Person-centred rehabilitation (PCR) means treating each service user undergoing rehabilitation as an individual.

There is a strong and committed drive to provide this type of rehabilitation both nationally and internationally.

To help rehabilitation teams improve their ability to deliver person-centred rehabilitation a training package was developed.

**Package Objectives**

1. Create a shared understanding of PCR in your service context.
2. Gain awareness of tools to grow PCR in your service context.
3. Consider how to assess PCR in your service context.
4. Ideas for sustaining PCR in your service context.

**Package Outline**

Four sessions of 45 minutes | Short exercises during sessions | Homework

- Session 1: Understanding PCR
- Session 2: Critical reflection and PCR
- Session 3: Empowerment | Shared decision making | Dealing with difference
- Session 4: Evaluating & sustaining PCR

- The package is designed to be undertaken in teams.
- The sessions will be delivered by a trained facilitator.
- Some examples of package content appear below.

**Key Sources:**

- Elwyn, G., Frosch, D.L., Kobrin, S., 2016. Implementing shared decision-making: consider all the consequences. Implement. Sci. IS 11, 114.

Development of this package was supported by The Hopkins Centre.
Self-management skills after stroke: What is the role of the hospital stroke team?

Kylie Bower & Carly Gomura
Princess Alexandra Hospital
Adjunct Research Fellows, Griffith University

Introduction
With an ageing population and improving acute care, increasing numbers of Australians are surviving stroke. However, 87% of stroke survivors face limitations to their activities of daily living, and over 30% are under 65 years of age, and potentially face decades living with the effects of their stroke.

To help stroke survivors navigate life after stroke, Stroke Guidelines recommend the use of generic or stroke-specific self-management programs in the community. However, there is currently little literature exploring the impact of incorporating aspects of self-management into hospital services. Recent research recommends that: ‘Research is required to explore a new model of stroke self management which is integrated across secondary, primary and community care and adopts a whole systems approach.’ (Parke et al, 2015, p 20)

Methods
- A descriptive, qualitative design will be used across 2 studies.
- Purposive sampling will be used to recruit staff (study 1), stroke survivors and carers (study 2) across three hospital stroke programs at the Princess Alexandra Hospital: the acute stroke unit, geriatric and rehabilitation unit, and the day hospital service.
- Semi-structured focus groups and interviews will explore elements of self-management represented in figure 1. Interviews will be recorded, transcribed and thematic analysis will be used to identify key themes.

Purpose
A program of research is planned to explore how self-management is conceptualised in a hospital service, which elements are currently evident, and whether there is value or feasibility in incorporating any other aspects of self-management. The links between these self-management components and those existing in community services will also be considered.

Planned studies
- Study one: staff perspectives across hospital stroke services
  - Acute MDT interviews
  - Rehab MDT interviews
  - Day Hospital MDT interviews
- Study two: patient/carer perspectives across hospital stroke services
  - Reflections on self-management interventions in the acute service.
  - Reflections on self-management interventions in acute and inpatient rehabilitation services.
  - Reflections on self-management interventions in acute, rehabilitation and day hospital services.
  - Reflections on self-management interventions across hospital and community experiences.

Expected Results
We expect to better understand how hospital stroke services contribute to a stroke survivor’s self-management journey.
We also hope to identify whether any changes to the hospital stroke services could optimise self-management outcomes for stroke survivors.

Progress
- Partnership developed with Griffith University School of Occupational Therapy.
- Ethics application commenced for study 1.

Literature Cited

Acknowledgement
We wish to acknowledge the contributions of Ms Amelia Greenway (honours student), Professor Louise Gustafsson, and Ms Kim Walder (Lecturer) of the Occupational Therapy Department, Griffith University, in the commencement of study one.
Introduction
Responsive, integrated, safe and effective (i.e., quality) care depends on individual nurses’ professional practice together with the practice environment (Stallings-Welden & Shirey, 2015).

A sense of personal wellbeing, shared decision-making, leadership, safety, quality and professional development are six service standards that support quality nursing care (Dans, Pabico, Tate & Hume, 2017).

Wellbeing has been linked with job satisfaction, personal assessment of own health, and measures of work engagement. Nurses’ wellbeing was identified by mental health nurses as a priority area for action within tertiary services.

Research Question
What workplace factors contribute to nurses’ wellbeing in tertiary mental health care?

Method
Scoping review as per Arksey and O’Malley (2005) and as refined by Colquhoun, Levac et al (2014).

Results
- Structuring care provision:
  Mental health nurses working in acute care environments appear more stressed than nurses working in other contexts. The way that care is organised within the system creates more or less possibilities for healthy and sustainable workplaces that do or do not support nurses’ wellbeing.

- Practice environment
  Five factors in the practice environment that influence nurses’ wellbeing were identified: (1) Scope and clarity of nurses’ roles; (2) Managerial leadership style; (3) Communication and appropriate flow of information; (4) Working conditions (staffing, pay, work hours, regulations, administration and opportunities for career advancement); and (5) Interdisciplinary relationships (i.e., strength of relationship, opportunities for career advancement).

- Caring work
  Identified factors that influence wellbeing and are associated with nurses’ care work include: (1) Patient characteristics; (2) Occupational violence; (3) Workload; and (4) Social support.

Discussion/Conclusion
Managerial support is key for optimising wellbeing outcomes of nurses. A transformational leadership style is encouraged.

A sense of being supported is important for nurses within organisational and direct care contexts. Appropriate personal and social support that extends beyond organisational and care contexts is necessary. Therefore, nurses need easy access to multiple support systems if their wellbeing is to be optimised.
CONSUMER ENGAGEMENT IN HEALTH SERVICES ACROSS QUEENSLAND

What roles do consumers and carers currently have in health services?

What impact does Health Consumers Queensland (HCQ) have on consumer engagement in Queensland?

WHO DID WE GET FEEDBACK FROM?

via interview

- HCP staff, board members and members of HCQ consumer advisory group
- Consumer and carer representatives across Queensland
- Health services staff

via survey

- Consumer and carer representatives across Queensland
- Health services staff across Queensland

WHO ARE THE CONSUMER & CARER REPRESENTATIVES?

- 77% Over 65 years of age
- 88% working in government service
- 35% 62 of 128 have been in role for 1-5 years

WHAT DID THE STUDY FIND?

Requests for HCQ services: HCQ staff reported increasing requests for their services.

Value for consumer engagement: Consumers reported feeling valued and validated by health service organisations.

Type of consumer engagement: Consumer engagement occurred on a continuum from simple passive activities to more meaningful and influential activities.

Remuneration of consumers and carers: is inconsistent and contested. A statewide policy for remuneration is required.

Depth of consumer engagement: Some qualitative evidence suggested that there were positive shifts in knowledge and use of consumer engagement in health services.

Accessibility of consumer engagement: Qualitative evidence showed that remuneration of consumer and carer representatives was one policy area that was inconsistent and contested.

Embedded consumer engagement: There was no consistent evidence to conclude that authentic consumer engagement was embedded in the health service system as standard practice.

WHAT WERE THE RESEARCH QUESTIONS?

- Requests for HCQ services?
- Value for consumer engagement?
- Type of consumer engagement?
- Remuneration of consumers and carers?

WHAT DID PEOPLE TELL US ABOUT THEIR INTERACTIONS WITH HCQ?

“HCQ have made an enormous impact”

Most useful engagement strategies:
- Training and workshop
- Electronic newsletter
- Networking sessions
- Annual forum

1/2 of participants attended training or workshops
44% received e-news
32% attended Annual Forum

Interesting result

Attending networking sessions and webinars was perceived to be most useful by people who were from an Aboriginal and Torres Strait or culturally and linguistically diverse background.

“HCQ helps me to connect a lot of dots”

“The more that HCQ helped individuals prepare for their role, the more likely they were to report higher levels of self-efficacy in their role.”

