Partnering in healthcare

A framework for better care and outcomes
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Foreword

We know better health outcomes are achieved when health professionals and services work in partnership with consumers, patients, carers and communities. Of course, some health services already do this, and do it well. But the focus on person-centred care is not consistent across the state – leading to great variation in consumer participation, experience and outcomes.

Consumers play a vital role in helping us to avoid making the same mistakes repeatedly. Hindsight shows that in nearly every instance of avoidable harm, consumer letters, emails and phone calls have attempted to warn us about system failures that contributed to that harm.

This document outlines a new partnership and consumer participation approach to drive positive change in healthcare.

As part of our methodology, we asked people what was most important to them. The Partnering in healthcare framework is the result.

This new framework aims to:

- bring consistency to how Victorians can participate in their own healthcare
- help health services involve consumers to deliver care that is safe, effective, person- and family-centred, equitable and clinically effective
- clearly describe consumer priorities for health services, Safer Care Victoria (SCV) and the Department of Health and Human Services (the department).

Partnering with consumers is a key driver behind our mission of outstanding healthcare for all Victorians. We work with health services to improve opportunities for consumers and community representatives to participate in quality and safety improvement activities.

We have two key priorities that guide our work. The first is to ensure that consumer voices and choices are central to their own care, and that consumer-defined outcome measures and improvement goals are being delivered at a health service level and within SCV.

The second is to ensure that consumer voices and experiences drive health service and health system improvement and improve people’s experiences and outcomes.

We are proud to have used a co-design approach to develop this framework. For the first time, we employed a consumer lead as part of our team, to ensure consumer voices were central in our work, and a sector lead to keep us connected with health services.

Thank you

The Partnering in healthcare framework could not have been developed without the participation of many consumers, health professionals and community members.

We will continue to work with you as we support the implementation of this framework across all public hospitals in the first half of 2019. We want you to inform a meaningful and easy way to measure our collective impact on improving consumer partnerships. The first step in this process will be a Partnering in healthcare forum in April 2019.

We look forward to you joining us.

Louise McKinlay
Director Consumers as Partners
Acknowledgements

Safer Care Victoria would like to acknowledge the contribution of the following individuals and groups who provided input in developing the Partnering in healthcare framework:

- Individual consumers and Voice at the Table for their valuable contribution, and making the priorities summit an inclusive consumer event.
- SCV Patient and Family Council members, Victorian health service staff, members of the Patient Experience Network, the Centre for Health Communication and Participation, the Centre for Culture Ethnicity and Health, and staff from the Department of Health and Human Services (the department), for providing feedback on the draft framework.
- BehaviourWorks Australia at Monash University, for their guidance and expertise in prioritisation methodology.
- SCV staff members – Louise McKinlay, Lidia Horvat, Belinda MacLeod-Smith, Kylie Foltin, Joanna Williams, Karen Hill, Gemma Cooper and Erin Pelly for their commitment and collaboration in implementing the engagement strategy, and the iterative development of the framework.
We consulted far and wide to develop the *Partnering in healthcare* framework. As it is rolled out to all public health services in Victoria, we hope you feel encouraged and supported to open up meaningful opportunities for consumers to partner in their healthcare.

**Helping you go the extra mile**

As you know, health services are obliged to meet key requirements for partnering with consumers under the Australian Commission on Safety and Quality in Health Care (ACSQHC) Standards. Achieving accreditation is a baseline measure for *Partnering in healthcare*.

This framework is designed for those health services that want to go beyond what is required. It describes suggested priorities and actions health services can take, and what SCV and the department will do, to deliver outstanding healthcare for Victorians. You can use it to enable innovation and identify future areas of work and improvement.

This framework will help you respond to the needs and expectations of Victorian consumers. We know this because we asked hundreds of people what was most important to them, and this framework is the result. For more about how we developed this framework, go to page 35.

**Rolling out the framework**

At a minimum, please identify two domains and the priorities you will choose to focus on in the next 12 months.

By **June 30 2019**, you need to:

- use the self-assessment tool
- complete the statement of intent outlining your two chosen domains and identified priorities.

We developed resources to support you to implement this framework. These are available at bettersafercare.vic.gov.au.

We will also hold a forum on **29-30 April 2019** to:

- celebrate what you are currently doing to progress consumers partnering in their healthcare
- share good practice examples
- generate new thinking and ideas
- identify the most useful ways to measure and report on progress with *Partnering in healthcare*
- identify how we can measure improvements in participation.

**This framework applies to all Victorian public health services**

This framework replaces *Doing it with us not for us: Strategic Direction 2010–2013* and the *Cultural Responsiveness Framework: guidelines for Victorian health services*. It states the expectations Victorians have about how to improve partnering with consumers to achieve better outcomes.

This is a living document and will be updated as we progress

To ask questions or provide feedback, please email: partnering@safercare.vic.gov.au
Partnering in healthcare has a clear and specific purpose:

To support practical strategies and partnerships, including people partnering in their own healthcare, in order to deliver higher quality care that is safe, person- and family-centred, equitable and clinically effective.

For more context, go to page 34.

Other services and sectors may learn from this

While this framework is designed for public health services, it may be relevant for:

- private hospitals
- ambulance services
- funded health services (such as community health services, mental health community services, alcohol and drug services and public residential aged care services)
- regulated health services such as Aboriginal health organisations
- primary health networks.

How to read this document

This document is structured around the five domains, to help you decide which are your immediate priorities for implementation. Each chapter clearly outlines:

- what consumers said
- what the domain means
- what matters in practice.

Comments from people who filled out our online survey are included throughout this document.

A note on terminology

We broadly use the term ‘consumer’ to refer to people, families, carers and communities who are current or potential users of health care services. This includes children, women and men, people living with a disability, people of diverse cultural, linguistic and religious experiences, socioeconomic status and social circumstances, sexual orientations, sexes, genders and gender identities, health and illness conditions.

The term also includes: people who choose to get involved in decision-making, health consumer representatives who provide advice on behalf of consumers, with the overall aim of improving healthcare, and carers who often have an important role in healthcare decision making and care giving.

