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Acknowledgments

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The AIHW would like to thank Professor Hugh Taylor and the Australian Bureau of Statistics for their valuable assistance in reviewing the first edition of this report.

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Abbreviations

ABS    Australian Bureau of Statistics
AIHW   Australian Institute of Health and Welfare
ASCED  Australian Standard Classification of Education
CATI   computer-aided telephone interviewing
CURF   confidentialised unit record file
DEEWR  Department of Education, Employment and Workplace Relations
DoHA   Department of Health and Ageing
DVA    Department of Veterans’ Affairs
FaHCSIA Department of Families, Housing, Community Services and Indigenous Affairs
GP     general practitioner
ICD-9-BPA International Statistical Classification of Diseases and Related Health Problems, Ninth Revision, British Paediatric Association
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
ICF    International Classification of Functioning, Disability and Health
ICO    International College of Ophthalmologists
ICPC   International Classification of Primary Care
MBS    Medicare Benefits Schedule
MDS    Minimum Data Set
NDS-IS National Data Standard for Injury Surveillance
NMDS   National Minimum Data Set
OATSIH Office for Aboriginal and Torres Strait Islander Health
PBS    Pharmaceutical Benefits Scheme
RFE    reason for encounter
SEIFA  Socio-Economic Indexes for Areas
SLA    Statistical Local Area
WHO    World Health Organization
Summary

This report updates *A guide to Australian eye health data*, first published in 2007. Since the publication of the first edition there have been a number of developments in the availability of eye health data, most notably sources of information about eye health among Aboriginal and Torres Strait Islander people. Data sources that have been included in this edition for the first time include:

- the National Indigenous Eye Health Survey
- the Longitudinal Study of Indigenous Children
- the National Sun Survey
- National Trachoma and Surveillance Reporting Unit data
- Northern Territory Emergency Response Child Health Check Initiative data.

As well as updating information from in the first edition, this report assesses how well the *National eye health framework* is informed by available eye health data. This includes the data sources included in the first edition as well as the new additions.
1 Introduction

1.1 Purpose of this report

In 2006, the Australian Government Department of Health and Ageing commissioned the Australian Institute of Health and Welfare (AIHW) to compile details of key Australian data sources that are relevant to assessing prevalence and outcomes of eye diseases and injuries and eye health-care utilisation (see Box 1).

The second edition of this report provides information on eye health data sources which have been made available since the publication of the first edition, as well as updating details of older data sources. Examples of new data sources in this guide include the National Indigenous Eye Health Survey, the National Sun Survey, the Longitudinal Study of Indigenous Children, National Trachoma Surveillance and Reporting data and Child Health Check Initiative data, among others.

Box 1: Background to this report

In response to the World Health Assembly resolution WHA56.26 on the elimination of avoidable blindness in member states, the Australian Health Ministers’ Conference endorsed the National Framework for Action to Promote Eye Health and Prevent Avoidable Blindness and Vision Loss (National Eye Health Framework or the Framework). The Framework focuses on eliminating avoidable blindness and vision loss in Australia, providing an outline for nationally coordinated action by governments, health professionals, non-government organisations, industry and individuals to work in partnership (Commonwealth of Australia 2005).

In the 2006 Federal Budget, the Australian Government allocated $13.8 million over 4 years to a new National Eye Health Initiative. This initiative supports a range of activities to raise public awareness of eye health issues and to strengthen the delivery of eye health care.

The Australian Government Department of Health and Ageing commissioned the AIHW through the National Eye Health Initiative to do this report. This follows on from work done for earlier publications: Vision problems among older Australians (released July 2005), Eye health in Australia: a hospital perspective (released August 2008), Eye health among Australian children (released November 2008), Eye-related injuries in Australia (released February 2009) and Eye health labour force in Australia (released September 2009). The first edition of this report was released in May 2007.

The key areas for action within the Framework provide a brief outline of the challenges to be tackled and a series of actions that might be used to meet these challenges.

Key action area 5 (Improving the evidence base) outlines a need to look at existing health data sets for relevance to eye health, and to identify eye health research gaps. This report informs this action area by presenting useful information for the development of future eye health performance indicators.
1.2 Structure of this report

This report is divided into three sections. Following this introduction, Section 2 presents detailed listings of each key eye health data source, as well as a brief analysis of the current most important eye health data sources in Australia that inform the Framework.

Section 3 presents a discussion on the definitional issues surrounding eye health terminology and includes a summary of eye health definitions in use in Australia.

1.3 Eye health in Australia

It is estimated that in excess of 161 million people worldwide are visually impaired. This includes people with significant loss of function and disability due to the impact of vision loss. The leading cause of this visual impairment is cataract. Other major causes of visual impairment worldwide are glaucoma, age-related macular degeneration, diabetic retinopathy and trachoma (Resnikoff et al. 2004).

Self-reported data on the prevalence of loss of sight is available through the Australian Bureau of Statistics (ABS) National Health Survey 2007–08. The survey indicates that 52% of the Australian population report eyesight problems, including long- and short-sightedness, as a long-term medical condition (ABS 2009).

As in other developed countries, the most prevalent causes of blindness and vision loss in Australia are those related to ageing. Eye health in Australia, the background paper to the National Eye Health Framework, reported that age-related macular degeneration, cataract, glaucoma, diabetic retinopathy, uncorrected or under-corrected refractive error, eye trauma and trachoma are the most prevalent causes of blindness and vision loss in Australia (Commonwealth of Australia 2005a). However, eye disorders are not only a problem for elderly adults. For example, eye disorders are among the most common long-term health problems experienced by Australian children (AIHW 2008).

In a recent analysis of pooled eye health data from population-based clinical studies, conducted both in Australia and internationally, it was estimated that 9.4% of Australians aged 55 or older are visually impaired and about 1.2% are blind (Figure 1). The combined impact of an ageing Australian population and the high age correlation of causes of vision loss indicate that the prevalence of visual impairment is set to increase over time in a policy-neutral environment (Access Economics 2004). It is also estimated that the absolute number of people with low vision or blindness will almost double by 2024 on the 2000 figure (Taylor et al 2005).

Certain groups within the Australian population are at greater risk of developing eye disease. These groups include older people, people with a family history of eye disease, people with diabetes, and marginalised to disadvantaged people (Commonwealth of Australia 2005a). Some Aboriginal and Torres Strait Islander communities have a high prevalence of trachoma, which is an infectious disease linked to poor hygiene that may lead to blindness (Tellis et al 2008).

People with diabetes have an increased risk of developing eye disease. The Australian Diabetes, Obesity and Lifestyle Study found that 15.3% of people with known diabetes and newly diagnosed diabetes had developed retinopathy (Tapp et al. 2003).

Eye disease and vision loss have considerable financial and social costs to the Australian community. Visual impairment can shorten life, increase the risk of other conditions, restrict
social participation and independence, and impair physical and mental health. In addition, people with visual impairment have a higher use of social services and a higher admission rate to nursing homes. It has been calculated that the total financial cost of visual impairment, both direct and indirect, was over $5.0 billion in 2004 (Access Economics 2004).

Figure 1: Estimated prevalence of eye disorders for people over age 55, Australia

Source: AIHW 2005a.

Note: These data are derived from pooling estimates from a number of surveys, including the National Health survey (2001, 1995), the Australian Diabetes, Obesity and Lifestyle Study (1999–2000), Melbourne Visual Impairment Project (1992–1996), and the Blue Mountains Eye Study (1992–1994).
2 Key data sources

This chapter describes the key data sources available that are relevant to assessing the prevalence and outcomes of eye diseases and injuries and eye health-care utilisation in Australia. The data sources contain information collected nationally, in a state or territory or regionally.

2.1 Methodology

Consultations with stakeholders and an extensive search of information provided a wide range of eye health data sources which indicate the prevalence and outcomes of eye diseases and injuries and eye health-care utilisation. A greater understanding of the data sources and their application to the monitoring of eye health issues was gained through consultation with the data sources’ primary contact officers. A similar approach was taken for updating the second edition of the data guide with new eye health data sources.

While identification of these data sources mainly focused on those within the health and welfare areas, relevant data collections in the areas of income, education and workforce were also explored. Data sources from these areas provide additional information to describe people with eye diseases and injuries. This report focuses predominately on population-survey-based and administrative national data collections within the public domain, that is, those that are available to the public either through published reports or via access to the data for research. However, the focus is not restricted exclusively to this domain. Jurisdictional or local-level data sources have also been included, depending on their relevance and quality.

Key data sources were documented individually according to a standard template. The template was designed to gather information on the purpose, collection methodology, data content, scope and details concerning eye health specific data, along with data custodian contact details.

Noted eye health researcher Professor Hugh Taylor was asked by the AIHW to verify the breadth and relevance of data sources referenced in the first edition of this publication and to provide comments on eye health definitions used in this report. The ABS also provided valuable comments.
2.2 Key Australian eye health data sources

Lists of national and regional data sources

National

Australian Cancer Database
Australian Congenital Anomalies System
Australian Corneal Graft Registry
Australian Longitudinal Study on Women’s Health
Australian National Diabetes Information Audit & Benchmarking
Bettering the Evaluation and Care of Health
Census of Population and Housing
Disability Services National Minimum Data Set
Disability Support Pension collection
Disease Expenditure Database
Growing Up in Australia: the Longitudinal Study of Australian Children
Higher Education Student Collection
Home and Community Care Program Minimum Data Set
Longitudinal Study of Indigenous Children
Macular Degeneration Foundation Database
Medical Labour Force Survey
Medicare Australia data
National Aboriginal and Torres Strait Islander Health Survey
National Data Set for Compensation-based Statistics
National Elective Surgery Waiting Times Data Collection
National Health Survey
National Hospital Morbidity Database
National Indigenous Eye Health Survey
National Mortality Database
National Outpatient Care Database
National Sun Survey
National Trachoma Surveillance and Reporting Unit
Nursing and Midwifery Labour Force Survey
Office for Aboriginal and Torres Strait Islander Health Service Activity Reporting system
Optometrists Association Australia Optometrists Database
Optometrist registration boards
Pharmaceutical Benefits Scheme data
Royal Australian and New Zealand College of Ophthalmologists Ophthalmology workforce database
Survey of Disability, Ageing and Carers
Treatment Accounts System

Regional

The 45 and Up Study
Association for the Blind of Western Australia Register of Blindness and Vision Impairment
Australian Longitudinal Study of Ageing
Blue Mountains Eye Study
Cataract Surgery and Age-related Macular Degeneration Study
Central Australia Ocular Health Study data/hospital records/Regional Eye Health Coordinator records
Dubbo Study
Florey Adelaide Male Ageing Study
International Centre for Eyecare Education/Aboriginal Health and Medical Research Council/VisionCare New South Wales Aboriginal project data sheets
Katherine Region Diabetic Retinopathy Study
Melbourne Longitudinal Studies in Healthy Ageing program
Melbourne Visual Impairment Project
New South Wales Population Health Survey
Northern Territory Emergency Response Child Health Check Initiative data collection
North West Adelaide Health Study
Queensland Injury Surveillance Unit Database
Royal Society for the Blind client database
South Australian Health Monitor survey
South Australian Health Omnibus Survey
South Australian Monitoring and Surveillance System
Spectacle assistance schemes
Sydney Myopia Study
Sydney Older Persons Study
Victorian Emergency Minimum Dataset injury surveillance collection
Victorian Population Health Survey
Western Australian Aboriginal Child Health Survey
National data sources

Australian Cancer Database

Description of data source
The Australian Cancer Database (ACD) receives data from individual state and territory cancer registries on cancer diagnosed in residents of Australia. This commenced with cases first diagnosed in 1982. The data items provided to the ACD by the state and territory cancer registries enable the compilation of statistics on cancer types by sex, age group and other variables.

Purpose(s)
The aim of the ACD is to foster the development and dissemination of national cancer statistics for Australia and specifically to:
- enable computation and publication of national statistics on cancer
- allow tracking of interstate movement of cancer cases via record linkage
- facilitate exchange of scientific and technical information between cancer registries and promote standardisation in the collection and classification of cancer data
- facilitate cancer research both nationally and internationally.

Collection methodology
It is a legal requirement for doctors to report new cases of cancer to state and territory cancer registries, who supply the data annually to the Australian Institute of Health and Welfare (AIHW).

Data content
The main variables collected are names, sex, date of birth, date and basis of diagnosis, SLA at time of diagnosis, site of cancer and cell type.

Data content related to eye health
Case information about people diagnosed with cancer of the eye and adnexa. Eye health-related data are medically verified.

Indigenous identification
Indigenous status is collected, however ascertainment is only satisfactory for Western Australia (WA), Northern Territory (NT) and Queensland.

Socioeconomic identification
No.

Scope/target population
All residents of Australia.
Geographical coverage
All states and territories, Australia.

Time period covered
Since 1982 to present (ongoing collection).

Publications and other outputs
The ACD produces reports of national incidence and mortality data. Periodically, analyses are undertaken of specific cancer sites, cancer histology, differentials in cancer rates by country of birth, geographical variation, trends over time and survival. The most recent publications are:


The cancer interactive data cubes include the number of new cases and age-specific rates for all registrable cancers by year of diagnosis, sex and 5-year age groups: <www.aihw.gov.au/cancer/data/datacubes/index.cfm> (viewed 19 August 2009).

Access
The ACD can make available a broad range of statistical data to researchers after a strict scientific and ethical review process. Research database linkage with the ACD requires approvals from the AIHW Ethics Committee and all the state and territory cancer registries, including their ethics committees.

Data management agency
Australian Institute of Health and Welfare (commissioned by Australasian Association of Cancer Registries)
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
**Australian Congenital Anomalies System**

**Description of data source**

The Australian Congenital Anomalies System contains information on congenital anomalies notified to birth defect registers and other data collections of Australian states and territories.

**Purpose(s)**

The purposes of the Australian Congenital Anomalies System are to monitor congenital anomalies at a national level, to fulfil international reporting requirements and to work towards the standardisation of definitions, classifications and collection methods among the states and territories.

**Collection methodology**

Data on congenital anomalies are collected in state and territory birth defect registers, perinatal data systems, hospital systems and other sources. These data are reported to the AIHW National Perinatal Statistics Unit (NPSU).

A National Minimum Data Set for congenital anomalies has been developed by NPSU. This will result in national standards for collection of congenital anomalies data.

**Data content**

For the mother, the following variables are collected: age, country of birth, Indigenous status, area of usual residence, place of birth, hospital of birth, hospital sector, admitted patient elected accommodation status, parity, gestation and plurality.

For the baby/infant, the following variables are collected: sex, birth weight, outcome and congenital anomaly. Anomalies are classified differently among the states and territories using either the International Statistical Classification of Diseases and Related Health Problems, Ninth Revision, British Paediatric Association (ICD-9-BPA), International Statistical Classification of Diseases and Related Health Problems, Ninth Revision, Clinical Modification (ICD-9-CM) or International Statistical Classification of Diseases and Related Health Problems, Tenth Revision, Australian Modification (ICD-10-AM).

**Data content related to eye health**

Congenital anomalies of the eye.

Eye health-related data are medically verified.

**Indigenous identification**

Yes, however the quality has not been determined.

**Socioeconomic identification**

Area of usual residence.

**Scope/target population**

Included within the collection are births of at least 20 weeks gestation or 400 grams birth weight with a major congenital anomaly notified up to 1 year of age, and terminations of pregnancy less than 20 weeks gestation and less than 400 grams birth weight with a major congenital anomaly.
The period of notification of a congenital anomaly varies among the states and territories. For some jurisdictions, the period of notification is the perinatal period or the birth episode. For others, the period of notification is up to 1, 5, 6 or 15 years of age. Only four jurisdictions provide data on terminations of pregnancy less than 20 weeks gestation or 400 grams birth weight.

**Geographical coverage**

All states and territories, Australia.

**Time period covered**

Data have been compiled for 1998 to 2005.

**Publications and other outputs**

- Data from the Australian Congenital Anomalies System also contributes to research projects.

**Access**

Access to the Australian Congenital Anomalies System can be requested through the Director, AIHW.

**Data management agency**

AIHW National Perinatal Statistics Unit  
Level 2 McNevin Dickson Building  
Randwick Hospital’s Campus  
Gate 6 Avoca St  
Randwick NSW 2031  
<www.npsu.unsw.edu.au>  
(02) 9382 1014
**Australian Corneal Graft Registry**

**Description of data source**

The Australian Corneal Graft Registry operates an Australia-wide register of human corneal transplants.

**Purpose(s)**

The purpose of the Australian Corneal Graft Registry is to collect information on corneal transplants performed in Australia, to measure graft and visual outcomes that will inform clinical practice and to identify risk factors for poor patient outcomes.

**Collection methodology**

Pro-formas completed by the treating ophthalmologist or other medical professional are returned by mail at the time of corneal transplantation and at follow-up.

Registration is one-off for a given corneal transplant. Follow-up is at approximately yearly intervals thereafter until recipient death, loss-to-follow-up or graft failure.

**Data content**

The initial register collects information including de-identified information on the corneal transplant recipient, the corneal donor, the eye bank procedures and the operative procedure.

The follow-up collects information on graft survival, post-operative course and management, complications and visual outcome.

**Data content related to eye health**

Corneal transplants performed in Australia, along with graft and visual outcomes.

Eye health-related data are medically verified.

**Indigenous identification**

No.

**Socioeconomic identification**

No.

**Scope/target population**

Any person in Australia with a corneal transplant.

**Geographical coverage**

All states and territories, Australia.

**Time period covered**

Since 5 May 1985 to present (ongoing collection).
Publications and other outputs

- Annual presentations at specialist meetings.
- Occasional papers in international literature.

