the UK. This website has resources in sign language, information that is easily accessible and when you press the “escape” key on the website, it will instantly change the screen to Google and erase any way of the perpetrator tracing back the website
- A school education program to be developed based on healthy relationships, domestic violence education and what is and isn’t acceptable
- A 24/7 hotline for Deaf people in domestic violence situations that is accessible for all people. This could be in the form of online chat.
Mary's Story- Finding my voice

I have been deaf since I was six weeks old and had the measles. Living with my small amount of hearing and not being validated the way it should have been has led me to having a life of heartache, inner pain and a severe social phobia. I was brought up in a very dysfunctional and abusive family. I was one of six children and as the only deaf one; it was easy for me to become invisible. My self-esteem was crushed from a very young age and it wasn’t until I was out of the abusive environment that I took myself to a doctor and told him I didn’t think I could hear properly. Finally my hearing loss was taken seriously and from then on I have had hearing aids in my ears. Of course this didn’t undo the trauma I had suffered at home and at school because nobody understood why I never spoke and why when I did it was a tiny whisper. I was forever being yelled at to speak up by my father. Life was a constant struggle with depression and anxiety but I was so young, I didn’t understand these feelings. I knew there was something wrong with me because I wasn’t like anyone else.

As an adult my marriage and bearing of three sons was tough. After fifteen years of marriage, I ended up along with my sons and trying to be mum, dad, friend and everything to them. I developed a severe eating disorder and depression. I wanted to disappear; I felt I was a burden to everyone, including my sons. I have had so many hospital admissions all over Australia, but have experienced some horrifying treatment and ignorance towards me because I was deaf. It was never taken into account that I was not just shy, I was deaf and although I had hearing they never completely fill the gaps.

I am lucky to still be here. Many times, I have given up, but continued to seek treatment until I found the right one for me. I found a terrific psychologist who would speak up, face me, repeat herself without complaint and treat me as if I am hard of hearing, not deaf and dumb as I grew up to believe. I relied on writing as being my voice for a long time, now I am working very hard to find and use my voice again.

Today, I still struggle with severe anxiety and anorexia nervosa, however I hope in time that I will beat this battle.
Alcohol and Substance Abuse in the Deaf Community

Debra Swann, a Deaf case manager with a lived experience of substance abuse, and Sandra Spadea, a person with a family member of lived experience, presented on this topic.

Deaf and hard of hearing people who abuse substances often face unique circumstances in accessing help:

- Deaf and hard of hearing people, as well as children of Deaf adults (CODA’s), are more vulnerable to the abuse of substances:
  
  “Commonly people, Deaf people, are at more risk of substance abuse because of lack of information. <We argue that> CODA’s are likely to be at a high risk also, because they’re not aware.”
  - Swann, 16th November 2015

- Alcohol Anonymous/Narcotics Anonymous/Eating Disorder groups are not eligible for funding of interpreters and captioning, therefore inaccessible for Deaf and hard of hearing people

- There is next to no information available in signed language, or with added captions online for Deaf and hard for hearing people who want to seek help.

  “Mental health professionals need to try to imagine that they find themselves in another country and they struggle with drug and alcohol issues. They go to try and access services that are presented in a different language- a language that is not provided to you via an interpreter. The information will be meaningless to you, especially if you’re feeling isolated and alone. That’s how Deaf people feel in their own country, information comes that is meaningless and complex.”
  - Spadea, 16th November 2015

- There are very few trained professionals in the field of substance abuse that have knowledge about working with Deaf people, hard of hearing people, deafblind people and CODA’s. Therapy sessions become about the patient educating the therapist about Deaf cultural issues, rather than treating the problem. Some children of Deaf adults feel that they are not able to access services because commonly, mental health workers are more interested in the family dynamic and Deaf heritage, so they’d ask questions about deafness, which meant they weren’t asking questions and treating the person that is ill.

Some recommendations from Swann and Spadea were:

- Resources and information on Drugs and Alcohol to be provided in sign language online

- Funding for Alcoholics Anonymous/Narcotics Anonymous/Eating Disorder groups and other community based support to be able to provide interpreting and captioning support for their meetings.

- Research into the effect of substance abuse on the mental health of Deaf people and CODA’s.

- Advocate for the provision of specialised training for Deaf and hard of hearing people, or people fluent in sign language, to become NA/AA support workers and counsellors
Rachels Story- My CODA brothers struggle with addiction.

I’m deaf and my parents are deaf. I’m a big sister to two younger brothers. We are one year apart in age and we share sign language as our first language. We are very close and my brother, Parker, was my best friend- he was always cheeky and made me laugh he was very talented with basketball, football and cooking. He struggled at school, but he was amazing at expressing himself. He left school at 13 after mixing with the wrong crowd.

We had moved to a new area and gone to a new school. He tried heroin and stole to feed his addiction. He ended up in a youth detention centre at the age of 15. As we got older he would always socialise with my deaf friend and I. He never saw us being different to him. He very much felt the same as us.

Parker spent 17 years in and out of prison and always felt that he was misunderstood. When I was visiting him in prison at one stage, I asked if he was seeing a psychologist or counsellor. His response was they had no understanding of him and they asked too many questions about deafness. He felt that he shouldn’t have to explain about deafness or explain himself. Parker was a CODA (Child of Deaf Adults) and he felt that the Deaf language and Deaf culture was very much a part of him. They were always asking him about deafness not about himself and his issues. He went for many years without access or support. The only information he received was through reading books or documentaries. I found it shameful.

When Parker was aged 32, this was the longest time he had been out of prison and free of his addiction. In a horrible twist of fate, he passed away in an accident.

Its time for things to change for Deaf and CODA addicts. The needs of CODAs have been ignored for far too long. I would like to see Deaf sector organisations put the needs of CODA's firmly on their agendas. There needs to be more research into the mental health and wellbeing of CODAs and their support needs.

Clara's Story- Forgiveness

I was sexually abused by 9 deaf boys as a young girl in a Catholic school over a number of years. I was also bullied by a couple of deaf students at the school. At the time, I did not know the word “sexual abuse” while growing up. My language was underdeveloped and when I was at school in the 70's and 80's, there was no education on these issues. I also abused alcohol and had no idea of the destructive and serious side effects of this. I blamed my troubles on the fact I was deaf, and it took me a long time to accept my deafness.

Eventually my troubles led me to a number of suicide attempts, which saw me enter the mental health system. After a long time, I found a counsellor skilled in Christian counselling and provided me with Auslan Interpreter access. I have been seeing this counsellor for eight years and I feel I am a changed person. I am a much better, happier and healthier person without alcohol. I used to abuse alcohol to escape from anger, hurt and pain. I have been sober now for 9 months however, I struggle in times of heightened anxiety and depression. Forgiving the 9 deaf boys has been a very long process, but forgiving people who have hurt us badly, is very healing and powerful and really helps us to leave our sad past behind and move forward. A lot of people stay bitter and are stuck with their lives, because they are not willing to forgive themselves and others every day or they don’t want to see a counsellor. Daily forgiveness is very important for everyone, because it gives us peace, love and joy and anger and bitterness can be very destructive for our mental and physical health.

I am able to enjoy life much more now.

Frasers Story- Growing up deaf in a small country town

I grew up in a small country town without any knowledge of sign language therefore some communication problems. As a young teenager, I became frustrated, bored and confused and this took me on a downward spiral to depression and binge drinking. At the time I didn’t know what “depression” meant and drinking was just my way to cope with the bad feelings.