Different health settings may use terms such as: patients, persons and families, carers, clients and residents. In this paper, ‘consumer’ includes these definitions.

See our glossary for more definitions. (page 37).
Navigating the framework

A summary of the Partnering in healthcare consultation

For Partnering in healthcare, we used a five-stage prioritisation process. Stages included desktop research, forming a project team that included consumer and sector leads, a statewide online survey that reached 180,000 Victorians, data analysis of more than 3,000 responses, and a face-to-face ‘priorities summit’ based on Cochrane Review prioritisation methodology. To elevate the consumer voice, the priorities summit deliberately included a 2:1 ratio of consumers to health sector representatives.

How many people did we reach?

180,000 people
reach through social media and other online networks

3,145 visits
to consultation web page

680 surveys
3,000+ responses

2:1 ratio
health consumers to health service reps at priorities summit

We need to ask patients and the people who support them what they want.

Who did we hear from?

74% female
10% speak a language other than English
26% frequent hospital users
39% have children at home
76% employed

58% under 54 years old
71% have worked in healthcare
32% identify as a carer
24% live outside Melbourne
74% educated to Bachelor’s degree or higher
The Partnering in healthcare framework consists of five domains. These represent collective focus areas, where work could improve consumer experience and outcomes.

Focusing on five domains for healthcare improvement
Each domain is underpinned by evidence and best practice and informed by extensive consultation with consumers and health services.

Using the domains as a reference point will help you identify where improvements may bring real benefits and impact, and therefore, where time and effort is best invested.

Figure 1: Five domains of Partnering in healthcare
Identifying priorities and actions for health services

We asked consumers and health professionals to identify issues important to them. These priority areas are where SCV, the Department and health services can make the most difference.

Under these priorities, we encourage health services to identify actions. The framework has been built to further strengthen and build on current initiatives and identify future areas of work and improvement. We recommend adapting and implementing these priorities in line with your existing initiatives, to best meet the needs of your consumers and organisation.

How consumers can participate

There are many opportunities for consumers to be meaningfully involved at all three levels of healthcare: the direct care level, the service level and the system level. The figure opposite highlights how consumers can participate at each level.
**Figure 2: Partnering in healthcare framework overview**

**Personalised and holistic**
- Individualised and connected care
- Compassion and respect

**Working together**
- Care is co-designed with patients, families and clinicians
- Coordination and continuity of care

**Shared decision-making**
- Use of decision aids, decision support coaching
- Increase use of patient reported outcome measures (PROMS), patient reported experience measures (PREMS)

**Equity and inclusion**
- Patient reported language services provision
- Cultural safety, diversity of consumer participation

**Effective communication**
- Respectful communication, health literacy, Ask Me 3, Teach Back

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**Direct care level**
- ‘I am respected and receive personalised care that treats and supports me as a whole person.’

**Service level**
- ‘I am included as a respected partner in learning about and improving healthcare.’

**System level**
- Consumers, carers, and communities participate in system-wide quality and safety improvement in healthcare organisations and the Department.
- Healthcare organisations partner with consumers in governance, planning and policy development, including through hospital boards and clinical governance.
- Government organisations partner with consumers in governance, planning and policy development.

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**Direct care**
- Consumers participate in their own healthcare, treatment, as do their family and carers.
- Health service enables and support consumers to be equal partners in their care, including through shared decision-making.

**Service**
- Consumers participate in service design, delivery and quality improvement.
- Health services partner with consumers in decisions about the design and delivery of services, including through community advisory committees, working groups, quality and safety committees, and service design working groups.

**System**
- Consumers provide feedback, ideas and personal experiences to drive change.
- ‘I receive personalised care where my healthcare holistically responds to my unique characteristics, cultures, beliefs and experiences – and those of my family and carers.’

**Accountability drivers**
- Health service strategic plan
- Health service quality improvement plan, policies and monitoring frameworks
- Victorian health services performance monitoring framework
- Statement of priorities
- Delivering high-quality healthcare: Victorian clinical governance framework
- Department policies
- Victorian Healthcare Experience Survey
- National Safety and Quality Health Service Standards (second edition)
1 Personalised and holistic

This domain includes what people can do for their own health and wellbeing, as well as what is done with and for people in a healthcare setting. This domain pivots on considering the whole person (or family). This means understanding their physical, cultural and social context, to identify what makes a difference to a person’s health, wellbeing and safety. It also recognises that people have agency when they are at the ‘centre’ of care being provided to them by healthcare professionals.

WHAT CONSUMERS SAID

- Actively engage consumers in partnership opportunities across all aspects of the healthcare system.
- Implement a statewide consumer network.
- Promote and support person-centred co-design as a method for improvement.
- Build empowering relationships.

Elements of this domain

Personalisation ‘by’ a person: how people shape their own lives, health and wellbeing.

Personalisation ‘for’ a person: practices in healthcare that treat an individual as a person and put people and families at the centre of – not fitting them into – services.

Individualised, connected and integrated care.

Compassion and respect.

Consumer-reported experience and outcome measures.

A broader perspective on illness, health and wellbeing.

Personalised health and wellness plans and networks.

Respecting people’s lived experience and values.

The whole person in their physical, social, cultural and linguistic context.

Taking into account people’s preferences and expressed needs.

Education and support for self-care.

Emotional support and empathy.

Involving family and friends.
'It is good if staff introduce themselves by name, and have a short chat, before getting down to the medical side of things. Consumers are worried, may be in pain, and are unwell. However, staff don’t seem to understand this, as it is normal for them to see people in this state. For most consumers, the hospital is abnormal, and outside their normal experience. It is a new world to be navigated. If the people they meet do not seem to care, and treat it as no-nonsense routine, it can be unbearable.'

WHAT MATTERS IN PRACTICE

**SCV**

**Priorities**
Deliver practical ways to improve person-centred care.

Enhance the utility of existing patient-reported experience measures and ensure they are publicly reported.

Work with consumers, health professionals and the Victorian Agency for Health Information (VAHI) to establish an agreed approach to collating patient-reported outcome measures (PROMS) and revised patient-reported experience measures (PREMS).