“HCQ have been really instrumental”

Assoc. Professor Carolyn Ehrlich, Dr Maddy Slattery, Ms Kelsey Chapman and Prof. Elizabeth Kendall

HCQ = Health Consumers Queensland
Integrating an Evidence-based Clinical Exercise Physiology Service Model into a Comprehensive Multiprofessional Rehabilitation Service for People with Brain Impairment: An Integrated Knowledge Translation Approach.

Clanchy, K., Tweedy, S., Kennedy, A., Jeffery, S., Gainforth, H.

**Introduction**

Investment in healthcare research continually produces significant breakthroughs for methods of treatment and care which can yield significant benefits for patients, financial value gains and returns on investment [1].

There are significant barriers to translating evidence into clinical practice including differences in the implementation environments, time and resource limitations of practitioners, insufficient training, lack of feedback and incentives for use of evidence-based practices, and limited organisation infrastructure to support translation [2].

Integrated Knowledge Translation (IKT) is a theory informed approach for knowledge translation that is characterised by the ongoing and collaborative involvement of stakeholders and the tailoring of knowledge to the needs of the individuals and organisations who will be implementing it [3, 4].

**Purpose**

The Acquired Brain Injury Transitional Rehabilitation Service (ABI-TRS) is currently developing its clinical Exercise Physiology service arm which requires the integration of current evidence regarding physical activity promotion in adults with an acquired brain injury in order to facilitate a best-practice model. The Adapted Physical Activity Program (APAP) is a physical activity promotion intervention that has been demonstrated to increase physical activity adoption in adults with brain impairment [5].

The aim of this project is to use an IKT approach to increase the prospects of successfully translating the APAP into ABI-TRS.

**Methods**

IKT has two components: 1) knowledge creation which includes the evaluation of an innovative idea and the synthesis of research findings into useable tools/products, and 2) action cycle including adaptation to the local context and assessing barriers and facilitators related to knowledge adoption. This information is used to develop a plan for the translation of knowledge into clinical practice [3]. The application of the IKT framework for the development of a plan for the translation of the APAP into ABI-TRS is presented in Figure 1.

**Expected Results**

A knowledge translation plan has been developed that addresses important considerations including: the willingness of stakeholders from multiple settings to adopt the APAP; the perceived fit of the APAP into ABI-TRS; and the ability of ABI-TRS staff to implement key components of the intervention in routine clinical practice [2]. This translation plan will be implemented in January 2019.

The effectiveness of the translation of APAP into ABI-TRS will be evaluated using the RE-AIM (Reach, Efficacy, Adoption, Implementation and Maintenance) Framework [6].

**Literature Cited**

5. Clanchy et al Neurorehabilitation and Neural Repair 30(9): 854-865.
Integrating technology into a new acquired brain injury rehabilitation service

Gisela Brittain 1, Elizabeth Beadle 1, Elizabeth Spitzer 1, Areti Kennedy 1,2
1. Acquired Brain Injury Transitional Rehabilitation Service, Division of Rehabilitation, MSH, Brisbane Australia
2. The Hopkins Centre, Division of Rehabilitation & Griffith University, Brisbane, Australia

Introduction
Integrating technology in rehabilitation after brain injury is common. This involves: (i) utilizing technology for service delivery, including telehealth, tablets and smartphones; and (ii) utilizing technology-enabled therapy, including web-based interventions and social media platforms.

The Acquired Brain Injury Transitional Rehabilitation Service (ABI TRS) commenced in Brisbane in 2017 and provides interdisciplinary community rehabilitation for clients with a brain injury.

The ABI TRS aimed to implement evidence-based technology to enhance rehabilitation for clients with ABI into a new community transitional rehabilitation service.

Method
A change management approach was used to establish the use of technology into the rehabilitation service. This included:

- Identified area of need; Literature review of current evidence-based practice
- Working group formed, consultation with inter-disciplinary team
- Exploration of affordability and type of technology suitable for the service
- Benchmarking with other community rehabilitation teams
- Clinician skill, knowledge and experience of use of technology was also explored

Results

**Identified area of need:** Recent systematic reviews1,2 identified several benefits of technology-based rehabilitation including: 1. **Intensity:** clinicians can provide patients with a greater intensity of sessions; 2. **Client-centred treatment:** Technology allows the clinician to tailor the tasks to clients impairments and goals for a more personalized rehabilitation program3; 3. **Improved rehabilitation outcomes:** studies have shown various degrees of improvement and greater independence for clients with cognitive1 and communication impairments.

**Clinician skill, knowledge & experience:** Team survey revealed that clinicians had skill and knowledge of technology usage, however lacked experience of using technology in a community rehabilitation setting. To address this barrier, clinical processes, training and education were provided and technology items have been successfully embedded within the community rehabilitation program.

**TRS team consultation:** Through ABI TRS service review, along with the results of the literature review, a need for internet usage in home and smart devices for therapeutic use was identified. Three iPads (2 with data) were purchased. Individual disciplines were consulted with regards to evidence based apps for: compensatory strategies (e.g., diary, communication aids); skill building (e.g., language therapy); education (e.g., brain information); leisure (e.g., games, music); support building (e.g., circles of support); and client safety/independence (e.g., GPS tracking; supported emergency calls). Additional dongles were purchased in order to access the internet when using laptops. Review of dongle use is underway.

**Benchmarking:** Community rehabilitation and inpatient services were consulted on their technology usage and resources. Community rehabilitation services reported using service laptops (each staff member) and otherwise the clients own devices, and inpatient services reported using service laptops, internet, and iPads.

In the last 12 months over 60% of ABI TRS clients have received technology based rehabilitation using service devices.

Conclusion
- Technology was successfully implemented into a new transitional rehabilitation service.
- Technology has enabled clinicians enhance client’s rehabilitation through using alternative modes of treatment and providing a tailored treatment program.
- Future research is needed in to how technology is being used by the interdisciplinary team with clients (e.g., specific apps and therapy treatment targets), as well as how clients use technology to enhance their recovery and wellbeing after injury. There is also a need to review both clinician and client confidence and experience with technology use.

References
Fostering self-management within an upper limb therapy group for stroke survivors and their caregivers

Carly Gomura & Kylie Bower
Princess Alexandra Hospital

Introduction

Self-management is recommended in the Australian Clinical Guidelines for Stroke Management 2017. It aims to help stroke survivors adopt strategies to manage changes in physical and cognitive ability, relationships, and to better participate in their community. 1

Opportunities exist for self-management care planning to commence within the acute inpatient episode of care, to best prepare stroke survivors and their caregivers to manage their chronic condition for the long-term. 2,3

Purpose

Existing models outline a process for which self-management behaviours may be developed and adopted. 4

This process requires the provision of effective therapeutic interventions to allow stroke survivors to acquire self-management skills and the self-efficacy to put these skills into practice.

Our project explored the impact of therapeutic interventions in imparting self-management skills, behaviours and self-efficacy for upper limb management post-stroke.

Methods

A stroke-specific upper limb education group in an inpatient rehabilitation setting was redesigned to target the development of self-management skills through the provision of the following interventions which are supported in the literature: 3,4:

Goal setting: impairment, activity and participation goals are set in collaboration with stroke survivors and their caregivers, as part of an established joint Occupational Therapy/Physiotherapy upper limb management process.

These goals are incorporated into education and performance, and goal performance is monitored at each group.

Active information provision: the group commences with education and discussion on a rotating list of topics, using an education resource that adopts a self-management approach. A variety of multimedia tools are incorporated, such as the EnableMe website.

Involving carers: caregivers are encouraged to attend the group, to develop their own knowledge-base and to develop skills to support the stroke survivors ongoing self-management care plan.

Brian & Catherine working on reaching for a fork, for the goal of eating steak.

Peer support: is actively facilitated. For example, once attendees of the group are familiar with the education topics and exercises, they are encouraged to lead discussion or demonstration of skills as a means of peer modelling and support.

