



CANCER SERIES Number 52

Ovarian cancer in Australia

An overview, 2010

Australian Institute of Health and Welfare and National Breast and Ovarian Cancer Centre

February 2010

Australian Institute of Health and Welfare Canberra Cat. no. CAN 48 The Australian Institute of Health and Welfare is Australia's national health and welfare statistics and information agency. The Institute's mission is better information and statistics for better health and wellbeing.

National Breast and Ovarian Cancer Centre (NBOCC) is Australia's authority and source of evidencebased information on breast and ovarian cancer. Funded by the Australian Government, NBOCC works in partnership with health professionals, cancer organisations, researchers, governments and those diagnosed to improve outcomes in breast and ovarian cancer. NBOCC plays a vital role in the translation of worldwide cancer research into meaningful and evidence-based information to guide the work of Australian health professionals, improve health service delivery, inform people with breast or ovarian cancer about all aspects of their diagnosis and treatment, inform policy and raise community awareness about these diseases. For more information, visit <www.nbocc.org.au>.

© Australian Institute of Health and Welfare 2010

This work is copyright. Apart from any use as permitted under the *Copyright Act 1968*, no part may be reproduced without prior written permission from the Australian Institute of Health and Welfare. Requests and enquiries concerning reproduction and rights should be directed to the Head, Media and Communications Unit, Australian Institute of Health and Welfare, GPO Box 570, Canberra ACT 2601.

This publication is part of the Australian Institute of Health and Welfare's Cancer series. A complete list of the Institute's publications is available from the Institute's website <www.aihw.gov.au>.

ISSN 1039-3307 ISBN 978 1 74024 996 6

Suggested citation

Australian Institute of Health and Welfare & National Breast and Ovarian Cancer Centre 2010. Ovarian cancer in Australia: an overview, 2010. Cancer series no. 52. Cat. no. CAN 48. Canberra: AIHW.

Australian Institute of Health and Welfare	National Breast and Ovarian Cancer Centre
Board Chair	Board Chair
Hon. Peter Collins, AM, QC	Dr Megan Keaney
Director	Chief Executive Officer
Dr Penny Allbon	Dr Helen Zorbas

Any enquiries about or comments on this publication should be directed to: Dr Adriana Vanden Heuvel Australian Institute of Health and Welfare GPO Box 570 Canberra ACT 2601 Phone: (02) 6244 1184 Email: a.vandenheuvel@aihw.gov.au

Published by the Australian Institute of Health and Welfare Printed by Blue Star Print Group

Foreword

Ovarian cancer in Australia: an overview, 2010 brings together in one volume the most up-todate statistical information available on the epidemiology, public health and health services impact of ovarian cancer in Australia. These data, collected through population-based cancer registries and other sources, are central to advancing our efforts to understand and ultimately control this disease. This report not only builds on previous monitoring reports but additionally provides data about the burden of disease due to ovarian cancer, as well as survival from ovarian cancer by Indigenous status and by histology types.

Ovarian cancer in Australia: an overview, 2010 also represents the significant contributions and the continuing partnership of National Breast and Ovarian Cancer Centre (NBOCC), the Australian Institute of Health and Welfare (AIHW) and the Australasian Association of Cancer Registries (AACR) and it highlights the importance of registries as a national resource. The report provides a nationwide snapshot of a major condition affecting a substantial number of Australian women.

The value of data and monitoring is its relevance to outcomes and its capacity to impact on change. This report identifies areas of significant gain over time and provides some predictions for the future. Our ability to plan for services and patient needs are predicated on this understanding of the impact of the disease as it affects our population.

We would like to thank the staff of the various cancer registries and data repositories. It is through their effort and diligence that these data are available to the Australian public. We anticipate that the information contained in *Ovarian cancer in Australia: an overview, 2010* will be used extensively to further reduce mortality from ovarian cancer and improve the wellbeing of women with the disease.

Dr Helen Zorbas Chief Executive Officer National Breast and Ovarian Cancer Centre Dr Penny Allbon Director Australian Institute of Health and Welfare

Contents

Ac	knowledgments	viii
Ab	breviations	ix
	Symbols	x
Exe	ecutive summary	xi
1	Introduction	1
	What is ovarian cancer?	1
	Purpose and structure of this report	2
	Data interpretation	2
	Data sources	4
2	Incidence of ovarian cancer	5
	Incidence in 2006	5
	Differences by age at diagnosis	6
	Trends	6
	Trends by age at diagnosis	8
	Risk of ovarian cancer and average age at diagnosis	9
	Projections	10
	Types of ovarian cancer	11
	Trends in types of ovarian cancer	12
	Incidence by stage at diagnosis	15
	Differences across groups	17
	Differences by geographical area	
	Differences by socioeconomic status	19
	Differences by Aboriginal and Torres Strait Islander status	20
	Differences by country of birth	21
	International comparisons	22
3	Mortality from ovarian cancer	25
	Mortality in 2006	25
	Differences by age at death	26
	Trends	27
	Trends by age at death	
	Risk of death from ovarian cancer and average age at death	
	Differences across groups	
	Differences by geographical area	
	Differences by socioeconomic status	31
	Differences by Aboriginal and Torres Strait Islander status	31
	Differences by country of birth	
	International comparisons	
	Ovarian cancer as an associated cause of death	34
4	Survival after a diagnosis of ovarian cancer	36
	Survival of those diagnosed between 2000 and 2006	

	Survival from ovarian cancer compared with other cancers	
	Differences by age at diagnosis	
	Trends	
	Trends by age at diagnosis	40
	Survival by type of ovarian cancer	41
	Survival by stage at diagnosis	44
	Differences by Aboriginal and Torres Strait Islander status	45
	International comparisons	46
5	Prevalence of ovarian cancer	48
	Prevalence in 2006	49
	Differences by age	50
	Differences across groups	50
6	Burden of disease due to ovarian cancer	53
	Burden of disease in 2003	53
	Differences by age	56
	Trends and projections	57
7	Hospitalisations for ovarian cancer	58
	Hospitalisations in 2007–08	58
	Differences by age	59
	Average length of stay	59
	Trends	60
	Trends in average length of stay	61
	Procedures undertaken during hospitalisations	63
8	Expenditure on ovarian cancer	64
	Expenditure in 2004–05	64
	Differences by age	65
	Trends	66
Ар	pendix A: Classifications	67
-	International Statistical Classification of Diseases and Related Health Problems	67
	International Statistical Classification of Diseases and Related Health Problems, Australian modification	67
	Australian Classification of Health Interventions	67
	International Classification of Diseases for Oncology	68
	Australian Standard Geographical Classification Remoteness Areas	68
	Index of Relative Socio-economic Disadvantage	68
	Standard Australian Classification of Countries	68
Ар	pendix B: Statistical methods and technical notes	69
-	- Age-specific rates	69
	Age-standardised rates	69
	Confidence intervals	70
	Definition of ovarian cancer	71
	What ICD codes are included in the definition	71
	Whether or not borderline tumours are included	72

Incidence projections	72
Mortality data differences	73
Mortality-to-incidence ratio	73
Relative survival analysis	74
Risk to age 75 and 85 years	75
Appendix C: Data sources	76
Australian Cancer Database	76
Non-melanoma skin cancers	77
Burden of disease data	78
Disease Expenditure Database	78
GLOBOCAN	79
National Death Index	79
National Hospital Morbidity Database	80
National Mortality Database	80
Population data	81
Appendix D: Additional tables	82
Additional tables for Chapter 2: Incidence of ovarian cancer	82
Additional tables for Chapter 3: Mortality from ovarian cancer	92
Additional tables for Chapter 4: Survival after a diagnosis of ovarian cancer	98
Additional table for Chapter 6: Burden of disease due to ovarian cancer	103
Additional tables for Chapter 7: Hospitalisations for ovarian cancer	104
Additional table for Chapter 8: Expenditure on ovarian cancer	106
Appendix E: Stage at diagnosis	107
FIGO staging system	107
TNM staging system	108
Summary staging system	108
Appendix F: Definition of ovarian cancer-related hospitalisations	109
Glossary	112
References	115
List of tables	121
List of figures	124

Acknowledgments

This report was commissioned by National Breast and Ovarian Cancer Centre (NBOCC). Advice on content was provided by Professor Tom Dodd and Professor Peter Russell, with additional input from Ms Margot Osinski. Special thanks are due to Dr Penny Webb and Clinical Associate Professor Peter Grant, who reviewed the report. The following NBOCC staff are acknowledged for their contribution to the development of this report: Professor David Roder; Dr Helen Zorbas; Ms Rosemary Wade; Ms Jane Francis; and Associate Professor Christine Giles. The work of NBOCC is funded by the Australian Government Department of Health and Ageing.

This report was prepared by staff in the Cancer and Screening Unit of the Australian Institute of Health and Welfare (AIHW). The main authors were Dr Adriana Vanden Heuvel and Ms Anne Ganner Bech. The main data analyst was Ms Candice Rabusa. Other AIHW staff who made a substantial contribution are, in alphabetical order, Ms Chun Chen, Dr Brett Davis, Ms Melissa Goodwin, Mr David Meere, Ms Galina Prosselkova, Dr Mark Short, Ms Christine Sturrock and Ms Kun Zhao. The authors would like to thank those AIHW staff who commented on earlier drafts of this report.

The support of the Australasian Association of Cancer Registries in both providing data and reviewing the draft report is gratefully acknowledged.

Abbreviations

AACR	Australasian Association of Cancer Registries
ABS	Australian Bureau of Statistics
ACD	Australian Cancer Database
ACHI	Australian Classification of Health Interventions
ACT	Australian Capital Territory
ASGC	Australian Standard Geographical Classification
AIHW	Australian Institute of Health and Welfare
AS	age-standardised
ASR	age-standardised rate
BCC	basal cell carcinoma
CI	confidence interval
CS	crude survival
DALY	disability-adjusted life year
DoHA	Australian Government Department of Health and Ageing
FIGO	International Federation of Gynecology and Obstetrics
IARC	International Agency for Research on Cancer
ICD-7	International Statistical Classification of Diseases and Related Health Problems, seventh revision
ICD-10	International Statistical Classification of Diseases and Related Health Problems, tenth revision
ICD-10-AM	International Statistical Classification of Diseases and Related Health Problems, tenth revision, Australian modification
ICD-O-2	International Classification of Diseases for Oncology, second edition
ICD-O-3	International Classification of Diseases for Oncology, third edition
IRSD	Index of Relative Socio-economic Disadvantage
MIR	mortality-to-incidence ratio
NBOCC	National Breast and Ovarian Cancer Centre
NHMD	National Hospital Morbidity Database
NMD	National Mortality Database
NMSC	non-melanoma skin cancer
No.	number
NOS	not otherwise specified
NSW	New South Wales
NT	Northern Territory
NZ	New Zealand
Qld	Queensland
RS	relative survival
SA	South Australia

SACC	Standard Australian Classification of Countries
SCC	squamous cell carcinoma
SEER	Surveillance Epidemiology End Results
Tas	Tasmania
TNM	a staging system based on size/extent of the tumour (T), lymph node involvement (N) and presence of distant metastases (M)
UICC	International Union Against Cancer
UK	United Kingdom
USA	United States of America
Vic	Victoria
WA	Western Australia
WHO	World Health Organization
YLL	years of life lost
YLD	years lost due to disability

Symbols

- .. not applicable
- % per cent
- < less than
- > greater than
- + and over
- n.a. not available
- n.p. not published (data cannot be released due to quality issues)

Executive summary

Although a relatively uncommon cancer, ovarian cancer is often diagnosed at a stage where the cancer has spread beyond the ovary. Such cases often have a poor prognosis.

Ovarian cancer in Australia: an overview, 2010 provides a comprehensive picture of national statistics on ovarian cancer using a range of data sources, with the latest available data and trends over time presented. Throughout this report, the term 'ovarian cancer' refers to invasive ovarian cancers; borderline tumours are not included.

The number of ovarian cancer cases is increasing

In 2006, ovarian cancer was the ninth most commonly diagnosed cancer among Australian women (excluding non-reportable skin cancers) and the second most commonly diagnosed gynaecological cancer, with a total of 1,226 ovarian cancer cases diagnosed. Ovarian cancer is mainly a disease of postmenopausal women, with six in ten (60%) cases diagnosed in women aged 60 years and over.

The number of ovarian cancer cases increased by 47% between 1982 and 2006 (from 833 cases to 1,226 cases) due to an ageing and growing population. It is anticipated that the number of new cases will continue to increase, with an estimated 1,434 women expected to be diagnosed with ovarian cancer in 2015.

Nonetheless, the age-standardised incidence rate of ovarian cancer decreased significantly by 14% between 1982 and 2006 (from 12.4 to 10.7 new cases per 100,000 females).

The rate of death from ovarian cancer has fallen

A total of 795 women died from ovarian cancer in 2006, making it the sixth most common cause of cancer-related death for Australian women, and the most common cause of gynaecological cancer death, representing over half (55%) of such deaths.

The age-standardised mortality rates for ovarian cancer decreased significantly by 26% between 1968 and 2006 (from 9.1 to 6.7 deaths per 100,000 females). In addition, the 2006 mortality rate was the lowest rate observed for any year to date. Possible reasons for the decrease in the mortality rate over time include the observed decline in the incidence rate, improvements in access to and quality of treatments, and change over time in the types of ovarian cancers occurring among women. However, the data also indicate that the decline in the mortality rate was not observed for all age groups, with the ovarian cancer mortality rate for older women (those aged 70 years and over at death) increasing rather than decreasing over the period considered.

The prognosis of women with ovarian cancer has improved

The prognosis for women with ovarian cancer is relatively poor. Women who were diagnosed with ovarian cancer between 2000 and 2006 were 40% as likely to live five years after diagnosis as their counterparts in the general population. Significantly poorer survival was seen for older women, with 5-year relative survival estimates ranging from a high of 86% for those aged less than 30 years when diagnosed with ovarian cancer to a low of 15% for those aged 80 years or older at diagnosis. Possible reasons for poorer survival of older women include a greater likelihood that these women were diagnosed with advanced stage cancer and/or with more-aggressive types of cancers, as well as a greater likelihood of

co-morbidities. Differences by age in the treatment provided to those with ovarian cancer are also believed to be a factor.

Improvement in the prognosis of those diagnosed with ovarian cancer has occurred over time, with the 5-year relative survival rate increasing significantly from 33% in 1982–1987 to 40% in 2000–2006. Nonetheless, the improvements in survival were focused on women in the middle age groups, with no significant change in the survival estimates over time for those aged less than 40 years and those aged 80 years and over.

1 Introduction

Ovarian cancer is often referred to as a 'challenging cancer'. Unlike breast and cervical cancer, no effective tests are currently available for population-based screening for ovarian cancer (NBOCC 2009). Further, the symptoms of ovarian cancer (such as abdominal swelling, abdominal or back pain, and intestinal and urinary symptoms) tend to be similar to the symptoms of many other common conditions. Thus, ovarian cancer is often diagnosed at an advanced stage.

What is ovarian cancer?

The ovaries are a pair of solid, oval-shaped organs that are part of the female reproductive system, with one ovary on each side of the uterus (Figure 1.1). Each ovary is around 3 centimetres long and 1 centimetre thick.

Ovarian cancer is a disease in which abnormal cells in the ovaries multiply and form an invasive (i.e. malignant) tumour. Such tumours can spread to other parts of the body and, if the spread is not controlled, can result in death. Benign tumours can also form in the ovaries; such tumours do not spread and, with very rare exceptions, are not life-threatening.



There are three main types of ovarian cancers, each of which begin in a different type of cell in the ovary. Epithelial cells form an outer covering over the ovary. Most ovarian cancers begin in this layer of cells and are called epithelial ovarian cancers. Germ cells are found inside the ovary and these cells eventually mature into the eggs (ova) that are released into the fallopian tubes. Cancers which develop in these cells are called germ cell ovarian cancers. Stromal cells release the hormones oestrogen and progesterone. Ovarian cancers which begin in these cells are called sex cord-stromal ovarian cancers. Similar to invasive epithelial ovarian cancers, borderline tumours (also known as tumours of low malignant potential) develop in the epithelium of the ovaries but these tumours are not as aggressive as other epithelial

ovarian tumours and there is less risk that they will spread or recur. As discussed in more detail later in this chapter, borderline tumours are not regarded as malignant tumours in the current version of the international coding standards for cancers and thus these tumours are not considered in this report.

While the cause of ovarian cancer is unknown, two of the main risk factors are advancing age and a family history of ovarian cancer, while important protective factors are increased parity and use of the contraceptive pill (ACN & NBCC 2004).

Purpose and structure of this report

The purpose of this report is to provide a comprehensive snapshot of national statistics on ovarian cancer in Australia. The aim is to increase the level of understanding about this disease and to inform decision-making, resource allocation and the evaluation of programs and policies. The report is aimed at a wide audience — including health professionals, policy makers, health planners, educators, researchers, consumers and the general public.

This report brings together the latest available statistics and trend data on the following topics:

- the number of cases of ovarian cancer diagnosed each year (Chapter 2)
- the number of women who die from this disease each year (Chapter 3)
- survival prospects for those diagnosed with ovarian cancer (Chapter 4)
- the number of women alive who have been diagnosed with ovarian cancer (Chapter 5)
- the burden of disease due to ovarian cancer (Chapter 6)
- the number of hospitalisations for ovarian cancer (Chapter 7)
- the extent of health care spending on ovarian cancer (Chapter 8).

Compared with the previous edition of this report (AIHW & NBCC 2006), this edition includes, for the first time, information on how incidence differs within Australia according to country of birth. It also provides additional information on how Australian ovarian cancer data compare globally and by Aboriginal and Torres Strait Islander status.

Data interpretation

In this report, the term 'ovarian cancer' is used to refer to primary ovarian tumours which are invasive (i.e. malignant). It does not encompass secondary ovarian cancers, nor does it include benign or non-invasive ovarian tumours. Furthermore, borderline tumours are not included. In the second edition of the International Classification of Diseases for Oncology (ICD-O-2), ovarian tumours of borderline malignancy were considered malignant. However, in the third edition of ICD-O (i.e. ICD-O-3), borderline tumours are considered to be of uncertain behaviour and are no longer considered malignant. For the 2006 version of this report (AIHW & NBCC 2006), tumours were classified according to ICD-O-2 and thus the results presented in that report are not strictly comparable to the results shown in this report, which are based on ICD-O-3. As shown in the earlier report, 6% of ovarian cancer cases diagnosed in 2002 were borderline tumours (AIHW & NBCC 2006).

In this report, ovarian cancer is defined, when possible, as those cancers classified as 'C56' in the tenth revision of the International Statistical Classification of Diseases and Related Health Problems (i.e. ICD-10). However, in some data sets used in this report (namely the burden of disease data, the expenditure data and the international GLOBOCAN data), data were not available for the ICD-10 code of 'C56' but instead were available for the ICD-10 codes of 'C56 and C57.0–C57.4' grouped together (see Appendix Table B.2). This grouping of cancers is referred to as 'ovarian and related cancers' in this report.

Information about the classifications referred to in this report, including ICD and ICD-O, is provided in Appendix A.

Information on tumour stage (i.e. extent of spread) at time of diagnosis is important in relation to both prognosis and determining the most appropriate type of treatment. Information on change over time in stage at diagnosis also assists in the monitoring of ovarian cancer control policies and programs. While some of the Australian states and

territories collect information on stage at diagnosis for ovarian cancer, not all do so and there are no nationally agreed standards for the collection of these data. While national data on these items are not available, some state-level and overseas data on incidence and survival are presented by stage at diagnosis in this report.

Information on the actual number of ovarian cancer cases and deaths is presented in this report, together with age-standardised rates. The use of age-standardised rates is important when making comparisons between groups and within groups over time in order to take into account differences in the age structure and size of the population. This is especially important in regard to ovarian cancer since the risk of this disease increases with age. Rates have been standardised to the Australian population at 30 June 2001 and are generally expressed per 100,000 females. In addition, for some of the key statistics and the international comparisons, age-standardised rates based on a World Standard Population are shown. The use of a world standard allows for the comparison of Australian data with those of other countries. Further information on age standardisation and other technical matters can be found in Appendix B. Note that all discussion of age-standardised rates in the text of this report pertains to rates that were standardised to the Australian population, with the exception of the discussion on international comparisons, which pertains to rates standard.

Confidence intervals (at the 95% level) are shown in graphs (as error bars) and tables in this report. As explained more fully in Appendix B, confidence intervals can be used as a guide when considering whether differences in rates may be a result of chance variation. Where confidence intervals do not overlap, the difference between rates may be greater than would readily be attributable to chance. While such differences may be regarded as 'significant' in statistical terms, they may or may not be 'significant' from a practical or clinical perspective.

In this report, comparisons are often made with international and state/territory-based data in order to help put the Australian situation into a broader context, and to give some indication as to whether the observed findings have been found in other studies as well. However, caution must be taken when making such comparisons since observed differences may be influenced not only by the underlying number of ovarian cancer cases (or the number of deaths when considering mortality data), but also by differences between Australia and individual jurisdictions or other countries in the following:

- cancer detection
- types of treatment provided and access to treatment services
- characteristics of the cancer such as stage at diagnosis and histological type
- coding practices and cancer registration methods, as well as the accuracy and level of cancer coverage of the data.

The last point is of particular relevance in regard to ovarian cancer since different jurisdictions and countries include different types of cancers within the definition of 'ovarian cancer'. In addition, ovarian borderline tumours are included in some data but excluded from others (see Appendix B for further details). Finally, difficulties in distinguishing between ovarian cancers and borderline tumours (Kricker 2002), as well as between ovarian cancers of cancers (such as peritoneum cancers or cancers of an unknown primary site) also contribute to observed variations in ovarian cancer rates between jurisdictions and countries, as well as over time.

Data sources

A key data source for this report was the Australian Cancer Database (ACD), which was previously known as the National Cancer Statistics Clearing House. The ACD is a database that holds information on 1.8 million Australian cancer cases diagnosed between 1982 and 2006. The AIHW compiles and maintains the ACD, in partnership with the Australasian Association of Cancer Registries (AACR), whose member registries provide data to the AIHW on an annual basis. Each Australian state and territory has legislation that makes the reporting of all cancers (other than two types of non-melanoma skin cancers (NMSC)) mandatory. Note that compared with past reports prepared by the AIHW, a different approach to the exclusion of non-melanoma skin cancers from the data was used when preparing this report. Additional information about this change, as well as about the ACD itself, can be found in Appendix C.

Another key data source was the National Mortality Database (NMD). This database contains information on the date and cause of death for all registered deaths in Australia from 1964 onwards. Depending on the coding version used (ICD-7 in 1964 through to ICD-10 at present), some diseases may not have had a unique code that identifies the disease separately from other closely-related diseases for some of the years. Ovarian cancer is an example of this, with data on mortality due specifically to ovarian cancer (rather than from ovarian and related cancers) available from 1968 onwards. Additional information about the NMD is provided in Appendix C.

In addition, several other data sources — including the National Death Index, the National Hospital Morbidity Database, the Disease Expenditure Database and the 2002 GLOBOCAN database — have been used to present a broad picture of ovarian cancer in Australia in this report. Information about each of these data sources can also be found in Appendix C.

Throughout this report:

- The term 'ovarian cancer' refers to primary, invasive ovarian cancers, with cancers classified as 'C56' in ICD-10 included (unless otherwise indicated). Borderline tumours are excluded.
- Differences that are described as 'significant' refer to a statistically significant difference. Such differences may or may not be significant from a practical or clinical perspective.

2 Incidence of ovarian cancer

Incidence data indicate the number of new cases of ovarian cancer diagnosed during a specified time period, usually one year. While these data refer to the number of *cases* diagnosed and not the number of *women* diagnosed with ovarian cancer, it is rare (although possible) that any one woman would be diagnosed with two or more primary ovarian cancers during a 1-year period. Thus, the annual number of new ovarian cancer cases is practically the same as the annual number of women newly diagnosed with ovarian cancer.

Details on the incidence of ovarian cancer over time are provided in this chapter, as is information on the projected number of new cases to 2015, the risk of a woman being diagnosed with ovarian cancer by the age of 75 and 85 years, and disparities in incidence among women according to age, geographical area, socioeconomic status, Aboriginal and Torres Strait Islander status and country of birth. Information on how Australia's ovarian cancer rates compare internationally is also shown.

As mentioned in Chapter 1, only those cases in which ovarian cancer was a primary, invasive cancer are considered. Additionally, to be counted, the case must be a 'new' primary cancer and not a reoccurrence of a previous primary cancer (IARC 2004).

The main data source for this chapter was the Australian Cancer Database.

Incidence in 2006

The ten most commonly diagnosed cancers among females in 2006 are shown in Table 2.1. Since two types of skin cancer – namely, basal cell carcinoma and squamous cell carcinoma – are not reported to cancer registries, data on these two types of cancer are not included in either the ACD or Table 2.1. Past research shows that these skin cancers are by far the most frequently diagnosed cancers in Australia in both females and males (AIHW & CA 2008).

In 2006, a total of 1,226 cases of ovarian cancer were diagnosed in Australia; this equates to an average of 3 women being diagnosed with this disease every day. Ovarian cancer was the ninth most commonly diagnosed cancer among females (excluding basal and squamous cell carcinomas of the skin) and the second most commonly diagnosed gynaecological cancer, after cancer of the uterus (1,860 cases). Ovarian cancer accounted for 3% of all reported cancer cases in women in 2006 and 29% of all gynaecological cancers.

The age-standardised rate of ovarian cancer incidence stood at 10.7 cases per 100,000 females in 2006. This compares with a rate of 112.4 cases per 100,000 females for breast cancer and 16.3 cases per 100,000 females for cancer of the uterus.

Cancer type (ICD-10 codes)	Number of cases	Per cent of all gynaecological cancer cases	Per cent of all cancer cases ^(a)	Age- standardised rate ^(b)	95% confidence interval
Breast (C50)	12,614		27.7	112.4	110.4–114.4
Bowel (C18–C20)	6,159		13.5	52.1	50.8–53.4
Melanoma of skin (C43)	4,275		9.4	38.2	37.1–39.4
Lung (C33–C34)	3,533		7.8	30.3	29.3–31.3
Lymphoma (C81–C85, C96)	1,961		4.3	17.2	16.4–18.0
Uterus (C54, C55)	1,860	43.8	4.1	16.3	15.6–17.1
Unknown primary site (C26, C39, C76, C80)	1,592		3.5	12.6	12.0–13.3
Thyroid (C73)	1,270		2.8	11.8	11.2–12.5
Ovary (C56)	1,226	28.9	2.7	10.7	10.1–11.4
All leukaemias (C91–C95)	1,111		2.4	9.7	9.1–10.3
All cancers ^(c)	45,534		100.0	396.3	392.6-400.0

Table 2.1: Incidence of the 10 most commonly diagnosed cancers^(a), females, 2006

(a) Excluding basal and squamous cell carcinomas of the skin.

(b) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females.

(c) Includes cancers coded in ICD-10 as C00–C97, D45, D46, D47.1 and D47.3 with the exception of those C44 codes which indicate a basal or squamous cell carcinoma.

Source: Australian Cancer Database, AIHW.

Differences by age at diagnosis

In 2006, the majority of cases of ovarian cancer were diagnosed among women aged 60 years and over (60%). In addition, one in fourteen cases (7%) were diagnosed in women under the age of 40 years, and one-third (33%) were diagnosed in those aged 40 to 59 years (Appendix Table D2.1). The mean age at first diagnosis was 63 years.

Figure 2.1 shows differences in the incidence rate of ovarian cancer by age group in 2006. The likelihood of being diagnosed with ovarian cancer increased with age. The rates were relatively low for those aged less than 40 years (i.e. less than 5 cases per 100,000 females for each of those age groups). In contrast, there was an observable increase between most age groups from the age of 40 to 44 years onwards, but these increases were not always statistically significant. The highest rate was observed for women aged 80 to 84 years (48.0 per 100,000), followed by those aged 85 years and over (44.5 per 100,000).

Trends

The number of new ovarian cancer cases diagnosed each year increased over the 25-year period from 1982 (the year in which national cancer incidence data were first available) to 2006 (Figure 2.2). In 1982, 833 new cases of ovarian cancer were diagnosed among Australian women compared with 1,226 in 2006, indicating an overall increase of 47% cases over this period. Furthermore, the number of ovarian cancer cases diagnosed in 2006 was slightly higher than the number diagnosed in the previous year (1,219 cases) but lower than the number diagnosed in 2004 (1,267 cases) when the largest number of ovarian cancer cases diagnosed to date was reported.





In contrast, there was a statistically significant decrease in the age-standardised incidence rate of ovarian cancer. In 1982, the incidence rate stood at 12.4 (per 100,000 females), while it was 10.7 (per 100,000 females) in 2006, indicating an overall decrease of 14%. This indicates that the increase in the absolute number of ovarian cancer cases over the years can be explained by the ageing and increasing size of the population.

The reason for the decline in the age-standardised rate is unclear, with possible reasons being change over time in the proportion of women with various ovarian cancer risk and protective factors (e.g. use of the contraceptive pill), changes in the classification systems and approaches used to diagnose borderline versus invasive tumours and, in some cases, the removal of precancerous lesions which prevented ovarian cancer from developing (Skirnisdottir et al. 2008).

The share of all reportable cancers diagnosed in females that were ovarian cancer decreased between 1982 and 2006, from 3.8% to 2.7% respectively (Appendix Table D2.2). Meanwhile, the proportion of all gynaecological cancers diagnosed in females that were ovarian cancer ranged over the years from 27.2% (1986) to 31.2% (2002).

Trends by age at diagnosis

While overall there was a slight decrease in the rate of ovarian cancer over the years considered, this did not apply equally to all of the age groups (Figure 2.3). Instead, the observed decrease in incidence rates over the years was centred on those aged 50 to 69 years, with rates decreasing significantly from 32.7 cases per 100,000 females in 1982 to 24.6 cases per 100,000 females in 2006; this equates to a decrease of 25%. The incidence rate of ovarian cancer for women aged less than 50 years also decreased somewhat from 1982 (3.9 cases per 100,000 females) to 2006 (3.1 per 100,000), but this decrease was not statistically significant. For women aged 70 years and over, incidence rates fluctuated considerably over the period, with no statistically significant difference found between the 1982 rate (35.8 per 100,000) and the 2006 rate (40.9 per 100,000).



Risk of ovarian cancer and average age at diagnosis

Table 2.2 shows the risk of a woman being diagnosed with ovarian cancer by the age of 75 years and 85 years (see Appendix B for an explanation of how these risks were calculated). While the risk of being diagnosed with ovarian cancer fluctuated somewhat between 1982 and 2006, an overall decline in the risk is evident. The risk of being diagnosed with ovarian cancer by the age of 75 years was 1 in 94 in 1982, compared with 1 in 116 in 2006. The corresponding values in the risk of being diagnosed by the age of 85 years was 1 in 71 (in 1982) to 1 in 77 (in 2006).

Table 2.2 also indicates change over time in the mean and median age at first diagnosis of ovarian cancer. Both the mean and median ages at diagnosis have been tending upwards over time, with the mean age increasing from 60 years in 1982 to 63 years in 2006.

Year	Risk to age 75 years	Risk to age 85 years	Mean age at first diagnosis	Median age at first diagnosis
1982	1 in 94	1 in 71	59.6	60.0
1983	1 in 98	1 in 70	59.8	61.0
1984	1 in 95	1 in 69	60.5	62.0
1985	1 in 101	1 in 67	60.7	62.0
1986	1 in 104	1 in 74	60.5	62.0
1987	1 in 98	1 in 72	60.6	62.0
1988	1 in 104	1 in 74	60.4	62.0
1989	1 in 93	1 in 66	61.2	63.0
1990	1 in 96	1 in 67	61.7	64.0
1991	1 in 100	1 in 68	62.1	64.0
1992	1 in 97	1 in 69	60.7	62.0
1993	1 in 97	1 in 66	61.3	64.0
1994	1 in 99	1 in 69	61.5	63.0
1995	1 in 105	1 in 67	62.1	64.0
1996	1 in 108	1 in 70	63.2	65.0
1997	1 in 113	1 in 74	62.9	64.0
1998	1 in 106	1 in 69	62.9	65.0
1999	1 in 109	1 in 74	62.8	64.0
2000	1 in 114	1 in 74	63.7	65.0
2001	1 in 120	1 in 76	64.3	65.0
2002	1 in 112	1 in 69	63.9	65.0
2003	1 in 118	1 in 77	63.3	64.0
2004	1 in 109	1 in 72	63.9	65.0
2005	1 in 120	1 in 75	64.1	65.0
2006	1 in 116	1 in 77	63.3	64.5

	Table 2.2: Risk a	ind average age af	diagnosis of	ovarian cancer,	1982 to 2006
--	-------------------	--------------------	--------------	-----------------	--------------

Source: Australian Cancer Database, AIHW.

Projections

To estimate the incidence of ovarian cancer from 2007 to 2015, data on the number of new cases of ovarian cancer diagnosed over the 10-year period from 1997 to 2006 were extrapolated (see Appendix B for further details on the methodology used). This estimation approach assumes that the trends in the incidence of ovarian cancer during that 10-year period will continue to 2015. Since it is impossible to anticipate and quantify future developments that might cause a change in the number of women diagnosed with ovarian cancer, these projections should be interpreted as only indicative of future trends. Note also that there is greater margin of error surrounding the projections for the later years than the earlier years.

The number of women diagnosed with ovarian cancer is expected to continue to increase in the future due to continued ageing and growth of the population (Figure 2.4). The projections suggest that 1,324 new cases of ovarian cancer will be diagnosed in 2010. By 2015, the number of new ovarian cancer cases diagnosed is estimated to be 1,434, which would be 17% higher than in 2006 (1,226 cases). If these projections are accurate, an average of 4 women would be diagnosed with ovarian cancer each day in 2015.

Figure 2.4 also indicates the projected age-standardised incidence rate for ovarian cancer from 2007 to 2015. When expected changes in the age structure and size of the population are taken into account, the results suggest that the rate at which new ovarian cancer cases are diagnosed will continue to decrease slightly through to 2015, reaching 10.1 new cases per 100,000 females in that year. However, due to the wide confidence interval around this estimate, the projected incidence rate for 2015 is not statistically different from the 2006 rate of 10.7 cases per 100,000 females.



3. For the years 2007 to 2015, grey lines around the age-standardised rates indicate the 95% prediction intervals.

4. The data for this figure are shown in Appendix Tables D2.2 and D2.4.

Source: Australian Cancer Database, AIHW.

Figure 2.4: Incidence of ovarian cancer, observed for 1997 to 2006 and projected for 2007 to 2015

Types of ovarian cancer

Ovarian cancer consists of a heterogeneous set of invasive tumours that arise from the different cell types in the ovary (as discussed in Chapter 1). Pathologists classify ovarian cancers into histological types, with each of the types associated with different genetic risk factors, patterns of transformation and responses to chemotherapy (Kobel et al. 2008). Due to the large number of histological types of ovarian cancer and the relative infrequency of many of these types in the Australian Cancer Database, similar types of tumours have been grouped into broader categories for the purposes of this report. As shown in Table 2.3, the three main categories are: 'group 1: carcinoma' (which is also referred to as epithelial tumours)'; 'group 2: sex cord-stromal tumours'; and 'group 3: germ cell tumours'. Two additional categories capture other specified types of ovarian cancers (group '4') and ovarian cancers that were unspecified (group '5'). One of the groups – namely 'group 1: carcinoma' – is further divided into seven subgroups. The histology types included in each group and sub-group are listed in Appendix Table D2.5. This system of grouping ovarian cancers was based primarily on documentation from the International Agency for Research on Cancer (Curado et al. 2007), with additional input from National Breast and Ovarian Cancer Centre. In 2006, 84% of the ovarian cancers were classified as carcinoma (group 1) with the most common type within this group being serous carcinoma (group 1.1) (52% of carcinomas), followed by adenocarcinoma not otherwise specified (group 1.5) (15% of carcinomas). Meanwhile, 4% of ovarian cancers were classified as germ cell tumours (group 3), 3% were coded as other specified malignant neoplasm (group 4) and 1% were coded as sex cord-stromal tumours (group 2). The histological type was *unspecified* (group 5) in 1 in 13 (8%) ovarian cancer cases.

