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Western Australia: patient outcomes in palliative care: July - December 2014

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Western Australia: patient outcomes in palliative care: July - December 2014

Abstract

The Palliative Care Outcomes Collaboration (PCOC) assists services to improve the quality of the palliative care they provide through the analysis and benchmarking of patient outcomes. In this PCOC report, data submitted for the July to December 2014 period are summarised and patient outcomes benchmarked to enable participating services to assess their performance and identify areas in which they may improve.

Keywords

PCOC, palliative, care, western, patient, australia, outcomes, july, 2014, december

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Western Australia

Patient Outcomes in Palliative Care

July – December 2014

March 2015



About the Palliative Care Outcomes Collaboration

The Palliative Care Outcomes Collaboration (PCOC) is a national program that utilises standardised clinical assessment tools to measure and benchmark patient outcomes in palliative care. Participation in PCOC is voluntary and can assist palliative care service providers to improve practice and meet the Palliative Care Australia (PCA) Standards for Providing Quality Palliative Care for all Australians. This is achieved via the PCOC dataset; a multi-purpose framework designed to:

- provide clinicians with an approach to systematically assess individual patient experiences,
- define a common clinical language to streamline communication between palliative care providers and
- facilitate the routine collection of national palliative care data to drive quality improvement through reporting and benchmarking.

The PCOC dataset includes the clinical assessment tools: Palliative Care Phase, Palliative Care Problem Severity Score (PCPSS), Symptom Assessment Scale (SAS), Australia-modified Karnofsky Performance Status (AKPS) scale and Resource Utilisation Groups – Activities of Daily Living (RUG-ADL).

PCOC has divided Australia into four zones for the purpose of engaging with palliative care service providers. Each zone is represented by a chief investigator from one of the collaborative centres. The four PCOC zones and their respective chief investigators are:



Each zone is also represented by one or more quality improvement facilitators, whose role includes supporting services to participate in PCOC and facilitating ongoing service development and quality improvement. The national team, located within the Australian Health Services Research Institute at the University of Wollongong, coordinates the patient outcomes reporting, education program, and quality activities across the four zones.

If you would like more information or have any queries about this report please contact your local quality improvement facilitator or contact the national office at pcoc@uow.edu.au or phone (02) 4221 4411.



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Introduction

The Palliative Care Outcomes Collaboration (PCOC) assists services to improve the quality of the palliative care they provide through the analysis and benchmarking of patient outcomes. In this PCOC report, data submitted for the July to December 2014 period are summarised and patient outcomes benchmarked to enable participating services to assess their performance and identify areas in which they may improve.

Patient outcomes are reported for a total of 18,310 patients, with 23,449 episodes of care and 53,467 palliative care phases. The information included in this report is determined by a data scoping method. See Appendix A for more information on the data included in this report.

Throughout this report, patient information for Western Australia is presented alongside the national figures for comparative purposes. The national figures are based on information submitted by 95 services, of which:

- 53 are inpatient services. Inpatient services include patients who have been seen in designated palliative care beds as well as non-designated bed consultations.
- 27 are community services. These services include primarily patients seen in the community as well as some patients with ambulatory/clinic episodes.
- 15 are services with both inpatient and community settings.

A full list of the services included in the national figures can be found at www.pcoc.org.au.

The Western Australian figures in this report are based on information submitted by 10 services. A list of these services is presented in Table 1 on the following page.

Interpretation hint:

Some tables throughout this report may be incomplete. This is because some items may not be applicable to Western Australian services or it may be due to data quality issues.

Please use the following key when interpreting the tables:

- na The item is not applicable.
- u The item was unavailable.
- s The item was suppressed due to insufficient data as there was less than 10 observations.



Table 1 List of Western Australian services included in this report

Service name	Setting of care
Bethesda Hospital	Inpatient
Northam Palliative Care	Both
Palliative Ambulatory Service North	Community
Ramsay Health Care Peel Health Campus	Inpatient
Royal Perth Hospital	Inpatient
Royal Perth Hospital - Nurse Practitioner	Inpatient
Silver Chain Hospice Care Service	Community
St John of God - Murdoch Community Hospice	Inpatient
St John of God Bunbury Hospital	Both
St John of God Geraldton Hospital	Inpatient



Section 1 Benchmark summary

1.1 Western Australia at a glance

Table 2 Summary of outcome measures 1 to 3 by setting

		Benchmark	Ir	patient	Community	
Outcome measure	Description		WA Score	Benchmark Met?	WA Score	Benchmark Met?
1. Time from ready for care to episode start	Benchmark 1: Patients episode commences on the day of, or the day after date ready for care	90%	96.0	Yes	96.8	Yes
2. Time in unstable phase	Benchmark 2: Patients in the unstable phase for 3 days or less	90%	92.5	Yes	96.5	Yes
3. Change in pain	Benchmark 3.1: PCPSS Patients with absent/mild pain at phase start, remaining absent/mild at phase end	90%	91.4	Yes	86.2	No
	Benchmark 3.2: PCPSS Patients with moderate/severe pain at phase start, with absent/mild pain at phase end	60%	50.8	No	66.6	Yes
	Benchmark 3.3: SAS Patients with absent/mild distress from pain at phase start, remaining absent/mild at phase end	90%	85.9	No	83.9	No
	Benchmark 3.4: SAS Patients with moderate/severe distress from pain at phase start, with absent/mild at phase end	60%	50.4	No	58.2	No

Table 3 Summary of outcome measure 4: Average improvement on the 2014 baseline national average (X-CAS)

Clinical Tool	Description	Average improvement on baseline	Benchmark met?
PCPSS	Benchmark 4.1: Pain	0.00	Yes
	Benchmark 4.2: Other symptoms	0.04	Yes
	Benchmark 4.3: Family/carer	0.09	Yes
	Benchmark 4.4: Psychological/spiritual	0.01	Yes
SAS	Benchmark 4.5: Pain	0.01	Yes
	Benchmark 4.6: Nausea	0.06	Yes
	Benchmark 4.7: Breathing problems	0.11	Yes
	Benchmark 4.8: Bowel problems	0.11	Yes

The benchmark for outcome measure 4 is zero.

For more information on the outcome measures and benchmarks, see Section 2.



1.2 National benchmark profiles

In this section, the national profiles for selected benchmarks are split by setting (inpatient or community) and presented graphically.

The selected benchmarks included are:

•	Benchmark 1	Patients episode commences on the day of or the day after date ready for care
•	Benchmark 2	Patients in the unstable phase for 3 days or less
•	Benchmark 3.1	PCPSS: Patients with absent/mild pain at phase start, remaining absent/mild at phase end
•	Benchmark 3.2	PCPSS: Patients with moderate/severe pain at phase start, with absent/mild pain at phase end
•	Benchmark 3.3	SAS: Patients with absent/mild distress from pain at phase start, remaining absent/mild at phase end
•	Benchmark 3.4	SAS: Patients with moderate/severe distress from pain at phase start, with absent/mild distress from pain at phase end

Interpretation hint:

The national profile graphs on the following pages allow services to see how they are performing in comparison to other palliative care services participating in PCOC. In each graph, the shaded region describes the national profile for that outcome measure. Western Australian services are highlighted as dots on the graph.

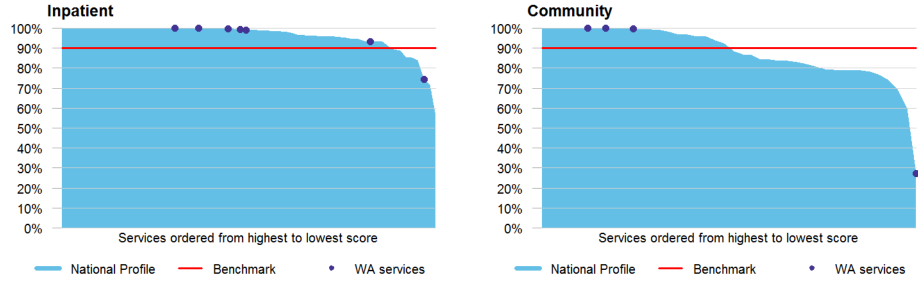
If no dots are present on a particular graph, this means that Western Australian services have not met the criteria for inclusion in this measure. This may be caused by insufficient data item completion, or not having any data falling into a particular category, for example, no phases starting with moderate/severe SAS pain.

The red line on the graph indicates the benchmark for that outcome measure.



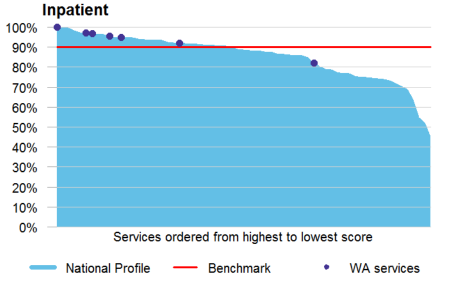
Outcome measure 1 – Time from date ready for care to episode start

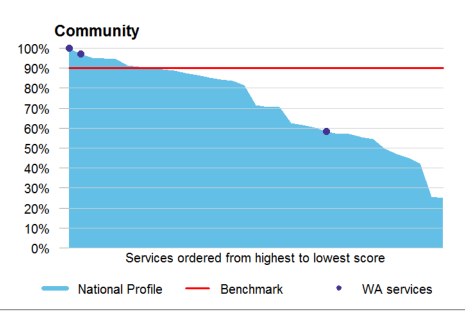
Figure 1 Percentage of patients with episodes started on the day of, or the day after date ready for care



Outcome measure 2 - Time in unstable phase

Figure 2 Percentage of patients in the unstable phase for 3 days or less







Outcome measure 3 - Change in pain

Figure 3 PCPSS: Percentage of patients with absent/mild pain at phase start, remaining absent/mild at phase end

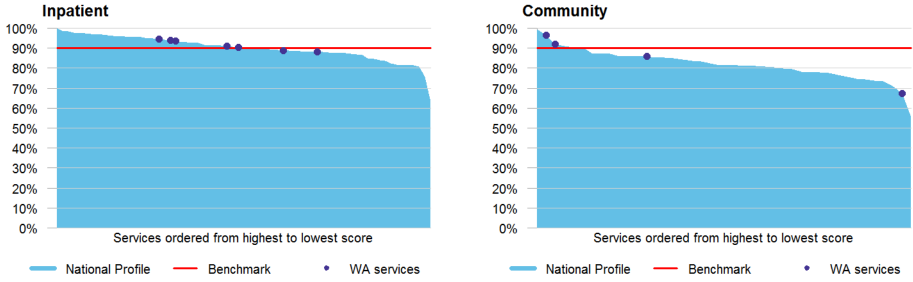
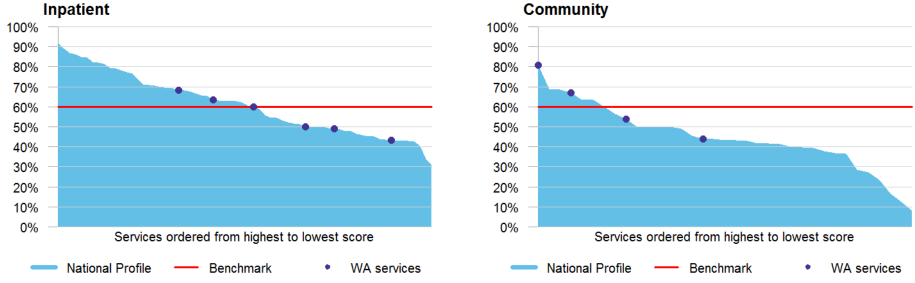


