‘People who have experience of using the system and who have done some thinking about this and know other people using the system can offer sensible advice about how to make it better. It’s about being experts by experience.’
The author would like to thank the members of the Research Reference Group, Priscilla Berkery (Tasmanian Mental Health Consumer Network), Jane Carlson (Anglicare), Kerrie Dissegna (Anglicare), Jo Flanagan (Anglicare), Gwyn Jones (Tasmanian Mental Health Consumer Network), Kath McLean (TasCOSS), Lynette Pearce (Mental Health Services) and Jeanette Sherington (Tasmanian Mental Health Consumer Network) for their advice on the development of the research and the recommendations which have grown from it. The author would particularly like to thank Gwyn Jones for writing the original proposal that led that led to this research and to Mental Health Services for their input and collaboration.

The research findings, conclusions and recommendations of this report are those of Anglicare and should not be attributed to any members of the reference group. Any errors in the report are the responsibility of the author.

The author would also like to thank all the mental health consumers from across Australia and internationally who gave up their time to participate in the research, to talk about their experiences and to freely express their views. Without them this research would not have been possible.

The title of this report, ‘Experts by Experience’, reflects the belief of the Research Reference Group that the ‘lived experience’ of mental illness has a vital role to play in the reform of mental health services. The title also reflects a concern to ensure that any development of mental health consumer activity in the state draws on the experiences of consumers across Australia and overseas.
Abbreviations

ABS       Australian Bureau of Statistics
ACHS      Australian Council on Healthcare Standards
ACROD     now known as National Disability Services
ACT       Australian Capital Territory
ACTCOSS   ACT Council of Social Service
AMHCN     Australian Mental Health Consumer Network
AMHWA     Association of Mental Health Consumers, WA
ARAFMI     Association of Relatives and Friends of the Mentally Ill
AusNet     Australian Network for Promotion, Prevention and Early Intervention for Mental Health
ACTMHCN   Australian Capital Territory Mental Health Consumer Network
CAAC      Consumer Affairs Advisory Council
CAG       Consumer Advisory Group or Consumer and Carer Advisory Group
CAGWA     Consumer Advisory Group, WA
CALD      Culturally and Linguistically Diverse
CAN       Consumer Activity Network, NSW
CDI       Consumer Developed Initiative
COAG      Council of Australian Governments
COMHWA    Consumer Organisation Mental Health, WA
CORE      Consumer Operated Research and Evaluation Support Group, VMIAC
CSO       Community Service Organisation
DohIFS    Commonwealth Department of Health and Family Services (now known as DoHA and FaHCSIA)
DHS       Department of Human Services, Victoria
DHHS      Department of Health and Human Services, Tasmania
DoHA      Commonwealth Department of Health and Ageing
FaHCSIA   Department of Families, Housing, Community Services and Indigenous Affairs
FTE       Full Time Equivalent
HCA       Health Consumers Alliance, SA
HCC       Health Consumers Council, WA
IMHIL     International Initiative for Mental Health Leadership
MHAC      Mental Health Advocacy Coalition, New Zealand
MHACA     Mental Health Association of Central Australia
MHAL      Mental Health Activity and Learning Centre, WA
MH-CoPES  Mental Health Consumer Perceptions and Experiences of Services
MHCA      Mental Health Council of Australia
MHCT      Mental Health Council of Tasmania
NCAGMH    National Community Advisory Group on Mental Health
NGO       Non-Government Organisation
NHS       National Health Service, UK
NMHCAG    National Mental Health Consumer and Carer Forum (previously NCCF, National Consumer and Carer Forum)
NMHETAG   National Mental Health Education and Training Advisory Group
NMHS      National Mental Health Strategy
NSW       New South Wales
NSWCAG    New South Wales Consumer Advisory Group
NT        Northern Territory
NTCAGMH   Northern Territory Community Advisory Group for Mental Health
NZ        New Zealand
PhaMs     Personal Helpers and Mentors Program
Qld       Queensland
QldCAG    Queensland Consumer and Carer Advisory Group
SA        South Australia
SACAG     South Australia Consumer and Carer Advisory Group
SARC      Social Action and Research Centre, Anglicare Tasmania
SIU       Social Inclusion Unit, South Australia
TASCAG    Independent Ministerial Advisory Committee on Mental Health, Tasmania
TasCOSS   Tasmanian Council of Social Service
TMHCN     Tasmania Mental Health Consumer Network
U & I     Understanding and Involvement Project
VMIAC     Victoria Mental Illness Awareness Council
WA        Western Australia
Executive Summary and Recommendations

This research explores the achievements and struggles of the mental health consumer movement across Australia in order to inform the development of consumer activities in one state, Tasmania. Using information collated from a literature review and one-to-one interviews with over 70 consumers and other stakeholders it maps consumer initiatives and activities nationally, describes the key themes and issues mental health consumers face both in Australia and overseas and presents some options for Tasmania.

The research demonstrates that both nationally and internationally the mental health consumer movement can claim a number of successes. Its two key aims – to transform mental health provision into a recovery-orientated service which is about wellness rather than illness and to ensure consumer participation is an accepted and routine part of service delivery and evaluation, policy and planning – are now accepted goals for those making decisions about mental health services. In Australia a recovery focus has become a cornerstone of mental health policy and mental health services are required to promote the participation of consumers at all levels. Twenty years ago consumers were excluded, their experiences of services were not valued and they were not involved in debate. Today the consumer movement is established, accepted and seen as being a good thing.

Across Australia the research found a complex jigsaw of consumer activities and participation mechanisms. These range from small consumer support and self-help groups and involvement in decisions about treatment through to developing a paid consumer workforce, consumer advocacy organisations and consumer-run services. The report identifies two major strategies: working within the system to promote cultural change and working outside the system in independent consumer organisations to advocate for systemic change and to role model alternative services. Each jurisdiction is different but most have witnessed a burgeoning of consumer initiatives on the ground accompanied by a push to develop state wide and national consumer run peaks and a consumer workforce. Victoria in particular is identified as demonstrating a positive model of mental health consumer involvement in public mental health services and in sustaining an independent consumer-run peak body.

However, the research also found that for many consumers and other stakeholders the consumer movement has not reached its full potential. Participation and advocacy activities are patchy, funding and resourcing are often inadequate and consumer run organisations and services struggle to survive. In many places there has been a failure to translate support for consumer initiatives and participation mechanisms into the financial resources and the capacity building required to make them work effectively. Some of the key issues for the movement identified in the literature review are about how best to promote a recovery agenda so that it becomes a reality for mental health consumers and how to progress the cultural change necessary to remove the barriers to effective participation in clinical services. There are also issues about the most effective way to develop a consumer workforce, the pros and cons of working with carers, the meaning of consumer leadership and representation and the establishment and sustainability of consumer-run organisations.

These issues are reflected in Tasmania where, despite having active and influential consumers, there have been low levels of consumer activity. The state now lags behind other jurisdictions. The main mechanism for consumer and carer participation has been TASCAG – an independent ministerial
advisory committee on mental health. In addition there have been two key consumer initiatives – a consumer consultancy project which began to develop a consumer workforce in the south of the state and the Tasmanian Mental Health Consumer Network which, for the past three years, has performed an important role in advocating for system improvement and improved community attitudes and worked to strengthen the mental health consumer community. There has also been a growth in small consumer support and self-help groups across the state. Nevertheless, despite these developments Tasmania’s small and dispersed population and a lack of financial and other resources have mitigated against state wide consumer activity and involvement and raised concerns about the sustainability of consumer-run initiatives. This has resulted in an on-going debate about how best to promote the consumer voice so that it informs policy, planning and service delivery. It has also resulted in an acknowledgement that there are no easy solutions and that there are different perspectives about how these issues should be addressed.

The research clearly demonstrates that the difficulties faced by Tasmanian consumers are not unique, that many mental health consumers and indeed many community organisations and movements have shared these struggles and that, although there is no ‘road map’, there are valuable lessons to be learnt from experiences in other jurisdictions and internationally. The research makes three major recommendations in order to improve the situation in Tasmania. It recommends the establishment of an independent and sustainable consumer-run organisation, the building of a consumer workforce and the creation of an appropriate supporting infrastructure.

**Recommendations**

**Recommendation 1:** That the Department of Health and Human Services provide funding to establish a state wide mental health consumer run organisation in Tasmania.

**Recommended 2:** That the Department of Health and Human Services support and facilitate the development of a consumer workforce in Tasmania in collaboration with mental health consumers.

**Recommendation 3:** That the Department of Health and Human Services use funding already allocated through the *Bridging the Gap* report to immediately recruit three regional consumer liaison workers to oversee the implementation of participation mechanisms and to build the capacity of consumers and carers to participate and of mental health providers to facilitate that participation.

**Recommendation 4:** That both public mental health services and those operated by CSOs implement positive discrimination practices in recruitment to mental health programs towards those with a lived experience of mental health problems.

**Recommendation 5:** That the Department of Health and Human Services provide opportunities for the training of consumers in order to build the capacity of the consumer movement.

**Recommendation 6:** That the Department of Health and Human Services make the consumer perspective intrinsic to all aspects of the education and training of the mental health service workforce through using consumer educators.

**Recommendation 7:** That the University of Tasmania through the Pro Vice Chancellor, Centre for the Advancement of Learning and Teaching, ensures that the consumer perspective and the use of consumer educators is embedded in teaching related to mental health issues.

**Recommendation 8:** That the Department of Health and Human Services sponsor a dedicated mental health consumer research fellowship at the University of Tasmania.

**Recommendation 9:** That the Department of Health and Human Services ensure mechanisms are available to develop and nurture consumer support and self-help groups across the state.

**Recommendation 10:** That the Department of Health and Human Services strengthen consumer engagement with Tasmania’s health and human service system by establishing a peak body to represent consumers and the community, to support engagement activities and to strengthen consumer networks and organisations.
**Recommendation 11:** That the Social Inclusion Commissioner advise the Premier to direct the Tasmanian Government’s Social Inclusion Unit to explore the civic participation of marginalised groups as a goal of the Tasmanian Social Inclusion Strategy.

**Recommendation 12:** That the Department of Health and Human Services ensure that performance indicators and outcome measures are developed and monitored for any consumer participation mechanisms or consumer initiatives in both public mental health services and in CSO provision and that this is undertaken in partnership with consumers and carers.

**Recommendation 13:** That the Department of Health and Human Services ensure that a requirement to involve consumers and/or to employ consumer workers is built into the contractual requirements for CSOs providing mental health services.

**Recommendation 14:** That CSOs delivering mental health services ensure that the existence of consumer engagement mechanisms are a key quality indicator in their organisation.
1. Introduction

1.1 Background

People with mental illness have battled for the past two decades to have their voice heard and their experiences used to drive reform of the mental health system. As a result consumer participation in mental health services has now been championed internationally for well over ten years and in many countries, including Australia, it has become the cornerstone of mental health policy (Steward et al. 2008). Consumer activities are seen as highly beneficial in offering opportunities to improve mental health services, tackle negative community attitudes, promote a better quality of life for consumers and assist in the recovery process. Indeed some would say that consumer activities are the key to transforming mental health services into a recovery-orientated service where wellness rather than responding solely to illness becomes the goal.

These perceived benefits have been endorsed in the Australian National Mental Health Strategy and there is now a clear requirement that mental health services should encourage the participation of consumers in the development, implementation, delivery and evaluation of services (Happell 2006). This has resulted in a range of activities from active participation at the micro level of individual decision making about treatment to more macro level involvement in service planning, evaluation, training and research. At one end of this spectrum are consumer driven and controlled advocacy services which offer leadership by and with people who share the experience of what it means to be a consumer. A number of jurisdictions in Australia now have independent mental health consumer organisations which are managed and governed by consumers and which oversee the development of the mental health consumer community. At the other end of the spectrum are participation activities, controlled by non-consumer stakeholders, where opportunities are made available for consumers to become involved in areas ranging from individual treatment plans through to decisions about policy and services. One manifestation of this is the employment of consumer consultants within mental health services to provide a consumer perspective.

Yet despite this significant political encouragement in reality consumer participation and advocacy in mental health services remains patchy, limited and under-funded. These difficulties are not exclusive to Australia but are global where there are concerns that mental health consumer involvement has not developed to its full potential. A recent Australian Senate inquiry (Standing Committee on Community Affairs 2008) highlighted the lack of priority given to the consumer voice in formulating policy and implementing programs. The inquiry emphasised that, despite hundreds of initiatives and projects across Australia, support for consumer advocacy, training, peer support and consumer-run services had yet to translate into resources and the capacity building required to assist consumers in these roles.

Compared to the rest of Australia Tasmania has seen lower levels of consumer activity and, despite a number of active and influential consumers, the consumer movement has struggled to make an impact. Some of the work has been undertaken in an embryonic form by the Tasmanian Mental Health Consumer Network, an organisation of consumers, for consumers. However, despite the existence of the Network and the formulation of a Consumer and Carer Participation Framework (DHHS 2006), there remains a dearth of activity and an ongoing debate about how best to encourage and support consumer participation.
to ensure that the consumer perspective is integrated into service development and delivery across the state.

This report contributes to that debate by examining the spectrum of consumer initiatives and experience both nationally and internationally, and exploring what lessons can be learnt for developing consumer initiatives in Tasmania.

1.2 Definitions

The terms consumer, consumer movement, consumer community, consumer participation and consumer advocacy are complex and multi-dimensional and require some clarity.

In Australia the most common term for someone who has used, is using or might use mental health services (and other health and human services) is a consumer (McInerney 2008). In some contexts it can be used to refer to both consumers and carers of those with mental illness. However in this report it refers solely to users of mental health services, not to carers. It is a useful term because it can be aligned with the wider consumer rights movement and accompanying expectations around rights and protection, being treated with respect and dignity, having regulations and standards and having an entitlement to redress for misrepresentation and poor quality services. However some question the use of a word which suggests choice and purchasing power and has connotations about taking rather than giving. For these reasons the terms service user, survivor, or patient/ex-patient are used in other countries. Advocates of the term survivor in the USA and some parts of Europe believe that it portrays a positive image of those who have had the strength to survive the mental health system.

The mental health consumer movement, also known as the user or survivor movement, is a diverse association of individuals and organisations, small local groups and national networks which campaigns for improved services and consumer-led alternatives. It developed during the 1970s and 1980s alongside the women's movement, disability movement and gay rights movement and internationally its catchphrase, like the disability movement, is 'nothing about us without us'. Some parts of the movement are anti-psychiatry so their objectives are to empower individuals through mutual support and create alternatives to the mental health service system. Others align themselves with the civil rights movement and emphasise a role in lobbying for the right to be part of decision making processes. This results in activities initiated by organisations or citizens to promote the right to participate. The largest part of the movement is consumerist and would like to influence the mental health system by getting inside it through consumer participation and involvement mechanisms, on boards and advisory committees or by obtaining employment in the system. Some consumers question whether the movement has become stronger in recent years or whether it has lost its focus and been pulled too much towards the agenda of service providers. However, both nationally and internationally, there is a general agreement that there are a growing number of consumer groups involved in a diverse range of activities and that consumers are increasingly working with professionals.

The term consumer community is problematic when applied to what is a heterogeneous and diverse population. Firstly, unlike other illnesses like cancer, there is not necessarily a solidarity between consumers who may not have much in common apart from difficulties in accessing appropriate services. For instance, someone with depression may hold the same prejudices as the general population about psychosis or personality disorder. There can also be tensions between well and ill consumers where judgements are made about whether people are sick or debilitated enough to be called consumers. There are those who use the public mental health system and those who use private services, those who use hospital services and those who have only used community-based services. This makes setting up structures to work across the mental illness community problematic. Secondly, there are many more people with mental illness than those who have contact with the service system and some may not even be seeking services. Almost two thirds of those with mental illness (ABS 2008) do not see a professional and some may rely entirely on their family or self help groups. This raises the issue of how to engage people with mental illness who do not access services. The vast majority of consumers remain unorganised and unconnected which presents a major challenge to the consumer movement as well as an opportunity.

Consumer participation can mean different things to different people which has led to confusion. It is variously defined as either voluntary or paid participation by consumers in all aspects of mental health care in order to improve the quality of services. This includes involvement in individual treatment
plans through to local, regional and state planning and policy decision making. Types of participation or engagement have been represented as a ladder or hierarchy moving towards an ideal goal of full engagement. More recently engagement has been described as a continuum offering sets of choices for policy makers (Health Canada 2000) and shaped by the policy problem and the resources available.

**Consumer advocacy** is the mechanism used to empower consumers and provide a consumer voice. There are several different types (Meagher 2002a) ranging from self advocacy where an individual or a group speaks for itself, individual advocacy where an advocate represents the perspective of an individual through to system advocacy where advocacy activities are directed towards the systems and structures which create injustice and inequity.

### 1.3 The Prevalence of Mental Health Problems

There are problems in estimating the number of people with mental illness (Cameron & Flanagan 2004) particularly as many people do not access mental health services.

The latest estimates come from the National Survey of Mental Health and Wellbeing (ABS 2008). They show that one in five Australians aged 16-85 had a mental disorder in 2007 and almost half or 45% will have a mental disorder at some point in their life. At any one time serious mental illness affects 3% of the adult population and a further 17% will experience problems in the next 12 months. Anxiety disorders like post-traumatic stress disorder and panic disorder are the most common and affect 14% of people. Affective disorders like depression affect 6%. However only two out of every five Australians with a mental health problem access services.

This means that the mental health community in Tasmania consists of approximately 60,000 people experiencing mild, moderate and severe disorders but with only about 24,000 people in touch with a mental health professional.

### 1.4 National Policy Framework

Since 1992 consumer (and carer) participation in mental health service development and delivery has been embedded in Australian policy frameworks at both a federal and state level. These policy frameworks are:

- **1991 Mental Health: statement of rights and responsibilities** (NMHS 1991). This aimed to ensure that consumers, carers, advocates, service providers and the community were aware of their rights and responsibilities and confident in exercising them. It followed the adoption of the United Nations principles for the protection of persons with mental illness (United Nations 1991) and included the right of consumers to live in the community and contribute to and participate as far as possible in the development of mental health policy, the provision of mental health care and the representation of mental health consumer interests.

- **Three National Mental Health Plans**. The first Plan (Australian Health Ministers 1992) officially recognised consumers as having a vital role in the mental health services system and that consumer and carer input is essential if improvements in service delivery are to be achieved. This commitment was operationalised at national and state levels by establishing a committee of consumers and carers – the National Community Advisory Group on Mental Health (NCAGMH) and state and territory based CAGs. The Second National Mental Health Plan (Australian Health Ministers 1998) reinforced this commitment. However by the third Plan (Australian Health Ministers 2003), although consumer and carer participation at all levels in policy, planning and treatment was identified as ‘the hallmark of a quality mental health system’, the Plan also stated that ‘participation in service planning and delivery across the spectrum of care from promotion and prevention to recovery has not yet been achieved’ (p. 24). A fourth plan is currently being drafted.

- **In 1997 the National Standards for Mental Health Services** (Australian Health Ministers 1997) were published as the key mechanism for assuring quality in mental health services and with a commitment from all jurisdictions to implement them. The Standards include the requirement for mental health services to have policies and procedures relating to consumer and carer participation and to maximise their roles and involvement. Implementation involves in-depth review of services against the standards by an external accreditation body.
The National Practice Standards for the Mental Health Workforce (NMHETAG 2002) were published in 2002. These Standards specify how services should involve consumers as active participants in their own care and in planning and evaluation and one of their key underpinning principles is that mental health professionals should be educated by consumers.

- The 2006-2011 Council of Australian Governments National Action Plan on Mental Health (COAG 2006) aimed to deliver a more seamless and connected care system. However, the Plan has been criticised for being a list of initiatives and programs rather than a vision for the future and for a lack of clarity about how it fits with the National Mental Health Plan. It is also criticised for not addressing how participation in state and service policy development and delivery should be addressed and for not prioritising a consumer and carer focus.

- Participation policy frameworks across health and human services parallel developments in the mental health sector and aim to place health consumers at the centre of health policy. A recent report (McInerney 2008) outlines some of these initiatives. The Australian Institute of Health Policy Studies is investigating current practice and developing new models for engagement in health policy. The Victorian Department of Human Services has produced a number of documents to promote participation as an essential principle of health development, community capacity building and development of social capital. The Australian Commission on Safety and Quality in Health Care is consulting on the development of a consumer engagement strategy and the Consumers’ Health Forum of Australia is working to involve consumers in health policy at a national level.

- The recently established national social inclusion agenda is also pushing for meaningful participation for people with mental health problems. Social inclusion is seen as an essential component of recovery which should be fostered through ensuring consumers, their families and carers have ongoing input into service delivery and planning. In addition they should have access to services tailored to individual needs and practical assistance with reintegration and there should be an increasing awareness of and a reduction in stigma in the community. In the UK consumers are beginning to articulate a model of social inclusion which does not mean fitting in and passing as normal but is about including the experience of madness as part of the social spectrum and recognising its positive aspects as well as the pain involved (Sayce 2000).

The National Mental Health Reports which monitor progress in mental health reform under the National Plans have identified that there have been substantial gains in consumer participation. The 2005 report (DoHA 2005) showed that the proportion of mental health organisations with a formal mechanism for consumer participation had increased from 53% in 1994 to 82% in 2003. Yet this also meant that 18% of mental health service organisations lacked a basic structural agreement for consumer and carer participation. It also found that the total expenditure on consumer and carer consultants was only a fraction of spending on wages in the mental health sector. The 2007 report (DoHA 2007) provided a summary of twelve years of reform in mental health services. It showed that there are pockets of good practice including an acceptance that consumer representatives should be remunerated and also included in all national planning groups. But progress has been uneven across jurisdictions and there is also an acknowledgement that there is a long way to go before satisfactory levels of consumer and carer participation are achieved. There is also confusion about how to transform theories of engagement into successful practice (McInerney 2008).

An inquiry into mental health services in Australia (Standing Committee on Community Affairs 2008) recommended that the Australian Government strengthen mental health consumer representation through funding consumer run organisations to provide independent advocacy at state, territory and Commonwealth levels. In particular it highlighted that awareness of the importance of consumer participation was not matched by funding and support to actually facilitate such involvement and that there were very few opportunities for consumers to develop the skills to be effective advocates.

1.5 Aims of the Research

This research aimed to formulate an evidence based model of effective consumer activity in order to progress the development
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of the mental health consumer community in Tasmania. More specifically it aimed to piece together the complex tapestry of consumer activities and initiatives across Australia and internationally, explore how this jigsaw fits together and draw out the lessons for development in Tasmania. It entailed:

- conducting a literature search of interstate and overseas models of mental health consumer community development;
- conducting a broad based consultation with mental health consumers and other stakeholders across Australia and internationally; and
- outlining possibilities for the development of models of mental health consumer community development for Tasmania.

The research was initiated by the Tasmanian Mental Health Consumer Network (TMHCN) and undertaken by the Social Action and Research Centre (SARC) at Anglicare Tasmania in partnership with the TMHCN. The work took place over a six month period between October 2008 and April 2009. It was not anticipated that the research would provide all the answers but that it would offer guidance for developments to be pursued by both mental health consumers and providers.

1.6 Research Methods

Globally there is a wealth of information and experience about mental health consumer activities and a growing literature. The research was designed to be able to draw on these experiences, on the successes and failures and on the lessons that can be learnt from them to inform developments in Tasmania. To this end the research proceeded in three stages:

- **collation of literature and documentation** locally, nationally and internationally about developing mental health consumer communities. This entailed conducting a step-by-step search for material on mental health consumer activities and participation through the internet and a range of other information sources including libraries, databases, government documents, conference proceedings and the ‘grey’ literature produced by organisations but not necessarily formally published.

- **interviews with key informants** in Tasmania, other states and territories and overseas to explore experiences of developing the mental health consumer community and map both consumer engagement processes and consumer run initiatives. Key informants were identified using a snow-balling approach whereby interviewees were asked to nominate further useful contacts. This ensured that the researcher was able to speak with key figures in each jurisdiction and tap into a range of perspectives.

In order to collect information in a systematic way interviews were conducted using a semi-structured interview schedule covering the history of consumer development, successes and pitfalls, the current situation and future directions. Although some interviews were conducted face-to-face the majority were carried out on the telephone and took between twenty minutes to one hour. They included:

- 7 interviews with informants generally recognised as consumer leaders in New Zealand, the UK and Canada;
- 42 interviews with consumer leaders, consumer workers and other stakeholders with an interest in promoting consumer activity across Australia; and
- 26 interviews with consumer leaders and other interested stakeholders in Tasmania. This included an open invitation to active and ex-active TMHCN members to participate. Where appropriate, remuneration was offered to mental health consumers for their participation in the research.

Some of the early telephone interviews were taped and transcribed. Subsequently comprehensive and often verbatim notes were taken and quotes from the interviews have been used to illustrate the text of the report.

- **engagement in concurrent development processes**. This entailed participant observation at consultation events and activities. It also entailed setting up communication mechanisms with development processes so that they were able to inform this research.

The research was guided by a research reference group with a fifty percent mental health consumer membership as well as representatives from Mental Health Services, mental health provider organisations and the Tasmanian Council of Social
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Service (TasCOSS). Consumers included both those involved in the TMHCN and non-TMHCN consumers. The reference group met four times during the course of the research.

The TMHCN produced the original research proposal and assisted the project by making literature and TMHCN documentation available and by identifying some of the key informants.