Access
Access to the Australian Corneal Graft Registry is restricted.

Data management agency
Australian Corneal Graft Registry
Department of Ophthalmology
Flinders Medical Centre
Bedford Park SA 5042
(08) 8204 5321
**Australian Diabetes, Obesity and Lifestyle study**

**Description of data source**

The Australian Diabetes, Obesity and Lifestyle (AusDiab) study is the largest Australian longitudinal population-based study examining the natural history of diabetes, pre-diabetes (in which glucose metabolism is impaired but not to the level to cause diabetes), heart disease and kidney disease.

**Purpose(s)**

The results provide a picture of the incidence of diabetes, cardiovascular disease and kidney disease, as well as the factors that increase the risk of these conditions.

**Collection methodology**

AusDiab is a field survey involving both physical testing as well as questionnaires. A team of researchers went to each of the 42 randomly selected testing sites around Australia to individually test each of the 11,247 individuals who participated in the baseline study in 1999–2000. Of these original participants, around 6,500 came back to participate in a similar survey five years later in 2004–2005. In addition, self-reported health information was obtained from more than 2,000 of those who could not attend the survey site.

**Data content**

AusDiab collects data in the following categories: biochemistry, demographic, medical history, disease status, survey procedures, anthropometric, smoking, exercise, clinical observations, self-efficacy, knowledge and opinion, socioeconomic, family medical history, alcohol and diet.

**Data content related to eye health**

Participants are asked the following questions:
- Do you think eye trouble or blindness is likely to occur in people with diabetes?
- Have you ever been treated for or suffered from trouble with the back of your eyes or retinopathy?
- Have you ever had the back of your eyes (retina) examined? Who conducted the eye examination? How long has it been since you last had the back of your eyes checked?

Eye health-related data are self-reported.

**Indigenous identification**

Participants are recorded as Aboriginal/Torres Strait Islander or neither.

**Socioeconomic identification**

Participants provide information about accommodation, education, income and employment.

**Scope/target population**

The baseline AusDiab (1999–2000) study was a population-based national survey of the Australian general population aged 25 years or older residing in 42 randomly selected urban areas.
and rural areas (census collector districts) in six states and the NT. AusDiab was designed to be representative of the general Australian population aged over 25 years.

**Geographical coverage**

Forty-two randomly selected urban and rural areas of Australia. A stratified cluster sample was used involving seven strata (the six states and the NT) and clusters were based on census collector districts.

**Time period covered**

Surveys were conducted in 1999–2000 and 2005. Pending further funding, a follow-up to AusDiab and a new cross-sectional survey will be conducted in 2009–10.

**Publications and other outputs**

Refereed publications, including those relevant to eye health:


**Access**

The AusDiab team welcomes approaches from research workers for access to the accumulated data and biological materials and for participation in ongoing and new data collection activities. Access will generally not be granted to research groups/institutions or individuals who are receiving support from the Australian Tobacco Research Foundation or other bodies associated with the tobacco industry.

**Data management agency**

International Diabetes Institute
250 Kooyong Road
Caulfield VIC 3162
<www.diabetes.com.au>
(03) 9258 5050
Australian Longitudinal Study on Women’s Health

Description of data source

The Australian Longitudinal Study on Women’s Health, also known as Women’s Health Australia, is a longitudinal population-based survey that examines the health of three large cohorts of Australian women over a 20-year period. It was first funded in 1995. The project was designed to explore factors that influence health among women who are broadly representative of the entire Australian population. The project goes beyond a narrow perspective that equates women’s health with reproductive and sexual health, and takes a comprehensive view of all aspects of health throughout women’s lifespans.

Purpose(s)

The aim of the project is to collect scientifically valid information that is relevant to the development of health policy, so that the Australia Government and state/territory government departments of health can base policy and practice in women’s health on current data and accurate information.

An overall goal of the project is to clarify cause-and-effect relationships between women’s health and a range of biological, psychological, social and lifestyle factors. Another goal is to assess the effects of changes in health policy and practice.

Collection methodology

Participants were initially selected by simple random sampling from the Medicare Australia database, with over-sampling for rural and remote areas. Three cohorts were recruited, with each cohort resurveyed every 3 years. The questionnaire is self-completed, however a telephone interview is provided for non-English-speaking respondents or when requested by participants.

Data content

The project assesses each participant’s:

• physical and emotional health (including wellbeing, major diagnoses, symptoms)
• use of health services (general practitioner, specialists and other visits, access, satisfaction)
• health behaviours and risk factors (diet, exercise, smoking, alcohol, other drugs)
• time use (including paid and unpaid work, family roles and leisure)
• sociodemographic factors (location, education, employment, family composition)
• life stages and key events (such as childbirth, divorce, widowhood).

Data content related to eye health

Participants have been asked the following questions:

• In the past 12 months have you had eyesight problems?
• Have you sought help for eyesight problems?
• Have you had eye surgery in last three years?
• In the last three years have you had cataract surgery?
• In the last three years have you had any other eye surgery?
• In the last three years have you been on a waiting list for cataract surgery?
• In the last three years have you been on a waiting list for other eye surgery?
• Do you have difficulty seeing print, even with glasses?
• Do you have difficulty recognising people across the road, even with glasses?

Eye health-related data are self-reported.

Indigenous identification
Participants are recorded as Aboriginal, Torres Strait Islander or neither.

Socioeconomic identification
Participants provide details of their postcode, highest educational qualification achieved and household income.

Scope/target population
Three age groups were selected in order to follow women through life stages that are critical to women’s health and wellbeing. Ages at recruitment were 18–23 years of age (younger), 45–50 years of age (mid-aged) and 70–75 years of age (older). The younger age group began to be tracked when they were in the early stages of transition from late adolescence to full adulthood; over time, most of them are moving into the workforce, entering adult relationships and becoming mothers. The mid-age group was selected to examine menopausal transitions and the social and personal changes of middle age. The older group was selected in their early 70s, to recruit older women who are generally still active, involved members of the community.

Geographical coverage
Urban, rural and remote areas, all states and territories, Australia.

Time period covered
Since 1996 to 2016.

Publications and other outputs
A number of annual reports and technical reports are available on the Women’s Health Australia website: <www.alswh.org.au/reports.html> (viewed 31 July 2009).

Access
Data may be made available to collaborating researchers where there is a formal request to make use of the material. Permission to use the data must be obtained from the Publications, Analyses and Substudies Committee of the Australian Longitudinal Study on Women’s Health. Where data are collected in collaboration with, by or for a community, community permission to use data is also necessary. Rules relating to data concerning Aboriginal and Torres Strait Islander communities, as set out by the National Health and Medical Research Council, apply.
Data management agency

Women’s Health Australia
Research Centre for Gender, Health and Ageing
University of Newcastle
Callaghan NSW 2308
<www.alswh.org.au>
(02) 4923 6872
whasec@newcastle.edu.au
**Australian National Diabetes Information Audit & Benchmarking**

**Description of data source**

The Australian National Diabetes Information Audit & Benchmarking (ANDIAB) project involves the collection of de-identified data on individuals with diabetes seen in diabetes centres and by specialist endocrinologists in private practice over a one-month period.

**Purpose(s)**

The purpose of ANDIAB is to collect information on individuals diagnosed with diabetes.

**Collection methodology**

Information is collected in survey format predominantly, plus some data from established databases.

**Data content**

Collected information includes:
* demographic details (date of birth, sex, Indigenous status)
* diagnosis details (year of diagnosis, type of diabetes, management method, current smoker, weight, height, blood pressure)
* comorbidity/complication details
* contact with health care providers (optometrists, ophthalmologists, podiatrists, diabetes educators and dieticians).

**Data content related to eye health**

Participants are asked the following questions:
* Have you seen an optometrist in the last 12 months?
* Have you been referred to an ophthalmologist in the last 12 months?
* Have you had a fundus examination in the last 12 months?
* Do you have a cataract currently present or previously removed?

They also have their visual acuity measured and are asked about comorbidities. Eye health-related data are self-reported and medically verified.

**Indigenous identification**

The patient is recorded to be either Aboriginal/Torres Strait Islander or neither.

**Socioeconomic identification**

No.

**Scope/target population**

Individuals diagnosed with diabetes who attend specialist diabetes services.

**Geographical coverage**

All state and territories, Australia, excluding WA and the NT in 2004.
Time period covered
Since 1998 to present (ongoing collection).

Publications and other outputs

Access
By application, contact:
Assoc. Prof. JR Flack FRACP MM
Bankstown-Lidcombe Hospital
Eldridge Road
Bankstown NSW 2200

Data management agency
Diabetes Centre, Bankstown-Lidcombe Hospital (managed on behalf of National Association of Diabetes Centres)
Bettering the Evaluation and Care of Health

Description of data source
The Bettering the Evaluation and Care of Health (BEACH) program continuously collects information about the patients seen, reasons people seek medical care, problems managed and treatments provided in general practice in Australia.

Purpose(s)
BEACH aims to:
- collect reliable and valid data about general practice which is responsive to the ever-changing needs of information users
- establish an ongoing database of general practitioner (GP)/patient encounter information
- assess patient-based risk factors and the relationship these factors have with health service activity
- provide accurate and timely data to a wide variety of users.

Collection methodology
BEACH uses a cross-sectional, paper-based data collection system developed over the past 27 years at the University of Sydney.

Data content
BEACH collects the following information from each GP/patient encounter:
- encounter characteristics
- GP characteristics—age, gender, years in general practice, number of sessions per week, country of graduation, size of practice, computer use, hours worked and on call each week, rurality of practice
- patient characteristics—age, sex, Indigenous status, Health Care Card and Veteran’s Affairs status, status to the practice (new/seen before)
- patient reasons for encounter (up to three)
- problems managed at encounter (up to four)
- management for each problem (pharmacological and non-pharmacological)
- referrals, tests and investigations for each problem.

Data content related to eye health
All problems managed by GPs that are labelled as eye problems and therefore classified in Chapter F of the International Classification of Primary Care (ICPC), version 2. Eye health-related data are medically verified.

Indigenous identification
Patients may self-report as Aboriginal and/or Torres Strait Islander.

Socioeconomic identification
Postcode of usual residence.
Scope/target population

Patients attending a GP who have been randomly selected for the study. The sample represents patients who see their GP. About 85% of the population visit the GP in any given year. However, the sample over-represents those people who attend more frequently, as they have more chance of being included.

Geographical coverage

All states and territories, Australia.

Time period covered

Since April 1998 to present (ongoing collection).

Publications and other outputs

A full list of publications can be found through the BEACH project website: <www.fmrc.org.au/beach.htm> (viewed 31 July 2009).

Access

The general public and researchers may request ad hoc analyses of BEACH data to be performed, a charge is made for access to the data and for work undertaken. Contact Associate Professor Helena Britt at the Australian General Practice Statistics and Classification Centre.

Data management agency

Australian General Practice Statistics and Classification Centre
Family Medicine Research Centre, Acacia House
Westmead Hospital
Westmead NSW 2145
<www.fmrc.org.au/agpscc>
(02) 9845 8151
gpscru@med.usyd.edu.au
Census of Population and Housing

Description of data source

The Census of Population and Housing is conducted every five years. It aims to provide a count of the number of people in Australia on the night of the Census, as well as characteristics of the population. The Census is the most comprehensive data source available for information regarding the Australian population.

Purpose(s)

The purpose of the Census is to enumerate all persons in Australia and external territories on the night of the Census. It is also used to collect important information about individuals and households, and to provide a benchmark of the size and composition of the population which can be used in sample surveys. Population estimates derived from the Census are also used to determine electoral representation and the distribution of government funding.

Collection methodology

The Census is conducted every five years. Census forms are delivered to and collected from every household in Australia by Census collectors, with each collector working in a collector district (CD). Each CD contains, on average, about 230 households. The Census is largely self-enumerated. Census forms may also be returned by mail, or completed online (eCensus).

Data content

A wide variety of information is collected in the Census. These include the age and population distribution of the population, occupation and earnings, education, number of children, birthplace, household characteristics, disability, transport use and voluntary work.

Data content related to eye health

Employed persons are asked to state their occupation.

Counts of ophthalmologists, optometrists, orthoptists, optical dispensers and optical mechanics are available, coded to the Australian and New Zealand Standard Classification of Occupations.

Indigenous identification

The standard ABS classification for Indigenous identification is used.

An Indigenous Enumeration Strategy is put into place to encourage Aboriginal and Torres Strait Islander people to participate in the Census.

Socioeconomic identification

Participants are asked about a range of socioeconomic indicators, including address, education, qualifications and income.

Scope/target population

All persons (including temporary visitors) resident in Australia on the night of the Census. Foreign diplomats and their families are excluded, as are Australian residents overseas.
Geographical coverage
All states and territories, Australia.

Time period covered
Since 1911 to present. Five-yearly since 1961.
Before Federation censuses were held by each colony, the first being in New South Wales (NSW) in 1828. In 1881, a census was held simultaneously by each colony for the first time.

Publications and other outputs
Data from the Census are used in a vast number of publications. An overview can be found in:
Census data are also available on the ABS website, such as through CDATA Online.

Access
For users who wish to undertake more detailed analysis of Census data, two confidentialised unit record files (CURFs) are available. A Basic CURF and expanded CURF (containing more detailed information than the basic CURF) are accessible through the ABS Remote Access Data Laboratory (RADL) system.

Data management agency
Australian Bureau of Statistics
Locked Bag 10
Belconnen ACT 2612
<www.abs.gov.au>
client.services@abs.gov.au
Disability Services National Minimum Data Set

Description of data source

The Disability Services National Minimum Data Set (DS NMDS) (previously the Commonwealth State/Territory Disability Agreement (CSTDA) National Minimum Data Set) is a national collection that provides information relating to specialist disability services funded under the National Disability Agreement (NDA) and the people receiving these services. The collection consists of a set of nationally significant data items that are collected in all Australian jurisdictions, with an agreed method of collection and transmission.

Purpose(s)

The purpose of the DS NMDS collection is to facilitate the annual collation of nationally comparable data about NDA-funded services, and to obtain reliable, consistent data with minimal load on the disability services field. Under the NDA, the disability administrators in all Australian jurisdictions are responsible for ensuring that ‘DS NMDS information will be comparable across all jurisdictions and years’.

Collection methodology

The DS NMDS is an administrative data set. Data are collected by disability support service agencies and forwarded to the appropriate state/territory or Australian Government department. All data are then forwarded to the AIHW on an annual basis for collation of a national data set.

Data content

Information collected on service users includes date of birth, sex, Indigenous status, country of birth, interpreter services required, communication method, living arrangements, postcode, residential setting, disability group (primary and ‘other significant’), support needs, informal carer arrangements, receipt of carer allowance, labour force status, main source of income, individual funding status, and receipt of service variables.

Information collected on service type outlets includes service type, postcode, statistical local area (SLA), funding jurisdiction, agency sector, full financial year of CSTDA funding, operation times (weeks, hours, days), staff hours, and number of service users.

Data content related to eye health

‘Vision’ is included as a disability group for two data items.

- primary disability group: the disability group that most clearly expresses the experience of disability by a person. The primary disability group can also be considered as the disability group causing the most difficulty to the person (overall difficulty in daily life, not just within the context of the support offered by the service).
- other significant disability group(s): disability group(s) (other than that indicated as being ‘primary’) that also clearly express the experience of disability by a person and/or cause difficulty for the person.

Eye health-related data are medically verified.

Indigenous identification

The standard ABS classification for Indigenous identification is used.
Socioeconomic identification

No.

Scope/target population

Under the NDA, ‘people with disabilities’ means people with disabilities attributable to an intellectual, psychiatric, sensory, physical or neurological impairment or acquired brain injury (or some combination of these) which is likely to be permanent and results in substantially reduced capacity in at least one of the following:

• self care/management
• mobility
• communication
• requiring significant ongoing and/or long-term episodic support and which manifests itself before the age of 65.

Geographical coverage

All states and territories, Australia.

Time period covered

The first DS NMDS collection is for 2009–10 financial year (ongoing collection). A redevelopment of the NMDS is likely to occur in the future in order to align with the objectives of the NDA.

The DS NMDS is a continuation of the CSTDA NMDS, which began in October 2002. Previously data were collected on a ‘snapshot day’ basis under the former CSTDA NMDS implemented in 1995 and conducted at a national level from 1996 to 2002.

Publications and other outputs

• Annual reports, most recently:
  – AIHW 2008. Disability support services 2006–07: national data on services provided under the CSTDA. Cat. no. DIS 52. Disability series. Canberra: AIHW.
• Indicators from this data set are also reported in the annual report on government services:
• Previous AIHW publications relating to the CSTDA NMDS data can be found on the AIHW website: <www.aihw.gov.au/publications/index.cfm/subject/7> (viewed 31 July 2009).
• AIHW also publishes online, interactive data cubes for the DS/CSTDA NMDS. These can be accessed from the AIHW website: <www.aihw.gov.au/disability/datacubes/index.cfm> (viewed 31 July 2009).
Access

The AIHW is the custodian of the national data set; jurisdictions are custodians of their own data. Therefore any jurisdiction-specific data analyses must be approved by the appropriate jurisdiction(s). Access to the DS NMDS data can be requested through the AIHW.

Data management agency

Australian Institute of Health and Welfare
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
**Disability Support Pension collection**

**Description of data source**
The Disability Support Pension collection is an administrative database.

**Purpose(s)**
The purpose of the Disability Support Pension collection is to collect information about recipients of the pension.

**Collection methodology**
Updated quarterly in Centrelink’s SuperStar database.

**Data content**
General demographic details including age, gender, geographical location by state (also including those currently overseas) and postcode.