Figure 4 PCPSS: Percentage of patients with moderate/severe pain at phase start, with absent/mild pain at phase end







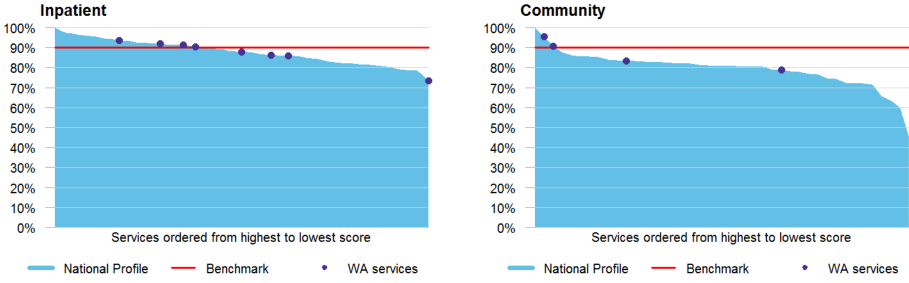
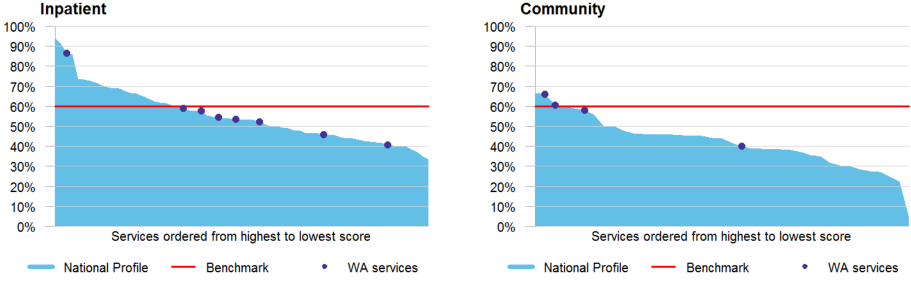


Figure 6 SAS: Percentage of patients with moderate/severe distress from pain at phase start, with absent/mild distress from pain at phase end





Section 2 Outcome measures in detail

2.1 Outcome measure 1 – Time from date ready for care to episode start

Time from date ready for care to episode start reports responsiveness of palliative care services to patient needs. This benchmark was set following feedback and subsequent consultation with PCOC participants. Service providers acknowledge that, whilst there is wide variation in the delivery of palliative care across the country, access to palliative care should be measured based on patient need rather than service availability. As a result, services operating five days a week (Monday to Friday) are not distinguished from services operating seven days a week (all services are being benchmarked together).

Benchmark 1:

This measure relates to the time taken for an episode to commence following the date the patient is available and ready to receive palliative care. To meet the benchmark for this measure, at least 90% of patients must have their episode commence on the day of, or the day following date ready for care.

Table 4 Time from date ready for care to episode start by setting

		Inpa	tient		Community				
Time (in days)	WA		All Services		WA		All Services		
	N	%	N	%	N	%	N	%	
Same day	1093	80.4	10,032	89.5	2108	96.0	8,240	82.2	
Following day	212	15.6	768	6.8	18	0.8	493	4.9	
2-7 days	53	3.9	380	3.4	51	2.3	953	9.5	
8-14 days	1	0.1	22	0.2	15	0.7	203	2.0	
Greater than 14 days	1	0.1	12	0.1	4	0.2	140	1.4	
Average	1.1	na	1.1	na	1.2	na	1.9	na	
Median	1	na	1	na	1	na	1	na	

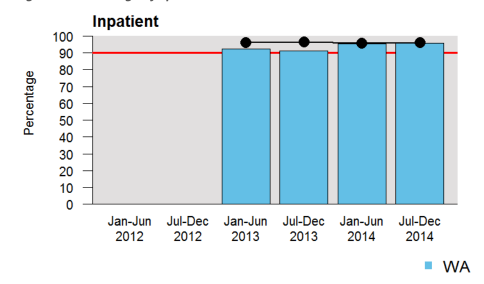
Note: Only episodes that started in this reporting period have been included in the table. Episodes where date ready for care was not recorded are excluded from the table. In addition, all records where time from date ready for care to episode start was greater than 90 days were considered to be atypical and were assumed to equal 90 days for the purpose of calculating the average and median time.

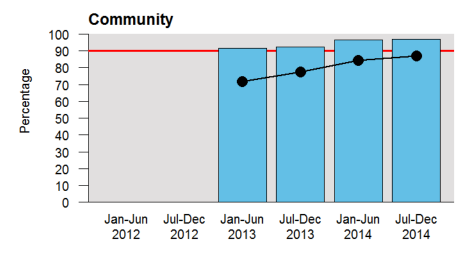
Interpretation hint:

Outcome measure 1 only includes episodes that have commenced in the reporting period. As a result, the number of episodes included in the calculation of this benchmark may not match the number of episodes in Appendix A. For more information on data scoping methods, see Appendix C.



Figure 7 Percentage of episodes that met outcome measure 1 over time







2.2 Outcome measure 2 – Time in unstable phase

The unstable phase type, by nature of its definition, alerts clinical staff to the need for urgent changes to the patient's plan of care or that emergency intervention is required. Those patients assessed to be in the unstable phase require intense review for a short period of time.

An unstable phase is triggered if:

- a patient experiences a new, unanticipated problem, and/or
- a patient experiences a rapid increase in the severity of an existing problem, and/or
- a patient's family/carers experience a sudden change in circumstances that adversely impacts the patient's care.

The patient moves out of the unstable phase in one of two ways:

- A new plan of care has been put in place, has been reviewed and does not require any additional changes. This does not necessarily mean that the symptom/crisis has been fully resolved. However, the clinical team will have a clear diagnosis and a plan for the patient's care. In this situation, the patient will move to either the stable or deteriorating phase.
- The patient is likely to die within a matter of days. In this situation, the patient will be moved into the terminal phase.

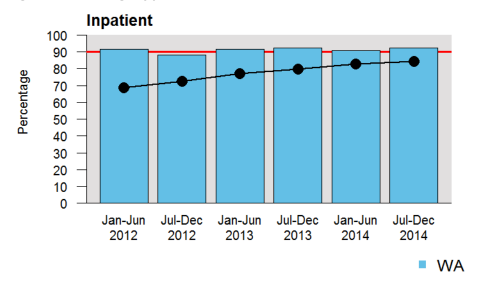
Benchmark 2: This benchmark relates to time that a patient spends in the unstable phase. To meet this benchmark, at least 90% of unstable phases must last for 3 days or less.

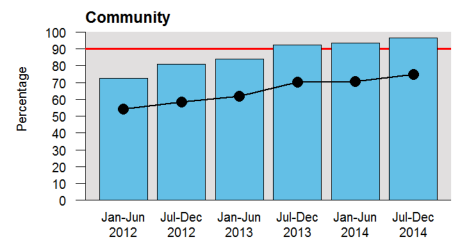
Table 5 Time in unstable phase by setting

	Inpatient				Community				
Length of unstable phase	WA		All Services		WA		All Services		
	N	%	N	%	N	%	N	%	
Same day	39	8.2	208	3.2	233	30.5	733	23.1	
1 day	273	57.1	2,917	44.6	407	53.3	1,078	33.9	
2 days	91	19.0	1,611	24.6	65	8.5	362	11.4	
3 days	39	8.2	776	11.9	32	4.2	208	6.5	
4-5 days	24	5.0	626	9.6	16	2.1	220	6.9	
6-7 days	8	1.7	225	3.4	7	0.9	175	5.5	
8-14 days	3	0.6	143	2.2	1	0.1	171	5.4	
Greater than 14 days	1	0.2	38	0.6	3	0.4	233	7.3	
Total	478	100.0	6,544	100.0	764	100.0	3,180	100.0	



Figure 8 Percentage of phases that met benchmark 2 over time







2.3 Outcome measure 3 – Change in pain

Pain management is acknowledged as a core business of palliative care services. The Palliative Care Problem Severity Score (PCPSS) and Symptom Assessment Scale (SAS) provide two different perspectives of pain. The PCPSS is clinician rated and measures the severity of pain as a clinical problem while the SAS is patient rated and measures distress caused by pain.

There are two benchmarks related to each tool: one relating to the management of pain for patients with absent or mild pain, and the other relating to the management of pain for patients with moderate or severe pain. Phase records must have valid start and end scores for the PCPSS and/or SAS clinical assessment tools to be included in the benchmarks.

Scores for PCPSS

0 absent

1 mild

2 moderate

3 severe

Scores for SAS

0 absent

1-3 mild

4-7 moderate

8-10 severe

Interpretation hint:

This outcome measure should be viewed in conjunction with **Error! Reference source not found.** Table 29 to Table 32 and Appendix B.

Benchmarks 3.1 and 3.3: These benchmarks relates to patients who have absent or mild pain at the start of their phase of palliative care. To meet these benchmarks, 90% of phases must end with the patient still experiencing only absent or mild pain.

Benchmarks 3.2 and 3.4: These benchmarks relates to patients who have moderate or severe pain at the start of their phase of palliative care. To meet these benchmarks, 60% of phases must end with the patient's pain reduced to being absent or mild.

Table 6 Summary of outcome measure 3

		Inpa	tient		Community			
Benchmark	WA		All Services		WA		All Services	
	N*	%	N*	%	N*	%	N*	%
Benchmark 3.1: PCPSS	1,310	91.4	15,589	90.9	6,741	86.2	14,943	84.8
Benchmark 3.2: PCPSS	545	50.8	5,346	57.1	1,333	66.6	3,933	50.1
Benchmark 3.3: SAS	1,368	85.9	13,526	88.1	6,322	83.9	13,991	82.7
Benchmark 3.4: SAS	838	50.4	6,541	52.8	1,699	58.2	4,879	45.4

^{*}Total number of phases included in this benchmark.