1.7 Limitations of the Research

There were some factors which limited the extent of this research. Firstly the research took place at the same time as other consultation and development processes. These were:

- a statewide review of mental health consumer and carer participation led by Mental Health Services. This entailed a consultation process with consumers, carers, families, providers and other stakeholders through public consultations, consumer focus groups, a questionnaire survey and individual submissions; and
- the Consumer Organisation Project (see page 63) which entailed a number of TMHCN-run development forums to develop a business plan for a new Tasmanian consumer organisation and the formation of a small coordination team to drive the project forwards.

Given this environment there was a high risk of ‘consultation fatigue’ and it was imperative to make every effort to collaborate with, rather than duplicate, other consultation exercises. To this end the researcher attended the TMHCN and review consultation forums as well as establishing boundaries and information exchange mechanisms with the Review process. This included inputting some questions particularly relevant to consumer-run activity into the consultation process and the presence of the lead officer on the research reference group. It was also determined that this research should focus specifically on consumer-run activities rather than exploring the full consumer participation spectrum.

Secondly, the fact that the TMHCN was a partner in the research meant that the starting point for the research was not a neutral one. This meant that every attempt was made to consult with a wide range of informants to ensure a balanced and objective view.

Thirdly, it was not the purpose of the literature review to comprehensively describe all the literature relating to the consumer movement and participation activities. Rather emphasis was given to identifying relevant discussions and examples of activity which could help inform future developments in Tasmania. So although the research draws upon a range of information sources and published material it is not a definitive summary.

Fourthly, despite high levels of activity where consumers are actively engaged with contributing to service development the documentation of this activity can be poor. This means some reliance on oral testimonies from a small number of informants and on the judgement of the researcher to gauge the accuracy and reliability of sources and to construct some sense of historical developments and current realities. As a consequence there may be some gaps in illustrating the breadth and depth, impact and outcome of consumer developments in particular jurisdictions. However, the research has been able to reflect the major initiatives, themes and developments in the consumer world.

Lastly, this is a snapshot picture at one point in time. In a sector where resources are tight but where there is significant pressure at a policy level to promote development, activities and initiatives can change rapidly. This should be taken into account in using the information in this report.
2. The Consumer Movement: Themes and Issues

This chapter combines an overview of the literature about the consumer movement and consumer participation activities with first hand accounts of key informants to create a picture of the major issues facing mental health consumers in their efforts to engage in the planning and delivery of mental health services and of services in trying to facilitate such engagement. It outlines what both existing research and consumer leaders themselves tell us about effective strategies and where developments and energies might be leading in the future.

Consumer involvement can be initiated by mental health services or by consumers with the former being the most common and where consumers are expected to contribute in a way which fits with bureaucratic boundaries (Gregory 2007). This chapter looks at the issues faced in developing consumer involvement mechanisms from both within and outside the mental health system. It covers the underlying theme of recovery as an organising principle for mental health reform and the drive towards consumer participation across the mental health sector including the issues of representation and leadership. It discusses particular strategies for consumer activity and involvement including consumer workers, consumers as educators and researchers, consumer organisations and working with carers. Finally it explores the literature on the effectiveness of consumer initiatives and participation mechanisms in influencing mental health provision.

2.1 Recovery

Until relatively recently it was assumed that most individuals with severe mental illness would never be well again (Kruger 2000). However a recovery focus in all mental health services has now become a cornerstone of Australian mental health policy (Rickwood 2004) and one of the key organising principles for services internationally. The National Mental Health Plan 2003-2008 states that ‘a recovery orientation should drive service delivery’ (Australian Health Ministers 2003, p.11). The concept emerged from the consumer movement and has increasingly informed policy and service development. It identifies that the purpose of services should be recovery and self determination and that adopting these values profoundly changes the way people think about madness and mental health and requires a radical transformation of mental health services.

Recovery is defined in terms of outcomes: for example the ability to lead a satisfying life despite illness or symptoms. It does not necessarily mean a cure or clinical recovery but rather is about ‘living well in the presence or absence of one’s mental illness’ (O’Hagan 2004). A paper providing an overview of the recovery literature (Casey 2008) identifies that it is underpinned by notions of hope and healing, positive identity, taking responsibility and control and choice and is a process or a journey which the individual undertakes. It is closely linked to the principles of consumer participation and empowerment where active consumers provide evidence to clinical workers that recovery is possible and are involved in decision making at all levels.

However, there are questions about how a recovery focus should be put into practice within services (Shepherd et al. 2008). Some research has been done to identify the perceptions of both case managers and of consumers about the key factors which contribute to recovery. The research identified these factors as collaborative treatment planning, strength based approaches, knowledge and acceptance, appropriate medication, spirituality and holistic support and the self monitoring of symptoms (Ellis & King 2003). Many of these
factors are now enshrined in policy and planning statements. Others have identified non-compulsory services, wide availability of talking therapies, consumer orientated outcome measures and an independent national agency to monitor quality (MHAC 2008) as being key to a recovery orientated service. Yet there are concerns about how far these ideals are reflected in workers’ actual practice. Although services are increasingly defining and implementing specific strategies to support recovery this process can be impeded by scepticism and a lack of understanding about which factors impact on recovery. This means it is far from being standard practice in services. Worldwide there are now a number of initiatives to develop recovery competency frameworks for mental health workers in order to embed these skills into services.

At the same time concerns have also been expressed that recovery has been ‘colonised’ and diluted by professionals and managers so that services now say they are doing it but in reality they are not (O’Hagan 2004, Wellesley Institute 2009). It is also being used to support other policy agendas. For example, one informant in the UK suggested that recovery was being used to support welfare to work initiatives:

Recovery is getting a bad name. We never managed to establish it as a service user concept in England. It was coming from America, Australia and New Zealand and was not a home grown concept. The government are now trying to use the term recovery to get people back into work in a punitive sense, to prevent people being out of the workforce too long. This is received with great suspicion by people who have been long term unemployed for mental health reasons. They are worried they will have their benefits taken away. So there is now a suspicion about the idea of recovery happening. (Consumer leader, UK)

2.2 Consumer Participation

There is a large body of literature about consumer participation both as it relates to mental health services and more broadly about methods of consumer engagement across the health and human services sector. This literature covers the rationale for facilitating consumer involvement, definitions of what it is and what it should encompass, models of effective practice, its impact and outcomes and the barriers and obstacles. This has generated a number of ‘how to do it’ guides.

Essentially mental health consumer involvement is about the ways and means of creating a partnership between consumers and providers to improve the quality of care and outcomes for consumers and to provide a mechanism to ensure accountability and recovery orientated services (Meagher 2002a). Using the ‘lived experience’ of consumers is increasingly accepted as making a unique contribution to improving care and outcomes through modelling recovery, instilling hope and providing support and practical help (Doughty & Samson 2005). It can be seen as a democratic right, as a vehicle for positive systems change, and as a new way of doing things which offers a different perspective, insights into mental distress and alternative approaches. Participation by consumers in mental health services can be seen as even more crucial than in other health fields in order to counteract the reduction in civil rights experienced by those deemed legally incompetent to make decisions about their own health and who are treated involuntarily (Roper 2003).

It is however still a challenge to define what is meant by involvement and what it would look like if it had been achieved. Nevertheless there is a consensus that when consumers are trained, paid and valued for their skills and expertise, have an active role in hospitals, community services and community service organisations (CSOs), have their ideas for change seriously considered and are involved in decision making structures the goal will have been reached. UK Mind\(^2\) (Mind 2009) states that consumer involvement means:

- equal citizenship;
- dignity and respect in mental health services;
- full information on treatment and rights;
- involvement in treatment and care;
- independent advocacy in every area;
- broad participation through equal opportunities in employment and service delivery practices;
- involvement in planning, running and evaluating services;
- policies to ensure it is safe to get involved;
- the training of workers by consumers; and

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\(^2\) UK Mind is a leading mental health charity in England and Wales which aims to influence policy, challenge discrimination and promote the development of quality services and inclusion.
practical commitment and resources for consumer involvement.

In practical terms consumer participation in mental health services operates at four different levels (DHHS 2006); at the personal level through involvement in care and treatment plans, the level of local services, the level of policy and the service system and the government level to improve provision state wide and nationally. At each level there are a range of mechanisms to facilitate involvement ranging from information provision and one-off consultations, opinion polls and questionnaires through to consumer representation on decision making bodies. Indeed in some jurisdictions the mental health system now routinely has consumers as salaried staff, consultants on policy, researchers and management committee members. This means that the lived experience is now operating at many levels from active participation in treatment upwards (Gregory 2007) and that consumers can be engaged in different ways to suit their level of interest. This spectrum of involvement has been described as a ladder of power sharing where at the highest level consumers are sharing decision making responsibility with policy makers. It is conceived as embracing three distinct conceptions of consumer involvement – as recipients, subjects of consultation and agents in control (Peck et al. 2002). To date activities, certainly in Australia but also globally, have focused on consultation rather than influence, partnership or control and there remains a general lack of genuine opportunities to participate at both a treatment level and in the broader system. Consumers have also raised concerns that although consumer participation in service planning and delivery attracts more attention one of the most useful ways to implement it is for consumers to be involved in the development of their own treatment plans and respected for their illness, their knowledge and their lived experience (Pinches 2003).

Three of the most common models of participation (Kroschel 2002, 2005) have been identified as:

- representatives on committees;
- involvement in short term projects without the establishment of a consumer participation infrastructure; and
- consumer groups developed through mental health service support.

The first is the most widespread and supported by national and state strategies. It is one of the easiest to implement which makes it the most feasible in terms of gaining the support of providers. However, it is also service controlled. The last model is, for many consumers, the most sustainable and productive model and allows consumers to set their own participation agenda. However, there is also a consensus that diversity is required: 

Consumers need several different avenues in order to tell their story, open groups where they can come and go when they feel well enough to engage and give their opinion, opportunities at service level and strategic levels to engage. The idea that one group, diverse and experienced as it may be can be the one point of reference is just not broad enough to realistically give a picture of what’s happening in mental health. They might be the point of engagement around particular things but you would need to engage in a broader and more open strategy. (Consumer leader, SA)

2.2.1 Community service organisations (CSOs)

Much of the literature about models of consumer participation focuses on public mental health services. However, CSOs have an increasing role in the delivery of mental health services and this is likely to expand as more services are contracted out. Although there may be an assumption that because they work at ground level and may have a history of being led by consumers and/or carers they are consumer aware and clear about consumer rights and participation mechanisms this is not necessarily the case. Yet the research did find many examples of good practice in CSOs where consumers held positions on boards and committees, where quality frameworks were designed to be inclusive of the lived experience and where there were initiatives to develop consumer worker roles and peer education. Informants reported a growing awareness of the need to have well developed roles, policies and mechanisms for consumers to participate in supported accommodation and day programs. As one informant said:
Some NGOs have really embraced consumer involvement and built it fundamentally into their core principles of operation. They offer ground breaking models of how to reengineer and redesign services to authentically take into account the consumer perspective and involvement. (Stakeholder, WA)

In one jurisdiction the peak body for the mental health non-government sector has developed a consumer caucus to promote involvement in strategic decision making:

When the Mental Health Coalition formed it established a consumer and carer caucus which was more than a consumer advisory body. It was a link to our Board of Management. The caucus nominates two consumers and two carers to sit on the Board and to raise the profile of issues with policy and service development. It really gives the potential for consumers to be involved in the organisation’s policy development and other program areas. This is a unique model for any peak body in Australia but it does require ongoing attention and support. (Stakeholder, ACT)

Despite significant levels of consumer activity in CSOs these models have not been reviewed and are rarely evaluated.

2.2.2 Barriers to consumer participation

Whatever model of participation is being used and whatever the rhetoric in policy and practice frameworks good practice consumer involvement in mental health is still essentially a minority activity (Tait & Lester 2005) and a range of obstacles are encountered which have been well documented in the literature nationally and internationally. The key barriers are:

- **lack of commitment to adequate resourcing** where consumer views are regarded as the icing on the cake rather than the cake itself and there is little practical support for consumers in terms of remuneration, training and welcoming environments (Bertram 2002). This is seen as the key reason for failure; for example when consumer participation is named up as a ‘money neutral priority’.
- **negative stereotyping of consumers** as incompetent, unreasonable and illogical, uninformed and potentially violent and where their views are considered to be a reflection of their psychopathology (Meagher 2002a).
- **professional attitudes and resistance** where people are understood according to their symptoms, where there is a lack of awareness about what can be gained from the consumer perspective and where experiences are seen as personal stories rather than having wider implications (Edan 2006).
- **a gap** between the requirement for participation at the political level and the lack of mandates on how processes or mechanisms should be facilitated. This leaves too much room for local interpretation and results in the absence of systemic approaches to facilitate participation.
- **developing a consistent response across a diffuse sector.** There are questions about how to promote consumer participation across a diverse sector which encompasses public and private services, GPs working autonomously and CSOs providing a range of community services.
- **lack of consumer confidence** where consumers suffer from self-doubt and a lack of faith in their ability to exert control or represent others and have difficulties in ‘coming out’ and taking on the risks of personal exposure.
- **professional control of participation structures** which are geared in favour of full time professionals. This can result in tokenism when there is little infrastructure to facilitate involvement, no training or support and no real transference of power.
- **conflicts within the consumer movement** which impairs the ability to present a united voice. Some consumers have described a crisis of integrity and ethics in the movement (Meagher 2002b).

These barriers mean that although consumers may be involved in the system and working in mental health settings their involvement is marginal because the status quo has not been challenged. Some have described a backlash
from professionals with consumer competence being challenged when participation gets close to the area of treatment which is traditionally the unquestioned domain of professionals (Roper 2003). As one consumer said:

 Sometimes we are just there as tokens. That's the feeling. Consumer reps are there and consumer consultants just so they can tick a box to say they've had consumer representation. That's why a lot of positions aren't paid and you only get paid travel sometimes or you don't get sitting fees. The leader of the mental health team said we can't have consumers on the Board because they don't understand how we work. My reaction is we don't have to. All we have to do is be an expert in our area which is being a consumer. If they wanted financial advice they wouldn't expect a worker to know how they ran their budget, it would be an expert on finance. (Consumer leader, SA)

Another consumer described how this had manifested as political correctness where a T-shirt designed by consumers and with the logo ‘batty is beautiful’ had been censored by professionals for reinforcing negative stereotypes (Happell 2008).

To overcome these barriers requires action to change cultures and attitudes, reverse low expectations, deal with the practical aspects and embark on education, training and awareness-raising.

2.3 The Consumer Workforce

There are increasing opportunities for those with ‘lived experience’ to go back into the mental health system as paid consumer workers performing a variety of roles. The three main roles are as:

- paid consumer consultants or advisors engaged in systemic advocacy work;
- paid peer support workers providing support, befriending, role modelling and hope for recovery to both consumers and the mental health workforce; and
- consumer representatives participating in advisory and decision making forums and paid sitting fees.

These roles can be established and integrated within traditional mental health services, in CSOs or in consumer-run organisations. There are also limited opportunities to work as consumer service auditors and mental health surveyors involved in surveying mental health services under the National Standards.

Internationally the value of consumer roles in mental health services has been widely accepted (Hansen 2003). They are seen as mechanisms for the transformation of mental health services towards a recovery orientation, as a vehicle for cultural change and as providing much needed employment opportunities for people with mental illness who suffer disproportionately from structural exclusion from employment (Bertram 2002). They can facilitate consumer participation activities, enhance the visibility of the consumer voice at both the individual treatment and service delivery end and provide general support and mentoring to other consumers.

In Australia five jurisdictions now employ consumer consultants, some also have paid peer support workers and all use consumer representatives. Many see these workers as an entirely new service located within traditional mental health services and an integral part of developing quality recovery-orientated services. The picture can be confusing because consumer workers are called different things in different places. There are 16 different job titles for employed consumers in Australia alone which means that little can be assumed about the role from the job title.

However research has also documented numerous problems in establishing consumer positions and sustaining them. They have evolved in an ad hoc fashion with an absence of guidelines or clarity about aims and functions and often minimal support and resourcing (Hansen 2003). There is a lack of standard training for consumer workers, confusion about roles, titles and expectations and no development of career pathways (Steward et al. 2008). An analysis of the situation in one jurisdiction (Watson 2007) found confusion between consumer positions and the mental health workforce generally with no recognition of the distinct role of the consumer, no appropriate award structure and no accredited training. In another state consumer consultants employed by area mental health services suffered from low rates of pay, lack of support and training, high workloads and high stress levels. This results in a high
turnover of personnel (Pinches 2004, 2005). They are employed
part-time for a few hours a week or month as cheap, unskilled
welfare workers and with no industrial protection (Edan
2006) and they are often seen as add-ons to mental health
services without the same entitlements as other employees.

You are piggy in the middle all the time. You are
neither staff nor consumer. Staff won’t treat
you as staff because you’re a consumer. You
are isolated and on your own. It’s a very, very
hard role. We have a workforce who already
have a disability and we’re asking them to
work in a hard and isolated role without much
support. We will have a lot of burnout and
staff turnover. We have people employed under
different awards and there are at least four or
five different descriptions of what a consumer
worker’s role may be in any given area. Also
there is no accredited training and it’s been
highlighted that consumer workers must have
a minimal set of standard training before they
hit the wards or they will fail, burn out, get
sick. They need to be well prepared and know
exactly what it is they are doing. Not even the
line managers or area directors really know
what the consumer worker’s job is. There is
therefore no real management or guidelines. It
seems to be whatever your area health service
deems it to be. (Consumer leader, NSW)

There are also difficulties associated with the blending and
overlap of consumer advocacy and peer support roles.

The position was supposed to be systems
advocacy. Obviously you can’t get away from
doing a bit of peer support but some positions
have turned into nothing but a peer support
role. We should be changing the system and
identifying the barriers and making a better
quality of life for the consumers. The touchy
feely peer support stuff is good and it’s wanted
but that wasn’t what consumer consultants
were supposed to be doing. Some are going
into the units and that’s all they’re doing, sitting
talking to consumers all day long and not doing
any advocacy at all. (Consumer leader, Qld)

The issue of training and qualifications has become paramount
and led to debate about formal qualifications, the benefits
this brings in terms of improved self esteem, credibility
and a reduction in stigma and how qualifications should
be balanced with lived experience and skills learnt on the
job. In the absence of any accredited training for consumer
workers the only option can be to qualify as mental health
workers and this can result in role confusion and an impact
on wellness (Watson 2007). This picture has been further
confused by recommendations from the Mental Health
Council of Australia that consumers should be encouraged
and assisted to become part of the mental health workforce:

There are national recommendations that
consumers should be part of the new mental
health workforce or substitute workforce
but without any proper distinction between
a consumer workforce and other workforce
issues. People are disappearing from the
workforce and we have people with mental
illness looking for work and all these vacancies.
How can we plug the hole? We put them in
as substitute mental health workers. This
approach was put forward for an award for
best practice in mental health and consumer
participation policy is used as the justification.
That is a big error. (Consumer leader, NSW).

This kind of approach has promoted a trend for some
consumer workers to become quasi mental health workers
but without a clinical caseload. There are concerns that they
are in situations where there are two lines of accountability
and divided loyalties to service managers and to consumers.
This creates difficulties in trying to maintain personal and
professional integrity with both groups. As one consumer said:

They had become subsumed by the system.
They were sitting in on clinical reviews and
having access to confidential information
without the consent of consumers. The
worst violations of rights by consumer
workers are the breaches of health
privacy rights. (Consumer leader, ACT)
This of course raises broader questions about how far being employed by the system traps consumer workers in reactive roles rather than being able to set agendas or do systemic advocacy. As one consumer said ‘there are difficulties in being an advocate in the service trying to initiate change and also protecting your own job’. Some would like to see consumer workers being employed by consumer-run organisations rather than the mental health sector.

Most of the consumer workforce don’t last more than ten years. They are put in positions to advocate against the people who fund them and there are many conflicts between the expectations and role and ethics of the consumer. It creates difficulties and can leave people in pieces. It happens all over Australia. (Consumer leader, NSW)

My biggest wish is that [the consumer organisation] take over running consumer workers with the area health services so they don’t get gagged. There have been situations where you report back and you get threatened by staff because you’re the whistle blower. If you bat for the consumer it feels you are then biting the hand that feeds you and you think will I have a job next week? (Consumer leader, NSW)

There are questions about how best to support an emerging consumer workforce and ensure that employees are resourced and well paid. Many consumers would like to see clear job descriptions, a common understanding of the role, adequate preparation, ongoing support and supervision and the development of accredited training for workers supported by a code of ethics and minimum performance standards (Steward et al. 2008). They wanted the consumer workforce seen collectively as a separate discipline – for instance like nursing – with external supervision and mentoring or apprenticeship accompanied by guidelines for employers to reduce inconsistencies and map career pathways. They also wanted to see ‘real jobs with real qualifications required and real expectations’. As one consumer said each 25 bed in-patient unit with an attached case management service requires one three day per week consumer consultant at a minimum.

This is certainly happening in New Zealand where peer support and consumer advisor services are run variously by consumers, district health boards and CSOs. A literature review undertaken to inform the development of a consumer workforce within mental health services found that successful services shared a clear philosophy, guidelines to separate peer work from traditional support roles, a good recruitment process, training consistent with the role and an effective supervision structure (Mind & Body Consultants 2008). There is now a push in New Zealand to establish national guidelines for consumer roles, build a culture of acceptance, reduce the stigma and discrimination they face and develop a training framework (Hansen 2003).

In Australia the Department of Health and Ageing (DoHA) has recently initiated a scoping study to look at the formal training available for consumer leaders, peer support workers and consumer consultants and to recognise them more formally as a profession with the mental health sector requiring specialist training. The study aims to explore how to develop a nationally consistent approach to better engage consumers in the mental health workforce, set national standards for consumer workers and develop accredited courses for consumer consultants and peer support workers. There are also moves to formalise and better support consumer representatives. The Consumer Affairs Advisory Council (CAAC 2002) published principles for the appointment of consumer representatives stating that it should be on merit, targeted at those with expertise in consumer affairs and linked to relevant consumer organisations.

The whole system needs to adjust to having well consumers involved in positions – representative or paid – when they might at any other time be using that system themselves as unwell consumers. Consumer representation provides a mechanism for showing that people can be well enough to participate. As a lot of clinicians only see people when they’re crook it’s about undoing the fear and stigma. (Consumer advocate, WA)

One of the key functions of developing a consumer workforce is to initiate the cultural change required in services. This can also be promoted by encouraging consumers to become part of the professional mental health workforce through using
positive discrimination and flexible working arrangements in the recruitment of staff to mental health programs. People who have used mental health services and who now hold reasonably paid employment both inspire other consumers and provide evidence to staff of what can be achieved. This challenges the “them and us” culture and the low expectations held by so many mental health professionals and works to reinforce social inclusion. Work undertaken in the UK (Seebohm & Grove 2006) provided a practical demonstration of how mental health trusts could lead by example in employing mental health consumers in the National Health Service (NHS) workforce while at the same time improving the working lives and job retention for all staff.

2.4 Consumers and Carers

Australia is unusual in having a number of initiatives which involve consumers and carers working together around a common agenda. Elsewhere consumers and carers have remained separate although they might come together to take up particular issues or campaigns. Commonly their agendas are seen as different; for example carers might want access to information or compulsory treatment to keep their relatives safe and ensure access to services while consumers might not want them to have access to information and feel that compulsory treatment severely impacts on their rights as citizens.

Working together in Australia was promoted by the first National Mental Health Plan in the early 1990s when there was a push to establish consumer and carer advisory groups in all jurisdictions. In 2002 the National Mental Health Consumer and Carers Forum was also established to provide a focus for joint work around national issues.

Some jurisdictions have continued to support structures where consumers and carers work together with joint agendas. In some cases this is due to preference while in others it is more a question of feasibility particularly when resources are too tight to be able to build separate structures. Informants described few differences of opinion and mechanisms which allowed separate platforms when required.

If you don’t have a group made up of consumers and carers and it’s just consumers it tends to fall over. A carer is a consumer of the service and we champion the cause of consumers so we work in partnership quite well. As long as there are not more carers than consumers it works. If there weren’t any carers in my group it would fold. If consumers are doing a lot of work they can put themselves under too much stress. (Carer leader, SA)

We are not advanced enough in our philosophy. Maybe there will be a split sometime in the future but I don’t think we have the luxury here and we don’t even have the forum for really having that discussion. (Consumer leader, Qld)

Other jurisdictions have maintained separate development. Carers are seen to dominate any mixed group and although sometimes it was mutually beneficial at other times it was seen to be destructive and had led to passionate discussion. As one consumer said ‘not having carers involved helps a lot. That’s just another extra layer of angst which we don’t need’.