Emotional / social focus: attendees are encouraged to share their experiences with managing their upper limb throughout their recovery. This is supplemented with a targeted education topic addressing basic social and emotional aspects to upper limb changes after a stroke.

Sharing achievements at the end of the group.

Evaluation of the upper limb education group was conducted via pre-post survey of participants, measuring their perceived knowledge, skills and confidence in caring for their stroke-affected upper limb. This outcome may represent the discrepancy between perceived and attained knowledge.

Results

Participants demonstrated improved self-ratings of knowledge, skill and confidence in upper limb self-management following participation in the group.

Participants identified a continued practice of upper limb self-management behaviours following completion of the group, and the benefit of peer support and encouragement that the group provided.

The separate cohort of stroke survivors not involved in this group displayed high levels of perceived knowledge, skills and confidence in caring for their stroke-affected upper limb.

Conclusion

A self-management framework can be successfully incorporated into a group setting to foster self-management knowledge, skills and the efficacy to translate these into ongoing self-management behaviours. A group setting provides an opportunity for peer support amongst participants, and can be an appropriate setting for consolidating self-management skills.

Acknowledgements

I wish to acknowledge the combined efforts of the PA Hospital GARU Occupational Therapy team and students for their work towards this project outcome over 2017-2018.

References

2. Pearce
Introduction

Adults undertaking rehabilitation for an acquired brain injury (ABI), spinal cord injury (SCI), or persistent pain (PP) condition (and their families) can experience a range of emotions and behaviours during the process of injury and adjustment, some of which may be experienced as concerning, confronting, or challenging for rehabilitation clinicians (Behaviours of Concern; BOC).

BOC have been widely examined in emergency, mental health and aged care settings [1, 2] however there is limited research regarding the type of BOC and their impact on staff working in frontline rehabilitation settings. An understanding of these issues is a necessary first step in developing local approaches for ensuring an environment in which rehabilitation professionals feel supported in working with clients and families who exhibit BOC.

Research Overview

As part of a larger program of research, a systematic review (Study 1) was conducted to identify and synthesise existing rehabilitation research with attention to determining the specific types of client and family BOC reported by rehabilitation clinicians and their impact.

Systematic Review Method

Four databases (PubMed, CINAHL, PsycINFO, and Web of Science) were systematically searched for eligible studies. Studies were included in the review if they:

- Reflected original, peer-reviewed research published between 2000 and 2018;
- Described the views of in-patient and/or community-based rehabilitation professionals working with adults with ABI, SCI, or PP conditions on one or more of the following:

1) Client or family BOC,
2) The impact of client or family BOC on rehabilitation staff

19 eligible studies were included in the review.

Systematic Review Results

The majority of eligible studies focussed on clients with ABI (n=17), with very few studies examining BOC for SCI (n=1), PP (n=1), and families (n=4).

1) Client / family BOC for rehabilitation staff:

<table>
<thead>
<tr>
<th>Behaviours of Concern</th>
<th>No. of studies</th>
<th>ABI</th>
<th>SCI</th>
<th>PP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Client behaviours:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Language / communication difficulties</td>
<td>7 ▶</td>
<td>▶</td>
<td>▶</td>
<td></td>
</tr>
<tr>
<td>Expressed emotion (e.g., emotional lability, non-expression of emotions, grief)</td>
<td>6 ▶</td>
<td>▶</td>
<td>▶</td>
<td></td>
</tr>
<tr>
<td>Aggression (e.g., physical, verbal or gestural)</td>
<td>5 ▶</td>
<td>▶</td>
<td>▶</td>
<td></td>
</tr>
<tr>
<td>Cognitive impairment (e.g., lack of awareness, impaired memory/ attention / concentration)</td>
<td>5 ▶</td>
<td>▶</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-compliance/ resistance to therapeutic approach</td>
<td>4 ▶</td>
<td>▶</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Impulsive, erratic or disinhibited behaviours</td>
<td>4 ▶</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Denial / minimising of limitations</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lack of motivation</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agitation</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Absconding</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High demand on staff</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Suicidal behaviours</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not taking feedback seriously/ making excuses for performance</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatigue</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fixed ideas about injury/ rehabilitation (e.g., difficulty accepting that psychological factors influence pain experience)</td>
<td>1 ▶</td>
<td>▶</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Family behaviours: | | |
| Intrusive behaviours | 3 ▶ | |
| Expression of grief | 1 ▶ | |
| Need for continual reassurance and explanation | 1 ▶ | |
| Verbal abuse | 1 ▶ | |
| Presence of drug paraphernalia in the home | 1 ▶ | |
| Inappropriate behaviour | 1 ▶ | |
| Presence of family member triggers challenging client behaviour | 1 ▶ | |
| Lack of support/ involvement | 1 ▶ | |

2) Impact of BOC on rehabilitation staff:

<table>
<thead>
<tr>
<th>Perceived Impact</th>
<th>No. of studies</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emotional</td>
<td>9</td>
<td>Feelings of disappointment, frustration, helplessness, anxiety, fear, overwhelm</td>
</tr>
<tr>
<td>Physical</td>
<td>2</td>
<td>Physical health risk, bruises, scratches, fractured bones, sore muscles</td>
</tr>
<tr>
<td>Loss of productivity / workforce</td>
<td>1</td>
<td>Sick leave, consider leaving / leave job</td>
</tr>
</tbody>
</table>

Interpretation

From the included studies (n=19):

- The majority of studies reported on BOC within the ABI setting with very few studies reporting on BOC in the SCI or PP setting.
- The majority of studies focussed on client behaviour, with fewer studies reporting on BOC experienced from families.
- 15 categories of client BOC and 8 categories of family BOC were identified.
- The majority of studies appeared to focus on ‘active’ or ‘overt’ behaviours (i.e., physical/ verbal aggression), with fewer studies reporting on ‘passive’ or ‘covert’ behaviours (i.e., self-neglect).
- 10 studies reported on the impact of BOC on staff. Emotional and psychological impacts were frequently reported with physical health impacts and reduced work productivity also described.

Conclusion

The current study has identified existing gaps in the rehabilitation literature and highlights the important need for continued research to inform service development and resourcing for supporting rehabilitation clinicians in working with clients and families who exhibit BOC following injury and adjustment.

Future Directions

As part of a wider program of research, further qualitative (Study 2) and quantitative (Study 3) studies are planned to examine the beliefs and expectations of rehabilitation clinicians in working with individuals who demonstrate BOC. It is anticipated that this research will further inform the development of local approaches for ensuring an environment in which frontline rehabilitation professionals feel supported in working with clients and families who exhibit BOC.

References


Acknowledgements

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Special thanks must be extended to Prof Michele Foster, Prof Elizabeth Kendall, Dr Sindhur Atreesh, A/Prof Annette Broome, Dr Harriet Bodimeade and Ray Quinn for their support and contributions to the current program of research.
Identifying the dispersion of disability services in Queensland priority areas

Lakhani, A.*, Parekh, S., Grimbeek, P., Gudes, O., Watling, D.P., Harre, P. & Stocker, J.

Introduction
• Disability support services that provide advocacy, information, and health and social care are integral to the health and wellbeing of people with disability.
• Research indicates that such services are essential in addressing individual comorbidities by linking clients with additional health services [1], improving community participation [2] and education and training outcomes [3], and supporting self-directed decision making [4].
• It is important that disability support services are proximately available. In Australia, the proximate availability of services has been questioned, even for those in metropolitan settings [5].

Purpose
• Recent research has established that areas in South-East Queensland with high proportions of people with disability are underserviced in terms of allied health services [6].
• Only a small fraction of research has focused on the use of Geographic Information System (GIS) technologies to clarify the geographic dispersion of disability services in relation to people with disability.
• The current study aimed to:
  1. Use spatial analysis to identify priority regions in Queensland with a significantly high number and proportion of people with disability.
  2. Use spatial analysis to estimate travel times from localities within these regions to disability services.