**Data content related to eye health**
Recipients identified as receiving the Disability Support Pension who are legally blind.
Eye health-related data are medically verified.

**Indigenous identification**
Recipients are identified as being Indigenous, non-Indigenous or unknown.

**Socioeconomic identification**
Information is collected about recipients’ postcode, country of birth, relationship status, home ownership and rent status.

**Scope/target population**
Australians who meet the eligibility requirements and are receiving the Disability Support Pension.

**Geographical coverage**
All states and territories, Australia.

**Time period covered**
Since 1999 to present (ongoing collection).

**Publications and other outputs**
Not applicable.

**Access**
Disability Support Pension data is available by request to Centrelink: datarequests@centrelink.gov.au.
Data management agency
Centrelink
<www.centrelink.gov.au>
Disease Expenditure Database

Description of data source
The report provides an overview of total health system expenditures on disease and injury in Australia during 2000–01, based on the best possible estimates from currently available data sources.

Purpose(s)
To provide a systematic analysis of Australian health expenditure by disease.

Collection methodology
The basic approach for direct costs of health services has been to take known aggregate expenditures on health care and apportion these to disease categories using Australian data.

Data content
Expenditure data are collected on hospitals, aged care homes, out-of-hospital medical services, pharmaceuticals, other professional services, dental and research (by age and sex).

Data content related to eye health
Expenditure data related to conditions of the eye and services relating to the treatment of these conditions.

Indigenous identification
No.

Socioeconomic identification
No.

Scope/target population
General Australian population.

Geographical coverage
All states and territories, Australia.

Time period covered

Publications and other outputs
• Publications, available on the AIHW website:
• Online data cubes on the AIHW website:

Access
Access is available through the AIHW or through the online data cubes.
Growing Up in Australia: the Longitudinal Study of Australian Children

Description of data source

Growing Up in Australia: the Longitudinal Study of Australian Children has been initiated and funded by the Australian Government Department of Families, Housing, Community Services and Indigenous Affairs (FaHCSIA) as part of its Stronger Families and Communities Strategy.

Purpose(s)

The Growing Up in Australia study aims to examine the impact of Australia’s unique social and cultural environment on the next generation to gain a further understanding of early childhood development, inform social policy debate and be used to identify opportunities for early intervention and prevention strategies in policy areas concerning children.

The longitudinal nature of the study aims to enable researchers to determine critical periods for the provision of services and welfare support and to identify the long-term consequences of policy innovations.

Collection methodology

Data are being collected over seven years from two cohorts every two years. The first main wave of data collection occurred in 2004, and two waves of data collection have now been completed. Wave 4 will be completed by mid-2010. The first cohort (Infant or B cohort) of approximately 5,000 children aged 0 to 1 years (when the sample was recruited in 2004) will be followed until they reach their mid-teens. The second cohort (Child or K cohort) comprising approximately 5,000 children aged 4 to 5 years (when the sample was recruited) will be followed until they reach 18 or 19 years of age. Study informants include the child (when of an appropriate age) and their parents, carers and teachers.

Data content

Participants provide information on child care; child care provider; child’s development; child’s diet and nutrition; child’s health; conception, pregnancy and birth; education; family demographics; finances; housing; neighbourhood; non-resident parent; paid work; parental health; parenting and social capital.

Data content related to eye health

Participants are asked to respond to the following:

• Does the study child have sight problems that have lasted, or are likely to last, for six months or more?
• Do the study child’s parents have sight problems that have lasted, or are likely to last, for six months or more?
• Are there concerns about the study child’s development, with regards to vision problems? (B cohort only)
• Do the study child’s siblings in the household have sight problems?
• Does the study child have ongoing problems with eyes or seeing properly?

Eye health-related data are self-reported.
Indigenous identification
The standard ABS classification for Indigenous identification is used.

Socioeconomic identification
Information is collected about the finances of the study child’s parents, their educational qualifications, their current work status and occupation, and home ownership. The ABS Socio-Economic Indexes for Areas (SEIFA) indices and other indicators are added to the data file for their neighbourhood.

Scope/target population
Children born from March 2003 to February 2004 (B cohort) and children born from March 1999 to February 2003 (K cohort). Children in about 40 per cent of remote postcodes were excluded.

Geographical coverage
All states and territories, Australia.

Time period covered

Publications and other outputs

Access
Data from Growing Up in Australia is warehoused at the Australian Institute of Family Studies and is available to researchers approved by the Department of Families, Community Services and Indigenous Affairs who must abide by strict security and confidentiality protocols. More information can be found on the Growing Up in Australia website: <www.aifs.gov.au/growingup/data.html> (viewed 17 August 2009).

Data management agency
Australian Institute of Family Studies
Level 20 485 La Trobe Street
Melbourne VIC 3000
(03) 9214 7888
lsacweb@aifs.gov.au
Higher Education Student Collection

Description of data source

The Higher Education Student Collection collects data from higher education providers in Australia.

It is a condition under the Higher Education Support Act 2003 (ss. 19–70) that a higher education provider must provide statistical information in respect of the provision of higher education by the provider.

Purpose(s)

The Higher Education Student Collection provides information required by the Australian Government Department of Education, Employment and Workplace Relations (DEEWR) as well as the Australian Tax Office (in relation to student loans). The collection provides data for analysis by government and non-government researchers which drives higher education policy formulation.

Collection methodology

Higher education providers report data in a series of data sets specified by DEEWR.

Data content

The Higher Education Student Collection collects data on enrolments, student load, courses, unit of study completions and course completions.

Data content related to eye health

The number of students studying the fields of study of optometry, optical technology and optical science not elsewhere classified, as classified by the Australian Standard Classification of Education (ASCE).

Indigenous identification

The standard ABS classification for Indigenous identification is used.

Socioeconomic identification

No.

Scope/target population

Higher education students from public and private higher education institutions that receive Higher Education Loan Program funding on behalf of students in the form of student Higher Education Loan Program loans from DEEWR.

Geographical coverage

All states and territories, Australia.

Time period covered

Since 1989 to present (ongoing collection).
Publications and other outputs

Available on the DEEWR website:

Access

Data from the most recent student collection are available on the DEEWR website in aggregated data sets:

Tabulations can also be generated to meet the needs of data users. Charges are incurred for these products.

Data management agency

Department of Education, Employment and Workplace Relations
GPO Box 9880
Canberra ACT 2601
<www.deewr.gov.au>
(02) 6240 9788
university.statistics@deewr.gov.au
Description of data source

The Home and Community Care (HACC) Minimum Data Set (MDS) is a collection of data about HACC clients (such as their age and living arrangements) and the amount and types of assistance being provided to them.

Purpose(s)

The objectives of the HACC MDS are to:

- provide HACC program managers with a tool to access data required for policy development, strategic planning and performance monitoring against agreed output/outcome criteria
- assist HACC service providers to provide high quality services to their clients by facilitating improvements in the internal management of HACC-funded service delivery
- facilitate consistency and comparability between HACC data and other aged, community and health data collections.

Collection methodology

Quarterly submission of data by service providers.

Data content

The HACC MDS collects information which includes a HACC linkage key, basic sociodemographic data (age, sex, country of birth, Indigenous status, main language spoken at home), care recipient circumstances (living arrangements, accommodation setting, carer details, pension status, state, suburb) and information about the service episode.

Data content related to eye health

The data element 'Assistance with goods and equipment' describes the number of goods/equipment received by the client during the HACC reporting period. It refers to the loan or purchase of goods and equipment to assist the person to cope with a disabling condition and/or maintain their independence. The data domain categories applicable to eye health are:

- support and mobility aids (white cane, sonic beam, laser cane, Optacon, guide dog)
- aids for reading (contact lenses, reading/magnifying glasses; Braille books/items/watch; books, large print/talking, cassette recorders/players; reading frame, page turners, reading aids not associated with loss of sight; other reading or sight aids; dwelling modification to aid communication-intercom; other aids for communication).

However, data specifically relating to eye health are limited as data are available only at the level of the aid categories above and are not further specified to indicate the specific aids provided. Data on support and mobility aids for the blind are only one component of the support and mobility category and therefore this category is not useful in its current form for identifying assistance that is solely related to eye health, as the collection does not distinguish an aid for a blind person from an aid for a sighted person. While the same is true for the category 'Aids for reading', the majority of aids in this category are aids to help people with impaired vision.
**Indigenous identification**
The standard ABS classification for Indigenous identification is used.

**Socioeconomic identification**
Area of residence (SLA) is collected.

**Scope/target population**
Elderly, frail or younger disabled people and their carers. The data collection covers all service recipients whether care recipient or clients with the exception of those who may be assisted anonymously or if unknown to the agency (for example, general advocacy for a group rather than a client, or group activities such as an information session where the agency does not have information about the participants).

**Geographical coverage**
All states and territories, Australia.

**Time period covered**
Since 2001–02 to present (ongoing collection).

**Publications and other outputs**

**Access**
Data are available as either aggregated data or unit record data. The conditions and protocols for access to HACC data are available on the DoHA website: <www.health.gov.au/internet/main/publishing.nsf/Content/hacc-mds-protocol.htm> (viewed 17 August 2009).

**Data management agency**
Australian Government Department of Health and Ageing
<www.health.gov.au>
acc@health.gov.au
Longitudinal Study of Indigenous Children

Description of data source

The Longitudinal Study of Indigenous Children (LSIC), also known as Footprints in Time, collects quantitative and qualitative information about the lives of Indigenous children. Interviews are conducted with the parents, carers and teachers of Indigenous children in urban, regional and remote communities around Australia.

Purpose(s)

The LSIC key research questions are:

• What do Indigenous children need to have the best start in life and to grow up strong?
• What is the difference between how Aboriginal and Torres Strait Islander children are raised compared to children of other cultural backgrounds?
• What is the importance of family, extended family and community in the early years of life?
• How can services and other types of support make a difference to the lives of Aboriginal and Torres Strait Islander children?

Collection methodology

The sample was recruited by locally-engaged, Indigenous Research Administration Officers using lists provided by Medicare and Centrelink of Indigenous children within the specified age cohorts, as well as their own connections to their local community. They also conducted community-engagement activities involving local community organisations, hospitals and child health centres to ensure elders and parents in the local area are aware of the study.

Data content

The project assesses:

• housing conditions
• child’s health (diet, nutrition, health conditions, injury, hospitalisations, sleeping patterns)
• parent’s health (health conditions, social and emotional wellbeing, smoking habits)
• child and family functioning (physical ability, language and development, parental warmth, major life events)
• demographic characteristics (parental language and culture, parental education, work, income, child support and maintenance, housing and mobility, child care and early education, activities).

Data content related to eye health

Parents are asked the following questions:

• In the past twelve months has the study child had any problems with eyes or eyesight?
• In the past twelve months have you had any problems with eyes or eyesight that made it difficult for you to look after the study child?

Teachers/centre-based carers are asked the following question:

• Does the study child have any vision problems that impact on his/her learning?
Eye health-related data are self-reported.

**Indigenous identification**

Yes.

**Socioeconomic identification**

Postcode, education of primary carer and household income.

**Scope/target population**


The sample is not designed to be representative of the Aboriginal and Torres Strait Islander child population.

**Geographical coverage**

Urban, rural and remote areas across Australia. Collection sites are located in all jurisdictions except the ACT and Tasmania.

**Time period covered**

Four annual waves from 2008–2011.

**Publications and other outputs**

Summary publications are planned to follow each data release.

**Access**

Organisations or persons needing access to the data will need to apply for a licence. More information on data access is available from the FaHCSIA website: <www.fahcsia.gov.au/lsic>.

**Data management agency**

FaHCSIA
LSIC Design and Analysis CW2
PO Box 7576
Canberra Business Centre ACT 2610
1800 106 235
LSIC@fahcsia.gov.au or LSICdata@fahcsia.gov.au
Macular Degeneration Foundation Database

Description of data source
The Macular Degeneration Foundation Database is a source of contact details for people within the community diagnosed with macular degeneration, their family, friends or carers, people with a general interest in macular degeneration and health-care professionals. The database is for the internal purposes of the Macular Degeneration Foundation. Within earlier collected data, it is not able to be determined whether a client has been diagnosed with macular degeneration or not.

Purpose(s)
The Macular Degeneration Foundation Database is an avenue through which people within the community diagnosed with macular degeneration or their carers/family can be contacted with educational information and invited to participate in surveys.

Collection methodology
People are placed on the database via phone enquiry, filling out a form at education seminars or by writing to the Macular Degeneration Foundation.

Data content
The information collected includes name and contact information; preferred choice of newsletter format; where they heard about the Macular Degeneration Foundation; if they have macular degeneration or if they are a family member, a friend or a person with a general interest. Specific details, such as the stage or type of the disease or treatment, are not collected. Health-care professionals are categorised into their field such as ophthalmologists, optometrists, GPs or aged care professionals.

Data content related to eye health
Clients registered with the Macular Degeneration Foundation who have been diagnosed with macular degeneration.
Eye health-related data are self-reported.

Indigenous identification
No.

Socioeconomic identification
No.

Scope/target population
Australians diagnosed with macular degeneration, their carers/family or people with a general interest.

Geographical coverage
All states and territories, Australia.
Time period covered
Since 2001 to present (ongoing collection).

Publications and other outputs

Access
The Macular Degeneration Foundation Database is for internal purposes of the Macular Degeneration Foundation only. The data are not released externally under any circumstances.

Data management agency
Macular Degeneration Foundation Limited
Suite 302, 447 Kent Street
Sydney NSW 2000
<www.mdfoundation.com.au>
(02) 9261 8900
info@mdfoundation.com.au
Medical Labour Force Survey

Description of data source
The AIHW Medical Labour Force Survey collects information on the characteristics of medical practitioners who are renewing their medical registration with medical boards in each state and territory.

Purpose(s)
The survey aims to provide comprehensive information about Australia’s medical labour force, such as demographics, employment characteristics, work locations and area of work. Along with other data sources, such as the Census and professional organisation data, the Medical Labour Force Survey is used in labour force projections and planning.

Collection methodology
Each state and territory medical board conducts an annual renewal of registration for medical practitioners eligible to practice in their jurisdiction. Questionnaires are sent to practitioners as part of the renewal process. Returned questionnaires are processed by the respective health authorities and then data is sent to the AIHW for collation into a national dataset.

Data content
The collection includes data elements about medical practitioners including medical qualifications, demographic information, employment status and type, location of work, hours worked and medical speciality.

Data content related to eye health
Characteristics of employed ophthalmologists.

Indigenous identification
Participants are recorded as Aboriginal, Torres Strait Islander or neither.

Socioeconomic identification
Postcode of usual residence and employment.

Scope/target population
Only practitioners who are on the register at the time of the survey, and who are required to renew their registration, receive a questionnaire. This excludes medical practitioners registering for the first time and those who are not required to renew their registration in the survey year. The estimated response rate in 2006 was 70%. Survey data is benchmarked to the number of registered practitioners in each state and territory.

Geographical coverage
All states and territories are included in the survey.
Time period covered
1997 to present (ongoing collection). From 2010, the National Health Workforce Taskforce will coordinate medical registration data under the National Registration and Accreditation Scheme.

Publications and other outputs
• Annual AIHW medical labour force reports are available on the AIHW website: <www.aihw.gov.au/labourforce/publications.cfm> (viewed 5 June 2009).

Access

Data management agency
Australian Institute of Health and Welfare
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
**Medicare Australia data**

**Description of data source**

Medicare Australia statistics cover services rendered on a ‘fee-for-service’ basis for which Medicare benefits were paid.

To be eligible for a Medicare rebate, the service must be listed in the Medicare Benefits Schedule (MBS).

**Purpose(s)**

Medicare statistics are a by-product of processing claims for Medicare benefits. The statistics are used by DoHA in undertaking policy development and review and program administration in relation to Medicare.

**Collection methodology**

Administrative by-product as a result of assessing Medicare claims.

**Data content**

Statistics can be compiled on: the number of services; fees charged; Schedule fees; benefits paid; bulk billing and Schedule fee observance down to MBS item level, by gender, age range, specialty of provider and by region of provider and region of patient.

**Data content related to eye health**

Eligible services provided by ophthalmologists and optometrists.

**Indigenous identification**

There is a voluntary Indigenous identifier in the Medicare claim record.

**Socioeconomic identification**

Statistics can be compiled by SEIFA index range, subject to availability of a postcode to SEIFA concordance file.

**Scope/target population**

All Australian residents are eligible for Medicare benefits. Medicare covers most medical and optometric services, and selected dental and allied health services. Excluded are details of services to public patients in hospital, to Veterans’ Affairs patients and some compensation cases.

**Geographical coverage**

All states and territories, Australia.

**Time period covered**

Since 1984 to present (ongoing collection).
Publications and other outputs


Access

Access to Medicare data is available through Medicare Australia.

Data management agency

Medicare Australia
PO Box 1001
Tuggeranong ACT 2901
<www.medicareaustralia.gov.au>
National Aboriginal and Torres Strait Islander Health Survey

Description of data source

The National Aboriginal and Torres Strait Islander Health Survey (NATSIHS) is a health survey of Indigenous Australians conducted by the ABS. This survey, which was conducted in remote and non-remote areas throughout Australia, was designed to collect a range of information from Indigenous Australians about health-related issues, including health status, risk factors and actions and socioeconomic circumstances.

Purpose(s)

The survey aims were to:

- provide broad information about the health of Indigenous Australians, by remoteness, and at the national and state/territory levels
- allow for the relationships across the health status, risk factors and health-related actions of Indigenous Australians to be explored
- provide comparisons over time on the health of Indigenous Australians
- provide comparisons with results for the non-Indigenous population.