We decided that even though we are on the same journey there are some areas that are different, in the same orchard but two different trees. I have met consumers where there are major issues with family that can exacerbate the mental illness. In some other states it’s quite successful. Here the carers have gone ahead in leaps and bounds, they are a very professional organisation. I am envious. But it has been difficult to prevent carers making remarks that are offensive to the consumers. (Consumer leader, WA)

In New Zealand there is a basic cry that the primary stakeholder is the consumer because if the consumer wasn’t there none of the other stakeholders would exist and the distinction between carers and consumers interests is very well accepted. There is a little more collaboration going on now partly because of the clarity around the differences and because consumers feel in a strong enough position that they won’t lose power. But what lies underneath this battle is an assumption of disability and that mental illness will be disabling. To try and
Experts by Experience: Strengthening the mental health consumer voice in Tasmania

get consumer initiatives started in a system
still working within this conventional paradigm
is very difficult. (Consumer leader, NZ)

2.5 Representation and Leadership

A common consumer participation strategy is to foster, support and remunerate consumer representatives to sit in decision making forums and provide a consumer perspective. This raises issues, usually among service providers but also among consumers, of how representative these active and involved consumers are of the broader and diverse mental health community (Happell & Roper 2006). Typically representatives do not represent an organisation or interest group but are unconnected individuals who rely on their own experiences. This means that it can be difficult for them to get beyond their personal view to represent the broader, generalised views of consumers or marginalised groups especially if they have had no training. In addition there is also a view that active consumers or consumer groups are unlikely to represent ordinary consumers. This raises questions about how to collate the broad views of the consumer community while avoiding the risk of sectional interests. These concerns have, as Mind UK identified, led to an endless search for the elusive typical user representative and to questions about who the ordinary consumer is: are they the most incapacitated or those who have been made better by treatment? Suggestions that those who advocate are not typical and therefore not representative is the ‘catch 22’ of consumer participation where consumer views are welcomed but not from those who are confident enough to give them. These issues are identified at the macro level of consumer participation engaged in systems reform rather than at the micro level of participation in individual treatment plans, where the issue of representation is not relevant as care is individually based (Lloyd & King 2003).

As one consumer said this emphasis on a requirement for representativeness means that instead of the consumer role being about introducing a consumer perspective, someone will always ask who you represent (Meagher 2002a). This is not a question asked of professionals and there is no known process to canvass the views of the broader profession. It means that professionals can make unrealistic demands but without offering the resources for consumers to develop the capacity to become representative.

Research (Crawford & Rutter 2004) sought to determine how far the views of an established mental health consumer group were representative of the broader community of consumers through comparing the rating given to the importance of specific priorities for service developments. The results were similar although the consumer group placed a higher importance on the priorities generally than other consumers. The research concluded that the assumption that the views of active consumers are out of step with others should be questioned and that in fact it was a discriminatory position which served to silence activism and diminish the importance of what consumers were saying. It also undermined the legitimacy of consumer roles and introduced an inherent contradiction in employing someone to provide a consumer perspective and then dismissing it as sectional or anecdotal.

This search for representativeness in consumer participation activities is leading, particularly in New Zealand, to a move towards the notion of consumer leadership where consumers participate as individual recipients and leaders. As one Australian consumer said:

We are now ditching the term representative completely. It’s a stupid term and you can never represent people. It takes you down a dead end. So we are talking about consumer leadership and we are moving on to thinking in quite different ways than what we were thinking ten years ago. (Consumer leader, Victoria)

Our current policy talks about consumer leadership. In New Zealand we got a little bit tired of consumer participation. We felt that was about being invited to someone else’s table according to their agenda and culture and roles. We felt if we talked about leadership then we would have our own table or at least a shared table. (Consumer leader, NZ)

A recent literature review and consultation about consumer leadership across the health sector (Victorian Quality Council 2007) found that it is organic and arises from a

3. See note 2 on page 11
community of interest like a consumer organisation or network. Its promotion and development relies on effective consumer participation mechanisms where it can grow and gain sustained support. These mechanisms can include having consumer representatives or consultants at all levels, including at a senior management level, to create an enabling environment and an investment in education, training, mentoring and support. The review outlined key leadership characteristics as having a strategic direction or vision, the ability to energise others and provide a role model, good communication, negotiation and relationship building and conflict resolution skills. The review concluded that the most effective strategies for developing leadership capability in the health system are formal learning, peer support and mentoring programs accompanied by guidelines for supporting consumer leadership in the health system.

Interestingly there is now a move in New Zealand to professionalise the representative role by employing representatives. The question then becomes how professional representatives or consumer leaders keep in touch with the broader constituency of mental health consumers. This points to having solid and credible consultation mechanisms:

4. See page 53 for a description of the work of Mind and Body Consultants.

**Mind and Body Consultants** in New Zealand do consumer representation but they have a number of employed consultants which creates a professional body. It is a different model with advantages and disadvantages. Are the representatives actually representative? Being a consultative body really is the key to doing a good job and a legitimate job rather than being representative. The argument then is that the representatives could be a more professionalised group and an employed group as long as the consultative processes are well established and sound. (Consumer leader, ACT)

It requires that consumers are clear about their links to mental health constituencies and proactive in representing disenfranchised groups to ensure a range of consumer views are heard. This requires the development of sophisticated and ongoing consultation mechanisms.

What happens with representative structures is they go to the government and they say, ‘you are just the strongest voice and just giving us your agenda. How do you know it’s what those other people out there are thinking? We need to develop a survey capacity and you can do that through technology easily via emails and websites. So in order to do advocacy we need to really get them good information and instead of going along with a few ideas from a committee it would be great to have a way of surveying consumer opinion very quickly. (Consumer leader, NZ)

Mad people are not a constituency whose interests can be represented like a political system. It is better to say that people who have experience of using the system and who have done some thinking about this and know other people using the system can offer sensible advice about how to make it better. So it’s about being experts by experience rather than representatives by mandate. This is a significant factor in development because it lays the groundwork for people in consumer positions having the same responsibility to be competent and deliver the goods as anyone else. (Consumer leader, NZ)

There are of course real issues about how far consumer participation mechanisms and consumer run organisations deal with diversity. It can be difficult to tap into the views of Aboriginal and culturally and linguistically diverse (CALD) groups, rural populations, older people and young people, all of whom may suffer particular disadvantage and have special issues and needs. For example, experiences of stigma can be compounded in rural environments (McColl 2007) by lack of services, isolation, confidentiality concerns and cultural differences.

In the Chinese community they don’t want to discuss mental illness. It’s seen as hereditary and will affect marriage prospects and family fortunes. It is not accepted by many circles.

5 In this document Aboriginal refers to Aboriginal and Torres Strait Islander people.
and they look down on you as crazy. Mental health in many CALD groups is not defined by the consumer but is a family matter and more of a community issue rather than just personal. (Consumer leader, NSW)

Organisations do describe working with a range of agencies and individuals to ensure the views of these groups are heard.

We sit on committees with Aboriginal people and know that if you need some specialist advice you can contact each other. We do have some Indigenous people who are quite involved as reps. When we go to in-patient facilities we see everybody, different ethnic backgrounds, Aboriginals are not exempt from problems. (Consumer leader, Victoria)

2.6 Consumers as Educators, Researchers and Evaluators

A core problem in increasing the involvement of consumers is the stigma and discrimination they face in the mental health system and the professional resistance to involvement activities which is a symptom of that. A major goal of the consumer movement has always been to change staff attitudes using the mantra of ‘treat us well and we will get better sooner’. In response to this consumers are increasingly acquiring a role as educators and trainers (Pinches 2003) and there is a push to make the consumer perspective intrinsic to all aspects of education and training (NMHETAG 2002). There are now examples of good practice where this has become a reality (Edan 2006) and where valuing the lived experience is an integral part of training.

It seems to work. A systematic review of the evidence of the impact of consumer run initiatives (Doughty & Samson 2005) did show that using consumers as trainers enhanced effectiveness and that post-training attitudes were more positive (see page 28). Delivering personal stories can impact on the values and attitudes of mental health clinicians. A number of consumers and consumer organisations are now engaged in this work across Australia.

We have been going to the medical schools to those who will end up being the future doctors. There has been work with current mental health service staff in various sessions and on a project called Consumers as Tutors which involved being engaged in training for staff. Mental Health ACT has got to the point where a lot of the training they provide invites consumers to participate so they end up playing an important role in educating staff that are also attending the training. So there are things happening on a more informal basis. We also did a couple of narrative sessions which brought consumers and mental health professionals together to talk about things like stigma or the doctor/patient relationship. Participants felt they have really opened minds and helped move on to a more open and equal relationship. (Consumer leader, ACT)

There are questions about how to build an enduring system where consumers are routinely involved in the development and presentation of training to clinicians.

Mental health services have nurse education units with nurses as educators. We want a consumer education worker in my directorate with input into the education program and curricula development. This should not be about story telling. It is much more complex than that and should be based on a philosophy of social change from a critical consumer perspective. So it’s not about being critical of the service but about developing services and reaching a situation where clinical staff always question what they are doing from a consumer perspective. (Consumer leader, Victoria)

There is also a trend towards consumer led research and consumer clinician research partnerships (Beresford 2002) although it is not routine in services and consumer-controlled research projects are still relatively rare in Australia. It is
challenging for clinicians to engage with the consumer perspective and there is often a lack of opportunities for consumers to have hands-on experience of research methods. Pinches describes a shift from non-patient researchers to patient researchers and the resourcing of consumers to evaluate services themselves (Pinches 2005). He identifies active consumers as ‘pathfinders’ who are providing new ways of seeing and doing things.

Consumers are increasingly involved in devising consumer-orientated outcome measures. Performance measures which use the recovery paradigm to improve clinical practice and policy development would have real benefits for consumers. This requires working in partnership with consumer consultants and advocates. There is also a move to establish consumer perspective research and teaching in universities.

One of the big things that’s happened in the last five years is this moving towards consumer studies as a discipline and thinking how to place the discourse somewhere where it can make a huge difference. We would like a consumer research unit attached to a university which would act rather like an Aboriginal educational research unit. It would be there to support consumers through university because people with mental health problems drop out or they are sick. It would collect the grey literature because so much of what we’ve written doesn’t get into Medline where you would go about finding literature on mental health. And it could act as a way of nurturing PhD students writing about consumer experiences. We want to challenge methodologies like what is knowledge in this area. (Consumer leader, Victoria)

Involvement in the evaluation of services has been promoted through the requirement to have consumers as part of the teams reviewing services against the National Standards for Mental Health Services. While one of the accrediting bodies reports difficulties in recruiting and maintaining a pool of consumer reviewers, the other has trained consumer surveyors and coordinates this workforce centrally. It has also set up annual surveyor training and support mechanisms entailing quarterly telephone conferences to establish an informal support network (ACHS 2007).

2.7 Mental Health Consumer Organisations

There are many different kinds of consumer groups and organisations. Firstly, there are local consumer support groups which are about sharing experiences, social contact and activities. They may target people with a particular disorder like eating difficulties or depression, and operate as mutual support and self help groups where members gain support from others living with the condition to promote recovery. They are small, commonly facilitated by volunteers and run by committees which may just be consumers but can also include carers and professionals.

Secondly, there are larger consumer groups which may or may not offer mutual support but also perform a range of other functions like information provision, individual or systemic advocacy, educating mental health professionals, establishing new consumer support groups and auspicing consumer projects or research. Some have also established web based groups and forums which provide support, information, debate and opportunities to feed back service experiences. They are largely run by consumers for consumers.

Thirdly, there are consumer run service provider groups which are driven by a desire to create alternatives to traditional mental health services at a grass roots level and to push for a recovery orientation. They may support patients in professionally-led services and provide drop-ins or alternatives to hospital admission as well as personal support and advocacy. Lastly, there are consumer-run businesses which can provide a range of employment opportunities for consumers outside the mental health sector.

Some classify these groups as ‘consumer developed initiatives’ (or CDIs) which is a broad phrase to describe any activities organised by consumers to provide mutual support or change society’s perception of mental health consumers. They are seen as providing a range of benefits including a support base, a sense of hope, opportunities for decision making, choice and the development of new skills and creating a cadre of knowledgeable consumers and leaders to press for change. They have increased in visibility and importance in the mental health systems of many countries in recent years and in the US, for example, now outnumber traditional mental
health providers by a considerable margin (Goldstrom et al. 2006). They are also seen as central to the development of a consumer workforce (Orwin & Burdett 2009).

Yet they also suffer from a range of problems in getting established and sustaining themselves and are constrained by both the resources available and the culture and understandings of mental health providers and systems. Research to pool the experiences and knowledge of consumer developed initiatives (Mowbray et al. 1997) describes their ideas, principles, guidelines and practices and the issues, challenges and barriers they face. These include a lack of support and funding, difficulties in dealing with internal conflicts and tensions, high burn out rates and a lack of stable workers to provide leadership and maintain organisational survival. Overall the research identifies the absence of a road map or established wisdom about how to avoid difficulties or resolve them and no mechanisms to share and accumulate knowledge as significant gaps.

2.7.1 Getting Started

The difficulties consumer organisations face in establishing themselves is well documented although they are not significantly different to the difficulties faced by many new community organisations. This research gathered a wealth of material on the experiences of setting up and sustaining consumer organisations both from the literature and from the first hand accounts of consumers. They all undergo a difficult struggle to exist, survive and be effective and, as one consumer said, the common experience is organisational crisis which is almost inevitable (Meagher 2002a). The factors contributing to this are a lack of adequate structures to maintain a shared purpose and direction, dominating charismatic leaders who can disempower others and lead to factions and breakdown and the difficulties reactive, activist groups have in sustaining a purpose and motivation. In terms of establishing organisations research informants spoke about three things – the shortfalls in capacity in the consumer community, the difficulties of gaining and maintaining independence and difficulties in defining a purpose.

Building a mental health consumer organisation can be more difficult than setting up other kinds of CSOs because of a lack of capacity within the consumer community:
One way of seeding a consumer organisation is to establish auspicing arrangements with a larger organisation which can provide accommodation, mentoring and support in the early days. A number of organisations who participated in the research had experienced auspicing. It had worked for some and not for others depending on the nature of the relationship with the auspicing organisation and clarity about what that relationship entailed.

We became incorporated and we got a small grant and established an office but there were difficult periods. Building up an organisation and structures is always a challenge and there were times when the network was auspiced by the local [Council of Social Service]. We had one part time coordinator a few hours a week and there was more than one instance when that person became unwell and there were difficulties just running it and keeping up with requirements and living up to a contract. There were times when it was fairly close to collapsing. (Consumer leader, ACT)

Organisations reported difficulties with auspicing arrangements, particularly when they were auspiced by organisations in the mental health sector. It meant that they were unable to escape the role of client or patient and the accompanying discrimination that this entailed.

It usually doesn’t work. There is some literature and a lot of discussion from around the world that this type of arrangement does not usually work well. Occasionally it does and that’s probably due to clarity around the relationship and a CEO who is not full of levels of prejudice because that is what we’ve got to face in the mental health sector. (Consumer leader, NZ)

It is very important to stay away from the mental health sector. Consumer run organisations do not get the understanding from mental health services. The clinical services are too ingrained in their practices to understand. They find it too risky, unsafe, they can’t actually see that there is another level of working. It’s really advisable to look at non mental health providers to support as auspicers or as an umbrella to make it easier to get contracts. If you work with a well established organisation getting contracts is easier. (Consumer leader, NZ)

This meant that the goal is often to become an independent, incorporated consumer-run organisation.

I have always seen that in having legitimacy and being a maturer organisation one has to be incorporated. I felt incorporation was a measure of independence and a demonstration that it’s a stand alone organisation. To a degree it added credibility to the organisation and it certainly acted as a spur for a number of others to become incorporated. (Consumer leader, Qld)

Although most consumer organisations are established initially to provide a voice for consumers and/or to offer support, in order to attract funding they need to be able to carefully define what kind of service or service mix they can realistically offer. It means choosing between a range of options from individual to systemic advocacy, support and recreational activities, education and awareness raising, information provision or consultancy. Some focus on systemic advocacy, others might find a balance between individual advocacy and broader lobbying and campaigning work.

Consultation is a key element in the work we do and it did grow organically with a desire to go and represent and be involved directly at different levels and there was a willingness from ACT Health to involve consumers. The government sees the consumer representative program as the key service and most of our contract revolves around that but there is also an element of awareness raising and education of consumers and the broader community and policy work. We are a systemic advocacy organisation so we try not to get too involved in individual advocacy. It would be logical to be involved with it and consistent with the aims and goals of the network but at the
same time it’s a very different dimension of work and individual advocacy is provided elsewhere. (Consumer leader, ACT)

It’s the individual advocacy that informs us about what is missing systemically. We do the individual advocacy, keep data on it and then we have group advocacy. We establish consumer groups in the areas in which people live, visit them on a monthly basis, have a meeting over a barbeque about how are you going, what are your issues? Those two activities inform us about the systemic issues. For example we know patients’ rights is a systemic issue right across this state, so is women’s safety, seclusion practices, the culture. We can say in this state what the major issues of concern are because we are out there visiting people in in-patient facilities, in the towns, listening to them, not telling them what the issues are but getting them to tell us. (Consumer leader, Victoria)

It is of course important to be delivering a service that people want and need and some organisations have had difficulties in defining a role that meets the desire to advocate and strengthen the voice of consumers in a way that attracts and involves a broad range of consumers:

We have an attitude that we are service providers. Every year we used to hold a meeting with consumers where we said you tell us what you want us to do. The services we deliver are the ones that they have identified as wanting. Other places don’t do that. You get a small core and they don’t do an analysis of what consumers need. They develop a service they want to deliver and that is comfortable for them. (Consumer leader, Victoria)

Some organisations may come together to promote a particular issue but when they achieve that goal have to restructure and define another purpose otherwise they cease to exist. It also means that success in one area can boost confidence and build the capacity of organisations to move on into bigger and broader fields.

It started out in the late 1990s with a group of people coming together after a workshop. We wanted a place to call our own that respected us in a different way to what we experience from mental health services. So the initial drive was to establish something based around the club house model. The Rainbow was established so there was a sense that we had achieved what we initially came for but also that as consumers getting together we can actually achieve things. From there it developed a broader perspective to have an input into the development of mental health services. (Consumer leader, ACT)

Lastly, a common problem faced by many organisations is the expectations of others about what they can or cannot achieve.

Mental health services here would like to have the group run before it can crawl and really it needs to evolve at its own pace. They want it running training for consumers, placing consumers on committees and supporting them in that work. All of that requires quite a solid structure and to be able to deliver a service means you have to have your house in order. It will take some time. (Consumer advocate, WA)

2.7.2 Governance and staffing

Governance structures vary but usually entail a management committee or board of trustees. Commonly consumer organisations require that all committee members are consumers and a number of research participants described the difficulties they had in building stable committees of consumers to govern organisations:

One of the problems at the beginning was how to develop any kind of governance structure for consumer involvement that was consumer led because a lot of the consumers who came into the system were dealing with their mental health problems and not a lot of them
necessarily had management or community governance experience. We had trouble recruiting people that were able to sustain their efforts for a range of different reasons. You have people at different stages of their own mental health and some may never be able to contribute a great deal, maybe their opinion now and again. So a lot falls on one or two shoulders. (Consumer leader, WA)

Getting a stable consumer board is quite an art in itself, one which will be there for the long haul and can create that stability. We have the capacity to have nine members on the board but we have never had the full nine. (Consumer leader, NSW)

This has led to exploring ways of building the capacity of consumers to participate through building up the membership. In 2007 the Health Issues Centre in Victoria undertook a project to determine the skills, knowledge and needs of mental health consumers and then to provide education and training to meet these needs and to support trainees to negotiate the mental health system. ‘Self stigma’ is an issue for most people with lived experience and is the result of internalising the negative messages and behaviour received from others. The project found that many consumers required support with their self-esteem and in dealing with stigma before they could participate. The Centre also conducts annual training to support consumer involvement in health care generally and many participants in the training have mental health issues.

At another level training is also required to equip consumers to participate equally with providers by acquiring skills in meeting procedures, public speaking, submission writing and advocacy. As one consumer said ‘it’s training people in the fundamentals of how a system works. It’s an apprenticeship’.

If you can get a membership core then the job is to educate people to a point where they take over. It allows you to train people up. We watch that person grow and over the years they end up becoming the president of your organisation. There’s a lot of education of consumers to be done to the point where they are confident enough to say things but not in a way that will alienate anybody. This has to be done well and by other consumers and they have to be funded. (Consumer leader, Victoria)

There has also been a more profound questioning of the appropriateness of committee structures for consumer-run organisations.

In some ways the traditional consumer organisation is really based on an old trade union structure. You have your representatives, they come together and make decisions. That is not a very 21st century structure. Who are you representing? And there isn’t always a body to represent because there aren’t local structures. So I think that model is dead, it’s disastrous. I would like to be able to do away with all that stuff and just have a group of people who may be governed through a company or a trust but who have accountability. (Consumer leader, NZ)

Many organisations are committed to employing only consumers. This can mean they have flexible arrangements to manage and accommodate illness. It can also mean recruiting from a very small pool of people with the appropriate skills, including relationship building which was seen as essential.

One of the reasons why this place has survived so well is because I have a management background. People are often appointed to be managers who have no education around these matters at all. I’m not talking about balancing a budget, I’m talking about managing people. Also people have become unwell from time to time. We have flexible working rules. Ninety nine per cent have never taken advantage of that flexibility. One per cent have and I’ve nipped it in the bud. (Consumer leader, Victoria)

If you can go into any meeting and make the people around you feel good you’re much better off. That is the fundamental responsibility of consumers if they want to do this work. But too many have an operational
style that is vitriolic and abusive. I don’t consider it okay to have a string of corpses that end up back in hospital because of the consumer movement. (Consumer leader, WA)

Some research informants felt that the requirement to employ only consumers should be revisited and that having the skills required should be the priority rather than the status as a consumer.

The shift was the realisation that it was necessary to run the office as a professional entity and that it might require employing someone who is not a consumer to do that. It’s about employing people for their skills rather than because they need a job. There was the experience of having a person from the group and the person simply not having the skills to do the job properly and at other times becoming unwell and suddenly the whole thing falls down. (Consumer leader, ACT)

Those experienced at employing staff all emphasised the vital importance of skilled recruitment to consumer positions.

I learnt very, very quickly that you should select on merit, not because the job would be good for someone but because they would be good for the job. It will be good for them but you give it to them because they can do it and that can never ever be stressed too much. We actually demand an incredibly high standard from our staff and they are more professional than the professionals. They get monthly external supervision and they use it. The amount of supervision workers need is more than people who don’t carry the scars of mental illness. The business model is important because you have to employ competent people because you have to deliver against contracts. (Consumer leader, NZ)

2.7.3 Dealing with conflict

Conflicts and tensions are common experiences for community organisations including mental health consumer organisations and some voiced the opinion that ‘too many consumers with too many egos’ had resulted in a fragmented consumer movement across Australia. As one consumer said ‘I don’t know of many consumer organisations that don’t threaten to self destruct every now and again, it’s par for the course’. These difficulties have been attributed to a lack of effective and appropriate leadership and to ‘disempowered individuals becoming empowered and then not knowing how to wield that power for the benefit of others’. Informants described situations where a core group of consumers had effective control of an organisation and became increasingly intolerant of others who did not share their views.

We have individuals who have a lot of long standing involvement but have taken a lot of power and aren’t particularly ethical in the way they operate. It’s very difficult to get rid of people who have set up empires and who have their own issues which they project into their role and who don’t have supervision. They are not acting with a strong sense of self reflection and are often unethical. They bully others out of the way. (Consumer leader, NSW)

We’ve had a great deal of difficulty with this issue of leadership because when you have a group of people who have been marginalised and experienced a lot of powerlessness you need particular types of leaders. Some of the consumer leaders haven’t quite got that and in fact if you spend a lot of time in the mental health system you might have learnt a few things about leadership which won’t be useful to you. Leaders are about helping people reach and live their values which is a very different philosophy to traditional mental health services. (Consumer leader, NZ)

Informants saw ways of addressing these issues as strictly enforcing policies, procedures and codes of conduct to eradicate bullying and abusive behaviour
and having organisational structures which ensured accountability. This can be a lengthy process but means that organisations can develop a culture of respect where everyone is heard and treated with dignity.

We remove bullying from the management of these organisations by strengthening the processes and insisting on processes no matter what. Processes are our safest bet. There should be no bullying and everybody should be treated with respect. Often people are very gifted, very knowledgeable and excellent representatives but the problem is they don’t allow for the empowerment of anybody else. One of the core values of a consumer organisation is that people must be enabled to have a voice. (Consumer leader, NSW)

Informants also identified the need to clearly separate the policy and strategic direction role from the administration and day-to-day management role.

With organisational structures, we need to divide the roles very cleanly and divide the administration and accountability role from the policy and direction role. Those two roles need to be equal and accountable to one another with the chair or president as a third party – a triumvirate. They should meet regularly, discuss the business and advise each other respectfully and be informed from the membership through the chair and the policy person. The membership as a whole, through strategic meetings and consultations, should be the people who make the directional decisions in discussion with the person managing the administrative and policy area. Everyone should have checks on everyone else that the processes are being followed. The real mistake is that people get to represent the organisation but the other members never hear anything about it. (Consumer leader, NSW)

2.7.4 Factors for Success

What determines whether a consumer-run organisation is successful or not in attracting funding, sustaining itself and providing an effective service? There are a complex array of factors but key informants variously identified the commitment of the group, a culture of learning, training for members and support from a range of sources. They also identified having capable and skilled staff and professionally delivered and credible services, having good consultation mechanisms, the capacity to build productive working relationships with all stakeholders and having independence as important for survival.