Methods
• The numbers, and proportions of people with disability across Queensland SA2 regions was sourced from the 2016 Australian Bureau of Statistics (ABS) census data. Localities within SA2 regions was sourced from the ABS, 2016 Urban Centre & Locality classification. The location of disability services in Queensland was provided by Health Direct.
• A Hot Spot Analysis (Getis-Ord Gi*) within ArcGIS was conducted to ascertain SA2 regions with significantly high numbers and percentages of people with disability in Queensland.
• The travel time between localities and disability service organisations was calculated via the network analysis feature in ArcGIS.

Findings
• Thirty-eight SA2 regions in Queensland were identified as having a high number and proportion of people with disability. These regions are presented in Figure 1. Sixty-two urban centre localities were located within these regions.
• Descriptive statistics relating to the travel time (in minutes) to three types of disability services has been included in the table below.

<table>
<thead>
<tr>
<th>Service Type</th>
<th>Mean (SD)</th>
<th>Min.</th>
<th>Max.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aids and Equipment</td>
<td>32.47 (17.56)</td>
<td>1.92</td>
<td>69.67</td>
</tr>
<tr>
<td>Information and Referral</td>
<td>22.53 (16.76)</td>
<td>0.90</td>
<td>70.34</td>
</tr>
<tr>
<td>Support and Advocacy</td>
<td>42.24 (25.45)</td>
<td>4.53</td>
<td>98.49</td>
</tr>
</tbody>
</table>
• On average, the greatest travel time for localities within priority areas was to Support and Advocacy services. The shortest travel time was to Information and Referral services.

Findings Continued...
• Friedman’s test (non-parametric equivalent to repeated measures ANOVA) was used to examine the extent of differences in travel time to the three types of disability services.
• The three travel times were significantly different. When the service with the shortest travel time (Information and Referral), was excluded, the travel times to Support and Advocacy services turned out to be significantly greater than travel times to Aids and Equipment services.

Points for Consideration
• The findings are consistent with travel times to Support and Advocacy disability support services for localities within priority areas being greater than the two other types of disability services, and consequently this type of service has the least proximate accessibility for localities considered.
• While travel time is important, the proximate availability of specific disability services might have less impact on access when telehealth or homeshare visits are available as options.
• Furthermore, there are potential services which have not been listed, and thus not included in the analysis.

Figure 1: Areas with significantly high levels of disability in Queensland

References
How is this working for you? Early insights from a masterclass to advance frontline rehabilitation nursing research

Letitia Burridge¹, Alison New², David Lewis³, Jenny Kohlhardt³, Tracey Tattam², Michele Foster¹, Melissa Legg¹
¹The Hopkins Centre, Griffith University; ²Division of Rehabilitation, Princess Alexandra Hospital; ³Division of Medicine, Princess Alexandra Hospital

Navigating the research process: from here to here

Background
- Nurses more than ever have higher education¹–², but are under-engaged with research vs. other health professionals³.
- Nurse-led research is impeded by lack of experience, knowledge or skills, lack of insight into the complexity of the research process, and the perennial problems of being time-poor and having inadequate resources and support⁴.
- Stronger research capacity and culture can enable nurse-led, practice-relevant research and its dissemination⁵,⁶,⁷.

Rationale
Nurse-led research can improve the quality of rehabilitation care, and showcase rehabilitation nursing as a specialty.

Purpose
To assess the value and impact and experience of a research masterclass program for rehabilitation nurses.

Method
Mixed methods, pre-post evaluation.

Sample and setting
11 rehabilitation nurses practising within Metro South Health, Queensland, Australia

Early results

<table>
<thead>
<tr>
<th>Aim</th>
<th>1. Value of Masterclass</th>
<th>2. Impact of Masterclass</th>
<th>3. Experiences of research process</th>
</tr>
</thead>
<tbody>
<tr>
<td>Focus</td>
<td>Satisfaction with workshop.</td>
<td>Engagement with research.</td>
<td>Participants’ experiences with a mentored research project.</td>
</tr>
<tr>
<td>Method</td>
<td>1-page self-administered survey at end of each workshop.</td>
<td>4-page self-administered survey at workshops; baseline demographics.</td>
<td>Group interview in second workshop, audio-recorded and transcribed for deductive/ inductive thematic analysis.</td>
</tr>
<tr>
<td>Measure</td>
<td>New world reaction sheet⁸: 21 statements across 9 domains, 4-point Likert scale to rate level of agreement.</td>
<td>Research Spider⁹: 4 domains (experience, confidence, interest, opportunity), each with 10 items, 5-point Likert scale to rate level of impact.</td>
<td>Open-ended questions: 1) expectations and experiences; 2) how mentoring influenced current involvement in research; 3) broader engagement with nursing research.</td>
</tr>
</tbody>
</table>

Results
- Fig. 1: Satisfaction with workshop (agreed/strongly agreed)
- Fig. 2: Changes in research engagement mean scores at workshop 2

Provisional conclusions
- The workshop evaluations suggest the Masterclass supports a keen pre-existing interest in and desire for nurse-led rehabilitation research.
- Participants remained highly interested in research, and their experience, confidence and opportunity increased between the workshops.
- Participants remained committed to their research goals in a landscape that included opportunities for colleagues and their practice specialty.

References

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1The Hopkins Centre, Griffith University; 2Division of Rehabilitation, Princess Alexandra Hospital; 3Division of Medicine, Princess Alexandra Hospital

Disclosure: The authors have no disclosures to make.
Introduction

Consumers of rehabilitation services (also known as service users) have a valuable role to play in the development and delivery of rehabilitation services and in conducting and translating rehabilitation research.

How to meaningfully partner with consumers in these activities is an important issue for the Division of Rehabilitation (DoR) and a key focus of The Hopkins Centre (THC).

The DoR and THC recognised that an appropriate and coherent strategy was required to effectively engage and partner with service users. A 12-month Consumer Engagement Strategy Project is in process.

The Project aims to design, plan and document a sustainable consumer engagement strategy which embeds into clinical, management, education and research activities of the DoR and THC.

Defining Consumers & Engagement

A consumer is someone who is using or has used a DoR service. Consumers are also referred to as service users.

Consumers can also be family members or carers of people using or who have used DoR services.

Consumer Engagement is meaningfully involving people (consumers) who have used DoR services in the planning, delivery & evaluation of DoR services & research.

Different Levels of Consumer Engagement

The IAP2’s Public Participation Spectrum is often used as the standard to define the consumer role in engagement processes. Different roles are appropriate for specific activities.

- **EMPOWER**
  - Users of a rehabilitation unit suggest changes to the existing service model in feedback provided to the hospital. Staff follow up on the feedback & ask the users to partner with them to develop a consumer-centred service model.

- **COLLABORATE**
  - A rehabilitation unit puts together a steering committee including staff members, researchers & service users to oversee the development of a new model of care. The steering committee is involved in all aspects of the decision making.

- **INVOLVE**
  - Staff from a rehabilitation unit put together a focus group of consumers who have used the unit to consider their concerns and preferences prior to the development of a new model of care. This information is incorporated into the new model.

- **CONSULT**
  - A rehabilitation unit has designed a new service for the region. They ask the hospital consumer advisory committee for feedback, some of which is evident in the final service model.

- **INFORM**
  - A hospital produces a fact sheet to let the community know about a new service being offered in the rehabilitation unit.

Methods

The Project involved four interconnected phases:

- **Initiation & Planning**
  - Develop project plan
  - Establish Reference Committee
  - Obtain ethics approval & governance authorisation

- **Scoping & Stakeholder Consultation**
  - Scan existing QH strategies
  - Rapid literature review
  - Stakeholder consultation – staff, researchers & service users

- **Strategy Development**
  - Develop draft strategy
  - Identify staff & consumer training needs

- **Review & Evaluation**
  - Next step

Stakeholder Consultation Outcomes

- Level of interest in being involved in consumer engagement activities >70% for all stakeholder groups.
- High level of agreement regarding important Consumer Engagement Strategy attributes.
- Staff & researchers identified skill areas to build, including: procedures & processes, communication skills, activity planning & partnering for service co-design & evaluation.
- Identified facilitators of consumer engagement included the following:
  - **Staff:**
    - Allocated time
    - Awareness of opportunities
    - Training / Education
  - **Consumers:**
    - Contribution is valued & respected
    - Accessibility & flexibility
    - Understandable & timely information
    - Training for effective participation
    - Feedback on engagement outcomes

Emerging Strategy Domains

The literature review & consultation outcomes have contributed to 6 potential strategy domains.