Figure 9 Trends in benchmark 3.1: PCPSS Patients with absent/mild pain at phase start, remaining absent/mild at phase end by setting



Figure 10 Trends in benchmark 3.2: PCPSS Patients with moderate/severe pain at phase start, with absent/mild at phase end by setting

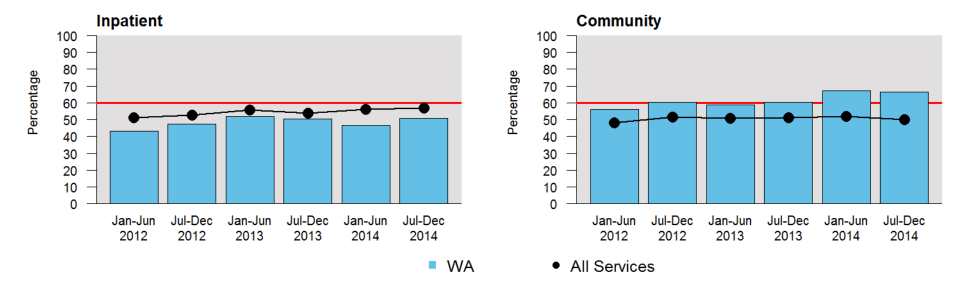




Figure 11 Trends in benchmark 3.3: SAS Patients with absent/mild pain at phase start, remaining absent/mild at phase end by setting

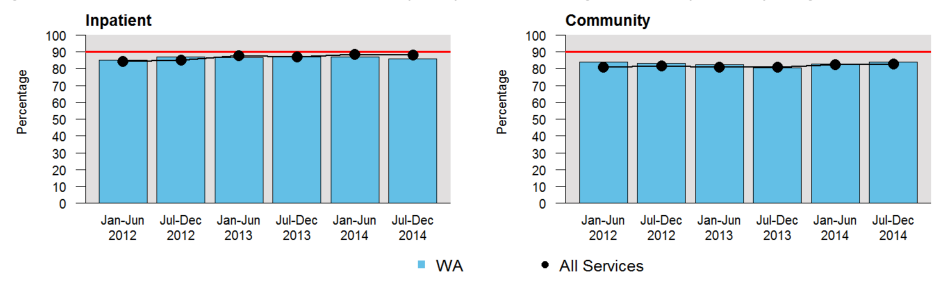
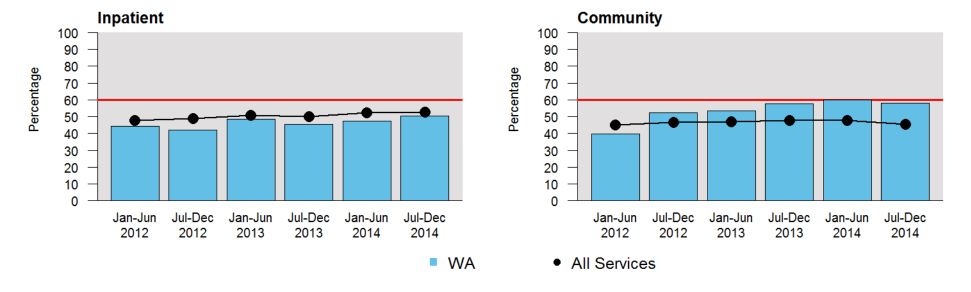


Figure 12 Trends in benchmark 3.4: SAS Patients with moderate/severe pain at phase start, with absent/mild at phase end by setting





2.4 Outcome measure 4 – Change in symptoms relative to the baseline national average (X-CAS)

Outcome measure 4 includes a suite of case-mix adjusted scores used to compare the change in symptoms for similar patients i.e. patients in the same phase who started with the same level of symptom. Eight symptoms are included in this report and the baseline reference period is January to June 2014. The suite of benchmarks included in outcome measure 4 are generally referred to as <u>X-CAS</u> – *CAS* standing for *Case-mix Adjusted Score*, and the *X* to represent that multiple symptoms are included. As X-CAS looks at change in symptom, they are only able to be calculated on phases which ended in phase change or discharge (as the phase end scores are required to determine the change).

Table 7 Summary of outcome measure 4

		V	/A		All Services				
Benchmark: Symptom	X-CAS	N phases included in measure	N phases at or above the baseline	% phases at or above the baseline	X-CAS	N phases included in measure	N phases at or above the baseline	% phases at or above the baseline	
4.1: PCPSS Pain	0.00	9,929	5,615	56.6	0.00	39,811	23,009	57.8	
4.2: Other symptoms	0.04	9,912	6,505	65.6	0.02	39,616	24,870	62.8	
4.3: Family/carer	0.09	9,687	6,534	67.5	0.01	39,198	23,943	61.1	
4.4: Psychological/spiritual	0.01	10,273	5,278	51.4	0.01	40,087	20,819	51.9	
4.5: SAS Pain	0.01	10,227	6,303	61.6	-0.01	38,937	23,873	61.3	
4.6: Nausea	0.06	10,101	8,624	85.4	0.02	38,580	31,707	82.2	
4.7: Breathing Problems	0.11	10,109	7,242	71.6	0.02	38,415	26,699	69.5	
4.8: Bowel Problems	0.11	9,886	7,631	77.2	0.03	38,024	27,132	71.4	

Interpretation hint:

The X-CAS measures are calculated relative to a baseline reference period, which has been updated for this report and is now the period January to June 2014. As a result:

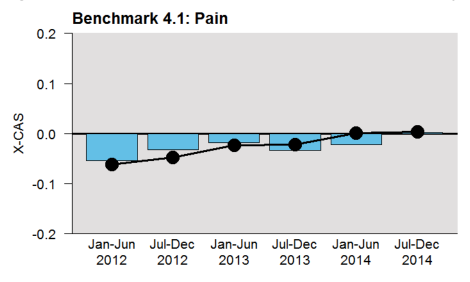
If X-CAS for WA is greater than 0 then on average, your patients' change in symptom was better than similar patients in the baseline reference period.

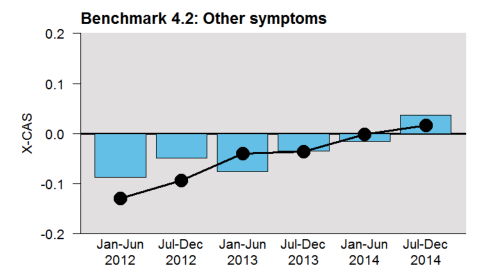
If X-CAS for WA is equal to 0 then on average, your patients' change in symptom was about the same as similar patients in the baseline reference period.

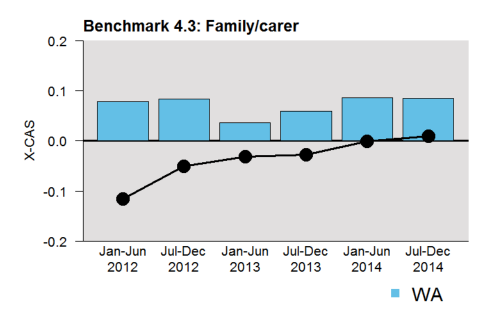
If X-CAS for WA is <u>less than 0</u> then on average, your patients' change in symptom was <u>worse than similar patients</u> in the baseline reference period.

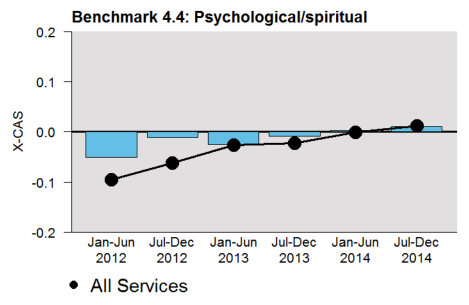


Figure 13 Trends in outcome measure 4 – Palliative Care Problem Severity Score (PCPSS)





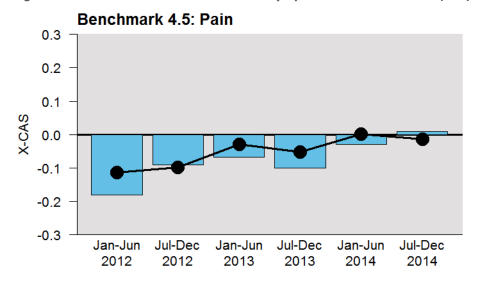


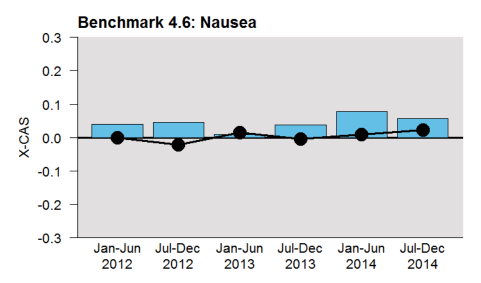


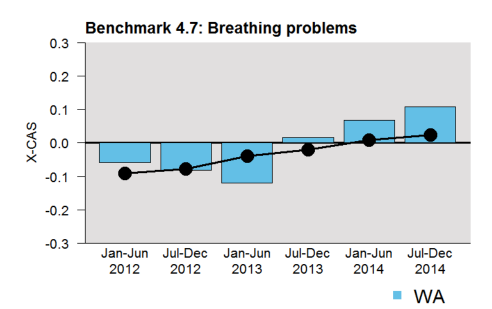
Note: Only services with 10 or more valid assessments are included in the above graphs.

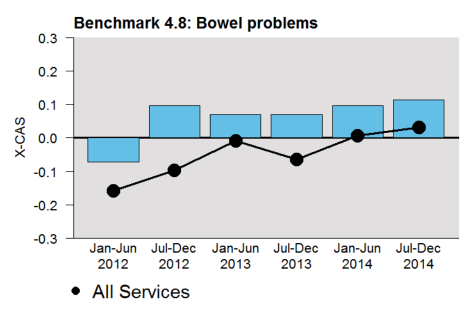


Figure 14 Trends in outcome measure 4 – Symptom Assessment Scale (SAS)









Note: Only services with 10 or more valid assessments are included in the above graphs.



Section 3 Descriptive analysis

This section provides descriptive information of the data submitted by Western Australian services at each of the three levels – patient, episode and phase.

Patient level information describes demographics such as Indigenous status, sex, preferred language and country of birth. This information about the patient provides a context to the episode and phase level information and enhances the meaningfulness of patient outcomes.

Episode level information describes the setting of palliative care service provision. It also includes information relating to the facility/organisation that has referred the patient, how an episode starts/ends and the setting in which the patient died.

Phase level information describes the clinical condition of the patient during the episode, using five clinical assessment tools. These are phase of illness, the patient's functional status and performance, pain and other common symptoms, the patient's psychological/spiritual and family/carer domain.

Summaries of the national data are included for comparative purposes.



3.1 Profile of palliative care patients

PCOC defines a patient as a person for whom a palliative care service accepts responsibility for assessment and/or treatment as evidenced by the existence of a medical record. Family/carers are included in this definition if interventions relating to them are recorded in the patient medical record.

Table 8 shows the Indigenous status for all the patients in Western Australia and nationally.

Table 8 Indigenous status

Indiana a status	W	'A	All Services		
Indigenous status	N	%	N	%	
Aboriginal but not Torres Strait Islander origin	42	1.3	179	1.0	
Torres Strait Islander but not Aboriginal origin	1	0.0	13	0.1	
Both Aboriginal and Torres Strait Islander origin	7	0.2	13	0.1	
Neither Aboriginal nor Torres Strait Islander origin	3,049	94.0	17,739	96.9	
Not stated/inadequately described	143	4.4	366	2.0	
Total	3,242	100.0	18,310	100.0	

Table 9 shows the breakdown of deaths for all patients in Western Australia and nationally for the reporting period. All inpatient deaths are reported in the hospital category while the community deaths are reported in the private residence and residential aged care facility categories.