Type of ovarian cancer ^(a)	Number of cases	Per cent of all ovarian cancers	Per cent of carcinomas	Mean age at first diagnosis	Median age at first diagnosis
1: Carcinoma (epithelial tumours)	1,026	83.7	100.0	63.5	64.0
1.1: Serous carcinoma	530	43.2	51.7	63.1	64.0
1.2: Mucinous carcinoma	85	6.9	8.3	62.2	62.0
1.3: Endometrioid carcinoma	103	8.4	10.0	56.8	53.0
1.4: Clear cell carcinoma	63	5.1	6.1	59.6	59.0
1.5: Adenocarcinoma NOS	152	12.4	14.8	69.4	73.0
1.6: Other specified carcinoma	30	2.4	2.9	56.8	57.0
1.7: Unspecified carcinoma	63	5.1	6.1	72.2	78.0
2: Sex cord-stromal tumours	16	1.3		51.9	50.0
3: Germ cell tumours	48	3.9		33.7	29.0
4: Other specified malignant neoplasm	39	3.2		64.2	65.0
5: Unspecified malignant neoplasm	97	7.9		77.1	82.0
Total	1,226	100.0		63.3	64.5

T 11 00 T 11	<i>c</i> •	1	4 11	• • •	<i>.</i> .	0 000
Lable 23. Incidence	re of ovarian cand	er and average	age at diagnos	is hv tvr	ne of ovarian d	cancer 2006
rubic 2.0. meruene	c of ovallall call	ci ana average	age at anaginos	10 0 9 1 9 1	c or ovariant	cancer, 2000

(a) All cases were coded as primary site, invasive ovarian cancers. Appendix Table D2.5 provides a list of the histology types included in each group.

Source: Australian Cancer Database, AIHW.

The average age at the first diagnosis of ovarian cancer differed by histology type (Table 2.3). The mean age of occurrence was highest for those cases in which the type of ovarian cancer was *unspecified* (group 5) (77 years). In contrast, the mean age at diagnosis was lowest for women diagnosed with *germ cell tumours* (group 3) (34 years) and *sex cord-stromal tumours* (group 2) (52 years). The mean age for women diagnosed with *carcinoma* (group 1) was 64 years, but considerable variation is seen across the various types of carcinomas. Those with an *unspecified type of carcinoma* (group 1.7) had a relatively high mean age at diagnosis (72 years), while those in 'group 1.6: *other specified carcinoma*' and 'group 1.3: *endometrioid carcinoma*' had a relatively low mean age (57 years).

Further information about the relationship between age and histological type of ovarian cancer in 2006 is provided in Table 2.4. For each of the three age groups, *carcinoma* (group 1) was the most commonly diagnosed type of ovarian cancer, although differences in the proportion of all ovarian cancers coded to this group of ovarian cancers are evident. Specifically, among those aged less than 50 years, three in four (75%) ovarian cancer cases were *carcinoma*, compared with 91% of those who were aged 50 to 69 years and 80% of those aged 70 years and over at diagnosis. Relative to the other age groups, the younger women were more likely to have been diagnosed with a *germ cell tumour* (group 3) (17% of those in the other age groups). One in 6 cases (16%) of ovarian cancer among those aged 70 years and over was coded as an *unspecified ovarian cancer* (group 5); this compares with 3% of those in the two other age groups.

Trends in types of ovarian cancer

Trends in the proportion of the various types of ovarian cancers are shown in Table 2.5, with the data grouped into four time periods. Caution should be exercised when interpreting these data since changes in histological assessment and coding practices may have affected the observed trends.

There was minimal change in the proportion of ovarian cancers that were classified as *carcinoma* (group 1), with values ranging from 86% (in 2000 to 2006) to 89% (in 1988 to 1993). However, within the group of *carcinomas*, there was change over time in the proportion of a number of the subgroups. In particular, three out of ten (30%) ovarian cancers were classified as *serous carcinoma* (group 1.1) in 1982 to 1987; this proportion had increased to 43% by 2000 to 2006. Over the same period, the proportion of ovarian cancers that were coded as *adenocarcinoma not otherwise specified* (group 1.5) fell from 25% to 14% of ovarian cancers.

For each of the time periods, the proportion of ovarian cancers that were classified as *germ cell tumours* (group 3) remained at around 3%, while the proportion of *sex cord-stromal tumours* (group 2) fell from 2% in 1982 to 1987 to 1% in 2000 to 2006, and the proportion that were coded as *other specified ovarian cancer* (group 4) increased from 2% to 4%. The proportion that was classified as *unspecified ovarian cancer* (group 5) fluctuated over the four time periods, with no clear pattern evident.

Further information on the histological types of ovarian cancers by age group is provided in Appendix Tables D2.6 to D2.8, with those tables showing trends from 1982–1987 to 2000–2006 for the three age groups of those less than 50 years, those 50 to 69 years, and those 70 years and over at diagnosis.

		Number of c	ases			Per c	ent	
Type of ovarian cancer ^(a) <	<50 years	50–59 years	70+ years	Total	<50 years	50–69 years	70+ years	Total
1: Carcinoma (epithelial tumours)	171	495	360	1,026	74.7	90.7	79.8	83.7
1.1: Serous carcinoma	76	283	171	530	33.2	51.8	37.9	43.2
1.2: Mucinous carcinoma	15	42	28	85	6.6	7.7	6.2	6.9
1.3: Endometrioid carcinoma	34	50	19	103	14.8	9.2	4.2	8.4
1.4: Clear cell carcinoma	16	34	13	63	7.0	6.2	2.9	5.1
1.5: Adenocarcinoma NOS	15	53	84	152	6.6	9.7	18.6	12.4
1.6: Other specified carcinoma	ω	18	4	30	3.5	3.3	0.0	2.4
1.7: Unspecified carcinoma	7	15	41	63	3.1	2.7	9.1	5.1
2: Sex cord-stromal tumours	ω	7	~	16	3.5	1.3	0.2	1.3
3: Germ cell tumours	38	თ	~	48	16.6	1.6	0.2	3.9
4: Other specified malignant neoplasm	5	18	16	39	2.2	3.3	3.5	3.2
5: Unspecified malignant neoplasm	7	17	73	67	3.1	3.1	16.2	7.9
Total	229	546	451	1,226	100.0	100.0	100.0	100.0
(a) All cases were coded as primary site, invasive ovarian ca Source: Australian Cancer Database, AIHW.	ancers. Append	lix Table D2.5 provide	s a list of the histology	types included in ea	ch group.			

		Number o	of cases			Per	cent	
Type of ovarian cancer ^(a)	1982–1987	1988–1993	1994–1999	2000-2006	1982–1987	1988–1993	1994–1999	2000-2006
1: Carcinoma (epithelial tumours)	4,587	5,355	5,766	7,140	87.8	88.6	88.2	85.7
1.1: Serous carcinoma	1,539	2,195	2,728	3,555	29.5	36.3	41.8	42.7
1.2: Mucinous carcinoma	597	673	627	586	11.4	11.1	9.6	7.0
1.3: Endometrioid carcinoma	522	559	557	629	10.0	9.3	8.5	8.2
1.4: Clear cell carcinoma	217	307	329	437	4.2	5.1	5.0	5.2
1.5: Adenocarcinoma NOS	1,309	1,161	1,067	1,170	25.1	19.2	16.3	14.0
1.6: Other specified carcinoma	45	68	64	178	6.0	1.1	1.0	2:1
1.7: Unspecified carcinoma	358	392	394	535	6.9	6.5	6.0	6.4
2: Sex cord-stromal tumours	98	88	87	77	1.9	1.5	1.3	0.9
3: Germ cell tumours	155	188	180	270	3.0	3.1	2.8	3.2
4: Other specified malignant neoplasm	119	190	203	293	2.3	3.1	3.1	3.5
5: Unspecified malignant neoplasm	265	221	298	550	5.1	3.7	4.6	6.6
Total	5,224	6,042	6,534	8,330	100.0	100.0	100.0	100.0
(a) All cases were coded as primary site, invasive	e ovarian cancers. Appen	dix Table D2.5 provi	des a list of the histo	ology types included in	each group.			
Source: Australian Cancer Database, AIHW.								

<u>4</u>

Incidence by stage at diagnosis

Stage at diagnosis refers to the extent or spread of cancer at the time of diagnosis. Such information is essential for a number of reasons, including determining an individual's prognosis, assisting in the planning and evaluation of treatment, and contributing to cancer monitoring and research (Odicino et al. 2008; Pecorelli et al. 2000). The fact that ovarian cancer is often diagnosed at an advanced stage is noted repeatedly in the literature and is considered to be one of the major contributors to the high mortality rate for this type of cancer (e.g. Laurvick et al. 2003; Menon & Jacobs 2001; Tracey et al. 2009).

A number of different staging systems are used to classify ovarian cancers, including the International Federation of Gynecology and Obstetrics (FIGO) system, the International Union Against Cancer (UICC) TNM system, and the Surveillance Epidemiology End Results (SEER) Summary Staging system (or 'summary staging system' for short). In both the FIGO and the TNM systems, ovarian tumours are given a value from I (indicating early disease with the tumours confined to the ovaries and of relatively small size) to IV (indicating clearly distant metastatic disease, with metastasis found beyond the peritoneal cavity such as inside the liver, the lungs or other organs). In the summary staging system, which is a simpler system, tumours are allocated to one of three categories: local (the tumour is confined to one or both ovaries); regional (the tumour has spread to surrounding tissue or nearby lymph nodes); and distant (the tumour has spread to distant organs or other parts of the body and has begun to grow at the new location) (Young et al. 2001). Further details about these staging systems are provided in Appendix E.

There is currently no national requirement in Australia for the collection of data on stage at diagnosis and not all states and territories collect this information. In addition, in the data that does exist, varying definitions of ovarian cancer are often used, with some including borderline ovarian tumours (which skew the results towards the less advanced stages) and others including only the staged cases (rather than all cases including those with an unknown stage), thus making comparison difficult with data that considered all cases.

To give an indication of the proportion of ovarian cancers diagnosed at various stages, Table 2.6 presents data from New South Wales (NSW) for 1980 to 2003 (Tracey et al. 2009) and for the United States of America (USA) for 1999 to 2005 (Horner et al. 2009). Although the time periods to which the data apply differ markedly from each other, the advantage of presenting data from both NSW and the USA is that ovarian cancer was defined in these studies in the same way in which it was defined for the purposes of this report (i.e. those cancers coded as 'C56' in ICD-10, with borderline tumours excluded).

Based on NSW data for 1980 to 2003, one-quarter (25%) of staged ovarian cancer cases were diagnosed when the tumour was localised, about two in nine (22%) were diagnosed when the tumour was at the regional stage, while over half (53%) were diagnosed when the tumour was at a distant stage. The 1999 to 2005 data from the USA also show that the majority of staged cases were at the distant stage when diagnosed, although the proportion is considerably higher (67%) than suggested by the NSW data. In addition, in 7% of the USA ovarian cancer cases, the stage at diagnosis was unknown.

Change over time in the proportion of ovarian and other female genital organ cancers (i.e. ICD-10 codes of C56 and C57.0 to C57.7) that are diagnosed at different stages in NSW are shown in Table 2.7 (Tracey et al. 2007). The proportion of cases that had an unknown stage at diagnosis was similar over the two time periods considered (13% and 12% respectively).

Stage at	New South W	ales (1980–2003) ^(b)	United States of America (1999–2005) ^(b,c)			
diagnosis ^(a)	No. of cases	% of staged cases	No. of cases	% of all cases	% of staged cases ^(d)	
Localised	1,763	25.4	n.a.	15	16	
Regional	1,522	21.9	n.a.	17	18	
Distant	3,649	52.6	n.a.	62	67	
Unknown	n.a.		n.a.	7		
Total	6,934	100.0	29,168	100	100	

Table 2.6: Incidence of ovarian cancer by stage at diagnosis, New South Wales and United States of America

(a) Based on the SEER summary staging system. 'Localised' tumours were defined as those that were confined to one or both ovaries, 'regional' tumours had spread to surrounding tissue or nearby lymph nodes and 'distant' tumours had spread to distant organs or other parts of the body and had begun to grow at the new location (see Appendix E).

(b) Includes ovarian cancers coded in ICD-10 as C56; borderline ovarian tumours are excluded.

(c) Data were from the SEER 17 areas which cover approximately a quarter of the USA (see Table 21.7 in Horner et al. 2009).

(d) These values are approximations that were calculated by the AIHW since only the percentage (rather than the exact number) of unknown cases was provided.

Source: Tracey et al. 2009; Horner et al. 2009.

Meanwhile, the proportion that was at the distant stage at diagnosis increased over time from 50% of staged cases in 1980–1998 to 60% of staged cases in 1999–2003. Tracey and associates (2009) suggest that this increase may reflect improvements in testing sensitivity due to advances in imaging and other diagnostic testing.

Studies on ovarian cancer based on Californian data (Morris et al. 2008) and Florida data (FDH 2009) also found an increase over time in the proportion of ovarian cancer cases diagnosed with distant metastases. In contrast, other studies have not found such an increase, with one example being a study using Danish data (Kjaerbye-Thygesen et al. 2005). In that study, the proportion of cases with distant metastases at diagnosis decreased significantly between 1978–1982 and 1998–2002. The authors of the Danish study suggest that the different trends observed internationally may be due to the use of different staging systems and practices for classifying stage at diagnosis in different countries.

1980–1998			1999–2003			
Stage at diagnosis ^(b)	No. of cases ^(a)	% of all cases	% of staged cases	No. of cases ^(a,c)	% of all cases	% of staged cases
Localised	1,422	23.1	26.5	368	19.0	21.7
Regional	1,266	20.5	23.6	302	15.6	17.8
Distant	2,685	43.5	50.0	1,023	52.9	60.4
Unknown	796	12.9		240	12.4	
Total	6,169	100.0	100.0	1,933	100.0	100.0

Table 2.7: Incidence of ovarian and other female genital organ cancers^(a) by stage at diagnosis, New South Wales, 1980–1998 and 1999–2003

(a) Includes ovarian and other female genital organ cancers coded in ICD-10 as C56 and C57.0–C57.7 (see Appendix Table B.2); excludes borderline ovarian tumours.

(b) Based on the SEER summary staging system. 'Localised' tumours were defined as those that were confined to one or both ovaries, 'regional' tumours had spread to surrounding tissue or nearby lymph nodes and 'distant' tumours had spread to distant organs or other parts of the body and had begun to grow at the new location (see Appendix E).

(c) The number of cases for 1999 to 2003 were derived from Table 25 in Tracey et al. 2007.

Source: Tracey et al. 2007.

Table 2.8 presents information on the stage of ovarian cancer according to age at diagnosis, using data from the USA for 1999 to 2005. Older women were more likely to be diagnosed with advanced stage ovarian cancer (69% of women aged 65 years and over) than other women (55% of those aged less than 65 years). This difference by age has also been observed in other research (e.g. Grossi et al 2002; Kosary 2007; WHC & NCRI 2006). In addition, the USA data indicate that an unknown stage at diagnosis was also more likely among older women than others (10% and 4% of cases, respectively).

		<65 years	;		65+		All ages		
Stage at diagnosis ^(b)	No. of cases	% of all cases	% of staged cases ^(c)	No. of cases	% of all cases	% of staged cases ^(c)	No. of cases	% of all cases	% of staged cases ^(c)
Localised	n.a.	21	22	n.a.	7	8	n.a.	15	16
Regional	n.a.	20	21	n.a.	13	14	n.a.	17	18
Distant	n.a.	55	58	n.a.	69	77	n.a.	62	67
Unknown	n.a.	4		n.a.	10		n.a.	7	
Total	16,000	100	100	13,168	100	100	29,168	100	100

Table 2.8: Incidence of ovarian cancer^(a) by stage and age at diagnosis, United States of America,1999-2005

(a) Includes ovarian cancers coded in ICD-10 as C56; borderline ovarian tumours were excluded. Data were from the SEER 17 areas which cover approximately a quarter of the USA (see Table 21.7 in Horner et al. 2009).

(b) Based on the SEER summary staging system. 'Localised' tumours were defined as those that were confined to one or both ovaries, 'regional' tumours had spread to surrounding tissue or nearby lymph nodes and 'distant' tumours had spread to distant organs or other parts of the body and had begun to grow at the new location (see Appendix E).

(c) These values are approximations since only the percentage (rather than the exact number) of unknown cases was provided.

Source: Horner et al. 2009.

Differences across groups

In this section, data on the incidence of ovarian cancer are provided according to geographical area, socioeconomic status, Aboriginal and Torres Strait Islander status and country of birth. In order to take into account differences in the age structures and the size of the groups being compared, age-standardised rates are provided for each of the comparisons. The data are presented for the 5-year period of 2002 to 2006 rather than for just 1 year since presenting the data for multiple years reduces random variation in the data. This is especially important for comparisons of small sub-groups (e.g. Indigenous women or women in smaller states and territories).

Observed differences by the characteristics examined in this section may result from a number of factors including variation in:

- population characteristics (e.g. a relatively greater proportion of Indigenous women in remote areas)
- the prevalence of risk and/or protective factors (e.g. reproductive patterns, use of the contraceptive pill)
- the availability of diagnostic services.

Differences by geographical area

In the 2002 to 2006 period, the largest average number of ovarian cancer cases diagnosed each year was in New South Wales (408 cases annually) and the smallest number in the Northern Territory (7 cases annually) (Table 2.9). When the age structure and size of the population in each state and territory was taken into account, the results indicate that the Northern Territory had the lowest incidence rate (9.1 cases per 100,000 females), followed by South Australia (9.4 per 100,000) and Tasmania (9.5 per 100,000). While the rates for the Northern Territory and Tasmania did not differ significantly from that of the other states and territories, the rate for South Australia was significantly lower than the rates for New South Wales (11.0 per 100,000), Victoria (11.8 per 100,000) and Western Australia (12.2 per 100,000). The highest incidence rates were observed for Western Australia and the Australian Capital Territory, with 12.2 cases of ovarian cancer diagnosed per 100,000 females in each of those jurisdictions. While the rate for the Australian Capital Territory was not significantly different from that of other states and territories, the rate for the Australian Capital Territory was not significantly different from that of other states and territories, the rate for Western Australia (10.5 and 9.4 cases per 100,000 females).

State or territory	Average annual number of cases ^(a)	Total number of cases	Age-standardised rate ^(b)	95% confidence interval
New South Wales	408	2,042	11.0	10.5–11.5
Victoria	327	1,633	11.8	11.2–12.4
Queensland	213	1,066	10.5	9.9–11.2
Western Australia	123	617	12.2	11.3–13.2
South Australia	89	446	9.4	8.6–10.4
Tasmania	28	138	9.5	8.0–11.3
Australian Capital Territory	19	93	12.2	9.8–15.0
Northern Territory	7	33	9.1	5.8–13.3
Total	1,214	6,068	11.1	10.8–11.3

Table 2.9: Incidence of ovarian canc	er by state and territory, 2002–2006
--------------------------------------	--------------------------------------

(a) Numbers may not sum to the total due to rounding.

(b) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females. The rates are based on the total number of cases over the 5-year period from 2002–2006.

Source: Australian Cancer Database, AIHW.

Age-standardised incidence rates according to level of remoteness of the area in which the women lived at diagnosis are shown in Figure 2.5. The Australian Standard Geographical Classification (ASGC) Remoteness Areas was used to categorise areas of Australia. Information about this classification is provided in Appendix A. While the incidence rate for women who lived in *Remote and very remote* areas at diagnosis (12.2 cases per 100,000 females) was higher than those for other women, the difference was not statistically significant.



Differences by socioeconomic status

The findings from studies that have addressed the associations between socioeconomic status and the risk of ovarian cancer have been inconsistent. For example, there was no significant association found using South Australian data for 1977 to 2001 (Cancer Council SA 2009) and Irish data for 2000 to 2004 (Donnelly et al. 2009), while disadvantaged women in Queensland were significantly more likely than more advantaged women to be diagnosed with ovarian cancer between 1996 and 2002 (Baade et al. 2005).

In this report, the Index of Relative Socio-economic Disadvantage (IRSD) is used to indicate socioeconomic status (ABS 2008a), with the first group (which is labelled '1') corresponding to geographical areas containing the 20% of the population with the lowest socioeconomic status according to the IRSD and the fifth group corresponding to the 20% of the population with the highest status. Appendix A provides further information about the IRSD.

As shown in Figure 2.6, there was no statistically significant association between the incidence of ovarian cancer and socioeconomic status in Australia in 2002 to 2006.



Differences by Aboriginal and Torres Strait Islander status

Across a range of health-related and socioeconomic indicators, Aboriginal and Torres Strait Islander people are disadvantaged relative to other Australians. They are more likely to live in remote areas of Australia and have a relatively younger age structure, with a median age of 21 years compared with 37 years for the non-Indigenous population (ABS & AIHW 2008). This age difference is thought to be largely due to higher rates of fertility, as well as a shorter life expectancy among the Indigenous population (ABS 2009g,h).

Reliable national data on the incidence of cancer for Indigenous women are not available. While all state and territory cancer registries collect Indigenous status information, the quality of the data in some areas is insufficient for analysis. In this report, data for four states and territories – Queensland, Western Australia, South Australia and the Northern Territory – are used to examine the incidence of ovarian cancer by Indigenous status. While the majority (60%) of Australian Indigenous women live in these four jurisdictions (ABS 2009f), the degree to which data for these jurisdictions are representative of data for all Indigenous women is unknown. Furthermore, due to the small number of Indigenous women who had been diagnosed with ovarian cancer in the four jurisdictions between 2002 and 2006 – namely, an average of 8 women per year – caution should be taken when making use of the data on ovarian cancer rates by Indigenous status.

For the four jurisdictions, the level of missing data on Indigenous status for ovarian cancer cases diagnosed between 2002 and 2006 was 3% (Table 2.10).

Although the age-standardised incidence rate for Indigenous women was slightly higher than that for non-Indigenous women (12.4 and 11.2 cases per 100,000 females, respectively), the difference was not statistically significant.

Table 2.10: Incidence of ovarian cancer by Indigenous status, Queensland, Western Australia, South Australia and the Northern Territory, 2002–2006

Indigenous status	Annual average number of cases ^(a)	Total number of cases	Age-standardised rate ^(b,c)
Indigenous	8	38	12.4
Non-Indigenous	410	2,050	11.2
Not stated	15	74	
Total	432	2,162	11.6

(a) Numbers may not sum to the total due to rounding.

(b) Indirectly age-standardised to the 2002–2006 non-Indigenous population for the four jurisdictions (see Appendix B) and based on the total number of cases over the 5-year period from 2002–2006.

(c) The 95% confidence interval around the age-standardised rate for Indigenous women is 8.8–17.0.

Source: Australian Cancer Database, AIHW.

A study using Northern Territory data for the years from 1991 to 2001 also found no significant difference in ovarian cancer incidence by Indigenous status (Condon 2004), as did a study using Queensland data for 1982 to 1996 (Coory et al. 2000). A study using New Zealand data for 2005 (NZ Ministry of Health 2009a, 2009b) also showed no significant difference by Indigenous status, whereas data from the USA indicated that American Indians and Alaska native women had a significantly lower incidence rate of ovarian cancer than their white counterparts (USCSWG 2009; Wiggins et al. 2008).

Differences by country of birth

Australia has one of the largest proportions of immigrant populations in the world. In 2006, it was home to 4.4 million overseas-born people and one in four (25%) residents was born outside of the country (ABS 2009a). Research has found that most migrants are at least as healthy, if not more so, as the Australian-born population. The 'healthy migrant effect' is believed to result from two main factors: a self-selection process in which those people who are physically and economically able to migrate are the ones who do; and government eligibility criteria for migrants based on health, education, language and job skills (AIHW 2008a). However, research suggests that this migrant health advantage decreases over time. In regard to ovarian cancer, it is thought that incidence rates for migrants also tend to converge with that of the host country over time, particularly so for any offspring (Kliewer & Smith 1995; Parkin & Iscovich 1997).

Immigrants are more likely than Australian-born people to live in urban areas (ABS 2009a); this often provides immigrants with relatively easier access to health care services. At the same time, though, language and cultural barriers may mean that some immigrants are less likely or able to access available services.

In the earlier edition of this report (AIHW & NBCC 2006), data on ovarian cancer incidence by country of birth were only available for New South Wales, whereas in this edition, national data are provided. Note that these data do not take into account the length of time the immigrants lived in Australia, although some groups – for instance, people from Asia – tend to be more recent immigrants while people from many European countries tend to have been in Australia for longer periods of time (ABS 2009a). In this report, country of birth data were classified using the Standard Australian Classification of Countries (SACC), second edition. Further information about this classification can be found in Appendix A. Information on the woman's country of birth was not available for 5% of the ovarian cancer cases diagnosed in Australia between 2002 and 2006.

Figure 2.7 shows that women living in Australia who were born in the Americas had the highest rate of ovarian cancer (13.7 cases per 100,000 females), with this rate significantly higher than the rates for women born in Australia (9.9 per 100,000). The lowest age-standardised incidence rate was seen among women who were born in North-East Asia (9.3 per 100,000). This rate was not significantly different from that of those who were born in Australia.



International comparisons

In this section of the report, the incidence rate of ovarian cancer in Australia is compared with the rate for other countries using data from the GLOBOCAN database – a database which is prepared by the International Agency for Research on Cancer (IARC) (Ferlay et al. 2004). The most recent GLOBOCAN estimates are for 2002, with these estimates based on cancer incidence rates from approximately two to five years earlier. The GLOBOCAN data for ovarian cancer pertain to 'ovarian and related cancers' (that is, the ICD-10 codes of C56 and C57.0–C57.4) and thus encompass a broader range of cancers than is generally considered in this report. However, in line with the definition of ovarian cancer used in this report, borderline ovarian tumours were not included in the 2002 GLOBOCAN data. See

Appendix C for further details about this database. As discussed in Chapter 1, caution must be taken when comparing data from different countries since observed differences may be due to a range of methodological factors, not just differences in the underlying rates.

The estimated number of new cases of ovarian and related cancers around the world in 2002 was approximately 204,500 (Appendix Table D2.12). Figure 2.8 shows the estimated incidence rates of these cancers by country (for Australia and New Zealand) and by region. The estimated age-standardised rate for Australia was 8.9 new cases per 100,000 females. Thus, Australia was estimated to have a significantly higher incidence rate than the average world rate (6.6 cases per 100,000 females) and than of all the African and Asian regions.



However, Australia's rate was significantly lower than the rates of all other Westernised countries and regions, including New Zealand (12.4 per 100,000), Northern America (10.7 per 100,000) and all of the European regions (which ranged from 9.7 to 13.3 cases per 100,000 females).

The differences by region and country in the incidence rates for ovarian and related cancers may be due to a number of factors, including differences in diagnostic and classification practices, completeness of cancer registration, the proportion of women with various risk and protective factors (e.g. parity and use of oral contraceptives) and genetic susceptibility (Bray et al. 2005; Colombo et al 2006; Kliewer & Smith 1995; Kricker 2002).
3 Mortality from ovarian cancer

The number of deaths from ovarian cancer in a given time period is a result of the incidence of ovarian cancer, as well as factors that affect the likelihood of fatality such as the characteristics of the ovarian cancers diagnosed (e.g. stage at diagnosis and histological type of ovarian cancer), and the nature and quality of treatments received.

In this report, mortality refers to the number of deaths for which the underlying cause was a primary site ovarian cancer. The ovarian cancer that led to the death may have been diagnosed many years previously, in the same year in which the person died or, in some cases, after death (e.g. at autopsy). Information on the underlying cause of death is derived from the medical certificate of cause of death which is issued by a certified medical practitioner.

The main data source used in this chapter was the National Mortality Database. This database contains information about deaths due specifically to ovarian cancer from 1968 onwards (see Appendix C for further information).

In this chapter, information is presented on the number of deaths attributed to ovarian cancer in 2006, as well as in previous years. In addition, differences in mortality rates according to age, geographical area, socioeconomic status, Indigenous status and country of birth are provided. Lastly, mortality rates for Australia and other countries are compared.

Mortality in 2006

A total of 795 women died from ovarian cancer in 2006 (Table 3.1); thus, across Australia, an average of 2 women died every day from this disease. Ovarian cancer was the sixth most

Cancer type (ICD-10 codes)	No. of deaths	% of all gynaecological cancer deaths	% of all cancer deaths	% of all deaths	ASR ^(a)	95% confidence interval
Lung (C33–C34)	2,683		15.7	4.1	22.7	21.8–23.6
Breast (C50)	2,618		15.3	4.0	22.1	21.3–23.0
Unknown primary site (C26, C39, C76–C80)	1,917		11.2	2.9	15.1	14.5–15.8
Bowel (C18–C20)	1,675		9.8	2.6	13.6	12.9–14.2
Pancreas (C25)	1,029		6.0	1.6	8.4	7.9–8.9
Ovary (C56)	795	54.7	4.6	1.2	6.7	6.2–7.2
All lymphomas (C81–C85,C96)	669		3.9	1.0	5.4	5.0–5.8
All leukaemias (C91–C95)	609		3.6	0.9	5.0	4.6–5.4
Melanoma (C43)	452		2.6	0.7	3.8	3.5–4.2
Stomach (C16)	448		2.6	0.7	3.6	3.3–4.0
All cancers ^(b)	17,123		100.0	26.3	141.0	138.9–143.2

Table	3.1:	The	10 most	common	types	of	cancer	deaths.	females.	2006
Iuvic	U.T.	THE	io most	common	ypco	OI.	current	acatilo,	remarco,	-000

(a) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females.

(b) Includes cancers coded in the ICD-10 as C00–C97, D45, D46, D47.1 and D47.3.

Source: National Mortality Database, AIHW.

common cause of cancer death among women. Mortality from ovarian cancer accounted for 5% of all female cancer deaths and 1% of all female deaths. Furthermore, ovarian cancer was the most common cause of gynaecological cancer death in 2006, representing over half (55%) of such deaths. This means that more women died from ovarian cancer than from all of the other gynaecological cancers combined.

The age-standardised mortality rate of ovarian cancer in 2006 was 6.7 deaths per 100,000 females. The corresponding rates for lung cancer and breast cancer were 22.7 and 22.1 (deaths per 100,000 females), respectively.

Differences by age at death

In 2006, 1% of deaths due to ovarian cancer occurred among women aged under 40 years, just over one-fifth (22%) occurred among those aged 40 to 59 years, while the over three in four deaths (77%) occurred among women aged 60 years and over (Appendix Table D3.1). Figure 3.1 presents mortality rates of ovarian cancer according to age at death for 2006. As a point of comparison, incidence rates by age at diagnosis are also shown. Similar to the incidence rates, the likelihood of women dying from ovarian cancer increased with age. The mortality rates were less than 2 deaths per 100,000 females for those aged up to and including 40 to 44 years. From that age group onwards, there was an increase in the mortality rates between most age groups (although in most cases the difference from one age group to the next was not statistically significant). The highest mortality rate of 48.7 deaths per 100,000 females was observed for the oldest age group (i.e. those aged 85 years and over), with this rate significantly higher than the rates for all age groups up to and including 70 to 74 years.



Trends

The first year for which national mortality data for ovarian cancer are available is 1968. In Figure 3.2, the number of deaths due to ovarian cancer and the corresponding age-standardised mortality rates are shown for 1968 to 2006. For comparison purposes, age-standardised incidence rates are also shown.

Despite some year-to-year fluctuations, the number of deaths from ovarian cancer increased over time. In 1968, 451 women died from ovarian cancer; in 1990, this number had risen to 705 women, while 795 women had died from ovarian cancer in 2006. Between 1968 and 2006, the number of deaths from ovarian cancer increased by 76%.

When the age-standardised rates are considered, some year-to-year fluctuations are again seen. However, overall, the mortality rate decreased significantly by 26% between 1968 (9.1 deaths per 100,000 females) and 2006 (6.7 per 100,000). In addition, the age-standardised mortality rate for 2006 was the lowest rate for any year to date (although it was not significantly lower than the rates observed for three of the preceding years).



Possible explanations for the decline in the mortality rate from ovarian cancer over time include the following:

- a decrease in the incidence rate of ovarian cancer (as discussed in Chapter 2)
- improvements in access to and quality of treatments, including greater sub-specialisation in gynaecological oncology, more aggressive surgery, the availability of more effective types of chemotherapy, and better multidisciplinary care (Kjaerbye-Thygesen et al. 2005; Oriel et al. 1999; Tracey et al. 2008)
- change over time in the histological types of ovarian cancers occurring among women (as discussed in Chapter 4, the prognosis is better for some types of ovarian cancers than others).

Trends by age at death

Trends in ovarian cancer morality rates by age for the 25-year period from 1982 to 2006 are shown in Figure 3.3. Between 1982 and 2006, the mortality rate due to ovarian cancer decreased slightly but significantly for those aged less than 50 years at death, from 1.7 per 100,000 females in 1982 to 0.8 per 100,000 females in 2006. The rate also fell significantly for women aged 50 to 69 years at death, with a decline of 39% (20.7 deaths per 100,000 females in 1982 to 12.7 per 100,000 in 2006). In contrast, for women aged 70 years and over at death, numerous year-to-year fluctuations in the mortality rate are observed, with an overall statistically significant increase in the ovarian cancer mortality rate of 9% from 1982 to 2006 (36.6 and 40.3 deaths per 100,000 females, respectively). Thus, the trend in the mortality rates for women aged 70 years and over was in the opposite direction to the trend for those aged less than 70 years.



While the trend of increasing ovarian cancer mortality rates for older women has also been observed elsewhere (e.g. in data from the USA reported by Oriel et al. 1999), the reasons for this increase are not clear. Part of the explanation may be that, unlike those aged 50 to 69 years, the ovarian cancer incidence rate did not fall for those aged 70 years and over (see Chapter 2). Second, it is possible that the benefits of improvements in treatment (such as the improved treatment of germ cell cancers) have been relatively smaller for older women than younger women (Bray et al. 2005; Cancer Council Victoria 2007; Oriel et al. 1999). Third, change over time in the relative proportion of older women (compared with other women) who are diagnosed with types of ovarian cancer that have a poorer prognosis may also help explain this trend.

Note that while it is well established that older women tend to be diagnosed at a later stage than younger women (see Chapter 2) and that older women are, in general, given less aggressive treatment than their younger counterparts (Maas et al. 2005; Petignat et al. 2004; WHC & NCRI 2006), these differences by age do not help explain the rise in mortality rates

among older women unless those trends have worsened over time (Oriel et al. 1999). In conclusion, the reasons for increasing ovarian cancer mortality rates for older women are not clearly understood and further investigation on this topic could be warranted.

Risk of death from ovarian cancer and average age at death

Based on 2006 data, the risk of a woman in the general population dying from ovarian cancer before the age of 75 years was 1 in 206; the corresponding risk for the age of 85 was 1 in 108 (Table 3.2). Although these risk levels have fluctuated over the years, the general pattern has been one of a decrease in the risk of a woman dying from ovarian cancer. For example, based on 1982 data, the risk of dying from ovarian cancer by the age of 85 years was 1 in 89 compared with the risk of 1 in 108 calculated from the 2006 data.

The average age at death due to ovarian cancer has increased over time. The mean age of death of women who died from ovarian cancer increased from 65 years in 1982 to 70 years in 2006, while the median age increased from 65 years to 72 years over the same period.

Year	Risk to age 75 years	Risk to age 85 years	Mean age at death	Median age at death
1982	1 in 145	1 in 89	64.9	65.0
1983	1 in 146	1 in 95	64.6	64.0
1984	1 in 140	1 in 90	65.0	66.0
1985	1 in 155	1 in 101	65.9	66.0
1986	1 in 144	1 in 94	65.1	66.0
1987	1 in 156	1 in 99	65.3	65.0
1988	1 in 166	1 in 100	65.4	66.0
1989	1 in 156	1 in 100	66.3	67.0
1990	1 in 142	1 in 88	66.5	68.0
1991	1 in 148	1 in 92	67.0	68.0
1992	1 in 161	1 in 101	66.6	68.0
1993	1 in 155	1 in 94	68.1	69.0
1994	1 in 159	1 in 92	66.2	68.0
1995	1 in 169	1 in 97	67.7	69.0
1996	1 in 162	1 in 90	67.8	70.0
1997	1 in 178	1 in 95	68.8	71.0
1998	1 in 176	1 in 95	69.0	71.0
1999	1 in 194	1 in 102	69.4	72.0
2000	1 in 178	1 in 100	69.2	71.0
2001	1 in 179	1 in 94	70.0	72.0
2002	1 in 173	1 in 93	69.2	71.0
2003	1 in 199	1 in 104	69.0	71.0
2004	1 in 173	1 in 94	69.6	71.0
2005	1 in 193	1 in 97	70.6	73.0
2006	1 in 206	1 in 108	70.3	72.0

Table 3.2: Risk of death from ovarian cancer and average age at death, 1982 to 2006

Note: The 1982 to 1996 data were adjusted from ICD-9 to ICD-10 standards using a factor of 0.98.