Promote and support the use of person-centred co-design as a method for improvement.

**Actions**
Implement a statewide consumer participation strategy (this Partnering in healthcare framework).

Support health services to recruit, develop and support consumer leaders.

Develop the Partnering in healthcare governance structure through the SCV Patient and Family Council.

**Department**

**Priorities**
Understand the evolving needs of patients, clients and victim survivors, by hearing their lived experiences and acknowledging their diverse needs when designing and delivering services to ensure people can access the support they need, when they need it.

Advance Aboriginal self-determination in health and human services by using Aboriginal-determined measures of success in health and wellbeing outcomes.

Improve use of patient- and client-reported experiences of care and treatment.

**Actions**
Improve integrated and coordinated care between health and community services, to support better outcomes for people with chronic and complex needs.

Implement the Korin Korin Balit-Djak: Aboriginal health, wellbeing and safety strategic plan 2017–2027.

Implement the Disability action plan 2018–2020.
**Health service**

**Suggested priorities**
- Treat consumers as a whole person and provide more individualised care.
- Provide hospital staff training on person-centred care.
- Optimise the time with consumers to understand their needs.
- Include family and carers in care planning.
- Improve access to care coordinators, pastoral care, counsellors and social workers to support people.
- Build a support network of consumers.
- Foster, promote and implement compassionate care strategies.
- Investigate the use of the Patient Activation Measure (PAM) to support consumers to engage in their healthcare.
- Provide education for consumers and healthcare professionals about the implementation of the Australian Charter of Healthcare Rights in Victoria.
- Support the unique characteristics and needs of the individual in the co-construction of the care process, from diagnosis to discharge.

**Actions**
- Identify at least two domains (and which priorities within them) you will focus on by **30 June 2019**.
- Use the self-assessment tool to map your health service’s current activities against the domains and priorities.
- Complete the statement of intent and share with SCV.
- Evidence for NSQHS Standards is a baseline measure for *Partnering in healthcare*.
- The *Partnering in healthcare* and NSQHS Standards resource highlights how the framework connects to the standards.

‘Reframe the conversation on the person, not the disease.’
Consultation

Other suggestions for improvement from our consultation

Ask consumers about what is most important to them.

Understand a consumer’s individual needs.

Build conversations with consumers.

Involve consumers’ families and carers in discussions and decisions, particularly when the consumer is vulnerable, or has language or cognitive difficulties.

Respond to an individual’s preferences for care.

Acknowledge and act on consumers’ and families’ knowledge and experience of their health and condition.

Include consumers’ understanding of their own condition and care as an integral component of daily ward rounds.

Address the isolation and lack of social supports experienced by some consumers.

Design healthcare systems to work for people.

‘Listen respectfully to consumers. Acknowledge that consumers live in their own skin, and experience their health condition 24/7, not just for 20 minutes when they see a specialist. Be prepared to adjust treatments based on consumer experience.’
Personalised and holistic care is possible when people work together in strong teams, partnerships and share knowledge. Knowledge transfer is a two-way street. Teams of health professionals should be connected and well informed about diagnostic techniques, the causes of disease, prognosis and health strategies. The consumer knows about his or her experience of the illness, social circumstances, values and culture. Without exchanging this information, the knowledge of each party may be limited, and the resulting care may be compromised.

**WHAT CONSUMERS SAID**

- Co-develop improvement and innovation activities.
- Co-develop PREMS and PROMS.
- Increase co-production opportunities.
- Include consumers in staff training.
- Build cultures of inclusion, trust and support.
- Provide capability assessment and support for boards and senior staff on consumer engagement.
- Provide flexible and diverse participation and feedback opportunities (variation in times and types of opportunities).
- Give adequate induction for consumers joining committees, and ongoing support to check how things are going.
- Fill consumer opportunities based on interests, preferences, experience, and suitability – ‘It should not be a matter of just filling a vacancy’.
- Support adequate handover time between staff to continue work with consumers.

**Elements of this domain**

High-quality relationships and interactions are at the core of people’s health and wellbeing.

Collaboration and engagement among consumers, families and health professionals.

Teams and partnerships and how teams work to improve genuine person-centredness.

Team support for a more coordinated and effective healthcare delivery system, improving outcomes and reducing errors.

Making sure there is continuity between and within services.

Shared knowledge and experience.

Shared responsibility.

Knowledge transfer.

Shared learning.

Transparency of health information across systems.
WHAT MATTERS IN PRACTICE

SCV

Priorities
Support health services to involve consumers in improvement and innovation activities.
Support health service leaders to develop their skills through team-based learning.
Support consumer participation in clinical governance.
Further develop leadership networks.

Actions
Develop a network with consumers and health services to facilitate the systematic sharing of knowledge and good practice in Partnering in healthcare implementation.
Facilitate a community of practice to support co-design and engagement activities across SCV and the department.
Refresh community advisory committee guidelines.
Embed consumers in clinical networks.
Provide education for boards and senior staff on consumer engagement.

Department

Priorities
Increase participation of service users, providers and staff in the design and delivery of services, policy and legislation.
Increase co-design and engagement with patients, clients, victim survivors of family violence, health professionals, practitioners and service providers.

Actions
Support the work of SCV.
Support rural and regional health partnerships.
Implement the mental health lived experience engagement framework.

‘Much of the physical design and care processes in hospitals are not designed for working together. There should be a focus on creating physical spaces for communication, through the care pathway.’
Health service

**Suggested priorities**
- Implement the *Delivering high-quality healthcare: Victorian clinical governance framework*.
- Ensure better coordination, integration and continuity of care through clear mechanisms.
- Support practical tools and strategies to improve collaboration and engagement among consumers, families and health professionals.
- Integrate medical records and improve transparency.
- Improve discharge communication and processes for consumers.
- Promote clinical and multi-disciplinary teamwork.
- Include consumers in staff training.
- Build cultures of inclusion, trust and support.
- Support clinician, consumer and family team meetings.
- Co-design care and services with consumers and families.
- Use feedback from consumers for quality improvement.
- Measure and report on quality, safety, consumer experience and outcomes data.