Table 9 Place of death

Place of death	V	/A	All Services		
Place of death	N	%	N	%	
Private residence	618	44.6	1,834	20.2	
Residential aged care facility	94	6.8	647	7.1	
Hospital	671	48.4	6,507	71.7	
Not stated/inadequately described	2	0.1	88	1.0	
Total	1,385	100.0	9,076	100.0	



The following two tables show the country of birth and the preferred language respectively for all patients in Western Australia and nationally. To allow for comparison with the broader Australian community the list of country of birth in Table 10 is in descending order of the most frequent country of birth according to the 2006 Census (e.g. Italy was the fifth most common country of birth in the 2006 Census). The same approach has been taken with Table 11 (e.g. Greek was the third most frequently spoken language in the 2006 census). All other countries and languages have been grouped together to form the categories 'All other countries' and 'All other languages' respectively.

Table 10 Country of birth

Connetting of himth	W	/A	All Services		
Country of birth	N	%	N	%	
Australia	1,743	53.8	11,458	62.6	
England	512	15.8	1,410	7.7	
New Zealand	88	2.7	363	2.0	
China	19	0.6	170	0.9	
Italy	117	3.6	679	3.7	
Vietnam	20	0.6	138	0.8	
India	45	1.4	139	0.8	
Scotland	79	2.4	280	1.5	
Philippines	9	0.3	76	0.4	
Greece	19	0.6	390	2.1	
Germany	34	1.0	233	1.3	
South Africa	36	1.1	91	0.5	
Malaysia	27	0.8	68	0.4	
Netherlands	49	1.5	194	1.1	
Lebanon	1	0.0	86	0.5	
All other countries	361	11.1	2,270	12.4	
Not stated/inadequately described	83	2.6	265	1.4	
Total	3,242	100.0	18,310	100.0	



Table 11 Preferred language

Duefamadian	W	/A	All Se	rvices
Preferred language	N	%	N	%
English	3,022	93.2	16,528	90.3
Italian	50	1.5	335	1.8
Greek	7	0.2	280	1.5
Chinese ^(a)	17	0.5	147	0.8
Arabic ^(b)	4	0.1	100	0.5
Vietnamese ^(c)	14	0.4	60	0.3
Spanish / Portuguese ^(d)	7	0.2	35	0.2
Filipino / Indonesian ^(e)	5	0.2	19	0.1
German ^(f)	6	0.2	31	0.2
Hindi ^(g)	6	0.2	22	0.1
Croatian / Macedonian ^(h)	20	0.6	116	0.6
Korean	1	0.0	16	0.1
Turkish ⁽ⁱ⁾	1	0.0	32	0.2
Polish ^(j)	7	0.2	30	0.2
Maltese	0	0.0	35	0.2
All other languages	74	2.3	519	2.8
Not stated/inadequately described	1	0.0	5	0.0
Total	3,242	100.0	18,310	100.0

(a) Chinese includes: Cantonese, Hakka, Mandarin, Wu and Min Nan; (b) Middle Eastern Semitic Languages includes: Hebrew, Assyrian Neo-Aramaic, Chaldean Neo-Aramaic, Mandaean (Mandaic); (c) Mon-Khmer includes: Khmer, Mon; (d) Iberian Romance includes: Catalan; (e) Southeast Asian Austronesian Languages includes: Bisaya, Cebuano, Ilokano, Malay, Tetum, Timorese, Tagalog, Acehnese, Balinese, Bikol, Iban, Ilonggo (Hiligaynon), Javanese, Pampangan; (f) German and Related Languages include: Letzeburgish, Yiddish; (g) Indo-Aryan includes: Bengali, Gujarati, Konkani, Marathi, Nepali, Punjabi, Sindhi, Sinhalese, Urdu, Assamese, Dhivehi, Kashmiri, Oriya, Fijian Hindustani; (h) South Slavic includes: Bosnian, Bulgarian, Serbian, Slovene; (i) Turkic includes: Azeri, Tatar, Turkmen, Uygur, Uzbek; (j) West Slavic includes: Czech, Slovak



Table 12 and Table 13 present a breakdown of malignant and non-malignant diagnosis for the patients seen by Western Australian services and at the national level. The primary diagnosis is the principal life limiting illness responsible for the patient requiring palliative care.

The primary diagnosis was not stated for 3 (0.1%) patients in Western Australia and was not stated for 58 (0.3%) patients nationally.

Table 12 Primary diagnosis - malignant

		WA			All Services			
Primary diagnosis	N	% malignant diagnosis	% all diagnosis	N	% malignant diagnosis	% all diagnosis		
Bone and soft tissue	29	1.2	0.9	220	1.6	1.2		
Breast	183	7.6	5.6	1,116	7.9	6.1		
CNS	50	2.1	1.5	281	2.0	1.5		
Colorectal	258	10.7	8.0	1,610	11.4	8.8		
Other GIT	240	10.0	7.4	1,406	10.0	7.7		
Haematological	144	6.0	4.4	837	6.0	4.6		
Head and neck	87	3.6	2.7	784	5.6	4.3		
Lung	617	25.6	19.0	3,083	21.9	16.8		
Pancreas	140	5.8	4.3	898	6.4	4.9		
Prostate	168	7.0	5.2	960	6.8	5.2		
Other urological	123	5.1	3.8	592	4.2	3.2		
Gynaecological	102	4.2	3.1	707	5.0	3.9		
Skin	124	5.2	3.8	528	3.8	2.9		
Unknown primary	77	3.2	2.4	404	2.9	2.2		
Other primary malignancy	62	2.6	1.9	494	3.5	2.7		
Malignant – not further defined	2	0.1	0.1	143	1.0	0.8		
All malignant	2,406	100.0	74.2	14,063	100.0	76.8		



Table 13 Primary diagnosis - non-malignant

		WA		All Services			
Primary diagnosis	N	% non-malignant diagnosis	% all diagnosis	N	% non-malignant diagnosis	% all diagnosis	
Cardiovascular disease	154	18.5	4.8	818	19.5	4.5	
HIV/AIDS	1	0.1	0.0	12	0.3	0.1	
End stage kidney disease	73	8.8	2.3	418	10.0	2.3	
Stroke	61	7.3	1.9	223	5.3	1.2	
Motor neurone disease	26	3.1	0.8	165	3.9	0.9	
Alzheimer's dementia	50	6.0	1.5	157	3.7	0.9	
Other dementia	67	8.0	2.1	242	5.8	1.3	
Other neurological disease	42	5.0	1.3	355	8.5	1.9	
Respiratory failure	126	15.1	3.9	749	17.9	4.1	
End stage liver disease	27	3.2	0.8	159	3.8	0.9	
Diabetes and its complications	2	0.2	0.1	19	0.5	0.1	
Sepsis	29	3.5	0.9	94	2.2	0.5	
Multiple organ failure	37	4.4	1.1	104	2.5	0.6	
Other non-malignancy	126	15.1	3.9	582	13.9	3.2	
Non-malignant – not further defined	12	1.4	0.4	92	2.2	0.5	
All non-malignant	833	100.0	25.7	4,189	100.0	22.9	



3.2 Profile of palliative care episodes

An episode of care is a period of contact between a patient and a palliative care service that is provided by one palliative care service and occurs in one setting – for the purposes of this report, either as an inpatient or community patient.

An episode of palliative care starts on the date when the comprehensive palliative care assessment is undertaken and documented using the five clinical assessment tools.

An episode of palliative care ends when:

- the patient is formally separated from the current setting of care (e.g. from community to inpatient) or
- the patient dies or
- the principal clinical intent of the care changes and the patient is no longer receiving palliative care.

Table 14 below presents the number and percentage of episodes by age group and sex for the patients seen by Western Australian services and at the national level. Age has been calculated as at the beginning of each episode.

Table 14 Age group by sex

	WA				All Services			
Age group	Male		Female		Ma	ale	Female	
	N	%	N	%	N	%	N	%
< 15	7	0.3	4	0.2	31	0.2	28	0.3
15 - 24	17	0.8	9	0.5	42	0.3	42	0.4
25 - 34	11	0.5	7	0.4	90	0.7	103	0.9
35 - 44	51	2.3	107	5.7	279	2.2	398	3.6
45 - 54	114	5.2	148	7.9	767	6.2	943	8.5
55 - 64	325	14.9	341	18.1	1,913	15.4	1,748	15.8
65 - 74	601	27.5	396	21.1	3,374	27.2	2,491	22.6
75 - 84	671	30.7	456	24.3	3,793	30.5	2,868	26.0
85+	389	17.8	411	21.9	2,128	17.1	2,411	21.9
Not stated/inadequately described	0	0.0	0	0.0	0	0.0	0	0.0
Total	2,186	100.0	1,879	100.0	12,417	100.0	11,032	100.0

Note: Records where sex was not stated or inadequately described are excluded from the table.



Referral source refers to the facility or organisation from which the patient was referred for each episode of care. Table 15 presents referral source by setting.

Table 15 Referral source by setting

		Inpa	tient		Community			
Referral source	W	/A	All Services		WA		All Services	
	N	%	N	%	N	%	N	%
Public hospital	993	67.9	6,397	52.3	1,129	43.4	5,458	48.6
Private hospital	75	5.1	1,534	12.5	433	16.6	1,206	10.7
Outpatient clinic	14	1.0	53	0.4	0	0.0	26	0.2
General medical practitioner	32	2.2	386	3.2	792	30.4	1,523	13.6
Specialist medical practitioner	38	2.6	605	4.9	34	1.3	384	3.4
Community-based palliative care agency	287	19.6	2,717	22.2	6	0.2	327	2.9
Community-based service	3	0.2	54	0.4	7	0.3	168	1.5
Residential aged care facility	7	0.5	99	0.8	175	6.7	927	8.3
Self, carer(s), family or friends	8	0.5	156	1.3	9	0.3	371	3.3
Other	2	0.1	163	1.3	13	0.5	302	2.7
Not stated/inadequately described	4	0.3	60	0.5	4	0.2	533	4.7
Total	1,463	100.0	12,224	100.0	2,602	100.0	11,225	100.0



Table 16 provides a summary of the time between referral to first contact by setting of care. The time from referral to first contact is calculated as the time from the date of referral received to either the date of first contact (if provided) or the episode start date.

Table 16 Referral to first contact by episode setting

		Inpa	tient		Community				
Time (in days)	WA		All Services		WA		All Services		
	N	%	N	%	N	%	N	%	
Same day or following day	1,361	93.0	11,383	93.1	1,379	53.0	5,932	52.9	
2-7 days	93	6.4	716	5.9	1,072	41.2	3,711	33.1	
8-14 days	5	0.3	61	0.5	114	4.4	883	7.9	
Greater than 14 days	4	0.3	61	0.5	37	1.4	697	6.2	
Average	1.2	na	1.2	na	2.4	na	2.8	na	
Median	1	na	1	na	1	na	1	na	

Note: Episodes where referral date was not recorded are excluded from the table. In addition, all records where time from referral to first contact was greater than 90 days were considered to be atypical and were assumed to equal 90 days for the purpose of calculating the average and median time.



Table 17 gives a summary of the length of episode for patients in Western Australia and nationally. Table 18 details the length of episode by setting. The length of episode is calculated as the number of days between the episode start date and the episode end date. Bereavement phases are excluded from the calculation and episodes that remain open at the end of the reporting period (and hence do not have an episode end date) are also excluded.