Source: National Mortality Database, AIHW.

Differences across groups

In this section of the report, differences in mortality of women from ovarian cancer are presented according to geographical area, socioeconomic status, Indigenous status and country of birth. Any observed differences among the groups compared may be due to a number of reasons, including differences in incidence rates of ovarian cancer, the characteristics of the ovarian cancers diagnosed (e.g. stage at diagnosis and type of tumour), and access to and quality of treatment. Similar to what was done to examine differences in incidence across groups, age-standardised rates for the 5-year period from 2002 to 2006 are compared.

Differences by geographical area

The average number of ovarian cancer deaths per year over the period from 2002 to 2006 ranged from 283 in New South Wales to 2 in the Northern Territory (Table 3.3). When the age-standardised rates are considered, the Northern Territory had the lowest rate (3.2 deaths per 100,000 females), with this rate significantly lower than the rates for Victoria (8.1 per 100,000 females), Western Australia (7.6 per 100,000) and New South Wales (7.3 per 100,000). The highest age-standardised rates were observed for the Australian Capital Territory (8.3 per 100,000) and Victoria (8.1 per 100,000). While the rate for the Australian Capital Territory did not differ significantly from that of other states and territories, the rate for Victoria was significantly higher than that observed for Queensland (6.8 per 100,000), South Australia (6.5 per 100,000) and the Northern Territory (3.2 per 100,000).

State or territory ^(a)	Average annual number of deaths	Total number of cases	Age-standardised rate ^(b)	95% confidence interval
New South Wales	283	1,414	7.3	6.9–7.7
Victoria	234	1,168	8.1	7.7–8.6
Queensland	139	693	6.8	6.3–7.3
Western Australia	77	385	7.6	6.8–8.4
South Australia	64	322	6.5	5.8–7.3
Tasmania	21	106	7.2	5.9–8.7
Australian Capital Territory	12	60	8.3	6.3–10.8
Northern Territory	2	10	3.2	1.2–6.3
Total	832	4,158	7.3	7.1–7.6

Table 3.3: Mortality	v from	ovarian	cancer	by state	and	territory	. 2002-	-2006
i ubic 0.0. montalit	, mom	0 v ul lull	cuncer	by blutt	unu	cerricol y		2000

(a) These data may not be comparable with data published in state and territory cancer reports since the data shown in this report relate to the place of residence at the time of *death*, not the place of residence at the time of *diagnosis* as is often shown in state and territory reports. Furthermore, the states and territory cancer registries tend to use a different methodology from that used by the AIHW to determine the cause of death (see Appendix B).

(b) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females. The rates are based on the total number of cases over the 5-year period from 2002–2006.

Source: National Mortality Database, AIHW.

In Figure 3.4, the ovarian cancer mortality rates for women living in different remoteness areas are presented. The mortality data are based on the remoteness of the usual place of residence of the women at time of death. During the 2002 to 2006 period, women living in *Remote and very remote* areas had the highest mortality rate (9.6 deaths per 100,000 females),

with this rate being significantly higher than the rate for those living in *Major cities* (7.2 per 100,000). This difference may be related to a number of factors including access to diagnostic and other health services in remote areas.



Differences by socioeconomic status

The socioeconomic status measure used in this report pertains to the characteristics of people in the area in which the women lived, rather than to the characteristics of the individual (see Appendix A). From 2002 to 2006, women living in areas with the highest socioeconomic status had the highest ovarian cancer mortality rate (7.8 deaths per 100,000 females) but this rate was not significantly higher than the rates for women living in other areas (Figure 3.5).

Differences by Aboriginal and Torres Strait Islander status

Information in the National Mortality Database (NMD) on Indigenous status for 2002 to 2006 is considered to be of sufficient quality for use for five jurisdictions: New South Wales, Queensland, Western Australia, South Australia and the Northern Territory. Almost nine in ten (89%) Indigenous women live in these five jurisdictions (ABS 2009f). In the NMD, the Indigenous status of 1% of the women who had died from ovarian cancer was not known (Table 3.4). Between 2002 and 2006, there was an annual average of 6 ovarian cancer deaths recorded for Indigenous women in the five jurisdictions. Due to the relatively small number of Indigenous women who died from ovarian cancer, caution should be taken when considering differences in mortality rates by Indigenous status.



There was no statistically significant difference in the age-standardised mortality rates from ovarian cancer for Indigenous women compared with non-Indigenous women (8.4 deaths per 100,000 females among Indigenous women and 7.9 per 100,000 among non-Indigenous women). This finding of a lack of significant difference by Indigenous status mirrors those of a number of other studies, including research using data from the Northern Territory (Condon 2004; Zhang et al. 2008), New South Wales (Supramaniam et al. 2006) and New Zealand (NZ Ministry of Health 2009a,b). In contrast, data from the United States of America indicated that American Indian and Alaska native women had a significantly lower age-standardised mortality rate from ovarian cancer than white American women (5 and 9 deaths per 100,000 females, respectively) (USCSWG 2009).

Table 3.4: Mortality from ovarian cancer by Indigenous status, New South Wales, Queensland, Western Australia, South Australia and the Northern Territory, 2002–2006

Indigenous status	Average annual number of deaths ^(a)	Total number of cases	Age-standardised rate ^(b,c)
Indigenous	6	30	8.4
Non-Indigenous	553	2,764	7.9
Not stated	6	30	
Total	565	2,824	8.0

(a) Numbers may not sum to the total due to rounding.

(b) Indirectly age-standardised to the 2002–2006 non-Indigenous population for the five jurisdictions (see Appendix B) and are based on the total number of cases over the 5-year period from 2002–2006.

(c) The 95% confidence interval around the age-standardised rate for Indigenous women is 5.7-12.0.

Source: National Mortality Database, AIHW.

Differences by country of birth

From 2002 to 2006, women living in Australia who were born in Sub-Saharan Africa had the highest age-standardised mortality rate (10.6 deaths per 100,000 females), with this rate significantly higher than the rate for women born in Australia (7.2 per 100,000) (Figure 3.6). The lowest mortality rates were observed for women born in the Americas (5.9 per 100,000), South-East Asia (6.0 per 100,000) and North-East Asia (6.0 per 100,000); these rates were not significantly higher than the rate for those born in Australia.



Figure 3.6: Mortality from ovarian cancer by country/region of birth, 2002-2006

International comparisons

As discussed in Chapter 1, caution must be taken when comparing international data on cancer mortality since observed differences may be due to a range of factors, not just differences in the underlying mortality rates. Data on ovarian cancer deaths from the GLOBOCAN database (Ferlay et al. 2004) are shown in Figure 3.7. The confidence intervals indicate the variation that would be expected by chance, assuming that the estimated mortality rates are accurate. Note that the GLOBOCAN data pertain to both 'ovarian and related cancers' (specifically, the ICD-10 codes of C56 and C57.0–C57.4) rather than to just 'ovarian cancer'. In addition, the data are estimates for 2002 and are based on information from around 3 to 5 years earlier. Further information about these data is provided in Appendix C.

The age-standardised mortality rate from ovarian and related cancers for Australia (4.9 deaths per 100,000 females) was significantly lower than that for a number of other



Westernised countries and regions including Northern Europe (7.9 per 100,000), New Zealand (6.4 per 100,000), Western Europe (6.3 per 100,000) and Northern America (6.1 per 100,000). Meanwhile, Australia's mortality rate from ovarian and related cancers was estimated to be significantly higher than the rate for each of the African and Asian regions (which ranged from 1.8 to 4.1 deaths per 100,000 females).

Ovarian cancer as an associated cause of death

The data presented thus far in this chapter apply to deaths of women for which the *underlying* cause of death was ovarian cancer. In addition to an underlying cause of death, *associated* causes of death can be listed on a death certificate. An associated cause of death is

any other condition or event that was not the underlying cause of death, but was considered to contribute to the individual's death. In this section, data are presented on deaths of women for which ovarian cancer was the associated (but not underlying) cause of death.

An annual average of 59 women who died between 2002 and 2006 had ovarian cancer recorded as an associated cause of death (Table 3.5). For most of these deaths (23 deaths or 38% of the relevant deaths), a circulatory system disease was the underlying cause of death. The second most common underlying cause of death for those deaths for which ovarian cancer was an associated cause of death was a cancer other than ovarian cancer (average of 15 deaths per year and 25% of the relevant deaths). The most common type of these 'other' cancers was 'cancer of independent (primary) multiple sites' (ICD-10 code of C97), followed by breast cancer (ICD-10 code of C50), with the corresponding average number of deaths being 4 and 2 per year, respectively.

Underlying cause of death	ICD-10 codes	Average annual number of deaths ^(a)	% of deaths
Circulatory system disease	100–199	23	38.4
Cancer (other than ovarian cancer)	C00–C55, C57–C97, D45–D46, D47.1, D47.3	15	24.9
Respiratory system disease	J00–J99	4	7.1
Nervous system disease	G00–G99	1	2.4
Endocrine, nutritional and metabolic disease	E00–E89	3	4.4
Digestive system disease	K00–K93	7	12.1
Mental and behavioural disorder	F00–F99	1	1.3
Other	all other ICD-10 codes (except for C56)	6	9.4
Total		59	100.0

Table 3.5: Underlying o	cause of death where	e ovarian cancer	was an associated	l cause, annual average
for 2002–2006				

(a) Numbers may not sum to the total due to rounding.

Source: National Mortality Database, AIHW

During 2002 to 2006, the majority (75%) of women who died with ovarian cancer as an associated cause of death were aged 70 years or older at death, while 24% were 50 to 69 years and 2% were less than 50 years (Table 3.6).

Table 3.6: Women who died with ovarian cancer as an associated cause by age at death, annual average for 2002–2006

Age at death	Average annual number of deaths ^(a)	Per cent of deaths
<50 years	1	1.7
50–69 years	14	23.7
70+ years	44	74.6
Total	59	100.0

(a) Numbers may not sum to the total due to rounding.

Source: National Mortality Database, AIHW.

4 Survival after a diagnosis of ovarian cancer

Information on the survival of those who are diagnosed with ovarian cancer provides an indication of the effect of cancer and the success of cancer control programs and treatments. Survival estimates provide information on the probability that a person will still be alive at a specified point in time (such as 1 or 5 years) after the diagnosis of cancer. Survival is influenced by a range of factors including the characteristics of those diagnosed with cancer (e.g. age, sex, additional illnesses and lifestyle), the nature of the tumours (e.g. stage at diagnosis and histology type), and the health-care system (e.g. screening, diagnostic and treatment facilities, and follow-up services) (Black et al. 1998; WCRF & AICR 2007).

Two different measures of survival from cancer can be presented, namely, *crude survival* and *relative survival*. Crude survival indicates the proportion of people alive at a specified point in time subsequent to diagnosis of cancer; it does not take into account the fact that some people diagnosed with cancer – for example, older persons – may have a relatively shorter lifespan than the rest of the population (regardless of their diagnosis of cancer) due to other illnesses. Relative survival takes this issue into account and it is thus a more meaningful measure of outcomes from cancer. Relative survival involves the comparison of the survival of people diagnosed with cancer (i.e. observed survival) with that experienced by a population of equivalent age, sex and calendar year in the general population (i.e. expected survival). The ratio of observed to expected survival is used to estimate the proportion of people who would have survived their cancer. As detailed more fully in Appendix B, relative survival can be calculated in a number of different ways; the 'cohort method' was used for this report.

Relative survival is generally presented as a proportion, with a value less than 100% suggesting that those with ovarian cancer had a lower chance of survival than the general population. For example, 5-year relative survival of 40% for females diagnosed with ovarian cancer means that these females had a 40% chance of surviving 5 years after diagnosis relative to comparable women in the general population.

Since relative survival estimates are based on the outcomes of a group of people with a diverse mix of ovarian cancer and other characteristics, they provide an indication of the *average* survival experience. They do not reflect an *individual's* chance of surviving since this is affected by specific characteristics of the individual and the cancer they have.

In this chapter, 1-year survival is shown, along with longer-term survival proportions such as 5- and 10-year survival, following a diagnosis of invasive ovarian cancer. One-year survival proportions might indicate the net short-term effectiveness of treatment and the stage at which the cancer was detected. In contrast, longer-term survival estimates might indicate:

- the effectiveness of treatment
- whether long-term side effects of cancer treatment are associated with additional mortality
- the number of people needing ongoing monitoring rather than cancer treatment
- milestones when there has been an arrest in the disease process or a slower progression.

In this chapter, change over time in relative survival estimates for those diagnosed with ovarian cancer are described, as are differences by age at diagnosis and by histological type

of ovarian cancer. In addition, selected findings on survival by stage at diagnosis are presented from the published literature. Relative survival proportions cannot be calculated according to Indigenous status and country of birth due to data limitations and the lack of necessary life tables. However, *crude* survival estimates can be calculated according to Indigenous status for women in four jurisdictions and the results from these calculations are shown in this chapter. In addition, international data on survival are provided.

The survival estimates shown in this chapter are based on the analysis of records of ovarian cancer cases diagnosed between 1982 and 2006 as held in the Australian Cancer Database. Data from the National Death Index on deaths (from any cause) that occurred up to 31 December 2008 were used to determine which women with ovarian cancer had died and when this occurred.

Survival of those diagnosed between 2000 and 2006

For women who were diagnosed with ovarian cancer between 2000 and 2006, 1-year relative survival was 74% (Table 4.1). The corresponding 5-year relative survival estimate was considerably lower at 40%. In other words, those women who were diagnosed with ovarian cancer between 2000 and 2006 were 40% as likely as comparable women in the general population to live 5 years after diagnosis.

	1-year relative s	urvival	5-year relative su	ırvival
	RS (%)	95% CI	RS (%)	95% CI
Ovarian cancer	73.7	72.7–74.7	40.0	38.8–41.2
Breast cancer	97.4	97.2–97.5	88.3	88.0-88.6

Table 4.1: Relative survival, ovarian and breast cancer, females, 2000-2006

Source: Australian Cancer Database, AIHW; AIHW & NBOCC 2009.

Survival from ovarian cancer compared with other cancers

To put the survival estimates for ovarian cancer into context, it is useful to compare these estimates with those of other cancers. Survival estimates for the period of 2000 to 2006 are currently available for one other type of cancer – breast cancer. As shown in Table 4.1, the prognosis for women diagnosed with breast cancer was better than that for those diagnosed with ovarian cancer for both the 1-year and the 5-year survival estimates. Specifically, the 1-year relative survival estimates were 74% for ovarian cancer and 97% for breast cancer (AIHW & NBOCC 2009). For the 5-year relative survival estimates, the estimate for breast cancer (88%) was more than double the estimate for ovarian cancer (40%).

In order to compare survival from ovarian cancer with a broader range of cancers, Figure 4.1 presents 1- and 5-year relative survival estimates (as sourced from a 2008 report by AIHW, CA and AACR) for the 10 most frequently diagnosed cancers among women (excluding non-melanoma skin cancer). The data pertain to women diagnosed with cancer between 1998 and 2004. For the 1-year relative survival estimates, the estimate for ovarian cancer (73%) is significantly lower than the estimate for all cancers combined (79%), and it ranked seventh lowest of the ten relative survival estimates shown. In regard to the 5-year relative survival estimates, the difference between the estimate for ovarian cancer (40%) and the estimate for all cancers combined is much more stark (64%), and ovarian cancer ranked eighth lowest of the ten cancers shown. These data indicate that the prognosis prospects for women

diagnosed with ovarian cancer are often poorer than those for women diagnosed with other frequently diagnosed cancers. The reasons for the poor survival outcomes for ovarian cancer include the relatively high proportion of diagnoses at an advanced stage (see Chapter 2) which may be affected by the non-specific nature of the symptoms of this type of cancer.



Differences by age at diagnosis

Figure 4.2 presents 1- and 5-year relative survival estimates by age at diagnosis for women diagnosed with ovarian cancer during 2000 to 2006. For 1-year relative survival, those in the youngest age group (those less than 30 years) had the highest survival estimate (96%). However, this estimate is not significantly higher than the estimates for women aged 30 to 39 years (91%) and those aged 40 to 49 years (92%). In contrast, for all of the age groups from 50 to 59 years onwards, there is a statistically significantly decrease in the relative survival estimates from one group to the next, with those aged 80 years and over having the lowest 1-year relative survival estimate (36%).

The 5-year relative survival estimates decreased steadily by age at diagnosis, with statistically significant differences observed between all of the age groups. The estimates ranged from 86% for those aged less than 30 years to 15% for those aged 80 years and over.

A large body of other research has also found that survival of women diagnosed with ovarian cancer at an older age is much poorer than it is for those diagnosed at a younger age; examples include studies using data from the USA (Horner et al. 2009), the United Kingdom (Cancer Research UK 2006) and Ireland (WHC & NCRI 2006). This difference by age in survival may be due to a number of different reasons, including differences in the histological type and stage at diagnosis of the tumours, a greater likelihood of co-morbidities and frailty among those diagnosed at an older age, and differences by age in treatments received and the inclusion in clinical trials (Chan et al. 2006; Maas et al. 2005; McMurdo et al. 2005; Petignat et al. 2004; South Australia Cancer Registry 2000; Tracey et al. 2009; Uyar et al. 2005; WHC & NCRI 2006).



Trends

Survival curves for ovarian cancer are presented in Figure 4.3 for four time periods from 1982–1987 to 2000–2006. For each of the four time periods, the relative survival estimates fell most sharply during the first five to six years following diagnosis, indicating that the relative risk of dying from ovarian cancer was highest during the initial years following diagnosis. In contrast, from about nine years following diagnosis onwards, the relative survival estimates were virtually stable. This suggests that the relative risk of dying from ovarian cancer was small for those women who survived for nine years or more following their ovarian cancer diagnosis.

A question of key interest is whether or not survival has improved over the years. The data indicate that improvement is evident when the entire time period from 1982–1987 to 2000–2006 is considered. For instance, between the first and the last of the four time periods considered, 1-year relative survival increased significantly from 63% to 74%, while 5-year relative survival increased significantly from 33% to 40% (Figure 4.3 and Appendix Table D4.3). However, much of the improvement in survival occurred during the earlier of the four time periods, rather than in the most recent ones. Specifically, no significant change in the 1-

year relative survival estimates was seen between 1994–1999 and 2000–2006 (72% and 74%, respectively); likewise, there was also no significant change in the 5-year estimates between the most recent two periods (39% and 40%, respectively).



Note that the method used to calculate the relative survival estimates shown in this chapter does not take into account differing age structures in the population over time. Since the average age of those diagnosed with ovarian cancer has increased somewhat over the years considered (see Table 2.2), the improvement over time in relative survival estimates may actually be somewhat greater than is indicated. However, determining whether this is the case is beyond the scope of this report. See Appendix B for further discussion on the age standardisation of relative survival estimates.

Possible reasons for the overall improvement in survival from ovarian cancer over time include the following (AIHW, CA & AACR 2008; ACN & NBCC 2004; Tracey et al. 2009):

- more accurate and effective investigation, diagnosis and staging of disease
- improvements in the speed and appropriateness of referral
- increasing subspecialisation in gynaecological oncology and the establishment of multidisciplinary teams
- advances in the effectiveness of treatment, including surgery and chemotherapy
- more widespread availability of treatment
- availability of evidence-based guidelines for the management of ovarian cancer.

Trends by age at diagnosis

Figure 4.4 illustrates 5-year relative survival curves by age at diagnosis for 1982–1987 to 2000–2006. These data indicate that the improvements in survival over time in Australia were centred on a subset of women – those in the middle age groups. While there was some improvement over the four time periods in the relative survival estimates for the two youngest age groups (i.e. from 81% to 86% for those under 30 years of age and 66% to 71%

for those 30 to 39 years at diagnosis), the difference was not statistically significant. Likewise, for the oldest age group, namely those aged 80 years and over, there was no statistically significant change between any of the four time periods, with estimates of 15% in both 1982–1987 and 2000–2006. In contrast, there was a significant increase in the 5-year relative survival estimates for women in the 'middle' age groups – women aged 40 to 79 years – over the periods considered. That is, the 5-year relative survival for those aged 40 to 49 years increased significantly from 45% in 1982–1987 to 61% in 2000–2006. The corresponding improvement for those aged 50 to 59 years was 36% to 50%, and for those aged 60 to 69 years, 27% to 41%. Meanwhile, although statistically significant, the improvement was not as marked for those aged 70 to 79 years, with the relative survival estimates increasing from 18% to 24% over the four time periods.



Survival by type of ovarian cancer

Five-year relative survival by histology types are shown for the period 2000 to 2006 in Figure 4.5 and Table 4.2. These data illustrate that ovarian cancer is a heterogeneous disease with widely varying clinical outcomes, depending on the type of ovarian cancer.

Women diagnosed with *germ cell tumours* (group 3) had the highest 5-year relative survival (91%), followed by those diagnosed with *sex cord-stromal tumours* (group 2) (84%). As shown in Chapter 2, younger women are more likely than older women to be diagnosed with these two types of ovarian cancers, especially *germ cell tumours*. (Also see Appendix Table D4.5 where 5-year relative survival is shown by age group for each of the histology groups for 1982 to 2006).

Women diagnosed with an *unspecified malignant neoplasm* (group 5) had the lowest 5-year relative survival (13%), with this estimate significantly lower than those calculated for each of the other major histology groupings. It has been suggested that women diagnosed with

this 'type' of ovarian cancer have poor survival prospects because they usually present with advanced disease or other factors that make them unsuitable for surgical treatment (Cancer Council Victoria 2007).



The 5-year relative survival for women diagnosed with *carcinoma* (group 1) was estimated to be 40%. However, this overall estimate masks the variation in relative survival estimates for the different subtypes of carcinomas, with the 5-year relative survival ranging from 77% for those diagnosed with *endometrioid carcinoma* (group 1.3) to 14% for those diagnosed with *unspecified carcinoma* (group 1.7). Five-year relative survival was also very low (15%) for those diagnosed with *adenocarcinoma not otherwise specified* (group 1.5).

A significant improvement in 5-year relative survival between 1982–1987 and 2000–2006 was found for some of the histology groups but not others (Table 4.2). Furthermore, the timing of the improvement varied. For women diagnosed with *sex cord-stromal tumours* (group 2), the 5-year relative survival estimates increased significantly from 62% in 1982–1987 to 84% in 1988–1993, with the estimates for the following two periods remaining in the low 80s. There was a significant increase over the four time periods in survival for those diagnosed with *germ cell tumours* (group 3) (80% to 91%, respectively), with the majority of change occurring between 1982–1987 and 1988–1993. Although less marked, there was also a significant increase in survival for those diagnosed with *carcinoma* (group 1), with these estimates increasing from 32% in the first period to 40% in the last period.

		1982–198	7		1988–199	3		1994–1999			2000–200	6
Type of ovarian cancer ^(a)	No. of cases	RS (%)	95% CI	No. of cases	RS (%)	95% CI	No. of cases	RS (%)	95% CI	No. of cases	RS (%)	95% CI
1: Carcinoma (epithelial tumours)	4,537	31.6	30.2–33.0	5,305	35.2	33.8–36.5	5,733	38.1	36.8–39.4	7,108	39.5	38.2-40.8
1.1: Serous carcinoma	1,533	30.4	28.0–32.8	2,195	32.7	30.7–34.8	2,728	35.6	33.7–37.4	3,555	36.7	34.9–38.6
1.2: Mucinous carcinoma	597	52.2	47.9–56.4	673	53.3	49.2–57.2	627	60.2	56.0-64.1	586	57.4	52.8-61.8
1.3: Endometrioid carcinoma	522	46.4	41.8–50.9	559	59.2	54.7-63.5	557	70.1	65.8-74.1	679	77.2	73.2-80.9
1.4: Clear cell carcinoma	217	44.8	37.9–51.6	307	51.4	45.4–57.2	329	62.2	56.3-67.5	437	67.5	62.3-72.2
1.5: Adenocarcinoma NOS	1,306	17.4	15.3–19.6	1,156	17.6	15.4–19.9	1,067	13.1	11.2–15.3	1,167	15.2	13.0–17.5
1.6: Other specified carcinoma	45	31.0	17.9-45.6	68	26.4	16.4–37.8	64	45.9	32.6–58.5	178	46.4	37.8–54.6
1.7: Unspecified carcinoma	317	24.2	19.6–29.2	347	20.4	16.4–24.8	361	16.7	13.1–20.7	506	13.9	10.9–17.3
2: Sex cord-stromal tumours	98	61.5	50.5-71.1	88	83.5	72.7–91.3	87	82.0	71.0-90.0	77	84.4	71.6–92.8
3: Germ cell tumours	155	79.6	72.2-85.3	188	89.1	83.6–93.0	180	87.4	81.4–91.6	270	90.8	86.1–94.2
4: Other specified malignant neoplasm	119	19.2	12.8-26.8	190	27.2	20.6–34.2	203	26.4	20.3–33.0	293	31.4	25.6–37.4
5: Unspecified malignant neoplasm	205	19.8	14.8–25.4	193	14.5	10.3–19.5	262	10.5	7.6–14.0	467	12.7	9.9–15.9
Total	5,114	33.0	31.6–34.3	5,964	36.8	35.5–38.1	6,465	38.8	37.5-40.0	8,215	40.0	38.8-41.2
(a) All cases were coded as primary site, inv	vasive ovariar	cancers. Ap	pendix Table D2.5	provides a lis	t of the histe	ology types include	d in each grou	Ċ				

Table 4.2: Incidence and 5-year relative survival by type of ovarian cancer, 1982-1987 to 2000-2006

Note: The number of cases equals the total number of diagnosed cases in the period considered.

Source: Australian Cancer Database, AIHW.

However, improvements over time within this group of carcinomas did not apply evenly. The most sizeable change was found for those diagnosed with *endometrioid carcinoma* (group 1.3), with 5-year relative survival estimates increasing from 46% in 1982–1987 to 77% in 2000–2006. The increase in the 5-year relative survival estimates for *clear cell carcinoma* (group 1.4) was also relatively large, with the estimates increasing from 45% to 68% between the first and the last of the four periods. In contrast, there was no significant improvement in survival for those diagnosed with some of the other types of carcinomas, such as *mucinous carcinoma* (group 1.2) and *adenocarcinoma not otherwise specified* (group 1.5).

Survival by stage at diagnosis

Existing research has consistently shown that stage at diagnosis of ovarian cancer is closely related to survival prospects, with advanced stage associated with poorer survival (e.g. Averette et al. 1995; Chan et al. 2006; Heintz et al. 2006; Laurvick et al. 2003; South Australia Cancer Registry 2000; Yang et al. 2008). Since no national data are available on stage at diagnosis in Australia, national relative survival estimates for ovarian cancer by stage at diagnosis cannot be calculated. However, to illustrate the trends, data from NSW and from the USA in which ovarian cancer was defined in the same way as in this report (i.e. ICD-10 code of C56) are shown. For information on the various staging systems used to classify ovarian cancers, see Appendix E.

As noted in Chapter 2, data on stage at diagnosis for ovarian cancer, based on the SEER summary stage system, are available for NSW (Tracey et al. 2009). According to those data, 5-year relative survival estimates for tumours that were diagnosed at the localised stage between 1980 and 2003 was 78% (Table 4.3). In contrast, the corresponding estimates for regional and distant tumours were significantly lower, at 34% and 18% respectively.

Stage at diagnosis ^(a)	Number of cases ^(b)	Per cent of staged cases	Relative survival (%) ^(c)	95% confidence interval
Localised	1,763	25.4	78.0	76.0–80.0
Regional	1,522	21.9	34.3	31.8–36.8
Distant	3,649	52.6	17.5	16.1–18.9
Unknown	n.a.		n.a.	n.a.
Total	6,934	100.0	36.7	35.7–37.7

Table 4.3: Five-year relative survival by stage at diagnosis, ovarian cancer, New South Wales, 1980–2003

(a) Based on the SEER summary stage system. 'Localised' tumours were confined to one or both ovaries, 'regional' tumours had spread to surrounding tissue or nearby lymph nodes and 'distant' tumours had spread to distant organs or other parts of the body and had begun to grow at the new location (see Appendix E).

(b) Includes ovarian cancers coded in ICD-10 as C56; borderline ovarian tumours are not included.

(c) The cause-specific method of calculating survival was used.

Source: Tracey et al. 2009.

Data for 1999 to 2005 from the USA are shown in Table 4.4. As was seen with the NSW data, there is a clear gradient in the survival estimates in the USA data according to stage at diagnosis with a 5-year relative survival estimate of 94% for those diagnosed with localised tumours and an estimate of 28% for those diagnosed with distant tumours (Table 4.4). In addition, the USA data provide an estimate of survival for those with an unknown tumour stage at diagnosis; according to these data, the 5-year relative survival for this group was 27%, which is similar to the estimate for those who were diagnosed with a distant tumour.

Stage at	<65 yea	rs	65+ yea	rs	All ages	6
diagnosis ^(a)	% of cases ^(b)	RS (%) ^(c)	% of cases ^(b)	RS (%) ^(c)	% of cases ^(b)	RS (%) ^(c)
Localised	21	93.9	7	93.5	15	93.8
Regional	20	79.7	13	56.7	17	72.8
Distant	55	34.9	69	20.0	62	28.2
Unknown	4	48.9	10	13.0	7	27.3
Total	100	56.5	100	29.8	100	45.9

Table 4.4: Five-year relative survival by stage at diagnosis and age group, ovarian cancer, United States of America, 1999–2005

(a) Based on the SEER summary stage system. 'Localised' tumours were confined to one or both ovaries, 'regional' tumours had spread to surrounding tissue or nearby lymph nodes and 'distant' tumours had spread to distant organs or other parts of the body and had begun to grow at the new location (see Appendix E).

(b) The numbers of cases were as follows: 16,000 cases of women less than 65 years old; 13,168 cases of women 65 years and over; and a total of 29,168 cases.

(c) The cohort method of calculating relative survival was used.

Note: Data are from the SEER 17 areas which cover approximately a quarter of the USA (see Table 21.7 in Horner et al. 2009).

Source: Horner et al. 2009.

The USA data also provide information on differences by age group in survival by stage at diagnosis. Overall, the survival prospects of women aged less than 65 years were much better than for older women (those aged 65 years and over), with 5-year relative survival estimates of 57% and 30%, respectively. This differential by age applied for all of the stages at diagnosis, with one exception. This exception pertains to women diagnosed with localised tumour – for women diagnosed with these 'early-stage' tumours, regardless of the age group, 5-year relative survival was 94%. In contrast, for example, for distant tumours, 5-year relative survival was 35% for those aged less than 65 years at diagnosis, while it was 20% for those aged 65 years and over at diagnosis. The differential by age was particularly marked for those diagnosed with an unknown stage at diagnosis – the 5-year relative survival estimate for those aged less than 65 years was 49% compared with 13% for those aged 65 years and over.

Differences by Aboriginal and Torres Strait Islander status

Relative survival estimates cannot be calculated for Indigenous women because of data issues and the lack of necessary life tables. However, 5-year *crude* survival estimates can be derived based on data from Queensland, Western Australia, South Australia and the Northern Territory. As discussed earlier in this chapter, crude survival estimates do not take into account the cause of death, nor do they compare observed survival with expected survival. Past research has shown that the life expectancy of Indigenous women is shorter than that of non-Indigenous women (ABS 2004, 2009e). At the same time, the mean age at which women were diagnosed with ovarian cancer differs by Indigenous status, with the Indigenous women being younger at diagnosis (mean of 53 years) than the non-Indigenous women (63 years) (Table 4.5). It is not known how these underlying differences may have affected the crude survival estimates presented.

Given the small number of ovarian cancer cases reported among Indigenous women, a 10-year time period from 1997 to 2006 is considered in these analyses. Despite this, the

number of cases of ovarian cancer among Indigenous women was still relatively small (68 cases) and this should be considered when making use of these data.

The crude 5-year survival estimate for ovarian cancer for Indigenous women was somewhat lower than for non-Indigenous women -35% and 38%, respectively – in the four jurisdictions; however, this difference was not statistically significant.

Table 4.5: Five-year crude survival by Indigenous status, ovarian cancer, Queensland, Western Australia, South Australia & Northern Territory, 1997–2006

	Indigenous	Non-Indigenous
Number of cases ^(a)	68	3,891
Crude survival (%)	35.1	38.1
95% confidence interval	23.1–47.4	36.5–39.8
Mean age at diagnosis	53.2	63.2

(a) Equals the total number of diagnosed cases in the period considered.

Source: Australian Cancer Database, AIHW.

International comparisons

In addition to the methodological challenges associated with comparing cancer statistics from different countries (as discussed in Chapter 1), additional uncertainties arise when comparing relative survival estimates. In particular, there tends to be wide variation across countries in: the years to which the relative survival estimates apply; the length of the follow-up period considered (e.g. 1-, 5-, 10-year and so forth); and the methods and age groups used to calculate the relative survival estimates. For these reasons, relative survival estimates for different countries are not compared in this report.

Although more rudimentary than relative survival estimates, the mortality-to-incidence ratio (MIR) is used in this report to make international comparisons. This ratio describes how many deaths there were in a particular year due to a particular disease, relative to the number of new cases diagnosed that year (using age-standardised data). For example, an MIR of 0.60 for ovarian cancer would indicate that there were 60 deaths for every 100 new cases of ovarian cancer diagnosed in that year (though the deaths need not relate to the same people as the cases). If survival tends to be lower in a particular country relative to others, then the MIR for that country generally would be expected to be higher (i.e. closer to 1.00). In contrast, if survival is higher, the ratio generally would be closer to zero. Appendix B provides further information about interpreting MIRs.

For this report, mortality-to-incidence ovarian cancer ratios were calculated using data from GLOBOCAN (Ferlay et al. 2004). The fact that the GLOBOCAN data were estimates for 2002 should be taken into account when interpreting the results shown in Figure 4.6. Note also that these data pertain to 'ovarian and related cancers' (i.e. the ICD-10 codes of C56 and C57.0–C57.4) rather than just 'ovarian cancer' (ICD-10 code of C56).

The GLOBOCAN data suggest that the MIRs for ovarian and related cancers varied markedly between different countries and regions, with survival from ovarian and related cancers poorest among women in South-Central Asia (MIR of 0.72), and best for women in Southern Europe (MIR of 0.46). The MIR for women in Australia was relatively low (0.55). This suggests that, in 2002, Australian women who were diagnosed with ovarian and related cancers had better survival prospects than their counterparts in many other countries and regions.

South Control Asia								
Eastern Africa								
Middle Africa								
Western Africa								
Northern Africa								
Western Asia								
Southern Africa								
World								
Caribbean								
Northern Europe								
Melanesia								
Central and Eastern Europe								
Micronesia								
Polynesia								
Northern America								
South-Eastern Asia								
Western Europe						1		
Australia								
_ New Zealand								
Central America								
Eastern Asia								
_ South America								
_ Southern Europe								
· -		0.2	03	0.4	0.5	0.6	0.7	0.8
	, 0.1	0.2	Mortality	-to-incide	ence ratio	0.0	0.7	0.0
			mortanty					
1. The ratios are based on est	imated incidenc	e and mortal	ity data for 2	002; those	estimates v	vere based	on data froi	m
approximately 3 to 5 years e	earlier.		.,	,				
2. The mortality-to-incidence ra	atio equals the a	age-standard	ised mortali	y rate divid	ed by the a	ge-standard	dised incide	nce rate.
		1) oo 1 'bk op	1 (5/ 1)_(5	/ / (coo /\n	nondiv R to	r further det	alle)	

Figure 4.6: International comparison of mortality-to-incidence ratios for ovarian and related cancers, 2002

5 Prevalence of ovarian cancer

Prevalence, or complete prevalence as it is sometimes called, is the number of people alive at a specified point in time who have ever been diagnosed with ovarian cancer regardless of how long ago. These people may or may not be undergoing treatment or be considered 'cured'. In contrast, 'limited-duration prevalence' provides information on the number of people alive who were diagnosed with ovarian cancer within a specified time period, such as the previous 1 or 5 years. One-year prevalence data, for example, would indicate the number of people alive on 31 December of a particular year who were diagnosed with ovarian cancer during that same year, while 5-year prevalence data would indicate the number of people alive on 31 December of a specified year who were diagnosed with ovarian cancer within the previous five years.