**Actions**
- Identify at least two domains (and which priorities within them) you will focus on by **30 June 2019**.
- Use the self-assessment tool to map your health service’s current activities against the domains and priorities.
- Complete the statement of intent and share with SCV.
- Evidence for NSQHS Standards is a baseline measure for *Partnering in healthcare*.
- The *Partnering in healthcare* and NSQHS Standards resource highlights how the framework connects to the standards.

‘Better frameworks to support consumer engagement at all levels. For example, goal setting for consumers, consumer representation in service planning, and feedback at point-of-care.’
Consultation

Other suggestions for improvement from our consultation

Ask consumers what matters to them and what they feel should be done differently, then act on what they say.

Improve collaboration and communication channels between different clinical teams and health professionals.

Build systems that work together – rather than being silos – within, across and outside health services.

Ensure continuity of care outside hospital, particularly in relation to discharge planning, follow-up and feedback to primary care providers.

Ensure continuity of care when people are transferred between metropolitan and rural or regional health services.

Develop strength-based planning that includes consumers.

Enhance opportunities for clinical team meetings with consumers.

Provide appropriate and accessible information and education for staff and consumers on consumers’ rights and responsibilities.

Co-design and co-produce improvement strategies with consumers.

Ensure there are enough family/carer meetings with health professionals prior to discharge.

Use consumer feedback to drive change.

‘Ensure comprehensive care planning. Discharge planning should be done by somebody outside the hospital, who is fully aware of available support services that can meet the needs of the consumer.’
Shared decision-making includes the process by which health decisions are made by consumers and health professionals, using the best available evidence and discussion of consumers’ preferences (Stacey et al., 2016). Key tools to support shared decision-making include decision aids and decision coaching.

WHAT CONSUMERS SAID

- Involve consumers at the service design, policy and governance levels.
- Use consumers’ expertise in care.
- Include shared decision-making in performance management frameworks.
- Provide consumer education opportunities, so they can best prepare for interactions with health professionals.
- Make consumers primary decision-makers about healthcare and treatment that affects them.
- Involve consumers in decisions about how healthcare and the health system can be improved.
- Implement collaborative decision-making that considers the views and needs of all parties.

Elements of this domain

Sharing power and responsibility in decision-making.

Acting on the principle of ‘nothing about us without us’.

Involving affected individuals in decisions.

Methods for making health decisions involving consumers and health professionals.

Information outlining treatment options, outcomes and uncertainties.

Routine use of decision aids in clinical practice.

Decision support counselling or coaching to clarify options and preferences.

Recording, communicating and implementing consumers’ preferences.

Community participation in healthcare governance and priority setting.
‘Health professionals should encourage consumers to ask questions, including regarding decisions they are unsure about. Some health professionals actively discourage effective feedback and questioning of their decisions. Power imbalance needs to be addressed in some way. Some consumers believe if they do ask questions, or question decisions, that the quality of care they receive will be negatively affected.’

WHAT MATTERS IN PRACTICE

SCV

Priorities
Deliver advice and support to promote the uptake of shared decision-making approaches, including decision aids and coaching.

Actions
Develop and implement a pilot to increase the use of consumer decision aids, decision coaching and question prompt lists in Victorian hospitals.

Department

Priorities
Increase client and patient choice concerning the services and treatment they receive.

Increase consumer participation in the design and delivery of services, policy, planning, regulation and legislation.

Actions
Implement the Public participation framework: Department of Health and Human Services, and its resource toolkit, to facilitate engagement and co-design approaches.
Health service

Suggested priorities

Develop shared decision-making as a priority goal for consumers and health professionals.

Provide reliable, balanced, evidence-based information outlining treatment options, outcomes and uncertainties.

Increase the routine use of decision aids and decision coaching in clinical practice to clarify options and preferences.

Record, communicate and implement consumers’ preferences.

Involve consumers at the service design, policy and governance levels.

Utilise consumers’ expertise in care.

Include shared decision-making in position descriptions and performance appraisals.

Ensure that shared decision-making interventions are customised to meet the needs of specific groups and populations.

Actions

Identify at least two domains (and which priorities within them) you will focus on by 30 June 2019.

Use the self-assessment tool to map your health service’s current activities against the domains and priorities.

Complete the statement of intent and share with SCV.

Evidence for NSQHS Standards is a baseline measure for Partnering in healthcare.

The Partnering in healthcare and NSQHS Standards resource highlights how the framework connects to the standards.

‘Have clear pathways and provide information to make shared decision-making – as opposed to making assumptions.’
Consultation

Other suggestions for improvement from our consultation

Provide information to consumers, and ensure they have the opportunity to ask questions about their rights and care options.

Work through situations in which consumer priorities/preferences are different from those of health professionals.

Involve consumers in planning services.

Create an environment which is respectful, safe and positive for consumers. This includes being friendly, respecting consumers’ decisions, and treating all people as equals.

Include shared decision-making in performance management frameworks.

Redress any power imbalance, so systems work better for consumers.

‘Consumers and loved ones need to be encouraged and empowered to ask questions and be part of decision-making. Allowing time to ask questions, and have staff answer them in an understandable way, will support this.’
Responding to diverse needs is complex. People want to be engaged and empowered in their healthcare experience and journey. Some groups and communities are often under-represented in healthcare participation opportunities and structures and over-represented in clinical risk. Informed by a definition of equity provided by the World Health Organization, equity is the absence of avoidable or remediable differences among groups of people, whether they are defined socially, culturally, linguistically, economically, demographically or geographically (HPH Task Force, 2014). Equity in healthcare means that all people receive care of equal quality that is safe, effective and person-centred. An equitable approach does not mean that everyone receives the same care, but that all people have their healthcare needs equally well met, including through removing differences in the quality, safety and accessibility of opportunities, services and rights between groups of people.

WHAT CONSUMERS SAID
- Provide accredited interpreters when needed.
- Engage diverse consumers in healthcare partnerships.
- Provide equitable access, care and treatment.
- Ensure participation in forums and consultations is available at a variety of times of day (for example during and after work hours).