References

2. International Association for Public Participation (IAP2) Public Participation Spectrum [https://www.iap2.org.au/About-Us/About-IAP2-Australasia-/Spectrum]
Filling the transitional care gap: First year outcomes of the ABI Transitional Rehabilitation Service Pilot Project

Areti Kennedy 1, Mandy Nielsen 1, David Borg 1, Kerrin Watter 1
1The Hopkins Centre and Division of Rehabilitation, Metro South Health

Introduction
The Acquired Brain Injury Transitional Rehabilitation Service (ABI TRS) is a 5-year pilot which aims to facilitate early community reintegration for individuals with acquired brain injury (ABI) and their families.

This service was established to address an identified unmet need in the continuum of ABI rehabilitation in Queensland. Funding for the pilot was received from the Motor Accident Insurance Commission (MAIC) in July 2016, with clinical services commencing in January 2017.

To determine the impact of ABI TRS on community re-integration, funding included an embedded service evaluation for the duration of the pilot.

Methods
People with ABI (n=71) completed the 12 week ABI TRS community-based program, comprising an intensive, individualised, goal directed program using an interdisciplinary, client-centred approach. Self and/or practitioner-rated measures were completed at 0 (baseline) and 3 months. Results were compared to a quasi-control (CON) where no transitional service existed. Data were analysed using linear mixed models in a Bayesian framework.

Results
1. Baseline ratings in all outcome measures were statistically worse for ABI TRS clients compared to CON i.e. the ABI TRS cohort was more impaired than the CON group at the first time point.

2. There were statistical improvements in measures of global function (MPAI-41) and psychological wellbeing (DASS-212) compared to CON.

3. Psychosocial function (SPRS-21) and health-related quality of life (EQ-5D1) were statistically improved at 3 months compared to baseline.

Conclusions
ABI TRS clients demonstrated statistically improved or matched outcomes at 3 months compared to the quasi-control group.

The ABI TRS sees a clinically diverse range of individuals, and early results indicate that clients generally exhibit superior outcomes following program completion.

1Mayo Portland Adaptability Inventory-4; 2Depression Anxiety Stress Scale 21; 3Sydney Psychosocial Reintegration Scale-2; 4EuroQuol Health Questionnaire
Addressing Client Needs Through Interdisciplinary Multi-site Groups in Acquired Brain Injury Transitional Rehabilitation

Nina Wegener¹, Sarah Jeffery², Sarah Kekki³, Elizabeth Beadle¹, Kerrin Watter¹,², Areti Kennedy¹,²
¹Acquired Brain Injury Transitional Rehabilitation Service, Division of Rehabilitation, MSH, Brisbane Australia
²The Hopkins Centre, Division of Rehabilitation & Griffith University, Brisbane, Australia

Introduction
Transition from hospital-to-home is a critical time for people with acquired brain injury (ABI) and their families. Recognised issues include changes to physical, cognitive and communication skills, which impact independence, reintegration and community access.¹ ² The Acquired Brain Injury Transitional Rehabilitation Service (ABITRS) is a new service that provides time limited community-based rehabilitation to people with ABI discharging home from hospital. The ABI TRS identified the clinical need to implement a new therapeutic model of care (MOC), which would:

- increase therapy intensity
- practise rehabilitation skills in real-life situations
- improve clients’ rehabilitation access in the community
- address clients’ goals of reintegration and independence
- assist with community participation & reintegration

Group-based rehabilitation was identified as a MOC that would address these areas. This included providing a tailored program with contextually relevant activities to address an individual’s goals. Group-based rehabilitation for adults with ABI has been shown to enhance learning experience, motivation and mood through peer interaction as well as being a cost-effective means of utilising health resources and increasing intensity in rehabilitation programs.³ ⁴ Emerging research also identifies that clients perceive group intervention to be beneficial for sharing experiences, reducing isolation, receiving help and feedback, and assisting with adjustment and adaptation to life after TBI.⁵ Having an emphasis on interventions targeting functional ‘real world’ activities in group settings should benefit people following TBI.⁶

Method
The Agency for Clinical Innovation (ACI) framework was used to develop a new model of care i.e., interdisciplinary group rehabilitation.

Results and Conclusion
The new MOC has been successfully integrated into ABI TRS. Consumer and stakeholder feedback has influenced ongoing program development and acknowledged the clinical benefit of the program. Using a service delivery model that involves the whole interdisciplinary team should support the ongoing implementation and translation to clinical practice and the long-term sustainability of the program.

A new service has been developed in a new model of care, to address the needs of people with ABI transitioning from hospital to home. The service uses an interdisciplinary approach, providing group-based rehabilitation in the community, focusing on improving clients' goals of reintegration and independence. The program evaluation includes feedback from consumers and stakeholders, influencing ongoing program development.

References
Online Education Resources for Spinal Cord Injury

Michael Todorovic1,2, Matthew Barton1,2, Steven Bentley3, Jenny Ekberg1,2, James St. John1,2

1. Menzies Health Institute Queensland, Griffith University, Gold Coast, Australia
2. Clem Jones Centre for Neurobiology and Stem Cell Research, Brisbane, Australia
3. Griffith Institute for Drug Discovery, Griffith University, Brisbane, Australia

We conducted a survey of 100 Australians within the SCI community...

Do online resources contain the information you are looking for? **NO (51.2%)**

What online resources are most important for people with SCI?

- **Education Resources**
  - Maintain Health & Wellbeing 22%
  - Body Physiology 4.9%
  - Equipment 11%
  - Community Resources/Services 11%
  - Current Research 17.1%

- **"Your" Injury**
  - 34.1%

- **Carer/Friend**
  - 67.1%
  - 33%

- **Spinal Cord Injury**
  - 57.3%
  - 42.7%

- **Other**
  - Cervical: 58.5%
  - Thoracic: 41.5%

- **Age**
  - 18-29 yrs: 12.2%
  - 30-39 yrs: 24.4%
  - 40-49 yrs: 24.4%
  - 50-59 yrs: 23.2%
  - 60 or older: 15.9%

- **Duration**
  - 1-5 yrs: 43.9%
  - 5-10 yrs: 20.7%
  - >10 yrs: 35.4%

- **Education**
  - Bachelor: 28%
  - High school: 13.4%
  - Tertiary: 32.9%
  - School: 12.2%
  - Post grad: 12.2%

**Points of Interest**
- 51.2% of participants could not find information online.
- Regardless of demographic, education resources on "your" injury are the most important.
- Specifically, treatment options directed at a specific injury-type are needed.

**What's Next?**
- Ask me!
Refugees in Pain: A Retrospective Chart Audit

Henderson S.1,2, Wan A.3, Kluver J.3, Siyambalapitiya S.1,2, Schmidt L.F.,4 Hills R.3
1. Griffith University; 2. Menzies Health Institute QLD; 3. Persistent Pain Clinic, Princess Alexandra Hospital, Metro South Health Service; 4. Refugee Health Service, Metro South Health Service

Introduction

Refugees often have experiences of violent and psychological trauma, leading to prevalence of mental health issues, physical disabilities, and persistent pain.

Management of persistent pain for refugees is complex as a result of cultural interpretations of pain, language barriers and distrust of formal health systems. In some cultures, persistent pain is considered to be a natural part of life, resulting in failure to seek treatment.

Clinicians at the Persistent Pain Clinic at Princess Alexandra Hospital sought to identify factors impacting on refugees using this service adequately.