The prevalence of a disease in a given population is influenced by the incidence of the disease, survival from the disease and the age at which people are diagnosed (i.e. older people are more likely to die sooner due to age-related morbidity and frailty).

Along with information on incidence, mortality and survival, prevalence is another indicator of the impact of ovarian cancer in our society both at the personal/familial level and societal level, particularly in terms of health care services. While the exact nature of health care needs can vary widely from one person to the next over the years following diagnosis, overall, different types and intensities of health care services may be required by those who were diagnosed with ovarian cancer recently (e.g. in the past year) compared with those diagnosed many years previously.

In Australia, as elsewhere, complete prevalence data are not available through cancer registry data collections since these collections do not hold data for a long-enough period. The only source of complete prevalence data in Australia is surveys, such as the National Health Survey, where prevalence estimates are based on self-reported information of a sample of Australians (ABS 2009c). However, since the National Health Survey excludes people in hospitals, hospices, and nursing and convalescent homes, those data are incomplete. An additional deficiency of those data may be the erroneous self reporting of benign or borderline ovarian tumours as invasive ovarian cancer.

In this report, limited-duration prevalence is presented using data from the Australian Cancer Database, with information on deaths (from any cause) sourced from the National Death Index. Since national incidence data on ovarian cancer are available from 1982 onwards, limited-duration prevalence data can be presented for a maximum of 25 years (from 1 January 1982 to 31 December 2006). In addition, information is provided in this chapter on differences in prevalence by age, geographical area and country of birth.

In this chapter, no international comparisons are made. Making such comparisons is difficult since prevalence data from other countries often differ from Australian data in the years to which they apply, the number of years considered (e.g. 5, 10, 25 years) and the analytical methods employed to calculate prevalence.

Unlike the incidence data, which pertain to the number of *cases* of ovarian cancer, the prevalence data presented in this report pertain to the number of *females* who have been diagnosed with ovarian cancer and are still alive. However, as mentioned in Chapter 2, since it is rare that any one woman would be diagnosed with more than one primary ovarian cancer during a 1-year period, the number of new *cases* of ovarian cancer in a particular year would be very similar to the number of *females* diagnosed with ovarian cancer in that year.

Prevalence in 2006

Of all females alive at the end of 2006, over 8,200 had been diagnosed with ovarian cancer in the previous 25 years (Table 5.1). This equates to 8 out of 10,000 females. At the same time, the 20-year prevalence for ovarian cancer was 7,500 women, the 10-year prevalence was 5,179 women and the 1-year prevalence was 1,016 women. The 1-year prevalence compares with an *incidence* of 1,226 cases for 2006 (Table 2.1). Note that those women who were both diagnosed with ovarian cancer and died in 2006 (approximately 210 women) may or may not have died as a result of ovarian cancer.

Time period	Number ^(a)	Per 10,000 population ^(b)
1-year prevalence	1,016	1.0
5-year prevalence	3,445	3.3
10-year prevalence	5,179	4.9
15-year prevalence	6,506	6.2
20-year prevalence	7,500	7.1
25-year prevalence	8,216	7.8

(a) Refers to the number of females, not cases.

(b) Based on the number of females in the Australian population at 31 December 2006.

Source: Australian Cancer Database, AIHW.

In order to compare prevalence across commonly diagnosed cancers in females, Table 5.2 presents data from the AIHW's 2008 publication on cancer survival and prevalence (AIHW, CA & AACR 2008). Those data pertain to the prevalence at the end of 2004 and thus 23 years of cancer incidence data were available.

Table 5.2: Limited-d	luration prevalence of th	e 10 most commonly	^r diagnosed cancers	s ^(a) , females, e	end
of 2004					

Cancer type (ICD-10 codes)	1-year prevalence	5-year prevalence	10-year prevalence	23-year prevalence
Breast (C50)	11,764	53,051	89,777	129,438
Bowel (C18–C20)	4,969	18,940	29,929	43,286
Melanoma of skin (C43)	4,151	18,697	33,303	56,235
Lung (C33–C34)	1,978	4,413	5,657	6,817
Uterus, body (C54)	1,630	6,665	11,244	17,720
Non-Hodgkin lymphoma (C82–C85, C96)	1,423	5,632	8,837	11,845
Unknown primary site (C26, C39, C76, C80)	632	1,511	1,943	2,690
Ovary (C56)	1,024	3,288	4,997	7,637
Thyroid (C73)	1,092	4,502	7,529	11,248
Leukaemia (C91–C95)	777	3,007	4,663	6,513
All cancers ^(b)	36,331	141,553	230,245	338,692

(a) Determined by the most commonly diagnosed cancers in 2004 and ordered accordingly; excludes non-melanoma skin cancer (C44).

(b) Includes cancers coded in ICD-10 as C00–C97 (except for C44), D45, D46, D47.1 and D47.3.

Note: Data refer to the number of females, not cases.

Source: AIHW, CA & AACR 2008.

When 23-year prevalence is considered, ovarian cancer was the seventh most prevalent type of cancer in women (excluding non-melanoma skin cancer) among the ten most commonly reported cancers. It was also the second most prevalent type of gynaecological cancer (after 'uterus, body' cancer). Of all females alive at the end of 2004, over 7,600 females had been diagnosed with ovarian cancer in the previous 23 years. In comparison, the 23-year prevalence for breast cancer was 129,438 women and for 'uterus, body' cancer, it was 17,720 women. For each of the other three prevalence durations considered, ovarian cancer was the eighth most prevalent type of cancer among women (excluding non-melanoma skin cancer).

Differences by age

Table 5.3 presents 25-year prevalence of ovarian cancer by age group. At the end of 2006, there were 2,051 women in the 60 to 69 year age group still alive who had been diagnosed with ovarian cancer in the previous 25 years, while there were a further 1,841 women aged 50 to 59 years who had been diagnosed with this type of cancer.

When the number of females diagnosed with ovarian cancer is compared with the number in the respective age group, the data indicate that the highest proportion exists among those aged 70 to 79 years, with a prevalence of 26 per 10,000 females.

Age group (years)	Number ^(a)	Per 10,000 population ^(b)
0–19	56	0.2
20–29	172	1.2
30–39	474	3.1
40–49	937	6.1
50–59	1,841	13.9
60–69	2,051	22.4
70–79	1,634	25.9
80+	1,051	22.6
Total	8,216	7.8

Table 5.3: Twenty-five-year prevalence of ovarian cancer by age group, end of 2006

(a) Refers to the number of females, not cases.

(b) Based on the number of females in the Australian population at 31 December 2006.

Source: Australian Cancer Database, AIHW.

Differences across groups

As noted earlier in this chapter, the prevalence of ovarian cancer is influenced by the incidence of the disease, survival rates and the average age at diagnosis. Since these factors can differ across sub-groups, prevalence may also differ. In this section of the report, prevalence data by state and territory, and by country of birth are presented.

Differences by geographical area

Table 5.4 presents prevalence data for the end of 2006 according to the state and territory in which the woman lived at the time of diagnosis. Since it is unknown whether the women lived in the same state and territory in 2006 as they did at the time of diagnosis, these data should be used with caution. During 1982 to 2006, just over 2,700 women had been diagnosed with ovarian cancer in New South Wales and were still alive at the end of 2006.

This compares with 25-year prevalence for ovarian cancer of 2,127 women for Victoria and 1,638 women for Queensland.

State or territory	1-year prevalence	5-year prevalence	10-year prevalence	25-year prevalence
New South Wales	315	1,154	1,731	2,707
Victoria	276	881	1,330	2,127
Queensland	196	636	1,009	1,638
Western Australia	114	382	507	759
South Australia	69	238	366	603
Tasmania	26	69	119	203
Australian Capital Territory	16	59	85	132
Northern Territory	4	26	32	47
Total	1,016	3,445	5,179	8,216

Table 5.4: Limited-duration prevalence of ovarian cancer by state and territory of diagnosis, end of 2006

Note: Data refer to the number of females, not cases.

Source: Australian Cancer Database, AIHW.

Differences by country of birth

The prevalence of ovarian cancer according to country or region of birth is shown in Table 5.5. The 25-year prevalence, as a proportion of the respective female population, was highest among women born in the European regions (13 per 10,000 females for both North-West Europe, and Southern and Eastern Europe). This compares with a figure of 6 per 10,000 for those born in Australia. The data also indicate that there was a relatively low proportion of females alive who had been diagnosed with ovarian cancer in the 25-year period among women born in North-East Asia and Sub-Sahara Africa (both 5 per 10,000 females).

	1-ye	ar	5-ye	ar	10-	year	25-1	year
Country/region of birth ^(a)	Number ^(b)	Per 10,000 population ^(c)						
North-West Europe	66	1.3	418	5.6	624	8.4	866	13.4
Southern and Eastern Europe	69	1.6	225	5.3	345	8.1	566	13.4
Americas	18	1.7	50	4.7	68	6.3	102	9.5
North Africa and the Middle East	14	1.0	52	3.7	67	4.8	98	7.1
Oceania and Antarctica, excl. Australia	24	0.9	89	3.2	128	4.5	194	6.9
South-East Asia	38	1.1	114	3.2	170	4.7	240	6.7
Australia	603	0.8	2,102	2.7	3,155	4.0	5,029	6.4
Southern and Central Asia	18	1.3	43	3.0	65	4.5	91	6.3
Sub-Saharan Africa	9	0.5	32	2.9	47	4.3	57	5.2
North-East Asia	21	0.8	61	2.4	96	3.7	128	5.0
Not stated	106	:	259	:	414	:	713	:
Total ^(b)	1,016	1.0	3,445	3.3	5,179	4.9	8,216	7.8
(a) Classified according to the Standard Australian Classifi	fication of Countrie	s, second edition (see	Appendix A). Countr	ies/regions of birth are	e ordered in descen	ding order according t	o the 25-year preva	alence

Table 5.5: Limited-duration prevalence of ovarian cancer by country/region of birth, end of 2006

5 ק proportions.

Refers to the number of females, not cases.

Based on the number of females in the Australian population born in each country/region as at 30 June 2006, except for the 'Total' which is based on the number of females in the Australian population at 31 December 2006. (c) (p)

Source: Australian Cancer Database, AIHW.

6 Burden of disease due to ovarian cancer

The effect of ovarian cancer on the health of the population can be summarised by using a number of different measures that combine information on both fatal and non-fatal health outcomes into a single number. Such measures can be used for a range of purposes including:

- comparing the burden associated with different diseases
- comparing the effect of a particular disease on different population groups or over time
- setting priorities for health planning, public health programs, as well as research and development (Murray et al. 1999).

Of the available summary measures, one of the most commonly used is the 'disabilityadjusted life year' (DALY), also commonly referred to as 'burden of disease'. The DALY combines information on the extent of:

- premature death which is measured by the years of life lost (YLL) due to disease or injury and
- non-fatal health outcomes which is measured by years of 'healthy' life lost (YLD) due to disease, disability or injury.

In order to combine these two health measures into a summary measure, the DALY uses time as a common 'currency'. Hence, the DALY is a measure of the years of life lost due to premature death (YLL) and years of healthy life lost due to disease, disability or injury (YLD), or a combination of the two. The more DALYs associated with a particular disease, the greater the burden. Further information about DALYs can be found in the AIHW report on the burden of disease and injury (Begg et al. 2007a).

In the report by Begg and associates, ovarian cancer was defined to include the ICD-10 codes of 'C56 and C57.0–C57.4', which we refer to as 'ovarian and related cancers' in this report. In this chapter, the burden of disease in Australia due to ovarian and related cancers is presented along with comparisons with other diseases that are also major contributors to the overall burden. The most recent burden of disease estimates for Australia are for 2003. These estimates, and the method by which they were derived, are detailed in an AIHW report by Begg and associates (2007a,b).

Burden of disease in 2003

The total burden of disease for females in Australia in 2003 was estimated to be more than 1.2 million DALYs and the burden due to cancer was 235,034 DALYs, which is 19% of the total burden. Table 6.1 presents the leading causes of disease burden for females, along with the five leading causes of cancer burden. Ovarian and related cancers ranked 25th in terms of the leading causes of burden of disease for women, and accounted for 1% of all female burden of disease. In terms of leading causes of burden due to cancer, ovarian and related cancers ranked fourth, with nearly 12,000 DALYs attributed to this disease. Ovarian and related cancers accounted for 5% of the female burden of disease due to all cancers and was the leading cause of burden from all gynaecological cancers.

Causa	ICD-10 codes	Disability- adjusted life	Per cent of total	Pank
Cause		years (DALTS)	DALIS	Nalik
Anxiety and depression	F30, 32–39, 40.0, 40.1, 41.0– 41.2, 42, 43.1, 93.0	126,464	10.0	1
Ischaemic heart disease	120–25	112,390	8.9	2
Stroke	G45; I60–69	65,166	5.1	3
Type 2 diabetes	E11–13	61,763	4.9	4
Dementia	F00–01, 02.0–02.1, 02.3, 03; G30, 31.0–31.1, 31.8–31.9	60,747	4.8	5
Cancer	C00–96	235,034	18.5	
Breast cancer	C50	60,520	4.8	6
Lung cancer	C33–34	33,876	2.7	8
Bowel cancer	C18–21	28,962	2.3	10
Ovarian and related cancers	C56, 57.0–57.4	11,994	0.9	25
Pancreas cancer	C25	11,246	0.9	27
Other cancers	All other 'C' codes	88,436	7.0	
Chronic obstructive pulmonary disease	I27.0, 27.8–27.9; J40–44	37,550	3.0	7
Asthma	J45–46	33,828	2.7	9
Total for all causes		1,268,156	100.0	

Table 6.1: Leading causes of burden of disease, including leading cancers, females, 2003

Source: Begg et al. 2007a.

Table 6.2 and Figure 6.1 show the extent of the cancer burden associated with the leading causes of cancer burden for females which were due to both premature death (YLL) and disease, disability or injury (YLD). For cancer, causes of years of healthy life lost to disability include side effects during and after treatment (e.g. during and after radiotherapy or chemotherapy) and the psychosocial affects of having gone through treatment.

Due to the relatively poor prognosis from many cancers compared with the majority of other diseases, most cancers contribute more years of life lost (YLL) than years of healthy life lost to disability (YLD). This is the case for ovarian and related cancers. In 2003, this disease resulted in 10,946 years of life lost due to premature mortality, which equates to 91% of the total estimated DALYs for this disease. This compares with an average of 82% for all cancers combined.

While ovarian and related cancers ranked 25th in terms of the causes of burden of disease for females when DALYs were considered, it ranked 11th in terms of the leading causes of mortality burden, and 75th in terms of causes of disability burden. Considering just the burden due to cancers, ovarian and related cancers accounted for 6% of all years of life lost from cancer and 2% of years lost due to disability.

33
, 20(
ales
fem
nts,
one
duud
) C
(YLI
Ital
n-fa
d no
) an
YLL
tal (
y fa
rs, b
nce
g ca
adin
g le:
ıdin
nclt
se, i
isea
of d
den
bur
s of
ause
ıg ci
adir
2: Le
e 6.2
[abl
- -

	Fata	l component		Non-fatal co	omponent		Total	% of	% of
Cause	Years of life lost (YLL)	Per cent of total YLL	Rank	Years of healthy life lost (YLD)	Per cent of total YLD	Rank	Disability-adjusted life years (DALYs)	DALYS due to YLL	DALYs due to YLD
Anxiety and depression	221	0.0	98	126,244	18.1	-	126,464	0.2	99.8
Ischaemic heart disease	89,152	15.7	-	23,238	3.3	5	112,390	79.3	20.7
Stroke	48,548	8.5	7	16,619	2.4	6	65,166	74.5	25.5
Type 2 diabetes	11,751	2.1	6	50,012	7.2	7	61,763	19.0	81.0
Dementia	16,009	2.8	7	44,738	6.4	с	60,747	26.4	73.6
Cancer	191,794	33.7	:	43,240	6.2	:	235,034	81.6	18.4
Breast cancer	40,080	7.0	ю	20,440	2.9	7	60,520	66.2	33.8
Lung cancer	31,551	5.5	4	2,325	0.3	51	33,876	93.1	6.9
Bowel cancer	23,735	4.2	5	5,227	0.7	31	28,962	82.0	18.0
Ovarian and related cancers	10,946	1.9	5	1,048	0.1	75	11,994	91.3	8.7
Pancreas cancer	10,984	1.9	10	262	0.0	120	11,246	97.7	2.3
Other cancers	74,498	13.1	:	13,938	2.0	:	88,436	84.2	15.8
Chronic obstructive pulmonary disease	21,025	3.7	Q	16,525	2.4	10	37,550	56.0	44.0
Asthma	2,423	0.4	44	31,405	4.5	4	33,828	7.2	92.8
Total for all causes	569,181	100.0	:	698,975	100.0	:	1,268,156	44.9	55.1
Source: Begg et al. 2007a.									



Differences by age

The leading causes of cancer burden for women by age are shown in Figure 6.2. In 2003, the distribution of the burden of ovarian and related cancers was flatter than that of a number of the other types of cancer shown, such as breast cancer and lung cancer.



The proportion of all DALYs from cancer that were due to ovarian and related cancers varied according to the age group considered, with the highest proportion found for those aged 15 to 19 years and those aged 20 to 24 years (9% and 8% of DALYs from cancer were due to ovarian and related cancers, respectively) (Appendix Table D6.1).

Trends and projections

Table 6.3 presents information on the burden of disease from ovarian and related cancers for 1993 and 2003, as well as the projected burden for 2013 and 2023. In 1993, the agestandardised burden of ovarian and related cancers was 123 DALYs per 100,000 females; it had decreased to 120 DALYs per 100,000 females by 2003. Projected trends to the year 2023 suggest that the age-standardised burden of ovarian and related cancers will decrease somewhat over time to 116 DALYs per 100,000 females in 2023. This indicates an overall estimated decrease of 3% from 2003 to 2023. In contrast, the total number of DALYs due to ovarian and related cancers is expected to continue to increase over the years from 11,994 in 2003 to 14,225 in 2023. This projected increase in the number of DALYs is due to a population that is both ageing and growing in size.

Year	Disability adjusted life years (DALYs)	Age-standardised rate
1993	10,918	123.1
2003	11,994	119.8
2013 ^(a)	13,331	119.2
2023 ^(a)	14,225	115.6

Table 6.5: Trends and projected burden of ovarian and related cancers, 1995 to 202	Table 6.3: Trends and	projected burden	of ovarian and related	cancers, 1993 to 2023
--	-----------------------	------------------	------------------------	-----------------------

(a) See Begg et al. 2007a for information on how the projections were derived.

Source: AIHW unpublished data.

7 Hospitalisations for ovarian cancer

Women with ovarian cancer may require hospitalisation as an admitted patient for a variety of reasons including diagnostic procedures and treatments (e.g. surgery, chemotherapy and the management of associated conditions). The number of such hospitalisations for ovarian cancer in any one year is related to a range of factors, including the number of women with ovarian cancer and the number of these requiring health services as an admitted patient in a hospital. Other factors include the availability of alternative health-care services, relative accessibility of hospital care, admission criteria and administrative policies.

In this chapter, details are provided on the number and characteristics of admitted patient hospitalisations that are related to the care and/or treatment of persons with invasive ovarian cancer, with the term 'hospitalisations' used interchangeably with 'separations'.

Due to the method in which the principal diagnosis for hospitalisations of cancer patients is coded (particularly in relation to same-day chemotherapy treatments), identifying those hospitalisations that are due specifically to ovarian cancer is not straightforward. As discussed in more detail in Appendix F, 'ovarian cancer-related hospitalisations' are defined in this report as those admitted patient hospitalisations in which:

- the principal diagnosis was ovarian cancer (i.e. ICD-10-AM code of C56) or
- ovarian cancer (i.e. ICD-10-AM code of C56) was recorded as an *additional* diagnosis and the principal diagnosis code related specifically to the treatment or care of a cancer patient.

The data source for this chapter was the National Hospital Morbidity Database (NHMD) which contains data on admitted patient separations. The most recent data available pertain to the 2007–08 financial year. Note that the data from the NHMD refer to hospitalisations and not individuals. Any one person may have multiple hospitalisations during the course of a year but data on the number of people hospitalised for a particular disease are not available. Further information about this data source can be found in Appendix C.

Over the course of the past decade, a number of hospitals (mainly in the public sector) in New South Wales, South Australia and the Australian Capital Territory changed their admissions practices so that not all patients who receive same-day chemotherapy services are admitted to hospital. Instead, these hospitals provide chemotherapy treatment on an outpatient (i.e. non-admitted patient) basis. This change in process, which is discussed in more detail in Appendix F, must be taken into account when examining change over time in the number of hospitalisations due to ovarian cancer. Because the change applies largely to same-day hospitalisations (and not to overnight ones), separate information is provided in this chapter on the number and rate of same-day and overnight hospitalisations. Ideally, data on the number of chemotherapy services provided to ovarian cancer patients on an outpatient basis would be included in this chapter, but such data are not available. In this chapter, rates of hospitalisations are presented per 1,000 females.

Hospitalisations in 2007–08

In the 2007–08 financial year, there were 14,277 hospitalisations due to ovarian cancer (Table 7.1); these accounted for 0.3% of all hospitalisations of women. The age-standardised rate of ovarian cancer–related hospitalisations was 1.2 (per 1,000 females).

	Number	Per cent of all hospitalisations	Age-standardised rate ^(a)	95% confidence interval
Ovarian cancer	14,277	0.3	1.2	1.20–1.24
All hospitalisations	4,149,381	100.0	370.0	369.60-370.31

Table 7.1: Hospitalisations for ovarian cancer and all reasons, females, 2007-08

(a) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 1,000 females.

Source: National Hospital Morbidity Database, AIHW.

Of the total number of hospitalisations for ovarian cancer, nearly eight out of ten (79%) were same-day hospitalisations (11,296), while the remainder (21%) were overnight hospitalisations (2,981).

Differences by age

Of all hospitalisations for ovarian cancer-related care in 2007–08, over half (61%) were of women aged 60 years and over, 34% were for those aged 40 to 59 years, while 5% were for those under the age of 40 years (Appendix Table D7.1).

Differences in the hospitalisation rate for ovarian cancer-related care according to age are shown in Figure 7.1. The rate of hospitalisation was less than 1 (per 1,000 females) for those under the age of 45 years, but rose to a high of 5 (per 1,000 females) for those aged 65 to 69 years and for those aged 70 to 74 years.



Average length of stay

Data on the total number of days that patients stayed in hospital are collected in the NHMD, with a length of stay of 1 day allocated to all same-day hospitalisations. By using those data, as well as information on the *number* of hospitalisations, the average length of stay (ALOS)

can be derived. In 2007–08, the average length of stay for ovarian cancer-related hospitalisations was 2.4 days (Table 7.2). When only those hospitalisations that included an overnight stay are considered, the average length of stay was 7.6 days.

Considering only those hospitalisations that involved an overnight stay, the average length of stay increased by age. For those aged less than 30 years, the average length of an overnight stay was 5.3 days, compared with 11.7 days for those aged 80 years and over.

Table 7.2: Average length of stay (ALOS) for ovarian cancer-related hospitalisations by age group,2007-08

Age group (years)	ALOS of overnight hospitalisations (days)	Total ALOS (days) ^(a)
<30	5.3	2.3
30–39	6.8	2.3
40–49	6.6	2.3
50–59	6.3	2.0
60–69	7.3	2.1
70–79	8.3	2.6
80+	11.7	4.1
Total	7.6	2.4

(a) Includes both overnight and same-day hospitalisations (with the latter, by definition, equal to1 day per hospitalisation).
Source: National Hospital Morbidity Database, AIHW.

Trends

The total number of hospitalisations for ovarian cancer increased by 35% between 1999–00 (10,604 hospitalisations) and 2007–08 (14,277 hospitalisations) (Table 7.3). While there was an overall decrease of 6% in the number of overnight hospitalisations over the period considered, the majority of change pertained to the number of same-day hospitalisations which increased by 52%. This is despite the fact that, as noted earlier, changes occurred in hospital admission procedures during the period considered such that by 2007–08, some

Year	Number of same-day hospitalisations	Number of overnight hospitalisations	Total number of hospitalisations
1999–00	7,434	3,170	10,604
2000–01	7,913	3,063	10,976
2001–02	9,095	2,978	12,073
2002–03	8,794	2,646	11,440
2003–04	9,407	2,710	12,117
2004–05	9,955	2,631	12,586
2005–06	10,317	2,942	13,259
2006–07	10,940	3,132	14,072
2007–08	11,296	2,981	14,277

Table 7.3: Hospitalisations for ovarian cancer by same-day and overnight status, 1999-00 to 2007-08

Source: National Hospital Morbidity Database, AIHW.
Cancer patients in three jurisdictions who received same-day chemotherapy were not classified as admitted patients and thus not included in the data (whereas in earlier years, they would have been included).

In Figure 7.2, trends in the age-standardised rate of ovarian cancer-related hospitalisations are shown. For all ovarian cancer-related hospitalisations, the rate increased slightly, but significantly, from 1.1 hospitalisations per 1,000 females in 1999–00 to 1.2 hospitalisations per 1,000 females in 2007–08. As shown, the overall increase over time was driven by changes in the number of same-day hospitalisations, while the rate of overnight hospitalisations decreased slightly but significantly between 1999–00 and 2007–08 (from 0.32 to 0.25 hospitalisations per 1,000 females, respectively).



Trends in the rate of hospitalisations for ovarian cancer by age group are shown in Figure 7.3. For the age group of those less than 50 years, the rates remained unchanged at 0.3 per 1,000 females over the period considered. For the other two age groups, the rates fluctuated somewhat over the years, but the general direction was one of an increase. Specifically, between 1999–00 and 2007–08, there was a 13% increase in the number of hospitalisations for ovarian cancer for those aged 50 to 69 years, and a corresponding increase of 22% increase for those aged 70 years and over.

Trends in average length of stay

Trends in the average length of stay of women who were hospitalised for ovarian cancer in 2007–08 are shown in Figure 7.4. For those hospitalisations that involved an overnight stay, the average length of stay for ovarian cancer–related hospitalisations was 7.3 days in 1999–00; it increased to 8.8 days in 2004–05 and then decreased to 7.6 days in 2007–08.







In contrast, the average length of stay for all ovarian cancer-related hospitalisations in 2007–08 was shorter than that in 1999–00 (2.4 and 2.9 days, respectively). The reduction of the overall average length of stay can be attributed to the increase of the number of same-day hospitalisations which, by definition, are shorter than those hospitalisations that involved an overnight stay.

Procedures undertaken during hospitalisations

Procedures undertaken in hospitals include surgical procedures, non-surgical procedures for investigative and therapeutic purposes (such as chemotherapy) and client support interventions (e.g. anaesthesia). One or more procedures can be reported for each hospitalisation, but procedures are not undertaken during all hospitalisations; thus, only some hospitalisations include data on procedures. The classification system that was used to code the 2007–08 data on procedures was the fifth edition of the Australian Classification of Health Interventions (ACHI) (see Appendix A).

Table 7.4 indicates the number of hospitalisations in which the indicated procedure was undertaken at least once during 2007–08. The majority of these hospitalisations included the 'administration of pharmacotherapy' (i.e. chemotherapy); this procedure was undertaken in almost two-thirds (62%) of ovarian cancer–related hospitalisations. In addition, 12% of the hospitalisations involved 'generalised allied health professions' and a further 11% included the 'loading of a drug delivery device'.

Procedure description (ACHI ^(a) block number)	Count of hospitalisations ^(b,c)	Per cent ^(c)
Administration of pharmacotherapy (1920)	8,793	61.6
Generalised allied health professions (1916)	1,691	11.8
Loading of drug delivery device (1921)	1,631	11.4
Transfusion of blood and gamma globulin (1893)	1,274	8.9
Cerebral anaesthesia (1910)	1,269	8.9
Vascular infusion device and pump (766)	839	5.9
Abdominal hysterectomy (1268)	551	3.9
Other procedures on female genital organs (1299)	423	3.0
Therapeutic interventions on cardiovascular system (1890)	422	3.0
Postprocedural analgesia (1912)	414	2.9
Immunisation (1884)	390	2.7
Division of abdomen adhesions (986)	330	2.3
Other excision procedures on abdomen, peritoneum or omentum (989)	269	1.9
Application, insertion or removal procedures on abdomen (983)	253	1.8
Salpingo-oophorectomy (1252)	251	1.8
Conduction anaesthesia (1909)	235	1.6
Appendicectomy (926)	191	1.3
Computerised tomography of abdomen and pelvis (1963)	183	1.3
Venous catheterisation (738)	180	1.3
Laparoscopy (985)	170	1.2
Total ovarian cancer-related hospitalisations	14,277	

Table 7.4: Hospitalisation	for ovarian cancer	by most common	procedures, 2007-08
----------------------------	--------------------	----------------	---------------------

(a) Classified according to the Australian Classification of Health Interventions, fifth edition (see Appendix A).

(b) Indicates the number of hospitalisations in which the indicated procedure was undertaken.

(c) The sum of the count of hospitalisations does not equal the total number of hospitalisations since no procedure, or multiple procedures, may be undertaken during each hospitalisation. For the same reason, the sum of the percentages does not equal 100.

Source: National Hospital Morbidity Database, AIHW.

8 Expenditure on ovarian cancer

Another measure of the impact of ovarian cancer is the health care costs incurred. Generally, when expenditure for various types of cancers is considered, estimates are provided on recurrent, direct health care costs — that is, expenditure on health goods and services spent by all levels of government, private health insurers, companies, households and individuals to diagnose and treat that type of cancer. This was the approach used in the recent report on breast cancer (AIHW and NBOCC 2009).

For ovarian cancer, however, only one component of such expenditure was considered to be of sufficient quality for publication – namely, hospital admitted patient services expenditure. This type of expenditure pertains to services provided to admitted patients, including private admitted patients, in hospitals. In contrast, estimates of out-of-hospital medical expenses and prescription pharmaceuticals for ovarian cancer patients were considered to be of insufficient quality to report due to insufficient numbers in the surveys on which the expenditure estimates are based.

The latest expenditure estimates that are available on hospital admitted patient services for ovarian cancer are for the 2004–05 financial year, with comparable data available for 2000–01. The data were sourced from the Disease Expenditure Database which is maintained by the AIHW. In this database, ovarian cancer was defined to include the ICD-10 codes of 'C56 and C57.0–C57.4', which we refer to as 'ovarian and related cancers'. This definition of ovarian cancer differs from the definition used in the majority of other chapters in this report (i.e. the ICD-10 code of 'C56').

In the Disease Expenditure Database (and unlike the approach taken in Chapter 7 of this report), the data on hospital expenditure for ovarian and related cancers pertain only to those hospitalisations for which the principal diagnosis was 'ovarian and related cancers'. Thus, expenditure related to same-day hospitalisations for the administration of chemotherapy, with ovarian and related cancer patients coded as an additional, rather than a principal, diagnosis is not included. As a result, the data shown are a minimum estimate of total hospital admitted patient services expenditure on ovarian and related cancer patients.

Further information about the Disease Expenditure Database and how the expenditure estimates were derived can be found in health expenditure reports produced by the AIHW (AIHW 2005, 2008b).

Expenditure in 2004–05

Expenditure on hospital admitted patient services for ovarian and related cancers was estimated to be \$25 million in the 2004–05 financial year (Table 8.1). The corresponding value for expenditure for all cancers for females was \$884 million and, for all diseases, it was \$12,688 million. Overall, ovarian and related cancers comprised 3% of all cancer-related hospital admitted patient services expenditure for females and 0.2% of such expenditure for all diseases.

	\$ (million)	Per cent of expenditure on all cancers ^(b)	Per cent of expenditure on all diseases
Ovarian and related cancers ^(c)	25	2.9	0.2
All cancers ^(b)	884	100.0	7.0
All diseases	12,688		100.0

Table 8.1: Hospital admitted patient services expenditure^(a) by disease, females, 2004–05

(a) Pertains to those hospitalisations for which the principal diagnosis was ovarian and related cancers. Does not pertain to hospitalisations for which ovarian and related cancers was an additional diagnosis, with the principal diagnosis relating specifically to the type of cancer treatment or care received.

(b) Includes cancers coded in ICD-10 as C00-C97. Does not include cancers coded as D45, D46, D47.1 and D47.3.

(c) Includes cancers coded in ICD-10 as C56 and C57.0–C57.4.

Source: Disease Expenditure Database, AIHW.

Differences by age

Differences by age in ovarian and related cancers expenditure for hospital admitted patient services is shown in Figure 8.1. Almost one quarter (23%) of the \$25 million was spent on women aged 54 to 64 years (\$6 million). As well, 23% was spent on women aged 65 to 74 years (\$6 million), while 21% (\$5 million) was spent on women aged 75 to 84 years and 16% (4 million) on those aged 45 to 54 years.



Average expenditure per hospitalisation for ovarian and related cancers was highest for women in the older age groups. In particular, average expenditure for those aged 85 years and over was \$10,379 per hospitalisation and, for those aged 75 to 84 years, it was \$8,945 per hospitalisation. In contrast, the lowest average expenditure per hospitalisation of \$6,790 was for those aged 54 to 65 years, followed by expenditure of \$6,860 for those aged less than 35 years.

Trends

Change over time in admitted patient expenditure on ovarian and related cancers is shown in Table 8.2. After prices were adjusted for inflation (with all prices shown in 2004–05 dollars), the data indicate that admitted patient expenditure on ovarian and related cancers grew by 15% from \$22 million in 2000–01 to \$25 million in 2004–05. Further investigation of the data indicated that this increase was largely due to increased expenditure on specialists. Table 8.2 also indicates that the overall increase in estimated expenditure on ovarian and related cancers (15%) is somewhat lower than the estimated increase of 19% for all cancers and 18% for all diseases.

Table 8.2: Hospital admitted patient services expenditure ^(a) by disease, constant prices ^(b) , fema	iles,
2000-01 and 2004-05	

Ovarian and related cancers ^(d) 22 25	Sector	2000–01 \$ (million) ^(b)	2004–05 \$ (million)	Change (%) ^(c)
	Ovarian and related cancers ^(d)	22	25	15.0
All cancers ^(e) 745 884	All cancers ^(e)	745	884	18.6
All diseases 10,739 12,688	All diseases	10,739	12,688	18.1

(a) Pertains to those hospitalisations for which the principal diagnosis was ovarian and related cancers (i.e. codes of C56 and C57.0–C57.4 in ICD-10). Does not pertain to hospitalisations for which ovarian and related cancers was an additional diagnosis, with the principal diagnosis relating specifically to the type of cancer treatment or care received.

(b) Constant price health expenditure for 2000–01 is shown in terms of 2004–05 dollars.

(c) These calculations were based on exact dollars.

(d) Includes cancers coded in the ICD-10 as C56 and C57.0-C57.4.

(e) Includes cancers coded in ICD-10 as C00–C97. Does not include cancers coded as D45, D46, D47.1 and D47.3.

Source: Disease Expenditure Database, AIHW.

Appendix A: Classifications

International Statistical Classification of Diseases and Related Health Problems

The International Statistical Classification of Diseases and Related Health Problems (ICD) is used to classify diseases and other health problems (including symptoms and injuries) in clinical and administrative records. The use of a standard classification system enables the storage and retrieval of diagnostic information for clinical and epidemiological purposes that is comparable between different service providers, across countries and over time.

In 1903, Australia adopted the ICD to classify causes of death and it was fully phased in by 1906. Since 1906, the ICD has been revised nine times in response to the recognition of new diseases (e.g. acquired immunodeficiency syndrome (AIDS)), increased knowledge of diseases and changing terminology in the description of diseases. Comparability factors are sometimes required between revisions to make comparisons valid if a disease definition changed between the revisions. For ovarian cancer, a comparability factor of 0.98 applies to convert ICD-9 mortality data to ICD-10 data (ABS 2007), while a comparability factor of '1' applies to convert such data from ICD-8 to ICD-9 standards (ABS 1981).