I ran away from home and drifted around all over place to find myself. I finally settled down in Bendigo and started doing a lot of exercise- mainly swimming- and joined a church group. They stopped me drinking, however it didn’t last because I had no interest in religion. I found out that there was a deaf club every Friday night in Melbourne, so I decide to go and check it out. I was shocked when I arrived to see them all signing to each other and slowly but surely I started picking it up. However I fell back in trap and started partying with deaf people and started drinking again.

Over time my health got worse, and my depression skyrocketed. The doctor tried giving me antidepressants but I didn’t understand the instructions for taking the medications ask I couldn’t hear or understand him at the appointment. I ended up taking too many tablets and found myself in hospital. The doctors there said that needed to go to a detox program in Footscray to stop drinking and dry myself out. I have now stopped drinking, however I was disadvantaged because Alcoholics Anonymous meetings do not have funding for interpreters or communication support. Recovery took longer for me because I couldn’t access this service.
Mental Health in the Workplace

Stephanie McDonald was born profoundly deaf and utilises a Cochlear Implant and hearing aid. She has worked in a variety of workplaces including Allied Health, Disability Employment Services (DES), and IT industries.

Working full-time in these challenging roles with a hearing loss was difficult, and caused some anxiety, high levels of stress and difficulty with coping with the large workload. McDonald shared her experiences about mental health issues in the workplace for people who are Deaf/hard of hearing and how to maintain mental health in an effective way.

A Mentally Healthy workplace is: “One that protects and promotes mental health and empowers people to seek help for depression and anxiety, for the benefit of the individual, organization and community.”

McDonald noted that, according to beyondblue, in the general Australian population:

- 45% of Australians between the ages of 16- 85 will experience a mental health condition in their lifetime
- 21% of Australians have taken time off work in the past 12 months because they felt stressed, anxious, depressed or mentally unhealthy
- It is estimated that untreated mental health conditions cost Australian workplaces 10.9 billion dollars

Anecdotal evidence arose from the Queensland Health Deafness and Mental Health Report, 2008. The evidence indicated that most depression and anxiety for Deaf people in the workplace stems from ill informed communication practices leading to reduced opportunity for the Deaf employee. This report notes that:

“Some barriers Deaf people face include communication issues, lack of accessible training opportunities, poorly defined career path, limited participation in the social milieu of the workplace and limited opportunity to engage with the public. Deaf people are often found to be overlooked for promotion and career enhancement opportunities which can contribute to feelings of worthlessness leading to an array of mental health disorders”
- Queensland Health, Deafness and Mental Health 2008


As a result of the Deafness and Mental Health report, the Australian government established the National Mental Health and Disability Employment Strategy in 2008. The strategy aimed to address the barriers faced by people with a disability and/or mental illness that make it harder for them to gain and keep work. From this, the Employment Assistance Fund was created, which helped to find and retain employment for Deaf and hard of hearing people, but doesn’t improve the social interaction, or workplace experience of being a deaf person in a hearing environment.
Carl's Story- A battle in the workplace

I have found myself employed as a disability expert advising the council on disabilities. Ironically, being a person that had to advise the council on disability access, I had difficulty getting the access that I required. The council decided in their wisdom that my access amounted to interpreters. And that was it. So, in their minds, providing interpreters was enough. When you have a look at the whole system of how councils work there are lots of things you have to do. I had deal with departments, individuals and community organizations. One of the difficulties I had in working with the department was that they wouldn’t commit to disability. Disability is not their core business. They don’t want to think about disability access unless they are forced to do it. For example, if I had a project with them on emergency management, and advise them what they need to include to be disability accessible, they will eventually say it’s too much for them and just simply stop responding to my requests. This happened with several departments. I tried to get the council to come on board and we meet halfway. I can’t have an interpreter with me all the time so I have to do things in a different way, such as emails. Some people were very good with emails, and others were not so good. What that meant was that there were often projects that I had to do that were delayed. My board would be on my back to finish them. There was one particular project where I had to do an online survey. My boss took 11 months to get back to me with information I required, and every month she was telling me to finish it. I was saying to them that without the online survey, I couldn’t finish the project and I was waiting on other people to respond to their emails.

One day I met with my boss and she said something that was really the responsibility of another department. I had been waiting on them for three or four months. She refused to understand what the issue was and said “Carl, this was left with you and this has been going on for 18 months or so”. Now, I’m a successful disability advocate, and have been for over 30 years. I can handle myself. However after months of this, I just couldn’t take any more. It got to a point where she said this was left on you. I said to her, “No. No more. I’m not wearing this anymore. That’s enough.” and I sat down at my desk and I was hyper ventilating. 18 months of this. I got an email from my boss that said “before you rudely stormed off ... I wanted you to forward me all of the emails that you sent to this particular person.” Now, the thing was that my boss had been CC’ed to every one of those emails so she knew what had happened. So I found all of the emails, 51 emails altogether and I sent them to her. I replied, “Deal with that.” And I was angry.

The next day, to cut a long story short, on my desk there was a letter of discipline. I took the letter, I went downstairs to meet with the Human Resources person as instructed. I walked in the door and I just sobbed. I bawled and sobbed. The Human Resources person was good and organised for me to see a counselor. Having to see a counselor at that point in time, I was lucky. I have extensive networks. I contacted a friend who is an interpreter, and within an hour came to me, and we went off to see the counselor.

I was lucky, because I know most people don’t have that luxury. If any other deaf person had just broken down, most people don’t have that luxury of calling somebody like I did. Over the weekend I basically just cried and cried and cried. By the Sunday, I said to my wife, “Jenny, I can’t deal with this on my own, I have to see a doctor.” The doctor was good but the problem was that Jenny, she is Deaf, and my doctor is not that easy person to lip read. My wife had to interpret for me.

He did the normal thing. He offered me antidepressants. I didn’t want that so he gave me diazepam to calm myself down. Every day I cried. Everybody supported me, particularly my wife and three boys.

Then, over time, something else happened where I got a phone call from the police because I had smashed somebody’s car. The mirror fell off and I didn’t know. The policeman came to my house and then that was it. We were all distressed. It was the straw that broke the camels back, trying to deal with what I call bullying, victimisation and all of that. I just broke down. I wanted to hit something. Not my family, but I was aiming at the wall. I could just imagine a big hole in the wall, to make me feel better just to damage something. I held it back. I was so angry. So frustrated.

In the end my wife, phoned Lifeline for me. I am lucky because I have good English through the national relay service, I could type back and forth with Lifeline and we had a chat. In the end we ended up in the hospital, and again my wife was the interpreter. I was in the hospital for about eight hours. At some point Jenny had to go home because the boys were on their home all alone. I was in there on my own. I had to face doctors, nurses and a social worker with cerebral palsy (which is not a problem, just very difficult to try and lipread). I was distressed and crying and it got to a point where communication and everything was really, difficult. I requested several times for an interpreter. Every time, they said an interpreter will come, but it never happened once. Jenny had to continue interpreting for me.

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I’m one of the lucky ones. I have a network. And if I didn’t have those networks where would I be? I would be lost I know people. I know interpreters. I know social workers and psychologists and I asked them for help to the hilt. Not everybody can do that. The whole process is hearing based. You come into the hospital, you have to meet the nurse, you have to talk to them. When you are distressed you don’t want to write notes. Your family help out and do things for you, however the stress on your family is enormous and its not ideal to rely on them.

Everything is so hearing focused and you have to wonder what sort of information gets exchanged between deaf patients and hearing professionals and whether it’s actually accurate or not. Are they making the correct diagnosis based on fractured information? Do they really have the full story?