Collection methodology

Data collection was undertaken by ABS interviewers using either computer-assisted interviewing (with a laptop) or pen and paper interviewing (paper questionnaire). Persons aged 18 years or more were interviewed personally, with the exception of persons who were too sick or otherwise unable to respond personally. Persons aged 15-17 years were interviewed (with consent from a parent or guardian) or their information was obtained from a parent or guardian. Information about children less than 15 years of age was provided by a responsible adult. A nominated household spokesperson was interviewed about household-level information.

While some information (for example, long-term conditions) was collected about Indigenous people of all ages, other data were only collected for relevant subpopulations, for example, self-assessed health for Indigenous people aged 15 years or over, and smoking and alcohol consumption for Indigenous people aged 18 years or over.

Data content

Information was collected about:

- health status, including long-term medical conditions and recent injuries
- use of health services such as consultations with health practitioners and visits to hospitals, health insurance (non-remote only), financial stress and other health-related actions
- health-related aspects of lifestyle, such as smoking, diet, exercise (non-remote only), substance use (non-remote only), alcohol consumption and financial stress
- demographic and socioeconomic characteristics.

Data content related to eye health

In the 2004–05 survey respondents provided the following information with regards to eye health:
• long-term eye and sight problems (including cataract, glaucoma, astigmatism, macular degeneration, presbyopia, short-sightedness, long-sightedness, blindness and other disorders, disturbances or diseases of the eye and adnexa)
• whether conditions were correctable by glasses or contact lenses
• whether they wear glasses or contact lenses
• whether eye/sight problem(s) were due to diabetes/high sugar level
• time since last consulted eye specialist or optometrist for diabetes-related eye condition
• time since last consulted eye specialist or optometrist for any eye/sight condition
• recent injuries to the eye (including type of injury, activity at time of injury, location of injury event and hospital/medical attendance).

Eye health-related data are self-reported.

Indigenous identification
Respondents are identified as being of Aboriginal, Torres Strait Islander or both Aboriginal and Torres Strait Islander origin.

Socioeconomic identification
The 2004–05 NATSIHS contains a range of socioeconomic indicators for persons and households.

Scope/target population
The NATSIHS 2004–05 was conducted on a sample of 10,439 Indigenous Australians (or about 1 in 45 of the total Indigenous population). The sample covered usual residents of private dwellings only.

Geographical coverage
Remote and non-remote areas of Australia.

Time period covered
The NATSIHS is to be repeated on a six yearly cycle. Previous surveys have been conducted for 1995, 2001 and 2004–05. In 2004–05 the NATSIHS was conducted over an eleven month period so that seasonal variations were captured. 1995 and 2001 data were collected as Indigenous components of the National Health Survey.

Publications and other outputs
• ABS 2006. National Aboriginal and Torres Strait Islander health survey 2004–05. Cat. no. 4715.0. Canberra: ABS.
• ABS 2006. National Aboriginal and Torres Strait Islander health survey 2004–05: users’ guide. Cat. no. 4715.0.55.004. Canberra: ABS.
• ABS 2006. National Aboriginal and Torres Strait Islander health survey, 2004–05: expanded confidentialised unit record file. Cat. no. 4715.0.55.001. Canberra: ABS.

Access

For users who wish to undertake more detailed analysis of the survey data, microdata from the 2004–05 NATSIHS are available in the form of an expanded CURF, National Aboriginal and Torres Strait Islander health survey, 2004–05: expanded confidentialised unit record file (cat. no. 4715.0.55.001). The expanded CURF is available via the ABS Remote Access Data Laboratory, which is a secure internet-based data query service.

Data management agency

Australian Bureau of Statistics
PO Box 10
Belconnen ACT 2612
<www.abs.gov.au>
client.services@abs.gov.au
National Data Set for Compensation-based Statistics

Description of data source

The National Data Set for Compensation-based Statistics is a national data set which provides details of accepted, pending and rejected workers compensation claims.

Purpose(s)

The purpose of the data set is to provide uniform, comparable national workers compensation data.

Collection methodology

Data are collected by jurisdictions on an ongoing basis, extracted annually and sent to the Safe Work Australia Council.

Before 1 July 2006, only records on accepted workers compensation claims were collected.

Data content

As of 1 July 2006, the following information is collected: record identifier, claim process details, claimant details, occurrence details, outcome of occurrence, compensation payments, payments for goods and services, non-compensation payments, premiums and remuneration, scheme assets and liabilities, scheme cost and disbursements, disputation and legal costs.

Data content related to eye health

Details of accepted, pending and rejected workers compensation claims for injuries or diseases affecting the eye.

Eye health-related data are medically verified.

Indigenous identification

No.

Socioeconomic identification

No.

Scope/target population

All new accepted, pending and rejected workers compensation claims made by or for an employee and which involved a death, a permanent incapacity or a temporary incapacity for which payments were made. It includes injuries and disease claims. It does not represent self-employed workers or employees of the defence forces.

Geographical coverage

All states and territories, Australia.

Time period covered

Since 1991 to present (ongoing collection).
Publications and other outputs

- Compensation-based statistics since 1997–98 are available in online data cubes (the National Online Statistics Interactive) at <nosi.ascc.gov.au> (viewed 19 August 2009).

Access

External access is available at <www.safeworkaustralia.gov.au> through the National Online Statistics Interactive.

Data management agency

Safe Work Australia
GPO Box 641
Canberra ACT 2601
<www.safeworkaustralia.gov.au>
(02) 6121 5317
National Elective Surgery Waiting Times Data Collection

Description of data source
The National Elective Surgery Waiting Times Data Collection (NESWTDC) is a collection of electronic confidentialised summary records for patients on elective surgery waiting lists and patients removed from waiting lists (for admission or another reason) in Australian public acute hospitals.

Purpose(s)
The NESWTDC is used to provide national information about the experience of patients on public hospital elective surgery waiting lists.

Collection methodology
Data are collected at each applicable hospital from patient administrative systems. Hospitals forward data to the relevant state or territory health authority on a regular basis (for example, monthly). State/territory health authorities forward these data to the AIHW annually.

Data content
The NESWTDC collects the following information about each elective surgery listing: listing date, removal date, waiting time at removal from elective surgery waiting list, indicator procedure, specialty of surgeon, clinical urgency, reason for removal from elective surgery waiting list, overdue patient, and extended wait patient.

Data content related to eye health
Elective surgery waiting time statistics where the indicator procedure is elected as ‘cataract extraction’ or the surgeon’s specialty is ‘ophthalmology’.
Eye health-related data are medically verified.

Indigenous identification
No.

Socioeconomic identification
No.

Scope/target population
The NESWTDC covers people added to elective surgery waiting lists in public acute hospitals only. However, some public patients treated under contract in private hospitals are included.

Geographical coverage
All states and territories, Australia, serviced by public acute hospitals.

Time period covered
Since 1995–96 to present (ongoing collection).
Publications and other outputs

A full list of elective surgery publications produced by the AIHW is available on the AIHW website: <www.aihw.gov.au/hospitals/waitingtime_data.cfm> (viewed 19 August 2009).

Access

The AIHW provides extracts of data from the NESWTDC on request. A charge may apply depending on the requirements and the complexity of the analysis undertaken. To utilise this data consultancy, email hospitaldata@aihw.gov.au.

Data management agency

Australian Institute of Health and Welfare
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
National Health Survey

Description of data source

The National Health Survey (NHS) is a triennial health survey which collects information about the health status of Australians, their use of health services and facilities, and health-related aspects of their lifestyle.

Purpose(s)

The survey is designed to obtain national benchmarks on a wide range of health issues and to enable changes in health to be monitored over time.

Collection methodology

Trained ABS interviewers conduct personal interviews with selected residents in sampled private dwellings. Persons included may vary from survey to survey: in the 2007–08 survey, one person aged 18 years and over in each dwelling was selected and interviewed about their health characteristics. Information was also collected about one child aged less than 18 years in sampled households; selected children aged 15–17 may have answered for themselves with parental consent, otherwise an adult, nominated by the household, was interviewed on behalf of the selected child.

Data content

The information collected in the NHS may vary from survey to survey. Information was collected in the 2007–08 survey about:

- the health status of the population, including long-term medical conditions experienced and recent injuries
- health-related aspects of people’s lifestyles, such as smoking, diet, exercise and alcohol consumption
- use of health services such as consultations with health practitioners, visits to hospital, days away from work, health insurance and other actions people have recently taken for their health
- demographic and socioeconomic characteristics.

Data content related to eye health

In the 2007–08 survey respondents were specifically asked about:

- long-term eye and sight problems (including cataract, glaucoma, astigmatism, macular degeneration, presbyopia, short-sightedness, long-sightedness, blindness and other disorders, and disturbances or diseases of the eye and adnexa)
- whether conditions are correctable by glasses or contact lenses
- whether they wear glasses or contact lenses
- whether eye/sight problem(s) was due to diabetes/high sugar level
- time since last consulted eye specialist or optometrist for diabetes-related eye condition
- time since last consulted eye specialist or optometrist for any eye/sight condition
- recent injuries to the eye (including type of injury, activity at time of injury, location of injury event and hospital/medical attendance).
Eye health-related data are self-reported.

**Indigenous identification**

No, data are unable to be disaggregated by Indigenous identification. A National Aboriginal and Torres Strait Islander Health Survey was conducted at the same time as the 2004–05 NHS.

**Socioeconomic identification**

The 2007–08 NHS contains a range of socioeconomic indicators for persons and households.

**Scope/target population**

The NHS 2007–08 was conducted on a sample of 15,800 dwellings in urban and rural areas in all states and territories; very remote areas were excluded.

The sample is designed to be representative of the population living in private dwellings other than those in very remote areas.

**Geographical coverage**

For the 2007–08 NHS, both urban and rural areas in all states and territories were included, but very remote areas of Australia were excluded.

**Time period covered**

The NHS is performed periodically: 1977–78, 1983, 1989–90, 1995, 2001, 2004–05 and 2007–08. In 2007–08 the NHS was conducted over an eleven month period so that seasonal variations were captured.

**Publications and other outputs**

National Health Survey 2007–08:


The survey user guide and state-specific results are also available on the ABS website, as well as other topic-specific publications.

**Access**

For users who wish to undertake more detailed analysis of the survey data, two CURFs are available. A basic CURF is available on CD-ROM, while an expanded CURF (containing more detailed information than the basic CURF) is accessible through the ABS Remote Access Data Laboratory (RADL) system.

**Data management agency**

Australian Bureau of Statistics
Locked Bag 10
Belconnen ACT 2616
<www.abs.gov.au>
client.services@abs.gov.au
National Hospital Morbidity Database

Description of data source
The National Hospital Morbidity Database (NHMD) is a collection of electronic confidentialised summary records for patients admitted in public and private hospitals in Australia.

Purpose(s)
The purpose of the NHMD is to provide information about care provided to admitted patients in Australian hospitals.

Collection methodology
Data are collected at each hospital from patient administrative and clinical record systems. Hospitals forward data to the relevant state or territory health authority on a regular basis (for example, monthly). State/territory health authorities forward these data to the AIHW annually.

Data content
The information collected falls into the following categories:
- establishment data (state/territory of the hospital; sector; rural, remote and metropolitan areas; accessibility/remoteness index of Australia; remoteness area)
- demographic (sex; date of birth, age at admission, country of birth; Indigenous status; state and local area of residence; remoteness area of residence)
- administrative variables (funding source; mode of admission; mode of separation; patient election status; urgency of admission)
- length of stay data (admission and separation dates; leave days; number of patient days; intended length of stay; same day flag)
- clinical variables (care type; principal diagnosis; additional diagnoses; morphology of neoplasm; procedures; external cause of injury or poisoning; place of occurrence of external cause; activity when injured; diagnosis-related group; major diagnostic category).

Data content related to eye health
Hospital separations with diagnoses for diseases and disorders of the eye, and procedures performed on the eye, in Australian hospitals.
Eye health-related data are medically verified.

Indigenous identification
Indigenous identification is based on the ABS standard for Indigenous status.

Socioeconomic identification
SEIFA based on provided SLA of individual.
Scope/target population
Episodes of care for admitted patients in all public and private acute and psychiatric hospitals, free-standing day hospital facilities and alcohol and drug treatment centres in Australia. Hospitals operated by the Australian Defence Force, corrections authorities and in Australia’s off-shore territories may also be included. Hospitals specialising in dental, ophthalmic aids and other specialised acute medical or surgical care are included.

Geographical coverage
Data are collected for almost all public and private hospitals for all states and territories of Australia.

Time period covered
Since 1993–94 to present (ongoing collection).

Publications and other outputs
- Data sourced from the NHMD contributes to many publications, reports and articles, both within the AIHW and by external researchers.

Access
The AIHW provides extracts of data from the NHMD on request. A charge may apply depending on the requirements and the complexity of the analysis undertaken. To utilise this data consultancy, email hospitaldata@aihw.gov.au.

Data management agency
Australian Institute of Health and Welfare
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
National Indigenous Eye Health Survey

Description of data source
The National Indigenous Eye Health Survey (NIEHS) is designed to provide essential baseline evidence to be used to plan and prioritise effective delivery of eye care for Indigenous Australians. It assesses the prevalence and main causes of vision impairment, as well as the utilisation of eye care services, barriers to health and the impact of vision impairment on people.

Purpose(s)
The objectives of the NIEHS are to:
• determine the prevalence and main causes of vision impairment (cataract, macular degeneration, glaucoma, diabetic retinopathy, refractive error and trachoma) in Indigenous Australians
• evaluate access to, and utilisation of, health-care services in Indigenous communities
• provide the evidence base for the planning of effective eye care programs for Indigenous people.

Collection methodology
Communities and/or geographic areas were randomly selected according to their location in urban, regional, remote or very remote areas, so that a comprehensive picture of vision loss could be compiled. The total sample size was 30 sites, with at least 50 adults (40 years and over) and 50 children (5 to 15 years) in each, giving a total sample of 3000 people. Questionnaires were self-completed where possible, otherwise a health worker or survey team member assisted.

Data content
Information was collected about:
• demographic information
• medical history and family eye health history
• utilisation of eye health services
• eye health status through examinations
• quality of life for people with impaired vision.

Data content related to eye health
For adults, eye examinations collected data on visual acuity using a standard E-chart, refractive error using an autorefractor, visual field analysis using frequency-doubling technology testing, neural and retinal disease testing (e.g. glaucoma, diabetic retinopathy), and presence of trachoma.
For children, data was collected on visual acuity and presence of trachoma.
Eye health-related data are medically verified.

Indigenous identification
All participants identified themselves as Aboriginal or Torres Strait Islander.
Socioeconomic identification
All participants recorded their address; adult participants recorded their highest educational qualification.

Scope/target population
Participants were eligible if they were Indigenous, lived within the boundaries of the randomly selected areas and were greater than or equal to 40 years of age or between 5 and 15 years of age.
The sample is designed to be representative of the Indigenous population.

Geographical coverage
30 urban and rural sites in all states and territories.

Time period covered
Data collection ran from February 2008 to December 2009.

Publications and other outputs

Access
Access to the data may be available with consent from the Centre for Eye Research Australia.

Data management agency
Centre for Eye Research Australia
32 Gisborne Street
East Melbourne VIC 3002
<www.cera.org.au>
(03) 9929 8360
cera-info@unimelb.edu.au
National Mortality Database

Description of data source
The National Mortality Database contains cause-of-death information for all deaths in Australia registered in Australia from 1964 onwards.
Death registration has been compulsory in all states and territories since the mid-1850s and this information is registered with the respective Registrars of Births, Deaths and Marriages.

Purpose(s)
To monitor the cause of all deaths in Australia.

Collection methodology
Information about a death is recorded on the death certificate and the death information form. Information is provided to the AIHW by the Registrars of Births, Deaths and Marriages in each state and territory and coded nationally by the ABS.
Additional demographic and administrative information is collected by funeral directors, usually through the next of kin or a person known to the deceased.

Data content
The collection includes data elements such as date of death, date of birth, underlying cause of death, multiple causes of death (1997 onwards), place of death by state and local government area and Indigenous status (for later years).

Data content related to eye health
Cases where a cause of death is associated with infections, cancers, disorders or injury of the eye.
Eye health-related data are medically verified.

Indigenous identification
Based on the ABS standard for Indigenous identification. The quality of Indigenous identification is not reliable for all states and territories.

Socioeconomic identification
SEIFA based on provided SLA of deceased.

Scope/target population
All deaths occurring in Australia. Deaths of Australians occurring overseas are not within scope of the collection.

Geographical coverage
All states and territories, Australia.

Time period covered
Since 1964 to present (ongoing collection).
Publications and other outputs


Access

Summary-level data are available publicly in the AIHW General record of incidence of mortality: interactive Excel workbooks (GRIM books), online at: <www.aihw.gov.au/mortality/data/grim_books.cfm> (viewed 16 November 2009). Data is also available as a consultancy. Email mortality_info@aihw.gov.au.

Data management agency

Australian Institute of Health and Welfare
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
National Outpatient Care Database

Description of data source
The National Outpatient Care Database (NOCD) is a collection of electronic aggregate records of the number of services provided in non-admitted non-emergency departments of principal referral and specialist women’s and children’s hospitals and large hospitals.

Purpose(s)
The NOCD is used to provide national information about the volume of services provided by non-admitted non-emergency departments of principal referral and specialist women’s and children’s hospitals and large hospitals.