Table 17 Length of episode (in days) summary by setting

Length of episode	Inpat	tient	Community		
Length of episode	WA	All Services	WA	All Services	
Average length of episode	8.7	10.6	38.3	35.8	
Median length of episode	5.0	6.0	26.0	24.0	

Note: Records where length of episode was greater than 180 days were considered to be atypical and are excluded from the average calculations. Only episodes ending during the reporting period are included.

Table 18 Length of episode (in days) by setting

		Inpa	tient			Comn	nunity		
Length of episode	W	/A	All Se	All Services		WA		All Services	
	N	%	N	%	N	%	N	%	
Same day	110	7.7	681	5.7	40	1.9	799	8.0	
1-2 days	311	21.7	2,304	19.1	134	6.3	597	6.0	
3-4 days	235	16.4	1,777	14.8	119	5.6	501	5.0	
5-7 days	258	18.0	2,044	17.0	165	7.7	745	7.5	
8-14 days	276	19.2	2,586	21.5	280	13.1	1,234	12.3	
15-21 days	119	8.3	1,106	9.2	222	10.4	906	9.1	
22-30 days	67	4.7	729	6.1	196	9.2	871	8.7	
31-60 days	50	3.5	659	5.5	406	19.0	1,736	17.4	
61-90 days	6	0.4	106	0.9	200	9.4	834	8.3	
Greater than 90 days	3	0.2	50	0.4	372	17.4	1,776	17.8	
Total	1,435	100.0	12,042	100.0	2,134	100.0	9,999	100.0	

Note: Only episodes ending during the reporting period are included.



Table 19 How episodes start – inpatient setting

Fuinada start mada	W	/A	All Services		
Episode start mode	N	%	N	%	
Admitted from community*	995	68.0	7,522	61.5	
Admitted from another hospital	380	26.0	3,088	25.3	
Admitted from acute care in another ward	71	4.9	1,343	11.0	
Change from acute care to palliative care – same ward	8	0.5	173	1.4	
Other**	8	0.5	92	0.8	
Not stated/inadequately described	1	0.1	6	0.0	
Total	1,463	100.0	12,224	100.0	

^{*} includes: admitted from usual accommodation, admitted from other than usual accommodation

Table 20 How episodes end – inpatient setting

Episode end mode	WA		All Services	
	N	%	N	%
Discharged to community*	556	38.7	4,331	36.0
Discharged to another hospital	96	6.7	877	7.3
Death	671	46.8	6,507	54.0
Change from palliative care to acute care**	4	0.3	74	0.6
Change in sub-acute care type	0	0.0	38	0.3
End of consultative episode – inpatient episode ongoing	26	1.8	98	0.8
Other	80	5.6	111	0.9
Not stated/inadequately described	2	0.1	6	0.0
Total	1,435	100.0	12,042	100.0

Note: Only episodes ending during the reporting period are included.

^{**} includes: change of sub-acute/non-acute care type and other categories

^{*} includes: discharged to usual accommodation, discharged to other than usual accommodation

^{**} includes: change from palliative care to acute care – different ward, change from palliative care to acute care – same ward



Table 21 How episodes start – community setting

Full and a start mode	W	'A	All Services		
Episode start mode	N	%	N	%	
Admitted from inpatient palliative care	1,714	65.9	4,137	36.9	
Other*	887	34.1	7,035	62.7	
Not stated/inadequately described	1	0.0	53	0.5	
Total	2,602	100.0	11,225	100.0	

^{*}includes: patient was not transferred from being an overnight patient

Table 22 How episodes end – community setting

Fuinada and mada	W	/A	All Se	rvices
Episode end mode	N	%	N	%
Admitted for inpatient palliative care	145	6.8	2,792	27.9
Admitted for inpatient acute care	976	45.7	2,677	26.8
Admitted to another palliative care service	60	2.8	133	1.3
Admitted to primary health care	9	0.4	611	6.1
Discharged/case closure	230	10.8	1,065	10.7
Death	714	33.5	2,569	25.7
Other	0	0.0	144	1.4
Not stated/inadequately described	0	0.0	8	0.1
Total	2,134	100.0	9,999	100.0

Note: Only episodes ending during the reporting period are included.



3.3 Profile of palliative care phases

The palliative care phase type describes the stage of the patient's illness and provides a clinical indication of the level of care a patient requires. The palliative care phase is determined by a holistic clinical assessment which considers the needs of the patients and their family and carers. A patient may move back and forth between the stable, unstable, deteriorating and terminal phase types and these may occur in any sequence. See Appendix D for more information on the definition of palliative care phase.

The clinical assessments are assessed daily (or at each visit) and are reported on admission, when the phase changes and at discharge.

Table 23 Number of phases by phase type and setting

		Inpa	tient		Community					
Phase type	hase type WA		All Services		W	/A	All Services			
	N	%	N	%	N	%	N	%		
Stable	733	24.6	7,330	25.8	3,408	36.0	9,334	37.2		
Unstable	478	16.0	6,544	23.0	764	8.1	3,180	12.7		
Deteriorating	1,182	39.7	8,978	31.6	4,496	47.5	10,424	41.6		
Terminal	587	19.7	5,557	19.6	798	8.4	2,120	8.5		
Total	2,980	100.0	28,409	100.0	9,466	100.0	25,058	100.0		

Note: Bereavement phases have been excluded due to inconsistent data collection and bereavement practices. Bereavement phases are not included in the total phases count.

Table 24 Average phase length (in days) by phase type and setting

Dhaca tura	Inpa	tient	Community			
Phase type	WA	All Services	WA	All Services		
Stable	5.9	6.9	13.9	19.8		
Unstable	1.7	2.3	1.4	4.4		
Deteriorating	4.9	5.5	8.6	12.7		
Terminal	2.3	2.1	2.6	3.0		

Note: Phase records where phase length was greater than 90 days were considered to be atypical and are excluded from the average calculations.



Table 25 presents information relating to the manner in which stable phases ended, both for Western Australian services and nationally. A stable phase will end if a patient moves into a different phase (phase change), is discharged or dies. Figure 15 summarises the movement of patients out of the stable phase for the inpatient and community settings. This movement from one phase to another is referred to as phase progression. The phase progression information is derived by PCOC.

Similar information is presented for the unstable (Table 26, Figure 16), deteriorating (Table 27, Figure 17) and terminal (Table 28, Figure 18) phases on the following pages.

Table 25 How stable phases end - by setting

		Inpa	tient		Community				
How stable phases end	WA		All Servi	All Services			All Services		
	N	%	N	%	N	%	N	%	
Patient moved into another phase	311	42.4	3,681	50.2	2,864	84.0	6,076	65.1	
Discharge/case closure	412	56.2	3,539	48.3	491	14.4	2,972	31.8	
Died	10	1.4	103	1.4	53	1.6	247	2.6	
Not stated/inadequately described	0	0.0	7	0.1	0	0.0	39	0.4	
Total	733	100.0	7,330	100.0	3,408	100.0	9,334	100.0	

Figure 15 Stable phase progression

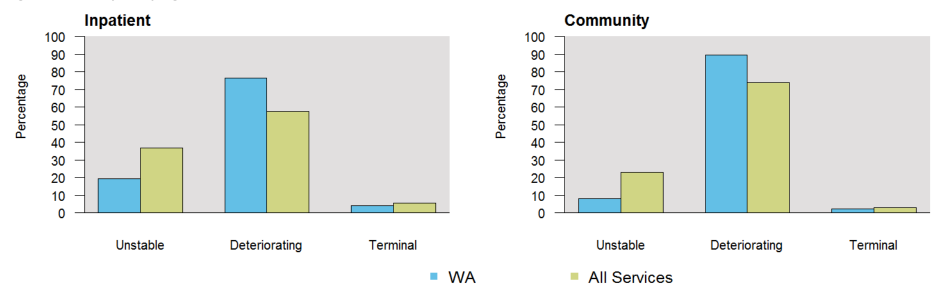




Table 26 How <u>unstable</u> phases end – by setting

		Inpa	tient		Community				
How unstable phases end	WA		All Servi	All Services			All Services		
	N	%	N	%	N	%	N	%	
Patient moved into another phase	430	90.0	6,022	92.0	595	77.9	2,147	67.5	
Discharge/case closure	35	7.3	367	5.6	150	19.6	963	30.3	
Died	13	2.7	148	2.3	19	2.5	63	2.0	
Not stated/inadequately described	0	0.0	7	0.1	0	0.0	7	0.2	
Total	478	100.0	6,544	100.0	764	100.0	3,180	100.0	

Figure 16 Unstable phase progression

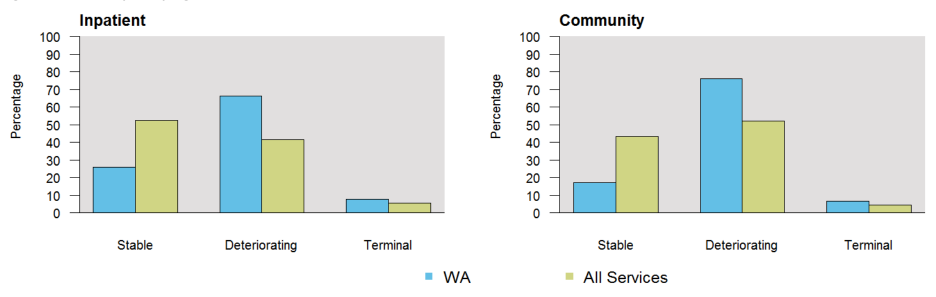




Table 27 How <u>deteriorating</u> phases end – by setting

		Inpa	tient		Community				
How deteriorating phases end	WA		All Servi	All Services			All Services		
	N	%	N	%	N	%	N	%	
Patient moved into another phase	777	65.7	6,444	71.8	3,603	80.1	6,442	61.8	
Discharge/case closure	278	23.5	1,501	16.7	733	16.3	3,209	30.8	
Died	126	10.7	1,027	11.4	160	3.6	762	7.3	
Not stated/inadequately described	1	0.1	6	0.1	0	0.0	11	0.1	
Total	1,182	100.0	8,978	100.0	4,496	100.0	10,424	100.0	