The mental health system is not user friendly. Not for deaf people. Not for everybody. But particularly not so for deaf people.
Suicide and Self-Harm

Alana Roy, a psychologist specialising in deaf and deafblind clients presented on suicide and self-harm. She reported that in the wider Australian community, there are:

- Approx. 9 suicides each day
- Women more likely to attempt suicide
- Men are three times more likely to complete suicide

In Roy’s PhD literature review, she searched extensively, with no limiters on years, and found no article with a focus on suicide or self-harm in deafblind people. There is a gap in research and understanding of self-harm and suicide prevention, rates, diagnosis and treatment. Roy explained that Turner et al conducted literature review from 1940’s- 2006 across multiple databases and only found 13 articles for inclusion with a focus on self-harm or suicide in Deaf people.

She did find that:

- Few studies focus on suicide or self harm
- Little evidence that suicide rates are higher
- Higher levels of depression and risk
- No firm evidence of effective suicide prevention in Deaf and deafblind people

Roy also found significant barriers in talking about mental health in the Deaf and deafblind community. Some of these barriers are:

- Stigma and shame e.g. losing face in the community: disclosure can be challenging. In these communities many worry about shame, stigma, not getting jobs, losing face, no longer being been seen as a role model or advocate
- Gossip and lack of trust
- Audism and medical model
- Lack of access to service and communication
- Difficult to access self harm and suicide support via text, Skype, interpreters and email

She noted that a number of mistakes that hearing and Deaf mental health professionals often make when working with a deaf and or deafblind clients in crisis:

- Pretending or assuming they can sign or understand the client
- Not having adequate training and skills in Deaf, deafblind trauma and risk assessment,
- Audism- assuming they know more or are correct because they can hear
- Not consulting with Deaf/deafblind specialists and community.

Specifically in the case of Deaf mental health professionals:

- Assuming they know the client e.g. because they grew up with them, case managed them for a long time
- May be able to communicate effectively but may not have clinical risk assessment skills,
- Assuming that Deaf and deafblind person wants or should want Deaf or deafblind professionals to support them and is reluctant to engage with hearing professionals.

She stressed that mental health professionals and the Deaf/hard of hearing/deafblind sector need to work together, respect each other’s differing points of view, and offer options and choices to those at risk.

The most important thing for the person who is self-harming or is suicidal is to access a professional with whom they have a good connection, can trust and who has the skills to work with risk. They should have access to hearing, Deaf and deafblind professionals and role models. This is a personal choice and should not be judged.
It was noted that some Deaf and deafblind do not want to be affiliated with an agency they know well and choose to have contact with hearing professionals for confidentiality and support outside of the Deaf and deafblind community. However, at this point in time, there is a lack of Deaf and deafblind professionals with Medicare rebate capacity, limiting the choices available.

Recommendations include:

- Funding for research into mental health issues in the Deaf and deafblind population
- Review of professional capacity of hearing, deaf and deafblind psychology and counselling services available, and access to the Medicare rebate and NDIS.

Patrick’s Story- Recovery and Male Pride

In 2006 my then wife gave birth to twin girls. On their birth, she developed severe postnatal depression. I was working full time at the time and she was the stay at home mum. Alarm bells rang for me when she texted me one day to tell me that she’s gone shopping and that the girls were on the floor with the dog looking after them. I rushed home from work, with my mother and mother in law both rushing over in a bid so that someone will get there in time, and luckily everything was fine. It was at that point, I had to act and make sure she got help. Both families offered immeasurable support. We went off to a sleep school to get the twins in a good sleeping habit, my mother and father, my sister and my mother in law all took turns coming over to help me settle, feed, bath and put the girls to sleep. My wife slept pretty much every chance she could get- she was ill and there was nothing we could do.

I had to juggle holding down a job, being an almost full time carer to baby twin girls and also look after my wife. Whilst my wife and the girls were getting all the support, love and help they needed, I felt like the forgotten one and I let my pride present to the world that I was strong and able to cope. When it was in the middle of the night, when I was trying to get the girls to sleep on my own, I was curled up in a ball crying with the two crying babies in my arms. And no one knew.

After the girls turned one, my wife decided that in order to get better, she wanted to return back to work full time, whilst I give up my job to become a full time father of the girls. However when she said she wanted to work, this meant she went to work, came home, bathed the girls and went to bed. I was left carrying the babies, so to speak. It was then that my five-year secret battle with depression began. I developed a severe skin infection that was related to stress, however I used to stand under the shower with hot water full blast (no cold water at all) and burn my skin, because the pain was a release from the pain I was going through inside of me. I would then take a bath with bleach to dry up the burnt patches all over my body. I now know, it’s an extreme form of self-harm. I started binge eating, resulting in a lot of weight gain. I was very unwell, yet to the outside world, they just saw me as a father of twins who was doing the best he could. No one had any idea.

After a particularly bad day, I had spent the whole day contemplating suicide. I was at the verge of planning the act when I looked at my beautiful girls and realized that I couldn’t do this anymore. I rang the Beyond Blue helpline and was told they couldn’t help me. In a distressed state, I built up the courage to tell a visiting friend, who long suspected I wasn’t well, what was going on and he sat me down and had a good talk to me about my options. He told me to see my GP, gave me the number of the local CAT team and encouraged me to tell my family. I told my family who were in varying states of shock and quickly made an appointment to see a GP and again a psychologist. I didn’t have an interpreter or any support with communication by choice. I felt I could lip-read and hear well enough to get by and I could. This was my choice.

I faced the fact that I was deeply unhappy in my marriage, and there was no hope of trying to fix it. I needed to talk about what I was feeling and find my happiness again. Eventually my wife and I divorced and arranged to share the care of the girls. I started telling my friends what was happening with me, starting with my friends at my football club. What followed was an outpouring of support and a number of other men telling me they were glad that I said something because they were secretly struggling too.

It has now been six years since the day that I decided to tell my family and friends my struggle. I am happy to say that despite what has been a very difficult six years, I have now found happiness again. My skin is now clear, my health has improved, and my weight is slowly but surely starting to come off. I have found love again and engaged to be married. I have a wonderful relationship with my now ten-year-old twin girls and enjoy life to its entirety.

My story is one of recovery, the power of male bonding and the need to bypass pride.
Deaf Indigenous Australians and Mental Health

Jody Barney is a leading Aboriginal Deaf consultant, a proud Birri/Gubba -Urangan/South Sea Islander woman from Queensland, she’s spent the last 30 years working with communities across Australia. Jody Barney has been granted permission and access to work with Aboriginal and Torres Strait Islander Deaf and Hard of Hearing people in justice, education, welfare, community development and health.

Therefore, the work done by Jody Barney highlights the need to have a robust and culturally engaging discussion at the Let’s Talk About…Mental Health and deaf people conference, on working effectively with Deaf and Hard of Hearing Aboriginal and Torres Strait Islander people in the Mental Health sector. The work done by Jody Barney is done with the utmost cultural respect where she ensures the cultural safety of all community members attending the session.

Deaf and hard of hearing Aboriginal and Torres Strait Islander people are a minority within a minority, and even more so are those who live, experience and manage their mental health issues.

Due to the high prevalence of hearing loss amongst Aboriginal and Torres Strait Islander communities due to chronic ear disease, poor housing, inadequate sanitation services, and lack of cultural safety accessing health services, many people often aren’t diagnosed with hearing loss until they reach school age or often older.

Culturally Aboriginal and Torres Strait Islander people are more astute to their environment, using visual modes of communications and signing systems to communicate between nations, during ceremonies, customs such as “LORE” or “Sorry Business” which are traditional times of healing, initiations, grieving and rituals such as participating in decision making practices, such as funerals and marriages. It is customary that all members of the families, clan and community have a role and duty to uphold traditionally, these are often cultural obligations that have no exemptions. Each member of the family must follow the culture of their family and adhere to the lore (law) held by the senior lore men and women. Therefore, the use of the spoken word was often forbidden, this could result in weeks or even months for some communities.