Aim

The aim of our study was to conduct a retrospective chart audit to gain an understanding of the characteristics of the refugee population attending the Persistent Pain Clinic at Princess Alexandra Hospital as a first step towards providing optimal pain management to refugees.

Methods

Retrospective chart audit of medical records of refugees who attended the clinic was conducted (2015-2017).

- Adapted Minnesota Complexity Assessment Method (MCAM) used.

Data Collected

- Demographics
- DASS21 & (Brief Pain Inventory) BPI pre and post scores
- Attendance details: No of appointments booked, No of Appointments kept, Failure to Attend (FNA)
- Discharge Information
- Information about Pain & Trauma
- Social context family & networks

Sample

N=37 (17 M, 20 F); Mean age: 44.7 years

![Birth Countries of Refugee Sample](Image)

Mean years in Australia: 9.4 years
- 29 patients required use of an interpreter.
- 8 Patients had functional use of English.

![Home Language](Image)

Work status: 18 unemployed, 3 working, 7 on Disability Support Program, 2 home duties, 4 studying English.

Results

Pain Duration: Mean 8.6 years

Most patients had PTSD (n=22) or PTSD Symptoms (n=11), often contributing to pain.

MCAM Measures:
- Very high levels of pain interfering with functioning;
- Very high levels of distress, distraction, preoccupation;
- Less than half displayed readiness to engage in treatment options;
- Almost all patients had restricted participation in social networks; modal interference in 2 domains.

Self-report measures:
- About two thirds completed questionnaires of intake & one third completed at end of treatment.

Discharges:
- 27 patients discharged
- 8 patients improved with treatment, especially in functioning.
- Other discharges equally split into active and passive withdrawal by patient, and doctor initiated discharge.

Issues that may lead to less than optimum pain management:
- Very high levels of pain, distress, and interference with functioning in refugee patients.
- Communication difficulties with clinicians despite use of interpreters.
- Inability to use functional English is a significant factor affecting attendance, completion of self-report measures and engagement in group programs.
- Cultural issues, including differing cultural views of pain, may interfere with patient treatment.

![Language Use and Attendance](Image)

References


Acknowledgement

This chart audit was supported by The Hopkins Centre Seeding Grant Funding for Translational Research; Research for Rehabilitation and Resilience, Princess Alexandra Hospital.

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Energy needs in people with spinal cord injury undergoing surgical repair of chronic pressure injuries

Amy Nevin1,2,4, Susan Urquhart5, Sridhar Atresh2,3,4 Timothy Geraghty2,3, Elizabeth Walter5, Angela Vivanti1,5, Leigh Ward6, Ingrid Hickman1,7

1Department of Nutrition and Dietetics, Princess Alexandra Hospital; 2The Hopkins Centre – Research for Rehabilitation and Resilience, Metro South Health and Griffith University; 3Spinal Injuries Unit, Division of Rehabilitation, Princess Alexandra Hospital; 4Faculty of Medicine, University of Queensland; 5School of Human Movement and Nutrition Sciences, University of Queensland; 6School of Chemistry and Molecular Biosciences, University of Queensland; 7Mater Research Institute, University of Queensland

Introduction

Consensus Dietetic guidelines for people with spinal cord injury (SCI) who have pressure injuries (PIs) recommend an increased energy intake to meet demands of wound healing, but evidence to support this is limited. The impact of PI surgical repair on energy needs is also unknown.

Purpose

The aims of this study were to:
1. Determine the resting energy needs of people with SCI admitted to hospital with established chronic PIs
2. Investigate the energy burden of PI surgical repair and healing.
3. Observe weight change from pre-surgery until discharge from hospital.

Methods

Data was collected at the bedside following an overnight fast of ≥8 hours. Resting energy expenditure (REE) was measured using indirect calorimetry (canopy hood) for ≥20 minutes one week pre-surgery, one week post-surgery, on removal of stitches and on return to unrestricted wheelchair sitting with complete wound healing (at discharge). Weight was measured one week pre-surgery and at discharge. Clinical energy prescription during admission was based on measured REE.

Conclusions

The presence of chronic PIs, surgical repair or subsequent healing do not appear to impact the energy needs of people with SCI and weight gain is common. Current guidelines recommending increased energy prescription conflict with these findings and may increase risk of unintentional weight gain over the course of healing.

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Understanding patient engagement in outpatient cardiac rehabilitation programs using the Model of Therapeutic Engagement

Sepideh Jahandideh, Elizabeth Kendall, Samantha Low-Choy, Kenneth Donald, Rohan Jayasinghe
Menzies Health Institute Queensland, Griffith University

Method

- Participants: 217 patients eligible for outpatient cardiac rehabilitation programs from the Cardiology ward at Gold Coast University Hospital
- Design of this study: a prospective study
- Data collection: paper surveys over three time intervals to cover all stages of the Model of Therapeutic Engagement
- Statistical method: Structural equation modeling to test the hypotheses proposed
- Nine hypotheses have been tested based on the Model of Therapeutic Engagement in Rehabilitation (significant relationships showed in red).

Results

Path analysis was carried out by ascertaining the significance of path (p-value).

- For stage 1, results revealed significant relationships between perceived need and patient intention to engage in CR programs ($\beta = 0.43; B = 0.63; SE = 0.29; p = 0.03$), and perceived self-efficacy and patient intention to engage in the CR program ($\beta = 0.23; B = 0.24; SE = 0.13; p = 0.05$).

- For stages 2 and 3, there were significant relationships between:
  - Intention to engage in CR program and CR preparation ($\beta = 0.37; B = 0.28; SE = 0.09; p = 0.001$), CR Initiation and Engagement ($\beta = 0.68; B = 0.20; SE = 0.03; p \leq 0.001$); Engagement and Analysis of Experience ($\beta = 0.38; B = 0.08; p \leq 0.001$); Analysis of Experience and Maintenance ($\beta = -0.58; B = -0.30; SE = 0.05; p \leq 0.001$); and Maintenance and Engagement ($\beta = 0.20; B = 0.56; SE = 0.29; p = 0.05$).

Expected Results

The Model of Therapeutic Engagement moves the field toward a more complete understanding of the process of engagement in cardiac rehabilitation programs and contributes knowledge about how to make this process more effective. It is suggested to embed this model into a broader socio-environmental context to have a comprehensive understanding of patient engagement in cardiac rehabilitation programs.

Literature Cited


Disclosure

Financial support for this study was received from Griffith University.

Conflict of interest The authors declare no conflict of interest for this study.
Introduction

- Self-rated health is an established marker of general health status and predicts long term morbidity and mortality.
- After Motor Vehicle Accidents, patients experience challenges with self-efficacy and return to work that extend well beyond the immediate time frame of discharge from acute health care.
- There is an abundance of literature on health during Acute Care or while receiving Rehabilitation after trauma. Information on long-term health status after Motor Vehicle Accidents for patients within the community, especially in Australia however is limited.
- In this context, we examined factors associated with general health status after Motor Vehicle Accidents from a survey of community dwelling Compulsory Third Party (CTP) insurance claimants in Queensland, Australia.

Purpose

Examine the association between demographic, injury related variables and general health status as assessed by responses to questions based on dimensions (i.e. health states) of the EuroQol-5D (EQ-5D).

Methods

Study Population

- Participants from a survey conducted by the Centre of National Research on Disability and Rehabilitation Medicine (CONROD), Griffith University and the Motor Accident Insurance Commission (MAIC).
- Mail-in-surveys distributed to a random sample of 2500 claimants. Criteria for inclusion: age ≥18 years, completion of claim payments and non-critical injuries (fatalities, AIS categories 5 & 6 excluded).
- Responses were returned within 2 months, a draw for a $500 shopping voucher was provided as incentive.
- A total of 426 responses (17%) were received. Those with unusable information (n=216), missing IDs (n=1) and non-specific injuries (n=4) were additionally excluded. Final analytic sample comprised of 205 participants (8.2%).