Figure 17 Deteriorating phase progression

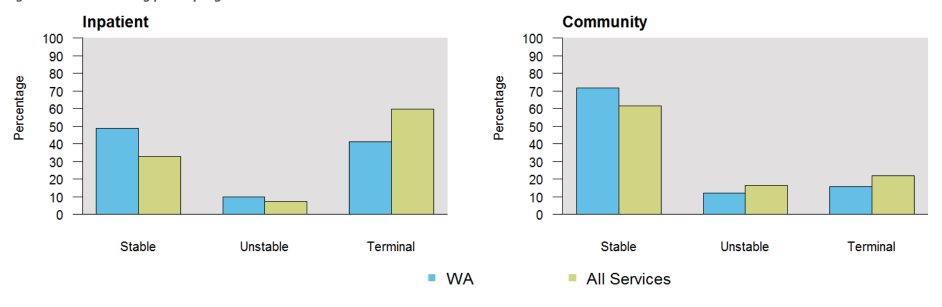
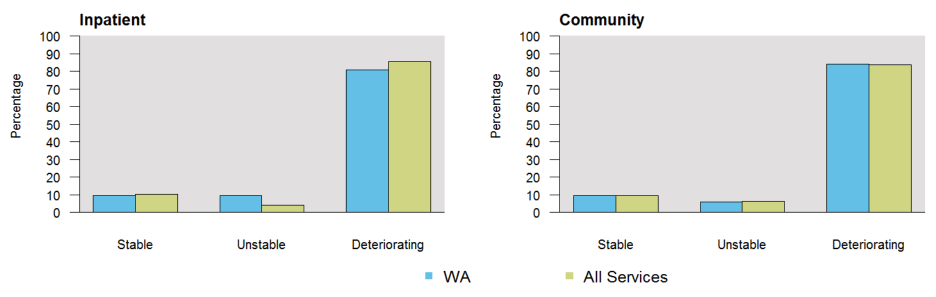




Table 28 How <u>terminal</u> phases end – by setting

		Inpa	tient		Community				
How terminal phases end	WA		All Servi	All Services			All Services		
	N	%	N	%	N	%	N	%	
Patient moved into another phase	21	3.6	215	3.9	229	28.7	330	15.6	
Discharge/case closure	38	6.5	106	1.9	46	5.8	241	11.4	
Died	528	89.9	5,236	94.2	523	65.5	1,548	73.0	
Not stated/inadequately described	0	0.0	0	0.0	0	0.0	1	0.0	
Total	587	100.0	5,557	100.0	798	100.0	2,120	100.0	

Figure 18 Terminal phase progression





The Palliative Care Problem Severity Score (PCPSS) is a clinician rated screening tool to assess the overall severity of problems within four key palliative care domains (pain, other symptoms, psychological/spiritual and family/carer). The ratings are: 0 - absent, 1 - mild, 2 - moderate and 3 - severe.

Table 29 and Table 30 show the percentage scores for the inpatient and community settings, respectively, for both Western Australian services and nationally.

Table 29 Profile of PCPSS at beginning of phase by phase type – inpatient setting (percentages)

Disease towns			N	/A			All Se	rvices	
Phase type	Problem severity	Absent	Mild	Moderate	Severe	Absent	Mild	Moderate	Severe
	Pain	51.2	39.5	8.6	0.7	48.8	37.6	11.0	2.5
Ctoble	Other symptoms	34.3	48.9	15.3	1.5	25.8	51.9	19.1	3.3
Stable	Psychological/spiritual	24.6	58.8	15.1	1.5	32.7	52.6	12.2	2.5
	Family/carer	34.0	44.1	18.1	3.8	40.6	43.1	12.8	3.4
	Pain	27.6	24.9	25.3	22.2	30.5	30.9	25.6	12.9
Haatabla	Other symptoms	20.9	23.8	33.6	21.8	13.8	34.1	38.3	13.8
Unstable	Psychological/spiritual	12.3	42.9	32.8	12.1	23.8	44.1	24.8	7.4
	Family/carer	17.0	33.7	36.2	13.1	26.1	40.8	24.2	8.9
	Pain	34.8	32.2	26.4	6.6	38.4	35.9	19.7	5.9
Deterioration	Other symptoms	20.6	30.7	37.0	11.8	15.3	40.8	33.5	10.4
Deteriorating	Psychological/spiritual	18.2	46.2	29.7	5.9	24.9	47.8	21.6	5.7
	Family/carer	19.8	40.2	32.2	7.8	27.6	41.6	23.0	7.8
Terminal	Pain	52.6	27.7	14.7	5.0	48.1	32.7	14.0	5.1
	Other symptoms	44.3	27.1	18.0	10.6	33.6	35.0	21.8	9.6
	Psychological/spiritual	52.0	27.4	15.4	5.1	51.1	31.8	12.5	4.6
	Family/carer	15.0	28.2	42.1	14.7	21.9	35.6	30.2	12.4



Table 30 Profile of PCPSS at beginning of phase by phase type –community setting (percentages)

Dhana tuna			N	<i>I</i> A			All Services				
Phase type	Problem severity	Absent	Mild	Moderate	Severe	Absent	Mild	Moderate	Severe		
	Pain	43.7	51.8	4.3	0.1	41.0	50.8	7.6	0.6		
Ctoblo	Other symptoms	14.8	72.7	12.3	0.1	14.9	66.3	17.5	1.3		
Stable	Psychological/spiritual	35.6	57.5	6.7	0.3	30.2	58.5	10.2	1.1		
	Family/carer	43.0	50.5	6.3	0.2	31.1	53.6	13.5	1.7		
	Pain	20.3	26.0	29.6	24.1	18.3	28.7	33.6	19.4		
Unstable	Other symptoms	4.3	27.5	49.8	18.3	5.4	28.0	48.4	18.3		
Unstable	Psychological/spiritual	11.9	43.6	38.7	5.8	12.1	45.6	34.4	7.9		
	Family/carer	19.9	35.4	39.5	5.3	14.1	34.8	39.3	11.8		
	Pain	30.7	48.7	18.7	2.0	28.4	49.3	19.5	2.8		
Deterieration	Other symptoms	7.8	53.4	36.7	2.1	7.0	48.7	39.2	5.1		
Deteriorating	Psychological/spiritual	22.5	57.9	18.0	1.6	18.4	58.0	20.8	2.9		
	Family/carer	28.9	49.9	19.6	1.6	19.1	47.9	28.5	4.5		
	Pain	40.5	41.7	15.3	2.5	35.7	44.4	15.8	4.1		
Torminal	Other symptoms	25.8	41.4	28.3	4.5	20.9	40.9	29.6	8.6		
Terminal	Psychological/spiritual	46.2	40.8	11.6	1.4	40.0	42.0	15.0	3.1		
	Family/carer	19.4	44.9	32.6	3.1	12.6	40.1	36.8	10.5		

The Symptom Assessment Scale (SAS) is a patient rated (or proxy) assessment tool and reports a level of distress using a numerical rating scale from 0 - no problems to 10 - worst possible problems. The SAS reports on seven symptoms, these being difficulty sleeping, appetite problems, nausea, bowel problems, breathing problems, fatigue and pain. It provides a clinical picture of these seven symptoms from the patient's perspective. The SAS scores are grouped in Table 31 and Table 32 on the following pages using the same categories as the PCPSS i.e. absent (0), mild (1-3), moderate (4-7) and severe (8-10). Additional information on the SAS profile by phase can be found in Appendix B.



Table 31 Profile of SAS scores at beginning of phase by phase type – inpatient setting (percentages)

Phase type	e of SAS scores at beginn			VA			All Se	ervices	
Phase type	Symptom distress	0 (Absent)	1-3 (Mild)	4-7 (Moderate)	8-10 (Severe)	0 (Absent)	1-3 (Mild)	4-7 (Moderate)	8-10 (Severe)
	Difficulty sleeping	62.3	20.8	13.1	3.8	67.8	18.1	11.5	2.6
	Appetite problems	56.4	19.7	18.9	5.1	55.0	23.3	17.7	4.0
	Nausea	80.3	11.9	6.8	1.0	79.7	13.2	6.0	1.0
Stable	Bowel problems	60.7	23.4	13.0	2.9	61.8	21.7	13.5	3.1
	Breathing problems	62.8	22.3	11.8	3.1	64.8	18.2	13.6	3.4
	Fatigue	27.3	22.3	36.4	14.0	27.3	25.0	38.4	9.3
	Pain	44.3	32.7	20.1	3.0	46.3	31.9	18.9	3.0
	Difficulty sleeping	45.8	17.4	23.7	13.1	57.6	17.5	18.7	6.1
	Appetite problems	45.0	17.3	24.0	13.6	41.8	22.4	25.8	10.0
	Nausea	63.4	16.7	12.0	7.9	68.3	14.3	12.2	5.3
Unstable	Bowel problems	51.1	18.3	21.1	9.5	50.9	21.2	20.7	7.3
	Breathing problems	46.3	17.4	21.0	15.3	55.2	17.2	18.6	8.9
	Fatigue	21.9	13.6	38.7	25.7	21.2	17.1	43.4	18.3
	Pain	24.2	16.5	35.8	23.5	30.9	24.6	31.8	12.7
	Difficulty sleeping	57.5	19.6	18.1	4.7	67.5	15.3	14.0	3.2
	Appetite problems	51.6	17.3	23.1	8.0	50.9	19.0	22.1	7.9
	Nausea	73.7	14.2	10.0	2.1	76.2	12.3	9.2	2.3
Deteriorating	Bowel problems	55.5	20.1	18.1	6.3	59.5	20.3	15.9	4.3
	Breathing problems	51.2	21.3	19.3	8.2	55.5	18.1	18.6	7.8
	Fatigue	25.2	13.5	36.9	24.3	24.8	14.7	41.0	19.5
	Pain	33.5	27.4	31.5	7.6	38.2	29.1	26.7	6.0
	Difficulty sleeping	86.3	5.8	6.2	1.7	90.0	4.8	4.2	1.1
	Appetite problems	89.5	2.9	3.1	4.5	87.8	3.7	5.0	3.5
	Nausea	92.0	4.1	2.4	1.5	93.0	3.7	2.4	0.9
Terminal	Bowel problems	87.5	4.5	5.8	2.2	84.4	7.7	5.8	2.2
	Breathing problems	65.1	13.5	15.4	6.0	67.6	12.4	13.4	6.6
	Fatigue	78.8	1.5	8.2	11.5	71.0	4.7	12.0	12.3
	Pain	57.3	20.3	17.6	4.8	56.3	22.4	16.9	4.3



Table 32 Profile of SAS scores at beginning of phase by phase type –community setting (percentages)