Traditionally those who were known to have hearing loss, were often observed by the Elders, and given appropriate names, totems, roles and song lines. Often many Deaf community members were seen to predict weather, notice changes to the landscape and were often given roles to seek food as their observation skills were more astute.

Jody Barney stressed that often the entrenched fear and transgenerational removal of family (since colonisation) has impacted all in the community today even the deaf members within communities. Where you see the ongoing displacement felt by those who are not with their kin or on their country can and has heightened the presents of mental health concerns for Deaf Aboriginal and Torres Strait Islander people today.

With the communities being able to travel frequently the presence of rural, remote Aboriginal Deaf people are popping up across the coastline communities and in the metro cities. When Aboriginal people move from across the country, there is vast differences to their use of “hand talk” or signing systems that don’t transfer into Auslan. These are more in relation to their totems, connection to kinship systems where by a “mum” can mean any kind of significant female in their lives: in this case, it could be their wife, sister, aunty or a friend.

Many urban Aboriginal Deaf people often have access to Auslan, learn spoken English and engage with their cultures daily or not at all. The various forms of the modern family unit can compound their mental health when trying to “fit in”, be accepted and even try to understand why they can’t explain how they feel. It is often difficult for the services to understand hearing Aboriginal cultural groups let alone Deaf members of the community. The struggles that service providers, interpreters, mental health workers, Deaf workers will face is accepting the identity of a Deaf or hard of hearing Aboriginal person, and their needs to return to wellness and treatment options.

Whilst the Deaf community is quite small, the Deaf Aboriginal and Torres Strait Islander (signing) Deaf community is even smaller, the prevalence of hearing loss amongst Aboriginal and Torres Strait Islander people’s is much higher, but the use of Aboriginal signing systems and hand-talk is much less. Therefore, the use of cultural ties, connections and awareness will greatly improve the health and well-being of Deaf and hard of hearing Aboriginal and Torres Strait Islander peoples. This is done by making sure that interpreters
who are booked have had cultural awareness training. That if they are known to the deaf person, they can assist in the message getting across correctly.

Often these ties can't be discussed, the confidentiality of their work often can restrict them in providing the best possible support and access. The conversations interpreters, health workers, practitioners can have will improve the outcomes of the person who's unwell. Often scenarios are used to explain a point, which Jody Barney did in this session, however, due to the cultural protocols they won't be put in this report. However, the message for readers is that without the connections or understanding of the importance of these connections and proper protocols the wellbeing of Deaf Aboriginal and Torres Strait Islander people will not improve.

The work that is being done in the mental health sector is based on a different way of doing. A different way of learning. Whilst practitioners and health workers will use the various taught theories, the impact for Deaf Aboriginal and Torres Strait Islander people is the opposite. That often the individual in western terms is the main focus, for Aboriginal and Torres Strait Islander people its community, a collective. It is hard for non-Indigenous people to understand this sometimes, especially around family structures and kinship systems and the strong connection to land.

What is culture being that the Deaf person who is unwell is focused on something that’s happening 400/500kms away and can't hurt them but it does, that’s when workers miss the importance of the culture and only focus on the person in front of them. Most Aboriginal people won’t even talk about themselves in general which is why they turn up to services with three or four people because those people are there to talk for them- this is the culture.

Most Aboriginal people actually go to mainstream services rather than Aboriginal services, and it's believed that because the community is close knit, they don’t want their community to know that they are in need of mental health services when they know the difference of being unwell. However, it is a catch 22 situation; the expertise in Indigenous Australians is not found in the mainstream services. Mainstream services are still struggling to work with Aboriginal people in general, let alone Aboriginal people that are Deaf or hard of hearing or have mental health issues or drug and alcohol issues.
Deaf Youth

Paula Zalcberg is a counsellor and family therapist who works specifically with Deaf and hard of hearing children, adolescents, adults and their families. She is fluent in Auslan and currently employed at St. Mary’s Primary School two days a week. She has her own private practice where she sees children, adolescents and adults for individual counselling, couple counselling and/or family therapy.

Zalcberg spoke about attachment theory. The child obtains physical and psychological safety through proximity to the parent, who is available, sensitive and responsive to the child’s various needs. The parent also has the role of repairing any relationship breaks that might threaten that sense of safety. Within attachment security, the parents serve as the secure base, which provides the child with the safety needed for it to explore and interact with the world. Attachment security enables the child to regulate fears in order to be free and learn from objects and events in his or her environment. As the child matures, physical proximity becomes less necessary, while psychological availability remains crucial throughout childhood and throughout adolescence. This is critical in the emotional and social development of the child.

Some parents develop a strong healthy attachment with their babies. Others have a lot of difficulties, often related to their own upbringing. Often when the family comes to counseling none of its members are feeling particularly safe. Parents feel they may be judged negatively and blamed for problems their child might be manifesting. The child may not feel safe because he or she may be told they are going to counseling because of their behaviors. The therapist’s first task is to facilitate a sense of safety among all the members of the family. The therapist needs to become the secure base.

Sometimes when parents receive news that their baby is Deaf it can elucidate feelings of loss for a perfect baby they no longer have or hoped for. This can cause some parents to become distant from their Deaf baby. This can make the bonding between mother and baby inconsistent and fragmented. As a result of this insecure attachment it may cause a Deaf child’s social and emotional development to also be insecure, delayed and fragile. This has a huge impact on the Deaf child’s understanding of the world, their ability to trust and their ability to have meaningful relationships. It’s important to note that 90 to 95 per cent of Deaf children are born to hearing parents who often had no experience of deafness before. Most of these parents will have had no contact with a Deaf person previously.

Parents see audiologists and the Cochlear implant team quite frequently as the child grows. The parents get used to handling hearing aids and moulds and processor bits and pieces and become familiar with the hearing aid jargon. What parents do not learn is the language that is related to the child’s emotional understanding about its deafness. It is so important that Deaf children, siblings and their parents also understand this aspect of deafness.

Often Deaf children come home from school and they say to their parents, “I hate being Deaf.” Often parents feel the child’s pain and offer words of encouragement like, “Don’t worry, darling, you’re the best runner in the class or you are really good at spelling or you are in the top reading group.” This does not actually validate what the child is feeling. Zalcberg encourages parents to say, “Yes, it must really suck when you can’t hear. It must be awful for you. You must get so frustrated and angry at times.” This is actually hearing the child and validating what he or she is feeling. She also encourages Deaf children to talk about their frustrations when they are constantly missing parts of conversations and feeling left out and confused.