It comes down to the dedication and commitment of the consumers involved. Funding does help and it does create more stability in an organisation. We have a very good culture of learning and we’ve received a lot of training over the years before incorporation. We also have the support of different people and the inaugural Board set up a reference group of interested individuals to advise and support the Board. So whilst there weren’t non consumers on the Board they had access to a group of non consumers to help support and advise them. That’s worked and they have always been there on tap for us. It’s also about the Board being proactive in meeting challenges and they have to be informed. If there are staff positions you must make sure the selection is full of integrity and that information doesn’t sit with one person. (Consumer leader, NSW)

The professionalisation of what we do has something to do with it and being focused on coming across as a professional body. There has also been a lot of support from the executive level of the public mental health service and on a broader level across Australia there has been a move towards more consumer participation. A lot has been achieved in terms of establishing the network and its consumer representative program as a credible player in the mental health sector in terms of being valued by other players for its expertise and input. (Consumer leader, ACT)
Having a receptive environment and particularly having a champion within the mental health sector was seen as essential by some informants:

_You have to have a champion within the health service who has some clout, who will listen to you and understand and to whom you can make points without them taking it personally._

_Without a champion when you criticise services they behave as if you’re attacking them and get very defensive. (Consumer leader, SA)_

There is a growing literature on the unique needs of consumer organisations and the development of programs to assist them to build capacity (Wituk et al. 2008). In the US there is a well established network of technical assistance through centres providing on site training and skill building in business management, leadership, communication and networking. Australia has recently seen the establishment of its first technical assistance centre, Our Consumer Place (see page 45), in Victoria and it is hoped that this will become a national resource. However like many other countries including New Zealand most consumer organisations have to find their own way and although funders may provide the financial resources they do little to develop the capacity of organisations (Orwin & Burdett 2009). This situation could benefit from in-depth study of successful and unsuccessful consumer enterprises and the cost factors involved in order to propose a suitable funding formula which recognises their particular needs.

2.8 Measuring the Impact

What is the impact of consumer participation and consumer run initiatives? If the aim is to move towards a better recovery-orientated mental health system how successful have they been? The evidence base is scarce and there is little literature on the effects or how widespread they are (Crawford et al. 2002). Although initiatives and organisations which receive public funds may be involved in independent evaluations of their work most studies are descriptive and there is no Australia-wide comparative study. People commonly say they are doing an effective job and having an impact but without the evidence base to prove it. As one consumer said ‘consumers are adverse to evaluative research about themselves and there is an attitude of I’m a consumer therefore I know what I’m doing’.

However what evidence there is indicates positive outcomes. Findings from research into consumer initiatives (Nelson et al. 2006) suggests that they can assist individuals with social support, empowerment, increased wellbeing, enhanced quality of life and reduced use of services. These outcomes can be achieved at a fraction of the cost of professional community programs. At the same time system level activism can result in changes in perceptions held by the public and professionals about the value of the consumer perspective and consumer run organisations and concrete change to service delivery, planning and funding.

A study designed to explore the effects of involvement (Simpson 2002) explored databases between 1966 and 2001 for systematic reviews of randomised controlled trials and other comparative studies. The research found five trials and a number of other studies where half considered that consumer involvement in managing treatment led to more satisfaction and less hospitalisation. The study found that providers trained by consumers had more positive attitudes to the lived experience. It also found that patients registered lower levels of satisfaction with services if they had been interviewed by consumers rather than non-consumers.

A systematic review was carried out in New Zealand (Doughty & Samson 2005) to identify the international evidence for the effectiveness of consumer-run activities compared to those run by mental health services. Consumer-run was defined as having self governance by consumers and consumer staffing and supervision, control of program policy and responsibility for program implementation. The review identified 26 primary data papers and two systematic reviews. Overall the research showed very positive outcomes for clients of consumer-run services including higher satisfaction with services, general wellbeing and quality of life. Although some studies did not show a significant difference between user-run and service-run activities, no studies reported any harm to users or that consumer-run services were in any way less effective. Despite these findings currently consumer run services worldwide receive very limited funding from mental health budgets.

Consumers themselves have raised concerns about whether consumer involvement, especially where the agenda is centrally controlled, actually detracts from improving
services and the quality of life. If consumer involvement mechanisms are tokenistic and do not necessarily impact on decisions and when there are insufficient resources for consumer priorities to be implemented is involvement a good use of consumers’ time and energy?

Some evidence also comes from outside mental health services, from the broader health sector. The Australian Government funded a project to review international and Australian evidence supporting consumer participation in generic health care (DoHA 2001). It concluded that participation in decision making at an individual treatment level results in improvements and access to good information helps decision making, supports people to manage their own health and results in more accessible and effective health services and a reduction in health inequalities. It also suggested that participation activities should use a range of methods.

What is required is systematic and comprehensive evaluations of processes and outcomes. The Cochrane Consumers and Communication Group7 is currently designing a number of reviews to find existing evidence for any positive or negative effects of consumer involvement in mental health service evaluation research, in consumer employment as providers of mental health services and in involving mental health consumers as trainers for professionals working in adult mental health services.

2.9 Future Directions

Key informants were asked how they would like to see consumer initiatives developing in the future. They identified broad changes in the environment which would impact on the consumer movement as well as developments in particular areas and a vision where consumer engagement becomes an integral part of practice in equal partnerships with practitioners. In line with international trends there will be a shift of mental health services into the community, into mental health promotion and prevention and into the development of partnerships and coalitions. This requires consumers to develop credibility and respect with professionals, agreement about what support is required, equality in status and a more united front. It also requires changing staff attitudes and staff culture.

Rather than using the language of participation and involvement we need to talk about partnerships. You have a better platform than if you talk about participation which is much more passive and about an agenda that has been set. Partnerships are the way of the future. It is a government imperative so there are ways in which it could be colonised. (Consumer leader, NSW)

These moves will be accompanied by a professionalisation of the consumer role with formalised representative roles, standardised peer support services, consumer-run and controlled services and a growth of consumer businesses. There were visions of an expansion in the consumer workforce, more specific roles in the education and training of clinicians, further inroads into consumer-controlled research and education programs for consumers to get them back on track in the education system. This will need to be accompanied by skills development so consumers can become more effective and by the establishment of technical assistance centres to assist in setting up and sustaining consumer organisations.

They will be funding us to provide information or services. We won’t be given money just to run groups but to produce goods or offer workers or take on contracts. That has been the pattern worldwide. We are getting far more professional than we were 20 years ago and governments won’t be funding us just to sit round and have talk-fests about advocacy. (Consumer leader, NSW)

As one consumer in New Zealand operating a consumer-run crisis intervention service said:

Having been involved for a long time, we made the decision that we would actually try and do something rather than just talking. We got fed up telling people what we wanted. We just started doing it. If we don’t like the services and they’re not listening let’s set up an

7 The Cochrane Collaboration is an international not-for-profit independent organisation dedicated to making up-to-date, accurate information available about the effects of healthcare worldwide. The Consumers and Communication Review Group coordinates the preparation and production of systematic reviews of interventions which affect consumers’ interactions with healthcare professionals, services and researchers.
alternative. It is really difficult to be accepted by the clinical-thinking funders that there is more needed to get people back on track than clinical services. It is really important to get people together who can start role modelling. That is how we got things funded, by showing them what could be done and starting off very small, showing them there was the ability and there were good results. (Consumer leader, NZ)

Some informants wanted to see a much more strategic approach and a closer alignment with other rights movements, with human rights organisations and particularly with the disability rights movement. They identified many shared issues, particularly in dealing with patriarchal systems and power imbalances, in moving towards a more rights based agenda, in focusing on the social rather than the medical model and in having strong organisational frameworks. However, internationally there have been difficulties in working collaboratively. Some consumers question whether mental illness is a disability which implies a long term unchangeable condition. At the same time there is a lot of “sanism” in the disability movement. And taking on both a patriarchal disability service structure and the mental health establishment is a big challenge.

The differences between disability and mental health are quite hard to negotiate. A lot of people with mental illness don’t see themselves as disabled and don’t see the advantage of campaigning on joint things around discrimination. Disability organisations don’t always get mental health and don’t want to be seen as mentally defective. All of us need a bigger picture and need to realise what the disability movement has done and how this could help us. We need to work out what we would gain if we called ourselves disabled and what would we lose. We just lack strategic thinking in the user movement. (Consumer leader, Victoria)

Consumers should be working with human rights organisations. The responsibility of the consumer movement is to try and make sure that consumers on the ground are aware of their human rights and to make sure people are treated properly. It is the one thing which makes a difference. (Consumer leader, UK)

2.10 In Summary

This chapter has illustrated the diversity of views, experiences and approaches in how best to engage with and use the lived experience of consumers to ensure that it informs a change in the way we think about mental health services. A key dilemma in this debate is whether, as consumers, it is better to work within services or remain independent from them. Informants illustrated both perspectives. Being employed within the system can offer distinct advantages in ‘subverting from within’ and having direct access to decision making forums. As two consumers employed at senior management and executive level said:

Initially I felt we should be employed outside the service in independent organisations. Now I have completely changed. When individual advocacy is the prime function you should be employed outside the service, peer support requires its own support structure or educating clinical staff where different skills are required. But if you work outside an organisation you are accused of not understanding the system. (Consumer leader, Victoria)

Some people said you can’t possibly be an advocate if you’re in the system but the job that I do is to subvert from within. I’ve been here two years now and I’m finding that I actually provide a lot of resources to staff. They say I am really interested in doing care planning and recovery plans or I want to know more about consumer participation but I don’t know how. So there is not so much a resistance as really not knowing how to do things. Being in house staff are more inclined to access me. From the outside trying to get in you get blocked off. I think I am able to say a lot more in the service than people
might have thought. There is real value in being at this level. (Consumer leader, WA)

There are also those who feel it is best to remain outside and that it is too easy to become subsumed by the system. Building consumer organisations which can provide autonomous and independent consumer services should be the goal.

Consumer participation has been imposed on services so if you’re a service which is not committed to it what you do is hire a consumer who you know will never be able to do the job. We have a few of them. Some start to identify with the service and then become apologists for it. They want to be colleagues with their colleagues and they have become colleagues of the clinicians. (Consumer leader, Victoria)

However many informants saw the advantages of both approaches and wanted to see models which combined the most positive aspects of working inside and outside traditional mental health services. As one informant in a consumer run organisation in New Zealand said:

The drawback is that we are outside the tent and our consumer advisers are external to the organisation. Elsewhere in NZ the district health board has set up peer support teams in the service itself with the intention of influencing the service culture whilst providing support services. They are more comfortable with someone they employ because they can control them. But how can one poor peer support worker in an in-patient unit change a culture with 150 years of history behind it? But it might have a greater impact on culture change than us as we are outside it maintaining our purity. (Consumer leader, NZ)
3. Mental Health Consumer Activity across Australia

This chapter maps the complex jigsaw of consumer activities and participation mechanisms across Australia and the structures at national level which draw these activities together. It has been collated from information provided by consumers and other stakeholders in all states and territories and from a range of policy, research and other material about particular initiatives. It highlights examples of good practice and particularly interesting and innovative approaches (which are boxed) and draws some comparisons between jurisdictions in terms of the history of consumer development and its current presentation.

3.1 The National Picture

National bodies can be important in providing a voice at the federal level, in setting national agendas, in promoting the consumer movement and in supporting developments locally. Without these structures there may be a lot of activity in individual jurisdictions but no mechanisms for bringing it together into a united voice. However, the ability to develop national structures is constrained by two issues. Firstly because the consumer movement has developed in a patchy and fragmented way across Australia there are questions about what infrastructure there is for any national bodies or coalitions to work with. Secondly while national organisations try to influence policy at the national level in reality most developments take place at a state or territory level. This means that it can be difficult to define concrete national agendas and to see positive outcomes spring from them.

Although there has been a growth in national advocacy organisations providing a voice for people with particular disorders, like SANE and the Schizophrenia Fellowship Council of Australia, the first all-embracing consumer and carer national body was the National Community Advisory Group on Mental Health (NCAGMH). This was established in the 1990s through the first National Mental Health Strategy as an influential committee of consumers and carers with representatives from each jurisdiction and its own secretariat, reporting to the Commonwealth Minister for Health. There was a parallel commitment at the state and territory level to establish CAGs of consumers and carers and give them direct input into policy discussions about mental health reform. It meant that for the first time consumers and carers were involved on every relevant federal committee and it led to the development of partnerships between mental health services and consumers and carers (Craze 2004). NCAGMH was disbanded in 1998 when the Mental Health Council of Australia (MHCA) was established.

In recent times the national scene has been dominated by four key bodies.

Australian Mental Health Consumers Network (AMHCN)
This was founded in 1996 as the only national consumer organisation for people with mental illness in Australia. It was an independent, incorporated body consisting of two delegates from each state and territory and aimed to provide a national voice in mental health service planning. It secured funding from the DoHA Mental Health Reform Branch to establish an office, a secretariat and a profile. During its life it organised conferences and events, issued a regular newsletter, initiated campaigns and was represented on government working parties, consultations and organisations at a national level. It also carried out a scoping study for the DoHA to explore workforce opportunities for consumers and the feasibility of establishing a national education and training strategy in consumer advocacy work.
AMHCN was defunded in 2008 due to concerns about its governance. There is an ongoing debate about developing a suitable successor to AMHCN and how to build on organised consumer activity in each jurisdiction to develop a new national peak consumer organisation.

**National Mental Health Consumer and Carer Forum (NMHCCF)***

This was established in 2002 with federal funding and is the only mechanism for bringing consumers and carers together at a national level. It aims to promote national partnerships at all levels, increase both participation opportunities and the capacity of consumers and carers to participate and provide a unified voice on consumer and carer issues. It also aims to identify best practice in participation and agree on national priorities for action in line with the National Mental Health Plan. The NMHCCF has one consumer and one carer representative from each jurisdiction and from national consumer and carer organisations. It operates under the auspices of the Mental Health Council of Australia (MHCA), meets four times a year and runs working groups to progress individual projects.

In 2004 the NMHCCF reviewed existing models of consumer and carer participation throughout Australia with a view to recommending the most practical working model at a national level (Craze 2004). It also published a guide for consumer and carer participation policy (NCCF 2004) which can be adopted by any organisation and which lays out the core best practice principles for defining the role, selection and employment of consumer and carer representatives. As well as setting up a website it is currently developing policy to improve its representative base, reviewing its strategic plan and identifying key priority areas. These include work around privacy and information sharing, seclusion and restraint and welfare to work.

**Mental Health Council of Australia (MHCA)**

This was set up in 1998 to provide representation from consumers, carers and providers at a national level. At the time many concerns were expressed that this would dilute the consumer voice because of domination by professional groups. It is now one of the first ports of call for those wanting mental health sector input at a national level, including consumer and carer input. As well as helping to establish the NMHCCF MHCA have been funded by DoHA to:

- support consumers and carers in surveying mental health services under the National Standards. The Standards require all public mental health services to conduct a survey by an external accreditation agency to review implementation of the Standards. MHCA organised a national training workshop in 2003 for consumer and carer surveyors and set up the Network of Consumer and Carer Surveyors to provide a national database of surveyors, a website, a quarterly newsletter, telephone contact, informal mentoring and an annual workshop.
- develop and trial mentoring and training opportunities for consumers and carers to strengthen their ability to promote issues and concerns at a national level. In 2007 this resulted in the development of:
  - the National Register of Mental Health Consumer and Carer Representatives. This involved selecting 50 consumers and carers from around Australia to attend a two-day workshop focusing on developing representation skills. It also provides ongoing training, support and access to representation opportunities.
  - the Mentoring for the Future Program. This gave opportunities to five consumers and five carers to participate in intensive training and mentoring to enhance their advocacy and leadership skills over a 10 month period. In August 2007 they travelled to the US and Canada to participate in the International Initiative for Mental Health Leadership (see page 51) exchange program. In addition they automatically become members of the Register and are linked with leaders with advocacy experience from other sectors for one-to-one mentoring.

**Private Mental Health Consumer and Carer Network**

Private mental health services account for approximately 16% of mental health beds and 43 hospitals across Australia. The Network was set up in 2002 with $13,000 annual funding from the Australian Medical Association, private health providers, the DoHA and the Royal Australian and New Zealand College of Psychiatrists. It has a committee in each state and in the ACT and aims to facilitate the sharing of experiences and concerns in order to lobby at a national level. The Network meets twice a year and communicates through a strong email network. There is no committee in the Northern Territory because it does not have any private hospitals.
The Network describes consumer participation mechanisms in the private sector as poor and attributes this to the absence of any mandate to establish them and the absence of accreditation processes requiring in-depth reviews against the National Standards as in the public sector. Some hospitals have consumer and carer advisory committees but they rarely have input into service design and delivery. There are also a small number of consumer consultants (six) working in private facilities and based in just two states. Tasmania is unique in having a consumer representative on a private hospital board. The Network is pushing for better involvement mechanisms in private hospitals.

**Other significant national initiatives**

These are:

- **The Kit.** This was a community development initiative which grew out of the National Mental Health Strategy and which, in 1998, resulted in the production of a comprehensive resource kit for consumers of mental health services and their families and carers (DoHFS 1998). It aimed to enhance advocacy activity through increasing knowledge and strengthening skills within the mental health community sector. MHCA was involved in disseminating The Kit and developing an infrastructure to enable its implementation. Five training modules were produced and consumers trained to deliver them. However, a subsequent evaluation found that although the resource had been disseminated Australia-wide there had been little training in how to use it in any of the jurisdictions. The Kit remains highly regarded in the mental health sector.

- **Australian Network for Promotion and Early Intervention for Mental Health (Auseinet).** This is a web resource which aims to inform, educate and promote good practice. It provides access to a comprehensive clearinghouse on best practice models, publications, a newsletter and links. These include access to consumer participation models and debates.

- **Consumers Health Forum of Australia.** This is a national body working to promote the consumer perspective and interests and to provide a voice for all health consumers. Although it does not have a mental health stream it does provide a useful source of information about models and issues in developing consumer participation across the health sector.

- **TheMHS Conferences** are held annually to create a learning network for improving mental health services in Australasia. They aim to promote collaborative research, practice, policy making and education as well as the active involvement and inclusion of all mental health stakeholders. They also advocate for and promote leadership in the mental health field and have an important role as vehicles for empowering and skilling up consumers.

Consumer representatives continue to sit on relevant federal committees and working groups. These include the recently established National Advisory Council on Mental Health set up to provide independent and confidential advice and expertise to Government on mental health issues as well as the advisory group developing the National Mental Health and Disability Employment Strategy.

### 3.2 Australian Capital Territory (ACT)

The ACT has a strong mental health consumer movement. Its size, both geographically and in population terms, and the close working relationships which have developed between organisations has fostered a strong consumer organisation which has quickly gained credibility and support across the sector. It also has a consumer and carer participation framework developed through a highly regarded, inclusive and open consultation process driven by the ACT Health Policy Unit where participants felt they had the opportunity to fully influence the process. The framework was launched in June 2007.

**ACT Mental Health Consumer Network (ACTMHCN)**

ACTMHCN began in 1997 with a drive to set up a place where consumers could feel safe. This resulted in the establishment of the Rainbow, based on the clubhouse model⁹ and offering a consumer space for sharing experiences and resources. This success motivated a push to set up a more advocacy focused organisation and the Network was formed. It incorporated in 1999 and acquired a small health promotion grant of $5,000 per annum to set up an office with a part-time

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⁹ The clubhouse model of psychosocial rehabilitation was set up in 1948 as a self-help organisation and is now an international movement providing non-clinical community based programs to assist people with mental illness to reclaim their lives. There are now over 300 clubhouses in 24 countries. Provision is focused on strengths and abilities not illness and members and staff work side-by-side to manage all operations.
It was initially housed with the Mental Health Foundation and there were difficult times and challenges when the coordinator was unwell and the organisation came close to collapse. For a while the Network was auspiced by ACTCOSS which provided staff mentoring, support and financial management. A new influx of members and ongoing funding increases from ACT Health allowed it to regain its independence in 2004 and establish a sound and credible organisational structure with a strong volunteer base.

ACTMHCN currently have a management committee of eleven consumers, a full time executive officer, two part-time administrative staff and a growing membership of 140 people. It aims to improve mental health services by providing systemic advocacy through regular consumer forums and consultative mechanisms to share information and identify common policy themes. Its key service is running the consumer representative program and most of the annual ACT Health grant of $260,000 is devoted to this work. The program provides one gateway for requests for consumer representatives. There is a rigorous nomination and selection process for the representative pool and ACTMHCN also provides training, ongoing support, accountability and the development of consumer policy positions. Following a New Zealand model (see page 53) there is now some debate within ACTMHCN about having employed representatives and creating a professional body backed by a sound consultative process.

Network informants describe a maturing of the organisation with the recent acquisition of additional funding to employ two more part-time staff to oversee the consumer representative program and develop more policy capacity. It would also like to invigorate its education program in medical schools and with mental health staff. This would support more informal work where mental health services have begun to routinely invite consumers to participate in training activities. The ACTMHCN has been facilitating narrative sessions to bring consumers and professionals together in power-free environments and it hopes to expand this area in future. The Network is now highly regarded in the sector although the downside of this is an increasing workload with limited resources.

There has been little emphasis on developing a consumer workforce in the ACT although two consumer consultants have been employed by the mental health agency for the past decade. When they were first employed they tried to do everything; systemic and individual advocacy as well as peer support. However, over the years their role has gained clarity and its core is seen as being systemic advocacy. They participate in policy forums and at an executive level in mental health services and are involved in educating staff and visiting community mental services to speak to consumers.

Like a number of other jurisdictions there is currently little paid peer support in ACT although it is suggested that ACT Health may be interested in funding a peer support pilot program. ACTMHCN is also interested in setting up peer support but without funding this would have to be on a voluntary basis. This raises concerns that using volunteers detracts from the possibilities of paid employment. The Mental Health Foundation currently operates a peer support program with seven peer support workers who are reimbursed at $40 per hour and are trained, de-briefed and given clinical supervision. Funding for the program has come through health promotion but the Foundation is hopeful that ACT Health will pick it up as they regularly express interest in this area.

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ACT Consumer and Carer Caucus

This was established by the Community Mental Health Coalition – the peak body for the mental health sector – and represents a unique model for ensuring consumer and carer participation in a peak mental health sector body. It was conceived as more than a consumer advisory body and has a link to the Board of Management whereby the Caucus nominates two carers and two consumers to sit on the board, raise policy and service development issues and be involved in the strategic direction of the organisation. It has been an important body in policy and planning issues and in bringing consumers and carers together and has run effectively for a number of years. It is now being reviewed by an independent consultant to reinvoke it, improve the participation structures and ensure meaningful involvement. This will also entail exploring ways of funding a coordinator position for the Caucus.
The ACT branch of Mental Illness Education Australia – the Mental Illness Education Group – works to raise awareness of mental health issues, reduce stigma and discrimination, encourage health seeking behaviour and promote the recovery message using the personal story rather than the collective experience. It educates the Canberra community about mental illness through presentations by consumer volunteers, websites, theatre projects and community consultations. The core business is the school education program which uses consumer and carer volunteers telling their personal stories. Volunteers go through a rigorous training and then attend practice sessions to observe until they are confident enough to join in. The Group is now also working with community programs tailored towards particular audiences as well as the police, mental health providers, bus drivers and prison workers. Consumers have a strong role in governing and running the organisation and 80% of the board are consumers. They are not seeking registered training organisation status and wish to remain a small organisation.

3.3 New South Wales (NSW)

NSW presents a diverse picture with a number of different centres of activity. Although there is currently no state wide framework for mental health consumer engagement there are now moves to develop one.

There is one state wide CAG which is engaged in systemic advocacy and CAGs operating at a local level which have become activity focused. Although some have become consumer-only groups including NSWCAG, many have retained the consumer and carer focus. For example, Mid-Western Consumers and Carers Advisory Group is an incorporated organisation which manages the O’Brien Centre in the grounds of Bloomfield Hospital. The Centre is open one day a week and provides a safe and supportive social environment offering a range of activities and acting as a stepping stone to the community.

**NSW Consumer Advisory Group – Mental Health Inc (NSWCAG)**

This is a statewide incorporated organisation providing an ongoing mechanism for consumer participation in mental health policy, service development and evaluation. It is governed by a board with a 50% consumer membership and a staff of one full-time executive officer, a part-time policy officer and two part-time administrative officers. It has gone through many changes and challenges over the years but is now seen by NSW Health as the peak body and has developed a good working relationship with mental health services. It receives $246,000 per annum from the Mental Health, Drug and Alcohol Office, the Centre for Mental Health and NSW Health with additional project funding. Although it represents the interests of both consumers and carers, given the existence of a strong carer organisation, the main focus has become the consumer perspective.

NSWCAG has a website, a weekly e-news, a quarterly newsletter and a mailing list of over 1,000 consumers, carers and other stakeholders. It maintains a pool of consumer representatives and has become a one-stop-shop for finding a consumer representative. It has produced a DVD on stigma and in 2007 worked with the research unit at Rozelle Hospital and consumer consultants to develop and trial four workshops for consumer workers and carers about helping people with co-morbidity issues. This was funded through a NSW Health Innovation Scholarship. In future it hopes to consolidate its role as the peak mental health consumer body in NSW, develop more formalised links with existing consumer networks and explore training gaps for consumers and consumer workers.

Like other jurisdictions there is a vigorous debate about the role of the 52 part-time consumer consultants employed by area mental health services in NSW. This debate has centred around the lack of role clarity and consistency in job titles, inappropriate awards and a lack of training and consistent guidelines. Because of the autonomy given to area mental health services to implement the role some areas have developed more strongly than others. For example, the Mental Health Consumer Participation Coordinator at St Vincent’s Hospital works 20 hours per week and is involved in sitting on committees, running issues workshops, community development and setting up a Recovery Working Group. This contrasts with other areas where there are serious concerns
that consumer consultants are unable to engage in systemic advocacy work due to fears about losing their jobs.