Dhana tuun			V	VA			All Se	ervices	
Phase type	Symptom distress	0 (Absent)	1-3 (Mild)	4-7 (Moderate)	8-10 (Severe)	0 (Absent)	1-3 (Mild)	4-7 (Moderate)	8-10 (Severe)
	Difficulty sleeping	76.2	18.0	5.4	0.4	63.5	25.9	9.6	0.9
	Appetite problems	63.8	26.1	9.6	0.5	48.4	33.3	16.4	1.9
	Nausea	89.6	9.6	0.8	0.0	80.5	15.9	3.3	0.3
Stable	Bowel problems	79.5	17.3	3.0	0.2	67.7	24.3	7.1	0.9
	Breathing problems	62.1	27.6	9.9	0.4	54.4	30.2	13.6	1.7
	Fatigue	19.3	32.9	45.4	2.4	15.6	34.3	44.6	5.5
	Pain	51.5	39.7	8.5	0.3	44.3	41.8	12.8	1.1
	Difficulty sleeping	53.7	17.8	21.9	6.5	44.7	25.6	23.8	5.9
	Appetite problems	48.5	18.8	24.5	8.3	34.9	26.4	29.5	9.2
	Nausea	61.2	12.2	18.5	8.2	61.0	17.6	15.1	6.2
Unstable	Bowel problems	65.2	17.0	12.8	4.9	52.9	26.4	15.4	5.3
	Breathing problems	58.5	21.4	14.6	5.5	47.2	26.2	20.0	6.6
	Fatigue	12.0	14.8	55.5	17.8	10.0	18.2	52.6	19.2
	Pain	23.7	19.8	31.1	25.3	20.3	24.1	35.9	19.7
	Difficulty sleeping	66.9	19.9	11.9	1.3	57.0	27.5	13.5	2.0
	Appetite problems	53.0	24.2	20.0	2.8	39.0	30.4	25.7	5.0
	Nausea	80.9	12.4	6.1	0.6	73.0	18.7	7.3	1.1
Deteriorating	Bowel problems	71.4	18.8	9.1	0.8	61.7	25.9	10.6	1.7
	Breathing problems	56.4	25.8	16.4	1.4	47.6	30.7	18.7	3.1
	Fatigue	14.2	20.7	57.7	7.3	10.6	22.3	54.5	12.7
	Pain	37.7	36.8	22.4	3.0	32.3	40.9	23.2	3.6
	Difficulty sleeping	77.8	10.8	9.9	1.5	75.7	13.6	8.4	2.2
	Appetite problems	89.3	3.0	4.3	3.4	78.1	6.6	7.3	8.0
	Nausea	88.9	4.1	6.4	0.6	85.0	8.9	4.9	1.2
Terminal	Bowel problems	85.9	8.2	5.5	0.4	74.1	15.7	8.4	1.8
	Breathing problems	59.7	20.4	16.4	3.5	55.5	22.8	17.0	4.6
	Fatigue	70.0	2.9	12.2	14.9	57.4	5.0	13.7	23.9
	Pain	45.8	31.4	19.6	3.1	40.9	35.5	19.6	4.0



The Australia-modified Karnofsky Performance Status (AKPS) is a measure of the patient's overall performance status or ability to perform their activities of daily living. It is a single score between 0 and 100 assigned by a clinician based on observations of a patient's ability to perform common tasks relating to activity, work and self-care. Table 33 shows the data for the AKPS at phase start.

Table 33 Australia-modified Karnofsky Performance Status (AKPS) at phase start by setting

		Inpa	ntient		Community				
AKPS assessment	WA		All Services		WA		All Services		
	N	%	N	%	N	%	N	%	
10 - Comatose or barely rousable	333	11.2	2,907	10.2	301	3.2	822	3.3	
20 - Totally bedfast and requiring extensive nursing care	609	20.4	6,243	22.0	959	10.1	2,428	9.7	
30 - Almost completely bedfast	404	13.6	3,860	13.6	557	5.9	1,682	6.7	
40 - In bed more than 50% of the time	456	15.3	5,151	18.1	897	9.5	3,161	12.6	
50 - Requires considerable assistance	606	20.3	4,855	17.1	1,935	20.4	5,864	23.4	
60 - Requires occasional assistance	430	14.4	2,947	10.4	2,769	29.3	6,317	25.2	
70 - Cares for self	100	3.4	706	2.5	1,760	18.6	3,322	13.3	
80 - Normal activity with effort	15	0.5	215	0.8	239	2.5	699	2.8	
90 - Able to carry on normal activity; minor signs or symptoms	6	0.2	56	0.2	35	0.4	155	0.6	
100 - Normal; no complaints; no evidence of disease	0	0.0	1	0.0	1	0.0	13	0.1	
Not stated/inadequately described	21	0.7	1,468	5.2	13	0.1	595	2.4	
Total	2,980	100.0	28,409	100.0	9,466	100.0	25,058	100.0	

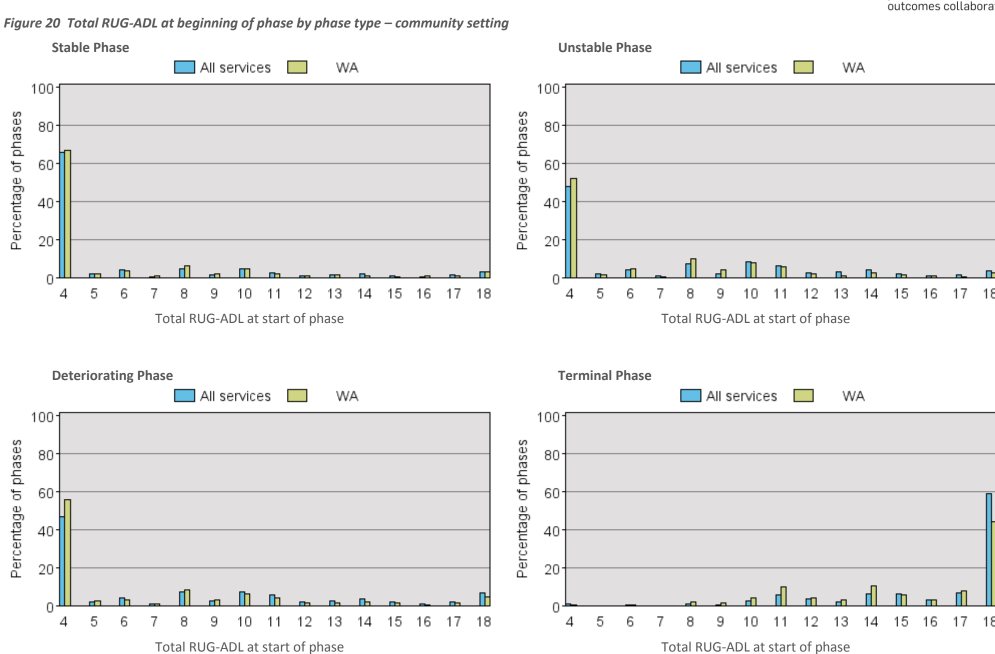
The Resource Utilisation Groups – Activities of Daily Living (RUG-ADL) consists of four items (bed mobility, toileting, transfers and eating) and assesses the level of functional dependence. The RUG-ADL are assessed daily (or at each visit) and are reported on admission, when the phase changes and at discharge. Figure 19 and Figure 20 on the following two pages summarise the total RUG-ADL at the beginning of each phase for inpatients and community patients. The total score on the RUG-ADL ranges from a minimum of 4 (lowest level of functional dependency) to a maximum of 18 (highest level of functional dependency).

AKPS & RUG-ADL can be used together to provide a profile of both patient dependency, equipment requirements, need for allied health referrals and carer burden/respite requirements.



Figure 19 Total RUG-ADL at beginning of phase by phase type – inpatient setting **Stable Phase Unstable Phase** All services WA All services WΑ 100 100 Percentage of phases Percentage of phases 80 80 60 60 40 40 20 20 n 14 15 16 17 18 5 6 10 12 13 5 6 11 12 13 14 15 16 Total RUG-ADL at start of phase Total RUG-ADL at start of phase **Deteriorating Phase Terminal Phase** All services All services WA WA 100 100 Percentage of phases Percentage of phases 80 80 60 60 40 40 20 20 16 17 18 18 5 10 12 13 14 4 8 9 13 17 Total RUG-ADL at start of phase Total RUG-ADL at start of phase







Appendix A Summary of data included in this report

A1 Data summary

During the reporting period, data were provided for a total of 18,310 patients who between them had 23,449 episodes of care and 53,467 palliative care phases. These total numbers are determined by a data scoping method. This method looks at the phase level data first and includes all phases that ended within the current reporting period. The associated episodes and patients are then determined (Appendix B contains a more detailed explanation of this process). Table 34 shows the number of patients, episodes and phases included in this report – both for Western Australian services and nationally.

A consequence of the data scoping method is that it is likely that not all phases related to a particular episode are included in this report. Hence, the average number of phases per episode calculation shown in Table 34 may be an underestimate (due to episodes that cross-over 2 or more reporting periods) as it only includes phases that ended within the current reporting period.

Table 34 Number and percentage of patients, episodes and phases by setting

	Inpa	tient	Comn	nunity	Total		
	WA All Services		WA	All Services	WA	All Services	
Number of patients*	1,272	10,311	2,044	8,963	3,242	18,310	
Number of episodes	1,463	12,224	2,602	11,225	4,065	23,449	
Number of phases**	2,980	28,409	9,466	25,058	12,446	53,467	
Percentage of patients*	39.2	56.3	63.0	49.0	100	100	
Percentage of episodes	36.0	52.1	64.0	47.9	100	100	
Percentage of phases	23.9	53.1	76.1	46.9	100	100	
Average number of phases per episode***	2.0	2.3	3.3	2.0	2.7	2.2	

^{*} Patients seen in both settings are only counted once in the total column and hence numbers/percentages may not add to the total.

^{**} Bereavement phases are excluded from this count.

^{***} Average number of phases per episode is only calculated for closed episodes that started and ended within the reporting period and excludes bereavement phases.



Table 35 shows the number of completed episodes and phases by setting for each month in the current reporting period for Western Australian services. This table allows a service to identify any change in patient numbers during the reporting period.

Table 35 Number of completed episodes and phases by month and setting

		Jul	Aug	Sep	Oct	Nov	Dec
Innations	No. of completed episodes	242	255	260	239	232	207
Inpatient	No. of completed phases	543	533	526	479	481	418
Community	No. of completed episodes	347	319	330	404	376	358
Community	No. of completed phases	1,614	1,493	1,604	1,557	1,538	1,660



A2 Data item completion

As shown in Table 36, Table 37 and Table 38 below, the rate of data completion is very high. In reviewing these tables, it is important to note that in some cases some data items are not required to be completed. For example, place of death is only required for patients who have died. Hence the complete column in the following tables only refers to the percentage of complete records where the data item was relevant.

PCOC strongly encourages services to complete and submit the whole data set on every patient as non-completion may result in services being excluded from relevant benchmarking activities or erroneous conclusions being drawn. Low completion of data items may also distort percentages and graphs in some sections.

Table 36 Item completion (per cent complete) - patient level

Data item	WA	All Services
Date of birth	100.0	100.0
Sex	100.0	100.0
Indigenous status	95.6	98.0
Country of birth	97.4	98.5
Preferred language	100.0	100.0
Primary diagnosis	99.9	99.7

Note: This table is not split by setting to be consistent with the patient level analysis throughout this report.

Table 37 Item completion by setting (per cent complete) - episode level

	Inpa	tient	Comn	nunity	Total		
Data item	WA	All Services	WA	All Services	WA	All Services	
Date of first contact	100.0	100.0	100.0	100.0	100.0	100.0	
Referral date	100.0	100.0	100.0	100.0	100.0	100.0	
Referral source	99.7	99.5	99.8	95.3	99.8	97.5	
Date ready for care	94.7	94.7	100.0	100.0	98.1	97.2	
Mode of episode start	99.9	100.0	100.0	99.5	100.0	99.7	
Accommodation at episode start	99.7	99.9	99.8	96.3	99.8	97.7	
Episode end date*	99.5	99.8	87.3	92.2	91.7	96.2	
Mode of episode end	99.9	100.0	100.0	99.9	99.9	99.9	
Accommodation at episode end	97.5	98.6	100.0	91.8	98.4	96.5	
Place of death	na	na	99.7	96.6	99.7	96.6	

Episode end date item completion may be affected by open episodes.