Methods

Study Variables

- We examined age, gender, marital status, country of birth, spoken language, education level, employment status, income, injury severity, time since injury and whether participants received funded rehabilitation care.

Outcomes: EuroQol-5D (EQ-5D) Health States

- Responses to the 3-level EQ-5D health states including: Mobility, Self-Care, Usual Activities, Pain or Discomfort and Anxiety or Depression.

Results

Table 1. Descriptive characteristics of the sample

<table>
<thead>
<tr>
<th>Variables</th>
<th>Sample (N=205)</th>
<th>Mean ± SD or N (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>48.4 ± 14.2</td>
<td></td>
</tr>
<tr>
<td>Gender</td>
<td>Men</td>
<td>78 (38.1%)</td>
</tr>
<tr>
<td></td>
<td>Women</td>
<td>127 (61.9%)</td>
</tr>
<tr>
<td>Marital Status</td>
<td>Single/Never Married</td>
<td>33 (16.1%)</td>
</tr>
<tr>
<td></td>
<td>Married/De Facto</td>
<td>129 (62.9%)</td>
</tr>
<tr>
<td></td>
<td>Separated/Divorced/Widowed</td>
<td>43 (21.0%)</td>
</tr>
<tr>
<td>Spoken Language</td>
<td>From English speaking country</td>
<td>180 (87.8%)</td>
</tr>
<tr>
<td></td>
<td>From Non-English speaking country</td>
<td>23 (11.2%)</td>
</tr>
<tr>
<td>Employment status</td>
<td>Unemployed</td>
<td>81 (39.5%)</td>
</tr>
<tr>
<td></td>
<td>Employed</td>
<td>122 (59.5%)</td>
</tr>
<tr>
<td>Abbreviated Injury Scale categories</td>
<td>Motor injuries</td>
<td>160 (78.0%)</td>
</tr>
<tr>
<td></td>
<td>Moderate injuries</td>
<td>33 (16.1%)</td>
</tr>
<tr>
<td></td>
<td>Severe injuries</td>
<td>12 (5.9%)</td>
</tr>
<tr>
<td>Time since Accident, years</td>
<td>3.4 ± 1.3</td>
<td></td>
</tr>
<tr>
<td>Received Rehabilitation Care</td>
<td>Yes</td>
<td>117 (57.1%)</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>71 (34.6%)</td>
</tr>
<tr>
<td>Mobility</td>
<td>No Problems with Mobility</td>
<td>137 (66.8%)</td>
</tr>
<tr>
<td></td>
<td>Have problems with Mobility</td>
<td>68 (33.2%)</td>
</tr>
<tr>
<td>Self-Care</td>
<td>No Problems with Self-Care</td>
<td>166 (81.0%)</td>
</tr>
<tr>
<td></td>
<td>Have problems with Self-Care</td>
<td>37 (18.1%)</td>
</tr>
<tr>
<td>Usual Activities</td>
<td>No Problems with Usual Activities</td>
<td>86 (41.9%)</td>
</tr>
<tr>
<td></td>
<td>Have problems with Usual Activities</td>
<td>118 (57.6%)</td>
</tr>
<tr>
<td>Pain or Discomfort</td>
<td>No Pain or Discomfort</td>
<td>51 (24.9%)</td>
</tr>
<tr>
<td></td>
<td>Have Moderate to Extreme Pain or Discomfort</td>
<td>152 (74.1%)</td>
</tr>
<tr>
<td>Anxiety or Depression</td>
<td>No Anxiety or Depression</td>
<td>119 (58.0%)</td>
</tr>
<tr>
<td></td>
<td>Have Anxiety or Depression</td>
<td>86 (41.9%)</td>
</tr>
</tbody>
</table>

The Hopkins Centre

Research for Rehabilitation and Resilience

Health Status after Motor Vehicle Accidents: Results from a survey in Queensland

Srinivas Teppala, Sanjoti Parekh, Elizabeth Kendall

The Hopkins Centre, Menzies Health Institute, Griffith University

Conclusion

- Gender, country of birth, employment status and time since injury were associated with individual health states of the EQ-5D.
- Employment and time since injury were consistently associated with overall health status (EQ-5D index score).
- Future prospective studies with larger representative samples are required to confirm our findings.
Housing for people with an acquired brain or spinal injury: Mapping the Australian funding landscape

Dr Courtney Wright, Dr Jacinta Colley, Ms Kate Knudsen, & Prof Elizabeth Kendall

The Hopkins Centre, Menzies Health Institute Queensland, Griffith University

**Background**
Due to several personal and systemic factors, individuals with an acquired brain injury (ABI) or spinal cord injury (SCI) are often unable to afford the extent of support required to address their housing needs. Funding is often sought from National or State/Territory-based insurance schemes to assist a person with ABI or SCI to make changes to their housing situation and improve their quality of life.

**Research Aim**
Given the crucial role of insurance schemes in contributing toward housing improvements for individuals with ABI and SCI, this research aimed to synthesise eligibility and funding contributions relating to housing infrastructure, tenancy, and service delivery for people who have sustained an ABI or SCI for the following schemes:
- National Disability Insurance Scheme (NDIS);
- Interstate National Injury Insurance Schemes (NIIS); and
- National and interstate workers compensation schemes.

**Results**
This review identified 20 major insurance schemes across Australia that provide housing-related funding to individuals who have sustained a moderate, severe, or profound ABI, or SCI resulting in incomplete or complete paraplegia or tetraplegia. The funding contributions of the 20 relevant schemes toward three main categories of housing support (i.e., infrastructure, tenancy, and service delivery) following ABI or SCI are outlined below:

**Major Australian Funding Schemes**

<table>
<thead>
<tr>
<th>National Schemes</th>
</tr>
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<tbody>
<tr>
<td>NDIS</td>
</tr>
<tr>
<td>Seacare</td>
</tr>
<tr>
<td>Comcare</td>
</tr>
<tr>
<td>DVA Scheme</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>State/Territory-based Schemes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Queensland</td>
</tr>
<tr>
<td>New South Wales</td>
</tr>
<tr>
<td>Australian Capital Territory</td>
</tr>
<tr>
<td>Victoria</td>
</tr>
<tr>
<td>Tasmania</td>
</tr>
<tr>
<td>Northern Territory</td>
</tr>
<tr>
<td>South Australia</td>
</tr>
<tr>
<td>Western Australia</td>
</tr>
</tbody>
</table>

**Implications for Consumers**
The findings highlighted the extent of missing information regarding funded housing supports across the NDIS, NIIS, and workers compensation schemes. Individuals with an ABI or SCI, and their families, may use this information to inquire about particular housing supports they may need to live as independently as possible at home and participate in the community, and that have not been mentioned in scheme guidelines, fact sheets, or other public documents. The review findings may therefore empower individuals and their families to pursue information about housing supports they may be entitled to, in order to improve their housing situation.

**Implications for Policy-makers**
The review identified inconsistencies across the 20 schemes regarding funded housing supports and eligibility criteria to access particular housing supports. This inconsistency raises equity issues that must be considered by policy-makers, particularly if individuals with ABI or SCI are unable to access the housing support they require from other government or non-government schemes. The development and communication of a unified, evidence-based framework for housing support that is tailored to ABI and SCI may inform future decision-making in this area and improve housing outcomes for consumers.
Problem

After a life-changing injury or illness, people often need to modify their home or find an alternative place to live. This is because their previous home is no longer suitable or equipped for their high level physical, mental, or support needs [1-4]. Despite the clear nature and extent of the problem (1.3 million Australians have a severe/profound disability requiring daily support) [4], many people remain in unsuitable living environments for years or are moved to institutional group homes or nursing facilities in an attempt to ‘solve’ the problem [2,5].

While there are some housing resources currently available to help consumers modify their homes, they are limited in scope (i.e., focused solely on physical accessibility design) and substance (i.e., static information mediums: websites, Microsoft Word© documents, or PDF documents simply listing information).