**The Consumer Activity Network (CAN)**

This developed from the Northern Beaches CAG and is the sole consumer-only organisation in the state. It aims to advance the empowerment and independence of consumers and facilitate participation in mental health services. CAN incorporated in 2005 and in order to diversify funding sources and attract philanthropic money acquired deductible gift recipient status. Its incorporation led to a withdrawal of health service support and a period of reliance on fundraising and volunteer resources and in the absence of core funding it has survived on project grants. Currently it is funded to deliver the DoHA Community Connections Project. This has two major components:

- Phone Connections (or the Warm Line) providing a national non-crisis, peer support and information line which runs for four evenings a week; and

- the Hospital-to-Home Team providing practical assistance and peer support to consumers within the first 28 days of discharge from in-patient units. A team leader and support worker provide the service five days a week with a very broad base of practical support including transport home, assistance with shopping, follow up phone calls, home visits and facilitating access to other organisations and support. It has supported 34 people in the last quarter and identified a noticeable decline in readmission rates.

CAN also runs the Pitane Recovery Centre – a mobile consumer-run recovery centre providing weekly activities with a focus on education, training and recovery. It has established the only peer support worker course in NSW and hosts the annual NSW Consumer of the Year Awards. Although it does not do advocacy work individual members may lobby on particular issues.

The organisation is run by a Board of Management but it is rarely able to fill its capacity for nine members and has had difficulties in sustaining a stable consumer Board. The inaugural Board did set up a reference group of interested individuals, including non-consumers, to provide advice and expertise which has been valuable. It has a staff of 3.5 full-time equivalents (FTEs) with the development manager being the only full-time worker and also employs some casual staff. Given the reliance on project funding CAN’s main issue is sustainability.

The consumer workers forum brings consultants together to exchange experiences and provide mutual support. The forum recently successfully lobbied the government to commission a report on the situation of consumer workers and a project worker will be employed to explore the role, awards and training issues and to set proper parameters for consumer worker positions, wages and minimum standards. There has also been some discussion about NSWCAG becoming the employing agency for consumer consultants across the state. Some informants consider that this would provide support, prevent isolation and burnout and reduce staff turnover as well as enabling better systemic advocacy work.

**The Mental Health Consumer Perceptions and Experiences of Services (MH-CoPES)**

This is a partnership which began in 2004 between NSW Health, the Centre for Mental Health and NSWCAG to develop a process for consumers to evaluate and improve public adult mental health services across the state based on consumer feedback. It has been led by a technical working group of eight consumers, three mental health staff and staff from the Centre for Mental Health and developed through a series of consultation forums. The team is currently completing the development of an instrument which uses a three-point rating scale measuring whether major or minor improvement is required to services (NSW Health 2006). The instrument is a questionnaire completed every three months by all consumers with versions being developed for in-patients, those using community mental health services, child and

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8 The Centre for Mental Health is responsible for developing, managing and coordinating NSW Health Department policy in relation to mental health services.
adolescent services, older people, Aboriginal and CALD populations. Ultimately the aim is to link MH-CoPES to the funding and accreditation process. However, it has encountered difficulties in gaining commitment from health services and there is now a realisation that a lead is required from within services to ensure surveys are distributed at ground level. It has been recommended that an independent consumer-directed organisation be established and funded to support MH-CoPES, to act as a central data collection point, undertake the analysis and publish and distribute the reports. However, it is also considered that the questionnaire is only part of the solution to genuine consumer participation in evaluation.

The NSW Institute of Psychiatry has provided a three day foundation course for consumer advocates and a three day extension course. The course was designed by consumers, has been running for five years and has trained over 200 people. It provides a basic grounding in the spectrum of advocacy work covering policies, legislation, discrimination and human rights, committee work and so on. Although it is not accredited, it is approved by the Institute and the feedback has been positive. However, because health services are no longer funding consumers to attend the course, numbers have dropped and this has threatened its viability.

Overall the general feeling in NSW is that despite the fact that there are formal mechanisms for consumer and carer participation in place, they do not necessarily comprise meaningful participation. In addition, the diversity of consumer initiatives has mitigated against a strong and united consumer voice in the state.

### 3.4 Northern Territory

The Northern Territory has a small population base in a geographically vast area with a large Aboriginal population. This presents logistical problems in ensuring involvement from Aboriginal people and rural and remote communities. There are differences between regional centres and although there is no consumer and carer participation framework the framework developed by the NMHCCF is used as a reference point.

The Territory has stayed with the CAG model. The Northern Territory Community Advisory Group on Mental Health (NTCAGMH), formed in 1992 to act as a ministerial advisory committee and provide strategic advice on the planning, delivery and evaluation of mental health services, still performs this role. It meets four times a year and aims to have 50% consumers and 50% carers as well as a balance of remote and Aboriginal members. It has an annual budget of approximately $20,000 and members have chosen not to be paid for their services although no one is allowed to be out of pocket. There have been issues in sustaining the CAG and questions have been raised about how far it is an effective mechanism for involvement. However, the past twelve months have seen a redevelopment and a renewed commitment from the Department to help sustain it and ensure that its role is not tokenistic. It is now seen as moving in a more positive direction and hopes to fill gaps in networking with key groups.

The only consumer-run organisation in the Northern Territory is the Top End Mental Health Consumer Organisation. This operates some 25 kilometres outside Darwin and was set up ten years ago by consumers to provide support, information, group activities, and individual advocacy to those leaving acute facilities. It incorporated seven years ago and is governed by a Board with a membership of about 150 people from across the Northern Territory. Last year it acquired funding of approximately $50,000 from NT Health to rent an office and employ a worker to manage the organisation and do administration work. The work also encompasses encouraging the development of support groups in other regional centres and especially rural areas. The organisation would like to attract more involvement from Aboriginal communities.

**The Mental Health Association of Central Australia (MHACA)**

This provides a number of recovery and housing programs, group activities, training and forums in Alice Springs. It also undertakes systems-based advocacy through representation on a number of local, state and national organisations and assists consumers to speak out. It has developed a consumer forum which meets monthly to discuss issues and how staff can improve services. It is run by consumers for consumers and, because it is held in the evenings, includes a dinner with
a regular attendance of eight to twelve people. There is also the Consumer Action Group Lunch, again meeting monthly, to put the ideas from the consumer forum into action. In addition consumer involvement is sought at all other levels from conducting a monthly feedback survey at the drop-in centre, offering casual work opportunities, having consumer representatives on the management committee, consumer participation in staff selection and supporting consumer attendance at conferences.

MHACA now wishes to develop peer support training based on the South Australian model (see page 42) and which could also offer mentoring and support as consumers move into employment. It has been proactive in encouraging participation from Aboriginal people and building up numbers through word of mouth approaches and translation of information into local languages. MHACA have found that consumer involvement work is complex, long term and requires flexibility, fluidity and constant change. However, it has also proved very motivational for other consumers.

3.5 Queensland (Qld)

Queensland is described by informants as having positive levels of activity and a number of good initiatives. A draft framework for consumer and carer participation was developed 18 months ago (Burge 2007). This is now being reworked for a launch in June 2009 which, it is hoped, will consolidate activities and cement a consumer workforce pathway. There are concerns that consumer developments have been delayed because of restructuring within the mental health sector.

The state has followed the CAG model with the establishment of both local and state wide CAGs. The Queensland Consumer and Carer Advisory Group (QLDCAG) was established in 1993 to provide strategic advice to the minister about the experiences of consumers and carers in the mental health system. It incorporated, became independent of the Department and operated as a focal point for local CAGs and assisted in their development. By 2001 there were over 30 local CAGs. QLDCAG was eventually defunded in 2004 leaving a large gap in participation mechanisms and with no centralised body with direct input into government. The local CAGs experienced problems with funding and support and diminished in number.

There is now a network of approximately 15 CAGs across the state at local and regional levels and the majority of health districts actively support them in some way. A few have also been able to attract external funding and run projects and two are now incorporated bodies. However, they are usually dependent on the goodwill of the district which means that some can act autonomously and others are tightly controlled. There is also no pivotal point which brings them together to collate experiences. Vanderburgh CAG was the first to incorporate in 2003 spurred on by a difficult history of auspicing by a variety of different organisations. It currently has no funding. Toowoomba CAG also incorporated after eight years of being auspiced by Queensland Health and changed its name to the Consumer Advisory Network. It now has a memorandum of understanding with the local mental health service and $50,000 per annum to provide the formal mechanism for consumer and carer participation in planning and development across the service and for quarterly meetings with the Director of Mental Health Services. They supply representatives for committees on request and employ eight part-time peer support workers. Cairns Consumer and Carer Advisory Group has substantial funding, is influential in the district and is now looking to incorporate.

Principal Officer, Consumer and Carer Participation

In 2006 the Mental Health Branch established this permanent post to provide leadership and support to the Minister’s office and to district mental health services around consumer and carer participation, national standards and accreditation. The position also aims to develop initiatives and support a state wide network of consumer and carer workers with monthly meetings and an annual conference. Although the post covers both consumer and carer issues the primary focus is the consumer perspective. Recently funding was acquired for a 12 month senior project officer and consumers are also regularly employed on a casual basis to do administrative work. A priority for the post has become the development of a consumer and carer workforce pathway within mental health services. It has also set
Experts by Experience: Strengthening the mental health consumer voice in Tasmania

The Consumer Companion Program

This was established by the Principal Officer, Consumer and Carer Participation in 2004 to support in-patients who were reporting loneliness, boredom and lack of support. This led to a state wide pilot funded by the Mental Health Branch to provide peer support — or consumer companions — based on the philosophy of recovery and including both training and evaluation. A positive evaluation and full support from mental health staff has meant that the program is now being rolled out across the state with funding of $500,000 per annum. This is being transferred to each district for local implementation but with support and guidance from the Principal Officer. Companions are remunerated at $18.50 per hour plus travel and receive a two day training package delivered locally by the Principal Officer. Currently there are 43 companions but the aim is to expand to 120 over the next 12 months.

In addition to the Consumer Companion Program there are currently approximately 39 (or 25 FTEs) consumer workers across the state working in all districts and known variously as consumer consultants or recovery support workers. They operate at both a strategic level and are also engaged in more practical support work. There is also a push to get a consumer consultant into every service as a systems advocate. Although there are no dedicated full-time peer support workers there are some paid peer support positions but it is confusing because some of the consumer consultants are also doing peer support work. Like other jurisdictions there is debate about a lack of clarity for the roles or the skill levels required and concerns about employment purely on the basis of having lived experience. There is now growing pressure on Queensland Health to employ a project worker to explore workforce pathways and how best to train and develop the skills needed for different specialisms including consumer representatives. There is a commitment in the State Mental Health Plan to build the consumer and carer workforce. However, decisions about how this should occur are the responsibility of the districts.

Developing a Consumer and Carer Organisation

There is currently a process to develop a consumer and carer-run state wide organisation in Queensland funded by Queensland Health. This is being fostered by the Queensland Alliance, the peak body for the mental health sector, and is called the Consumer Voice Project. It has entailed a state wide consultation with consumers and carers to develop an accountable mechanism for participation, a draft constitution and a business plan for the new organisation. The project has now attracted $320,000 from the Mental Health Branch for two years to develop the organisation, build membership and create an ongoing structure. This pays for a full time worker, some travel time and the capacity to get people together once a year. It has a group of 25 consumers and carers acting as a wider reference group and a core group of six people who undertake most of the work.

There have been difficulties in working with project officers and building the capacity of the committee but the intention is to have it initially auspiced by Queensland Alliance but moving towards incorporation within the first year. It aims to undertake systemic advocacy and provide consumer and carer mechanisms for engaging at a high level with government and departments. Part of the core work is also envisaged as supporting other participation mechanisms in the community, for instance CAGs and consumer and carer support groups.

The Brook Recovery and Empowerment Centre (BrookRed)

The present site of the BrookRed Centre was originally purchased and used by the local hospital as a mental health out-patients clinic and drop-in centre in the early 1990s. In the late 1990s workers at the building put in a funding submission to have their services auspiced by a non-government organisation. BrookRed was the result of that process. The Centre incorporated in 2002 and now offers a peer run, peer support service providing a safe space for people to meet and be themselves whatever the state of their health and a variety of activities and groups chosen by and run with people who attend. In 2007 DoHA granted funding for a two year project under
the Day-to-Day Living Program to extend the running of peer support at the Centre. Through this funding approximately 60 people were able to obtain individual peer support from a trained peer worker of their choice. The Centre operates a five day-a-week service with additional support from Disability Services Queensland. It has a total of 180 people on its books and regularly sees up to 90 people. The Centre uses a range of in-house and international training for peer support workers based on the intentional peer support model\(^{11}\).

All of the membership and most of the Board have a lived experience of mental illness. The Board is supported by a ‘think tank’ consisting of members of the community who offer advice on governance and direction.

3.6 South Australia (SA)

Mental health is a major area of reform in South Australia initiated by the Social Inclusion Unit’s Plan for Mental Health Reform 2007-2012\(^{(SIU \text{ 2006})}\). This pushes for participation at all levels of the system and requires ‘innovative ways of ensuring meaningful involvement in planning, organising and evaluating services’. The Department has chosen to interpret this in many different ways including writing a requirement to use the lived experience into funding contracts. However, currently there is no state wide participation framework.

Health regions across the State have developed their own participation frameworks and a key mechanism has been the establishment of local CAGs reporting to a regional CAG. There are currently approximately 12 CAGs in both metropolitan and rural areas, with the majority attached to psychiatric hospital facilities. South Australia has a history of consumers and carers working together and the majority of CAGs represent both. Recently each region was given $30,000 for participation activities. This was interpreted in different ways with some regions inputting funds into local CAGs; for example South Adelaide Health Service now has $12,000 per year to pay consumer representatives and support local CAGs.

There is no peak consumer/carer organisation in South Australia although the Social Inclusion Mental Health Consumer Advisory Group, which meets quarterly to advise the Social Inclusion Board, has been discussing forming a peak consumer group. In the past there was a South Australian CAG (SACAG) which operated as a peak body until the establishment of the Health Consumers Alliance (HCA). As one consumer said ‘the wheels fell off’ the consumer movement when the SACAG was replaced. The HCA covers the health spectrum as the peak body for all consumer complaints and advocacy and was set up to ensure resources were not dissipated through the duplication of participation mechanisms. It is required to ensure that mental health consumer and carer participation is active in the development of relevant services and policies and to this end established the Mental Health Taskforce which was used as the state peak body for mental health consumers and carers.

Permanent Half-Time Consumer and Half-Time Carer Consultant

These posts were recently established in the Mental Health Unit to explore mechanisms for participation and to counter the lack of a procedural approach to consumer representation where active consumers were ‘tapped on the shoulder’. These appointments are seen as demonstrating a real commitment from the Department to participation. The posts hope to move beyond a committee representative approach and ensure the consumer voice is heard at a strategic level. They are currently forging links with consumers and looking at how consumer and carer participation frameworks used regionally and locally might be developed into a state wide framework.

South Australia has a number of consumer and carer initiated and run rural Mental Health Activity and Resource Centres providing a range of recreational and recovery activities. Most are run by volunteers with little funding but some have incorporated and are able to support paid coordinators.

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\(^{11}\) Intentional peer support is a process where people use the peer support relationship to support and challenge each other and to look at things from new angles. This process has been championed by consumer leaders in the USA.
Murray Mallee CAG and Mental Health Activity and Learning Centre

The CAG incorporated in 1997 and is the only rural CAG to do so. It aims to work with the health service to improve services, support consumers with individual problems, mentor embryonic groups, evaluate services and promote the mental health consumer movement in South Australia. It has also been providing representatives on a range of local mental health forums, distributing a monthly newsletter, organising events during mental health week, running writing competitions in local schools and organising a community and business leaders breakfast. It has no paid staff and is run by a management committee of 10 who meet monthly. It reported a slow process spread over a number of years of developing a relationship of trust with the local health service who, by 2002, were providing in kind support – printing the newsletter, providing meeting rooms and covering postage costs. The CAG is supported by $4,000 from Country Health.

Over a five year period it was successful in attracting funding to develop a Mental Health Activity and Learning Centre (or MHAL) – a psychosocial centre providing a drop-in three days a week and a range of activities including a consumer camp. The MHAL receives a grant of $52,000 over two years through the DoHA Day-to-Day Living Program to provide 15 drop-in places a year with each worth 100 hours of contact time. It regularly sees 15 consumers and carers a day. The grant also funds the administrative work and reporting requirements. However there are currently no paid staff and the program is run by trained volunteers. It has recently received $40,000 from the MHCA to develop a marketing plan and do some development work and Country Health is considering funding a coordinator.

One of the most developed of the Activity and Resource Centres is the Station Centre, established in 1998 and auspiced by the Mood Disorders Association of SA. It is now incorporated and governed by a management committee of one professional, three to four community members or carers and three to four consumers. It receives $35,000 from Country Health and other one-off grants to run activities supported by 26 coordinator hours per week and administrative support.

The One Voice Network has recently been set up and incorporated to operate as a peak body and network for the Mental Health Activity and Resource Centres. It aims to present a united voice and provide a platform for advocacy, consumer involvement and consumer driven recovery orientated services delivered in local communities. The Network is currently lobbying Country Health for funding for a secretariat.

South Australia has a consumer workforce backed by a departmental commitment that they are an integral part of a quality mental health service. The term or title most commonly used in the State is peer specialist worker or consumer consultant and they are seen as vehicles for providing a consumer voice in policy, assisting in changing the culture and helping with staff development. They also offer psychosocial, non-clinical support and are involved in quality control mechanisms. There are currently approximately 80 consumer consultants and peer specialists based in acute public mental health services and in CSO-run community rehabilitation facilities in metropolitan areas, including a mix of paid and volunteer staff. Most major in-patient units have some consultants working on a part-time basis of up to 20 hours a week. As elsewhere there is much debate about their role and about the kind of training they should receive.

Training and Support for Peer Workers

The Mental Illness Fellowship and Baptist Care SA have been funded to run training and support for peer workers. This includes a “train the trainers” package as well as working with organisations so that they are ready to employ peer workers. They are currently looking to match the training with Certificate IV. They have recently developed an internet site for peer workers and employers to share experiences and offer online support with information about training, peer meetings, job vacancies, events and online mentoring. The website also offers employers a toolkit to guide the employment of peer workers, advice about roles, job and person specifications, recruitment and workplace support. The project has recently been positively accepted and is establishing new peer services with the MHCA and Country Health.
evaluated and the importance of organisational leadership for peer work identified as significant, especially during the transitional phase. The training is now identified as an example of good practice and attracted interest from other states. Unlike other training it is unique in offering ongoing mentoring and support to peer workers as they move into employment.

However, overall consumers in South Australia feel the state is lagging behind other jurisdictions. They would like to see more independent advocates working outside the system to counter concerns about tokenistic participation and a more genuine partnership at a case management level or where consumers actually experience the service. They want to see an expansion in consumer employment and clear pathways for engagement up to ministerial level rather than just ticking the consumer box.

3.7 Victoria

Victoria is often held up as the most positive model of mental health consumer involvement in Australia and involvement activities are well advanced in public mental health services compared to other jurisdictions. This is against a backdrop of a more systematic approach generally to building and enhancing consumer participation throughout all public health services. This has been pushed by the Department of Human Services’ (DHS) consumer participation policy, Doing It With Us Not For Us (DHS 2006). However, despite the development of a draft mental health consumer and carer participation action plan two years ago the plan was never ratified and was not well received and although each area mental health service has been encouraged to develop a policy around consumer and carer participation, as one consumer said ‘this generally meant rubber stamping old documents’. Currently there no state wide policy or framework for mental health consumer participation and there is now a push from consumers for the DHS to develop one.

The Victorian Mental Illness Awareness Council (VMIAC)

This was established in 1982 and incorporated in 1985. By 1988 it had attracted recurrent funding from the DHS Mental Health Branch and now receives approximately $464,000 per annum from the State and a further $73,000 from the Commonwealth. It has seen an interesting development from its origins as an advocacy body to branching out into service delivery, peer support and consumer research. It now operates as Victoria’s peak consumer body and provides a mix of individual, group and systemic advocacy, mutual support, education, training and information. It is also involved in development work to establish consumer groups and there are now 82 across the State. It conducts and auspices consumer driven projects and consumer perspective research and supports and educates consumer representatives sitting on 57 different committees. Eighteen months ago, in partnership with DHS and Carers Network Victoria, VMIAC conducted a pilot evaluation project about consumer/carer experience in eight mental health services across Victoria and disseminated the results to clinical and non-clinical sectors. It is hoped that this will now become a permanent activity. It also supports CORE – the Consumer Operated Research and Evaluation Support Group – which is open to all those interested in research and evaluation and funded by MHCA.

VMIAC is staffed by a director, one rural and one metropolitan advocate running groups in in-patient facilities, an information officer also doing telephone advocacy and support, an office manager and volunteer staff. There is also a young persons’ group and the membership operates as a training ground for future active consumers and as a pool of volunteers. Like other consumer-run organisations its history has been volatile and it passed through a period when it was not successfully engaging with its membership and saw a number of short lived appointments to the Director position. This led to the threat of defunding. However, for the past 15 years it has been under the same directorship which has stabilised the organisation as
has the development of an internal code of conduct
to promote a respectful organisational culture. VMIAC
is currently approaching the Department for funding
for an advocate for each area mental health service.
It would also like to employ a full time education
officer and to provide alternative health therapies
like massage, yoga and meditation classes.

Achievements in Victoria have been attributed to a strong
history of health participation work generally and of CSO
development across the board. Two manifestations of this
have been firstly the early establishment of VMIAC and
secondly the establishment of the Health Issues Centre twenty
years ago. This is a state funded CSO promoting consumer
perspectives in health care. It runs a consumer nominee
program that trains and links consumers to government and
to health services to increase consumer participation.

Unlike other jurisdictions there are no CAGs in Victoria. During
the 1990s agreements stipulated that all states must establish
a CAG in order to attract Commonwealth funding. The existence
of strong organisations like VMIAC meant a duplication of roles
and the CAG model was abandoned. However, all metropolitan
public health services and five regional health services are
required by legislation to establish community advisory
committees and develop community participation plans.

During the mid-1990s VMIAC auspiced two projects which
have been described as ‘breakthroughs’ and ‘landmark’
pieces of consumer research in Australia – the Understanding
and Involvement Project (ed. Wadsworth 2001) and the
Lemon Tree Learning Project (Epstein & Shaw 1997).

The Understanding and Involvement Project

This was a three year research project funded by
the Victorian Government to develop a set of built in
processes for consumer participation within mental
health services which could become an ongoing part
of planning, development, evaluation, staff training and
delivery. It presented a model for the development and
implementation of consumer consultancy in clinical
settings which was comprehensive and systematic,
and which placed emphasis on consumer participation
being intrinsically linked to the discourse of quality
assurance, staff-consumer communication and cultural
change. Attention was given to the importance of
consumers acting as consultants to staff as well as to
consumers. It identified ten rules for the successful
employment of consumer consultants and resulted in the
publication of a Job Manual for consumer consultants
in mental health services (Epstein & Wadsworth 1996).

In late 1996 the first four consumer consultants were
employed at the Royal Melbourne Hospital and a year
later DHS provided what became recurrent funding for
the state wide introduction of consumer consultants in
every area mental health service in Victoria. In recognition
of the influence of the U & I Project, a compendium
of findings was published (ed. Wadsworth 2001). The
research also identified ten key methods of consumer
participation – end of stay reviews, suggestion boxes,
complaints procedures, individual advocacy, spot
surveys, community ward meetings, involvement in
treatment plans, sensitive interpersonal conversation,
obtaining consumer input into staff training and involving
consumers as part of management structures.

The Lemon Tree Learning Project

This was consumer run, auspiced by VMIAC and
funded through the National Mental Health Strategy.
It aimed to develop an effective model of training for
consumer participation through building on the U
& I project and was the precursor to the idea of the
consumer perspective in the education and training
of the workforce. It used the analogy of a lemon
tree to demonstrate the connectedness between the
roots of consumer empowerment and a healthy tree
of service structures and policies. It identified:

- root learning where training nourishes an
  infrastructure to support consumer participation;

- branch learning where training adds to or
  modifies existing knowledge, skills and attitudes
  around consumer participation; and
• leaf learning where training provides information about consumer participation where little or no infrastructure to support it exists.

To build consumer participation requires root work which is labour intensive, time consuming and about developing trust and relationships so that the consumer perspective is experienced as valuable. The project developed ways in which consumers can work as educators of staff to promote effective participation. These include using consumer facilitated discussion groups and deep-dialogue learning through small groups of consumers and staff meeting on a regular basis. It used story telling as a powerful way to get a point across and developed a board game to help staff learn about the consumer perspective.

Victoria has one of the most developed consumer workforces in Australia and there are currently approximately 60 consumer consultants employed by local adult clinical services across the state. Most operate part time – one or two days a week – and work in pairs. Their role is to monitor the consumer experience and advocate for change through committee work at the local service level.

However, across the consumer consultant program and as in other jurisdictions, there is a continuing debate about the lack of role clarity, poor employment conditions and wage structures, lack of career pathways and the common expectation that consumer consultants are ‘expected to do everything’. So although the program has raised the profile of consumer needs and meant that they are now better addressed at local service delivery level the consensus is that it was never properly implemented as conceived in the U & I project. This is attributed to the fact that local areas were given autonomy over how they established the role which led to a range of interpretations. It has also led to some debate about consumer consultants being employed by an independent organisation although there is also strong support for retaining the program within mental health services.