Elements of this domain
- Fairness, dignity and respect.
- Responding to everybody’s common and unique needs.
- Seeing, hearing, valuing, including and engaging others.
- Understanding how social determinants of health impact people’s healthcare access, outcomes and experiences.
- Pursuing shared, equitable and socially just outcomes.
- Cultural safety, and culturally responsive and inclusive care.
- Accredited interpreters are provided when needed.
- Addressing avoidable or remediable differences among groups of people whether they are defined socially, culturally, linguistically, economically, demographically or geographically.
- All people have their healthcare needs equally well met.
- Diverse groups and communities are involved in participatory structures.
‘Interpreters are provided to people who require them along their care journey.’

WHAT MATTERS IN PRACTICE

SCV

Priorities
Develop and promote consumer engagement guidelines, with an emphasis on engaging diverse consumers from a broader demographic of the population.

Actions
Work with partner organisations to develop resources which foster and enable diverse consumer engagement, and more inclusive consultation strategies.

Department

Priorities
Increase participation in universal and early-intervention services – especially by Aboriginal Victorians.

Reduce unexplained variation in the care people receive – especially disadvantaged groups.

Increase consumer participation in design and delivery of services.

Actions
Implement a review of the Improving care for Aboriginal and Torres Straits Islander Patients (ICAP) program.


Reform the department’s language services policy and guidelines, to strengthen quality and accessibility of language services.

Improve equity of access to high-quality and safe healthcare for regional and rural Victorians.

Monitor and review regional and rural consumers’ access to the Victorian Patient Transport Assistance Scheme (VPTAS).

Investigate ways to benchmark equity, diversity and responsiveness.
Health service

Suggested priorities

Ensure accredited interpreters are provided when needed.

Monitor consumer-reported accredited interpreter provision through the Victorian Healthcare Experience Survey (VHES).

Provide cultural safety and cultural responsiveness training for staff.

Provide advocates for those with limited or no personal supports.

Ensure diversity, culture and inclusion are organisational priorities.

Build diverse consumer representation at all levels.

Value consumers, including through remuneration.

Develop more inclusive community consultation strategies.

Ensure information is translated into various languages.

Promote consumer awareness of healthcare rights.

Ensure regional and rural consumers have access to VPTAS.

Actions

Identify at least two domains (and which priorities within them) you will focus on by 30 June 2019.

Use the self-assessment tool to map your health service’s current activities against the domains and priorities.

Complete the statement of intent and share with SCV.

Evidence for NSQHS Standards is a baseline measure for Partnering in healthcare.

The Partnering in healthcare and NSQHS Standards resource highlights how the framework connects to the standards.

‘Shifting care processes from a ‘one size fits all’ to ‘one size fits one’.'
Consultation

Other suggestions for improvement from our consultation

Ensure people who need accredited interpreters have access to them.

Provide critical health information in relevant languages and formats.

Respond effectively to diversity and difference.

Increase awareness of (and services/facilities for) a person’s cultural, religious and social needs.

Ensure greater access to accredited interpreters for consumers with limited English proficiency, including information in their preferred language.

Create environments that are respectful and safe for all.

Ensure adequate training in specific areas such as cultural responsiveness, mental health, providing and using accredited interpreters, and checking for understanding.

Address barriers for particularly at-risk communities or population groups.

Ensure data is collected – and acted upon – regarding who is accessing services, and their experiences and outcomes.

Ensure participation across diverse groups and communities, which reflect the demographics of the communities served.

‘Different systems are required for people in different stages of life and health. People with a disability, people who are older, and people who are disenfranchised in the hospital system, need to be supported. Not all people are able to articulate their health issues, nor do all people have carers who can advocate for them.’
Effective communication

Health communication refers to interactions that occur during the process of improving health and healthcare. Effective health communication is essential for public health strategy and practices. Poor quality communication, and the mistakes associated with it, are a major cause of error in diagnosis and treatment. Health literacy is an enabler of communication and participation in healthcare. It is also the product of good communication between health professionals and consumers, and of health systems that are responsive to consumer needs (Phillips, 2016).

WHAT CONSUMERS SAID

- Always use clear, jargon-free and accessible language.
- Co-develop information about healthcare conditions, processes and pathways that are easy to understand and act upon.
- Ensure communication is consistent across all points of a patient’s journey.
- Provide quality health information and in relevant languages.
- Promote the use of ‘Teach back’ by health professionals.

Elements of this domain

Quality of communication in all healthcare relationships.

Open disclosure.

Healthcare experience feedback mechanisms for consumers.

Active listening.

Health literacy.

Developing and providing health information.

Opportunities to use health information and services.

Online interactions.

Communication skills and capabilities of health professionals.

‘I receive high-quality information that I can readily understand and act upon.’

‘Place value on better communication skills and kindness. Change KPIs so these skills matter.’
WHAT MATTERS IN PRACTICE

SCV

Priorities
Develop and promote tools to improve health literacy.
Support open disclosure between consumers and health services when things go wrong.
Provide alternative pathways for consumers to escalate their concerns when they are worried.
Provide different ways for consumers to provide feedback about their experiences in healthcare.
Support training for consumers to participate in serious event investigations.
Support training for health professionals to enhance communication.

Actions
Disseminate best practice health literacy guidance to health services (in partnership with the Centre for Health Communication and Participation).
Implement and review a pilot of the Patient Opinion feedback tool.
Implement a consumer-initiated escalation of care program.
Roll out communication skills training for healthcare professionals, including clinical communication skills, and effective communication for person-centred care.
Identify and deliver training that builds compassionate workplaces.

Department

Priorities
Increase consumer participation in design and delivery of services.

Actions
Implement a range of reforms to the department’s language services policy and guidelines to strengthen the quality and accessibility of language services.
Suggested priorities
Listen to consumers to ensure mutual understanding.
Promote friendly, supportive interactions.
Facilitate open and timely communication with consumers.
Develop clear written communication.
Provide training for staff on respectful communication.
Improve hospital environments to facilitate effective communication.
Support strategies to improve health literacy.
Provide training for staff on health literacy (for example Teach-back, Ask Me 3 and Asking the Right Questions Matter).
Enable consumers to provide feedback.
Use technology to communicate with consumers when that method is right for them.
Provide information for consumers (in multiple formats) before, during and after consultations.
Improve communication, so it is responsive to cultural, linguistic, cognitive and other needs.