The version currently in use, ICD-10 (WHO 1992), was endorsed by the 43rd World Health Assembly in May 1990 and officially came into use in World Health Organization (WHO) member states from 1994.

International Statistical Classification of Diseases and Related Health Problems, Australian modification

The Australian modification of ICD-10, which is referred to as the ICD-10-AM (NCCH 2008b), is based on ICD-10. ICD-10 was modified for the Australian setting by the National Centre for Classification in Health (NCCH) with assistance from clinicians and clinical coders. Compatibility with ICD-10 at the higher levels (i.e. up to 4 character codes) of the classification has been maintained. ICD-10-AM has been used for classifying diagnoses in hospital records in all states and territories since 1999–00 (AIHW 2000).

Australian Classification of Health Interventions

The current version of the ICD does not incorporate a classification system for coding health interventions (i.e. procedures). In Australia, a health intervention classification system was designed to be implemented at the same time as the ICD-10-AM in July 1998. The system was based on the Medicare Benefits Schedule (MBS) coding system and was originally called MBS-Extended. The name was changed to the Australian Classification of Health Interventions (ACHI) with the release of the third revision of the ICD-10-AM in July 2002 (NCCH 2008c). ACHI and ICD-10-AM are used together for classifying morbidity, surgical procedures and other health interventions in Australian hospital records.

International Classification of Diseases for Oncology

Cancers were originally classified solely under the ICD classification system, based on topographic site and behaviour. However, during the creation of the ninth revision of ICD in the late 1960s, working parties suggested the creation of a separate classification for cancers that included improved morphological information. The first edition of the International Classification of Diseases for Oncology (ICD-O) was subsequently released in 1976 and, in this classification, cancers were coded by both morphology (histology type and behaviour) and topography (site).

Since the first edition of the ICD-O, a number of revisions have been made, mainly in the area of lymphomas and leukaemias. The current (third) edition was released in 2000 (Fritz et al. 2000) and is currently used by most state and territory cancer registries in Australia, as well as by the AIHW in regard to the Australian Cancer Database.

Australian Standard Geographical Classification Remoteness Areas

The Australian Standard Geographical Classification (ASGC) Remoteness Areas was used to assign areas across Australia to a remoteness category (ABS 2001). This classification divides all areas of Australia into five categories – namely, *Major cities, Inner regional, Outer regional, Remote* and *Very remote* (AIHW 2004). For the purposes of this report, the categories of *Remote* and *Very remote* were collapsed due to the small number of cases in these two subgroups.

Index of Relative Socio-economic Disadvantage

The Index of Relative Socio-economic Disadvantage (IRSD) is one of four Socio-Economic Indexes for Areas (SEIFAs) developed by the Australian Bureau of Statistics (ABS 2008a). This index is based on factors such as average household income, education levels and unemployment rates. Rather than being a person-based measure, the IRSD is an area-based measure of socioeconomic status in which small areas of Australia are classified on a continuum from disadvantaged to affluent. This information is used as a proxy for the socioeconomic status of people living in those areas and may not be correct for each person living in that area. In this report, the first socioeconomic status group (labelled '1') corresponds to geographical areas containing the 20% of the population with the lowest socioeconomic status according to the IRSD and the fifth group corresponds to the 20% of the population with the highest socioeconomic status.

Standard Australian Classification of Countries

The Standard Australian Classification of Countries (SACC) is the Australian statistical standard for statistics classified by country (ABS 2008c). It is a classification of countries which is essentially based on the concept of geographic proximity. In its structure, it groups neighbouring countries into progressively broader geographical areas on the basis of their similarity in terms of social, cultural, economic and political characteristics. The first edition of the SACC was published in 1998, while the second edition – the one used in this report – was released by the ABS in 2008.

Appendix B: Statistical methods and technical notes

Age-specific rates

Age-specific rates provide information on the incidence of a particular event in an age group relative to the total number of people at risk of that event in the same age group. It is calculated by dividing the number of events occurring in each specified age group by the corresponding 'at risk' population in the same age group and them multiplying the result by a constant (e.g. 100,000) to derive the rate. Age-specific rates are often expressed per 100,000 population.

Age-standardised rates

A crude rate provides information on the number of, for example, new cases of cancer or deaths from cancer relative to the number of people in the population at risk in a specified period. No age adjustments are made when calculating a crude rate. Since the risk of cancer is heavily dependent on age, crude rates are not suitable for looking at trends or making comparisons across groups in cancer incidence and mortality.

More meaningful comparisons can be made by the use of age-standardised rates, with such rates adjusted for age in order to facilitate comparisons between populations that have different age structures (e.g. between Indigenous and other Australians). This standardisation process effectively removes the influence of age structure on the summary rate.

There are two methods commonly used to adjust for age: direct and indirect standardisation. In this report, the direct standardisation approach presented by Jensen and colleagues (1991) is used with two exceptions. The exceptions are the calculation of incidence and mortality rates by Indigenous status and by country/region of birth. There are relatively small numbers of cases of, and deaths from, ovarian cancer among Indigenous women and among Australian women born in some overseas regions; indirect age-standardisation is commonly used in such circumstances.

To age-standardise using either the direct or the indirect method, the first step is to calculate population numbers and numbers of cases (or deaths) in age ranges – typically 5-year age ranges. If direct standardisation is used, a key step is to multiply the age-specific population numbers for the reference population (e.g. the Australian population as at 30 June 2001 or the WHO 2000 World Standard Population) by the age-specific incidence rates (or death rates) for the population of interest (such as those in a certain socioeconomic status group or those who lived in *Major cities*). This is then used to derive a standardised incidence rate (or death rate) for the population of interest.

When indirect standardisation is used, a key step is to estimate an 'expected' incidence rate (or death rate) for the population of interest (such as Indigenous women) from the agespecific rates in the reference population (e.g. non-Indigenous women) and the age-specific population numbers for the population of interest. Details of the age-standardisation method used and the reference population for each of the relevant incidence and mortality analyses are summarised in Table B.1 (and noted in the footnotes to the relevant tables and graphs).

Variable	Age-standardisation method used	Reference population
State and territory	Direct	Australian population as at 30 June 2001
Remoteness area	Direct	Australian population as at 30 June 2001
Socioeconomic status	Direct	Australian population as at 30 June 2001
Indigenous status	Indirect	Non-Indigenous population
Country of birth	Indirect	Australian-born population

Table B.1: Age-standardisation method and reference population for analyses of differences in incidence and mortality rates by group

Confidence intervals

An observed value of a rate may vary due to chance, even where there is no variation in the underlying value of the rate. A confidence interval provides a range of values that has a specified probability of containing the true rate or trend. The 95% (*p*-value = 0.05) confidence interval is used in this report; thus, there is a 95% likelihood that the true value of the rate is somewhere within the stated range. Confidence intervals can be used as a guide to whether or not differences are consistent with chance variation. In cases where no values within the confidence intervals overlap, the difference between rates is greater than that which could be explained by chance and is regarded as statistically significant.

Note, however, that overlapping confidence intervals do not necessarily mean that the difference between two rates is definitely due to chance. Instead, an overlapping confidence interval represents a difference in rates which is too small to allow differentiation between a real difference and one which is due to chance variation. It can, therefore, only be stated that no statistically significant differences were found, and not that no differences exist. The approximate comparisons presented might understate the statistical significance of some differences, but they are sufficiently accurate for the purposes of this report.

As with all statistical comparisons, care should be exercised in interpreting the results of the comparison of rates. If two rates are statistically significantly different from each other, this means that the difference is unlikely to have arisen by chance. Judgement should, however, be exercised in deciding whether or not the difference is of any practical significance.

With one exception, the confidence intervals presented in this report were calculated using a method developed by Dobson and associates (1991). This method calculates approximate confidence intervals for a weighted sum of Poisson parameters.

The one exception applies to the confidence intervals that were calculated for the international comparisons of incidence and mortality using GLOBOCAN data. For those data, the lack of the required data meant that the Dobson method could not be used and the AIHW approximated the confidence intervals using the following formula:

95% CI approximation = AS rate $\pm 1.96 \times \sqrt{\frac{\text{AS rate}}{\text{Number of cases}}}$

Since the GLOBOCAN data are based on the estimates of the number of new cases and deaths from ovarian cancer, the associated confidence intervals indicate the range of random variation that might be expected, should those estimates be 100% accurate.

Note that statistical independence of observations is assumed in the calculations of the confidence intervals for this report. This assumption may not always be valid for episode-based data (such as data from the National Hospital Morbidity Database).

Definition of ovarian cancer

Definitions of ovarian cancer differ considerably in the literature, with the following two factors varying between definitions: what ICD codes are included in the definition; and whether or not borderline tumours are included. These two factors are discussed below.

What ICD codes are included in the definition

In the 9th revision of the ICD, ovarian cancers were grouped with cancers of 'other uterine adnexa' and coded as '183'. Examples of 'other uterine adnexa' cancers are cancers of the fallopian tube and of the parametrium. While the actual ICD-9 name for code '183' is 'Malignant neoplasm of ovary and other uterine adnexa', this grouping of cancers is often simply referred to as 'ovarian cancer' in the literature.

In the 10th revision of the ICD, ovarian cancers were coded separately and assigned the code of 'C56', with the remaining cancers from the ICD-9 code of '183' grouped with a number of other cancers in 'C57'. The code of 'C57' is labelled 'malignant neoplasm of other and unspecified female genital organs' (see Table B.2).

Table B.2: Codes for ovarian and associated cancers i	in the International Statistical Classification of
Diseases and Related Health Problems, tenth revisio	on (ICD-10)

ICD-10 codes	Type of cancer
C56	Malignant neoplasm of ovary
C57	Malignant neoplasm of other and unspecified female genital organs
C57.0	Fallopian tube (oviduct, uterine tube)
C57.1	Broad ligament
C57.2	Round ligament
C57.3	Parametrium (uterine ligament NOS)
C57.4	Uterine adnexa, unspecified
C57.7	Other specified female genital organs (Wolffian body or duct)
C57.8	Overlapping lesion of female genital organs (malignant neoplasm of female genital organs whose point of origin cannot be classified to any one of the categories C51–C57.7, C58) (tubo-ovarian, utero-ovarian)
C57.9	Female genital organ, unspecified (female genitourinary tract NOS)

Source: WHO 1992.

Cancers coded as 'C56' in ICD-10 are included for the analyses of ovarian cancer shown in this report that utilise the Australian Cancer Database (Chapters 2, 4 and 5), the National Mortality Database (Chapter 3) and the National Hospital Morbidity Database (Chapter 7). This is the same definition as has been used for ovarian cancer by the cancer registries in Victoria (Thursfield et al. 2009), Queensland (Queensland Cancer Registry & Cancer Council Queensland 2008) and Tasmania (Dalton et al. 2008). However, different definitions for ovarian cancer have been used by other state and territory registries. For example, New South Wales and the Australian Capital Territory include the ICD-10 codes of 'C56 and C57.0–C57.7' (ACT Health 2007; Tracey et al. 2006, 2008), Western Australia includes the ICD-10 codes of 'C56 and C57.0–C57.9' (Threllfall et al. 2005) and South Australia includes the ICD-9 code of '183' (South Australia Cancer Registry 2008).

For the analyses shown in this report that are based on GLOBOCAN data (in Chapters 2, 3 and 4), the burden of disease report (Chapter 6), and the Disease Expenditure Database (Chapter 8), ovarian cancer was defined as the ICD-10 codes of C56 and C57.0–C57.4 (see Table B.2). This grouping of cancers is referred to as 'ovarian and related cancers' in this report.

Whether or not borderline tumours are included

In the second edition of the ICD-O, ovarian tumours of borderline malignancy were considered malignant. However, in the third edition of the ICD-O, they are considered to be of uncertain behaviour and are no longer considered malignant. Thus the number of cases that are considered to be ovarian cancer will be less when ICD-O-3 coding rules are used rather than ICD-O-2 rules.

The third edition of ICD-O was released in 2000, and has been implemented in cancer registries from the early 2000s onwards. Furthermore, each of the cancer registries has recoded their cancer data holdings for all years based on the ICD-O-3 coding rules. Thus the Australian data shown in this report for all years are comparable over time and exclude borderline cases of ovarian cancer.

For the earlier edition of this report (AIHW & NBCC 2006), tumours were classified according to ICD-O-2 and thus the data presented in that report are not strictly comparable to the results shown in this report. As shown in the earlier report, 6% of the ovarian cancer cases diagnosed in 2002 were borderline tumours.

Due to the varying approaches used to define ovarian cancer in the literature, comparisons of data from different sources must be done with care.

Incidence projections

To calculate the incidence projections shown in Chapter 2, ovarian cancer incidence data for females for the 10-year period from 1997 to 2006 were divided into 18 series – one for each 5-year age group. The incidence numbers were divided by the age-specific mid-year populations to obtain the age-specific incidence rates. Least squares linear regression was used to find the straight line of best fit through the 1997 to 2006 rates and to compute the various quantities needed for the 95% prediction intervals. The projected incidence rates were then multiplied by the estimated resident population to obtain the projected incidence numbers. The populations used were the ABS projected populations from Series 29(B) (ABS 2008b).

Mortality data differences

The state and territory data on mortality due to ovarian cancer that are shown in this report may not be comparable with data published by individual state and territory cancer registries for a number of reasons, including the following (Cancer Council Queensland 2009; Tracy et al. 2008):

- The state and territory mortality data presented in this report refer to the place of a person's residence at the time of *death*. In contrast, the state and territory cancer registries generally present mortality information based on a person's place of residence at the time of *diagnosis*. In these latter data, the deaths may or may not have occurred in the state or territory indicated.
- Different approaches were used to assign cause of death. In this report, data on mortality
 for each jurisdiction were derived from the National Mortality Database (see
 Appendix C). Information on cause of death in the NMD is sourced from the ABS which
 makes use of death certificate information to assign cause of death. In contrast, the state
 and territory cancer registries tend to make use of information from a number of
 different sources, including pathology reports and other notifications, to assign a cause
 of death.

Mortality-to-incidence ratio

Both mortality-to-incidence ratios (MIRs) and relative survival ratios can be used to estimate survival from a particular disease, such as ovarian cancer, for a population. Although MIRs are the cruder of the two ratios, MIRs do not have the same comparability and interpretation problems associated with them when attempting to make international comparisons (see Chapter 4). Thus, the MIR is considered to be a better measure when comparing survival between countries.

The MIR is defined as the age-standardised mortality rate divided by the age-standardised incidence rate. For example, an MIR of 0.42 in a given year for all types of cancers means that for every 100 new cancer cases diagnosed that year, there were 42 deaths due to cancer in the same year (though the deaths need not be of the same people as the cases). If people tend to die relatively soon after diagnosis from a particular cancer (that is, the death rate is nearly as high as the incidence rate for that cancer), then the MIR will be close to 1.00. In contrast, if people tend to survive a long time after being diagnosed, then the MIR will be close to zero. The MIR only gives a valid measure of the survival experience in a population if:

- cancer registration and death registration are complete or nearly so, and
- the incidence rate, mortality rate and survival proportion are not undergoing rapid change.

The incidence and mortality data used to calculate the MIRs in Chapter 4 were extracted from the 2002 GLOBOCAN database (Ferlay et al. 2004).

Relative survival analysis

Relative survival estimates compare the survival of persons diagnosed with ovarian cancer (i.e. the observed survival) with the survival of the entire Australian population of the same sex and age in the same calendar year as the cancer cohort (i.e. the expected survival). Note that the actual cause of death (whether it is from ovarian cancer or another cause) is not of importance in these analyses. Thus, relative survival is defined as follows:

relative survival = <u>observed survival for cancer cohort</u> expected survival for 'matched' population

The resulting value is usually given as a proportion. For example, if the observed 5-year survival of a particular cohort diagnosed with ovarian cancer was 0.60 (that is, 60% of them were still alive 5 years after diagnosis) and their expected survival, based on Australian lifetables, was 0.90 (that is, 90% of people with the same age- and sex-profile as the cohort would be expected to be alive 5 years later), then the 5-year relative survival would be 0.6/0.9 = 0.67 or 67%. One way to interpret this figure is that the 'average' person in the cancer cohort has a 67% chance of being alive 5 years after diagnosis *relative to others of the same sex and age*.

In order for the relative survival estimate to be a valid approximation of the probability that a person will not die of their diagnosed cancer within the given time interval, the presence of the cancer is assumed to be the only factor that distinguishes the cancer cohort from the general population (Ries et al. 2008). The degree to which this is true is not known.

Relative survival proportions have traditionally been calculated using the 'cohort method', and NBOCC preferred the use of that method for this report. In the cohort method, a cohort of people diagnosed with cancer is followed over time to estimate the proportion surviving for a selected time frame (e.g. 1, 5 or 10 years). An alternative approach to calculating relative survival is the period method, which was developed by Brenner and Gefeller (1996). This method examines the survival experience of people who were alive at the beginning of a particular recent calendar period and who were diagnosed with cancer before this period. Therefore, the period method might provide more up-to-date estimates of survival, especially in the presence of temporal trends affected by improvements in cancer detection and treatment. However, the cohort method is thought to provide more precise estimates (i.e. estimates with narrower confidence intervals).

An alternative to the calculation of relative survival proportions is to use the 'cause-specific model' to derive survival estimates. This model calculates survival based on deaths due to cancer-related causes alone. There are various advantages and disadvantages to using the cause-specific model (Le Teuff et al. 2005). Because the 2006 version of the Australian Cancer Database that was utilised for this report included a limited amount of cause of death information, this approach could not be used to calculate survival estimates.

Data from the ACD on the incidence of ovarian cancer were used to calculate observed survival proportions. These incidence data were linked to the National Death Index in order to obtain information on those people with ovarian cancer who died and the date on which this occurred (see Appendix C for more information on the data sets). In order to calculate the expected survival belonging to the age-, sex- and calendar-year matched population, ABS life tables for the population under study were used (ABS 2009b).

When comparing relative (or crude) survival estimates over time and/or between population subgroups it would seem appropriate to age-standardise the figures in order to

remove potential confounding by different age-structures. However, there are some undesirable features of doing so, as well as some difficulties in interpreting directly agestandardised survival estimates (Brenner et al. 2004; Brenner & Hakulinen 2003). For example, when numbers are small the age-specific survival for a certain age group and population subgroup may be undefined; hence the age-standardisation procedure breaks down. Also, the calculation of age-standardised survival can produce a figure that differs substantially from the unadjusted survival, even if it is adjusted to the original age distribution of the study population. In light of these and related shortcomings in the procedure, it was decided not to age-standardise the survival estimates produced for this report.

The software used to calculate the relative survival proportions was written by Dickman (2004). It uses the Ederer II method of calculating the interval-specific expected survivals. Further details on the approach used to calculate the relative survival estimates, including rules which were applied during data preparation, can be found in the 2008 report prepared by the AIHW on cancer survival and prevalence (AIHW, CA & AACR 2008).

Risk to age 75 and 85 years

The calculations of risk shown in this report are measures that approximate the risk of developing (or dying from) ovarian cancer before a given age, assuming that the risks at the time of estimation remained throughout life. It is based on a mathematical relationship with the cumulative rate. Note that in these risk factors, no account is taken of specific ovarian cancer risk factors. Further details on how the risks were calculated can be found in the 2008 *Cancer in Australia* report (AIHW & AACR 2008).

Appendix C: Data sources

To provide a comprehensive picture of national ovarian cancer statistics in this report, a range of data sources were used, including AIHW and external data sources. These data sources are described in this appendix.

Australian Cancer Database

The Australian Cancer Database (ACD) is a database that holds information about 1.8 million cancer cases of Australian residents who were diagnosed with cancer (other than basal call and squamous cell carcinomas of the skin) between 1982 and 2006. Data from this source are used in Chapters 2, 4 and 5.

The AIHW compiles and maintains the ACD, in partnership with the Australasian Association of Cancer Registries (AACR), whose member registries provide data to the AIHW on an annual basis. Each Australian state and territory has legislation that makes the reporting of all cancers (excluding basal call and squamous cell carcinomas of the skin) mandatory. Pathology laboratories and Registrars of Births, Deaths and Marriages across Australia must report on cancer cases, as do hospitals, radiation oncology units and nursing homes in some (but not all) jurisdictions.

The data provided to the AIHW by the state and territory cancer registries include, at a minimum, an agreed set of items that provide information about the individual with the cancer, and the characteristics of the cancer (see Table C.1). In addition to the agreed set of items, registries often provide other data which are also included in the ACD. For example, data on ductal carcinoma in situ (DCIS) are not part of the agreed ACD data set but are regularly provided by the state and territory registries.

Once the data are received from the state and territory cancer registries, the AIHW assembles the data into the ACD. Internal linking checks are undertaken to identify those who had tumours diagnosed in more than one state or territory; this process reduces the degree of duplication within the ACD to a negligible rate. The ACD is also linked with information on deaths (from the National Death Index) in order to add information on which people with cancer have died (from any cause). Any conflicting information and other issues with the cancer data are resolved through consultation with the relevant state or territory cancer registry.

The registration of cases of cancer is a dynamic process such that records in the state and territory cancer registries may be modified if new information is received. Thus, records in the cancer registries are always open and they are updated as required. In order for these changes to be incorporated into the ACD, a new complete file for all years of cancer data is provided by each of the jurisdictions annually. As a result, the number of cancer cases reported by the AIHW for any particular year may change slightly over time and, in addition, data published by a cancer registry at a certain point in time may differ to some extent from what is published by the AIHW.

Person-level attributes	Tumour-level attributes
Person identification number (assigned by the state/territory)	Tumour identification number (assigned by the state/territory)
Sumame	Date of diagnosis
First given name	Date of diagnosis flag
Second given name	Age at diagnosis
Third given name	ICD-O-3 ^(a) topography code
Sex	ICD-O-3 ^(a) morphology code
Date of birth	ICD-10 ^(b) disease code
Date of birth flag	Most valid basis of diagnosis
Indigenous status	Statistical local area at diagnosis
Country of birth	Postcode at diagnosis
Date of death	Melanoma thickness
Age at death	
Cause of death	

Table C.1: Agreed set of items to be provided by the states and territories to the AIHW for inclusion in the Australian Cancer Database

(a) International Classification of Diseases for Oncology, 3rd edition (see Appendix A).

(b) International Statistical Classification of Diseases and Related Health Problems, 10th revision (see Appendix A).

Source: AIHW 2009b.

Non-melanoma skin cancers

Data on all types of cancer, other than two types of non-melanoma skin cancer (NMSC), are reportable and collected by the state and territory registries. The two most common types of NMSC – namely, basal cell carcinoma (BCC) and squamous cell carcinoma (SCC) – are not reportable and are thus not generally recorded in cancer registries in Australia. These two types of skin cancers are by far the most frequently diagnosed cancers in Australia for both males and females (AIHW & CA 2008). A number of other, rarer types of cancer also fall within the NMSC category (e.g. Merkel cell lesions, Kaposi sarcoma and cutaneous lymphoma) and these are reportable cancers.

In the past, the agreed approach was to exclude all NMSC cases from the cancer incidence data produced by the AIHW. However, for the first time this year, a new approach was used whereby all cases that pertained to reportable forms of NMSC were included in the incidence data; as previously, no data on BCC and SCC were included. To implement this change, the state and territory registries were asked to supply, along with the usual data, information on all NMSC cases other than BCC and SCC for 2006 and for all previous years, where possible. All of the registries were able to provide such data for 2001 to 2006, with only some being able to provide such data for earlier years. Thus the data on non-melanoma skin cancers other than BCC and SCC may be incomplete before 2001.

Burden of disease data

Information on the burden of disease from ovarian cancer is shown in Chapter 6 of this report.

The first study that provided a comprehensive overview of disease and injury burden in Australia was published in 1999 (AIHW: Mathers et al. 1999). The second and most recent such study was published in 2007, and it provides burden of disease information in relation to 2003 (Begg et al. 2007a,b). The summary measure used in that study is the disabilityadjusted life year or DALY, with this term used interchangeably with 'burden of disease'. The DALY quantifies the gap between a population's actual health status and some 'ideal' or reference status, with time (either lived in health states or lost through premature death and illness) being the unifying 'currency' for combining the impact of mortality and non-fatal health outcomes.

A DALY for a disease or health condition is calculated as the sum of the years of life lost due to premature mortality (YLL) in the population and the equivalent 'healthy' years lost due to disability (YLD) for incident cases of the health condition such that:

```
DALY = YLL + YLD
```

where YLL = number of deaths x standard life expectancy at age of death and

YLD = incidence x duration x severity weight.

Further information about how the DALY was derived, as well as further information on interpretation of burden of disease data, can be found in Begg and associates (2007a).

In the burden of disease study, ovarian cancer was defined to include the ICD-10 codes of 'C56 and C57.0–C57.4'; this set of codes is referred to as 'ovarian and related cancers' in this report (see Appendix B).

Disease Expenditure Database

Expenditure data are used in Chapter 8 of this report to describe health expenditure on ovarian cancer. These data were obtained from the Disease Expenditure Database which is maintained by the AIHW.

Since 1984, the AIHW has had responsibility for developing estimates of national health expenditure. Data for this purpose are obtained from a wide variety of sources in the public and private sectors, with most of the data being provided by the ABS, the Australian Government Department of Health and Ageing, and state and territory health authorities. Other major sources are the Department of Veterans' Affairs, the Private Health Insurance Administration Council, Comcare, and the major workers compensation and compulsory third-party motor vehicle insurers in each state and territory.

The definition of ovarian cancer used in this database is the ICD-10 codes of 'C56 and C57.0–C57.4', which we refer to as 'ovarian and related cancers' (see Appendix B). Expenditure data for just the ICD-10 code of 'C56' were not available.

In the Disease Expenditure Database (and unlike the approach taken in Chapter 7 of this report), ovarian and related cancer hospitalisations are defined as those hospitalisations for which the *principal diagnosis* was ovarian and related cancer. Therefore, hospitalisations that involved same-day chemotherapy administration for ovarian and related cancer patients (with ovarian cancer coded as an *additional diagnosis* rather than a principal diagnosis) are not

included. In turn, any spending related to these latter hospitalisations is not included in the expenditure data for hospital admitted patient services for ovarian and related cancers. Thus, the data shown are a minimum estimate of total admitted patient services expenditure on ovarian and related cancer patients. Note that in future expenditure analysis work done by the AIHW, further work to identify the costs of chemotherapy that are due to specific types of cancers may be undertaken.

The definition of 'all cancers' used in Chapter 8 is somewhat different from that used in earlier chapters, as it only includes the ICD-10 'C' codes and excludes those malignant cancers with the ICD-10 'D' codes (such as polycythaemia vera). Separate expenditure data were not readily available for the required subset of ICD-10 'D' cancers. Since the forms of malignant cancers covered by the ICD-10 'D' codes are not common (see AIHW & AACR 2008), their exclusion is not expected to have a large effect on the health expenditure estimates shown in this report.

Further information about the Disease Expenditure Database can be found in the annual health expenditure reports published by the AIHW (AIHW 2008b).

GLOBOCAN

One of the main sources of internationally comparable data on cancer is the GLOBOCAN database which is prepared by the International Agency for Research on Cancer (IARC) (Ferlay et al. 2004). The IARC collates cancer incidence and mortality data from cancer registries around the world and uses those data to produce estimates for a 'common year'. The most recent GLOBOCAN estimates for which data could be obtained are for 2002, with these estimates based on cancer incidence rates from approximately 3 to 5 years earlier. GLOBOCAN data are shown in Chapters 2, 3 and 4 of this report.

For the GLOBOCAN data, ovarian cancer was defined as those cancers that were coded as 'C56 and C57.0–C57.4' in ICD-10. Thus the definition used in those data is broader than that used in most other sections of this report. While not clearly stated, we presume that borderline ovarian tumours have not been included since IARC indicates that the third edition of ICD-O was used. As noted in Chapter 1, in the third edition of ICD-O (and unlike the previous edition), borderline ovarian tumours were not considered to be malignant tumours.

In the GLOBOCAN database, age-standardised incidence and mortality rates are provided, with the data standardised to the Doll et al. (1966) World Standard Population. However, the database does not include confidence intervals. In order to provide some guidance in terms of whether the differences were statistically significant, the AIHW calculated approximate confidence intervals (with the methodology for doing so explained in Appendix B).

National Death Index

Cancer incidence data were linked to the National Death Index (NDI) in order to provide survival and prevalence information (Chapters 4 and 5). The NDI is a database that is maintained by the AIHW; it contains information on all deaths that have occurred in Australia since 1980.

The NDI database comprises the following variables for each deceased person: name; alternative names (including maiden names); date of birth (or estimated year of birth); age at death; sex; date of death; marital status; Indigenous status; state or territory of death

registration; and death registration number. Cause of death information in a coded form is also available. For records to 1996, only the code for the underlying cause of death is available. For records from 1997, the codes for the underlying cause of death and all other causes of death mentioned on the death certificate are available.

This database exists solely for research linkage purposes, such as gaining epidemiological mortality information on individuals in a particular cohort, or with a known disease state. Ethics approval is required for the NDI to be utilised for any particular research project.

National Hospital Morbidity Database

Data from the National Hospital Morbidity Database (NHMD) are used in Chapter 7 to examine the number of ovarian cancer-related hospitalisations. The NHMD contains demographic, diagnostic, procedural and duration of stay information on episodes of care for patients admitted to hospital. This annual collection is compiled and maintained by the AIHW, using data supplied by state and territory health authorities. Information from almost all hospitals in Australia is included in the database: public acute and public psychiatric hospitals; private acute and psychiatric hospitals; and private free-standing day hospital facilities. The database is episode-based and it is not possible to count patients individually.

Data are held in the NHMD for the years from 1993–94 to 2007–08. However, around 1998–99, hospitals across Australia began to implement a change in the classification system used to code the diagnosis for hospitalisations (i.e. from ICD-9-AM to ICD-10-AM). The first full year for which national data are available using ICD-10-AM is 1999–00. Hence, in Chapter 7, data from 1999–00 onwards are presented.

The hospitalisations data presented in this report exclude those hospitalisations for which the care type was reported as *newborn*, *hospital boarder* or *posthumous organ procurement*. Thus, it includes all other admitted care hospitalisations including those with a care type of *acute care*, *rehabilitation care* and *palliative care*.

Comprehensive hospital statistics from the NHMD are released by the AIHW on an annual basis (AIHW 2009a). Further information about this data source is available in those reports.

National Mortality Database

Data from the National Mortality Database are used in Chapter 3 to provide statistical information on mortality in Australia due to ovarian cancer.

The registration of deaths has been compulsory since the mid-1850s and this information is registered with the relevant state and territory Registrar of Births, Deaths and Marriages. Since 1906, the Commonwealth Statistician has compiled the information collected by the Registrars and published national death information.

The National Mortality Database, which is maintained by the AIHW, currently contains information for all deaths in Australia registered from 1964 to 2006.

The information on deaths from the Registrars is coded nationally by the ABS according to rules set forward in various versions of the ICD. Deaths are coded to reflect the underlying cause of death. As well, since 1997, multiple causes of death have been added to the mortality data.

Over time, changes have been made to the coding and processing of mortality data and these have affected the comparability of the data. For instance, data holdings on cause of death for

1987 to 1996 were manually coded using the ninth revision of the ICD, while the corresponding data for 1997 onwards were coded using ICD-10, using an automated system with slightly different coding rules. The change to the coding and processing of mortality data introduced a break in the time series. Where possible, the ABS has developed comparability factors so that a time series may still be derived (ABS 2009d). As noted in Appendix A, for ovarian cancer, the comparability factor for ICD-9 to ICD-10 is 0.98.

Note, though, that due to changes in classifications over time and the way in which diseases were grouped in these classifications, data on deaths due to ovarian cancer are only available from 1968 onwards. Before 1968, the data on deaths due to ovarian cancer were grouped together with deaths due to cancers of 'other uterine adnexa' (which are coded separately from C56 in ICD-10).

In the National Mortality Database, information on the dates of death is provided in two different ways: one is based on the year in which people *died* and the other is based on the year in which the deaths were *registered*. For the purposes of this report, mortality data are shown based on the year of *death*, except for the most recent year (namely, 2006) where the number of people whose death was *registered* in that year is used. Previous investigation has shown that the year of death and year of registration, for the most part, coincide. However, in some instances, deaths at the end of each calendar year may be held over until the following year, as are deaths whose cause requires further examination by a coroner (e.g. possible suicides). Thus, year of death information for the latest available year generally underestimates the true number of deaths, with the number of deaths registered in that year being closer to the true value.

Population data

Throughout this report, population data were used to derive rates of, for example, cancer incidence and mortality. The population data were sourced from the ABS Demography section using the most up-to-date estimates available at the time of analysis.

To derive estimates of the resident populations, the ABS uses the 5-yearly Census of Population and Housing data as follows:

- all respondents to the Census are coded in relation to their state or territory, statistical local area and postcode of usual residence; overseas visitors are excluded
- an adjustment is made for persons missed in the Census (approximately 2%)
- Australians temporarily overseas on Census night are added to the usual residence Census count.

Estimated resident populations are then updated each year from the census data using indicators of population change, such as births, deaths and net migration. More information is available from the ABS website <www.abs.gov.au>.

For the Indigenous comparisons presented in this report (Chapters 2, 3 and 4), the most recently released Indigenous experimental estimated resident populations, as released by the ABS, were used (ABS 2009f). Those estimates were based on the 2006 Census.

Appendix D: Additional tables

Additional tables for Chapter 2: Incidence of ovarian cancer

Age group (years)	Number of cases	Age-specific rate ^(a)	95% confidence interval
<20	13	0.5	0.3–0.8
20–24	10	1.4	0.7–2.5
25–29	13	1.9	1.0–3.2
30–34	16	2.1	1.2–3.5
35–39	33	4.3	3.0–6.0
40–44	45	5.8	4.3–7.8
45–49	99	13.1	10.7–16.0
50–54	128	18.7	15.6–22.2
55–59	137	21.5	18.1–25.5
60–64	119	24.1	20.0–28.9
65–69	162	41.0	34.9–47.8
70–74	120	36.7	30.4–43.8
75–79	119	39.7	32.9–47.5
80–84	115	48.0	39.6–57.6
85+	97	44.5	36.1–54.3
Total ^(b)	1,226	10.7	10.1–11.4

Table D2.1:	Incidence of	ovarian	cancer by	age at	diagnosis,	2006
			J	0		

(a) Number of cases per 100,000 females.

(b) The rate shown in this row was age-standardised to the Australian population as at 30 June 2001 and is expressed per 100,000 females.

Source: Australian Cancer Database, AIHW.

Year	Number	ASR (A) ^(a)	95% confidence interval	ASR (W) ^(b)	95% confidence interval	Per cent of gynaecological cancer cases ^(c)	Per cent of female cancer
1092	01 04000	10.4	11 6 12 2		0.2 10.6	20.2	2.0
1902	033	12.4	11.0-13.3	9.9	9.2-10.6	20.3	3.0
1903	07	12.4	11.0-13.3	9.0	9.1-10.5	20.0	3.0 0 -
1984	875	12.5	11.7–13.4	9.8	9.2–10.5	27.9	3.7
1985	881	12.3	11.5–13.2	9.7	9.0–10.4	27.7	3.6
1986	873	11.9	11.1–12.7	9.4	8.7–10.0	27.2	3.5
1987	905	12.1	11.3–12.9	9.5	8.9–10.2	27.3	3.4
1988	895	11.7	10.9–12.5	9.2	8.6–9.9	27.5	3.3
1989	1,015	12.8	12.0–13.6	10.0	9.4–10.7	30.2	3.6
1990	1,006	12.6	11.8–13.4	9.8	9.2–10.4	29.6	3.5
1991	1,013	12.4	11.7–13.2	9.6	9.0–10.3	28.3	3.3
1992	1,036	12.5	11.8–13.3	9.9	9.3–10.5	29.0	3.3
1993	1,077	12.6	11.9–13.4	9.9	9.3–10.5	29.8	3.3
1994	1,064	12.3	11.6–13.1	9.6	9.1–10.3	27.5	3.1
1995	1,079	12.2	11.5–13.0	9.4	8.9–10.0	29.2	3.0
1996	1,076	11.9	11.2–12.6	9.1	8.5–9.7	29.6	3.0
1997	1,057	11.4	10.7–12.1	8.7	8.2–9.3	29.4	2.9
1998	1,123	11.8	11.1–12.5	9.1	8.6–9.7	30.2	3.0
1999	1,135	11.6	11.0–12.3	9.0	8.4–9.5	30.6	3.0
2000	1,137	11.3	10.7–12.0	8.6	8.1–9.2	29.8	2.8
2001	1,125	10.9	10.3–11.6	8.3	7.8–8.8	29.8	2.7
2002	1,228	11.7	11.0–12.3	8.9	8.4–9.4	31.2	2.9
2003	1,128	10.6	9.9–11.2	8.2	7.7–8.7	28.8	2.7
2004	1.267	11.5	10.9–12.2	8.8	8.3-9.4	30.3	2.9
2005	1 219	10.9	10 2-11 5	8.3	78-88	28.9	2.0
2006	1,226	10.7	10.1–11.4	8.3	7.8–8.8	28.9	2.7

Table D2.2: Inci	idence of	ovarian	cancer	. 1982 to) 2006
	Mentee or	0 / mi imii	cancer	, _, _,	

(a) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females.