Deaf young adults also need the skills to be able to advocate for themselves. A lot of kids struggle with accepting that you can feel that being Deaf is okay, and it should not be a barrier to following your dreams. Zalcberg notes that a lot of her work is with teenagers who are oral and often they get overlooked in terms of the issues to do with being oral. Some examples are:

One teenage oral Deaf male suffered from high anxiety especially in social situations. He was going on an excursion with his hearing school mates and he needed to tell them to face him when they spoke to him so he could lip read. He was quite horrified at the thought of telling someone he was Deaf. - Zalcberg, 16th November 2015

An oral 25-year-old young lady who had a mild to moderate hearing loss and was also very shy and withdrawn was suffering from depression when I met her. She had gone through high school as the only hearing-impaired student in the class. When she finished year 12, took off her hearing aids and didn’t tell anyone she was Deaf. This included tertiary teachers. She struggled through several courses, not completing any of them. When she came to me, she was suffering from depression, had symptoms of anorexia and was incredibly socially isolated. I had a session with her, her sister, and parents. Very soon into the session, it became evident that
the family had never really accepted their daughter’s hearing loss. Even some of the extended family did not
know at age 25 that she had a hearing loss. So the first work I did with this young lady, was to talk about what
her hearing loss meant to her. Again, the feelings of enormous shame and embarrassment came up. Bit-by-
bit, I encouraged her to start wearing her hearing aids, role-playing job interviews and work situations with her.
During the session with this young lady’s family, I asked them what they understood about her hearing loss.
They replied that it was the same as wearing glasses. This is often an analogy that people make that is actually
quite wrong. The parents were both medical doctors. – Zalcberg, 16th November 2015

Teenagers strive to be accepted, fit in and most importantly they do not want to be different. This is
particularly hard for deaf kids. It’s very hard to fit in when you have a hearing loss, because that makes
you different from the start. We can see from these stories that a common theme in oral-based deaf youth
is extreme difficulties in understanding and accepting their own deafness. They have intense feelings of
shame. They often feel that hearing people, especially within their peer group, are constantly judging them.
It is common that they feel that they are not good enough and get very angry and upset about this. These
can be kids who are academically equal to their peers, however still feel other people judge them as stupid.
They often walk around with so much frustration and resentment towards hearing kids who they feel don’t
understand what it feels like to be Deaf.

Deaf kids tend to learn to use different strategies to protect themselves from the feelings of rejection. One
of these strategies what we call the ‘Deaf nod’. When they nod and smile or laugh and hope its at the right
time in the conversation. They do this in an attempt to save face and fool the hearing person into thinking
that they know what they are talking about. At time this works, but in the long term, what does that do to
a teenager’s self-esteem, always second-guessing and never being 100 per cent sure that they have all the
information?

Deaf kids often want to know what is happening or said in a conversation. They are regularly dismissed with
“Don’t worry about it, I’ll tell you later.” Unfortunately this rarely happens.

Many Deaf kids have some degree of language delay. For many the delay is quite severe. The impact of
deafness is influenced by various factors such as quality of family environment, parental adaption, and
coping with deafness. In a lot of cases in hearing families of Deaf children, discuss the day’s events at
dinnertime. Often the deaf child does not get full access to these conversations and they sit quietly eating
their dinner. This lack of communication can cause a high incidence of anxiety and depression in the Deaf
child. This can affect their ability to develop close and meaningful relationships.

Exploring friendships is such an important part of adolescence. Without appropriate role models this can be
fraught with difficulties. Anger and resentment build up over the years and the adolescent either becomes
depressed or withdrawn. A common complaint from Deaf children is, “I feel really lonely at school but I feel
even lonelier at home when I can’t understand my family talking: I’m always the last one to know what’s
going on.”

Most oral teenagers don’t identify with the Deaf community. When they meet signing Deaf people they often
feel intimidated and reluctant to communicate. Many Deaf kids speak well and one would never know from
their speech that they are, in fact, Deaf. Often oral Deaf kids experience identity confusion. They do not
identify with their Deaf peers yet they feel alienated from their hearing peers. Most oral Deaf kids see their
deafness and hearing loss primarily as a disability, impairment and a physical disorder. For many Deaf kids
that have grown up using Auslan, being Deaf means being part of a unique culture with its own language,
traditions and values. Zalcberg encourages all her oral clients to learn Auslan and find a place where they
feel accepted as a Deaf person.

Mentoring is very important for Deaf kids. We need to encourage this to happen more, particularly for Deaf
students at a vulnerable age of years 8 and 9. They will benefit immensely from seeing and meeting kids of
year 11 and 12 or at university who are doing okay for themselves. This can be of particular value when they
are really starting to wonder about friendships, relationships and their future.

Some recommendation include:

- A review of the newborn screening program to ensure a whole of wellbeing approach so there is improved
  social/ emotional support to parents at the point of diagnosis
- Explore the growth of peer group programs, such as Hear For You, teenaged Oral Deaf and signing Deaf peer
  support programs.
Amanda: my son Thomas and his journey with autism

I got married in 2002 to an amazing man who comes from a deaf family. I had my second child (Thomas) in 2004. When Thomas was born he had 3 newborn screening tests. Finally at 10 weeks old they confirmed he was deaf. That red stamp ‘FAILED’ still creeps up in my mind sometimes. I had this nagging feeling inside me that something was not right about Thomas. He was such a difficult baby with feeding, handling and touching and would never sleep. He was always rigid.

We went to a sleep clinic and it was an awful experience, We were looked down as ‘deaf parents’; the nurses’ body language was so easy to read. They scowled at us for signing to Thomas, they scolded at us for not being ‘strict enough’. We didn’t last long there. We had access to an interpreter, however they would not respect us.

Thomas achieved his milestones way earlier than expected; his signing and speech was advanced. I still had that nagging feeling that something wasn’t right. Thomas was diagnosed as having ADHD at 18 months old, however i still felt something wasn’t right. He attended 4 different schools and every single time he would get himself in trouble. I knew there was an explanation but I was told ‘he’s lashing out because he is frustrated with communication’ among other theories. He was suspended many times. I refused to give up and allow anyone to tell me that I had a ‘naughty little deaf boy who was demonstrating typical deaf behaviour’.

We saw 11 psychologists and psychiatrists – each one of them said “He is demonstrating typical deaf behaviour”. It didn’t matter how many times I tell them that I have 2 other deaf children and worked with deaf children and I knew this wasn’t the case. Then we were told it is our ‘parenting skills’- they said that because we are ‘Deaf parents’, he is frustrated because of communication we use (we sign at home). So we did numerous courses on parenting deaf children as well a specialised language program from Germany. We were humiliated each time.

Labelling became a thing to every appointment, meeting and session. I felt I wasn’t good enough to be a mother because everyone was telling us we were the problem not them.

I refused to give up, while seeking an answer I tried everything including the Gap Diet, the Failsafe Diet, behaviour management…. Endless stuff. We were and still are his 24 hour carers.

Finally we tried a new psychiatrist - he said my son is autistic and there is no doubt. He was almost 8.

Since the diagnosis, we were too late to register for any early intervention programs or access the Better Start Funding and I was furious. We had 2 x 3 hours respite in 11 years and no other support or help. I have only met one other deaf child with autism and he is in a special school. I have another friend who is deaf with a deaf autistic son and they live in America. We share our pains via Facebook messages.

Thomas is a kid of many words; he would talk endlessly about dinosaurs but very rarely open his heart. When he does, it kills a little bit of me and breaks my heart more because I did not know how to help him and I didn’t know where to go for help. Thomas completed a survey for the Bonds University to assist their research, he wrote that he wished he was dead.

There is no one in Australia who is experienced in deafness and ASD- especially someone who could sign. Autism QLD rarely contact us to offer support even we are a member and get news but we get no help from them or any strategies.

Today he is in a better emotional situation he is happier and we are working very hard to maintain a safe and calming environment for him at home and at school. I am afraid of what it will be like when he starts high school.
Amanda: My daughter Louise and her journey with the Cochlear Implant

I had a baby girl, Louise, who was born profoundly deaf. My husband Simon asked the ENT specialist on the same day she was diagnosed if she can have a cochlear implant and I was shocked. My initial reaction was no. I had fears that she will lose her ‘Deaf identity’. My husband and I agreed to six months of research. Within me, I had made up my mind and that was a no. Simon's fear was the surgery itself.