Consumer Consultancy Work in the Southern Health Region

The Southern Health Region is the largest health care provider in Victoria with six main hospital sites and 25 primary care sites. Here a Director of Consumer and Carer Relations was appointed in 2005 in an executive level management position to oversee a unit composed of 10 consumer and carer workers. The post required tertiary qualifications and the role was to manage the workforce and provide a lead on consumer and carer participation at executive level. The post has been filled by a consumer who has faced challenges in working within a professional clinical environment and managing both consumers and carers. The unit engages in service and systemic advocacy and supports self-advocacy and access to individual advocacy. It is currently developing training for workers and for clinical and non-clinical staff, embedding participation in service delivery and lobbying for more funding.

Many CSOs in Victoria also have consumer consultants. Some informants felt that being able to operate in non clinical environments had been more successful although there is also debate about their role and the divide between individual and systemic advocacy.

There are no paid peer support workers in Victoria although some consumer support groups and VMIAC have volunteer peer workers. There are also non-government organisations which have established consumer advisory groups which may involve some peer support work; for example the Mental Illness Fellowship runs a consumer advisory group where members are paid to attend meetings.

Our Consumer Place

This was established in 2008 with funding from the Mental Health Branch. Its origins lay in an interest by the Branch in developing consumer delivered services and they engaged a consumer to put together a proposal for developing a model for Victoria. This recommended establishing a resource centre for consumer services and a needs analysis to find out what groups already existed.
and what resources were required to sustain them. Our Consumer Place now provides technical assistance to established and fledgling mental health consumer groups initially in Victoria but with the intention of growing into a national resource. It operates under the auspices of Our Community – a technical advice bureau working across the community sector – and it was considered important to locate it with the mainstream community sector and not in a ‘mental health ghetto’. It is modelled on the technical advice bureaus of the US and Canada and aims to offer information and advice to individuals and groups who are setting up and providing consumer developed initiatives (or CDIs). These can range from advocacy and self help groups and peer support through to fledging and for profit businesses. Through its three workers it offers advice on incorporation, governance and organisational development and provides opportunities to swap experiences with others and to access national and international resources, training and events. It has developed its own website and resources including help sheets and a newsletter and is moving into community development work.

To date, Our Consumer Place has been working with fledgling groups which are struggling to survive. It has also been surprised at the interest voiced by consumers in developing small businesses which generate income under the Disability Support Pension earnings cut-off point.

Overall Victoria has experienced a rapid blossoming of consumer activity and policy over the past decade. However, many now feel the lack of any subsequent investment has stalled growth and limited the development of opportunities.

**Consumer Academic**

Victoria has the first consumer academic appointed in 2000 at Melbourne University alongside a carer academic and funded by the Mental Health Branch. Although there may be other consumers contributing on a sessional basis in other universities, this remains the only permanent post in Australia. The post operates from a belief that the consumer perspective, as it applies to research and to the education of mental health practitioners, is a discrete and separate discipline with its own philosophy and critical theoretical approach. The post holder is involved in teaching students at the School of Nursing using a consumer perspective, inputting into the psychiatric nursing curricula, establishing a consumer academic program in mental health and a website and doing collaborative work both nationally and internationally. The post has also sustained the Psychiatric Action and Training Group. This is a group of consumer and senior nurse educators which acts as a mutual reference and support group and has been developing training workshops to enhance consumer skills in research and training. One success has been ensuring that research fellowships include a dedicated consumer researcher and that all applications require a demonstration of how consumers will be involved in the conduct and development of the research.

### 3.8 Western Australia

The history of consumer activism in WA has seen a number of peaks and troughs with initiatives driven by a pool of experienced and knowledgeable consumers with determination and commitment. Early on a decision was taken that the best way forward was to separate consumers and carers. Carers have since gone on to develop a successful professional organisation that provides training, a carer representative program, newsletters and systemic advocacy work. Developments in and for the consumer community have been more mixed.

One of the first initiatives was auspiced by the Health Consumers Council (HCC). The Council was established 15 years ago to provide a consumer perspective on health, to support participation through representatives and to establish an individual advocacy program. In 1996 a recommendation from a Taskforce on Mental Health (WA Health Department 1996) established a Mental Health Consumer Advocacy Training Program within the HCC for consumers and carers. This brought consumers (and carers) together, trained them, politicised them and developed a collective awareness of the need for reform and the potential role of consumers in influencing that. The Program had $140,000 per annum, three part-time workers...
and involved 120 active consumers. However, the Program was de-funded abruptly in 2003 and because there was a perception that the funding had been used to develop the Carers Association WA it created a divide between consumers and carers. Some informants consider that ‘WA has floundered ever since in realistically trying to involve consumers’. At the time of closure funding was released by the Office of Mental Health to add a further component to the Program – the Consumer Consultant Trial – reliant on the existing infrastructure. This was loosely modelled on the Victorian U & I Project (see page 44). It employed three part-time workers and aimed to enhance consumer participation by employing consumer consultants to input into developing effective strategies and practices in partnership with mental health services. It lasted for two years but was eventually de-funded in 2005 leaving an unfortunate legacy of divisiveness and mistrust.

In an attempt to clarify the situation the Office of Mental Health employed a consumer worker to prepare an integrated model of consumer participation for Western Australia.

A Model of Mental Health Consumer Participation for Western Australia

This model was developed in 2005 through wide consultation with consumers and carers and involved investigating models used elsewhere as well as the national policy documents and standards. It proposed a model consisting of a number of elements which would be implemented in three stages over the next five years. The elements were:

- consumer consultants, peer support workers, representatives and consumer participation coordinators in every service;
- a Consumer Education and Training Centre staffed by consumers;
- consumer academic positions linked with universities and TAFE;
- consumer researchers;
- consumer run services and independent consumer operated self-help and recovery services;
- a consumer peak body providing information as well as individual advocacy;
- a CAG in every service;
- a network for the coordination of consumer participation; and
- consumer-community liaison workers.

Stage one entailed the development of a Statewide Advocacy, Education, Training and Information Service (or SAETIS) staffed by consumers and managed by area mental health services. It also entailed the development of a peak body and network of consumer coordinators as well as a code of practice for employing consumers. Stage two would see trained and supported consultants and coordinators in every adult service, more commitment to the development of consumer operated services and the establishment of academic positions. Stage three involved developing research opportunities, the introduction of peer support workers in every service and the ongoing development of the rest of the model. The model was certainly ambitious and to date has not been implemented.

Thirteen CAGs are currently seen as the key consumer participation mechanism across WA. A number have grown out of local disorder specific support groups. They are usually linked to local clinics, run by a committee with monthly meetings and supported in kind by local mental health services who provide office space and some basic equipment. Although, in theory, there should be a CAG in every service this has not happened and consumers complain that they are not necessarily seen as a priority or valued by local services. At times services will recruit representatives from among their own clients rather than approaching the local CAG.

A consumer workforce has been slow to develop in WA and although there are three project workers called consumer consultants they are working on specific projects – for example to improve consumer outcome measures – rather than having a broader systemic advocacy role. A report undertaken by the University of Western Australia about the physical health and well being of people with mental illness (Lawrence et al. 2001) identified that they had the same mortality rates as Aboriginal
people. This led to funding for a pilot training program for peer support workers to encourage consumers to maintain their health. There is now a peer support program where eight peer workers are paid to work in the discharge lounge of one hospital linking consumers to GPs and encouraging them to maintain their physical health. Another hospital has now asked for a similar program and it is anticipated that this approach will expand state wide. There are also peer workers employed in CSOs, some of which have embraced consumer involvement and built it into their core principles.

Establishing a Senior Consumer Position

In 2006 a permanent senior consumer position was created within Mental Health Services to implement consumer participation across the state, coordinate the existing CAGs and assist in developing new ones. The post has established a quarterly forum for CAG representatives (CAGWA) to share information and input into training and hopes to nurture interest-based CAGs – for example for youth, CALD groups, older adults and Aboriginal people – to become pools of knowledge and expertise for consultation purposes. The post has also instigated quarterly meetings with area directors to input CAG feedback and provide a clear pathway into decision-making. It has established a database for expressions of interest from potential consumer representatives and there is work being done to expand the hourly rate for representatives to include a broader range of tasks, for example participation in evaluations and reviews. However, there is now a push to move away from a consumer representative approach and to bring issues to the CAGWA which would then set up working groups to work with committees and decision-making forums. The CAGWA is currently reliant on generic training available through the HCC but this is not specifically geared to mental health issues. The senior consumer position is also involved in post-graduate nursing and social work training and acts as a resource to mental health staff about how to do care planning and consumer participation.

Developing a Consumer Organisation

The Consumer Organisation Mental Health WA (COMHWA) has developed over the past few years using the VMIAC model (see page 43) and building on the foundations laid by an earlier organisation called The Association of Mental Health Consumers WA. It is incorporated, has deductible gift recipient status and has successfully overcome the divisions created by the demise of the AMHCWA. It aims to provide a vehicle for consumer advocacy and provide a one-stop-shop for training and education services and support for consumer representatives. It is governed by a management committee with occasional sub-committees and is supported by the Richmond Fellowship which provides an office, equipment and some administrative support. It has recently attracted a small grant to employ a project officer to prepare a proposal for government funding.

An election promise was made in September 2008 to create a funded CSO peak mental health consumer-run organisation. However, an anticipated cut to core mental health funding put the promise on hold. The Auditor General is currently reviewing funding in the community mental health sector and it is expected that this will indicate where money for COMHWA might come from. Consumers aim to launch the organisation in June 2009 and undertake a membership drive. There are concerns about expectations and that they will be expected to ‘run before they can walk’.

Consumers in Western Australia realise that they are some way from achieving the model for consumer participation put forward in 2005. Nevertheless they would like to see real money for consumer participation and champions to promote this, paid consumer consultant and peer support workers in every service and proper payment for doing representation work.
3.9 In Summary

Individual states and territories host a range of levels of consumer and/or carer organisational development and most states feel others are doing better. As one consumer said ‘we look to Victoria and say they’re well resourced and they look to New Zealand who look to the USA’.

Within jurisdictions there is a common experience of having diverse initiatives and pockets of good practice and different health regions having different approaches. Not all developed models are in urban areas and it can be the more rural communities where there is a general lack of services where consumer driven service delivery flourishes. Jurisdictions also share a debate about the increasing demand for involvement and representation from services so that they can fulfil their statutory obligations combined with a general inadequacy of resources and what this means in terms of sustainability. As one consumer said ‘it is not that tenable to work state wide with a 0.5 position’. Consumer initiatives have struggled to develop in isolation and to gain financial support which is often limited to project rather than core funding. Many also commented on the difficulties of implementing policy at ground level even when there was executive level commitment to change. They also commented on a small number of key individuals being instrumental in creating the impetus for change or for continuing with the status quo.

Two jurisdictions – Victoria and New South Wales – have developed mixed models where there is a well developed consumer consultant workforce, combined with a peak or state wide consumer organisation and consumer developed initiatives. Queensland will shortly join them if the new consumer and carer-run advocacy organisation manages to consolidate. Elsewhere CAGs at local, regional and state level have remained a key tool of consumer participation. Three states are sustaining state wide CAGs – New South Wales, the Northern Territory and Tasmania – and Western Australia is currently developing CAGWA as a state wide mechanism. NSW CAG has incorporated and is now operating as a peak body for consumers involved in information provision, consultation and policy formation. Elsewhere CAGs have remained as ministerial advisory groups with a more limited focus. There is a tendency to establish consumer positions as a ‘good thing’ in themselves without necessarily any rigorous analysis of the process, appropriate support structures and the impact of these positions. The consumer workforce is most developed in the Southern Health region of Victoria with the establishment of a consumer and carer consultant unit headed by an executive-level manager who is a consumer.

Energies put into developing a consumer workforce have varied widely across Australia, from non-existent to a strong push to integrate consumer workers into area mental health services. There has been a tendency to establish consumer positions as a ‘good thing’ in themselves without necessarily any rigorous analysis of the process, appropriate support structures and the impact of these positions. The consumer workforce is most developed in the Southern Health region of Victoria with the establishment of a consumer and carer consultant unit headed by an executive-level manager who is a consumer.

Although consumer consultants engaged in systemic advocacy are now reasonably well established, paid peer support work is less well developed and almost non-existent in many jurisdictions. The exceptions are Queensland with the rolling out of the consumer companion program across the state and in South Australia with a government initiative to train and employ peer workers. Western Australia is now piloting a peer support workforce in one hospital and an incorporated CAG in New South Wales offers peer support worker training. These initiatives are combined with volunteer peer support work and peer initiatives in CSOs. The absence of any accredited training for consumer workers remains a major gap.

There are pockets of good practice in developing the role of consumers as educators, trainers, evaluators and researchers. This includes the establishment of a consumer academic post and the promotion of consumer research activities hosted by the mental health consumer peak body in Victoria. There
have also been some interesting moves to create ongoing, formalised consultation and evaluation mechanisms and one example is the MH-CoPES project in New South Wales.

Beyond the more formalised consumer initiatives and participation mechanisms there are a vast array of consumer support and self help groups, organisations providing individual advocacy for mental health consumers, official visitors’ schemes and CSO initiatives. Some states — in particular Western and South Australia — also have health consumer peak bodies working to consolidate a broader health consumers’ lobby and engaging in capacity building activities to boost the capacity of consumers to participate and of providers to welcome that participation through support, training and education.

Overall most jurisdictions have witnessed a burgeoning of consumer initiatives on the ground accompanied by a push to develop state wide consumer-run peaks and a consumer workforce.
4. The International Picture

Like Australia, New Zealand, Canada and the UK have all seen a move towards mental health reform and recovery orientated services. They have also seen questions raised about how best to build the consumer movement and see the future in terms of fully integrating a consumer perspective into service planning and delivery and developing independent consumer-run advocacy, service and business organisations.

Although this research did not include direct contact with consumer leaders in the USA, the US does demonstrate a rich history of consumer activity. Today the National Empowerment Center\(^\text{12}\) estimates that there are over 7,460 mental health mutual support groups, self help organisations and consumer operated services across the USA. At a national level there is the National Mental Health Consumers Self Help Clearing House which provides information and practical assistance, including on site consultation, to consumer organisations. The National Coalition of Mental Health Consumer/Survivor Organisations was established in 2006 and consists of 31 state wide consumer organisations and four national consumer run technical assistance centres. It aims to provide, for the first time, a national voice for consumers in decisions made by the federal government and is the result of several attempts to form a single national organisation and bridge the divide between those who see themselves as consumers and those who see themselves as survivors. It is now seeking adequate funding, organising peer run networks and services and drafting model legislation.

4.1 International Initiatives

There are two key initiatives which shape the consumer movement at an international level. Firstly, there is the recent development of Interrelate, a new international coalition of national consumer organisations launched in Canada in 2008. The current membership is seven English-speaking countries including Australia, Canada, England, Ireland, New Zealand, Scotland and the USA. Interrelate aims to forge a strong international recovery movement and meets monthly by teleconference to share information, offer support across national borders and formalise collaboration. The mission and values statements are being finalised and it hopes to become a legal entity in one jurisdiction so that it can attract funds to sustain the work and offer support to consumers in other countries.

Secondly, there is the International Initiative for Mental Health Leadership (IMHL) established in 2003. It grew out of a realisation that mental health leadership is just as vital to the success of community based services as effective clinical practices but that currently there was an absence of resources and supports for key leaders in the mental health sector. IMHL offers support and technical assistance to countries and their leaders, including consumers and carers, in adapting to rapid changes in the field. It provides an international infrastructure to identify and exchange information about effective leadership, management and operational practices in the delivery of mental health services. Countries take turns in hosting leadership exchanges every 18 months which involve a conference and the matching of participants with leaders in the hosting countries. Other initiatives organised by IMHL have included an international trailblazer program conducting joint training between mental health and primary care organisations.

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12 The National Empowerment Center Inc. is an advocacy and peer support organisation run by consumers to promote an empowerment-based recovery model of mental disorder. It operates a free information and referral line, organises conferences, research and publications and is involved in many national policy debates. It keeps a database of consumer run organisations across the US, operates as a technical assistance centre and produces educational and self-help resources.
care practitioners, work in developing countries and an international social inclusion network which links leaders involved in developing more socially inclusive environments.

4.2 New Zealand

New Zealand is often put forward in Australia as a model of good practice for consumer development and as one consumer said ‘New Zealand is five miles ahead of us’. How far is this true? Certainly the emerging consumer movement profoundly shaped mental health service reform in New Zealand. Some of the key factors identified as influencing this have been a powerful Maori revival during the late 1980s where debates about treaties, reconciliation and self determination encouraged an acceptance of others which softened the way for the mental health consumer movement. There was also an early and radical program of deinstitutionalisation which opened the door to look at new ways of doing things. By the mid-1990s every district health board employed mental health consumer advisors to advise management. The year 1996 saw the establishment of the Mental Health Commission with the recommendation that one of the three commissioners should have a service user or family/carer background. This led to the appointment of a consumer and meant that from an early date there were vocal people in leadership roles in mental health structures.

In 1998 the Blueprint for Mental Health Services in New Zealand was published (Mental Health Commission 1998) and required all mental health services to use a recovery approach. This was accompanied by a national policy to support implementation which placed emphasis on social and personal responsibility and citizenship as well as breaking down stigma and discrimination. It also identified a need to expand the consumer workforce. A recovery approach was reinforced in 2001 with the identification of the recovery competencies required by mental health workers (Mental Health Commission, 2001) accompanied by a teaching resource kit to embed a recovery orientated system. The kit is used in mental health education, in job descriptions and by services for quality improvement. New Zealand has also supported an 11 year anti-discrimination campaign – Like Minds, Like Mine – aimed at reducing the stigma and discrimination faced by people with experience of mental illness. The program is funded by the Ministry of Health with a budget of approaching $NZ5 million per annum and involves a national public education program, conferences, research and work with regional providers of community education and anti-discrimination support, many of which are consumer run agencies.

New Zealand now has a flourishing consumer workforce with consumers employed at various levels and positions throughout the mental health system. Like Australia there has been debate about a lack of clarity about the role, employment conditions, remuneration and career pathways. There is also an ongoing debate about whether it is best to have a consumer workforce within the system which can limit the ability to voice opinions or to locate a workforce in independent organisations outside the system or whether a balance between the two is required and what the most productive balance should be.

In New Zealand I don't think we're doing that well with consumer initiatives and we probably had a few more in the late 80s than now even. One of the issues is getting that balance between people inside and outside the system. Some of these organisations are small and under-funded, they were run by people who were inexperienced and they folded. Also a lot of people got drawn into jobs inside the mental health system, like consumer advisory jobs and they are usually better paid and more secure. For a while it turned the other way and we seemed to have everyone inside the system and not many outside it. That is changing a bit. (Consumer leader, NZ)

There is a network of consumer-run services providing support and an alternative to traditional mental health services working within district health boards, within CSOs and independently. A recent review (Doughty & Samson 2005) identified 35 such services across New Zealand and there has been a move to develop regional structures and a national network to ensure a collaborative approach and the sharing of experiences as well as a peer support forum to showcase the work. However, there is also a lack of definition of what constitutes a peer-run initiative and how they relate to the rest of the mental health sector.
Mind and Body Consultants

This is a registered company established twelve years ago by a mental health consumer. It has grown substantially and now offers:

- a peer support service contracted by Mental Health Services and available to all service users in Central and West Auckland;
- a training arm registered as a separate company and providing New Zealand’s only comprehensive Level 4 training program for peer support workers. The course was recently accredited;
- a research and development team involved in service development and evaluation;
- a team of consumer advisers providing specialist advice to mental health services; and
- a local provider of Like Minds Like Mine, the national anti-discrimination campaign.

Altogether Mind and Body Consultants now employ almost 50 people and have a turnover of $2.5 million per annum. Most recently it was successful in contracting for an intensive support service for those living in supported accommodation and which meant doubling in size overnight. The service provides support using the peer support model and is split with a CSO provider offering support in more traditional ways. It has recently been contracted to establish a similar service in Christchurch.

The Director described difficult periods for the organisation and a steep learning curve particularly with recruiting suitable employees, clarifying roles and philosophies and designing appropriate training and supervision structures. However it now offers a unique model in NZ providing exemplar services founded on different assumptions to those which underpin traditional services.

A key issue for the future is how to get consumer run initiatives in the advocacy, support or educational fields treated as a core service. This has enormous implications for the kind of funding they should receive, the kind of training workers should get and a general professionalisation of initiatives. It also requires a clarity which is currently absent about what consumer-run initiatives are and what they can offer.

I would like to see peer run services be seen as a core service. In order to do that you need an awful lot of workforce development and service development. I would like to see a career path emerge through that. So I am for formalising it although I can see there are always risks with that. If you’re going to go down the track of saying we think this should be a core service you have to do those things like get a qualification, have a set of agreed standards. I imagine those first psychiatrists who started defining themselves in the asylums were not an organised, well defined group of people at the start. But there is a lack of definition about what is a peer run initiative, what are the range of things it can do. There is really no formal definition or consensus about the values it operates from, what are their purpose, how do they relate to the rest of the mental health sector? (Consumer leader, NZ)

There are particular challenges presented by the evolution of community-based services and the implications this has for consumer development in working within different locations, cultures and expectations. And there is always a need for more consumers to take on leadership roles within the system to encourage more people to ‘come out’. Finally there is now a focused attempt to develop an infrastructure at national or regional level to undertake information gathering and advocacy work which could supply an ongoing stream of information about consumer experiences to feed into decision making processes. But as one consumer said:

The consumer movement has been the networking, political activism stuff. The more recent developments are the consumer-providers and there are quite a
few organisations of various sizes providing services of varying sophistication around New Zealand and this is the direction we’re shifting in. With the networking function people are starting to wonder what the value of it is – a bunch of mad people getting together to do what. If the Ministry had its way it would probably get rid of all the networks and be happy to talk to people in consumer-provider organisations who are very experienced and knowledgeable and reliable and use them to get input. But that’s not politically palatable yet. What we will end up with is some kind of consumer lobby which will probably be predominately provided by consumer-providers. (Consumer leader, NZ)

COMPASS

This is a mental health peer advocacy and support service contracted by the Nelson Marlborough District Health Board to support mental health consumers through a mix of advocacy and information so that consumers are empowered to speak up for themselves and make informed choices. Its origins lie with a small group of consumers who during the 1990s decided to take action because they were not satisfied with the way services were provided locally. As well as working in the mental health unit, there are contracts with individual CSOS to provide peer support. COMPASS also operates a consumer-run community centre, the White House, and a 24 hours a day, seven days a week 0800 phone line answering questions from the community about mental health issues. It contracts Health Action, a community-based health promotion trust, to provide administrative and managerial support. Most recently it has established a crisis intervention residential facility run by consumers as an alternative to hospital. It provides training internally for peer workers and will support them to gain the national mental health certificate.

It described a slow process of gaining the acceptance of clinical services. At the beginning it was assumed that their services would not work and it was difficult to attract funding. Now clinical services openly state that these services are a great asset and actually make the work of clinicians easier. The crisis house is able to work with people the clinical services find very problematic because it offers a very different approach. COMPASS now identifies a population of consumers locally who have very little to do with traditional mainstream services and are totally supported through consumer-run services.

4.3 United Kingdom

The split in government between England, Wales, Scotland and Northern Ireland means that in the UK health and social care field there are regional differences in mental health legislation, user involvement mechanisms and the way in which mental health consumers have organised themselves. However, unlike most countries which have insurance based health services, the existence of the National Health Service (NHS) means that the private sector in mental health has remained very small and the majority of people in the UK access mental health services in the same way, through the public system.

At a policy level patient and public involvement in healthcare has become one of central tenets of the NHS modernisation agenda. This has been formalised in legislation which requires all NHS organisations to engage with users in service planning, evaluation and decision making in treatment. Patient choice is now a topic of heated debate and a ‘Patient Tsar’ has been appointed to advise on how the experience of users can be improved. However, there is no legislation to guide how consultation and user involvement should happen and, as elsewhere, there is still a long way to go in making service user involvement a reality. In the mental health field a survey of service users and providers of psychiatric services in London found that none of the Mental Health Trusts had systems for involving users that met national standards. (Tait, 2005). There has been some progress in involving consumers in research and the Department of Health has taken a lead in promoting this approach.

13 ‘User’ is the preferred UK term for consumers.
14 The word Tsar is used to refer to an appointed official tasked to regulate or oversee a specific area.
England has a network of consumer organisations and a systematic survey of their extent, scope and impact on reshaping mental health services (Wallcraft et al. 2003) identified over 500 organisations. It showed that most groups were small with an average membership of 50, formed in the last 15 years, poorly funded and with little representation from black and ethnic minority populations. They were providing a mix of mutual and social support, acting as consultants and representatives in decision-making forums, involved in training staff, in service evaluation and in user led research. They were also providing a mix of education and training for members, creative activities and individual advocacy. Some had become service providers and were offering services like drop-ins, befriending, help-lines, social enterprises and recovery support. Over half had paid staff and two-thirds sat on planning bodies. Their funding came from a mix of local and health authorities, NHS trusts and charitable bodies. In addition there were also consumer organisations established within larger CSOs. For example one mental health network run by consumers and providers was auspicing a user-controlled training organisation.