Actions
Identify at least two domains (and which priorities within them) you will focus on by **30 June 2019**.
Use the self-assessment tool to map your health service’s current activities against the domains and priorities.
Complete the statement of intent and share with SCV.
Evidence for NSQHS Standards is a baseline measure for Partnering in healthcare.
The Partnering in healthcare and NSQHS Standards resource highlights how the framework connects to the standards.

‘Nurses often provide a large amount of consumer education. However, they must find resources to implement and reinforce the concepts discussed. Finding credible and appropriately succinct information can often take more time than the actual education itself.’
Consultation

Other suggestions for improvement from our consultation

Listen to and act on consumers’ needs.

Avoid complex technical jargon and provide information in a clear and accessible way.

Provide clear explanations of conditions and treatment options.

Ensure information and educational resources are readily available for consumers.

Improve communication between hospital and non-hospital health professionals, including primary healthcare providers.

Use a variety of materials that include visual, spoken and written approaches.

Implement policy and service reforms for language services provision.

‘Introduce tools that enable people to communicate their needs and preferences (for example, a bedside whiteboard). Support people to record these. These needs and preferences should be clearly documented in the health record, and clearly linked to their healthcare, treatment, rehabilitation and discharge plan.’
Partnering in healthcare is an ongoing process. It involves all stakeholders in assessing issues and co-developing potential solutions.

Implementing the framework involves five steps:

1. **Review the domains and suggested priorities for health services in this framework**

   Partnering in healthcare domains and priorities align with many initiatives currently underway within health services. This framework is designed to build on current initiatives, identify future areas of work and improve innovation.

2. **Use the Partnering in healthcare self-assessment tool**

   Use the Partnering in healthcare self-assessment tool to assess current strengths and challenges against each domain.

3. **Identify areas in need of improvement**

   This will help identify where improvements may bring real benefits and impact, and therefore, where time and effort is best invested.

4. **Identify at least two domains and the priorities you will choose to focus on in the next 12 months**

   Make a decision about which two domains and priorities to focus on in line with your existing initiatives, to best meet the needs of your consumers and organisation.

5. **Complete the Partnering in healthcare statement of intent and share with SCV by 30 June 2019**

   Complete the statement of intent outlining your two chosen domains and identified priorities. This should be endorsed by your health service board. Email to: partnering@safercare.vic.gov.au.

   The statement of intent can help to monitor implementation, progress and achievements.

‘Our organisation fosters mutual learning, and embeds co-design in healthcare governance, planning, implementation, monitoring and evaluation.’

Health services should identify at least two domains to focus on by 30 June 2019. Your statement of intent should be endorsed by your health service board.
Health services will be able to monitor and track their progress through a mix of existing internal and external mechanisms. These include:

- health service strategic plans
- health service quality improvement plans
- health service organisational policies, plans, improvement goals and monitoring frameworks
- statement of priorities
- departmental policy and funding requirements
- Victorian health services performance monitoring framework
- Delivering high-quality healthcare: Victorian clinical governance framework
- NSQHS Standards (second edition)
- VHES data analysis.

**Partnering in healthcare forum**

In advancing and supporting this framework, we will host a **Partnering in healthcare** forum on **29-30 April 2019**. It will bring consumers and health services together to:

- celebrate what you are currently doing to progress consumers partnering in their healthcare
- share good practice examples
- generate new thinking and ideas
- identify the most useful ways to measure and report on progress with **Partnering in healthcare**
- identify how can we measure improvements in participation.

The knowledge and lessons from this event will be documented and shared with all Victorian health services.

Further stages will include working with health services to develop ways to systematically share knowledge and good practice in implementing **Partnering in healthcare**.

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**Partnering in healthcare is everybody’s business**

This includes:

- SCV
- the department
- consumers
- health service boards
- chief executive officers
- executive and managers
- clinical leaders
- consumer experience and engagement professionals
- quality and safety professionals.

‘We partner in each aspect of decision-making with the people who use or deliver our services. We provide training and support for people to be involved in their care.’
‘I have hope that the health system will improve, that consumers are being heard and their contributions valued. Consumer involvement is integral to creating a responsive health system.’

Figure 3: Implementation timeline

What we will deliver in 2019–20

**FEBRUARY**
- Launch of *Partnering in healthcare* framework document and planning resources

**MARCH**
- Shared decision-making to feature in Better Care Victoria Innovation Fund

**APRIL**
- Inaugural *Partnering in healthcare* forum held 29–30 April

**JUNE**
- 30 June deadline for health services to submit at least two domains and priority areas for action

**JULY**
- Identify baseline data for *Partnering in healthcare* implementation
  - Develop evaluation strategy

**DECEMBER**
- Develop resources to support shared decision-making strategies
- Launch resources to support equitable and diverse consumer engagement

**2020**
- *Partnering in healthcare* outcomes summit held
- Shared decision-making projects launched in health services
EVALUATION AND REVIEW

In supporting the healthcare improvement process, we want to know what has improved, or what has changed. Tracking and documenting improvement processes as they evolve allows us to learn what works and build the evidence base.

To help measure and monitor progress, SCV will develop an evaluation strategy with consumers and health services which considers:

- how will we track and analyse the improvement process as it evolves?
- which information, data and evidence need to be collected, by whom, how, and when?
- what insights about healthcare improvement are being produced through Partnering in healthcare?

SCV will support the evaluation and review of Partnering in healthcare through the SCV Patient and Family Council. The council aims to involve consumers in a meaningful and effective partnership. It is one of the key mechanisms to ensure consumers and community needs and perspectives are represented in health service:

- development plans
- improvement goals
- program implementation and evaluation.

This approach provides a governance and reporting structure that will help to:

- review achievements in the five domains through the identified priorities and actions
- develop an evaluation strategy to capture and analyse learnings from the improvement effort.

Annual outcomes summit

We propose to hold an annual outcomes summit to support consumers and health services to share Partnering in healthcare achievements, learnings and good practice strategies. This will be documented and shared with participants.