We noticed she had issues with her vision, and after testing, were told she had a DVM. My worries about her vision did alter my decision toward the cochlear implant. After a lot of research, we agreed to go ahead with the implant. It worked well for her.

5 years later, Louise came home from school one day said she wanted a second cochlear implant. Shocked, I asked why. She said everyone got two she wanted the same. We said no. We felt she wasn’t mentally prepared for the hard work that an implant at a later age will require. A year later she kept on asking, saying she wanted to hear from her other ‘ear’. We discussed it with her in depth and made the decision to let her go ahead with it.

She struggled with the switch on for 6 months. She hated it. She felt she couldn’t make it work and because we put so much pressure on her to work hard to make it happen, she felt she wasn’t good enough. I started to wish I didn’t allow it to happen because she was too old for the second implant – it's better as a baby or when she was mentally prepared for the hard work it takes. She was neither. Louise struggled so hard with the second implant and after about 8-10 months she finally got used to it. Although she was happy and could settle down, her self-esteem was already low and I felt so responsible for that.

I felt I couldn’t reach out to the Deaf community about my little girl's wellbeing because they still held largely negative views on cochlear implants and would blame that instantly. There was no one in the mental health system who is a deaf person that is able to both speak and sign so we were unable to access the support we needed.

Amandas: my son Kurt, the CODA

My little man Kurt, was a very much wanted baby. I already had three children, and they were all deaf. I secretly hoped that he would be hearing because we had to battle for the right access for the first three. It took medical professionals 7 years to really confirm that Kurt is completely hearing and will not lose any hearing unless something happens in the future.

Kurt always said and still says today that he wished he were deaf. He feels like he missed out because he is the odd one in the family. It doesn’t matter how many times we assure him that everyone is equal; He knows we are different and he knows he ‘hears’ everything when we don’t. He is carrying that little burden on his shoulders as a hearing child in a deaf family and it doesn’t matter how many times I tell him not to, He still feels that is in his life plan.
Positive Psychology of Deaf People

Paul Jacobs has lived with a profound hearing loss since he was five years old. Dr Jacobs' Deaf-specific research has covered mental health, proactive thinking, social skills and advocacy by parents of children who are Deaf and teachers of the Deaf.

Jacobs presented on how Deaf-specific research influences therapeutic and educational practice. Research tends to focus on the negatives and the problems. He argued that this is not helpful because it does little to inform proactive courses of action. Research needs to focus on the positives that demonstrate how people living with deafness can socially and professionally participate. Jacobs believes such research promotes a healthy mental wellbeing.

Jacobs explained that people often perceive the term “Mental Health” as a negative. Mental health means depression, suicidal thoughts, drug abuse, and alcoholism amongst other largely negative illnesses. Mental health can be a positive thing. The word “health” means life. It means enjoyment and happiness. Jacobs spoke about the concept of Positive Psychology, and argued that the focus on negatives can influence poor self perception and there was a need to consider positive psychology as means of promoting positive self-image.

When Jacobs explored this concept, he profiled famous Deaf people and their characteristics. One main theme he found was approach anxiety- the fear of approaching or being approached. It affects people who live with deafness in a big way, as they are unable to predict what will happen next. Deaf people tend to seek out and learn a strategy to enable them to predict what could happen if or when they are approached or they have to approach someone. There is nothing more intimidating to a Deaf person than to be approached randomly by someone talking about something that is out of context.

Jacobs stated that when we talk about mental health and deafness we focus mostly on the medical model and ‘fixing’ deafness and dealing with ‘loss’ of hearing. However, he argued if society just removed barriers to allow Deaf people to participate fully and equally, then there will be a positive change in mental wellbeing.

Research on the effect of being Deaf

Dr Anthony Hogan’s work addresses issues of social identity and wellbeing across a variety of settings. Dr Hogan has completed projects on hearing and social identity, stigma and mental health, disability and employment and the social wellbeing of children with hearing loss.

Hogan has done extensive research on the life effect of being Deaf. He argues that the oppression of Deaf people, due to audism is affecting the lifespan of Deaf people across Australia. He believed that the challenge is the larger society has what he calls “unconscious bias”. This is a way of saying “I realise I’m doing the wrong thing, but I am going to do it anyway”.

He argues that one of the challenges for Deaf individuals is to come to the realization that they need to stop internalizing the oppression, self-blame and feeling anxious, to recognizing that it is actually the other person that needs to change the way they interact with them. The Deaf individual is not at fault; it is the inaccessible environment that is at fault. This internalization leads to people trading in their own identity, who they truly believe they are, for an identity that they think other people want and will accept. This brings on depression, anxiety and all that comes with it.

One of the big problems we have in Australia is that our major intervention for adults is primarily focused on devices; hearing aids and Cochlear Implants. The very first professionals that we deal with from early age with our deafness, and throughout our lifetime, are not equipped to work with us on our psychosocial wellbeing. Your physical or mental wellbeing is your sense of belonging and connectivity, not your level of hearing. Your confidence and ability to manage your hearing loss is closely related to your quality of life.
Whilst we continue as a public to fund audiology services alone, things will not change. The provision of hearing aids and specialist services to children is vital and should still continue. Specialist services for kids will still continue. However, we need to see a model that is inclusive of mental health and whole of wellbeing services.

**Current Mental Health Model in the United Kingdom.**

Dr Brendan Montiero presented on the current mental health model for Deaf people in the UK. Dr Montiero has been a psychiatrist for Deaf people since 1984 and set up three specialist Deaf services for Deaf people.

Rampton Hospital is a high security hospital based in Nottingham. It has 10 beds for Deaf patients in one ward. It has a fence that’s about 5.3m high, which means it has highly secure conditions. Deaf people there get a range of services. There are also medium and low secure services at Saint Andrews in Northampton (developed in 2001 with four beds), Saint Maoris (four beds) and All Saints (six beds). They provide a medium security Deaf specific service. Lastly, St Mary’s has four Deaf beds in a ward of 16 hearing and Deaf patients. The other option is, there are open hearing wards with visiting Deaf services.

Deaf people come in with all different types of communications needs. There are some that speak, some that sign, some that use both. There is a critical phase of language development. If you miss out on the critical phase of language development, which is usually when a child is quite young, then picking up language and the later stage of life is far more difficult. It makes it difficult for such individuals to acquire complete fluency. This is one of the situations that arise for many Deaf patients.

Quite often, problems arise when they leave the security of home, and go into the world, and suddenly they are left lonely and cannot engage properly with the hearing world. Particularly this impacts on forming relationships, getting jobs and further education. There are numerous situations where they may experience discrimination and mental health issues.

In other cases, there are some of causes of deafness that can also cause learning disabilities, including very poor emotional environments and deprivation. In this context, “deprivation” means deprivation of language and communication. It causes a domino effect; if you are deprived of language, and cannot speak to your parents, siblings or peers, then you can’t be educated properly using conventional language systems. When that happens, you wind up not being able to absorb age-appropriate concepts. This makes it difficult to make sense of the world as an adult and participate appropriately.