The survey made recommendations about how best to improve the ability of the consumer movement to influence mental health reform. It advocated for the development of national good practice guidelines for user involvement, refocusing mental health services towards recovery and increased support and funding for consumer groups to build the capacity of the movement. It also urged the formation of a national network to encourage good practice and build capacity. This has led to the foundation of the National Survivor User Network (England) set up in 2007. This links local user groups and provides a clearing house for information and assistance with an online database and newsletters. There is also the UK Advocacy Network which was set up in 1990 to provide a national voice for mental health service users, link users groups together and provide them with practical assistance. However, as one consumer leader said these national organisations have struggled to establish themselves:

We do need a national voice, it's very important. Our government have not supported national organisations and we've had to go ourselves to charity funding. It's difficult to do anything without a national voice because service users are so disparate. We have some strong organisations for manic depression, people with personality disorder but they can't come together very easily. I can't see what's going to make it happen unless our new national network can start to really function. All that work that was put into setting it up and money – we won't be able to start again because we won't have any credibility with funding organisations. We could set ourselves back a lot of years if the network doesn't work (Consumer leader, UK)

In Scotland the Scottish Executives program for improving mental health and wellbeing is driving a recovery agenda aiming to implement recovery orientated practice. The Scottish Recovery Network is now working to develop a recovery competencies framework for mental health workers (Dorrer 2008) by outlining the skills, values and knowledge required to drive recovery orientated practice. The national voice of service users in Scotland – VOX – has developed an extensive anti-discrimination educational program and made an agreement with the media not to portray people with mental illness in a negative light. This has been boosted by the recent ‘See Me’ anti-discrimination campaign run by a coalition of mental health CSOs and based on the New Zealand ‘Like Minds’ model.

However across the UK there are concerns about a loss of focus in the user movement despite considerable activity on the ground at grass roots level and a continuing battle to get appropriate remuneration for representation and consultation activities.

4.4 Canada

Canada is a federation of ten provinces and three territories. Each carries responsibility for health services and has its own Mental Health Act. This means that the mental health consumer movement in individual jurisdictions has different experiences, levels of funding and sophistication, rather like Australia.

At a national policy level there is no Mental Health Plan and no legislation promoting a recovery approach. In 1997 the Federation of Provincial and Territory Advisory Committees on Mental Health made recommendations about funding consumer groups and initiatives as a key element of good practice in
mental health. More recently Canada has established a well-funded and prestigious Mental Health Commission which has a stated enthusiasm for consumer-initiated projects and aims to become a national focal point for making progress on mental health reform. The Commission is funding a national study of consumer activity which is due to report at the end of 2009. It is also running an anti-stigma campaign based on models in Australia, New Zealand and Scotland and calling for a radical transformation of services to make them recovery orientated.

The largest province, Ontario, has a mature consumer/survivor sector. Twenty years ago the provincial government mandated that each of the psychiatric hospitals in Ontario should have a patient council to engage in systemic advocacy and operate as the voice of the patient. They were given funding to employ staff and there is now an Association of Patient Councils which operates across the Province. There are also over 50 peer organisations engaged in a variety of activities and mainly funded by health but also through employment and education budgets as well as charitable funding and business profits. Most commonly they undertake peer support and social and recreational activities but they also do systemic advocacy work and a number run businesses and social enterprises that provide employment for members. One consumer controlled organisation is now a community mental health service provider. Another runs a successful courier business using 65 consumers to courier documents across Toronto using the public transport system. This organisation also runs another three businesses and, interestingly, is becoming a player in the more generic field of anti-poverty advocacy and is impacting on policy.

Ontario has also made progress in taking consumer knowledge into the university with a number of academics who have been open about their mental health backgrounds and who are developing courses about ‘mad people’s history’. There is also consumer influenced development of employment and education. One community college now offers training in the construction industry or ‘augmented education’ for those with a mental health history. The training is stretched over a longer time period than the standard course and is accompanied by job placement and support and the general building of confidence levels.

The National Network for Mental Health is the only non-diagnosis specific consumer driven national mental health organisation. It is based in Ontario, survives on project funding and works to eradicate stigma and promote income security, employment, mental health reform and quality of life. It has developed a supportive employment program, BUILT, and is working on a curriculum for training consumers to become peer specialists.

However generally there are concerns about a lack of government investment in this area. A recent review of consumer/survivor initiatives in Ontario (Casey 2008) found that most had not received any new investment and that some had lost funding or organisational autonomy. An international consultancy team has been working with consumer/survivor initiatives and other stakeholders to gather information about how to strengthen and support these initiatives in Ontario and to clarify their role in the health care system.

4.5 In Summary

This section has only been able to provide a brief outline of major developments overseas. It demonstrates the commonality of issues, experiences and struggles across national borders, the difficulties mental health consumers in all countries have had in developing national structures and attracting adequate funding and the push to develop consumer-run alternatives to traditional mental health services and consumer businesses. There is also a shared experience of difficulties in defining and establishing the role of a consumer workforce.

Interestingly Australia is the only country which has pushed structures for consumers and carers to work together. Elsewhere there has either been a slower development of carer issues and mechanisms or a split into a consumer leadership and a carer or family leadership with few organisations or networks to bring the two perspectives together.
5. Tasmania

This chapter collates information from Tasmanian consumers and other stakeholders, policy documents and consultation processes to present a picture of the consumer movement and consumer participation and involvement activities in Tasmania. It outlines the policy environment, summarises the history of the consumer movement and participation activities and outlines some options for the future development of consumer activity in the state.

5.1 The Policy Environment

The Bridging the Gap Report (DHHS 2004) reviewed the state of mental health service delivery in Tasmania and made 33 recommendations to improve it. This resulted in a government commitment of $47 million spread over four years to improve mental health services, including defining a recovery framework and progressing consumer and family/carer involvement. The latter entailed making funding available to establish four consumer and family/carer advisory positions – a senior consumer and carer consultant and three regional posts.

In 2006 a Consumer and Carer Participation Framework was formulated (DHHS 2006) through consultation and guided by a Project Reference Group of consumers, carers and mental health staff. It aimed to set the direction for participation in mental health services and bring together a number of processes and mechanisms into one coordinated system of participation, provide clarity about what participation means and offer a roadmap as to how it should occur.

The 2006-2011 Mental Health Strategic Plan (DHHS 2005) aims to create ‘a Tasmanian model of care for mental health that is centred on consumers, promotes their recovery and is provided equitably and efficiently through a strategy which includes involving consumers, carers, partners and others in the development of the model’. This makes participation and partnership one of six strategic priorities for mental health services in the next five years and crucial to the development of better services. The Plan aims to ‘include consultation and participation strategies in all major service activities and projects’ by progressively implementing the Consumer and Carer Participation Framework and to ensure that all consumers have individual plans developed collaboratively with them. A key outcome measure is a significant increase in participation levels by consumers and carers.

An evaluation of the implementation of the Bridging the Gap recommendations (KPMG 2008) identified that there had been difficulties in filling the consumer and carer positions and they remained vacant. This was partially overcome in 2008 with the appointment of a Senior Liaison Officer, Consumer and Carer Participation. The officer has led a Consumer and Carer Participation Review to identify an optimum model for participation, inform the implementation of the Framework and turn it into a practical and effective working system. The Review has been overseen by a steering committee of representatives from Mental Health Services and an advisory committee of consumer and carer groups, individuals and community sector organisations. It has entailed public consultations, regional focus groups with consumers, a widely disseminated questionnaire and individual submissions. The Review process has acknowledged that implementation of the Framework will involve establishing systems, processes and a supporting structure and is likely to take some years.

The Review is running parallel to DHHS-wide work to develop a consumer engagement strategy in order to involve communities and consumers in health and human services planning, policy
development, service design and decisions about their own care (DHHS 2009). The DHHS wishes to build on existing processes and draw from the growing base of research and experiences elsewhere. It will make decisions about the focus of engagement, type and techniques and it is anticipated that the strategy will provide an overarching framework for DHHS policy and operational units and lead to legislative change. It places a responsibility on units to improve the way in which they engage with consumers and communities and to make decisions about what kind of engagement suits them best. A consultation process is currently underway.

Also of relevance is the current review of Tasmanian DHHS-funded peak bodies (DHHS 2008). The review has explored the number and type of peak bodies and is developing a peak body strategic framework. The review notes the absence of a ‘body which has the capacity to represent the voice of [health and human services] consumers in a strategic, considered and well-researched way’ as well as current debates specifically in relation to mental health consumers. It recommends establishing an appropriate body to undertake peak functions on behalf of consumers of health and human services.

Overall the involvement of consumers and carers in the delivery, design and planning of mental health services is now a key priority for the Department. Yet currently, and certainly in comparison to many other jurisdictions, Tasmania has low levels of consumer activity.

5.2 Mental Health Consumer Activity

Unlike most other states and territories Tasmania has no fully operational consumer run organisations and a very small paid consumer workforce. Among the few paid consumer positions are a two day-a-week post to undertake the ground work involved in setting up a consumer organisation and a full time peer support worker in Anglicare funded through the Personal Helpers and Mentors Program (PHaMs)15. The State does have a participation framework and a state wide ministerial advisory group of consumer and carers, TASCAG. It also has a number of active and influential consumers who have pushed for better participation mechanisms and a stronger voice for consumers.

They have been involved as consumer representatives on state and national bodies and policy forums, working with professionals and establishing consumer-led initiatives including an embryonic consumer advocacy organisation, the Tasmanian Mental Health Consumer Network (TMHCN).

Tasmania also has a number of consumer support and self-help groups which bring people together to provide mutual support and social and recovery orientated activities. Although this research did not attempt a comprehensive mapping of these groups it did seek to throw some light on their numbers, size and range of activities. Some are convened and run under the auspices of larger CSOs, others remain independent of mental health services. Some are very short term; for example Family Based Care Association North West oversees a group engaged in craft activities for those with depression, a men’s group, and a support group for those discharged from the local in-patient ward. These groups have a life span of a few months at most. Others are longer term and either run on an entirely voluntary basis or have acquired varying levels of funding from the State Government to cover rental and running costs. One of the earliest was Breakthrough, established in March 1986 as a self-help group for psychiatric patients providing diversion, support, education and therapy for up to 80 members. Breakthrough was subsequently subsumed into Eureka Clubhouse.

Table 1 lists the more established and longer term consumer support groups. In addition there are two eating disorder groups and a post natal depression group operating in Hobart. There may also be additional disorder specific and youth groups operating in the state and consumer groups in residential and social support programs.

Table 1

<table>
<thead>
<tr>
<th>Consumer Group</th>
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</thead>
<tbody>
<tr>
<td>Name</td>
</tr>
<tr>
<td>Breakthrough</td>
</tr>
<tr>
<td>Eureka Clubhouse</td>
</tr>
<tr>
<td>Family Based Care Association North West</td>
</tr>
<tr>
<td>Eating Disorder Group</td>
</tr>
<tr>
<td>Post Natale Depression Group</td>
</tr>
</tbody>
</table>

These consumer groups are not advocacy organisations. Although they might undertake individual advocacy activities on behalf of their members and get involved in consumer participation and consultation processes, their main function is to support their members rather than to undertake more systemic advocacy work.

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15 PHaMs programs, delivered by CSOs, provide one-on-one support to assist people with mental health problems who have severe limitations to manage their daily activities and access services. It is one of the key initiatives of the Australian Government’s mental health reform package to promote community support and social connection.
### Table 1: Consumer Support Groups in Tasmania

<table>
<thead>
<tr>
<th>Name of group or auspicing organisation</th>
<th>Activities</th>
</tr>
</thead>
<tbody>
<tr>
<td>GROW</td>
<td>This is a national organisation which runs 10 weekly recovery-focused groups across Tasmania using a 12-step approach. Groups are facilitated by two fieldworkers in the north and south and average attendance is eight people.</td>
</tr>
<tr>
<td>Anglicare</td>
<td>Club Haven is a psychosocial centre run with direct input from consumers who make decisions about activities. There are currently approximately 40 members.</td>
</tr>
<tr>
<td>ROPES – Reaching Out to People in Emotional Stress</td>
<td>This has a core membership of approximately 25 people and has been meeting twice a week for over 20 years to provide social contact and activities in the Glenorchy area. It is funded by Mental Health Services. Membership is mostly female, long-term and in the 50+ age bracket.</td>
</tr>
</tbody>
</table>
| ARAFMI – Association of Relatives and Friends of the Mentally Ill | ARAFMI runs three support groups with $2,000 from the Department of Premier and Cabinet and smaller sums from charitable organisations. They are:  
• HOT – Helping Ourselves Together – meeting monthly in Hobart and offering information, shared experiences and mutual support;  
• a bi-polar support group in Launceston; and  
• a wellbeing group meeting fortnightly in the North. |
| Richmond Fellowship | Richmond Fellowship have set up a number of recreational groups most of which have been meeting for over three years. They are:  
• a fishing group with a membership of seven people;  
• a women’s group of approximately 13 people targeting the geographically or socially isolated;  
• a mother and baby group used as a stepping stone into more mainstream mother and baby activities;  
• a group for in-patients; and  
• a mountain biking group with a membership of eight. |
| Eureka Clubhouse | This is based on the clubhouse model and was established in Tasmania 14 years ago. It has an attendance of up to 30 per day, is currently auspiced by Colony 47 and funded by DHHS and has a staffing of 3.2 FTEs. It runs Total Quality Maintenance – a gardening and cleaning business employing up to 20 people with mental health problems. It also has a contract with DHHS to provide maintenance services to Housing Tasmania properties. |

Overall, despite small scale initiatives and a growth in consumers coming together to support each other, the history of the consumer movement in Tasmania has been one of struggle in order to get government and service providers to recognise the benefits. Explanations for this vary but it is commonly believed among both consumers and many service providers that an entrenched ‘Royal Derwent’ culture is partly to blame. This culture is characterised by a strong attachment to the medical model and pharmaceutical approaches and an ageing workforce. Being an island state there has been little input from the experiences of mainland mental health services and consumers. This isolation has promoted an ingrained culture and difficulties in grasping the complexities of consumer participation. As one informant said:

\[\text{The Royal Derwent was the economic, employment and social centre. Young people would leave school and become ward cleaners and then do the Royal Derwent training for psychiatric nursing, like their parents, and with no general nursing training. Many lived in the grounds with patients mowing their lawns and coming to barbeques. There was no flow through and as Tasmania also underpays very few staff come from the mainland. The Royal Derwent culture permeates today on the wards and among staff. (Stakeholder, Tasmania)}\]

This has led to a lack of priority for and under-resourcing of consumer activities. Indeed the National Mental Health Report (DoHA 2007) identified that Tasmania in 2005 was the worst performing jurisdiction in Australia for both consumer and carer participation in public mental health services.

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16 The Royal Derwent Hospital was the main institution for psychiatric patients in Tasmania. It was closed in 2001.
Table 2: Indicators of Consumer and Carer Participation in Public Sector Mental Health Service Organisations in States and Territories in 2004-05

<table>
<thead>
<tr>
<th>Indicator</th>
<th>ACT</th>
<th>NT</th>
<th>NSW</th>
<th>Qld</th>
<th>SA</th>
<th>Tas</th>
<th>Vic</th>
<th>WA</th>
<th>Aust</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percentage of services with level one participation*</td>
<td>100</td>
<td>100</td>
<td>54</td>
<td>54</td>
<td>46</td>
<td>0</td>
<td>53</td>
<td>33</td>
<td>51</td>
</tr>
<tr>
<td>Consumer consultants per 1000 clinical FTEs</td>
<td>4.4</td>
<td>0</td>
<td>4.1</td>
<td>2.4</td>
<td>2.4</td>
<td>0</td>
<td>3.8</td>
<td>0</td>
<td>2.9</td>
</tr>
<tr>
<td>Carer consultants per 1000 clinical FTEs</td>
<td>0</td>
<td>0</td>
<td>0.7</td>
<td>0.2</td>
<td>0</td>
<td>0</td>
<td>1.8</td>
<td>0</td>
<td>0.7</td>
</tr>
</tbody>
</table>


* Level one participation refers to agencies where consumers and carers are given a formal place in local decision making structures OR where a specific mental health consumer/carer advisory group is established to advise on all aspects of service delivery.

Table 2 describes the situation at a particular point in time after the demise of the consumer consultancy project and before the establishment of the TMHCN. However, it does demonstrate the low base from which Tasmania has been working.

There has recently been the development of a state wide, incorporated carer and family organisation, ARAFMI Tasmania (Association for Relatives and Friends of People with a Mental Illness) provides support for families and those with mental illness through workshops, a website, a 24-hour helpline, inputting into policy debates and overseeing the development of some consumer support groups. It is run by a Board which sets the strategic direction and has recently appointed an executive officer. There are also two part-time workers in the north and south. It has a membership of approaching 200 people and sends newsletters to over 1,000 families. A recent MHCA capacity development grant has allowed it to employ a consultant to review its current service delivery and the gaps, improve its community profile and survey members’ views about possible future directions. ARAFMI now wishes to consolidate its position, enhance its role as a specialist information and support service, build effective partnerships and ensure its future financial viability.

5.3 Consumer Participation Mechanisms

Current participation mechanisms in Tasmania include individual petitions, ad hoc consultation processes, TASCAG and initiatives in individual CSOs. In the private sector the Hobart Clinic has a consumer and carer advisory group and a consumer representative sitting on the hospital Board of Management. This is unique in the private sector in Australia.

There are two further mechanisms. The Official Visitors Scheme, established in 2000, has the power to monitor and investigate standards of care and treatment issues in acute facilities under the Mental Health Act 1996 and report on these to the Mental Health Tribunal. There is also Advocacy Tasmania which has one mental health advocate operating state wide to provide individual advocacy and with a secondary consultation role supporting the advocacy work of other services (Advocacy Tasmania 2008).

At a broader level the Health Complaints Commissioner investigates complaints related to the provision of health services in the public and private sectors. However, there is no capacity to specialise in mental health issues and unlike other jurisdictions there is no general health consumer organisation in Tasmania. Elsewhere this kind of organisation has provided a focus for engagement work as well as access to training and resources for health consumers.

5.3.1 TASCAG - Independent Ministerial Advisory Committee on Mental Health

TASCAG has been the main mechanism for consumer and carer participation in Tasmania. It was established in 1993 through the first National Mental Health Strategy as a ministerial advisory group to provide an independent mechanism for consumer and carer input into mental health policy and decision making at a state level. It consists of 13 delegates from around
the state who are appointed by the Minister for Health on an annual basis with a tenure of two years. Members must be consumers, carers or interested community members. It meets a minimum of four times a year in various locations and members are paid sitting fees and reimbursed for their costs. It aims to provide advice to the Minister for Health, assist mental health services in the development of policy, plans and initiatives, promote consumer and carer participation at all levels and sustain a pool of consumer and carer representatives.

TASCAG was not established as a representative structure and has no formal links with services or the broader consumer community. This has led to a lack of clarity about its role which was described by some research participants as ‘shadowy’. Over the years it has managed difficulties caused by a shortage of skilled representatives and some tensions in trying to bridge what can be conflicting agendas between carers and consumers. There have also been difficulties with remuneration for representative roles. Departmental guidelines (DHHS 1999) covering the process for appointing representatives and remuneration have set rates which now fall well below those established nationally. This has led to some pressure to increase them.

5.3.2 Consumer consultancy project

This was one of the few consumer-led initiatives in Tasmania. It was established in the south in 1996 with $30,000 funding through the National Mental Health Plan and auspiced by the Tasmanian Association for Mental Health. It employed a number of part-time consultants (three hours a week) to work in pairs on in-patient wards in the Royal Derwent and Royal Hobart hospitals and provide consumer consultancy. Consultants were supported and trained by a 16 hour per week coordinator post. In 2000 the project designed an information resource for in-patients about hospital and community services which was positively evaluated by staff and patients. There was also some discussion about extending the project to provide support to patients when they were discharged into the community.

However, like consumer workers in other jurisdictions there was some confusion about the role and concerns that it was providing befriending and a ‘listening ear’ rather than a more systemic consumer consultancy role. It was also limited in its geographical coverage because of transport issues. In 2002 it was defunded pending a review by the DHHS to assess the potential for a state wide consumer consultant program which would address concerns about access and equity across Tasmania. The review did not occur. However, consumers involved in the project were able to initiate Champs Camps, based on a Victorian model, which provides support for children with parents who are affected by mental health issues. The camps continue to operate in Tasmania and are now organised by CSOs.

5.3.3 The Mental Health Council of Tasmania (MHCT)

MHCT is the peak body representing the interests of the community mental health sector. It provides a public voice for people affected by mental illness and the community organisations which work with them. It aims to influence mental health policy and maximise the participation of consumers, carers and the community. The Board consists of representatives from each of the key stakeholders — consumers, carers and providers. However, it has struggled to fill consumer and carer positions especially in the north of the state and this has led some to concerns about the dilution of the consumer voice. The MHCT has in the past auspiced a number of organisations including the Official Visitors Scheme, ARAFMI, TASCAG and the TMHCN.

5.3.4 Initiatives in Community Service Organisations (CSOs)

The last few years have seen both an expansion in CSOs providing community mental health services and in initiatives to promote and encourage consumer involvement in those services. Some of the key CSOs in this field are Aspire, Anglicare and the Richmond Fellowship.

Aspire runs a three-year rehabilitation and recovery program across the state which works with over 400 people at any one time. Each of the six geographically based services has an advisory or ‘In-the-Loop’ group of consumers to provide feedback and advice on services. Average attendance is approximately six consumers and each group also produces a newsletter. In-the-Loop groups come together annually to share experiences and a consumer sits on Aspire’s Committee of Management in Victoria. Aspire also recently
funded a visit from a consumer leader in New Zealand to train providers about the experience of hearing voices.

Anglicare has actively promoted employment pathways for consumers and, as well as employing a number in professional positions in mental health programs, has one full time peer support worker in its Personal Helpers and Mentors program in the South. It also employed a consumer for a 12 month project officer role to assist in the redevelopment of one of its social support programs — Club Haven, a psychosocial centre. A second social support program has an informal consumer support group and all members are encouraged to participate in six-weekly planning meetings. However there are no designated mental health consumer positions on decision making bodies within Anglicare.

The Richmond Fellowship provides residential rehabilitation and independent living skills and recreational programs state wide. At a program level is has promoted the involvement of consumers through full engagement in individual treatment planning and reviews and through weekly community meetings where consumers make decisions about activities, program issues and policies and procedures. Consumers are also involved in staff recruitment and interviewing. Richmond Fellowship is now proactively exploring how to develop a more systematic approach through researching initiatives elsewhere and through setting up a consumer reference group which can operate state wide.

5.3.5 Independent consumer representatives

There are a small number of consumers working independently as consumer representatives and some have acquired high levels of expertise and skill in particular areas, for example mental health legislation. This has given them credibility with service providers and reinforced a process whereby the increasing demand for consumer representation has meant particular individuals being ‘tapped on the shoulder’ to participate rather than following any agreed process for nomination to committees and decision making forums, for example through TASCAG.

Research informants described a lack of skilled consumer representatives in Tasmania and a need for regular support meetings and training for those involved in consumer representation across the state.

5.4 Tasmanian Mental Health Consumer Network (TMHCN)

The TMHCN is the only example of a consumer-run advocacy organisation in Tasmania. It grew out of the voluntary efforts of consumers during the 1990s to bring consumers together, produce a newsletter and progress funding for a consumer network. It was officially formed by a group of consumers in 1999 to provide a mechanism for Tasmanians to provide input to the national body, the Australian Mental Health Consumer Network. Despite several attempts the founders were unable to attract any significant funding but a group of active consumers continued to build the foundations for a network by holding social events, attending forums, providing information and building relationships with consumers and other organisations. It was involved in the consultation process associated with the development of the Consumer and Carer Participation Framework and undertook the analysis of questionnaires completed during the consultation.

In 2005 and with a formal membership of approximately seven people, the network was granted $30,000 per annum for three years by Mental Health Services to fund a coordinator post for two days a week. The size of the grant was determined by the wish to ensure funding equality with Tasmania’s two mental health carer organisations which existed at the time. Although auspicing arrangements had been explored with Volunteering Tasmania, TasCOSS and ACROD, an auspicing agreement was agreed with Mental Health Council of Tasmania for 12 months with the expectation that the Network might move towards incorporation at the end of that period. A member of the core group was appointed to the coordinator position and a job description subsequently compiled using NSWCAG as a model. There followed a number of discussions to clarify the nature of the relationship between TMHCN and MHCT and eventually a Letter of Intent was issued by the MHCT Board outlining the nature of the auspicing relationship and financial accountability. A further grant of $4,000 from Bridging the Gap monies was used to pay for office furniture and a computer.

17 See note 15 on page 58

18 ACROD is now known as National Disability Services — the national industry association for disability services with an office in every state and territory.
The year 2006 saw the formal launch of the TMHCN, the official opening of the TMHCN office, the production of a newsletter and a program of visits to mental health organisations around the state to build partnerships. Success in acquiring funding and being able to undertake more activities meant that the Network began to attract more interest and more members. By 2007 it had a membership of 44 people with approximately 10 of those actively involved in the committee. During the year considerable energies were devoted to formalising governance structures, policies and procedures and a strategic and operational plan. The Plan outlined strategies for systemic advocacy activities, supporting representatives on committees, awareness raising within the community, developing a media profile, and establishing effective governance and communication through a newsletter and website. A leaflet to advertise the TMHCN was designed and an advocacy forum organised which gained good media coverage. There was increasing support for the organisation including a TasCOSS recommendation that there should be adequate funding for a representative body for Tasmanian mental health consumers and to strengthen the capacity of existing health consumer advocacy groups in the state as a Budget priority.

However, despite the achievements of 2007 there were difficulties. There were concerns about a deteriorating relationship with the auspicing organisation and difficulty in establishing regular communications with Mental Health Services senior management. This created anxieties among active TMHCN members that Mental Health Services was unaware of the Network’s successes and achievements. In addition, and despite a small but functioning committee, the increasing workload was falling on fewer shoulders.