Collection methodology
Data are collected at each applicable hospital from patient administrative systems. Hospitals forward data to the relevant state or territory health authority on a regular basis (for example, monthly). State/territory health authorities forward these data to the AIHW annually.

Data content
The NOCD collects the following information: organisation identifier, outpatient clinic type, number of group sessions and number of occasions of service.

Data content related to eye health
Outpatient statistics are available where the outpatient clinic service is elected as ‘ophthalmology’.

Eye health-related data are medically verified.

Indigenous identification
Not applicable.

Socioeconomic identification
Not applicable.

Scope/target population
The scope includes all arrangements made to deliver specialist care to non-admitted, non-emergency department patients whose treatment has been funded through the hospital, regardless of the source from which the hospital derives these funds.

Geographical coverage
All states and territories, Australia, serviced by principal referral and specialist women’s and children’s hospitals and large hospitals.

Time period covered
Since 2005–06 to present (ongoing).
Publications and other outputs

Australian hospital statistics has included NOCD data since 2005–06. A full list of publications is available on the AIHW website:

Access

The AIHW will be able to provide extracts of data from the NOCD on request. A charge may apply, depending on the extract requirements and the complexity of the analysis undertaken. To utilise this data consultancy, email hospitaldata@aihw.gov.au.

Data management agency

Australian Institute of Health and Welfare
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
National Sun Survey

Description of data source
The National Sun Survey provides information about people’s attitude and behaviour regarding ultraviolet (UV) radiation exposure.

Purpose(s)
The survey aims to provide comprehensive national data on Australian adolescents’ and adults’ risk and preventive behaviours for sun exposure, as well as sunburn history. The survey primarily aims to provide an information base for skin cancer prevention studies.

Collection methodology
Interviews are conducted in weekly cross-sectional telephone surveys over 8 weeks of summer. The interviews are held on Monday and Tuesdays evenings, with respondents asked to report on their outdoor activities for the previous Saturday and Sunday. The 2006–07 survey included approximately 5,000 adults (18–69 years) and 700 adolescents (12–17 years).

Data content
Respondents are asked in detail about their outdoor activities, sun protection behaviour and sunburn incidence during peak UV times during the previous weekend. Demographic characteristics are also recorded.

Data content related to eye health
Respondents are asked whether they wore sunglasses and/or hats while outdoors.

Indigenous identification
No.

Socioeconomic identification
Postcode, income and occupation.

Scope/target population
All persons resident in Australia over 12 years of age. The sample is designed to be representative of the Australian population.

Geographical coverage
All states and territories, Australia.

Time period covered
Two surveys have been completed in 2003–04 and 2006–07. Additional data on sunglasses use is also available for 2008–09.

Publications and other outputs


Access

Access to the data is available pending permission from the Cancer Council Australia and Cancer Australia.

Data management agency

Cancer Council Australia
GPO Box 4708
Sydney NSW 2001
<www.cancer.org.au>
(02) 8063 4100
info@cancer.org.au
National Trachoma Surveillance and Reporting Unit

Description of data source
The National Trachoma Surveillance and Reporting Unit (NTSRU) is the central collection point for trachoma data collected from state and territories where trachoma has been identified as present.

Purpose(s)
The NTSRU was established in 2006 to ensure compliance with the Communicable Disease Network Australia’s (CDNA) guidelines for the public health management of trachoma. The NTSRU analyses and reports on trachoma prevalence and control activities and monitors antibiotic resistance to azithromycin.

Collection methodology
All jurisdictions use the WHO’s trachoma grading criteria to detect and grade trachoma. Each jurisdiction categorises communities as being ‘at risk’ or ‘not at risk’ for trachoma, and further categorised them as having been ‘screened’ or ‘not screened’. CDNA guidelines state that screening in ‘at risk’ communities should be conducted annually until the prevalence of active trachoma is less than 5% for five consecutive years.

Data on antibiotic resistance in Indigenous communities since 2007 is collected by the Institute of Medical Veterinary Science, Northern Territory Government Pathology Service and the Western Diagnostics Pathology Service.

Data content
The information collected includes:
• trachoma prevalence among Indigenous children
• prevalence and treatment for trichiasis among Indigenous adults
• treatment strategy used by each community to distribute azithromycin
• implementation of trachoma control activities relating to the SAFE strategy (surgery for trichiasis, antibiotic treatment, facial cleanliness and environmental improvement).

Data content related to eye health
Information is collected on the prevalence of:
• trachomatous inflammation—follicular
• trachomatous inflammation—intense
• trachomatous scarring
• trachomatous trichiasis
• treatment distribution for trachoma
• implementation of components of SAFE strategy.

Indigenous identification
Yes.
Socioeconomic identification

No.

Scope/target population

CDNA guidelines recommend that surveillance focus on Indigenous children aged 1 to 9 years. Additionally, it is recommended that in communities ‘at risk’ for trachoma, all Indigenous adults aged 40 to 54 years should be examined every two years and those aged 55 years and over should be examined annually for trichiasis.

Geographical coverage

Trachoma was identified in Indigenous communities in the NT, South Australia (SA) and WA.

Time period covered

November 2006 to present (ongoing collection).

Publications and other outputs

An annual Trachoma Surveillance Report is submitted to DoHA, Office for Aboriginal and Torres Strait Islander Health. An annual summary is submitted to the journal *Communicable Disease Intelligence*.


Access

Access to the data collection in restricted.

Data management agency

Centre for Eye Research Australia (CERA)
32 Gisborne Street
East Melbourne VIC 3002
<www.cera.org.au>
(03) 9929 8360
btellis@unimelb.edu.au
Nursing and Midwifery Labour Force Survey

Description of data source
The Nursing and Midwifery Labour Force Survey collects information on the characteristics of those in nursing professions in Australia.

Purpose(s)
The survey aims to provide comprehensive information about Australia’s nursing labour force, such as demographics, employment characteristics, work locations and area of work. The Nursing and Midwifery Labour Force Survey is used in labour force projections and planning.

Collection methodology
Each state and territory nursing/midwifery board conducts an annual renewal of registration and enrolment. Questionnaires are sent to nurses and midwives as part of the renewal process. Returned questionnaires are processed by the respective health authorities, and then data is sent to the AIHW for collation into a national dataset.

Data content
The collection includes data elements concerning registration/enrolment details, demographic characteristics, current study, employment status and type, place of work, hours of work, and area of medical practice.

Data content related to eye health
Characteristics of ophthalmic nurses.

Indigenous identification
Participants are recorded as Aboriginal, Torres Strait Islander or neither.

Socioeconomic identification
Postcode of usual residence and employment.

Scope/target population
All nurses who were registered or enrolled with the nursing/midwifery board in each state/territory at the time of the survey. It excludes nurses who registered or enrolled for the first time in the year prior to the survey and those who were not required to renew their registration. The response rate in 2005 was 55%.

Geographical coverage
All states and territories, Australia.

Time period covered
1995 to present (ongoing collection). From 2010, the National Health Workforce Taskforce will coordinate medical registration data under the National Registration and Accreditation Scheme.
Publications and other outputs


Access


Data management agency

Australian Institute of Health and Welfare
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
Office for Aboriginal and Torres Strait Islander Health Service Activity Reporting system

Description of data source
The Office for Aboriginal and Torres Strait Islander Health (OATSIH) (in conjunction with the National Aboriginal Community Controlled Health Organisation) Service Activity Reporting (SAR) system contains service-level data from all Aboriginal and Torres Strait Islander primary health care services in Australia that are in receipt of Australian Government funding.

Purpose(s)
Administrative collection of Aboriginal health services which receive federal funding.

Collection methodology
Service-level data on health care covering a twelve-month period are collected by questionnaire and completed by Aboriginal and Torres Strait Islander primary health care service staff.

Data content
The data collected in the SAR cover three main areas: health oriented activities, staffing and client contacts. Many of the services receive additional funds from sources other than the Australian Government. The data collected in the SAR relate to health activities resulting from all funding sources.

Data content related to eye health
Information is provided on:
- number of client contacts with eye health workers
- full time equivalent eye health staff
- number of visiting eye health workers
- whether optometry/ophthalmology specialist eye services were provided
- whether eye disease screening was provided.

Indigenous identification
Yes.

Socioeconomic identification
No.

Scope/target population
All Aboriginal and Torres Strait Islander primary health care services that receive Australian Government funding to facilitate access to primary health care, including health promotion, dental care and counselling services (151 services nationwide in 2005-06).

Geographical coverage
All states and territories, Australia.
Time period covered
1997 to present (ongoing collection).

Publications and other outputs
A SAR ‘Key results’ report is produced annually and can be found on the DoHA website: <www.health.gov.au>.

Access
The SAR database is not distributed externally. However, de-identified analyses of the data may be provided upon request to the data manager.

Data management agency
Office for Aboriginal and Torres Strait Islander Health (OATSIH)
Australian Government Department of Health and Ageing
GPO Box 9848
Canberra ACT 2601
<www.health.gov.au>
1800 678445
oatsih.enquiries@health.gov.au
Optometrists Association Australia Optometrists Database

Description of data source
The Optometrists Association Australia Optometrists Database (OAAOD) includes both members and non-members of the Optometrists Association Australia and is continually updated with information received from registration boards, state divisions of the association and optometrists. The database is able to be manipulated to eliminate duplications, accounting for optometrists who are registered in more than one state, and to determine how many optometrists registered in Australia are in clinical practice.

Purpose(s)
The purpose of the OAAOD is to provide information about the workforce of the optometric sector.

Collection methodology
Approximately every 2 years, the association mails a survey form to all optometrists registered in Australia to correct errors and omissions in the database.

Data content
The most recent survey collected information on age, gender, practice addresses, hours spent practising optometry, states where registered, employment type (clinical practice/academia/research etc.) and employment status (employee, sole owner or partner).

Data content related to eye health
Statistics on the optometric labour force.

Indigenous identification
Not applicable.

Socioeconomic identification
Not applicable.

Scope/target population
Optometrists registered to practice in Australia.

Geographical coverage
All states and territories, Australia.

Time period covered
Studies have been conducted in 1988, 1992, 1996 and 2005.

Publications and other outputs
Access

The OAAOD is for the internal purposes of the Optometrists Association Australia and is not for public access.

Data management agency

Optometrists Association Australia
PO Box 185
Carlton South VIC 3053
<wwww.optometrists.asn.au>
(03) 9663 6833
oaanat@optometrists.asn.au
**Optometrist registration boards**

**Description of data source**

State/territory optometrist registration boards are currently established under legislation in their state/territory. In order to practice as an optometrist in a state/territory, an optometrist must currently comply with the conditions of registration for that particular state/territory. Renewal of registration occurs every financial year.

This arrangement is scheduled to end from July 2010 when optometrists will be included in the National Registration and Accreditation Scheme (NRAS).

**Purpose(s)**

State/territory optometrist registration boards, among many things, have the functions to:

- administer the registration of optometrists
- keep a register of, and records relating to, registrants
- supervise the practice of optometry in a particular state/territory
- monitor standards of optometrical education and training.

**Collection methodology**

Optometrists wishing to apply for registration to practice optometry in a state/territory must apply through the state/territory’s board and comply with the board’s conditions of registration. The board verifies the applicant’s identity and qualifications and officially registers the applicant. A registered optometrist must renew their registration annually. An optometrist may be registered to practice in more than one state/territory.

**Data content**

Name, address, phone number, qualifications, employment status.

**Data content related to eye health**

The number of optometrists registered to practice in each state/territory.

**Indigenous identification**

Not applicable.

**Socioeconomic identification**

Not applicable.

**Scope/target population**

A graduate of an institution accredited by the Council of Optometry Registration Authorities or overseas-registered optometrists who have undertaken a qualifying examination conducted by the Optometry Council of Australia and New Zealand.

**Geographical coverage**

All states and territories, Australia.
Time period covered
Varies among states/territories.

Publications and other outputs
Not applicable.

Access
Not applicable.

Data management agency
Each registration board has an individual state/territory office.
Pharmaceutical Benefits Scheme data

Description of data source

The Pharmaceutical Benefits Scheme (PBS) is an Australian Government initiative that subsidises around 80% of prescriptions dispensed in Australia through ‘community pharmacies’. It does not cover pharmaceuticals delivered to most inpatients in hospitals or delivered through alternative means such as through Aboriginal and Torres Strait Islander medical services.

Purpose(s)

Payment system.

Collection methodology

Data are collected on a bill payment basis and patient refund through Medicare Australia.

Data content

The PBS collects the following data:

- dates of prescription, supply and processing
- identifiers for doctors, patients and pharmacists
- basic patient demographic details (date of birth, gender, postcode and patient eligibility)
- pharmacy details
- prescriber details (main specialty)
- drug item number, benefit paid, patient contribution etc.

Data content related to eye health

Data on medicaments used in treating ophthalmological problems. It is also possible to identify other drugs throughout the schedule that may be related to eye health and identify the specialty of the prescribing doctor or optometrist.

Eye health-related data are medically verified.

Indigenous identification

Indigenous people can elect to have their data used for research and statistical purposes. Approximately 160,000 Indigenous people have given their consent.

As stated above, pharmaceuticals delivered through Aboriginal and Torres Strait Islander medical services are not included in PBS. These amount to approximately $23 million expenditure across all drug types.

Socioeconomic identification

There is the potential to match the postcode to the SEIFA index.

Scope/target population

All Australians, though the difference in subsidy depending on eligibility means that data for health-care card holders is likely to be more complete than for other Australians.
Geographical coverage
All states and territories, Australia.

Time period covered
While the PBS has been in place since 1949, the data are only available in machine readable form from July 1990 (ongoing collection).

Publications and other outputs
- Various DoHA publications, for example:

Access
Access to the data is controlled by the National Health Act 1953 (as amended). Data can be requested through the Data and Estimates Unit, DoHA.

Data management agency
Pharmaceutical Benefits Division
Australian Government Department of Health and Ageing
GPO Box 9848
Canberra ACT 2601
<www.health.gov.au>
(02) 6289 7945
pbsstats@health.gov.au
Royal Australian and New Zealand College of Ophthalmologists ophthalmology workforce database

Description of data source

The ophthalmology workforce database provides statistics on the numbers and distribution of the ophthalmology workforce in Australia and New Zealand, as collected by the Royal Australian and New Zealand College of Ophthalmologists (RANZCO).

Purpose(s)

The database is used for the purposes of planning, continuing professional development, referral and membership services.

Collection methodology

Information is collected from members of RANZCO.

Data content

Name, address, age, gender, qualifications, activities.

Data content related to eye health

Service provision.

Indigenous identification

Not applicable.

Socioeconomic identification

Not applicable.

Scope/target population

Australian and New Zealand ophthalmologists. The database represents 95% of the Australian and New Zealand ophthalmological workforce.

Geographical coverage

Australia and New Zealand (including some overseas members).

Time period covered

Since 2004 to present.

Publications and other outputs

• Statistical analysis of the database is published in the membership journal.

Access

Access to the ophthalmology workforce database may be obtained through RANZCO.
Survey of Disability, Ageing and Carers

Description of data source

This survey collects information on three population groups: people aged 60 and over, people with disabilities and their carers. There are other sources of information about services provided to these groups, but this survey is the only source of information on the assistance requirements of those groups, the extent to which these requirements are met and the characteristics of those with unmet need. It also provides information on participation in economic and community activities. The survey is the major source of national statistics on carers and primary carers: numbers and characteristics of carers, care relationships, activities for which informal care is provided and, for primary carers, support available and required and the effects of the caring role on their lives.

Purpose(s)

The aims of the survey undertaken in 2003 were to:

• measure the prevalence of disability in Australia
• measure the need for support of older people and those with a disability
• provide a demographic and socioeconomic profile of people with disabilities, older people and carers compared with the general population
• estimate the number of and provide information about people who provide care to older people and people with disabilities.

Collection methodology

Different collection methodologies were used for people in households and people in care accommodation:

• Trained interviewers conducted computer-assisted personal interviews for the household component of the survey. A set of screening questions was asked of a responsible adult in the household to establish whether the household included people with a disability, people aged 60 years or over or people who were carers. Individuals identified as being in one of these populations were interviewed personally, where possible. Proxy interviews were conducted for children aged less than 15, children aged 15–17 whose parents did not permit them to be personally interviewed and those whose disability prevented them from having a personal interview.

• The care accommodation component of the survey was collected in two steps using a mail-based methodology sent to directors of selected establishments. First, the name of a contact officer, the current number of occupants and the type of establishment were established in a contact information form. Second, the contact officer selected occupants, based on provided instructions. A separate questionnaire was completed for each selected occupant meeting the coverage requirements. The range of data collected in this component was smaller than in the household component.

Data content

The survey collected the following information:

• People with a disability were asked about their long-term health conditions, need for and receipt of assistance for self-care, mobility, communication, cognition or emotion, health care, housework, property maintenance, meal preparation, paperwork and
transport activities; use of aids; equipment such as wheelchairs and hearing aids; and participation in community activities. They were also asked questions relating to computer and internet use and participation in community activities. Those aged 5–20 years (or their proxies) were asked about schooling restrictions, and those aged 15–64 years about employment restrictions.

• Older people (i.e. those aged 60 years and over) were asked about their need for and receipt of assistance with housework, home maintenance, meal preparation, paperwork, and transport activities; and participation in community activities. They were also asked questions relating to computer and internet use and participation in community activities.

• People who care for persons with a disability and older people were asked about the type of care they provide; the support available to them; and the characteristics of carers and some effects that the caring role has on their lives. A self-enumeration form was completed by these people during the interview which collected information about their attitudes to, and experience of, their caring role.

• Basic demographic and socioeconomic information was collected for all other people. Most of this information was provided by a responsible adult in the household rather than by each person individually.