Partnering in healthcare is an ongoing improvement strategy. SCV will continue to develop and improve Partnering in healthcare priorities, actions and resources. This will be done in partnership with consumers and health services, the department, as well as other healthcare, education and peak organisations.

‘Involving people in all aspects of care is a priority for our organisation. Managers take a leadership role, encouraging staff to involve people as much as they would like.’

‘Our board starts each meeting with a consumer story.’
Background

The role of the healthcare consumer is changing. Consumer participation in their own treatment is now a key indicator of healthcare performance and quality (Conway et al., 2006). This places different expectations on our healthcare system and requires health services to develop new ways to ensure consumers are equal and active partners.

Reducing variation in consumer participation and experience

All Victorians should be able to:

- access high-quality, safe and effective healthcare, and have their healthcare needs equally well met
- meaningfully participate in decision-making about their health and wellbeing
- meaningfully participate in planning, delivering and evaluating healthcare at both a service and system level.

There have been significant advances in the quality and safety of healthcare. However, finding the best ways to respond to people’s unique needs, and to address healthcare inequities, remains a challenge. As a result, not all Victorians participate in their health in an equal way, or have their healthcare needs equally well met.

Our research and data tell us that there is variation in:

- how health services promote and engage with consumers to partner in their own care
- the quality of consumer experience
- people’s access to an accredited interpreter and information in their own language
- experiences of person-centredness, participation, respect and quality of care
- capacity of health services to partner with consumers in policy development and governance
- capacity of health services to address diversity and equity in participation mechanisms.

The changing healthcare environment

Our priorities shift with significant changes in healthcare, such as:

- growing preference for more personalised services and treatment, more choice, greater co-design, and shared decision-making
- individuals becoming partners in their own care, which ultimately translates to better outcomes and more effective services
- ensuring that services and products are tailored to people’s needs
- better ways to organise services to break down artificial barriers within and between health services
- making it easier for people to access and navigate services
- supporting health professionals and consumers to access better data which informs conversations and practice
- integrating health and social care
- strengthening devolved governance (where decision-making happens at all levels of the system, not just ‘the top’)
- understanding the social determinants of ill health and its context
understanding and responding to the diversity of consumers, nurturing cultural safety, and ensuring freedom from bias and discrimination (real or perceived)

- progressing Aboriginal self-determination in the design and delivery of services

- use of digital media and technology

- being at the frontier of genomics, digital health, telehealth and big data.

DEVELOPING THE FRAMEWORK

In 2017, SCV set about creating a new framework for consumer participation in healthcare. This built on the previous work undertaken by the department.

The framework comprised five key areas, drawn from the best evidence, practice and research, through which work could make a real difference. To test this framework, SCV consulted widely about how best to improve healthcare in Victorian hospitals. The main activities included:

- hosting an online consultation and survey, receiving 680 responses from Victorian consumers, and those who work in our healthcare system. Through this process, SCV learnt about individuals’ experience of health services, and their suggestions for improvement

- hosting a priorities summit workshop with 31 people (consumer-health professional ratio of two to one). It was a special opportunity to explore, learn and document what people thought was most important in improving healthcare when in hospital, and what could be done differently.

Over the past year, we have consulted with more than 750 consumers and health professionals on our draft framework, and on developing priorities for improving quality and safety of healthcare in Victoria.

Why we developed this framework

Victoria needed to develop a framework for consumer participation in healthcare, to improve equity in treatment and care options, and reduce healthcare variation across Victoria.

This framework supports the strategic priorities of both SCV and the department. It was also informed by recommendations in:

- Consumer participation in the health system (Victorian Auditor-General’s Office, 2012)

- KPMG recommendations regarding the summative evaluation of policy frameworks:
  - Doing it with us not for us: Strategic direction 2010–2013 (2009)
  - Cultural responsiveness framework (2009)

- Targeting zero: Report of the Review of Hospital Safety and Quality Assurance in Victoria (Department of Health and Human Services, 2016), which recommended that SCV adopt consumer engagement and consumer experience as a priority improvement goal for the hospital system.
Partnering in healthcare is connected
This framework supports the numerous policies of the department, as well as work already underway in Victorian health services.

It supports the department’s Public participation framework (2018a) and Stakeholder engagement toolkit (2018b), which outline the department’s overarching stakeholder engagement, and public participation vision and processes.

This framework complements accountability and performance reporting
Health services currently have multiple accountability and reporting mechanisms that drive participation, quality and safety in healthcare. In addition, health services are required to report on many quality and safety performance measures at a local, statewide and national level.

Clinical governance framework
Partnering in healthcare outlines expectations for implementing the consumer partnerships domain described in Delivering high-quality healthcare: Victorian clinical governance framework. That is, the consumer is at the centre of care, and is viewed as a critical partner in healthcare design and delivery.

NSQHS Standards
From 1 January 2019, health services are implementing the NSQHS Standards (second edition). In this new edition, there is an increased focus on partnering with consumers.

The Partnering with Consumers Standard remains an overarching standard, but introduces new and specific core actions, with the expectation that health services will provide evidence of continuous improvement in:

- actively involving consumers in their own care.
- meeting consumers’ information needs.
- shared decision-making.

A shift in direction now requires health services to show how healthcare initiatives and continuous improvement programs influence the quality and safety of person-centred care. Including consumers is key to the evaluation process, in order to learn from consumer feedback and experience.

Evidence for NSQHS Standards is a baseline measure for Partnering in healthcare.

The Partnering in healthcare and NSQHS Standards resource highlights how the framework connects to the standards.

Victorian health experience survey
The Partnering in healthcare domains and priorities align with the VHES questions. You can use your VHES results to track and monitor progress in the domains and priorities.
Accredited interpreter
It is Victorian Government policy that wherever possible, organisations should engage interpreters and translators accredited at the professional level by the National Accreditation Authority for Translators and Interpreters (NAATI). NAATI accreditation is the only qualification officially accepted for the profession of translation and interpreting in Australia. Accredited interpreters and translators act in accordance with the Australian Institute of Interpreters and Translators (AUSIT) Code of Conduct and Code of Ethics. NAATI also provides an online directory of accredited interpreters and translators. Multilingual language skills can also be verified through a NAATI language aide or interpreter test.