(d) The age-standardised rates were standardised using the WHO 2000 World Standard Population and are expressed per 100,000 females.

(c) Includes cancers coded in ICD-10 as C51–C58.

(b) Includes cancers coded in ICD-10 as C00–C97, D45, D46, D47.1 and D47.3 with the exception of those C44 codes which indicate a basal or squamous cell carcinoma of the skin. Due to changes over time in which cancers were reportable, the data on cancers that begin with an ICD-10 code of 'D' may be incomplete before 2003 and data on C44 codes other than basal or squamous cell carcinomas may be incomplete before 2001.

Source: Australian Cancer Database, AIHW.

	<50 y	vears	50–6	9 years	70+	years	All a	ges
Year	ASR ^(a)	95% CI	ASR ^(a)	95% CI	ASR ^(a)	95% CI	ASR ^(a)	95% CI
1982	3.9	3.3–4.5	32.7	29.7–35.9	35.8	31.1–41.0	12.4	11.6–13.3
1983	3.7	3.2–4.3	30.7	27.9–33.8	41.3	36.3–46.7	12.4	11.6–13.3
1984	3.8	3.3–4.5	30.6	27.7–33.6	41.6	36.7–46.9	12.5	11.7–13.4
1985	4.1	3.5–4.7	28.6	25.8–31.5	42.2	37.3–47.5	12.3	11.5–13.2
1986	4.0	3.5–4.6	27.9	25.3–30.8	39.2	34.6-44.2	11.9	11.1–12.7
1987	3.5	3.0-4.0	30.7	27.9–33.7	39.6	35.1–44.6	12.1	11.3–12.9
1988	3.8	3.3–4.3	28.0	25.4–30.9	38.8	34.4–43.7	11.7	10.9–12.5
1989	3.6	3.1–4.1	31.9	29.1–34.9	44.2	39.4–49.3	12.8	12.0–13.6
1990	3.9	3.4-4.5	28.1	25.5–30.9	46.7	41.9–51.9	12.6	11.8–13.4
1991	3.6	3.1–4.1	29.8	27.1–32.7	44.9	40.2–49.9	12.4	11.7–13.2
1992	4.1	3.6–4.6	29.7	27.1–32.6	41.9	37.5–46.6	12.5	11.8–13.3
1993	4.0	3.5–4.5	28.8	26.2–31.6	45.3	40.8–50.2	12.6	11.9–13.4
1994	3.5	3.1–4.0	30.5	27.8–33.3	42.5	38.2–47.2	12.3	11.6–13.1
1995	3.5	3.1–4.0	28.1	25.5–30.8	46.5	42.1–51.3	12.2	11.5–13.0
1996	3.2	2.8–3.7	26.4	24.0–29.1	48.4	43.9–53.2	11.9	11.2–12.6
1997	3.1	2.7–3.6	25.9	23.6–28.5	44.7	40.5–49.3	11.4	10.7–12.1
1998	3.1	2.7–3.5	27.6	25.2–30.2	45.8	41.6–50.4	11.8	11.1–12.5
1999	3.1	2.7–3.6	27.1	24.8–29.7	45.1	41.0–49.5	11.6	11.0–12.3
2000	3.0	2.6–3.4	25.5	23.2–27.9	46.6	42.4–51.0	11.3	10.7–12.0
2001	2.7	2.4–3.2	24.7	22.5–27.0	45.4	41.4–49.7	10.9	10.3–11.6
2002	3.2	2.8–3.7	25.4	23.2–27.8	48.4	44.2–52.8	11.7	11.0–12.3
2003	2.9	2.5–3.3	24.5	22.4–26.8	40.6	36.8–44.7	10.6	9.9–11.2
2004	3.2	2.8–3.7	25.3	23.2–27.6	47.0	43.0–51.4	11.5	10.9–12.2
2005	2.9	2.5–3.3	24.1	22.1–26.3	44.9	40.9–49.1	10.9	10.2–11.5
2006	3.1	2.7–3.5	24.6	22.6–26.8	40.9	37.1–44.9	10.7	10.1–11.4

Table D2.3: Incidence of ovarian cancer by age at diagnosis, 1982 to 2006

(a) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females. Source: Australian Cancer Database, AIHW.

Year	Projected number of cases ^(a)	95% prediction interval	Age- standardised rate (A) ^(b)	95% prediction interval	Age- standardised rate (W) ^(c)	95% prediction interval
2007	1,265	1,166–1,363	10.8	10.0–11.6	8.2	7.6–8.9
2008	1,284	1,177–1,390	10.7	9.8–11.5	8.2	7.5–8.8
2009	1,304	1,189–1,419	10.6	9.7–11.5	8.1	7.4–8.8
2010	1,324	1,200–1,449	10.5	9.6–11.4	8.0	7.3–8.7
2011	1,345	1,210–1,480	10.4	9.5–11.4	8.0	7.2–8.7
2012	1,368	1,223–1,514	10.3	9.3–11.4	7.9	7.1–8.7
2013	1,390	1,233–1,548	10.3	9.2–11.3	7.8	7.0–8.7
2014	1,412	1,243–1,581	10.2	9.0–11.3	7.8	6.9–8.6
2015	1,434	1,252–1,616	10.1	8.9–11.3	7.7	6.8–8.6

Table D2.4: Projected ovarian cancer incidence, 2007 to 2015

(a) The projections were based on ovarian cancer incidence data for 1997 to 2006.

(b) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females.

(c) The age-standardised rates were standardised using the WHO 2000 World Standard Population and are expressed per 100,000 females.

Source: Australian Cancer Database, AIHW.

Table D2.5: Grouping of ovarian cancer histology types

Type of ovarian cancer ^(a)	Corresponding ICD-O-3 histology codes
1: Carcinoma (epithelial tumours)	All codes included in groups 1.1 to 1.7
1.1: Serous carcinoma	8441, 8442, 8450, 8460–8463, 9014
1.2: Mucinous carcinoma	8470–8490, 9015
1.3: Endometrioid carcinoma	8380–8383, 8560, 8570
1.4: Clear cell carcinoma	8310–8313, 9110
1.5: Adenocarcinoma not otherwise specified	8140–8147, 8170–8190, 8211–8231, 8260, 8384, 8440, 8576
1.6: Other specified carcinoma	Includes 8041, 8050, 8070, 8120, 8323, 9000 and all other specified carcinomas
1.7: Unspecified carcinoma	8010–8035
2: Sex cord-stromal tumours	8590–8671, 8810
3: Germ cell tumours	8240-8246, 8340, 9060-9102, 9473, 9501
4: Other specified malignant neoplasm	Includes 8800, 8890, 8930, 8935, 8950, 8951, 8980 and all other specified tumours
5: Unspecified malignant neoplasm	8000–8005

(a) For the purposes of this study, the grouping of ovarian cancer histology types was based primarily on those recommended by the International Agency for Research on Cancer (Curado et al. 2007) with additional input from National Breast and Ovarian Cancer Centre. All cases included in each of the groups were coded by state and territory cancer registries as primary site, invasive ovarian cancers.

9
8
й
7
З
ō
2
0
-
5
õ
-
ų.
òò
6
Γ.
S,
. <u>.</u>
9
E
• H
σ
Ľ
ų,
ğ
N.
5
6
Ĩ,
at
Ĕ
Ŧ
ŝ
ő
Ē
q
. Б
8
2
E
Ē
Ξ
2
<u>ک</u>
Ъ
ē
2
ar
Ü
드
5
.Е
ğ
2
0
~
å
5
. ب
₹ A
9
e S
ž
e
σ
.2
ă
H
ö
і,
$\tilde{\Box}$
le l
-

		Number of	cases			Per ce	nt	
Type of ovarian cancer ^(a)	1982–1987	1988–1993	1994–1999	2000-2006	1982–1987	1988–1993	1994–1999	2000–2006
1: Carcinoma (epithelial tumours)	895	1,094	1,062	1,189	81.0	82.3	82.3	79.3
1.1: Serous carcinoma	327	454	444	522	29.6	34.2	34.4	34.8
1.2: Mucinous carcinoma	157	207	222	173	14.2	15.6	17.2	11.5
1.3: Endometrioid carcinoma	123	149	163	202	11.1	11.2	12.6	13.5
1.4: Clear cell carcinoma	43	82	92	120	3.9	6.2	7.1	8.0
1.5: Adenocarcinoma NOS	185	143	94	91	16.7	10.8	7.3	6.1
1.6: Other specified carcinoma	11	17	15	47	1.0	1.3	1.2	3.1
1.7: Unspecified carcinoma	49	42	32	34	4.4	3.2	2.5	2.3
2: Sex cord-stromal tumours	32	30	27	31	2.9	2.3	2.1	2.1
3: Germ cell tumours	131	172	161	216	11.9	12.9	12.5	14.4
4: Other specified malignant neoplasm	16	20	22	34	1.5	1.5	1.7	2.3
5: Unspecified malignant neoplasm	31	13	18	29	2.8	1.0	1.4	1.9
Total	1,105	1,329	1,290	1,499	100.0	100.0	100.0	100.0