Data content related to eye health

Participants provided information about the following issues:

• whether person has loss of sight—no loss, loss of sight corrected by wearing glasses or contact lenses, partial loss of sight not corrected by wearing glasses or contact lenses, total loss of sight

• disability type/main disability/disability that causes most problems—loss of sight

• disability type as a result of head injury—loss of sight

• disability type as a result of stroke—loss of sight

• disability type as a result of other brain damage—loss of sight

• type of long-term condition reported—retinal disorders/defects, glaucoma, sight loss, other diseases of the eye and adnexa

• condition(s) producing main disability (multiple response)—retinal disorders/defects, glaucoma, sight loss, other diseases of the eye and mastoid

• main condition causing loss of sight

• type of difficulty experienced at school or institution because of condition—hearing or sight problems

• reasons for inability to use some or all forms of public transport—sight problems.

Eye health-related data are medically verified and self-reported.

Indigenous identification

Not included. Detailed data on disability in the Australian Aboriginal and Torres Strait Islander population are contained in the National Aboriginal and Torres Strait Islander Social Survey 2002.
**Socioeconomic identification**

Data items related to socioeconomic factors: main source of income; cash income amount (personal, family, income unit and household level); cash income decile; receipt of government benefits (such as Disability Support Pension, Carers Allowance or Parenting Payment); labour force participation; education levels achieved.

**Scope/target population**

The survey included people in both private and non-private dwellings, including people in care accommodation establishments but excluding those in jails and correctional institutions. The sample comprised of 36,241 people in households (both private and non-private) and 5,145 people in cared accommodation. Visitors to private dwellings, certain diplomatic personnel of overseas governments, overseas residents in Australia and members of non-Australian defence forces stationed in Australia (and their dependants) were also excluded.

**Geographical coverage**

People in both urban and rural areas in all states and territories, except for those living in remote and sparsely settled parts of Australia.

**Time period covered**

1998 and 2003. The next survey was conducted between April and December 2009.

**Publications and other outputs**


**Access**

All organisations and individuals within organisations who wish to access the CURF are required to read the Responsible access to ABS CURFs training manual and sign an undertaking to abide by the legislative restrictions on use. These documents are available to read or download on the access to ABS CURFs page on the ABS website: <www.abs.gov.au>. The Australian Statistician’s approval is required for release of the CURF to an organisation and individuals. Use of the data for unauthorised purposes may render the organisation and/or user liable to severe penalties. Advice about the propriety of any particular intended use of the data is available from curf.management@abs.gov.au.

**Data management agency**

Australian Bureau of Statistics
Locked Bag 10
Belconnen ACT 2616

<www.abs.gov.au>

client.services@abs.gov.au
**Treatment Accounts System**

Description of data source
The Treatment Accounts System is administered by Medicare Australia on behalf of the Department of Veterans’ Affairs (DVA). It is the system which will pay accounts for most medical and hospital services. A major exclusion is public hospital expenditure. It captures all relevant details of the services rendered.

Purpose(s)
Payment of health services for veterans with treatment cards, i.e. gold, white or orange cards. It is DVA’s primary source of health services data.

Collection methodology
Data on services rendered and paid are collected from payment invoices and associated documentation, following accepted health data standards.

Data content
The list of data content is available through metadata and data dictionaries. It covers the full spectrum of data on health services.

Data content related to eye health
In addition to the data gathered through the claims, which use service type, item type and description codes, DVA also records the medical condition in respect of its Disability Pension claims using the ICD-10 codes. Eye health-related data are medically verified.

Indigenous identification
No.

Socioeconomic identification
It can be linked to DVA’s beneficiary population which contains some socioeconomic data (for example, income and assets).

Scope/target population
As covered under the *Veterans’ Entitlements Act 1986*, veterans with treatment cards, i.e. gold, white or orange cards.

Geographical coverage
All states and territories, Australia.

Time period covered
Available from the late 1980s. However, the recent database holds readily accessible information since 2001.
Publications and other outputs

No external publications. However, the data are made available to researchers for health studies.

Access

Access to the Treatment Accounts System is available, subject to privacy principles and the consent of DVA’s Ethics Committee.

Data management agency

Department of Veterans’ Affairs
PO Box 21
Woden ACT 2606
<www.dva.gov.au>
Regional data sources

The 45 and Up Study (NSW)

Description of data source
The 45 and Up Study is a long-term health study in Australia of men and women aged 45 and over. A key feature of the 45 and Up Study is its linkage to a range of established health data sets such as GP services, medication use, procedures and hospitalisations.

Purpose(s)
The primary aim of the 45 and Up Study is to investigate the main factors that determine healthy ageing in the population.

Questions are included relating to common diseases and health conditions that people may encounter as they grow older, including cancer, cardiovascular disease, obesity, arthritis, diabetes and depression.

Collection methodology
Over 250,000 men and women aged 45 and over have been randomly selected across NSW and invited to join the 45 and Up Study. They received an information leaflet and a 45 and Up Study baseline questionnaire for either men or women. They will receive a follow-up questionnaire every five years.

Both men’s and women’s questionnaires have the same number and type of questions about health, lifestyle and experience. All questions have been carefully selected by leading NSW health professionals and researchers.

Participants are asked to complete the questionnaire as accurately as possible and to sign the accompanying consent form before enclosing it in the provided envelope and returning it. In addition to the follow-up questionnaires, their health will be followed via linkages to already routinely collected data sources such as records of hospital admissions and medical and surgical procedures, registers of deaths, cancers and infectious diseases and use of medications and aged care facilities.

Participation in the study is entirely voluntary.

Data content
The 45 and Up Study questionnaire contains demographic, social, health, lifestyle, screening and other variables. Copies of questionnaires are available on the 45 and Up website: <www.45andup.org.au>. A wide range of other variables is available via data linkage. Biological samples will be collected and relevant data will be stored in study records. In addition, participating studies will be required to send key data collected or analysed by them to the 45 and Up Study for storage.

Data content related to eye health
The baseline questionnaire gathers information on:

- self-rated eyesight (excellent, very good, good, fair or poor)
- any operations
- serious illnesses
- medication use.
Records are able to be linked to relevant eye-related data on GP services, medication use, procedures and hospitalisations.
Note that due to the nature of the self-administered questionnaire, it could be more difficult for people with significant vision problems to participate in the 45 and Up Study.
Eye health-related data are medically verified and self-reported.

**Indigenous identification**
Participants are asked if they are Aboriginal and/or Torres Strait Islander origin.

**Socioeconomic identification**
Participants are asked about their address, highest educational qualification achieved, household income and current work status.

**Scope/target population**
Men and women aged 45 or over, living in NSW and on Medicare Australia’s records. The eligible population is highly representative of the Australian population aged 45 and over.

**Geographical coverage**
All of NSW.

**Time period covered**
The recruitment of participants commenced in February 2006 and will continue until at least 2011.

**Publications and other outputs**

**Access**
Individuals or groups may access the 45 and Up Study dataset, subject to appropriate approvals.

**Data management agency**
The Sax Institute
PO Box 123
Broadway NSW 2007
[www.saxinstitute.org.au](http://www.saxinstitute.org.au)
(02) 9514 5950
directorate@saxinstitute.org.au
Association for the Blind of Western Australia Register of Blindness and Vision Impairment

Description of data source
A collection of demographic and diagnostic information relating to vision loss in adults who are referred to the Association for the Blind of Western Australia’s Low Vision Clinic.

Purpose(s)
To collect data on the incidence of blindness and vision impairment in WA.

Collection methodology
Referring specialists provide the diagnostic information. The Association supplements this with data from the initial assessment process.

Data content
Information is collected on basic demographic data, bilateral diagnosis, other medical conditions, visual acuity, visual field, pension eligibility and name of referring specialist.

Data content related to eye health
Bilateral diagnosis, visual acuity, visual fields, associated ocular and medical conditions.
Eye health-related data are medically verified.

Indigenous identification
No.

Socioeconomic identification
No.

Scope/target population
Adults referred to the Association’s Low Vision Clinic.

Geographical coverage
Western Australia.

Time period covered
Data have been collected since 1987 (ongoing collection).

Publications and other outputs
Yong VK, Morgan WH, Cooper RL, Shaw M, Bremner AP, Bulsara M & Yu D 2006. Trends in registered blindness and its causes over 19 years in Western Australia. Ophthalmic Epidemiology 13:35–42.

Access
For access to the data, contact the Association for the Blind of Western Australia Inc.
Data management agency
Association for the Blind of Western Australia Inc.
PO Box 101
Victoria Park WA 6979
<www.abwa.asn.au>
(08) 9311 8202
mailbox@abwa.asn.au
Australian Longitudinal Study of Ageing (SA)

Description of data source
The Australian Longitudinal Study of Ageing is a population-based bio-psycho-social and behavioural study of older adults. The longitudinal data collected in the course of the study comprise the most comprehensive longitudinal database yet assembled on ageing Australians.

Purpose(s)
The research has a number of specific objectives. These include:

• determination of levels of health and functional status of an older population and to track the changes in these characteristics over time
• identification of factors which promote and maintain health and wellbeing in an ageing population
• identification of risk factors for major morbid conditions and social, behavioural and other problems among an ageing population
• assessment of the effects of disease processes on functional status and the demand for health-care services and both informal and formal long-term care.

Collection methodology
Data from older people residing in SA in the community and residential care are collected using a mixture of personal and telephone interviews. Waves 1, 3, 6, 7 and 9 used comprehensive home interviews, functional assessments and optional mail-back questionnaires. Waves 2, 4, 5 and 8 used short telephone interviews of about 20 minutes duration examining major life events, accommodation changes, health and daily living activities. Data were also gathered during Wave 1 from medical practitioners and service providers (for example, domiciliary care, Meals on Wheels).

Data content
Demographic characteristics, health, formal and informal support systems, chronic conditions, mental health, housing, living arrangements, social activities, lifestyles, labour force participation and economic circumstances, nutrition, psychological wellbeing.
Physical tests measured include anthropometric parameters, blood pressure, visual and auditory acuity, tremor sway, grip strength, respiratory function and dental health.
Neuropsychological tests assessed memory, language and speed of information processing.
Laboratory investigations cover basic haematology, 20 channel biochemical analysis, lipid profiles and hormone analysis. Urine samples were tested for calcium, phosphate, sodium and creatinine.

Data content related to eye health
Data are collected about a participant’s chronic conditions; a measurement of visual acuity is also taken (both using the Snellen and LogMar charts).
Eye health-related data are medically verified and self-reported.
Indigenous identification
No.

Socioeconomic identification
Information is collected concerning housing, labour force participation, highest level of education achieved and economic circumstances.

Scope/target population
SA residents aged 70 years and over at Wave 1 of the study.

Geographical coverage
South Australia.

Time period covered
1992 (Wave 1) to 2007 (Wave 9); further waves of the study are dependent on future funding.

Publications and other outputs
• A comprehensive overview of the survey can be found in The Australian Longitudinal Study of Ageing: 15 years of ageing in South Australia: <flinders.edu.au/socsci/fcas/alsa/alsa_home.cfm> (viewed 24 August 2009).
• A full list of publications can be found on the Flinders Centre for Ageing Studies website: <flinders.edu.au/socsci/fcas/alsa/bibliography.cfm> (viewed 24 August 2009).

Access
Access to the Australian Longitudinal Study of Ageing can be obtained through the Flinders Centre for Ageing Studies.

Data management agency
Flinders Centre for Ageing Studies
School of Psychology
GPO Box 2100
Adelaide SA 5001
<flinders.edu.au/socsci/fcas/fcas_home.cfm>
(08) 8201 2041
fcas@flinders.edu.au
Blue Mountains Eye Study (NSW)

Description of data source
The Blue Mountains Eye Study was a large population-based assessment of visual impairment and common eye diseases of a representative older Australian community sample.

Purpose(s)
To determine the prevalence of eye disease in older Australians.

Collection methodology
In late 1992 the target population was identified by a door-to-door census of all dwellings in the two postcode areas. A total of 3,654 residents, aged 49–97, were examined during the period 1992–1994 (an overall response of 82.4% of non-institutionalised residents).
During 1997–1999, all surviving participants were invited to attend a 5-year follow up examination, for which 2,334 persons returned (75% of survivors).
A second census of the same postcode areas was conducted in 1999, identifying a further 1,510 residents now eligible to participate of which 1,206 were examined as part of an extension study during 1999–2000.
In 2002, participants were invited to return for 10-year follow-up exams, and during 2002–2004 1,952 original participants were re-examined. All participants were requested to partake in a 15-year follow-up in 2007.

Data content
Data collected included medical history (self-reported), socio-economic status, food frequency questionnaire, quality of life, eye examination, hearing examination, smell test (for Wave 3 only).

Data content related to eye health
The prevalence of eye disease in older Australians.
Eye health-related data are medically verified.

Indigenous identification
Information not provided.

Socioeconomic identification
Data was collected on marital status, income source and housing.

Scope/target population
All residents aged 49 years or older at baseline.

Geographical coverage
The project was conducted in an area that included Katoomba, Leura, Medlow Bath (postcode 2780) and Wentworth Falls (postcode 2782), NSW.
Time period covered

The first wave of the study, with the initial recruitment of participants, was conducted over 1992–1994. Wave 2 was conducted over 1997–1999, Wave 3 over 2002–2004 and Wave 4 over 2007–2008.

Publications and other outputs

- In excess of 300 peer-reviewed papers have been published to date, and abstracts regularly presented at conferences.
- Data from the Melbourne Visual Impairment Project and the Blue Mountains Eye Study have been combined and extrapolated to the entire Australian population. Results of this study were presented in the paper, Taylor HR, Keefe JE, Vu HTV, Wang JJ, Rochtchina E, Pezzullo ML & Mitchell P 2005. Vision loss in Australia. Medical Journal of Australia 182: 565–8.

Access

Access is granted with permission of the data management agency.

Data management agency

Centre for Vision Research
Department of Ophthalmology
Westmead Hospital, Hawkesbury Road
Westmead NSW 2145
<www.cvr.org.au>
(02) 9845 9077
paul_mitchell@wmi.usyd.edu.au
Cataract Surgery and Age-related Macular Degeneration Study (NSW)

Description of data source
The Cataract Surgery and Age-related Macular Degeneration Study is a longitudinal follow-up study.

Purpose(s)
The purpose of the study is to determine whether the risk of age-related macular degeneration increases following cataract surgery.

Collection methodology
Patients are examined pre-operatively and then re-examined at specified intervals post-operatively to determine whether the risk of age-related macular degeneration increases following cataract surgery.

Data content
Information is collected on medical history and quality of life data. An eye examination is conducted, including retinal photographs to document the state of the macula.

Data content related to eye health
The risk of age-related macular degeneration following cataract surgery.
Eye health-related data are medically verified.

Indigenous identification
Yes, but only a very small proportion are Indigenous participants.

Socioeconomic identification
No.

Scope/target population
Patients aged 65 and over undergoing cataract surgery at a large public hospital (Westmead Hospital) and in private ophthalmologists rooms.

Geographical coverage
Western Sydney suburban area.

Time period covered
Recruitment for the study was initiated in 2004; the study is currently ongoing.

Publications and other outputs
Access

Access to the data is not yet available as data collection is ongoing.

Data management agency

Centre for Vision Research
Department of Ophthalmology
Westmead Hospital, Hawkesbury Road
Westmead NSW 2145
<www.cvr.org.au>
(02) 9845 9077
paul_mitchell@wmi.usyd.edu.au
Central Australia Ocular Health Study data / hospital records / Regional Eye Health Coordinator records (NT)

Description of data source
Data sheets and clinical records summaries.

Purpose(s)
Data is used for coordination purposes and reporting.

Collection methodology
Completed by optometrists and ophthalmologists on bush visits, hospital clinics and town Aboriginal Medical Service clinics.

Data content
Data collected include demographic, diagnosis, referral and follow-up information.

Data content related to eye health
Summary screen of eye assessment parameters.
Eye health-related data are medically verified.

Indigenous identification
Yes.

Socioeconomic identification
No.

Scope/target population
Any patient seen by the Regional Eye Health Team or Visiting Optometrist.
More than 90% of remote community patients are Indigenous, while 50% of hospital clinic patients are Indigenous.

Geographical coverage
All of Central Australia and the Barkly region.

Time period covered
Since 2001 to present; summary data from 2001 and detailed data sheets from 2004 (ongoing collection).

Publications and other outputs
Some publications have followed specific research projects on central corneal thickness and optic disc appearances. However, there is extreme sensitivity about publishing any data without fully informed consent or ethical committee approval.

Access
Access to the data collected is restricted to clerical and administrative staff only.
Data management agency
NT Department of Health and Community Service/Regional Eye Program
Alice Springs Hospital
(08) 8951 7777
Dubbo Study

Description of data source

The Dubbo Study is a longitudinal community study of people born before 1930 living in the town of Dubbo, NSW.

Purpose(s)

The study has classic epidemiological goals, namely to identify patterns and predictors of mortality, hospitalisation and need for residential care. It is a biomedical and social science investigation of healthy ageing, service use, delay of disability and diseases like cardiovascular diseases and dementia.

Collection methodology

A group of 2,805 residents (1,233 men and 1,572 women) were first interviewed in 1988 and have been followed to the present with continuous ‘cold pursuit’ of death, hospitalisation and residential care data. There have been one-off collections of community services and other data as part of separately funded projects. From 1987, eligible subjects were identified through a reconciliation of GP and electoral records. They attended a centrally located office for medical and psychosocial assessment. The examinations extended from late August 1988 over a period of 13 months.