In terms of literacy, there is plenty of scope for misunderstandings. There is plenty of scope for the treating person to get the construct of the communication wrong and thereby going down a line that a person may think the Deaf person is paranoid. An example of this:

> “Remember Maggie Thatcher? (In 1985) There were lots of cuts and very strict fiscal policies (in UK). Disability was sort of ignored. I saw this guy who came into the outpatient clinic. He said that the government is very cruel. I said, “I agree with you to a certain extent.” He said, “No, they are specifically cruel because they are breaking ducks’ legs and putting them on ice.” I thought that this sounded psychotic. I’m trying to explore this further, and my consultant John Denmark was in the next room. Otherwise he seemed perfectly sane, communicating very well. But he said the government is breaking ducks’ legs and putting them on ice. My consultant said, “Where did you get this from?” He said it was in the newspaper. The headline was: Government puts lame duck policy on ice. So this is the way even somebody who has training in the field, somebody who has some understanding can go around and misunderstand situations.”- Monterio, 17th November 2015

If you look at the assessment of Deaf people and why there is misdiagnosis of intellectual impairment in Deaf people, it’s largely because of public ideas and public awareness. If that person then goes for assessments, and a psychologist who does not have Deaf awareness, or understand sign language, uses intelligence tests are based on verbal tests and performance tests without them understanding it properly, they will underscore because they don’t understand the words. If you give them performance tests they tend to score according to their performance ability. In the UK system, they always look at the performance tests as indicative of their true innate intelligence. Deaf people are often misdiagnosed as being intellectually impaired, and this explains why there is a higher incidence of intellectual impairment in the Deaf population.

- So, how do the doctors in the UK system communicate with Deaf people who are mentally ill?
- At the very first instance, assess their first mode of communication; are they more comfortable hearing/speaking
Do not ask vague questions about anything that you can’t make sense of. Ask very specific questions about: how is it affecting you? Why are you angry? Why are you upset?

Often you do not get an appropriate time line with particularly minimal Sign Language using Deaf people.

Clarify name signs right from the beginning.

Use a qualified interpreter; often there are times where something might not make sense in the context, however interpreters are qualified to pick this up. This could mean the difference in diagnosing someone as having a mental illness or just being misunderstood.

Always check understanding because deaf people are known to nod even when they don’t understand.

Dr Montero mentioned that it is also critical to understand the family dynamic. There are instances where families reject or try to overprotect or try to blame. The Deaf person has different relationships with peers and all that needs to be explored. Sometimes what appears like a mental health problem like anger, aggression, acting out might be because the person is upset and does not have an emotional vocabulary to describe it i.e. “I was upset with my brother because he took my X-Box and played with it and I wanted it and couldn’t get it so I punched him.”

Assessments on Deaf people at an intellectual level needs to be done by psychologists who are Deaf aware and can communicate using Sign Language. When you are working with Deaf people who have minimal Sign Language, you also need Deaf relay interpreters. Deaf relay interpreters assist to break the language down even further.

Deaf people suffer from the same range of problems as hearing people. However, maladjustment, anxiety and depression, stress related issues are far higher in the Deaf population compared to the hearing population. This is because their life can be far more stressful. Anything that is likely to be environmentally determined; stress, difficulties of adjustment, discrimination, problems of finding employment, problems in relationships, is likely to be higher in the Deaf population than the hearing population.

When doing risk assessments of Deaf people it is important that a number of other issues are taken into account. When you do formal risk assessments on Deaf people, there are items like the history of early employment. Deaf people might not have the employment, history, however you can’t put that down as a risk factor. There are also items such as history of anger problems and temper tantrums. You can’t put that down as risk factor because it might be part of their developmental phase. So there are a number of things that you have to be cautious about and this has to be reflected in the report in order to make an accurate assessment

In Deaf people, generally, personality disorders and behavioral adjustment problems are increased. That’s not surprising because many of the environmental factors are stressors and are factors that can be naturally difficult for Deaf people. Their personalities can take a slightly different pathway. They can appear antisocial, paranoid and appear to be isolated and uncooperative. The difference between a Deaf person with those personality characteristics and a hearing person with those personality characteristics is that the Deaf person is often treatable. Often it’s about their personality changing because of environmental issues. You can bring about change by changing the environmental barriers, however in hearing people, who have had all the opportunities, it is deeply ingrained in their personalities and it is very difficult to change.

In a Deaf treatment model, Dr Montero talked about the “habilitation” model that is used, rather than a ‘rehabilitation’ model. Habilitation is a concept of developing skills in people that are lacking. Rehabilitation is a concept where you have lost the skills because of something and we teach you to regain the skills. In the UK model, most Deaf people came in without the skills. Their personalities can take a slightly different pathway. They can appear antisocial, paranoid and appear to be isolated and uncooperative. The difference between a Deaf person with those personality characteristics and a hearing person with those personality characteristics is that the Deaf person is often treatable. Often it’s about their personality changing because of environmental issues. You can bring about change by changing the environmental barriers, however in hearing people, who have had all the opportunities, it is deeply ingrained in their personalities and it is very difficult to change.

To do this, Dr Montiero says that first they need to educate the patients about terms, the meaning of signs, their knowledge of the world, etc. Then move into specific types of management - i.e.: anger, trauma, substance abuse, sex abuse, violence etc. Then you move into recovery program and relapse. But the important thing is, these stages are not fixed. They can move up and down between stages. So you might get to recovery and then they don’t understand something about sex offences so you come back to the
management. All assessments are completed in the Deaf persons preferred language and at their level of communication. The staff all use sign language to some degree, however not everyone is absolutely fluent. The program uses cognitive behavior therapy approaches and psycho education. All of these programs, sex offender treatment, group or individual, substance misuse, they are adapted for Deaf people.

As mentioned previously in the Audism chapter, the treatment of Deaf people costs more and takes longer. In the UK the Government has sex offender programs that need to be completed within nine months to a year. For Deaf people the program can last between three and five years.

The environment is very important. The UK Services are run from a Deaf cultural perspective. You don’t just need hearing psychiatrists and professionals to come in with Sign Language interpreters if you have a Deaf service. It’s very important that staff can sign and have good Deaf awareness. In the UK system, all staff have an induction course of a week of Deaf awareness as well as varying levels of sign language training. There’s a signing policy in the hospital, with a fine that goes to a Deaf charity if you don’t use signs or if hearing people speak to each other in corridor. You need interpreters with expertise in mental health, because there are particular signs that can be used or to explain a concept and need expert translation. However it’s still important that the environment has people signing.

The UK Government often says “Why don’t we send them to a hearing hospital with Sign Language interpreters?” That’s what we call a pseudo team. What you have got is a range of professionals who understand nothing about deafness, who have no idea about the range of problems they are going to have and you are putting in an interpreter and expecting the interpreter to solve it all. That gives the impression that you are treating the person. It actually does not mean real treatment.

A real team is like the team you can have in a Deaf service where everybody working has an understanding of deafness, can communicate at a particular level, can understand the variations of minimal Sign Language to highly eloquent Sign Language and can get into the patient’s individual experience. A highly effective team, that’s been recognised by the UK government and they have funded it for the past 22 years. Three services have been developed in that time and a lot of lessons have been learnt.

Guidelines for working with Auslan Interpreters- Ryan Tuema

Dr Ryan Tuema is a psychologist who established a private practice called Vitalogy Consulting, devoted to the Deaf and deafblind community. He regularly works with Victorian Deaf Society, Queensland Deaf Society, Deaf Association New Zealand and Able Australia. He has spent the last 4.5 years working at the Victorian College for the Deaf in collaboration with Deaf Children Australia and the Victorian Deaf Education Institute.