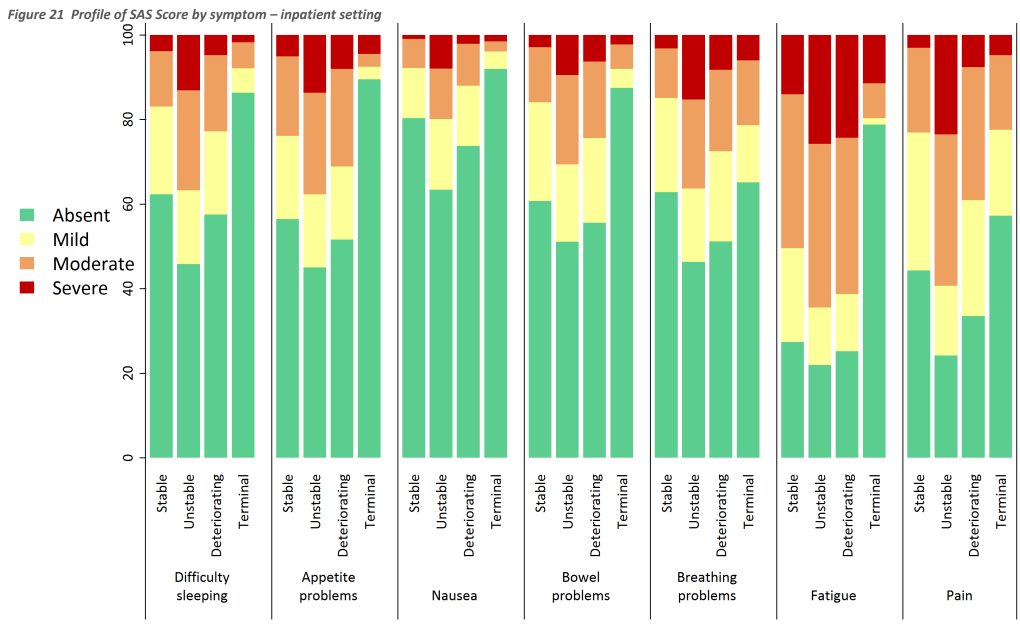
Table 38 Item completion by setting (per cent complete) - phase level

	Cula Catamani	At phase start					At discharge						
	Sub-Category	Inpatient		Comr	nunity	Total		Inpatient		Community		Total	
Data item (where a	(where applicable)	WA	All Services	WA	All Services	WA	All Services	WA	All Services	WA	All Services	WA	All Services
	Bed mobility	99.8	99.7	99.0	97.4	99.2	98.6	91.2	92.3	55.6	63.2	68.1	75.7
DUC ADI	Toileting	99.8	99.7	98.7	97.3	99.0	98.6	91.2	92.3	55.5	63.2	68.0	75.6
RUG-ADL	Transfers	99.8	99.7	97.4	96.7	98.0	98.3	91.2	92.3	55.5	63.2	68.0	75.6
	Eating	99.7	99.5	94.7	95.6	95.9	97.7	91.2	92.3	54.2	62.8	67.1	75.4
	Pain	82.7	97.8	99.9	97.5	95.8	97.6	82.0	91.1	55.9	62.9	65.0	75.0
PCPSS	Other symptom	82.3	97.6	99.8	96.7	95.7	97.2	81.7	91.1	56.0	62.5	65.0	74.7
	Psychological/spiritual	99.0	99.4	99.9	97.2	99.7	98.4	90.0	92.2	56.0	62.8	67.9	75.4
	Family/carer	79.8	97.3	99.5	96.0	94.8	96.7	60.3	88.0	56.0	62.2	57.5	73.2
	Difficulty sleeping	98.8	92.9	98.3	93.8	98.4	93.3	89.1	81.3	55.4	60.4	67.2	69.3
	Appetite problems	98.4	93.1	96.5	95.0	97.0	94.0	89.4	81.5	54.4	61.7	66.7	70.2
CAC	Nausea	99.0	93.2	99.0	96.4	99.0	94.7	90.0	81.7	55.7	62.4	67.7	70.6
SAS	Bowel problems	98.4	93.0	97.7	95.0	97.9	93.9	89.8	81.5	55.2	61.3	67.3	69.9
	Breathing problems	99.1	93.2	99.0	96.0	99.0	94.5	90.2	81.6	55.8	62.0	67.8	70.4
	Fatigue	98.7	93.2	98.0	96.1	98.2	94.5	89.6	81.6	55.3	62.3	67.3	70.5
	Pain	99.3	93.2	99.6	97.3	99.5	95.1	90.0	81.6	55.8	63.0	67.8	71.0
AKPS	-	99.3	94.8	99.9	97.6	99.7	96.1	90.8	89.7	56.0	63.2	68.2	74.5

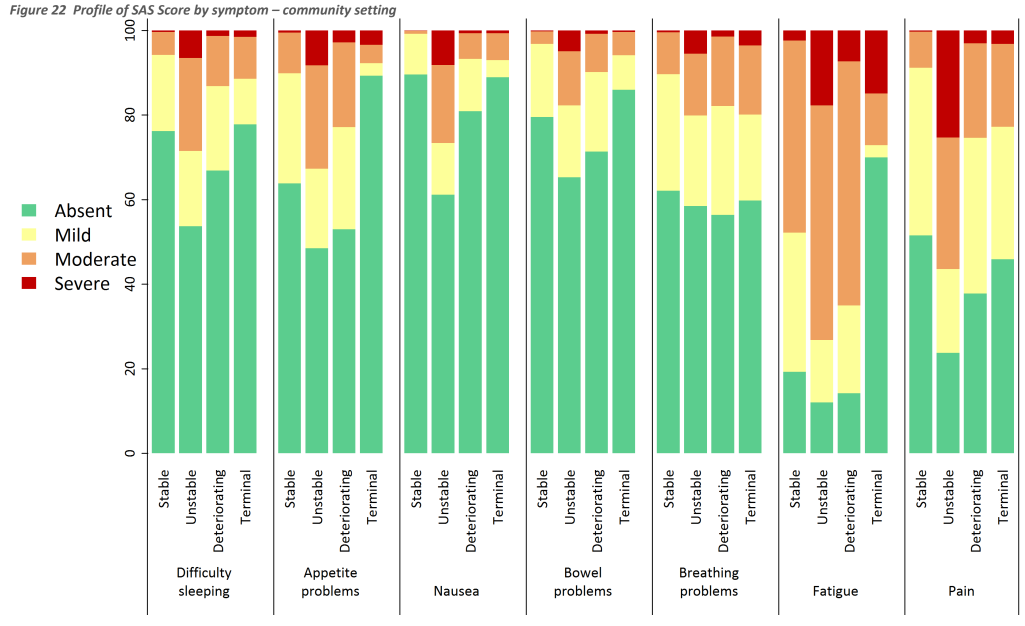
	Inpa	itient	Comn	nunity	Total	
Data item	WA	All Services	WA	All Services	WA	All Services
Phase End Reason	100.0	99.9	100.0	99.7	100.0	99.8



Appendix B Additional information on profile of SAS scores





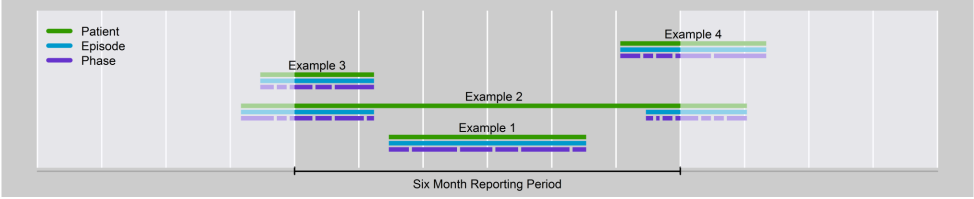




Appendix C Data scoping method

The method used to determine which data is included in a PCOC report looks at the phase level records first. All phase records that <u>end</u> within the 6 month reporting period are deemed to be "in scope" and would be included in the report. The episode and patient records associated with these phases are also deemed to be "in scope" and hence would also be included in the report. Figure 23 below displays four examples to help visualize this process.





In <u>Example 1</u>, the patient (represented by the green line) has one episode (represented by the blue line). This episode has six phases (represented by the purple line segments). All six phases would be included in the report as they all end within the reporting period. Hence, the episode and patient would also be in the report.

In <u>Example 2</u>, the patient has two episodes - the first having six phases and the second having seven phases. Looking at the phases associated with the first episode, the last four will be included in the report (as they end within the reporting period). The first two phases would have been included in the previous report. For the phases relating to the second episode, only the first three end within the reporting period, so only these would be included in the report. The following four phases would be included in the next report. Both of the episode records and the patient record would also be included in the report.

In <u>Example 3</u>, the patient has one episode and five phases. Only the last three phases will be included in the report as they are the only ones ending within the reporting period (the first two phases would have been included in the previous report). The episode and patient records would be included in the report.

In <u>Example 4</u>, the patient again has one episode and five phases. This time, only the first three phases will be included in the report (the last two phases will be included in the next report). Again, the episode and patient records would be included in the report.



Appendix D Palliative Care Phase definitions

START	END
1. Stable	
Patient problems and symptoms are adequately controlled by established plan of care and Further interventions to maintain symptom control and quality of life have been planned and Family/carer situation is relatively stable and no new issues are apparent.	The needs of the patient and / or family/carer increase, requiring changes to the existing plan of care.
2. Unstable	
 An urgent change in the plan of care or emergency treatment is required because Patient experiences a new problem that was not anticipated in the existing plan of care, and/or Patient experiences a rapid increase in the severity of a current problem; and/or Family/ carers circumstances change suddenly impacting on patient care. 	 The new plan of care is in place, it has been reviewed and no further changes to the care plan are required. This does not necessarily mean that the symptom/crisis has fully resolved but there is a clear diagnosis and plan of care (i.e. patient is stable or deteriorating) and/or Death is likely within days (i.e. patient is now terminal).
3. Deteriorating	
 The care plan is addressing anticipated needs but requires periodic review because Patients overall functional status is declining and Patient experiences a gradual worsening of existing problem and/or Patient experiences a new but anticipated problem and/or Family/carers experience gradual worsening distress that impacts on the patient care. 	 Patient condition plateaus (i.e. patient is now stable) or An urgent change in the care plan or emergency treatment and/or Family/ carers experience a sudden change in their situation that impacts on patient care, and urgent intervention is required (i.e. patient is now unstable) or Death is likely within days (i.e. patient is now terminal).
4. Terminal	
Death is likely within days.	 Patient dies or Patient condition changes and death is no longer likely within days (i.e. patient is now stable or deteriorating).
5. Bereavement – post death support	
 The patient has died Bereavement support provided to family/carers is documented in the deceased patient's clinical record. 	 Case closure Note: If counselling is provided to a family member or carer, they become a client in their own right.



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PCOC has made every effort to ensure that the data used in this report are accurate. Data submitted to PCOC are checked for anomalies and services are asked to re-submit data prior to the production of the PCOC report. We would advise readers to use their professional judgement in considering all information contained in this report.

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