Survivors were re-interviewed in 2001 as part of the Asset and Health Dynamics phase of the study. At the end of 2000, there were 1,767 survivors of whom 1,537 remained in Dubbo and 230 lived outside Dubbo. Both groups were re-interviewed and there were 989 full interviews and 314 short interviews completed. Those missed in the follow-up interviews were generally unable to participate in interviews because of dementia and serious health limitations.

Data content

Briefly, the baseline examinations comprised demographic, psychosocial and standard cardiovascular risk assessments.

The medical examination included anthropometry; blood pressure (mean of two readings, phase V diastolic, after 10 minutes’ seated rest); resting electrocardiography; peak expiratory flow rate (best of two attempts, measured by a Wright peak flow meter); and 12-hour fasting levels of plasma lipids, lipoproteins and glucose.

A questionnaire explored measures of social support, depression status, education level, cognitive function (Short Portable Mental Status Questionnaire), alcohol and tobacco use, medications, medical history, myocardial infarction and chest pain, physical activity (walking, gardening and sporting activity, coded as daily, weekly or less often), self-rated health, and physical disability.

Data content related to eye health

Participants answered the following questions:

• Do you wear glasses?
• Do you have good sight?
• Do you have poor sight?
Eye health-related data are self-reported.

**Indigenous identification**
No.

**Socioeconomic identification**
Participants provided information on education and economic status.

**Scope/target population**
The eligible study population comprised all non-institutionalised residents of the Dubbo local government area who were born before 1 January 1930 (i.e. those 60+ years) and who attended for baseline assessment.

The study population was broadly representative of the Australian population born before 1930 by gender, age, employment, socioeconomic status, housing tenure, tobacco usage, mean blood pressure and other variables. However, 90% of the study population were Australian-born compared to 73% in NSW.

**Geographical coverage**
Dubbo, NSW.

**Time period covered**

**Publications and other outputs**

**Access**
Access to the Dubbo Study can be requested through the study directors.

**Data management agency**
Associate Professor Leon Simons
(02) 8382 2301
l.simons@notes.med.unsw.edu.au

Professor John McCallum
(03) 9199 4255
john.mccallum@vu.edu.au
Florey Adelaide Male Ageing Study

Description of data source

The Florey Adelaide Male Ageing Study (FAMAS) is a major study into the health and ageing of Australian men. It is believed to be the first of its kind in Australia to attempt to identify the wide range of factors that contribute to Australian men’s health (reproductive, physical and emotional) and their relationship with the ageing process. It is a longitudinal study involving biomedical assessments, self-reported measures and data linkage with established national databases.

Purpose(s)

FAMAS aims to identify the causes of men’s poorer physical and mental health in an effort to develop and implement measures to promote a healthy and active lifestyle, prevent diseases and guide the development of appropriate health services and policy.

Collection methodology

Random sampling using the Electronic White Pages (telephone directory) was used to recruit 1,200 men aged 35–80 from Adelaide’s north-west suburbs.

Biomedical clinic investigations occur began in 2002. Follow-up questionnaires are completed annually. Participants are invited on occasion to participate in substudies with selected collaborators.

Data content

Data collected includes utilisation of health services, including preventive and early intervention services; endocrinology of ageing, especially male sex hormones and the insulin growth factor axis; and prostate health and function in relation to the hormonal changes of ageing.

Chronic medical and psychological conditions such as obesity, cancer, diabetes, cardiovascular disease and mental health are also investigated.

Data content related to eye health

Information collected includes the following:

- vision-based quality-of-life (self-report)
- ophthalmic-based history (includes previous ocular trauma or surgery, topical medication use and ocular symptoms)
- ophthalmic examination (approx. 45 mins) involving
  - visual acuity testing (using a Snellen chart, photographs of eyes and adnexae)
  - slit-lamp examination of the anterior segment and lens, and tonometry
  - fundus photographs (pupils are dilated with tropicamide 1%)
  - automated refraction
  - A-scan ocular ultrasound
  - keratometry
  - pachymetry
  - optical coherence tomography.
Eye health-related data are medically verified and self-reported.

**Indigenous identification**
As per ABS Census 2001.

**Socioeconomic identification**
As per ABS Census 2001 and other statutory sources.

**Scope/target population**
Males aged 35–80 years, residing in the northern and western regions of Adelaide.
Participants’ data were compared with the 2001 Australian Census in order to assess, where possible, the representativeness of the cohort with the local and national populations. Overall, FAMAS participants were comparable to men in the same age group from the northwest Adelaide and Australian regions. The men in our cohort were found to be under-represented in the younger age group and over-represented in the older bracket, a common finding in studies of this type. In addition, there was an under-representation of men who had never married, reflecting previous findings that men without live-in support systems tend not to participate in health studies. The FAMAS cohort displayed both a higher proportion of men with some form of post-school qualification (specifically, trade and tertiary qualifications) as well as men from the lowest income bracket, both previously demonstrated as predictors of study involvement.

**Geographical coverage**
The ABS Northern and Western Statistical Divisions of Adelaide, SA.

**Time period covered**
Initial investigations began in 2002. Funding dependent, the study will continue for 15–20 years.

**Publications and other outputs**
The Ophthalmic Division of the Florey Adelaide Male Ageing Study is the largest substudy of FAMAS. Publication of results is pending.

**Access**
Submissions for collaboration are reviewed at FAMAS Investigator meetings as appropriate. A final decision on collaboration is made by the FAMAS Executive Committee.

**Data management agency**
Data Management and Analysis Centre
A joint data management centre of the University of Adelaide’s departments of Public Health and General Practice
<www.dmac.adelaide.edu.au>
(08) 8303 3564
International Centre for Eyecare Education / Aboriginal Health and Medical Research Council / Visioncare NSW Aboriginal project datasheets

Description of data source
International Centre for Eyecare Education (ICEE) / Aboriginal Health and Medical Research Council (AHMRC) / Visioncare NSW Aboriginal project data sheets comprise the data collected from the project to deliver culturally appropriate eye-care services to Aboriginal people through Aboriginal controlled communities.

Purpose(s)
For Aboriginal Medical Services internal utilisation and ICEE/AHMRC/Aboriginal Medical Services planning activities.

Collection methodology
Eye health coordinators complete standard forms.

Data content
Demographical data, eye conditions and spectacle wear history.

Data content related to eye health
Records incidence of diabetic retinopathy, glaucoma, cataract, trachoma, spectacles (whether had them before/Visioncare NSW spectacles), referrals to GP, optometrist or ophthalmologist. Self-reported diabetes.
Eye health-related data are medically verified.

Indigenous identification
Clients of the service are Aboriginal only.

Socioeconomic identification
No.

Scope/target population
Aboriginal people presenting at an ICEE/AHMRC eyecare clinic, approximately 75 locations in NSW.

Geographical coverage
New South Wales, where there are ICEE/AHMRC eyecare clinics.

Time period covered
Since 2000 to present (ongoing collection).

Publications and other outputs
Data are used for internal purposes only (e.g. planning). Data may be used on occasion in media reports (e.g. numbers seen).
Access

Access to the data is restricted.

Data management agency

International Centre for Eyecare Education
Level 4, North Wing
Rupert Myers Building
Gate 14 Barker Street
University of New South Wales NSW 2033

<www.icee.org>
(02) 9385 7435
icee@icee.org
Katherine Region Diabetic Retinopathy Study

Description of data source
The Katherine Region Diabetic Retinopathy Study was carried out in the lower top end of the NT between 1993 and 1996 as part of the Northern Territory Eye Health Program. It investigated diabetic eye conditions and its determinants in the Aboriginal Australian population of the region.

Purpose(s)
The study’s primary objectives were to:

• describe diabetic eye disorders in Aboriginal people living in the Katherine region by documenting the prevalence and incidence of diabetic eye disease in the known diabetic population of the region and monitoring the progression of diabetic retinopathy
• describe the visual acuity in the known diabetic population of the Katherine region and describe the effect of various ocular pathologies (both diabetic and non-diabetic eye disorders) on vision
• address the issue of service delivery in the Katherine region of the NT.

Collection methodology
The study population consisted of all known Aboriginal diabetic patients who were residents at the time of community visits in Katherine region. An initial study was performed by the Mobile Eye Unit in 1993 and then followed up in 1996.

Data content

• Personal section: data on the personal section were of primary importance in identifying the Aboriginal patients. This could only be done by filling in of all the information requested in the personal section. Name, date of birth, sex, community seen in, community patient lived in, date of examination, Medicare number and ethnicity were recorded.

• Medical history section: the medical records of the patients and all relevant past eye history and past medical history were recorded and reviewed. The diabetic medications actually taken by the patient, not necessarily those prescribed, were recorded. The presence of hypertension was also recorded.

• Ophthalmological examination:
  - visual acuity
  - intraocular pressure
  - anterior segment and posterior segment examinations.

• Collection of biochemical data.

Data content related to eye health
Diabetic eye conditions and its determinants.
Eye health-related data are medically verified.
Indigenous identification
The study population was limited to Aboriginal people. Aboriginal identification was accepted on the basis of the patient viewing him/herself as such.

Socioeconomic identification
No.

Scope/target population
The study population consisted of all known Aboriginal diabetic patients who were residents at the time of community visits in Katherine region.

Geographical coverage
Katherine region, NT.

Time period covered
Strictly, the data collection was covered between 1993 and 1996. For cultural reasons, the denominator data could not be compiled until 2001.

Publications and other outputs

Access
Not applicable.

Data management agency
Not applicable.
Melbourne Longitudinal Studies on Healthy Ageing program

Description of data source
The Melbourne Longitudinal Studies in Healthy Ageing (MELSHA) program was conducted initially by La Trobe University in partnership with the University of Sydney. Monash University now houses the study in collaboration with the University of Sydney. It was a series of three linked studies that formed a large longitudinal study on healthy ageing.

Purpose(s)
The aim of this baseline study (begun in 1994) was to investigate the health, health behaviours and service use of a representative group of older people living in Melbourne. The aim of the project is to identify avenues for changing individual actions and social environments that influence the health and wellbeing of older people.

Collection methodology
Data collected for Study 1 form the baseline for the MELSHA program. Data collection involved a face-to-face interview (with a proxy if necessary) to gather data on health, health behaviours and service use; a self-completed questionnaire which covered attitudes and life histories; a clinical examination which measured variables such as weight, height, eyesight and hearing; a brief mail-out questionnaire (with a phone call or visit, where necessary) to collect change and limited outcome variables and CATI to collect follow-up data on core variables.

Participants have been followed up every two years since 1994.

Data content
Medical conditions including continence, musculoskeletal and cardiovascular disease, falls and injuries, confusion, pain, medication use, depression and mental health, health behaviours (nutrition, exercise, social activity), wellbeing and service use (health services, community services, and residential care), functional ageing, health and service outcomes.

Data content related to eye health
The program collected the following information: cataracts, glaucoma, self-reported eyesight, eyesight check by optometrist, as well as visual acuity as measured by a Snellen chart.

Eye health-related data are medically verified and self-reported.

Indigenous identification
There were no Indigenous participants in the program.

Socioeconomic identification
Yes.

Scope/target population
Participants aged 65 and over living in Melbourne.

Geographical coverage
Melbourne, Victoria.
Time period covered

Publications and other outputs
- Journal articles, including:
- Books, book chapters and research reports, including:

Access
Access is currently restricted to investigators only. Data will be archived for restricted access in the future.

Data management agency
Prof. Colette Browning
Health Ageing Research Unit
Monash University
<www.mihsr.monash.org/centres/healthage.html>
(03) 9594 7560
collette.browning@med.monash.edu.au
Melbourne Visual Impairment Project

Description of data source
The Melbourne Visual Impairment Project was a population-based study of urban, rural and institutionalised residents of Victoria aged 40 years or older.

Purpose(s)
To determine the prevalence study of eye disease.

Collection methodology
Participants completed a standardised questionnaire covering sociodemographic characteristics, symptoms of eye disease, medical history and medication use. They were also invited to a local examination centre for a standard ophthalmic examination. Home visits were conducted for participants unable to attend the local examination centre.

Data content
The examination included presenting visual acuity (which is measured with participant’s spectacles, if usually worn for distance viewing), visual field, intraocular pressure, slit-lamp examination, dilated ophthalmoscopy and photography of the lens and fundus. Best-corrected visual acuity was measured for those with vision less than 6/6, using a standardised refraction.

Data content related to eye health
The prevalence of eye disease.
Eye health-related data are medically verified.

Indigenous identification
No.

Socioeconomic identification
No.

Scope/target population
Participants (4,744) were recruited from randomly selected pairs of adjacent Census collector districts (nine in urban Melbourne and four in rural communities in Victoria), had lived in their homes for at least 6 months and were aged 40 years and older. A further 403 participants were recruited from 12 nursing homes and 2 hostels randomly selected from 104 nursing homes and hostels located within a 5 km radius of the nine selected urban pairs of census districts.

Geographical coverage
Urban and rural Victoria.

Time period covered
The urban study was carried out from 1992 to 1994, the nursing home and hostel study in 1995 and the rural study in 1996.
Publications and other outputs

- Data collected from the Melbourne Visual Impairment Project have been published in many papers.
- Data from the Melbourne Visual Impairment Project and the Blue Mountains Eye Study have been combined and extrapolated to the entire Australian population. Results of this study were presented in the paper: Taylor HR, Keeffe JE, Vu HTV, Wang JJ, Rochtchina E, Pezzullo ML & Mitchell P 2005. Vision loss is Australia. Medical Journal of Australia 182:565–8.

Access

Access to the Melbourne Visual Impairment Project can be obtained through the Centre for Eye Research Australia.

Data management agency

Centre for Eye Research Australia
Department of Ophthalmology
University of Melbourne
Locked Bag 8
East Melbourne VIC 8002
<www.cera.org.au>
(03) 9929 8360
cera-info@unimelb.edu.au
New South Wales Population Health Survey

Description of data source

The NSW Population Health Survey is an ongoing telephone survey of state residents that is one of the main mechanisms through which NSW Health monitors population health and reports on performance indicators.

Purpose(s)

The NSW Population Health Survey objectives are to:

• monitor changes over time in self-reported health behaviours, health status, health service use, satisfaction with health services and other factors that influence health
• support the planning, implementation and evaluation of health services
• collect health information that is not available from other sources
• respond quickly to emerging needs for health information
• promote research.

Collection methodology

The NSW Population Health Survey is conducted continuously between February and December each year and covers the whole state population from birth upwards.

Households are contacted using list-assisted random digit dialling. One person from the household is randomly selected for inclusion in the survey. Carers or parents of children aged 0–15 years are interviewed on their behalf.

Trained telephone interviewers carry out the interviews. Most interviews are conducted in English but the survey is also conducted in five other languages: Arabic, Chinese, Greek, Italian and Vietnamese.

The sample is weighted to adjust for differences in the probabilities of selection among subjects, and for differences between the age and sex structure of the sample and ABS mid-year population estimates for NSW. This enables calculation of prevalence estimates for the state population rather than for the respondents selected.

Data content

The survey has collected information in modules in the areas of alcohol, asthma, cancer screening (breast and cervical), cancer screening (colorectal), community health centres, demographics, diabetes, difficulties getting health care, emergency departments, hearing, hospitals, immunisation, injury (youth violence), injury prevention, mental health, nutrition, oral health, overweight and obesity, physical activity, public dental services, self-rated health, sexual health, smoking, sun protection and vision.

Data content related to eye health

The vision module asked the respondent the following questions:

• When did you last have your eyesight checked? (Less than one year ago/1 year ago to less than 2 years ago/2 years ago to less than 5 years ago/5 or more years ago/Never/Don’t know/Refused)
• As far as you know, do you have normal vision in both eyes? (Yes/No/Don’t know/Refused)
• Do you currently wear glasses or contact lenses? (Yes/No/Don’t know/Refused)
• Are you wearing glasses for reading or close work, distance or both? (Reading/Distance vision/Both/Don’t know/Refused)
• Even when wearing glasses or contact lenses, do you have any difficulty reading or doing close work? (Yes/No/Don’t know/Refused)

Eye health-related data are self-reported.

**Indigenous identification**

Indigenous status is collected according to the ABS standard.

**Socioeconomic identification**

Socioeconomic details are collected, including the highest achieved education level, employment, household income and postcode.

**Scope/target population**

The target population is all NSW residents living in households with private telephones. The target sample is approximately 1,500 people in each area health service (a total sample of 12,000).

**Geographical coverage**

New South Wales.

**Time period covered**

The NSW Population Health Survey is a continuous survey which commenced in January 2002 (ongoing collection).

**Publications and other outputs**

The survey reporting plan includes:
• an annual report on adult health for the whole state
• a monthly report on adult health for the whole state
• an annual report on adult health for each health area
• a biennial report on child health for the whole state
• a triennial report on adult health for each division of general practice
• occasional reports on the health of older people, Indigenous people, people from culturally and linguistically diverse backgrounds and other specific populations.

Survey reports, which contain an analysis of the collected data, can be downloaded from the NSW Health website: <www.health.nsw.gov.au>.

**Access**

Survey data sets are available for further analyses. The NSW Department of Health encourages as many people as possible to access these data sets through the Health Outcomes Information Statistical Toolkit (HOIST). For users who do not have access to HOIST, data sets are available through specific requests to the Chief Health Officer.
publication that reports survey data must be approved by the Chief Health Officer before publication.