Using his knowledge he developed a list of Best Practice Principles for mental health clinicians when working with Level 3 NAATI qualified interpreters

Strategies & Guidelines for effective communication in forensic interviews with Deaf clients: For assessing Mental Health clinicians:

a. Talk to the interpreter before you meet in-person about your experience or lack of experience of working with interpreters. Ask the interpreter about their experience of facilitating mental health assessments.

b. Request all previous, if any, psychological, psychiatric, correctional and educational assessments.

c. With the clients consent, talk to relevant caseworkers, doctors, teachers, parents or extended family members to gain as much information as possible about the clients general functioning and communication strengths and weakness.

d. Be aware that most psychometric assessment tools are not standardized for deaf populations. To avoid misdiagnosis, liaise with your local Deaf society about ways to ensure your testing is relevant and accessible for the client, and conduct a literature search on psychological assessments of deaf persons.

e. On the day of assessment meet with the interpreter before hand and discuss the nature of the assessment, topics to be covered, the language used and potential risks and sensitive information.

f. Set the room up so that you, the interpreter and the client have clear sight of each other. Ensure that the
room is well-lit, private from onlookers, has good ventilation, comfortable chairs and a table at a height that is practical for all concerned.

g. Do not shout at the client; be aware of your facial gesture at all times. Speak clearly and at an even pace. Regularly check in with the interpreter to see if you are speaking too fast or too slow. Pay attention to your pitch and choice of language as there are many words in the discipline of psychology that do not have an Auslan equivalent. However, having said this, a competent interpreter will bridge the gap by interpreting the meaning or intent of what is being spoken where necessary.

h. Ensure that you explore with the client their experience of a mental health assessment. Take time to build rapport and trust. Explain slowly and clearly what your role is and what you want to achieve. Explain why you have been asked to provide a psychological assessment. Ask the client if they have any questions before you start getting personal.

i. Explain to the client your experience in working with deaf people. If it is extensive this will greatly add to the client’s sense of ease and comfort. If it is limited be upfront about it. Ask the client to assist the process by seeking clarification and providing advice as to where you can improve.

j. It is essential to conduct a basic language assessment and explore how they best communicate such as sign language, lip reading, use of hearing aids, gesture, reading, or are they oral (using whatever intelligible speech they have.) If they wear glasses, ensure they are wearing them during the assessment. If they are wearing hearing aids, check if they have been adjusted to suit the environment.

k. Explore the client’s perception of their deaf identity and sense of enmeshment to the deaf culture. It is not uncommon for a deaf person to be uncomfortable about their deafness. When conducting a genogram ask if they are the only deaf person in their family. Explore who are their Deaf role models. Explore whether other family sign fluently. Explore how they communicate with their family members who are not deaf.

l. Explore their education in detail. It is not uncommon for a deaf child to be placed in mainstream schools with hearing children. Ask whether they were subject to bullying or ridicule because of their deafness. Explore their social circle and if they have deaf friends. Where did they go to school and what was their relationship like with their teachers and who helped them with their homework.

m. If the client does not respond to a question or seems to be having difficult comprehending, try to rephrase the message, instead of repeating it exactly.

n. Avoid abrupt topic changes and explain when you are changing the topic or the focus of inquiry.

o. Use visual aids wherever possible. Have a large notebook and thick markers so you can write down key words to guide your inquiry. If the client cannot read at all, draw a picture in simple style. There are a number of pictorial based cards that depict emotions, feelings, event, and behaviors, which can be found on the Internet. These learning aids are concrete, clear and highly visual.

p. Schedule breaks every 30 minutes and check in with the interpreter on how the client comprehends the information. Discuss with the interpreter if your pitch and vocabulary is too high or too low or if you are talking too fast or slow. Explore with the interpreter if they felt comfortable with the level of detail and if there were any confusing or ambiguous questions or responses.

q. If it is a forensics interview, it is sometimes useful to explore details of an offence after you have gathered information about social and development background. Deaf culture and identity, psychosexual development, and drug and alcohol history, suicide and self harm behaviors, issues with aggression and violence for example. This allows the client to feel like they are being understood as a person not as a criminal. On occasion, the client may not want to start the interview with the offence. In this situation be guided by the client and then return to the offence after you have gathered all relevant background information to clarify specific details.

r. Learn about a healthy psychological deaf presentation in contrast to a health psychological hearing presentation. Generally it is uncommon for hearing practitioners like us to be exposed to a healthy deaf psychological presentation, simple because of deaf population is a minority in Australia. This exposure could occur through contact with various deaf conferences and workshops provided in your state deaf society or deaf organization. Learn about possible Deaf adolescent mental health issues for example: Loneliness.
as a result of being left out of social interactions; Social isolation, Inferiority and feeling like a lesser person than hearing people; Frustration from not being understood or listened too; Aggressive behaviors; Fearful of rejection; Low self esteem; Shame resulting from being taught to behave and act like a hearing child; Depression and despair that their life will always be a struggle; Trauma from sexual abuse, rape, bulling and ridicule; Poor self expression from overly controlling parents; Sexual development problems from delayed, confused and poor sex education and deaf gay/lesbian issues; Suicide and self injury – very hard to spot warning signals and potential for high incidence.

s. When formulating a diagnosis for a client for example: Depression, Obsessive Compulsive Disorder, Post Traumatic Stress Disorder; do not rely on the DSM-IV (Diagnostic and Statistical Manual for Mental Disorders) as it is culturally biased towards the western hearing population. Misdiagnosis can lead to a number of problems including an inappropriate treatment plan, inaccurate medication regime, client confusion and frustration, and further traumatizing the client. Inaccurate forensics assessments have the potential to significantly disadvantage the deaf client. For example: the over estimation of risk of re-offending, the over estimation of dangerousness to the community, they may be detained in higher security cells unnecessary, and held imprisoned longer than is necessary.

t. Learn some basic signs such as “Hello how are you?” “Would you like a break?” “toilet break” for example. This will demonstrate to the client that you respect and have taken a basic step to understand their language. If you make a mistake be prepared to laugh at yourself and learn from the client and the interpreter (Farrugia, 1988)
Recommendations

Recommendation one – Development of a Community Based Support model for deaf mental health:

It was clear from many speakers that support for people who are deaf in the current mental health system in Australia is lacking. Systems are designed for people who are hearing and adjustments are not readily made for people who are deaf who need mental health support.

It is recommended that:

1. Research be conducted to identify “best practice” community based support models for deaf mental health support throughout the world.
2. Continuing consultation occur with key stakeholders in Australia to identify needs and improvements required for deaf mental health support.
3. From data collected a report be drafted with key recommendations for the development of a community based response to Deaf mental health in Australia.

Recommendation 2 – Development of training and support to increase the skills and awareness in working with deaf people within the mental health system for communication support professionals such as interpreters, relay interpreters, deafblind interpreters etc.

It is clear that interpreting and communication support for Deaf mental health support is a specialized field. Training needs to be developed around this to increase skills and awareness of communication support people to work with deaf people receiving mental health support.

Recommendation 3 – Development of accessible resources for people who are deaf who are accessing the mental health support system.

It is clear that many of the resources, both printed and online are not accessible for people who are deaf. Plain English versions and Auslan versions of printed material are required. Online videos need to be captioned and Auslan versions developed. This needs to be developed across the spectrum of mental health support including, but not limited to:

1. Domestic violence
2. Substance abuse
3. General mental health information such as Beyond Blue websites and printed information.
Recommendation 4 – Training and awareness programs be developed for mainstream mental health professionals such as psychiatrists, counselors, nurses and other support people to increase awareness of specific issues surrounding mental health support for people who are deaf.

It is clear that there is a lack of awareness among professionals surrounding specific issues of the deaf populace in regard to mental health support. This needs to be addressed through the development of training and resources to increase awareness of the requirements of people who are deaf accessing mental health support.

Recommendation 5  - Development of programs that focus on prevention through the development of positive mental health activities and strategies.

Evidence was presented that suggested that deaf people are subject to many negative stereotyped that focus on ‘cures’ and fixing. This can lead to the development of poor self-concept.

It is recommended that programs be developed that focus on developing “positive life skills” through exposure to strong role models.