In 2008 the Network successfully applied for $40,000 from the MHCA in NGO Capacity Development Grants to undertake projects to develop a business plan, a mental health profile plan and a consumer empowerment, engagement and professional development plan, all of which would contribute towards developing a consumer run incorporated organisation. The Network subsequently facilitated and housed the Consumer Organisation Project to undertake this planning process using external consultants and a small coordination team to develop the paperwork required for incorporation.

The DHHS review of mental health consumer and carer participation was accompanied by funding for both TMHCN and TASCAG until July 2009 with continued funding subject to the outcome of the review.

5.5 The Mental Health Consumer Organisation Project

This Project, initiated and managed by the TMHCN, aimed to develop a business plan for a state wide consumer organisation and make a case for funding. It brought together both consumers involved in the Network and other consumers in a number of forums during October and November 2008 to discuss what a strong, effective and well-funded consumer organisation in Tasmania would look like.

There was a clear message from participants that any organisation should be consumer driven and run with the purpose of providing a strong and informed consumer voice with the motto ‘nothing about us without us’. Ideally it should provide consumer leadership, strong relationships, communication and advocacy across the state as well as opportunities to share experiences through social contact and activities and pursue the development of consumer run services. It would not provide individual advocacy but would assist with the development of consumer groups, undertake media work, campaigning and lobbying and act as an information and resource centre. This would include establishing a website. Other possible functions for future discussion were identified as facilitating peer support and mutual support mechanisms, education work in clinical services and schools and creative projects. Tackling stigma and encouraging employment for consumers were seen as longer term goals. Overall the vision was about feeding back the experiences of consumers into the system and promoting the importance of the lived experience or as one informant said ‘consumers as the canaries of society’. Discussions centred around the need to match the role of any consumer initiative with gaps in services in order to avoid duplication with other organisations.

The Project also focused on governance issues and the need to separate strategic control and direction functions from the content and day-to-day operations of the organisation. This would be best achieved through establishing a Board of Trustees consisting of senior, experienced people who have undertaken leadership roles and who are selected through...
an independent nominations committee. Rather than having regional representation, regional issues would be dealt with through a regional issues sub-committee. It was anticipated that a more professional structure would immediately attract a broader membership. Membership would be open to those who self identified as mental health consumers and who were resident in Tasmania with an annual fee of $5. An associate membership would apply to non-residents and to non-consumer supporters. It was suggested that the aim should be a core membership of approximately 50 people.

The Project also discussed process issues including the necessity to ‘walk before running’ by undertaking clearly defined, achievable tasks to boost confidence and prove capability and credibility to both consumers and non-consumers. Potential funding sources, minimum budgets and how to deal with a transitional period before incorporation were also discussed. Project participants were invited to join a small coordination team to drive the project forward and to produce a newsletter to keep interested stakeholders informed about progress.

5.6 Lessons from the TMHCN

Over its three year history the TMHCN can claim a number of achievements. These include the establishment of an office and building up a resource library, providing consumer representatives for working groups, projects and committees and preparing public policy submissions; there have been seven major submissions over a two year period. It has engaged in a number of successful promotional activities including organising a launch and advocacy forum, producing two newsletters, coordinating email distribution lists, holding a photographic display and achieving face-to-face contact with mental health organisations across the state. The organisation of the advocacy forum in particular has been described as a ‘breakthrough’ for such a small group. It also organised social events, networking and information exchange opportunities. It has devoted energies to organisational development by drafting policies and procedures, formalising governance structures and experimenting with developing mechanisms for a broader input from the membership. It has established sub-committees for focused work on public speaking skills, social and promotional activities, auspicing and incorporation. It has tried to develop a volunteer program to underpin operations and accessed training and skill development opportunities for staff, volunteers and the membership. Overall it has laid the foundations for building an incorporated consumer run organisation in Tasmania.

Nevertheless it has also encountered difficulties along the way and the lessons which can be learnt from these difficulties are an important part of its legacy. These lessons are about:

- **auspicing.** When the TMHCN was first funded the consumers involved expressed a wish to be auspiced outside the mental health sector. This followed concerns expressed by the consumer movement both nationally and internationally that auspicing within the sector means that consumers can never escape the role as patient or client with their work being seen in terms of psycho-social rehabilitation. This can have a negative impact on both credibility and self esteem. When the Network was first established a number of organisations were approached as possible aupisers including Volunteering Tasmania. This was favoured because it would have meant devoting energies to establishing a volunteer program engaging a broader section of the membership and building up their skills and capacity to run the organisation. However, auspicing by the MHCT became part of the funding conditions.

- **membership capacity and sustainability.** Members of the Network identified as mental health consumers and ranged from those who expressed an interest in the work through to those who wanted to play a proactive part in defining the direction of the organisation. The latter remained a small percentage of the total membership – between seven and 10 individuals. As a number of informants pointed out identification as a consumer can be stigmatising and this is perceived to have impacted on the ability of the Network to build up a strong membership base and to recruit members particularly from the north and north-west of the state to ensure effective organisational functioning. There is also a perception that many consumers have a limited understanding of the value of advocacy and of the links between advocacy and the recovery process. All these factors contributed to a situation where the capacity to undertake a growing body of work was limited.
support levels. Developing a state wide organisation to represent the views of consumers through a two day-a-week coordinator post is challenging. The expectations and the workload required high levels of support which could have been met through a mix of regular supervision, paid mentoring and relevant training. Not all of these mechanisms were in place.

clarity about role and functions. The TMHCN was established with no formal guidelines about how it should operate alongside other participation mechanisms, for example TASCAG and the MHCT. Although some clarity around these issues developed over the course of the Network’s history and through its own attempts to develop an operational framework confusions remained which hampered its development and ability to build a profile within the sector. This caused tensions in the mental health consumer community particularly around the balance between providing a mechanism for systemic advocacy and providing a network to promote communication and support across a community.

5.7 Options for the Future

The research shows that mental health consumers and many other stakeholders in Tasmania want change. They want to see a mental health system based on a partnership between consumers and providers in a respectful and dignified environment and which offers strategies for recovery. As one consumer said services ‘need to ooze hope’. An essential aspect of such a system is having a range of opportunities to both hear and build on the consumer experience. How should this be achieved?

The factors that should be taken into account in thinking about developments in Tasmania are:

a small dispersed population in an island state. Although Tasmania is not such a geographical challenge as many parts of Australia the population is heterogeneous, decentralised and isolated from many mainland initiatives and good practice. This has contributed towards problems in sharing positive developments in service knowledge and practice among many mental health staff, problems in sharing consumer experiences across the state, the development of regional cultures among a fragmented consumer community and difficulties in creating equitable state wide access to any consumer initiatives.

the history of consumer development and participation initiatives in the state. Previous attempts to organise consumer-run initiatives, like the TMHCN, have provided a vehicle for skilling up mental health consumers, testing the waters and learning what works and what does not work. However, they also demonstrate the difficulties stemming from a combination of under-resourcing, high expectations, lack of clarity about roles and auspicing within the mental health sector. This has created pessimism in some quarters about the capacity and sustainability of consumer organisations generally.

lack of resources. A small community and a shortage of resources means there is pressure to combine a number of functions into one organisational structure. This can result in good outcomes like efficiencies and cost effectiveness but can also mean inbuilt structural conflicts, for example pushing consumers and carers together into one representative structure or participation mechanism.

development of a carers’ organisation. ARAFMI is the only organisation in Tasmania providing peer support to carers, relatives and friends as well as supporting a number of consumer support groups. It has recently become a state wide provider and is exploring its longer term sustainability. Currently it offers a range of ways for people to get involved – through direct services, in community education, through advocacy and in running the organisation. Interestingly the most popular way to participate is through helping others and this has meant that there have been no problems in recruiting for the Helpline and for peer led support groups. This provides a useful model for any consumer-run organisation.

There is a temptation to seek an instructional ‘how to do it manual’ (Kroschel 2002) or to emulate others where there has been evidence of success. However, every jurisdiction operates differently. Tasmania has particular and unique characteristics but it also shares many of the same issues and tensions which have been experienced elsewhere in developing consumer activities and initiatives. This means that it is neither a question
of adopting one particular model or of reinventing the wheel but rather using the lessons learnt elsewhere to develop a Tasmanian approach which can do three things: establish a sustainable consumer-run organisation, build a consumer workforce and create an appropriate supporting infrastructure.

1. Establishing a consumer-run organisation
Establishing community organisations generally, can be difficult. Informants involved in consumer-run organisations both interstate and overseas were asked what their key messages would be for others. They were:

- **keep it simple** and ask what an organisation can do for people rather than what people can do for an organisation. Experiences overseas demonstrate that the future may be about service provision rather than, as one consumer put it, ‘having talk-fests about advocacy’;
- **employ skilled people** able to do the job and provide them with adequate supervision and support. Several organisations identified one of the biggest mistakes as employing people because it would be good for them not because they would be good for the job;
- **if possible having one person who is full time** is an enormous asset. Having more than one employee can counter difficulties with isolation;
- as one consumer said **building relationships** with all stakeholders is ‘the fundamental responsibility of consumers in this work’. The research suggests that it is also important not to be afraid to get support from non-consumers with particular skills and expertise. For example, some organisations had used reference groups of non-consumers to support their establishment;
- **consumers and carers** can work together once consumers feel strong enough to maintain their position. Otherwise the involvement of carers can operate as an extra layer of complexity which impedes development;
- **introduce mechanisms** to provide a safe and healthy working environment which is free from discrimination, harassment and bullying. Working relationships and standards of behaviour between employees, and between employees and Boards of Management and the policies and procedures which underpin these must reflect best practice in community service organisations and contribute to a culture of cooperation, learning, training and support;
- **set priorities and think in ‘project-sized pieces’**. It can be difficult to meet the high expectations of other stakeholders, including consumers, and tempting to try to do everything at once;
- **clearly divide** administrative and accountability functions from the policy and strategic direction role;
- **auspicing** can be a useful development tool as long as mutual requirements are clear. However auspicing within the mental health sector can mean an inability to escape the role of ‘patient’; and
- **adequate resourcing is essential** to develop a professional and credible service but a mental health consumer organisation has to be seeded and funded in a different way to other community organisations. Currently most consumer organisations have to find their own way and although funders may provide the financial resources they do little to assist with developing capacity. Organisations require security of funding (for example three year grants) and clear service agreements.

2. Building a consumer workforce
This is seen as a mechanism for transforming mental health services so that they fully adopt a recovery orientation. Many people also see it as an entirely new service seeking professional recognition. However, a number of jurisdictions have experienced difficulties with establishing consumer positions without an appropriate infrastructure. They demonstrate that whether consumers are employed as peer support workers, consumer consultants or advocates, consumer representatives or as consumer educators, researchers or evaluators they require:

- clarity about roles – is it peer support or systemic advocacy, is it a representative or ‘leadership’ role?;
- clearly framed position descriptions;
- employment conditions which mirror those in the mental health workforce;
- training, mentoring and support and particularly training which can mentor people into employment; and
- the development of workforce pathways.

3. Creating an infrastructure
Jurisdictions have different infrastructures available to them to assist in building consumer activities and initiatives. Those identified as the most significant are:

- **consumer (and carer) participation frameworks**.
  Some jurisdictions, like Tasmania, have them; some are developing them. Although interstate consumers
described them as useful and important tools in holding the mental health system accountable, they also identified that the process involved in developing them is valuable, particularly if all stakeholders have been involved. They can also incorporate guidelines for supporting and building consumer leadership in the mental health sector;

- **developments in CSOs** which in some states and territories have led the way in using innovative approaches and in facilitating the involvement of consumers in decision making;

- **training and mentoring opportunities** for consumers to better participate, to build the skills of consumer workers and those involved in consumer-run organisations and to promote the cultural change required in the mental health workforce. Some jurisdictions have relied on generic health consumer training provided by health consumer peaks or similar bodies. Elsewhere there has been a development of specific training programs, for example, training for peer support workers in South Australia and New South Wales. There has also been an ad hoc involvement of consumer educators in the training of clinical and non-clinical mental health providers. All informants emphasised the vital importance of relevant training to progress consumer participation and the consumer movement more generally.

- **health consumer peak bodies.** These have operated as important supports to mental health consumer activity in some states by offering training and support for consumer workers and consumer-run organisations and generally boosting leadership capability in the health system. This can be particularly important in the absence of technical assistance centres for consumer initiatives on the US model;

- a lack of **monitoring and evaluation** of consumer activity is a significant obstacle to development and to disseminating lessons learnt from the experiences of others. Informants were keen to emphasise the importance of prioritising the monitoring and evaluation of processes and outcomes as an integral part of any consumer initiative; and

- **having a champion** within the health service to drive the consumer agenda forwards was seen as essential in a number of jurisdictions.

Future initiatives in Tasmania will need to build on these interstate and overseas experiences as well as those which are unique to Tasmania. This includes addressing the pressing issues of state wide access and state wide communication in any developments.
6. Conclusions and Recommendations

6.1 Summary

This research shows that across Australia and despite policy commitments at both a federal and state/territory level there is still a long way to go in order to achieve a satisfactory level of consumer participation. Increasingly the structural or formal arrangements are in place, there are guarantees in many documents and reports and pockets of good practice. There is also an acceptance that consumer representatives should be remunerated, that consumers should be included in planning groups and that there is a valuable role for a consumer workforce and independent consumer-run organisations. Yet, at the same time, the resourcing and support to fully endorse these trends has not been available and has led to a fragmented and stop-start approach. There is now a situation where, although the system officially endorses consumer participation, at the same time it often resists and denies the value of it, defunds it and prematurely ends initiatives. Tasmania reflects this national experience and indeed is falling behind other jurisdictions in both the range and level of consumer activity which it supports. However, opinions about how to proceed are diverse.

Evidence from both national and international sources suggests that models effective in driving the process of cultural change and recovery-orientated mental health reform require both independent advocates outside the mental health system as well as opportunities for people to be consumer advocates within the system. Ideally, and following experiences in New Zealand and elsewhere, it also involves the development of consumer-run services. Evidence suggests that the process of establishing an effective model can be hard, will take time and involves ensuring solid foundations, the setting of priorities and a staged approach. These are lessons which have been learnt locally through the recent work of the TMHCN.

6.2 Recommendations

Overall the consumer movement requires financial and organisational support if its contribution to mental health reform is to be fully developed and utilised. Given that promoting the involvement of consumers is a key priority for Mental Health Services in Tasmania this should be demonstrated through a commitment to ensure that there are a range of opportunities for the consumer voice to be heard.

In order to progress the development of consumer activity in Tasmania this report makes a number of recommendations which have emerged from the research.

6.2.1 Establishing a consumer-run organisation

The research shows that having an independent consumer-run organisation providing systemic advocacy and operating outside mental health services has significant benefits. These include providing a support base for consumers, fostering a sense of hope and inspiration and creating a cadre of knowledgeable consumers and leaders who can press for change. Three jurisdictions – ACT, Victoria and NSW – already have such an organisation and other states are now pressing to develop one. Such an organisation in Tasmania would be a strategic investment by the Department of Health and Human Services. It would strength the capacity of the Tasmanian community, particularly health and human services, to respond to the concerns of mental health consumers. It would improve policy development, provide effective systemic advocacy and an informed voice. It would also provide the opportunity to create a database of mental health consumer experiences and mechanisms for providing information and support to consumers across the state.
The key elements of this organisation are:

- an office base and paid employees with the ability to tour the state, market the organisation and have face-to-face contact with consumer groups, mental health organisations and individual consumers;
- a governing body providing strategic direction and support;
- consultation mechanisms to collate consumers’ experiences and views of services and feed them into policy and decision making structures; and
- an interactive website and hard copy newsletter to link, inform and support consumers state wide.

This fits with discussions which have already taken place in the Consumer Organisation Project. It also opens the way for any consumer organisation to acquire a wider remit in the longer term. This might entail supporting a consumer representative program (like the ACT Mental Health Consumer Network), providing consumer educators to work in clinical and non-clinical settings, developing consumer-run services (like the peer support provided by the Consumer Activity Network in NSW) or undertaking community development work to nurture and support new consumer groups (like VMIAC in Victoria).

Although the consumer movement is characterised by an extraordinary resilience in the face of setbacks the research has demonstrated that the development of consumer organisations across Australia has been halting due to a range of factors including funding. Consumer organisations are subject to the pressures all growing community organisations face as well as the possibility of cyclical illness among staff, volunteers and the governing body. It is also clear that inadequate funding for a Tasmanian organisation will undermine its capacity and sustainability. Funding levels must be adequate to attract and retain appropriately skilled staff to positions which will require community development, communications, community sector policy and programme expertise.

This research does not recommend a specific funding allocation for such an organisation. However, experiences in other jurisdictions and a growing literature demonstrate that a mental health consumer organisation may need to be seeded and funded in a different way to other community organisations. Not only do funders have to provide adequate financial resources but they also need to ensure opportunities for developing the capacity of the organisation so that it is not set up to fail. It requires a funding formula which recognises these particular needs which is clearly defined and agreed through a Service Agreement with annual performance and reporting requirements. Funding must be adequate for core functions which include the maintenance of communications mechanisms with the membership, state-wide coverage and policy development. It should be noted that industry sectoral peaks in Tasmania with governing bodies, professionally trained staff and a membership base receive funding of around $150,000 for core functions.

Recommendation 1: That the Department of Health and Human Services provide funding to establish a state wide mental health consumer run organisation with the above elements in Tasmania.

The research has clearly shown that if incorporation as an independent body is not immediate, auspicing outside the mental health sector should be considered. This avoids a situation where consumers have difficulties in escaping the role of ‘patient’ or ‘client’ and the accompanying discrimination that this can entail.

6.2.2 Developing a consumer workforce

The research has demonstrated that both in Australia and internationally the value of consumer roles in mental health services has been widely accepted. Consumer workers are seen as mechanisms for the transformation of services towards a recovery orientation, as vehicles for cultural change and for inspiring hope and as providing much needed employment opportunities for people with mental health problems. They can facilitate consumer participation activities and provide general support and mentoring to other consumers. Ideally consumer workers should be integrated into mental health services at several levels:

- as peer support workers in clinical and non-clinical environments to provide support and ‘befriending’;
- as consumer consultants, advisors or advocates in local services to oversee feedback processes and provide systemic advocacy;
- as consumer representatives to participate in decision making forums;
• as consumer educators in mental health workforce training; and
• in management positions to develop and coordinate consumer participation mechanisms at a regional or state wide level.

Lessons learnt from other jurisdictions point to a necessity for clarity about consumer workforce roles and responsibilities and a need to ensure appropriate remuneration, employment conditions, training, support and career pathways. There is also a recognition that developing a consumer workforce can take time and will need to have both short term and longer term goals. A starting point might be the piloting of consumer consultancy and/or peer support positions in Mental Health Services which are rigorously monitored and evaluated.

Recommendation 2: That the Department of Health and Human Services support and facilitate the development of a consumer workforce in Tasmania in collaboration with mental health consumers.

The Bridging the Gap report (DHHS 2004) made funding available for four consumer and family/carer advisory positions – a senior consumer and carer consultant and three regional posts. The regional posts remain unfilled. This report recommends recruiting to these posts as soon as possible to create ‘consumer/carer liaison positions’. These positions would ensure engagement systems are in place and operational, organise consultation mechanisms, provide consumers with information about their rights and foster the development of consumer support groups. This would be carried out in collaboration with any new consumer-run organisation and with CSOs. Again given the difficulties consumer workers have experienced in other jurisdictions these posts need to be accompanied by an appropriate supporting infrastructure.

Recommendation 3: That the Department of Health and Human Services use funding already allocated through the Bridging the Gap report to immediately recruit three regional consumer liaison workers to oversee the implementation of participation mechanisms and to build the capacity of consumers and carers to participate and of mental health providers to facilitate that participation.

Recommendation 4: That both public mental health services and those operated by CSOs implement positive discrimination practices in recruitment to mental health programs towards those with a lived experience of mental health problems.

6.2.3 Capacity building

Establishing a consumer-run organisation and developing a consumer workforce requires firm foundations and implementing the basic building blocks of consumer participation and empowerment. This means setting up mechanisms to:

• ensure the involvement of consumers in care planning and treatment at an individual level;
• provide information about mental health services state wide. (This process has begun with the development of a booklet for consumers, carers and family members);
• support the development of consumer support and self-help groups across the state;
• provide training for consumers to better negotiate the mental health service system, to build self esteem, to acquire the specialist skills required to fill peer support, consumer consultancy, consumer representative and
consumer educator roles and to boost expertise in the governance of consumer-run organisations; and
• provide training for clinical and non-clinical staff to understand the benefits of consumer involvement and raise awareness of the consumer experience.

Recommendation 5: That the Department of Health and Human Services provide opportunities for the training of consumers in order to build the capacity of the consumer movement.

Research demonstrates that the attitudes of staff are one of the major blocks to participation activities. It also shows that using consumers as trainers and educators enhances effectiveness and leads to more positive post-training attitudes towards consumers and involvement activities. However, although consumers are involved in the training and education of the mental health workforce in Tasmania there is no consistent approach which ensures that a consumer perspective is embedded in in-service training programs, in vocational and educational programs and at a tertiary level at the University of Tasmania.

Recommendation 6: That the Department of Health and Human Services make the consumer perspective intrinsic to all aspects of the education and training of the mental health service workforce through using consumer educators.

Recommendation 7: That the University of Tasmania through the Pro Vice Chancellor, Centre for the Advancement of Learning and Teaching, ensures that the consumer perspective and the use of consumer educators is embedded in teaching related to mental health issues.

As well as the need to establish consumer perspective teaching, there is also a need to establish more consumer perspective research.

Recommendation 8: That the Department of Health and Human Services sponsor a dedicated mental health consumer research fellowship at the University of Tasmania.

Both this research and the Review of Consumer and Carer Participation conducted by Mental Health Services has shown that many consumers want support and self-help groups where they can gain mutual support and acceptance from others, promote recovery and engage in social activities. There are currently a number of such groups across the state operating independently as well as being supported by Mental Health Services and by CSOs. However there is also a significant unmet demand for these groups.

Recommendation 9: That the Department of Health and Human Services ensure mechanisms are available to develop and nurture consumer support and self-help groups across the state.

Unlike some jurisdictions Tasmania has no peak representative body for consumers in the health system. Elsewhere these have operated as mechanisms for capacity building in the mental health consumer community, for example, the Health Consumers Alliance in South Australia and the Health Consumers Council in Western Australia. They have provided support for mental health consumers, offered training to promote the ability of consumers to become active, built a health consumer leadership and in some instances auspiced specific mental health consumer activities. A peak body in Tasmania should provide a range of advocacy and information services for consumers as well as a focal point for training initiatives and leadership building.

However, current proposals for such a body in Tasmania suggest that the consumer peak would represent the interests of clients of health and human services potentially broadening its remit to include a wide diversity of interests (such as clients of homelessness services, public housing tenants, ex-prisoners and so on). Representing such a breadth of concerns would clearly be difficult and would require strong linkages with a membership of effective consumer organisations.
Recommendation 10: That the Department of Health and Human Services strengthen consumer engagement with Tasmania’s health and human service system by establishing a peak body to represent consumers and the community, to support engagement activities and to strengthen consumer networks and organisations.

The Social Inclusion Unit is currently developing a social inclusion strategy for Tasmania which will outline a set of guiding principles. Anglicare has already, in its submission to the consultation process on the strategy (Anglicare 2009) advocated for a human rights framework and for local and service user accountability whereby any strategy must ensure a voice for people at the local level. Although many people with mental illness fully participate in the community, there are a proportion who experience severe disadvantage and exclusion and whose voices are not heard. This is particularly true of those with severe mental illness.

Recommendation 11: That the Social Inclusion Commissioner advise the Premier to direct the Tasmanian Government’s Social Inclusion Unit to explore the civic participation of marginalised groups as a goal of the Tasmanian Social Inclusion Strategy.

This recommendation supports that made by the TMHCN in their submission for mechanisms to support the civic participation for people with a mental illness (TMHCN 2009).

6.2.4 Oversight, monitoring and evaluation

Across Australia the evidence base for mental health consumer participation and consumer-run initiatives is sparse. What evidence there is is overwhelmingly positive, but there is also a need to ensure both routine monitoring and systematic and comprehensive evaluation. Whatever initiatives or mechanisms are established in Tasmania it is imperative that their processes, impact and outcomes are measured. As well as public mental health services provided by the state there are now an increasing and diverse range of community mental health services delivered by CSOs. This means that CSOs must also be involved in any monitoring and evaluation exercises.

The National Standards for Mental Health Services require services to have policies and procedures relating to consumer and carer participation. These are reviewed by external accreditation bodies. However despite this requirement there is a lack of mandates on how processes or mechanisms should be facilitated. This leaves room for local interpretation and fosters an environment where responses are ad hoc and unsystematic.

Recommendation 12: That the Department of Health and Human Services ensure that performance indicators and outcome measures are developed and monitored for any consumer participation mechanisms or consumer initiatives in both public mental health services and in CSO provision and that this is undertaken in partnership with consumers and carers.

Recommendation 13: That the Department of Health and Human Services ensure that a requirement to involve consumers and/or to employ consumer workers is built into the contractual requirements for CSOs providing mental health services.

Recommendation 14: That CSOs delivering mental health services ensure that the existence of consumer engagement mechanisms are a key quality indictor in their organisation.
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