5
~
2
2
Ņ
1
2
2
2
2
0
Ŧ
N
5
~~~
<u> </u>
<u> </u>
~
2
ŝ
5
à
ŝ
0
ē
50
ച്ചാ
<u>, , , , , , , , , , , , , , , , , , , </u>
1
9
Ę
5
Ś
Ξ.
ъ
e
≻
÷.
5
9
0
Ŧ
Ś
2
ц)
5
- ŭ
50
2
ā
ē
9
0
5
>
- H
e.
2
9
_
σ.
G
n ca
nn ca
ian ca
rian ca
arian ca
varian ca
ovarian ca
ovarian ca
of ovarian ca
of ovarian ca
e of ovarian ca
pe of ovarian ca
vpe of ovarian ca
type of ovarian ca
r type of ovarian ca
y type of ovarian ca
by type of ovarian ca
e by type of ovarian ca
ce by type of ovarian ca
ice by type of ovarian ca
nce by type of ovarian ca
lence by type of ovarian ca
dence by type of ovarian ca
idence by type of ovarian ca
ncidence by type of ovarian ca
incidence by type of ovarian ca
Incidence by type of ovarian ca
?: Incidence by type of ovarian ca
.7: Incidence by type of ovarian ca
2.7: Incidence by type of ovarian ca
<b>32.7: Incidence by type of ovarian ca</b>
D2.7: Incidence by type of ovarian ca
e D2.7: Incidence by type of ovarian ca
le D2.7: Incidence by type of ovarian ca

Type of ovarian cancer ^(a) 1982–1987         1988–1993         1994–1999         2000           1: Carcinoma (epithelial tumours)         2,396         2,545         2,594         2,594           1: Carcinoma (epithelial tumours)         2,396         2,545         2,594         2,1422           1: Serous carcinoma         826         1,119         1,422         1,422           1: Serous carcinoma         289         300         238         252           1: Serous carcinoma         289         287         252         252           1: Serous carcinoma         289         287         252         252           1: Serous carcinoma         141         164         182         263           1: Set cord-stroma NOS         169         23         28         382           1: Set cord-stroma NOS         169         23         28         382           1: Set cord-stroma NOS         169         124         90         37           1: T: Unspecified carcinoma         143         124         90         37           1: T: Unspecified carcinoma         15         11         37         37           2: Set cord-stromal tumours         15         12         90         37	1982–1987     1988–1993       2,396     2,545       2,396     2,545       826     1,119       289     300       289     300       289     287       141     164       692     528	<b>1994–1999</b> 2,594 1,422 238 252	2000-2006			1	
1: Carcinoma (epithelial tumours)       2,396       2,545       2,594         1: Carcinoma (epithelial tumours)       826       1,119       1,422         1: Serous carcinoma       829       300       238         1: Serous carcinoma       289       300       238         1: Sendometrioid carcinoma       289       300       238         1: Sendometrioid carcinoma       289       287       252         1: A: Clear cell carcinoma       141       164       182         1: A: Clear cell carcinoma       141       164       182         1: 5: Adenocarcinoma NOS       692       528       382         1: 6: Other specified carcinoma       16       23       28         1: 7: Unspecified carcinoma       143       124       90         2: Sex cord-stromal tumours       15       12       90         3: Germ cell tumours       15       15       11	2,396 2,545 826 1,119 289 300 287 141 164 692 528	2,594 1,422 238 252		1982–1987	1988–1993	1994–1999	2000-2006
1.1: Serous carcinoma       826       1,119       1,422         1.2: Mucinous carcinoma       289       300       238         1.3: Endometrioid carcinoma       289       287       252         1.3: Endometrioid carcinoma       289       287       252         1.4: Clear cell carcinoma       141       164       182         1.5: Adenocarcinoma NOS       692       528       382         1.5: Adenocarcinoma NOS       143       164       182         1.5: Adenocarcinoma NOS       692       528       382         1.6: Other specified carcinoma       16       23       28         1.6: Other specified carcinoma       143       124       90         2: Sex cord-stromal tumours       15       12       11         3: Germ cell tumours       15       12       90	826 1,119 289 300 287 141 164 692 528	1,422 238 252	3,268	92.5	93.0	93.2	92.5
1.2: Mucinous carcinoma       289       300       238         1.3: Endometrioid carcinoma       289       287       252         1.3: Endometrioid carcinoma       289       287       252         1.4: Clear cell carcinoma       141       164       182         1.4: Clear cell carcinoma       692       528       382         1.5: Adenocarcinoma NOS       692       528       382         1.6: Other specified carcinoma       16       23       28         1.6: Other specified carcinoma       143       124       90         2: Sex cord-stromal tumours       15       12       90         3: Germ cell tumours       15       12       90	289 300 289 287 141 164 692 528	238 252	1,872	31.9	40.9	51.1	53.0
1.3: Endometrioid carcinoma       289       287       252         1.4: Clear cell carcinoma       141       164       182         1.5: Adenocarcinoma NOS       692       528       382         1.5: Adenocarcinoma NOS       692       528       382         1.6: Other specified carcinoma       16       23       28         1.7: Unspecified carcinoma       143       124       90         2: Sex cord-stromal tumours       15       12       11         3: Germ cell tumours       15       12       11	289 287 141 164 692 528	252	240	11.2	11.0	8.6	6.8
1.4: Clear cell carcinoma       141       164       182         1.5: Adenocarcinoma NOS       692       528       382         1.5: Adenocarcinoma NOS       692       528       382         1.6: Other specified carcinoma       16       23       28         1.7: Unspecified carcinoma       143       124       90         2: Sex cord-stromal tumours       48       39       37         3: Germ cell tumours       15       12       11	141 164 692 528		343	11.2	10.5	9.1	9.7
1.5: Adenocarcinoma NOS       692       528       382         1.5: Other specified carcinoma       16       23       28         1.7: Unspecified carcinoma       143       124       90         2: Sex cord-stromal tumours       48       39       37         3: Germ cell tumours       15       12       11	692 528	182	234	5.4	6.0	6.5	6.6
1.6: Other specified carcinoma1623281.7: Unspecified carcinoma143124902: Sex cord-stromal tumours4839373: Germ cell tumours1512114. Other conclused and increational conclused and increased	0	382	387	26.7	19.3	13.7	11.0
1.7: Unspecified carcinoma     143     124     90       2: Sex cord-stromal tumours     48     39     37       3: Germ cell tumours     15     12     11       4. Other consisted maliament complexes     60     60     60	16 23	28	91	0.6	0.8	1.0	2.6
2: Sex cord-stromal tumours 48 39 37 3: Germ cell tumours 15 12 11	143 124	06	101	5.5	4.5	3.2	2.9
3: Germ cell tumours 15 12 11	48 39	37	35	1.9	1.4	1.3	1.0
1. Other encodered medication according and a contract of the	15 12	11	35	0.6	0.4	0.4	1.0
	60 09	83	129	2.3	3.6	3.0	3.7
5: Unspecified malignant neoplasm 72 42 57	72 42	57	67	2.8	1.5	2.1	1.9
Total 2,591 2,737 2,782	2,591 2,737	2,782	3,534	100.0	100.0	100.0	100.0

(a) All cases were coded as primary site, invasive ovarian cancers. Appendix Table D2.5 provides a list of the histology types included in each group. Source: Australian Cancer Database, AIHW.

90
20
ģ
200
5
51
198
2
198
Š
osi
Ē
lia
ato
EL 6
OVE
q
an
ars
ye
2
g
age
en
E
M
er,
nc
Ca
an
ari
0
of
pe
ţ
by
ICe
len
cic
In
8
D
le

Type of ovarian cance ¹⁶¹ 1982–1937         1983–1935         1994–1939         2000–2006         1383         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1393         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1313         1		Per cent	
1: Carcinoma (epithelial turnours)       1,296       1,716       2,683         1.1: Serous carcinoma       386       622       862       1,161         1.2: Mucinous carcinoma       151       166       167       173         1.2: Mucinous carcinoma       151       166       167       173         1.2: Mucinous carcinoma       110       123       142       173         1.3: Endometrioid carcinoma       33       61       55       83         1.4: Clear cell carcinoma       33       61       55       83         1.4: Clear cell carcinoma       33       61       55       83         1.5: Adenocarcinoma NOS       432       490       591       692         1.5: Other specified carcinoma       18       28       21       40         1.7: Unspecified carcinoma       166       226       272       400         2: Sex cord-stromal turnours       9       4       8       19         3: Germ cell turnours       9       4       8       19         4: Other specified malignant neoplasm       165       216       130	-2006 1982-1987 19	88–1993 1994–1999	2000-2006
1.1: Serous carcinoma       386       622       862       1,161         1.2: Mucinous carcinoma       151       166       167       173         1.2: Mucinous carcinoma       151       166       167       173         1.3: Endometrioid carcinoma       110       123       142       173         1.4: Clear cell carcinoma       33       61       55       83         1.5: Adenocarcinoma NOS       432       490       591       692         1.5: Adenocarcinoma NOS       18       28       21       40         1.6: Other specified carcinoma       18       28       21       40         1.7: Unspecified carcinoma       166       226       272       400         2: Sex cord-stromal tumours       18       19       23       11         3: Germ cell tumours       9       4       8       11         4: Other specified malignant neoplasm       165       166       233       130	2,683 84.8	86.8 86.8	81.4
1.2: Mucinous carcinoma       151       166       167       173         1.3: Endometrioid carcinoma       110       123       142       134         1.4: Clear cell carcinoma       33       61       55       134         1.4: Clear cell carcinoma       33       61       55       83         1.5: Adenocarcinoma NOS       432       490       591       692         1.6: Other specified carcinoma       18       28       21       40         1.6: Other specified carcinoma       166       226       272       400         1.7: Unspecified carcinoma       166       226       272       400         2: Sex cord-stromal tumours       18       19       23       11         3: Germ cell tumours       9       4       98       130         4: Other specified malignant neoplasm       165       223       454	1,161 25.3	31.5 31.5	35.2
1.3: Endometrioid carcinoma       10       123       142       134         1.4: Clear cell carcinoma       33       61       55       83         1.5: Adenocarcinoma NOS       432       490       591       692         1.5: Adenocarcinoma NOS       18       28       21       40         1.6: Other specified carcinoma       18       28       21       40         1.7: Unspecified carcinoma       166       226       272       400         2: Sex cord-stromal tumours       18       19       23       11         3: Germ cell tumours       9       4       98       19         4: Other specified malignant neoplasm       16       23       130         5: Unspecified malignant neoplasm       16       23       16       130	173 9.9	8.4 8.4	5.3
1.4: Clear cell carcinoma       33       61       55       83         1.5: Adenocarcinoma NOS       432       490       591       692         1.6: Other specified carcinoma       18       28       21       40         1.7: Unspecified carcinoma       18       28       27       40         1.7: Unspecified carcinoma       166       226       272       400         2: Sex cord-stromal tumours       18       19       23       11         3: Germ cell tumours       18       19       23       11         4: Other specified malignant neoplasm       43       71       98       130         5: Unspecified malignant neoplasm       162       166       223       454	134 7.2	6.2 6.2	4.1
1.5: Adenocarcinoma NOS       432       490       591       692         1.6: Other specified carcinoma       18       28       21       40         1.7: Unspecified carcinoma       166       226       272       400         2: Sex cord-stromal tumours       18       19       23       11         3: Germ cell tumours       9       4       98       19         4: Other specified malignant neoplasm       16       23       130         5: Unspecified malignant neoplasm       162       166       223       454	83 2.2	3.1 3.1	2.5
1.6: Other specified carcinoma       18       28       21       40         1.7: Unspecified carcinoma       166       226       272       400         2: Sex cord-stromal tumours       18       19       23       11         3: Germ cell tumours       9       4       8       19       19         4: Other specified malignant neoplasm       43       71       98       130         5: Unspecified malignant neoplasm       162       166       223       454	692 28.3	24.8 24.8	21.0
1.7: Unspecified carcinoma       166       226       272       400         2: Sex cord-stromal tumours       18       19       23       11         3: Germ cell tumours       9       4       8       19         4: Other specified malignant neoplasm       43       71       98       130         5: Unspecified malignant neoplasm       16       223       454	40 1.2	1.4 1.4	1.2
2: Sex cord-stromal tumours       18       19       23       11         3: Germ cell tumours       9       4       8       19         4: Other specified malignant neoplasm       43       71       98       130         5: Unspecified malignant neoplasm       162       166       223       454	400 10.9	11.4 11.4	12.1
3: Germ cell tumours       9       4       8       19         4: Other specified malignant neoplasm       43       71       98       130         5: Unspecified malignant neoplasm       162       166       223       454	11 1.2	1.0 1.0	0.3
4: Other specified malignant neoplasm4371981305: Unspecified malignant neoplasm162166223454	19 0.6	0.2 0.2	0.6
5: Unspecified malignant neoplasm 162 166 223 454	130 2.8	3.6 3.6	3.9
	454 10.6	8.4 8.4	13.8
Total 1,528 1,976 2,462 3,297	3,297 100.0	100.0 100.0	100.0

Table D2.9: Incidence of ovarian cancer b	y remoteness area, 2002–2006
-------------------------------------------	------------------------------

Remoteness area ^(a)	Annual average number of cases ^(b)	Total number of cases	Age-standardised rate ^(c)	95% confidence interval
Major cities	842	4,211	11.4	11.0–11.7
Inner regional	244	1,218	10.4	9.8–11.0
Outer regional	105	527	10.3	9.4–11.2
Remote and very remote	22	109	12.2	9.9–14.7
Not stated	1	4		
Total	1,214	6,068	11.1	10.8–11.3

(a) Classified according to the Australian Standard Geographical Classification (ASGC) Remoteness Areas (see Appendix A).

(b) Numbers may not sum to the total due to rounding.

(c) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females. The rates are based on the total number of cases over the 5-year period from 2002–2006.

Source: Australian Cancer Database, AIHW.

Socioeconomic status ^(a)	Annual average number of cases ^(b)	Total number of cases	Age-standardised rate ^(c)	95% confidence interval
1 (lowest)	254	1,269	11.6	10.9–12.2
2	241	1,204	10.5	9.9–11.1
3	238	1,189	11.2	10.5–11.8
4	226	1,128	10.8	10.2–11.4
5 (highest)	253	1,266	11.3	10.7–11.9
Not stated	2	12		
Total	1,214	6,068	11.1	10.8–11.3

(a) Classified using the ABS Index of Relative Socio-economic Disadvantage (see Appendix A).

(b) Numbers may not sum to the total due to rounding.

(c) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females. The rates are based on the total number of cases over the 5-year period from 2002–2006.

Source: Australian Cancer Database, AIHW.

Country/region of birth ^(a)	Annual average number of cases ^(b)	Total number of cases	Age-standardised rate ^(c,d)	95% confidence interval ^(c)
Americas	14	70	13.7	10.7–17.3
Sub-Saharan Africa	11	55	12.1	9.1–15.8
Southern and Central Asia	15	74	11.8	9.2–14.8
North-West Europe	163	814	11.2	10.4–12.0
Oceania and Antarctica, excl. Australia	26	130	10.7	8.9–12.7
South-East Asia	32	161	10.6	9.1–12.4
North Africa and the Middle East	15	75	10.3	8.1–12.9
Australia	767	3,833	9.9	
Southern and Eastern Europe	92	461	9.6	8.7–10.5
North-East Asia	18	92	9.3	7.5–11.4
Not stated	61	303		
Total	1,214	6,068	10.7	

#### Table D2.11: Incidence of ovarian cancer by country/region of birth, 2002-2006

(a) Classified according to the Standard Australian Classification of Countries, second edition (see Appendix A).

(b) Numbers may not sum to the total due to rounding.

(c) Indirectly age-standardised to the 2002–2006 Australian-born population (see Appendix B).

(d) The rates are expressed per 100,000 females and based on the total number of cases over the 5-year period from 2002–2006. Countries/regions of birth are ordered in descending order according to the age-standardised rate.

Source: Australian Cancer Database, AIHW.

Country or region	Estimated number of cases	Age-standardised rate ^(c)	95% confidence interval ^(d)
Northern Europe	10.531	13.3	13.0–13.6
New Zealand	320	12.4	11.0-13.8
Western Europe	17 650	11.3	11 1_11 5
Northern America	25 162	10.7	10.6-10.8
	23,102	10.7	10.1 10.3
Southorn Europa	11 640	0.7	0.5 0.0
	1 225	9.7	9.5-9.9
	1,235	0.9 7 7	<b>0.4-9.4</b>
South America	12,794	1.1	7.6–7.8
Polynesia	18	7.7	4.1–11.3
Central America	4,009	7.2	7.0–7.4
South-Eastern Asia	16,880	7.2	7.1–7.3
World	204,499	6.6	6.6-6.6
Melanesia	153	6.6	5.6–7.6
Micronesia	12	6.0	2.6–9.4
Eastern Africa	4,706	5.8	5.6–6.0
Western Asia	4,058	5.3	5.1–5.5
South-Central Asia	32,559	5.3	5.2–5.4
Southern Africa	1,003	5.2	4.9–5.5
Western Africa	3,601	4.6	4.4-4.8
Caribbean	838	4.3	4.0-4.6
Eastern Asia	30,617	3.7	3.7–3.7
Middle Africa	1,182	3.3	3.1–3.5
Northern Africa	1,892	2.6	2.5–2.7

## Table D2.12: International comparison of estimated incidence of ovarian and related cancers^(a), 2002^(b)

(a) The data pertain to cancers coded in ICD-10 as C56 and C57.0–C57.4.

(b) The data were estimated for 2002 by the International Agency for Research on Cancer (IARC) and are based on data from approximately 3 to 5 years earlier.

(c) The age-standardised rates were standardised by the IARC using the Doll et al. (1966) World Standard Population and are expressed per 100,000 females. Countries or regions are ordered in descending order according to the age-standardised rate.

(d) The confidence intervals are approximations and were calculated by the AIHW (see Appendix B).

Source: Ferlay et al. 2004.

## Additional tables for Chapter 3: Mortality from ovarian cancer

Age group	Number of deaths	Age-specific rate ^(a)	95% confidence interval
<20	1	0.0	0.0–0.2
20–24	1	0.1	0.0–0.8
25–29	1	0.1	0.0–0.8
30–34	2	0.3	0.0–1.0
35–39	6	0.8	0.3–1.7
40–44	14	1.8	1.0–3.0
45–49	35	4.6	3.2–6.5
50–54	39	5.7	4.0–7.8
55–59	83	13.1	10.4–16.2
60–64	69	14.0	10.9–17.7
65–69	95	24.0	19.4–29.4
70–74	107	32.7	26.8–39.5
75–79	124	41.4	34.4–49.3
80–84	112	46.7	38.5–56.3
85+	106	48.7	39.9–58.9
Total ^(b)	795	6.7	6.2–7.2

Table D3.1: Mortality from ovarian cancer by age at death, 2006

(a) Number of deaths per 100,000 females.

(b) The rate shown in this row was age-standardised to the Australian population as at 30 June 2001 and is expressed per 100,000 females.

Source: National Mortality Database, AIHW.

#### Table D3.2: Mortality from ovarian cancer, 1968 to 2006

Year	Number of deaths	Age-standardised rate (A) ^(a)	95% confidence interval	Age-standardised rate (W) ^(b)	95% confidence interval
1968	451	9.1	8.2–10.0	7.1	6.4–7.8
1969	434	8.6	7.8–9.4	6.6	6.0–7.3
1970	433	8.3	7.6–9.2	6.5	5.9–7.2
1971	436	8.1	7.3–8.9	6.2	5.6–6.8
1972	484	8.8	8.0–9.6	6.8	6.2–7.4
1973	501	8.9	8.1–9.7	6.8	6.2–7.4
1974	493	8.6	7.9–9.4	6.7	6.1–7.3
1975	497	8.5	7.7–9.3	6.5	6.0–7.2

(continued)

Voar	Number of	Age-standardised rate (Δ) ^(a)	95% confidence	Age-standardised	95% confidence
1076	513	2 7	70.05		60.72
1970	545	0.7	82.08	6.0	63.75
1977	533	9.0	7.0.04	6.4	5.9.7.0
1978	505	8.7	7.9-9.4	0.4	5.9-7.0
1979	527	0.4	7.7-9.1	0.4	5.0-0.9
1960	559	0.0	0.0-9.5	0.5	5.0-7.1
1901	549	0.4	7.7-9.1	0.4	5.0-0.9
1982	5/5	8.6	7.9–9.3	6.4	5.9-7.0
1983	583	8.5	7.8–9.2	6.4	5.9-6.9
1984	624	8.8	8.1–9.6	6.6	6.1–7.2
1985	549	7.6	7.0–8.3	5.7	5.2–6.2
1986	623	8.5	7.8–9.2	6.3	5.8–6.9
1987	612	8.2	7.5–8.8	6.1	5.6–6.6
1988	590	7.6	7.0–8.3	5.7	5.3–6.2
1989	624	7.9	7.3–8.5	5.8	5.4–6.3
1990	705	8.8	8.1–9.5	6.5	6.0–7.0
1991	696	8.4	7.8–9.0	6.2	5.7–6.7
1992	650	7.7	7.1–8.3	5.7	5.2–6.2
1993	690	7.9	7.3–8.5	5.7	5.3–6.2
1994	715	8.2	7.6–8.9	6.1	5.6–6.6
1995	702	7.8	7.2-8.4	5.6	5.2–6.1
1996	768	8.3	7.8–9.0	6.0	5.6–6.5
1997	729	7.7	7.1–8.3	5.5	5.1–5.9
1998	750	7.7	7.2–8.3	5.5	5.1–5.9
1999	731	7.3	6.8–7.8	5.2	4.8–5.6
2000	780	7.6	7.1–8.1	5.4	5.0–5.8
2001	837	7.9	7.4–8.4	5.6	5.2–6.0
2002	842	7.8	7.3–8.4	5.6	5.2–6.0
2003	781	7.1	6.6–7.6	5.1	4.7–5.5
2004	852	7.6	7.1–8.1	5.4	5.1–5.8
2005	888	7.6	7.1–8.1	5.3	5.0–5.7
2006	795	6.7	6.2–7.2	4.7	4.4–5.1

Table D3.2 (continued): Mortality from ovarian cancer, 1968 to 2006

(a) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females.
 (b) The age-standardised rates were standardised using the WHO 2000 World Standard Population and are expressed per 100,000 females.

Note: The 1968 to 1996 data were adjusted from earlier ICD standards to ICD-10 standards using a factor of 0.98.

	<50 y	ears	50-69	9 years	70+	70+ years		ages
Year	ASR ^(a)	95% CI	ASR ^(a)	95% CI	ASR ^(a)	95% CI	ASR ^(a)	95% CI
1982	1.7	1.3–2.2	20.7	18.4–23.2	36.6	31.8–41.9	8.6	7.9–9.3
1983	1.2	0.9–1.6	22.3	19.9–25.0	36.3	31.6–41.5	8.5	7.8–9.2
1984	1.4	1.1–1.8	21.6	19.2–24.1	39.4	34.6-44.6	8.8	8.1–9.6
1985	0.9	0.6–1.2	20.1	17.8–22.5	33.5	29.2–38.3	7.6	7.0–8.3
1986	1.7	1.3–2.1	18.8	16.6–21.2	39.7	35.0–44.8	8.5	7.8–9.2
1987	1.5	1.2–1.9	20.3	18.0–22.7	34.2	29.9–38.9	8.2	7.5–8.8
1988	1.4	1.1–1.7	18.2	16.1–20.5	34.2	30.0–38.8	7.6	7.0–8.3
1989	1.2	0.9–1.6	18.6	16.5–20.9	37.2	32.9-42.0	7.9	7.3–8.5
1990	1.5	1.2–1.8	19.5	17.3–21.8	43.4	38.7–48.4	8.8	8.1–9.5
1991	1.3	1.0–1.7	18.9	16.8–21.2	41.0	36.6–45.8	8.4	7.8–9.0
1992	1.2	0.9–1.5	18.1	16.1–20.4	36.2	32.1-40.6	7.7	7.1–8.3
1993	1.0	0.7–1.2	17.5	15.6–19.7	41.3	37.0–46.0	7.9	7.3–8.5
1994	1.3	1.0–1.6	19.1	17.0–21.4	39.1	35.0–43.6	8.2	7.6–8.9
1995	1.2	0.9–1.5	16.8	14.8–18.8	40.4	36.3-44.9	7.8	7.2–8.4
1996	1.2	0.9–1.4	17.7	15.7–19.9	44.7	40.4–49.4	8.3	7.8–9.0
1997	1.0	0.8–1.3	15.6	13.8–17.6	42.9	38.8–47.4	7.7	7.1–8.3
1998	0.8	0.6–1.0	16.8	15.0–18.9	42.4	38.4–46.8	7.7	7.2–8.3
1999	0.9	0.7–1.2	14.6	12.9–16.5	41.5	37.6–45.8	7.3	6.8–7.8
2000	0.9	0.7–1.2	15.7	14.0–17.6	42.5	38.5–46.7	7.6	7.1–8.1
2001	0.9	0.7–1.2	15.5	13.8–17.4	46.5	42.4–50.9	7.9	7.4–8.4
2002	0.9	0.7–1.1	16.8	15.0–18.7	43.1	39.2–47.3	7.8	7.3–8.4
2003	1.0	0.8–1.3	14.4	12.8–16.1	39.0	35.3–42.9	7.1	6.6–7.6
2004	0.8	0.6–1.0	16.0	14.3–17.8	43.0	39.1–47.1	7.6	7.1–8.1
2005	0.9	0.7–1.1	14.6	13.1–16.3	45.2	41.3–49.4	7.6	7.1–8.1
2006	0.8	0.6–1.0	12.7	11.3–14.3	40.3	36.6-44.3	6.7	6.2–7.2

Table D3.3: Mortality from ovarian cancer by age at death, 1982 to 2006

(a) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females. The 1982 to 1996 data were adjusted from ICD-9 to ICD-10 standards using a factor of 0.98.

#### Table D3.4: Mortality from ovarian cancer by remoteness area, 2002-2006

Remoteness area ^(a)	Average annual number of deaths ^(b)	Total number of cases	Age-standardised rate ^(c)	95% confidence interval
Major cities	549	2,744	7.2	6.9–7.4
Inner regional	185	926	7.6	7.1–8.1
Outer regional	82	408	7.8	7.1–8.6
Remote and very remote	15	76	9.6	7.5–12.0
Not stated	1	4		
Total	832	4,158	7.3	7.1–7.6

(a) Classified using the ABS Australian Standard Geographical Classification (ASGC) Remoteness Areas (see Appendix A).

(b) Numbers may not sum to the total due to rounding.

(c) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females. The rates are based on the total number of cases over the 5-year period from 2002–2006.

Source: National Mortality Database, AIHW.

## Table D3.5: Mortality from ovarian cancer by socioeconomic status, 2002–2006 Average annual Total number of cases Age-standardised 95%

Socioeconomic status ^(a)	number of deaths ^(b)	of cases	Age-standardised rate ^(c)	95% confidence interval
1 (Lowest)	164	819	7.2	6.7–7.7
2	185	926	7.7	7.2–8.2
3	156	779	7.1	6.6–7.6
4	143	716	6.8	6.3–7.3
5 (Highest)	180	899	7.8	7.3–8.3
Not stated	4	19		
Total	832	4,158	7.3	7.1–7.6

(a) Classified using the ABS Index of Socio-economic Disadvantage (see Appendix A).

(b) Numbers may not sum to the total due to rounding.

(c) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 100,000 females. The rates are based on the total number of cases over the 5-year period from 2002–2006.

Country/region of birth ^(a)	Annual average number of deaths ^(b)	Total number of cases	Age- standardised rate ^(c,d)	95% confidence interval ^(c)
Sub-Saharan Africa	8	42	10.6	7.6–14.3
North-West Europe	128	640	8.4	7.7–9.0
North Africa and the Middle East	10	50	7.4	5.5–9.8
Southern and Central Asia	9	43	7.4	5.3–9.9
Southern and Eastern Europe	75	377	7.3	6.6–8.1
Australia	553	2,767	7.2	
Oceania and Antarctica, excl. Australia	15	74	7.2	5.7–9.0
North-East Asia	10	52	6.0	4.5–7.8
South-East Asia	15	74	6.0	4.7–7.5
Americas	5	26	5.9	3.8–8.6
Inadequately described, not stated or unknown	3	13		
Total	832	4,158	7.3	

#### Table D3.6: Mortality from ovarian cancer by country/region of birth, 2002-2006

(a) Classified according to the Standard Australian Classification of Countries, second edition (see Appendix A).

(b) Numbers may not sum to the total due to rounding.

(c) Indirectly age-standardised to the 2002–2006 Australian-born population (see Appendix B).

(d) The rates are expressed per 100,000 females and based on the total number of cases over the 5-year period from 2002–2006. Countries/regions of birth are ordered in descending order according to the age-standardised rate.
Region or country	Estimated number of deaths	Age-standardised rate ^(c)	95% confidence interval ^(d)
Northern Europe	7,188	7.9	7.7–8.1
New Zealand	184	6.4	5.5–7.3
Western Europe	12,162	6.3	6.2–6.4
Northern America	16,005	6.1	6.0–6.2
Central and Eastern Europe	15,243	6.0	5.9–6.1
Australia	772	4.9	4.6–5.2
Southern Europe	6,431	4.5	4.4–4.6
Polynesia	11	4.4	1.8–7.0
Eastern Africa	3,340	4.1	4.0-4.2
South-Eastern Asia	9,262	4.1	4.0-4.2
World	124,860	4.0	4.0-4.0
Melanesia	87	3.9	3.1–4.7
South-Central Asia	22,813	3.8	3.8–3.8
South America	6,108	3.7	3.6–3.8
Central America	1,901	3.6	3.4–3.8
Micronesia	6	3.5	0.7–6.3
Western Asia	2,484	3.4	3.3–3.5
Western Africa	2,551	3.2	3.1–3.3
Southern Africa	612	3.2	2.9–3.5
Caribbean	496	2.6	2.4–2.8
Middle Africa	841	2.3	2.1–2.5
Northern Africa	1,343	1.8	1.7–1.9
Eastern Asia	15,019	1.8	1.8–1.8

Table D3.7: International comparison of estimated mortality from ovarian and related can	cers ^(a) ,
2002 ^(b)	

(a) The data pertain to cancers coded in ICD-10 as C56 and C57.0–C57.4.

(b) The data were estimated for 2002 by the International Agency for Research on Cancer (IARC) and are based on data from approximately 3 to 5 years earlier

(c) The age-standardised rates were standardised by the IARC using the Doll et al. (1966) World Standard Population and are expressed per 100,000 females. Countries or regions are ordered in descending order according to the age-standardised rate.

(d) The confidence intervals are approximations and were calculated by the AIHW (see Appendix B).

Source: Ferlay et al. 2004.

# Additional tables for Chapter 4: Survival after a diagnosis of ovarian cancer

	1-year relativ	ve survival	5-year relat	ive survival
Cancer type (ICD-10 codes)	RS (%)	95% CI	RS (%)	95% CI
Breast (C50)	97.2	97.1–97.4	87.8	87.5–88.1
Bowel (C18–C20)	80.0	79.5–80.4	62.4	61.8–63.1
Melanoma of skin (C4)	98.4	98.2–98.6	94.1	93.6 –94.6
Lung (C33–C34)	38.8	38.1–39.4	14.0	13.4 –14.5
Uterus, body (C54)	92.6	92.1–93.2	82.1	81.1–83.0
Non-Hodgkin lymphoma (C82–C85, C96)	77.5	76.7–78.3	62.6	61.5–63.6
Unknown primary site (C26, C39, C76, C80)	15.8	15.2–16.4	7.6	7.1–8.0
Ovary (C56)	73.2	72.2–74.2	39.8	38.6–41.0
Thyroid (C73)	96.9	96.4–97.3	95.3	94.5–96.0
Leukaemia (C91–C95)	65.7	64.6–66.8	47.3	46.0-48.6
All cancers ^(b)	78.7	78.5–78.8	64.1	63.9–64.3

Table D4.1: Relative survival, 10 most commonly diagnosed cancers^(a), females, 1998–2004

(a) Determined by most commonly diagnosed cancers in 2004 and ordered accordingly. Excludes non-melanoma skin cancer (C44).

(b) Includes cancers coded in ICD-10 as C00–C97 (except for C44), D45, D46, D47.1 and D47.3.

Source: AIHW, CA & AACR 2008; Australian Cancer Database, AIHW.

Age at diagnosis		1-year re	lative survival	5-year relative survival		
(years)	Number of cases ^(a)	RS (%)	95% CI	RS (%)	95% CI	
<30	240	95.9	92.4–97.8	86.4	81.0–90.3	
30–39	321	91.1	87.4–93.7	70.6	64.8–75.6	
40–49	938	91.5	89.5–93.1	61.0	57.5–64.4	
50–59	1,712	88.7	87.1–90.2	49.8	47.1–52.4	
60–69	1,812	81.4	79.5–83.1	41.1	38.5–43.6	
70–79	1,842	60.4	58.2–62.6	23.5	21.3–25.7	
80+	1,350	35.8	33.4–38.2	14.8	12.5–17.3	
All ages	8,215	73.7	72.7–74.7	40.0	38.8-41.2	

#### Table D4.2: Relative survival by age at diagnosis, ovarian cancer, 2000–2006

(a) Equals the total number of cases diagnosed in the period considered.

Source: Australian Cancer Database, AIHW.

Voars after	1982	2–1987	1988	-1993	1994	<b>1</b> –1999	2000	-2006
diagnosis	RS (%)	95% CI	RS (%)	95% CI	RS (%)	95% CI	RS (%)	95% CI
1	63.1	61.8–64.4	67.4	66.2–68.6	72.0	70.9–73.1	73.7	72.7–74.7
2	46.1	44.7–47.5	51.9	50.6–53.2	57.4	56.1–58.6	60.0	58.9–61.0
3	38.6	37.2–40.0	43.8	42.5–45.1	47.9	46.6–49.1	51.0	49.8–52.1
4	34.8	33.5–36.2	39.5	38.2–40.8	42.0	40.8–43.3	44.3	43.1–45.4
5	33.0	31.6–34.3	36.8	35.5–38.1	38.8	37.5–40.0	40.0	38.8–41.2
6	31.5	30.1–32.8	34.6	33.3–35.9	36.3	35.1–37.6	37.0	35.7–38.2
7	30.1	28.8–31.5	33.5	32.2–34.7	34.7	33.4–35.9	34.8	33.5–36.2
8	29.3	28.0-30.7	32.4	31.1–33.7	33.3	32.1–34.5	32.8	31.3–34.3
9	28.6	27.2–29.9	31.5	30.3–32.8	32.3	31.1–33.6	31.2	29.3–33.1
10	28.2	26.9–29.6	30.6	29.3–31.9	31.6	30.4–32.9		
11	27.9	26.5–29.2	30.2	28.9–31.5	31.1	29.9–32.4		
12	27.2	25.9–28.6	29.8	28.5–31.1	30.5	29.2–31.8		
13	27.2	25.8–28.6	29.3	28.0-30.6	30.3	29.0–31.7		
14	26.9	25.5–28.3	29.0	27.7–30.4	29.8	28.4–31.2		
15	26.5	25.2–28.0	28.6	27.3–29.9	29.1	27.4–30.8		
16	26.5	25.1–27.9	28.2	26.9–29.6				
17	26.2	24.8–27.7	27.9	26.6–29.3				
18	25.8	24.4–27.3	27.7	26.3–29.1				
19	25.9	24.4–27.4	27.6	26.1–29.0				
20	26.0	24.5–27.5	27.1	25.5–28.6				
21	25.7	24.2–27.2	26.7	24.8–28.7				
22	25.7	24.2–27.3						
23	25.8	24.2–27.4						
24	25.9	24.3–27.6						
25	26.0	24.4–27.8						
26	26.1	24.3-28.0						
27	26.2	24.1–28.4						

Table D4.3: Relative survival by period of diagnosis, ovarian cancer, 1982–1987 to 2000–2006

Source: Australian Cancer Database, AIHW.

		1982–198	7		1988–1993			1994–1995			2000-2006	
Age at diagnosis (years)	Number of cases ^(a)	Relative survival (%)	95% confidence interval									
<30	192	81.4	75.1–86.2	226	85.6	80.3-89.6	220	85.6	80.2-89.6	240	86.4	81.0-90.3
30–39	281	66.2	60.3-71.5	329	71.0	65.7–75.6	298	70.3	64.8-75.2	321	70.6	64.8-75.6
40-49	627	44.7	40.8-48.6	771	51.3	47.6–54.8	770	59.0	55.4-62.4	938	61.0	57.5-64.4
50-59	1,185	35.6	32.9–38.4	1,120	44.2	41.2-47.1	1,338	45.5	42.8-48.2	1,712	49.8	47.1–52.4
60-69	1,383	27.0	24.6–29.4	1,602	29.2	26.9–31.5	1,434	37.1	34.5–39.7	1,812	41.1	38.5-43.6
70–79	1,027	18.4	15.9–21.0	1,307	22.7	20.3–25.2	1,542	24.2	22.0–26.6	1,842	23.5	21.3–25.7
80+	419	14.7	10.8–19.3	609	13.4	10.4–16.9	863	11.1	8.8-13.7	1,350	14.8	12.5–17.3
All ages	5,114	33.0	31.6–34.3	5,964	36.8	35.5-38.1	6,465	38.8	37.5-40.0	8,215	40.0	38.8-41.2
(a) Equals the tot	al number of di	agnosed case.	s in the period consi	dered.								
<i>Source:</i> Australian C	Cancer Databas	e, AIHW.										

Table D4.4: Five-year relative survival by age at diagnosis, ovarian cancer, 1982-1987 to 2000-2006

	v	50 years		-20-	69 yea	rs	20	)+ year:	S	4	ll ages	
Type of ovarian cancer ^(a)	Number of cases ^(b)	RS (%)	95% CI	Number of cases ^(b)	RS (%)	95% CI	Number of cases ^(b)	RS (%)	95% CI	Number of cases ^(b)	RS (%)	95% CI
1: Carcinoma (epithelial tumours)	4,236	58.7	57.1-60.2	10,778	38.5	37.5-39.5	7,669	20.8	19.8–21.9	22,683	36.7	36.1–37.4
1.1: Serous carcinoma	1,747	52.5	50.1-54.9	5,237	34.5	33.2–35.9	3,027	24.0	22.3–25.8	10,011	34.8	33.8–35.8
1.2: Mucinous carcinoma	759	74.4	71.0-77.4	1,067	53.1	49.9–56.1	657	36.9	32.6-41.2	2,483	55.8	53.7-57.9
1.3: Endometrioid carcinoma	637	74.0	70.3-77.4	1,171	63.9	60.9–66.8	509	51.5	46.0–56.9	2,317	64.2	62.0-66.3
1.4: Clear cell carcinoma	337	62.7	57.2-67.8	721	58.8	54.9-62.5	232	49.6	41.7–57.4	1,290	58.3	55.3-61.1
1.5: Adenocarcinoma NOS	512	37.2	33.0-41.4	1,987	19.9	18.2–21.8	2,197	7.3	6.2–8.5	4,696	16.0	14.9–17.1
1.6: Other specified carcinoma	06	49.9	38.8–60.1	158	44.5	36.0–52.7	107	26.1	17.1–36.6	355	40.6	35.0-46.2
1.7: Unspecified carcinoma	154	54.9	46.7–62.4	437	27.9	23.7–32.2	940	8.2	6.5-10.1	1,531	18.4	16.5–20.4
2: Sex cord-stromal tumours	120	84.3	76.2–89.9	159	75.7	67.5–82.3	71	67.0	51.6-80.7	350	77.0	71.6-81.7
3: Germ cell tumours	680	91.1	88.6–93.1	73	76.4	64.0-85.4	40	40.7	23.3–59.4	793	87.5	84.9-89.8
4: Other specified malignant neoplasm	92	47.4	36.8–57.3	371	30.5	25.7–35.5	342	18.5	14.1–23.5	805	27.5	24.3–30.8
5: Unspecified malignant neoplasm	85	72.8	61.8-81.2	205	35.5	29.2-42.0	837	4.0	2.9–5.4	1,127	13.8	12.0–15.8
Total	5,213	63.5	62.1-64.8	11,586	39.0	38.0-39.9	8,959	19.5	18.6-20.5	25,758	37.7	37.1–38.3

Table D4.5: Five-year relative survival by type of ovarian cancer and age at diagnosis, 1982-2006

All cases were coded as primary site, invasive ovarian cancers. Appendix Table D2.5 provides a list of the histology types included in each group. Equals the total number of diagnosed cases in the period considered.

(a)

Source: Australian Cancer Database, AIHW.

Region or country	Mortality: ASR	Incidence: ASR	Mortality-to-incidence ratio
South-Central Asia	3.8	5.3	0.72
Eastern Africa	4.1	5.8	0.71
Middle Africa	2.3	3.3	0.70
Western Africa	3.2	4.6	0.70
Northern Africa	1.8	2.6	0.69
Western Asia	3.4	5.3	0.64
Southern Africa	3.2	5.2	0.62
World	4.0	6.6	0.61
Caribbean	2.6	4.3	0.60
Northern Europe	7.9	13.3	0.59
Melanesia	3.9	6.6	0.59
Central and Eastern Europe	6.0	10.2	0.59
Micronesia	3.5	6.0	0.58
Polynesia	4.4	7.7	0.57
Northern America	6.1	10.7	0.57
South-Eastern Asia	4.1	7.2	0.57
Western Europe	6.3	11.3	0.56
Australia	4.9	8.9	0.55
New Zealand	6.4	12.4	0.52
Central America	3.6	7.2	0.50
Eastern Asia	1.8	3.7	0.49
South America	3.7	7.7	0.48
Southern Europe	4.5	9.7	0.46

Table D4.6: International comparison of mortality-to-incidence ratios for ovarian and related cancers, 2002

Notes

1. The data pertain to cancers coded in ICD-10 as C56 and C57.0–C57.4 (see Appendix B for more details).

2. The mortality and incidence rates were derived from estimates of the number of new ovarian and related cancer cases and deaths for 2002; those estimates were based on data from approximately 3 to 5 years earlier.

3. The age-standardised rates were standardised by the IARC using the Doll et al. (1966) World Standard Population and are expressed per 100,000 females.

4. The mortality-to-incidence ratio equals the age-standardised mortality rate divided by the age-standardised incidence rate.

Source: Ferlay et al. 2004.

# Additional table for Chapter 6: Burden of disease due to ovarian cancer

	Breast cancer	Lung cancer	Bowel cancer	Pancreas cancer	Ovarian and related cancers	Total from all cancers	Ovarian and related cancers
(years)		Number (	Disability-ac	ljusted life yea	rs (DALYs))		cancers
<1	0	0	0	1	0	99	0.0
1–4	0	0	0	0	0	417	0.0
5–9	0	0	1	0	2	605	0.3
10–14	0	1	1	0	8	456	1.8
15–19	0	0	11	0	70	804	8.7
20–24	25	30	41	0	94	1,122	8.4
25–29	260	33	78	2	101	2,116	4.8
30–34	1,132	90	370	3	278	3,855	7.2
35–39	2,680	208	473	35	308	6,459	4.8
40–44	4,971	1,042	963	422	465	11,651	4.0
45–49	6,878	1,553	1,671	426	898	16,642	5.4
50–54	8,382	3,031	2,240	659	1,089	22,157	4.9
55–59	9,135	3,926	2,533	1,186	1,619	26,676	6.1
60–64	7,618	4,965	3,364	1,439	1,672	28,003	6.0
65–69	5,835	4,813	3,599	1,711	1,395	27,127	5.1
70–74	4,610	5,125	3,913	1,312	1,236	26,702	4.6
75–79	4,102	4,890	4,028	1,821	1,492	26,951	5.5
80–84	2,686	2,735	3,112	1,267	801	18,896	4.2
85–89	1,524	1,121	1,779	728	338	10,140	3.3
90–94	554	271	654	200	108	3,445	3.1
95–99	115	36	124	33	19	653	2.9
100+	14	5	7	1	0	58	0.0
All ages ^(a)	60,520	33,876	28,962	11,246	11,994	235,034	5.1

Table D6.1: Leading cancer causes of burden of disease by age group, females, 2003

(a) Values may not sum to the total due to rounding.

Source: Begg et al. 2007b.

# Additional tables for Chapter 7: Hospitalisations for ovarian cancer

Age group (years)	Number of hospitalisations	Age-specific rate ^(a)	95% confidence interval
<20	120	0.04	0.04–0.05
20–24	139	0.19	0.16–0.22
25–29	37	0.05	0.04–0.07
30–34	180	0.24	0.21–0.28
35–39	229	0.29	0.25–0.33
40–44	449	0.59	0.53–0.64
45–49	806	1.04	0.97–1.11
50–54	1,441	2.04	1.94–2.15
55–59	2,177	3.39	3.25–3.54
60–64	2,384	4.35	4.18–4.53
65–69	2,208	5.33	5.11–5.56
70–74	1,635	4.82	4.59–5.06
75–79	1,266	4.26	4.02-4.50
80–84	846	3.47	3.24–3.71
85+	360	1.53	1.38–1.70
Total ^(b)	14,277	1.22	1.20–1.24

Table D7.1: Hospitalisations	s for ovarian o	cancer by age group,	2007-08
------------------------------	-----------------	----------------------	---------

(a) Number of cases per 1,000 females.

(b) The rate shown in this row was age-standardised to the Australian population as at 30 June 2001; it is expressed per 1,000 females.

Source: National Hospital Morbidity Database, AIHW.

Table D7.2: Hospitalisations	for ovarian cancer	by same-day and	overnight status,	1999–00 to
2007-08				

	Same-da	y hospitali	sations	Overnigh	t hospitali	sations	Total h	ospitalis	ations
Year	Number	ASR ^(a)	95% CI	Number	ASR ^(a)	95% CI	Number	ASR ^(a)	95% CI
1999–00	7,434	0.77	0.76–0.79	3,170	0.32	0.31–0.34	10,604	1.10	1.08–1.12
2000–01	7,913	0.80	0.78–0.82	3,063	0.31	0.30-0.32	10,976	1.11	1.09–1.13
2001–02	9,095	0.90	0.88–0.92	2,978	0.29	0.28–0.30	12,073	1.19	1.17–1.21
2002–03	8,794	0.85	0.84–0.87	2,646	0.25	0.24–0.26	11,440	1.11	1.09–1.13
2003–04	9,407	0.89	0.87–0.91	2,710	0.25	0.24-0.26	12,117	1.14	1.12–1.16
2004–05	9,955	0.92	0.90–0.94	2,631	0.24	0.23–0.25	12,586	1.16	1.14–1.18
2005–06	10,317	0.93	0.91–0.95	2,942	0.26	0.25–0.27	13,259	1.19	1.17–1.22
2006–07	10,940	0.96	0.94–0.98	3,132	0.27	0.26-0.28	14,072	1.23	1.21–1.26
2007–08	11,296	0.97	0.95–0.99	2,981	0.25	0.25–0.26	14,277	1.22	1.20–1.24

(a) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 1,000 females.

Source: National Hospital Morbidity Database, AIHW.

	<50	years	50-69	9 years	70+	years	All	ages
Year	ASR ^(a)	95% CI						
1999–00	0.28	0.27–0.29	3.10	3.02–3.19	3.19	3.08–3.31	1.10	1.08–1.12
2000–01	0.28	0.27–0.29	2.97	2.89-3.05	3.59	3.48–3.72	1.11	1.09–1.13
2001–02	0.26	0.25–0.27	3.51	3.43-3.60	3.52	3.40-3.64	1.19	1.17–1.21
2002–03	0.26	0.25–0.27	3.21	3.13–3.29	3.20	3.09–3.31	1.11	1.09–1.13
2003–04	0.26	0.25–0.28	3.26	3.18–3.34	3.50	3.38–3.62	1.14	1.12–1.16
2004–05	0.30	0.28–0.31	3.23	3.15–3.31	3.51	3.39–3.63	1.16	1.14–1.18
2005–06	0.29	0.28–0.30	3.34	3.26-3.42	3.67	3.55–3.79	1.19	1.17–1.22
2006–07	0.27	0.26-0.29	3.55	3.47-3.63	3.79	3.67–3.91	1.23	1.21–1.26
2007–08	0.26	0.25–0.27	3.49	3.41–3.56	3.90	3.78-4.02	1.22	1.20–1.24

Table D7.3: Hospitalisations for ovarian cancer by age group, 1999-00 to 2007-08

(a) The age-standardised rates were standardised to the Australian population as at 30 June 2001 and are expressed per 1,000 females.

Source: National Hospital Morbidity Database, AIHW.

Table D7.4: Average length of stay (ALOS) for ovarian cancer-related hospitalisation	s by same-day
and overnight status, 1999–00 to 2007–08	

Year	ALOS of same-day hospitalisations (days)	ALOS of overnight hospitalisations (days)	Total ALOS (days)
1999–00	1.0	7.3	2.9
2000–01	1.0	7.5	2.8
2001–02	1.0	8.2	2.8
2002–03	1.0	8.3	2.7
2003–04	1.0	8.7	2.7
2004–05	1.0	8.8	2.6
2005–06	1.0	7.8	2.5
2006–07	1.0	7.7	2.5
2007–08	1.0	7.6	2.4

Source: National Hospital Morbidity Database, AIHW.

# Additional table for Chapter 8: Expenditure on ovarian cancer

Table D8.1: Hospital admitted patient services expenditure and number of hospitalisations for ovarian and related cancers by age group, 2004–05

Age group	Hospital admitted	patient expenditure	Number of admitted	Expenditure per	
(years)	(\$ million)	Per cent	patient hospitalisations	hospitalisation (\$)	
<35	1	3.6	131	6,860	
35–44	2	6.0	210	7,237	
45–54	4	16.2	546	7,506	
54–64	6	23.4	872	6,790	
65–74	6	23.4	841	7,029	
75–84	5	20.8	588	8,945	
85+	2	6.5	159	10,379	
Total	25	100.0	3,347	7,547	

*Note:* Data pertain to those hospitalisations for which the principal diagnosis was ovarian and related cancers (i.e. codes of C56 and C57.0–C57.4 in ICD-10). Does not pertain to hospitalisations for which ovarian and related cancers was an additional diagnosis, with the principal diagnosis relating specifically to the type of cancer treatment or care received.

Source: Disease Expenditure Database, AIHW.

### **Appendix E: Stage at diagnosis**

A number of staging systems are used to classify ovarian cancers. These systems are described in this appendix.

#### FIGO staging system

The International Federation of Gynecology and Obstetrics (FIGO) system is the conventional measure used to stage ovarian cancer, as well as other types of gynaecological cancers, with staging decisions mainly based on findings from surgical exploration. Ovarian tumours are given a value from I to IV, with each of the first three stages divided further into three sub-stages (Odicino et al. 2008; Pecorelli et al. 2000). Table E.1 provides a description of each of these stages and sub-stages.

Stage	Description
Stage I	Growth limited to the ovaries
la	Growth limited to one ovary: no ascites present containing malignant cells. No tumour on the external surface; capsule intact
lb	Growth limited to both ovaries: no ascites present containing malignant cells. No tumour on the external surfaces; capsules intact
Ic ^(a)	Tumour either Stage Ia or Ib, but with tumour on surface of one or both ovaries, or with capsule ruptured, or with ascites present containing malignant cells, or with positive peritoneal washings
Stage II	Growth involving one or both ovaries with pelvic extension
lla	Extension and/or metastases to the uterus and/or tubes
llb	Extension to other pelvic tissues
llc	Tumour either Stage IIa or IIb, but with tumour on surface of one or both ovaries, or with capsule(s) ruptured, or with ascites present containing malignant cells, or with positive peritoneal washings
Stage III	Tumour involving one or both ovaries with histologically-confirmed peritoneal implants outside the pelvis and/or positive retroperitoneal or inguinal nodes. Superficial liver metastases equals Stage III. Tumour is limited to the true pelvis, but with histologically-proven malignant extension to small bowel or omentum
Illa	Tumour grossly limited to the true pelvis, with negative nodes, but with histologically-confirmed microscopic seeding of abdominal peritoneal surfaces, or histologic proven extension to small bowel or mesentery
IIIb	Tumour of one or both ovaries with histologically confirmed implants, peritoneal metastasis of abdominal peritoneal surfaces, none exceeding 2 cm in diameter: nodes are negative
IIIc	Peritoneal metastasis beyond the pelvis N2 cm in diameter and/or positive retroperitoneal or inguinal nodes
Stage IV	Growth involving one or both ovaries with distant metastases. If pleural effusion is present, there must be positive cytology to allot a case to Stage IV. Parenchymal liver metastasis equals Stage IV
(a) In order	to evaluate the impact on prognosis of the different criteria for allotting cases to Stage Ic or IIc, it would be of value to know if

Table E.1: Carcinoma of the ovary: FIGO nomenclature (Rio de Janeiro 1998)

(a) In order to evaluate the impact on prognosis of the different criteria for allotting cases to Stage Ic or IIc, it would be of value to know if rupture of the capsule was spontaneous or caused by the surgeon, and if the source of malignant cells detected was peritoneal washings or ascites.

Source: Heintz et al. 2006.

### **TNM staging system**

Ovarian cancers can also be staged according to the TNM system which was initially developed by the International Union Against Cancer (UICC). This staging system describes the size of the primary tumour (T), the absence or presence of metastasis to nearby lymph nodes (N) and the absence or presence of distant metastasis (M). The TNM system is considered to be virtually identical to the FIGO system (ACS 2009; Pecorelli et al. 2000). Further information about the TNM staging system can be found on the UICC website (UICC 2009).

### Summary staging system

The Surveillance Epidemiology End Results (SEER) Summary staging system (or 'summary staging system' for short) is a simpler method to stage ovarian cancers. According to Tracey and associates (2009), this summary measure is preferred by a number of cancer registries overseas and in Australia (such as the New South Wales registry) since the required information can be sourced more readily from the pathology and clinical reports to which the registries have access. In this staging system, tumours are allocated to one of three categories, as well as an 'unknown' category, as shown in Table E.2.