There is no resource currently available to systematically and comprehensively assist consumers in designing or re-designing their own home.

Solution

The “My Accessible Home” resource (see https://www.myaccessiblehome.com.au/) is a comprehensive, online decision tool (with the potential of being a ‘one-stop-shop’) for people with disability and their families to access relevant housing information and plan for the future.

Importantly, the information provided includes, but is not limited to, physical access considerations. In contrast to other housing resources currently available, the My Accessible Home tool prompts consumers to not only think about functional design, but to also think more broadly in terms of what they would like their home to include, and where and how they would like to live.

For the first time, consumers are able to navigate through a comprehensive and detailed resource, watch informative videos, browse pictures, and collate (save) their ideas in a virtual space representing ‘My Dream Home’. The resultant report may then be shared with individuals in a position to help the person improve their housing situation (e.g., friends, family, advocates, architects, designers, builders, occupational therapists) via print form, email, or social media. Importantly, the information comprising the technological platform is evidence-based and informed by consumers’ voices.

Impact

Traditionally, consumers have not had the financial capacity to build, buy, or modify their own home unless they had been able to access compensation, inheritance, or grant funding. This situation changed in 2016, when the National Disability Insurance Scheme (NDIS) and National Injury Insurance Scheme (NIIS) commenced in Queensland (and has also been rolled-out across Australia).

The individualised funding packages provided to consumers through these schemes have enabled consumers to invest in their housing situation, subsequently compelling the sector to respond to a new consumer market. The user-friendly, interactive nature, and comprehensive information provided by the My Accessible Home resource will empower consumers to make important decisions in early design (or purchase) processes.

It is anticipated that the resource will improve housing suitability and liveability for people with complex disability, reducing the number of forced relocations of people into group homes and nursing facilities.

References


Acknowledgements

The My Accessible Home resource was produced by a collaborative partnership between The Hopkins Centre (Menzies Health Institute Queensland, Griffith University) and Youngcare. The original idea for the resource was conceived by Dr Courtney Wright.

Dr Wright’s PhD thesis informed the resource content. The partnership sincerely thanks Elizabeth Ainsworth (Occupational Therapist; Access Consultant) for her additional intellectual contributions and Winning Media for developing the website. The partnership are also grateful for the financial contributions of Perpetual through the 2016 IMPACT Philanthropy Program and acknowledge the Recovery Injury Research Centre (The University of Queensland) for their support.

Dr Wright’s PhD thesis (cited below) was supervised by Prof Heidi Zeeman and Prof Elizabeth Kendall: Wright, C. (2016). Likes, dislikes, must-haves, and must-nots: An exploratory study into the housing preferences of adults with neurological disability (Doctoral dissertation). Brisbane, Queensland: Griffith University.

The My Accessible Home resource would not have been possible without the people who shared their stories so willingly in the research. Dr Wright’s deepest thanks are extended to the research participants who gave so generously of their time.
This study aimed to investigate the perceived costs and benefits to Australian society if all new homes were built to an agreed Universal Design ('Livable Housing Design') standard. Uptake of Universal Housing Design has been met with resistance to implement by the housing and construction sector. This has resulted in an agreed Universal Housing Design standard by 2020 [1]. Owing to a perceived lack of consumer demand, however, the voluntary industry-led voluntary approach over 10 years, a national guideline, and a strategic plan with the aspirational goal that “all new homes will be of an agreed Universal Housing Design standard by 2020” [1].

In response to persistent advocacy from people with disability and their supporters, the Australian Government established the National Dialogue on Universal Housing Design (NDUHD) in 2009, to address the lack of inclusive housing in Australia. NDUHD argued for an overarching themes and limit interpretation bias.

The results also highlighted the perceived need for a comprehensive education and awareness strategy to accompany regulation, so that all stakeholders understand why the regulation of Livable Housing Design is important for them, their businesses, and for a more inclusive Australian society.

Method
An online mixed-methods survey was available for completion over a three month period. Convenience and snowball sampling strategies were used to recruit a total of 1,329 participants. The sample predominately reflected home owners and people whose housing needs are not met through the current mainstream housing system.

Quantitative component:
Participants were asked to indicate their perspectives on how (1) costly and (2) beneficial it would be to Australian society if all new homes were built to Livable Housing Design standards, using a five point Likert scale (i.e., no cost to massive cost; no benefit to massive benefit). Data was analysed according to frequency counts and the percentage of participants who endorsed each possible response.

Qualitative component:
Participants were asked to explain their reason/s for their multiple choice response. Data was thematically analysed using a text analysis software package, Leximancer (Version 4.5, 2016), to identify a full list of emerging concepts. Following the software-supported text analysis, a manual thematic analysis was completed by two researchers to develop overarching themes and limit interpretation bias.

Quantitative Findings

**Perceived costs to Australian society**

<table>
<thead>
<tr>
<th>Percentage of Participants (N=1,054)</th>
<th>No cost</th>
<th>Minimal cost</th>
<th>Some cost</th>
<th>Significant cost</th>
<th>Massive cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants (%)</td>
<td>60%</td>
<td>35.39%</td>
<td>35.29%</td>
<td>12.43%</td>
<td>3.42%</td>
</tr>
</tbody>
</table>

**Perceived benefits to Australian society**

<table>
<thead>
<tr>
<th>Percentage of Participants (N=1,054)</th>
<th>No benefit</th>
<th>Minimal benefit</th>
<th>Some benefit</th>
<th>Significant benefit</th>
<th>Massive benefit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants (%)</td>
<td>60%</td>
<td>1.23%</td>
<td>5.79%</td>
<td>37.57%</td>
<td>54.93%</td>
</tr>
</tbody>
</table>

Qualitative Findings

**Theme 1: Right to choice, autonomy and participation**

The majority of participants were in favour of mandating Livable Housing Design standards. Participants believed that mandating Livable Housing Design in all new builds would ultimately promote the physical, psychosocial, and emotional health and quality of life of individuals and families. These benefits were perceived to relate to all Australians (either now or in the future) since “we are all ageing as well as vulnerable to illness, accidents or injuries which could lead to incapacity of varying degrees at any age or stage of life” (Response 0674).

A smaller number of participants suggested that mandating Livable Housing Design would infringe on peoples’ right to choose their housing design and features: “Socially boring!!! ... Lack of human-rights for individuals to choose to live in the built environment they prefer” (Response 0036).

**Theme 2: Perceived financial impact of change**

Participants’ perceptions regarding the financial impact of mandating Livable Housing Design standards varied. Several participants believed that implementing livable design features following regulation would cost little or no more than current housing options since changes would be made during the design stage. Others believed that implementation would cost a lot more than current housing designs. As one person suggested, more space is needed regardless and “space costs money” (Response 0369). Others noted the initial cost of changing from current designs and learning how to do things differently (Response 0874). However, most participants believed that mandating Livable Housing Design standards would come with an initial cost, but that the cost would be outweighed by social benefits and/or recovered over time.

Conclusion

The majority of participants considered mandating Livable Housing Design in all new builds a cost-effective venture for Australia’s housing future. The research findings indicated that any cost-benefit exercise must go beyond the immediate impacts on the housing industry and the housing market, to consider the public interest in Australia’s housing infrastructure in the long-term, and the significant Government investment to assist people to be socially and economically included and to participate in family and community life.

The results also highlighted the perceived need for a comprehensive education and awareness strategy to accompany regulation, so that all stakeholders understand why the regulation of Livable Housing Design is important for them, their businesses, and for a more inclusive Australian society.

References


Acknowledgements

This research project was conceived and conducted by the Australian Network for Universal Housing Design (ANUHD). The Australian Building Codes Board also assisted with participant recruitment. Two Research Fellows from The Hopkins Centre, Griffith University analysed the data. The project team are sincerely grateful to the many organisations and individuals who forwarded the online survey to others in their networks.