Agency
In order to bring about improvements in health and healthcare, people’s agency may include:

- the ability of an individual or group to act independently and make their own choices
- how they shape their own health and healthcare in partnership with health professionals
- the emotional resources and support to do so in the face of difficulty or uncertainty.

Co-design
Is a method of human-centred design, where new approaches to services are created with the people who use or deliver those services. They are typically participatory processes that collaboratively explore consumer and staff experiences and ideas, consider how these relate to the care journey, develop and implement improvements, and review what difference these make.

Decision aids
Consumer decision aids are tools that help people become involved in decision-making. They explicitly identify the decision that needs to be made, provide information about the options and possible outcomes, and clarify personal values. They are designed to complement, rather than replace, counselling from a health professional.

Equal partners
This term describes consumers being partners in their own care, to the extent that they choose to be.

Equity
Informed by a definition of equity provided by the World Health Organisation, equity is the absence of avoidable or remediable differences among groups of people, whether they are defined socially, culturally, linguistically, economically, demographically or geographically (HPH Task Force, 2014).
Equity in healthcare

Equity in healthcare means that all people receive care of equal quality that is safe, effective and is responsive to their individual needs, culture, language, ability, experience and preferences. This does not mean that everyone receives the same care, but rather that all persons have their healthcare needs equally well met, including through removing differences in the quality, safety and accessibility of opportunities, services and rights between groups of people.

Consumer

Consumers include people, families, carers and communities who are current or potential users of health services. This includes children, women and men, people living with a disability, patients, carers, clients, people of diverse cultural, linguistic and religious experiences, socioeconomic status and social circumstances, sexual orientations, sexes, genders and gender identities, health and illness conditions. The term also includes: people who choose to get involved in decision-making; health consumer representatives who provide advice on behalf of consumers, with the overall aim of improving healthcare; and carers who often have an important role in health care decision making and care giving. Different health settings use terms such as: patients, people/persons, families, carers, clients and residents.

Health consumer representative

A health consumer representative is a health consumer who has taken up a specific role to provide advice on behalf of consumers, with the overall aim of improving healthcare. A consumer representative is often a consumer member of a committee, project or event, who voices consumer perspectives and takes part in decision-making on behalf of consumers. A health consumer representative may be nominated by, and accountable to, a consumer organisation. Usually the person works with a health service or consumer organisation, but they may also operate independently in some activities.

NSQHS Standards (second edition)

The National Safety and Quality Health Service (NSQHS) Standards were developed by the Australian Commission on Safety and Quality in Health Care with the Australian Government, state and territory partners, consumers and the private sector. The primary aim of the NSQHS Standards is to protect the public from harm, and improve the quality of healthcare. They describe the level of care that should be provided by health service organisations, and the systems that are needed to deliver such care. The second edition of the NSQHS Standards was released in November 2017. Health service organisations will be assessed against the standards in the second edition from January 2019.
Partnering with consumers
Partnering with consumers is about healthcare organisations, healthcare providers and policy-makers actively working with people who use the healthcare system to ensure that health information and services meet people’s needs. Essentially, partnerships with consumers exist when consumers are treated with dignity and respect, information is shared with them, and their participation and collaboration is supported.

Patient Activation Measure
Patient (consumer) activation is a behavioural concept. It captures several core components of consumer involvement, each of which is important for active engagement and participation. It is defined as an individual’s knowledge, skill, and confidence in managing their health and healthcare. Consumers with high levels of activation understand their role in the care process and feel capable of fulfilling it. Individuals with long-term conditions who are more highly activated are more likely to engage in positive health behaviours, and to manage their health conditions more effectively.

Patient Opinion
Patient Opinion was founded in the UK in 2005. Since then it has grown to be the UK’s leading independent non-profit feedback platform for health services. Patient Opinion Australia (POA) was established in 2012 and, like its UK counterpart, is registered as an independent not-for-profit charitable institution. Patient Opinion is about honest and meaningful conversations between consumers and health services.

Patient-reported experience measures (PREMs)
PREMs are questionnaires used to obtain consumers’ views and observations on aspects of health services they have received. This includes their views on the accessibility and physical environment of services (for example, waiting times and the cleanliness of consultation rooms and waiting spaces) and aspects of consumer–clinician interactions (such as whether the clinician explained procedures clearly, or responded to questions in a way the consumer could understand).
Patient-reported outcome measures (PROMs)

PROMs are questionnaires which consumers complete. They ask for the consumers’ assessments of how health services and interventions have, over time, affected their quality of life, daily functioning, symptom severity, and other dimensions of health which only consumers can truly know. PROMs promise to fill a vital gap in knowledge about outcomes, and about whether healthcare interventions make a difference to people’s lives.

Social determinants of health

The social determinants of health are the conditions in which people are born, grow, live, work and age that can and do influence their health. These circumstances are shaped by the distribution of money, power and resources at global, national and local levels. The social determinants of health are mostly responsible for health inequities - the unfair and avoidable differences in health status seen within and between countries.

Victorian Agency for Health Information (VAHI)

VAHI was created as part of Victorian Government reforms to overhaul quality and safety across Victoria’s public health system. VAHI produce regular reports for health services to monitor safety and performance through data (such as rates of potentially preventable infections and readmissions). Health services can see their performance against relevant targets and compare this to the performance of similar health services. VAHI also collect and report on consumers’ experiences of Victoria’s public health services, and work with other stakeholders to produce reports on selected topics of public interest.

Victorian Healthcare Experience Survey (VHES)

The Victorian Healthcare Experience Survey (VHES) collects data from a range of users of Victorian public health services. The survey is conducted on behalf of the department by Ipsos (an independent contractor). The survey is completed by a random sample of eligible adults aged 16 or over who have used public health services. Results cover key factors that contribute to consumers’ experience with a health service, such as waiting time, how clean the bathrooms were, and the information they were given when discharged.
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Departmental policies and documents


Useful resources