Stage	Description
Localised	A malignancy limited to the organ of origin; it has spread no farther than the organ in which it started. There is infiltration past the basement membrane of the epithelium into the functional part of the organ, but there is no spread beyond the boundaries of the organ
Regional	There is tumour extension beyond the limits of the organ of origin. There is invasion through the entire wall of the organ into surrounding organs and/or adjacent issues or by direct extension or contiguous spread to nearby lymph nodes
Distant metastases	Tumour cells that have broken away from the primary tumour, have travelled to other parts of the body and have begun to grow at the new location. Distant stage is also called remote, diffuse, disseminated, metastatic or secondary disease. In most cases there is no continuous trail of tumour cells between the primary site and the distant site
Unknown	There are cases for which sufficient evidence is not available to adequately assign a stage. Examples include occasions when the patient dies before workup is completed, when a patient refuses a diagnostic or treatment procedure, and when there is limited workup due to the patient's age or a simultaneous contraindicating condition. If there is insufficient information the case cannot be assigned a stage

Table E.2:	Summary s	taging systen	n – extent of	disease at	diagnosis
I WOIC LIL	Summing 5	mania oyoten		anocaoe at	anagnoono

Source: Tracey et al. 2006.

## Appendix F: Definition of ovarian cancer– related hospitalisations

Due to the method in which the principal diagnosis for hospitalisations of cancer patients is coded, it is insufficient to simply select those hospitalisations for which ovarian cancer was the principal diagnosis. Most importantly, when a patient receives same-day chemotherapy as a treatment for cancer, the Australian Coding Standards (NCCH 2008a) indicate that the principal diagnosis is to be coded to reflect the fact that the patient received chemotherapy, with the type of cancer listed as an additional diagnosis. The same coding practice is used for a number of other same-day cancer-related interventions – such as the implanting of chemotherapy ports. Hence, the number of hospitalisations would be greatly underestimated if only those for which the principal diagnosis was listed as ovarian cancer (i.e. ICD-10-AM code of C56) were included.

Thus, for the purposes of examining the number of admitted patient separations that arose specifically due to invasive ovarian cancer and were directly related to treatment/care for ovarian cancer, 'ovarian cancer-related hospitalisations' were identified in this report as follows:

- either a *principal* diagnosis of ovarian cancer (ICD-10-AM code of C56)
- **or** an *additional* diagnosis of ovarian cancer (ICD-10-AM code of C56) and a principal diagnosis of one of the following ICD-10-AM 'Z' codes (with these Z codes falling within ICD-10-AM Chapter 21 'Factors influencing health status and contact with health services'):
  - follow-up examination after treatment for malignant neoplasms (Z08)
  - prophylactic immunotherapy (Z29.1)
  - other prophylactic immunotherapy (Z29.2)
  - prophylactic surgery for risk-factors related to malignant neoplasm ovary (Z40.01)
  - adjustment and management of drug delivery or implanted device (Z45.1)
  - adjustment and management of vascular access device (Z45.2)
  - radiotherapy session (Z51.0)
  - pharmacotherapy session for neoplasm (Z51.1)
  - convalescence following radiotherapy (Z54.1)
  - convalescence following chemotherapy (Z54.2).

Using data from the National Hospital Morbidity Database (NHMD) for 2007–08, Table F.1 shows the number of hospitalisations for each of the relevant Z code principal diagnoses, as well as for those hospitalisations in which ovarian cancer was the principal diagnosis.

The number of hospitalisations that pertain to each of the inclusions in the definition of ovarian cancer-related hospitalisations is shown in Table F.1. The principal diagnosis was 'ovarian cancer' for one in four (26%) of all ovarian cancer-related hospitalisations. Thus, if one were to define ovarian cancer hospitalisations based solely on this disease being classified as the principal diagnosis, 74% of hospitalisations due to this disease would be missed. For almost two in three (65%) ovarian cancer-related hospitalisations, the principal diagnosis was 'pharmacotherapy session for neoplasm' (e.g. chemotherapy) with ovarian cancer listed as an additional diagnosis.

	Same-day hosp	italisations	Overnight hospi	italisations	Total hospita	lisations
Diagnosis (ICD-10-AM code)	Number	Per cent	Number	Per cent	Number	Per cent
Ovarian cancer as principal diagnosis (C56)	836	7.4	2,938	98.6	3,774	26.4
Ovarian cancer as additional diagnosis (C56) AND principal diagnosis of:						
Follow-up examination after treatment for malignant neoplasms (208)	0	0.0	0	0.0	0	0.0
Prophylactic immunotherapy (Z29.1)	0	0.0	0	0.0	0	0.0
Other prophylactic immunotherapy (Z29.2)	0	0.0	0	0.0	0	0.0
Prophylactic surgery for risk-factors related to malignant neoplasm—ovary (Z40.01)	0	0.0	0	0.0	0	0.0
Adjustment and management of implantable infusion device or pump (Z45.1)	640	5.7	15	0.5	655	4.6
Adjustment and management of vascular access device (245.2)	499	4.4	Q	0.2	505	3.5
Radiotherapy session (Z51.0)	ς	0.0	0	0.0	б	0.0
Pharmacotherapy session for neoplasm (Z51.1)	9,314	82.5	14	0.5	9,328	65.3
Convalescence following radiotherapy (Z54.1)	2	0.0	0	0.0	N	0.0
Convalescence following chemotherapy (Z54.2)	2	0.0	8	0.3	10	0.1
Total ovarian cancer-related hospitalisations	11,296	100.0	2,981	100.0	14,277	100.0
Source: National Hospital Morbidity Database, AIHW.						

Table F.1: Hospitalisations for ovarian cancer by same-day and overnight status, 2007–08

As noted in Chapter 7, not all hospitals in all states and territories formally admit patients for same-day chemotherapy services. Instead, in three states and territories, some patients are provided same-day chemotherapy on an outpatient (or non-admitted patient) basis. Such services are not captured in the NHMD. In particular, during the 1990s, hospitals in New South Wales began to apply this change in admission processes. In addition, hospitalisations data for the Australian Capital Territory from approximately 2003–04 reflect changed admission practices, as do data for South Australia from 2007–08. Thus, the recorded data on this type of admitted patient service is not comparable over time.

To illustrate the effect on the data of this change in admission processes, data on the number of hospitalisations for same-day chemotherapy sessions (referred to as 'pharmacotherapy sessions for neoplasms' in ICD-10-AM) for ovarian cancer are shown for each state and territory over time in Table F.2. While the number of such sessions increased by more than 50% between 1999–00 and 2007–08 in some of the jurisdictions – including Tasmania (186%), the Northern Territory (667%), 95% in Victoria (95%) and Western Australia (76%) – the number of chemotherapy sessions decreased by 8% in New South Wales over the period considered. Furthermore, the drop in the number of such sessions in the Australian Capital Territory is evident from 2004 –05 onwards, as is the more recent drop in South Australia between 2006–07 and 2007–08.

Table F.2: Number of ovarian cancer-related hospitalisations for same-day 'Pharmacotherapy sessions for neoplasm'^(a) by state and territory, 1999–00 to 2007–08

Year	NSW	Vic	Qld	WA	SA	Tas	ACT	NT	Total
1999–00	1,227	1,861	1,867	843	653	63	110	6	6,630
2000–01	1,210	1,878	2,217	779	767	15	197	26	7,089
2001–02	877	2,168	2,594	1,166	789	114	160	84	7,952
2002–03	760	2,701	2,083	1,138	738	124	157	105	7,806
2003–04	938	3,341	2,044	1,010	593	131	211	59	8,327
2004–05	917	3,782	1,943	1,193	737	81	97	60	8,810
2005–06	940	3,582	2,098	1,212	704	250	86	88	8,960
2006–07	980	3,526	2,208	1,256	855	265	101	68	9,259
2007–08	1,133	3,624	2,284	1,486	441	180	120	46	9,314

(a) ICD-10-AM code of Z51.1.

Source: National Hospital Morbidity Database, AIHW.

# Glossary

This section provides a general description of the terms used in this report. The terms have been defined in the context of this report; some terms may have other meanings in other contexts.

Additional diagnosis: a condition or complaint either coexisting with the principal diagnosis or arising during the episode of care.

Administrative databases: observations about events that are routinely recorded or required by law to be recorded. Such events include births, deaths, hospital separations and cancer incidence. Administrative databases include the Australian Cancer Database, the National Mortality Database and the National Hospital Morbidity Database.

Admitted patient: a person who undergoes a hospital's formal admission process to receive treatment and/or care. Such treatment or care can occur in hospital and/or in the person's home (as a 'hospital-in-home' patient).

**Age-specific rate:** a rate for a specific age group. The numerator and denominator relate to the same age group.

**Age-standardisation:** a method of removing the influence of age when comparing populations with different age structures. This is usually necessary because the rates of many diseases vary strongly (usually increasing) with age. The age structures of the different populations are converted to the same 'standard' structure. The disease rates that would have occurred with that structure are then calculated and compared.

Associated cause of death: any other condition or event that was not related to the underlying cause of death but was still considered to contribute to the individual's death.

**Average length of stay:** the average (mean) number of patient days for admitted patient episodes. Patients admitted and separated on the same date are allocated a length of stay of 1 day.

**Benign:** non-cancerous tumours that may grow larger but do not spread to other parts of the body.

**Cancer (malignant neoplasm):** a large range of diseases in which some of the body's cells become defective, begin to multiply out of control, can invade and damage the area around them, and can also spread to other parts of the body to cause further damage.

**Carcinoma:** a cancer that begins in the lining layer (epithelial cells) of organs such as the ovary.

**Confidence interval:** a statistical term describing a range (interval) of values within which we can be 'confident' that the true value lies, usually because it has a 95% or higher chance of doing so.

**Crude rate:** the number of events in a given period divided by the size of the population at risk in a specified time period.

**Crude survival:** the proportion of people alive at a specified point in time subsequent to the diagnosis of ovarian cancer.

**DALYs (disability-adjusted life years):** the sum of years of life lost due to premature mortality (YLL) in the population and the equivalent years of 'healthy' life lost due to disability (YLD).

Death due to cancer: see Mortality due to cancer.

**Heath expenditure:** includes expenditure on health goods and services (e.g. medications, aids and appliances, medical treatment, public health, research) which collectively are termed 'current expenditure' and on health-related investment which is often referred to as 'capital expenditure'.

Hospitalisation: see *Separation*.

**Incidence:** the number of new cases (of an illness or event, and so on) occurring during a given period.

**International Statistical Classification of Diseases and Related Health Problems:** the World Health Organization's internationally accepted classification of death and disease. The tenth revision (ICD-10) is currently in use. ICD-10-AM is the Australian modification of ICD-10; it is used for diagnoses and procedures recorded for patients admitted to hospitals (see Appendix A).

#### Invasive: see Malignant.

**Length of stay:** duration of hospital stay, calculated by subtracting the date the patient was admitted from the day of separation. All leave days, including the day the patient went on leave, are excluded. A same-day patient is allocated a length of stay of 1 day.

**Limited-duration prevalence:** the number of people alive at a specific time who have been diagnosed with ovarian cancer over a specified period (such as the previous 5 or 25 years).

**Malignant:** a tumour with the capacity to spread to surrounding tissue or to other sites in the body.

Metastasis: see Secondary cancer.

**Mortality due to cancer:** the number of deaths which occurred during a specified period (usually a year) for which the underlying cause of death was recorded as cancer.

**Mortality-to-incidence ratio:** the ratio of the age-standardised mortality rate for ovarian cancer to the age-standardised incidence rate for ovarian cancer.

New cancer case: see Incidence.

**Neoplasm:** an abnormal ('neo', new) growth of tissue. Can be 'benign' (not a cancer) or 'malignant' (a cancer). Also known as a tumour.

**Overnight patient:** an admitted patient who receives hospital treatment for a minimum of 1 night (that is, is admitted to, and separates from, hospital on different dates).

**Patient days:** the total number of days for admitted patients who separated during a specified reference period. A same-day patient is allocated a length of stay of 1 day.

**Population estimates:** official population numbers compiled by the Australian Bureau of Statistics at both state and territory, and statistical local area levels by age and sex, as at 30 June each year. These estimates allow comparisons to be made between geographical areas of differing population sizes and age structures (see Appendix C).

**Prevalence (or complete prevalence):** the total number of people alive at a specific date who have ever been diagnosed with a particular disease such as ovarian cancer.

Primary cancer: a tumour that is at the site where it first formed (also see Secondary cancer).

**Principal diagnosis:** the diagnosis established after study to be chiefly responsible for occasioning an episode of admitted patient care.

**Procedure:** a clinical intervention that is surgical in nature, carries a procedural risk, carries an anaesthetic risk, requires specialised training and/or requires special facilities or equipment available only in the acute care setting.

**Relative survival:** the ratio of observed survival of a group of persons diagnosed with ovarian cancer to expected survival of those in the corresponding general population after a specified interval following diagnosis (such as 5 or 10 years).

**Risk factor:** any factor that represents a greater risk of a health disorder or other unwanted condition or event. Some risk factors are regarded as causes of disease, others are not necessarily so. Along with their opposites, namely protective factors, risk factors are known as 'determinants'.

**Same-day patient:** a patient who is admitted to, and separates from, hospital on the same date.

**Secondary cancer:** a tumour that originated from a cancer elsewhere in the body. Also referred to as a metastasis.

**Separation:** An episode of care for an admitted patient which may include a total hospital stay (from admission to discharge, transfer or death) or a portion of a hospital stay that begins or ends in a change of type of care (e.g. from acute to rehabilitation). In this report, separations are also referred to as hospitalisations.

**Statistical significance:** an indication from a statistical test that an observed difference or association may be significant or 'real' because it is unlikely to be due just to chance. A statistical result is usually said to be 'significant' if it would occur by chance only once in twenty times or less often (see Appendix B).

**Stage:** the extent of a cancer in the body. Staging is usually based on the size of the tumour, whether lymph nodes contain cancer, and whether the cancer has spread from the original site to other parts of the body (see Appendix E).

Symptom: any indication of a disorder that is apparent to the person affected.

**Underlying cause of death:** the disease or injury that initiated the sequence of events leading directly to death.

**YLD (years of healthy life lost due to disability):** for each new case, YLD equals the average duration of the disease (to remission or death) multiplied by a severity weight for that disease (which depends upon its disabling effect over the disease duration).

**YLL (years of life lost):** for each new case, YLL equals the number of years between premature death and the standard life expectancy for the individual.

### References

ABS (Australian Bureau of Statistics) 1981. Causes of death, Australia, 1979. ABS cat. no. 3303.0. Canberra: ABS.

ABS 2001. Information paper: ABS views on remoteness. ABS cat. no. 1244.0. Canberra: ABS.

ABS 2004. Experimental estimates and projections, Aboriginal and Torres Strait Islander Australians, 1991 to 2009. ABS cat. no. 3238.0. Canberra: ABS.

ABS 2007. Causes of death, Australia, 2005. ABS cat. no. 3303.0. Canberra: ABS.

ABS 2008a. Information paper: an introduction to Socio-Economic Indexes for Areas (SEIFA) 2006. ABS cat. no. 2039.0. Canberra: ABS.

ABS 2008b. Population projections, Australia 2006 to 2101. ABS cat. no. 3222.0. Canberra: ABS.

ABS 2008c. Standard Australian Classification of Countries (SACC), Australia, 2nd edn. ABS cat. no. 1269.0. Canberra: ABS.

ABS 2009a. A picture of the nation: the statistician's report on the 2006 Census. Canberra: ABS.

ABS 2009b. Life tables, Australia. ABS cat. no. 3302.0.55.001. Canberra: ABS. Viewed 15 December 2009, <www.abs.gov.au>.

ABS 2009c. National Health Survey: summary of results, Australia, 2007–08. ABS cat. no. 4364.0. Canberra: ABS.

ABS 2009d. Causes of death, Australia, 2007. ABS cat. no. 3303.0. Canberra: ABS.

ABS 2009e. Experimental life tables for Aboriginal and Torres Strait Islander Australians, Australia, 2005–2007. ABS cat. no. 3302.0.55.003. Canberra: ABS.

ABS 2009f. Experimental estimates and projections, Aboriginal and Torres Strait Islander Australians, 1986 to 2006. ABS cat. no. 3238.0. Canberra: ABS.

ABS 2009g. Births 2008, Australia. ABS cat. no. 3301.0. Canberra: ABS.

ABS 2009h. Deaths 2007, Australia. ABS cat. no. 3302.0. Canberra: ABS.

ABS & AIHW (Australian Institute of Health and Welfare) 2008. The health and welfare of Australia's Aboriginal and Torres Strait Islander peoples. ABS cat. no. 4704.0 and AIHW cat. no. IHW 21. Canberra: ABS & AIHW.

ACN (Australian Cancer Network) and NBCC (National Breast Cancer Centre) 2004. Clinical practice guidelines for the management of women with epithelial ovarian cancer. Sydney: NBCC.

ACS (American Cancer Society) 2008. Cancer facts and figures 2008. Atlanta: ACS.

ACS 2009. Detailed guide: ovarian cancer. Atlanta: ACS. Viewed 17 September 2009, <www.documents.cancer.org/114.00/114.00.pdf>.

ACT Health 2007. Cancer in the ACT, 1998–2004. Health series no. 42. Canberra: ACT Government.

AIHW 2000. Australian hospital statistics 1998–99. Health services series no. 15. Cat. no. HSE 11. Canberra: AIHW.

AIHW 2004. Rural, regional and remote health: a guide to remoteness classifications. Cat. no. PHE 53. Canberra: AIHW.

AIHW 2005. Health system expenditures on cancer and other neoplasms in Australia, 2000–01. Cat. no. HWE 29. Canberra: AIHW.

AIHW 2008a. Australia's health 2008. Cat. no. AUS 99. Canberra: AIHW.

AIHW 2008b. Health expenditure Australia 2006–07. Health and welfare expenditure series no. 35. Cat. no. HWE 42. Canberra: AIHW.

AIHW 2009a. Australian hospital statistics 2007–08. Health services series no. 33. Cat. no. HSE 71. Canberra: AIHW.

AIHW 2009b. National Cancer Statistics Clearing House protocol 2009. Canberra: AIHW.

AIHW & AACR (Australasian Association of Cancer Registries) 2008. Cancer in Australia: an overview, 2008. Cancer series no. 46. Cat. no. CAN 42. Canberra: AIHW.

AIHW & CA (Cancer Australia) 2008. Non-melanoma skin cancer: general practice consultations, hospitalisation and mortality. Cancer series no. 43. Cat. no. 39. Canberra: AIHW.

AIHW, CA & AACR 2008. Cancer survival and prevalence in Australia: cancers diagnosed from 1982 to 2004. Cancer series no. 42. Cat. no. CAN 38. Canberra: AIHW.

AIHW & NBCC 2006. Ovarian cancer in Australia: an overview, 2006. Cancer series no. 35. Cat. no. CAN 30. Canberra: AIHW.

AIHW & NBOCC (National Breast and Ovarian Cancer Centre) 2009. Breast cancer in Australia: an overview, 2009. Cancer series no. 50. Cat. no. CAN 46. Canberra: AIHW.

AIHW: Mathers C, Vos T & Stevenson C 1999. The burden of disease and injury in Australia. Cat. no. PHE 17. Canberra: AIHW.

Averette HE, Janicek MF & Menck HR 1995. The national cancer data base report on ovarian cancer. Cancer 76:1096–103.

Baade PD, Fritschi L & Aitken JF 2005. Geographical differentials in cancer incidence and survival in Queensland: 1996 to 2002. Brisbane: Viertel Centre for Research in Cancer Control, Queensland Cancer Fund.

Begg S, Vos T, Barker B, Stevenson C, Stanley L & Lopez AD 2007a. The burden of disease and injury in Australia, 2003. Cat. no. PHE 82. Canberra: AIHW.

Begg S, Vos T, Barker B, Stevenson C, Stanley L & Lopez AD 2007b. Annex tables for the burden of disease and injury in Australia, 2003. Viewed 4 September 2009, <www.aihw.gov.au/publications/hwe/bodaiia03/bodaiia03-x02.pdf>.

Black RJ, Sankaranarayanan R & Parkin DM 1998. Interpretation of population-based cancer survival data. IARC Scientific Publication 145:13–7.

Bray F, Loos AH, Tognazzo S & La Vecchia C 2005. Ovarian cancer in Europe: cross-sectional trends in incidence and mortality in 28 countries, 1953–2000. International Journal of Cancer 113:977–90.

Brenner H, Arndt V, Gefeller O & Hakulinen T 2004. An alternative approach to age adjustment of cancer survival rates. European Journal of Cancer 40:2317–22.

Brenner H & Gefeller O 1996. An alternative approach to monitoring cancer patient survival. Cancer 78(9):2004–10.

Brenner H & Hakulinen T 2003. On crude and age-adjusted relative survival rates. Journal of Clinical Epidemiology 56:1185–91.

Breslow N & Day N 1987. Statistical methods in cancer research. Volume II: the design and analysis of cohort studies. Lyon: IARC.

Cancer Council Queensland 2009. Cancer in Queensland: incidence and mortality 1982 to 2006, statistical tables. Spring Hill: Cancer Council Queensland.

Cancer Council SA (South Australia) 2009. Statistics: ovarian cancer. Eastwood: Cancer Council SA.

Cancer Council Victoria 2007. Canstat: ovarian cancer. Carlton: Cancer Council Victoria.

Cancer Research UK 2006. CancerStats: ovarian cancer survival statistics. London: Cancer Research UK. Viewed 4 September 2009,

<www.info.cancerresearchuk.org/cancerstats/types/ovary/survival>.

Chan JK, Urban R, Cheung MK, Osann K, Husain A, Teng NN et al. 2006. Ovarian cancer in younger vs older women: a population-based analysis. British Journal of Cancer 95:1314–20.

Colombo N, Van Gorp T, Parma G, Amant F, Gatta G, Sessa C et al. 2006. Ovarian cancer. Critical Reviews in Oncology/Hematology 60:159–79.

Condon J 2004. Cancer, health services and Indigenous Australians. Aboriginal and Torres Strait Islander Primary Health Care Review. Consultant report no. 5. Canberra: Office for Aboriginal and Torres Strait Islander Health.

Coory M, Thompson A & Ganguly I 2000. Cancer among people living in rural and remote Indigenous communities in Queensland. Medical Journal of Australia 173(6):301–4.

Curado MP, Edwards B, Sin HR, Storm H, Ferlay J, Heanue M & Boyle P (eds) 2007. Cancer incidence in five continents, vol. IX. IARC Scientific Publications no. 160. Lyon: IARC.

Dalton M, Veen A, Albion T, Otahal P & Blizzard L 2008. Cancer in Tasmania: incidence and mortality 2006. Hobart: Menzies Research Institute.

Dickman P 2004. Estimating and modelling relative survival using SAS. Stockholm: Karolinska Institutet. Viewed 8 May 2007, <www.pauldickman.com/rsmodel/sas_colon>.

Dobson AJ, Kuulasmaa K, Eberle E & Scherer J 1991. Confidence intervals for weighted sums of Poisson parameters. Statistics in Medicine 10:457–62.

Doll R, Payne P & Waterhouse J (eds) 1966. Cancer incidence in five continents: a technical report. Berlin: Springer–Verlag (for UICC).

Donnelly DW, Garvin AT & Comber H 2009. Cancer in Ireland 1994–2004: a comprehensive report. Ireland: Northern Ireland Cancer Registry/National Cancer Registry.

Florida Department of Health 2009. Ovarian cancer in Florida. Tallahassee: Bureau of Epidemiology. Viewed 5 September 2009,

<www.doh.state.fl.us/disease_ctrl/epi/cancer/Ovarian_Report.pdf>.

Ferlay J, Bray F, Pisani P & Parkin DM 2004. GLOBOCAN 2002: cancer incidence, mortality and prevalence worldwide. IARC CancerBase no. 5 version 2.0. Lyon: IARC Press.

Fritz A, Percy C, Jack A, Shanmugaratnam K, Sobin L, Parkin DM et al. 2000. International Classification of Diseases for Oncology, 3rd edn. Geneva: World Health Organization.

Grossi M, Quinn MA, Thursfield VJ, Francis PA, Rome RM, Planner RS et al. 2002. Ovarian cancer: patterns of care in Victoria during 1993–1995. Medical Journal of Australia 177(1):11–16.

Heintz APM, Odicino F, Maisonneuve P, Quinn MA, Benedet JL, Creasman WT et al. 2006. Carcinoma of the ovary. International Journal of Gynecology & Obstetrics 95(0):S161–S192.

Horner MJ, Ries LAG, Krapcho M, Neyman N, Aminou R, Howlader N et al. (eds) 2009. SEER cancer statistics review, 1975–2006. Bethesda, MD: National Cancer Institute. Viewed 21 September 2009, <www.seer.cancer.gov/csr/1975_2006/index.html>. IARC (International Agency for Research on Cancer) 2004. International rules for multiple primary cancers (ICD-O 3rd edn). Lyon: IARC. Viewed 8 May 2009, <www.iacr.com.fr/MPrules_july2004.pdf>.

Jensen OM, Parkin DM, MacLennan R, Muir CS & Skeet RG (eds) 1991. Cancer registration: principles and methods. IARC scientific publications no. 95. Lyon: IARC.

Kjaerbye-Thygesen A, Huusom LD, Frederiksen K & Kjaer SK 2005. Trends in the incidence and mortality of ovarian cancer in Denmark 1978–2002: comparison with other Nordic countries. Acta Obstetricia et Gynecologica Scandinavica 84:1006–12.

Kliewer EV & Smith KR 1995. Ovarian cancer mortality among immigrants in Australia and Canada. Cancer Epidemiology, Biomarkers and Prevention 4:453–8.

Kobel M, Kalloger SE, Boyd N, McKinney S, Mehl E, Palmer C et al. 2008. Ovarian carcinoma subtypes are different diseases: implications for biomarker studies. PLoS Medicine 5(12):1749–59.

Kosary CL 2007. Cancer of the ovary. In: Ries LAG, Young JL, Keel GE, Eisner MP, Lin YD & Horner MJ (eds). SEER survival monograph: cancer survival among adults: US SEER Program, 1998–2001, patient and tumour characteristics. NIH pub. no. 07–6215. Bethesda, MD: National Cancer Institute.

Kricker A 2002. Ovarian cancer in Australian women. Camperdown, NSW: NBCC.

Laurvick CL, Semmens JB, Leung YC & Homan CDJ 2003. Ovarian cancer in Western Australia (1982–1998): trends in surgical intervention and relative survival. Gynecologic Oncology 88:136–40.

Le Teuff GL, Abrahamowicz M, Bolard P & Quantin C 2005. Comparison of Cox's and relative survival models when estimating the effects of prognostic factors on disease-specific mortality: a simulation study under proportional excess hazards. Statistics in Medicine 24:3887–909.

Maas HAAM, Kruitwagen RFPM, Lemmens VEPP, Goey SH & Janssen-Heijnen MLG 2005. The influence of age and co-morbidity on treatment and prognosis of ovarian cancer: a population-based study. Gynecologic Oncology 97:104–9.

McMurdo ME, Witham MD & Gillespie ND 2005. Including older people in clinical research. British Medical Journal 331:1036–7.

Menon U & Jacobs IJ 2001. Ovarian cancer screening in the general population: current status. International Journal of Gynecological Cancer 11(Supplement 1):3–6.

Morris CR, Rodriquez AO, Epstein J & Cress RD 2008. Declining trends of epithelial ovarian cancer in California. Gynecologic Oncology 108:207–13.

Murray CJL, Salomon JA & Mathers C. 1999. A critical examination of summary measures of population health. Global programme for evidence for health policy discussion paper series no. 2. Geneva: World Health Organization.

NBOCC (National Breast and Ovarian Cancer Centre) 2009. Population screening and early detection of ovarian cancer in asymptomatic women: NBOCC position statement. Surry Hills, NSW: NBOCC. Viewed 16 January 2010, <a href="https://www.nbocc.org.au/our-organisation/position-statements/population-screening-and-early-detection">https://www.nbocc.org.au/our-organisation/position-statements/population-screening-and-early-detection</a>>.

NCCH (National Centre for Classification in Health) 2006. Australian Classification of Health Interventions, 5th edn. Sydney: University of Sydney.

NCCH 2008a. Australian coding standards for ICD-10-AM and ACHI. Sydney: University of Sydney.

NCCH 2008b. The International Statistical Classification of Diseases and Related Health Problems, 10th revision, Australian modification (ICD-10-AM): tabular list of diseases. Sydney: University of Sydney.

NCCH 2008c. Australian Classification of Health Interventions (ACHI): tabular list of interventions, 6th edn. Sydney: University of Sydney.

National Cancer Institute 2009. Anatomy: the cervix and nearby organs, NCI visuals online. Bethesda, MD: National Institutes of Health, United States Department of Health and Human Services. Viewed 21 August 2009,

<visualsonline.cancer.gov/details.cfm?imageid=4350>.

NZ (New Zealand) Ministry of Health 2009a. Cancer: new registrations and deaths 2005, revised edn. Wellington: Ministry of Health.

NZ Ministry of Health 2009b. Personal communication with staff at the NZ Ministry of Health.

Odicino F, Pecorelli S, Zigliani L & Creasman WT 2008. History of the FIGO cancer staging system. International Journal of Gynecology and Obstetrics 101:205–10.

Oriel KA, Hartenbach EM & Remington PL 1999. Trends in United States ovarian cancer mortality, 1975–1995. Obstetrics & Gynecology 93(1):30–3.

Parkin DM & Iscovich J 1997. Risk of cancer in migrants and their descendants in Israel: II. Carcinomas and germ-cell tumours. International Journal of Cancer 70(6):654–60.

Pecorelli S, Ngan HY & Hacker NF (eds) 2000. Staging classification and clinical practice guidelines for gynaecological cancers. International Journal of Gynecology & Obstetrics 10(2):207–312.

Petignat P, Fioretta G, Verkooijen HM, Vlastos AT, Rapiti E, Bouchardy C et al. 2004. Poorer survival of elderly patients with ovarian cancer: a population-based study. Surgical Oncology 13:181–6.

Queensland Cancer Registry & Cancer Council Queensland 2008. Cancer in Queensland. Incidence and mortality 1982 to 2005. Queensland: The Cancer Council Queensland.

Ries LAG, Melbert D, Krapcho M, Stinchcomb DG, Howlader N, Horner MJ et al. (eds) 2008. SEER cancer statistics review, 1975–2005. Bethesda, MD: National Cancer Institute.

Skirnisdottir I, Garmo H, Wilander E & Holmberg L 2008. Borderline ovarian tumours in Sweden 1960–2005: trends in incidence and age at diagnosis compared to ovarian cancer. International Journal of Cancer 123:1897–1901.

South Australia Cancer Registry 2000. Epidemiology of cancer in South Australia. Incidence, mortality and survival 1977 to 1999. Incidence and mortality 1999 analysed by type and geographical location. Twenty-three years of data. Adelaide: South Australia Cancer Registry, Epidemiology Branch, Statewide Division, Department of Human Services.

South Australia Cancer Registry 2008. Cancer in South Australia 2006 – with projections to 2009. Adelaide: South Australia Department of Health.

Supramaniam R, Grindley H & Pulver LJ 2006. Cancer mortality in Aboriginal people in New South Wales, Australia, 1994–2002. Australian and New Zealand Journal of Public Health 30(5):453–6.

Threllfall TJ, Thompson JR & Olsen N 2005. Cancer in Western Australia: incidence and mortality 2003 and Mesothelioma 1960–2003. Perth: Department of Health.

Thursfield V, Farrugia H & Giles G (eds) 2009. Cancer in Victoria 2006. Carlton: Cancer Epidemiology Centre, Cancer Council Victoria.

Tracey EA, Chen S, Baker D, Bishop J & Jelfs P 2006. Cancer in New South Wales: incidence and mortality 2004. Sydney: Cancer Institute New South Wales.

Tracey EA, Barraclough H, Chen W, Baker D, Roder D, Jelfs P & Bishop J 2007. Survival from cancer in NSW: 1980 to 2003. Sydney: Cancer Institute New South Wales.

Tracey E, Alam N, Chen W & Bishop J 2008. Cancer in New South Wales: incidence and mortality 2006. Sydney: Cancer Institute New South Wales.

Tracey EA, Roder D, Francis J, Zorbas HM, Hacker NF & Bishop J 2009. Reasons for improved survival from ovarian cancer in New South Wales, Australia, between 1980 and 2003: implication for cancer control. International Journal of Gynecological Cancer 19(4): 591–9.

UICC (International Union against Cancer) 2009. Geneva: UICC. Viewed 30 November 2009, <www.uicc.org/index.php>.

USCSWG (United States Cancer Statistics Working Group) 2009. United States cancer statistics: 1999–2005, incidence and mortality web-based report. Atlanta: U.S Department of Health and Human Services, Centre for Disease Control and Prevention, and National Cancer Institute. Viewed 21 September 2009, <www.apps.nccd.cdc.gov/uscs>.

Uyar D, Frasure HE, Markman M & Von Gruenigen VE 2005. Treatment patterns by decade of life in elderly women (≥ 70 years of age) with ovarian cancer. Gynecologic Oncology 98:403–8.

WCRF (World Cancer Research Fund) & AICR (American Institute for Cancer Research) 2007. Food, nutrition, physical activity and the prevention of cancer: a global perspective. Washington: AIRC.

Wiggins CL, Espey DK, Wingo PA, Kaur JS, Wilson RT, Swan J et al. 2008. Cancer among American Indians and Alaska natives in the United States, 1999–2004. Cancer 113(5): 1142–52.

WHC (Women's Health Council) & NCRI (National Cancer Registry Ireland) 2006. Women and cancer in Ireland 1994–2001. Dublin: The Women's Health Council.

WHO (World Health Organization) 1992. International Statistical Classification of Diseases and Related Health Problems, 10th revision. Volume 1. Geneva: WHO.

Yang L, Klint A, Lambe M, Bellocco R, Riman T, Bergfeldt K et al. 2008. Predictors of ovarian cancer survival: a population-based prospective study in Sweden. International Journal of Cancer 123:672–9.

Young JR, Roffers SD, Ries LAG, Fritz AG & Hurlbut AA (eds) 2001. SEER summary staging manual – 2000: codes and coding instruction. NIH pub. no. 01–4969. Bethesda, MD: National Cancer Institute. Viewed 17 September 2009, <www.seer.cancer.gov/tools/ssm/>.

Zhang X, Cordon J, Dempsey K & Garling L 2008. Cancer incidence and mortality, northern Territory 1991–2005. Darwin: Department of Health and Families.

# List of tables

Table 2.1:	Incidence of the 10 most commonly diagnosed cancers, females, 2006	6
Table 2.2:	Risk and average age at diagnosis of ovarian cancer, 1982 to 2006	9
Table 2.3:	Incidence of ovarian cancer and average age at diagnosis by type of ovarian cancer, 2006	11
Table 2.4:	Incidence by type of ovarian cancer and age at diagnosis, 2006	13
Table 2.5:	Incidence by type of ovarian cancer, 1982–1987 to 2000–2006	14
Table 2.6:	Incidence of ovarian cancer by stage at diagnosis, New South Wales and United States of America	16
Table 2.7:	Incidence of ovarian and other female genital organ cancers by stage at diagnosis, New South Wales, 1980–1998 and 1999–2003	16
Table 2.8:	Incidence of ovarian cancer by stage and age at diagnosis, United States of America, 1999–2005	17
Table 2.9:	Incidence of ovarian cancer by state and territory, 2002–2006	18
Table 2.10:	Incidence of ovarian cancer by Indigenous status, Queensland, Western Australia, South Australia and the Northern Territory, 2002–2006	21
Table 3.1:	The 10 most common types of cancer deaths, females, 2006	25
Table 3.2:	Risk of death from ovarian cancer and average age at death, 1982 to 2006	29
Table 3.3:	Mortality from ovarian cancer by state and territory, 2002–2006	30
Table 3.4:	Mortality from ovarian cancer by Indigenous status, New South Wales, Queensland, Western Australia, South Australia and the Northern Territory, 2002–2006	32
Table 3.5:	Underlying cause of death where ovarian cancer was an associated cause, annual average for 2002–2006	35
Table 3.6:	Women who died with ovarian cancer as an associated cause by age at death, annual average for 2002–2006	35
Table 4.1:	Relative survival, ovarian and breast cancer, females, 2000–2006	37
Table 4.2:	Incidence and 5-year relative survival by type of ovarian cancer, 1982–1987 to 2000–2006	43
Table 4.3:	Five-year relative survival by stage at diagnosis, ovarian cancer, New South Wales, 1980–2003	44
Table 4.4:	Five-year relative survival by stage at diagnosis and age group, ovarian cancer, United States of America, 1999–2005	45
Table 4.5:	Five-year crude survival by Indigenous status, ovarian cancer, Queensland, Western Australia, South Australia & Northern Territory, 1997-2006	46
Table 5.1:	Limited-duration prevalence of ovarian cancer, end of 2006	49
Table 5.2:	Limited-duration prevalence of the 10 most commonly diagnosed cancers, females, end of 2004	49
Table 5.3:	Twenty-five-year prevalence of ovarian cancer by age group, end of 2006	50
Table 5.4:	Limited-duration prevalence of ovarian cancer by state and territory of diagnosis, end of 2006	51
Table 5.5:	Limited-duration prevalence of ovarian cancer by country/region of birth, end of 2006	52
Table 6.1:	Leading causes of burden of disease, including leading cancers, females, 2003	54

Table 6.2:	Leading causes of burden of disease, including leading cancers, by fatal (YLL) and non-fatal (YLD) components, females, 2003	55
Table 6.3:	Trends and projected burden of ovarian and related cancers, 1993 to 2023	57
Table 7.1:	Hospitalisations for ovarian cancer and all reasons, females, 2007-08	59
Table 7.2:	Average length of stay (ALOS) for ovarian cancer-related hospitalisations by age group, 2007–08	60
Table 7.3:	Hospitalisations for ovarian cancer by same-day and overnight status, 1999–00 to 2007–08	60
Table 7.4:	Hospitalisations for ovarian cancer by most common procedures, 2007–08	63
Table 8.1:	Hospital admitted patient services expenditure by disease, females, 2004-05	65
Table 8.2:	Hospital admitted patient services expenditure by disease, constant prices, females, 2000–01 and 2004–05	66
Table B.1:	Age-standardisation method and reference population for analyses of differences in incidence and mortality rates by group	70
Table B.2:	Codes for ovarian and related cancers in the International Statistical Classification of Diseases and Related Health Problems, tenth revision (ICD-10)	71
Table C.1:	Agreed set of items to be provided by the states and territories to the AIHW for inclusion in the Australian Cancer Database	77
Table D2.1:	Incidence of ovarian cancer by age at diagnosis, 2006	82
Table D2.2:	Incidence of ovarian cancer, 1982 to 2006	83
Table D2.3:	Incidence of ovarian cancer by age at diagnosis, 1982 to 2006	84
Table D2.4:	Projected ovarian cancer incidence, 2007 to 2015	85
Table D2.5:	Grouping of ovarian cancer histology types	85
Table D2.6:	Incidence by type of ovarian cancer, women aged less than 50 years at diagnosis, 1982–1987 to 2000–2006	86
Table D2.7:	Incidence by type of ovarian cancer, women aged 50 to 69 years at diagnosis, 1982–1987 to 2000–2006	87
Table D2.8:	Incidence by type of ovarian cancer, women aged 70 years and over at diagnosis, 1982–1987 to 2000–2006	88
Table D2.9:	Incidence of ovarian cancer by remoteness area, 2002-2006	89
Table D2.10:	Incidence of ovarian cancer by socioeconomic status, 2002-2006	89
Table D2.11:	Incidence of ovarian cancer by country/region of birth, 2002-2006	90
Table D2.12:	International comparison of estimated incidence of ovarian and related cancers, 2002	91
Table D3.1:	Mortality from ovarian cancer by age at death, 2006	92
Table D3.2:	Mortality from ovarian cancer, 1968 to 2006	92
Table D3.3:	Mortality from ovarian cancer by age at death, 1982 to 2006	94
Table D3.4:	Mortality from ovarian cancer by remoteness area, 2002-2006	95
Table D3.5:	Mortality from ovarian cancer by socioeconomic status, 2002-2006	95
Table D3.6:	Mortality from ovarian cancer by country/region of birth, 2002-2006	96
Table D3.7:	International comparison of estimated mortality from ovarian and related cancers, 2002	97
Table D4.1:	Relative survival, 10 most commonly diagnosed cancers, females, 1998-2004	98
Table D4.2:	Relative survival by age at diagnosis, ovarian cancer, 2000–2006	98
Table D4.3:	Relative survival by period of diagnosis, ovarian cancer, 1982–1987 to 2000–2006	99

Table D4.4:	Five-year relative survival by age at diagnosis, ovarian cancer, 1982–1987 to 2000–2006	100
Table D4.5:	Five-year relative survival by type of ovarian cancer and age at diagnosis, 1982–2006	101
Table D4.6:	International comparison of mortality-to-incidence ratios for ovarian and related cancers, 2002	102
Table D6.1:	Leading cancer causes of burden of disease by age group, females, 2003	103
Table D7.1:	Hospitalisations for ovarian cancer by age group, 2007–08	104
Table D7.2:	Hospitalisations for ovarian cancer by same-day and overnight status, 1999–00 to 2007–08	104
Table D7.3:	Hospitalisations for ovarian cancer by age group, 1999-00 to 2007-08	105
Table D7.4:	Average length of stay (ALOS) for ovarian cancer-related hospitalisations by same-day and overnight status, 1999-00 to 2007-08	105
Table D8.1:	Hospital admitted patient services expenditure and number of hospitalisations for ovarian and related cancers by age group, 2004–05	106
Table E.1:	Carcinoma of the ovary: FIGO nomenclature (Rio de Janeiro 1998)	107
Table E.2:	Summary stage – extent of disease at diagnosis	108
Table F.1:	Hospitalisations for ovarian cancer by same-day and overnight status, 2007-08	110
Table F.2:	Number of ovarian cancer-related hospitalisations for same-day 'Pharmacotherapy sessions for neoplasm' by state and territory, 1999–00 to 2007–08	111

# List of figures

Figure 1.1:	The ovaries and nearby organs	1
Figure 2.1:	Incidence of ovarian cancer by age at diagnosis, 2006	7
Figure 2.2:	Incidence of ovarian cancer, 1982 to 2006	7
Figure 2.3:	Incidence of ovarian cancer by age at diagnosis, 1982 to 2006	8
Figure 2.4:	Incidence of ovarian cancer, observed for 1997 to 2006 and projected for 2007	
	to 2015	10
Figure 2.5:	Incidence of ovarian cancer by remoteness area, 2002–2006	19
Figure 2.6:	Incidence of ovarian cancer by socioeconomic status, 2002–2006	20
Figure 2.7:	Incidence of ovarian cancer by country/region of birth, 2002-2006	22
Figure 2.8:	International comparison of estimated incidence of ovarian and related cancers, 2002	23
Figure 3.1:	Ovarian cancer incidence and mortality by age group, 2006	26
Figure 3.2:	Ovarian cancer incidence and mortality, 1968 to 2006	27
Figure 3.3:	Mortality from ovarian cancer by age at death, 1982 to 2006	28
Figure 3.4:	Mortality from ovarian cancer by remoteness area, 2002-2006	31
Figure 3.5:	Mortality from ovarian cancer by socioeconomic status, 2002-2006	32
Figure 3.6:	Mortality from ovarian cancer by country/region of birth, 2002-2006	33
Figure 3.7:	International comparison of estimated mortality from ovarian and related cancers, 2002	34
Figure 4.1:	Relative survival, 10 most commonly diagnosed cancers, females, 1998-2004	38
Figure 4.2:	Relative survival by age at diagnosis, ovarian cancer, 2000–2006	39
Figure 4.3:	Relative survival by period of diagnosis, ovarian cancer, 1982–1987 to 2000–2006	40
Figure 4.4:	Five-year relative survival by age at diagnosis, ovarian cancer, 1982–1987 to 2000–2006	41
Figure 4.5:	Five-year relative survival by type of ovarian cancer, 2000–2006	42
Figure 4.6:	International comparison of mortality-to-incidence ratios for ovarian and related cancers, 2002	47
Figure 6.1:	Leading causes of burden of disease, including leading cancers, by fatal and non-fatal components, females, 2003	56
Figure 6.2:	Leading cancer causes of burden of disease by age group, females, 2003	56
Figure 7.1:	Hospitalisations for ovarian cancer by age group, 2007–08	59
Figure 7.2:	Hospitalisations for ovarian cancer by same-day and overnight status, 1999–00 to 2007–08	61
Figure 7.3:	Hospitalisations for ovarian cancer by age group, 1999–00 to 2007–08	62
Figure 7.4:	Average length of stay for ovarian cancer-related hospitalisations by same-day and overnight status, 1999–00 to 2007–08	62
Figure 8.1:	Hospital admitted patient services expenditure on ovarian and related cancers by age group, 2004–05	65