**Data management agency**
New South Wales Health Survey Program
(02) 9424 5759
catiadmin@doh.health.nsw.gov.au
Northern Territory Emergency Response Child Health Check Initiative data collection

Description of data source
The database contains information collected at each child’s Northern Territory Emergency Response (NTER) Child Health Check (CHC). Doctors and other health professionals perform voluntary health checks for eligible children, including a comprehensive medical check and referrals to any follow-up services required. This information is recorded and transmitted to the AIHW for inclusion in the collection.

Purpose(s)
• to manage and analyse information recorded at CHCs funded through the NTER
• to help assess the extent and type of follow-up care provided to children who had a CHC, by acting as a central collection whose content can be compared against the other three Child Health Check Initiative (CHCI) follow-up collections.

Collection methodology
The collection is a by-product of a clinical process whereby NTER health teams provide checks to children whose families volunteered for CHCs. Information collected at these health checks is recorded on a standard form and sent to the AIHW as a paper-based document. Information is entered into the CHC database manually by AIHW staff.

Data content
Topics include current health status, medical history, family history, living conditions and CHC referrals.

Data content related to eye health
All children are asked about any concerns about their vision. For children aged 6–15 years, data is collected concerning visual impairment and trachoma.

Eye health-related data are based on an assessment by a health professional. However, those collecting the data may not have had specific eye health training, and therefore data relating to trachoma should be treated with caution.

Indigenous identification
Yes.

Socioeconomic identification
No.

Scope/target population
Aboriginal and Torres Strait Islander children aged 15 years or less who live in the prescribed areas of the NT, who have not had a previous CHC or an MBS item 708 (Aboriginal and Torres Strait Islander child health check) in the last 9 months.
Geographical coverage
Prescribed areas of the NTER, including the OATSIH Planning Regions of Alice Springs Urban, Barkly, Borroloola, Central Australia, Darwin Rural, Darwin Urban, East Arnhem, Katherine East, Katherine Urban, Katherine West, Maningrida, Tiwi, Top End West and West Arnhem.

Time period covered
From July 2007 (ongoing collection).

Publications and other outputs
• Monthly summary and monitoring reports provided for DoHA (not for public release).

Access
Access to the data source is restricted.

Data management agency
Australian Institute of Health and Welfare
GPO Box 570
Canberra ACT 2601
<www.aihw.gov.au>
(02) 6244 1000
info@aihw.gov.au
North West Adelaide Health Study

Description of data source

The North West Adelaide Health Study is a representative biomedical population cohort study of approximately 4,000 adults aged 18 years and over in the northern and western regions of Adelaide. It is an ongoing epidemiological research collaboration between the South Australian Department of Health, the University of Adelaide, the University of South Australia, The Queen Elizabeth Hospital and Health Service, the Lyell McEwin Health Service and the Institute of Medical and Veterinary Science.

Purpose(s)

Its primary aim is to improve the regional conditions that help people maintain and promote health. It achieves these goals through the mobilisation of existing and new resources, the formation of new health partnerships and applied research that monitors and identifies community health activity to improve health outcomes for specific groups.

Collection methodology

Stage 1 of the study established baseline self-reported and biomedically measured information on those who were at risk of these conditions, previously undiagnosed with these conditions and diagnosed with these conditions.

The longitudinal nature of the study (Stage 2 onwards) allows examination of: incidence of these conditions; changes in patterns of severity, management costs and utilisation of resources; the impact of existing and future guidelines; and the potential for other health researchers to access a database for value-added chronic disease or related research.

Data content

Issues covered in the recruitment questionnaire included various health conditions, mental health, smoking status, reason for not wanting to participate in the study (if applicable) and demographics. The clinic tests conducted included blood pressure measurement, height and weight measurement, waist and hip measurement, blood test (fasting plasma glucose, lipids, glycated haemoglobin), allergy skin test and spirometry breathing test.

Data content related to eye health

Participants are asked: How many times in the last 12 months have you used the health services of an eye specialist/ophthalmologist in SA?

Indigenous identification

Yes.

Socioeconomic identification

Education, household income, occupation, pension/benefit, work status.

Scope/target population

Residents, aged 18 years and over, of the northern and western regions of Adelaide.
Geographical coverage
North-west suburbs of Adelaide, SA.

Time period covered
Since 1999 to present (ongoing collection).

Publications and other outputs
Reports available online: <www.nwadelaidelhealhstudy.org/results.asp> (viewed 24 August 2009).

Access
Access to the North West Adelaide Health Study can be obtained through the Population Research and Outcome Studies Unit.

Data management agency
Population Research and Outcome Studies Unit
Department of Health, Government of South Australia
PO Box 287
11 Hindmarsh Square
Adelaide SA 5000
(08) 8226 7042
pros@health.sa.gov.au
Queensland Injury Surveillance Unit database

Description of data source
The Queensland Injury Surveillance Unit (QISU) currently collects data from participating hospitals in Queensland which comprise three sample regions: metropolitan (Brisbane), regional (Mackay and Moranbah Health Districts) and remote (Mt Isa). These data characterise and measure the frequency of injuries, provide demographic information and detail activity, location and main product or factor involved.

Purpose(s)
QISU’s core business is injury data collection, data analysis and data dissemination. QISU supports injury prevention through the provision of injury surveillance data collected from metropolitan, regional and remote areas of Queensland. QISU’s data provide a unique database of intentional and unintentional injury presentations. The QISU database provides information to develop and evaluate injury prevention programs at a state level.

Collection methodology
Participating emergency departments provide data either electronically or on standardised forms which are then coded in accordance with the National Data Standard for Injury Surveillance (NDS-IS) and stored on the QISU database.

Data content
Data items include:
- age, sex, postcode
- country of birth, language
- time and date of injury event
- injury text description
- cause of injury (e.g. fall, poisoning)
- intent of incident (unintentional, assault etc.)
- place of injury (e.g. bedroom in boarding house)
- activity (e.g. playing cricket)
- nature of injury and body location or ICD-10 code
- mechanism and major injury factor (e.g. grinder)
- triage category (indication of severity)
- admission status.

Data content related to eye health
Injuries occurring to the eye, coded according to the NDS-IS. Eye health-related data are medically verified.

Indigenous identification
Indigenous status is collected. However identification of these cases is reliant upon participating hospitals.
Socioeconomic identification
Postcode of residence.

Scope/target population
Patients presenting to participating emergency departments in Queensland with an injury.

Geographical coverage
Areas of Queensland serviced by the participating hospitals.

Time period covered
Since 1998 to present (ongoing collection).

Publications and other outputs

Access
Access to the QISU database can be obtained through QISU.

Data management agency
Queensland Injury Surveillance Unit
Mater Health Services
Level 1 Whitty Building
Raymond Terrance
South Brisbane QLD 4101
<www.qisu.org.au>
(07) 3163 8569
mail@qisu.org.au
Royal Society for the Blind client database (SA)

Description of data source
The Royal Society for the Blind (RSB) client database is a client collection which holds information about the client and services provided to them.

Purpose(s)
The purpose of the Royal Society for the Blind client database is to collect data on clients of the Royal Society for the Blind. In addition, to monitor staff productivity and service delivery.

Collection methodology
Information is captured and input directly into the database.

Data content
Demographic details, eye impairments, additional health information, carer details, referrals and services delivered.

Data content related to eye health
Eye impairments, including primary and secondary eye conditions, degree of vision impairment and visual acuity.
Eye health-related data are medically verified.

Indigenous identification
The standard ABS classification for Indigenous identification is used.

Socioeconomic identification
Postcode and pension details are collected.

Scope/target population
SA residents with vision impairment. In 2007, the RSB estimated that the total vision-impaired population of SA to be 0.75% of the total SA population, equating to 11,660 people. The RSB active client base at the time was approximately 10,480, equating to 90% of the total estimated.

Geographical coverage
South Australia.

Time period covered
Since 1993–94 to present (ongoing collection).

Publications and other outputs
• Royal Society for the Blind strategic planning reports.
• Royal Society for the Blind annual reports.
• Reports to management and board, on topics including key performance indicators, service delivery and staff productivity.
• Benchmarking projects.
• Internal and external research projects.
• Government and Commonwealth State/Territory Disability Agreement National Minimum Data Set.

Access
Permission to access the Royal Society for the Blind client database must be made through the Royal Society for the Blind management and board.

Data management agency
The Royal Society for the Blind
Knapman House
230 Pirie Street
Adelaide SA 5000
<www.rsb.org.au>
(08) 8232 4777 or (08) 8232 2444
South Australian Health Monitor survey

Description of data source

Health Monitor is a telephone survey of the SA population (although other areas may also be surveyed). It is conducted three times a year and surveys approximately 2,000 households. The Health Monitor supplements the Health Omnibus Survey, which is conducted annually (see next entry). Health Monitor can be used by health professionals and policy makers for planning and development of strategies.

Purpose(s)

The goal of the Health Monitor survey is to collect, analyse and interpret data, which can be used to plan, implement and monitor health programs and other initiatives. Health Monitor is a ‘user-pays’ service. Each organisation pays only for survey questions that have direct relevance to their information requirements. Alternatively, surveys that focus on a specific topic may be commissioned by a sole organisation.

Collection methodology

Health Monitor uses a computer-aided telephone interviewing (CATI) system. Households to be included in the survey are randomly selected from the Electronic White Pages (telephone directory). Within households, the person who was last to have a birthday (aged 18 years or over) is usually selected to participate in the survey. Selected persons are non-replaceable, hence, if the selected person is not available, interviews are not conducted with alternative household members. At least six call-backs are made to each household before the selected individual is classified as a non-contact.

Data content

The Population Research and Outcome Studies Unit of the Department of Health assists clients to formulate questions and coordinates the preparation of the survey questionnaire. Each questionnaire includes approximately 60 questions. Ten background demographic questions are included free of charge to users.

Data content related to eye health

The following questions were included in the March 2000, July 2000 and March 2002 Health Monitor surveys:

- When did you last see an optometrist (who does eye testing for glasses)?
- Who referred you to the optometrist?
- When did you last see an ophthalmologist (an eye specialist)?
- Who referred you to the ophthalmologist?

Indigenous identification

Yes.

Socioeconomic identification

Employment status, pension received, education, household income.
Scope/target population
SA residents aged 18 years and over living in a household listed in the Electronic White Pages. Health Monitor can also survey populations outside of SA if requested to do so. The survey represents the population of SA aged 18 years and over.

Geographical coverage
South Australia.

Time period covered
Since 1999 to present (ongoing collection).

Publications and other outputs
See the Population Research and Outcome Studies Unit website for full details of publications and reports.

Access
Access to the Health Monitor survey can be obtained through the Population Research and Outcome Studies Unit.

Data management agency
Population Research and Outcome Studies Unit
Department of Health, Government of South Australia
PO Box 287
11 Hindmarsh Square
Adelaide SA 5000
(08) 8226 7042
pros@health.sa.gov.au
South Australian Health Omnibus Survey

Description of data source
The Health Omnibus Survey is a face-to-face survey conducted annually since 1991 for government and non-government organisations responsible for servicing the health needs of the SA community.

Purpose(s)
The goal of the Health Omnibus is to collect, analyse and interpret data that can be used to plan, implement and monitor health programs and other initiatives. The idea of an omnibus survey is that several organisations share the cost of conducting a survey, with each organisation paying only for those questions that are of direct relevance to their information requirements.
The Health Omnibus provides organisations in the health sector with an opportunity to generate health statistics at an affordable cost.

Collection methodology
A sample of 4,400 households is selected. One adult (non-replaceable) is interviewed in each household, on a face-to-face basis. The Health Omnibus aims to achieve a response rate of 70% or higher.

Data content
Approximately 10 background demographic questions (for example, age, gender, household income, marital status, education, country of birth) are included without charge and all clients’ data can be cross-tabulated against these.

Data content related to eye health
The following questions were included in the 2004 Spring Survey:
• Have you ever been told by a doctor that your vision has been affected because of your diabetes?
• Have you ever had laser therapy on your eyes because of your diabetes?
• Have you ever had cataract surgery?
Eye health-related data are self-reported.

Indigenous identification
Yes.

Socioeconomic identification
Household income, education, occupation.

Scope/target population
SA residents, aged 15 years and over.

Geographical coverage
South Australia.
Time period covered
Since 1991 to present (ongoing collection).

Publications and other outputs
See the Population Research and Outcome Studies Unit website for full details of publications and reports.

Access
Access to the Health Omnibus Survey can be obtained through the Population Research and Outcome Studies Unit.

Data management agency
Population Research and Outcome Studies Unit
Department of Health, Government of South Australia
PO Box 287
11 Hindmarsh Square
Adelaide SA 5000
(08) 8226 7042
pros@health.sa.gov.au
South Australian Monitoring and Surveillance System

Description of data source

The South Australian Monitoring and Surveillance System (SAMSS) was established in 2002 by the Population Research and Outcome Studies Unit of the SA Department of Health. SAMSS monitors population trends in state and national risk factors and chronic diseases so that the Department of Health has appropriate, timely and valid population health information to monitor health status, respond to population changes and support planning, implementation and evaluation of health services and programs.

Purpose(s)

The objectives of SAMSS are to:

• provide representative data
• characterise the problem or topic by time
• detect epidemics or changes in the topic occurrence
• identify high-risk groups or risk factors associated with the problem or topic and to suggest hypotheses for further investigation
• estimate the burden of the problem or topic
• highlight gaps in information and services that affect South Australians’ general health and wellbeing
• disseminate findings to professionals and administrators within the Department of Health, and other human services professionals or organisations in SA and Australia
• project future health-care needs
• set priorities for allocation of resources
• strengthen the network for surveillance and monitoring of issues relevant to the Department of Health to improve information gathering and exchange.

Collection methodology

SAMSS is a relatively simple, efficient and timely system—that is, data collection is quick without compromising the quality of the information collection. The survey is conducted by telephone utilising the CATI system.

SAMSS collects data each month. Approximately 600 interviews are conducted with people of all ages (surrogates are used for those under 16 years of age).

Data content

Data collected include demographics, health conditions, physiological risk factors, behaviours, health service utilisation, medication usage, social capital, quality of life and socioeconomic indicators.

Data content related to eye health

Participants respond to the following question: in the last four weeks, how many times [have you/has child’s name] used the health services of an optometrist in SA?
Indigenous identification
Yes.

Socioeconomic identification
Yes.

Scope/target population
SA residents of all ages.

Geographical coverage
South Australia.

Time period covered
Data collection started in June 2002 and as of March 2009, over 47,000 interviews had been conducted in SA for all ages.

Publications and other outputs
The following SAMSS reports have been published by the South Australian Department of Health, Population Research and Outcome Studies Unit:

- Population Research and Outcome Studies Unit 2006. Prevalence of self reported arthritis in the SA health regions.

Access
Access to SAMSS can be obtained through the Population Research and Outcome Studies Unit.

Data management agency
Population Research and Outcome Studies Unit
Department of Health, Government of South Australia
PO Box 287
11 Hindmarsh Square
Adelaide SA 5000
(08) 8226 7042
pros@health.sa.gov.au
**Spectacle assistance schemes**

**Description of data source**

Spectacle assistance schemes exist in each Australian state and territory to provide assistance with the purchase of spectacles to residents who satisfy particular eligibility requirements. Availability of data from these schemes is not standard across states and territories.

**Purpose(s)**

The schemes provide low cost spectacles (or other vision correction) to low income and disadvantaged residents. Each state/territory has different eligibility requirements and provides a different level of benefit.

**Collection methodology**

Collection of information differs between states and territories. Generally, information is collected at the dispensing point and then forwarded to be processed centrally in each jurisdiction.

**Data content**

While each state and territory differs, generally the following are collected: name, date of birth, Medicare number, pension details, prescription provider details, details of prescription.

**Data content related to eye health**

The number of people receiving assistance to purchase spectacles and the cost of providing this assistance.

**Indigenous identification**

Some schemes collect information concerning Indigenous status.

**Socioeconomic identification**

Not strictly, however benefits of the scheme are restricted to eligible residents according to their low income status.

**Scope/target population**

Generally residents of the applicable state and a recipient of the Commonwealth pension.

**Geographical coverage**

States and territories, Australia.

**Time period covered**

Varies.

**Publications and other outputs**

Reports are generally produced for internal purposes.
Access

Access availability differs from state to state.

Data management agency

Arrangements vary between states and territories. Contact the relevant state/territory health department.
Sydney Myopia Study

Description of data source

Funded by the National Health and Medical Research Council, the Sydney Myopia Study looks at factors contributing to the development of myopia.

Purpose(s)

The Sydney Myopia Study aims to document the prevalence of myopia in a large, representative sample of Sydney school children aged 6 and 12 (school years 1 and 7).

Collection methodology

The first phase of the Sydney Myopia Study was completed during the period June 2003 to September 2004. A comprehensive eye examination was conducted of 1,740 6-year-old children attending Year 1 from 34 schools, with an overall response of 78.9%. The second phase exams of 12-year-old children attending Year 7 was completed in December 2005, and saw 2,353 children from 21 schools, an overall response of 75.3%.

Data content

A complete eye test was conducted on each student. A questionnaire was used to collect information on students’ demographics, self-reported vision and hearing problems, utilisation of eye health services, use of glasses, general medical details, medications, birth history, home environment, activities and family history.

Data content related to eye health

Prevalence of myopia in school-aged children.
Eye health-related data are medically verified.

Indigenous identification

The ethnic origins of the student’s biological parents are recorded, with the ability to indicate as Indigenous Australian.

Socioeconomic identification

Yes.

Scope/target population

All year 1 (6-year-old) and year 7 (12-year-old) children from a random selection of primary and high schools in Sydney, balanced by socioeconomic status.

Geographical coverage

Sydney metropolitan area.

Time period covered

Baseline data was collected over 2002–2004. Follow up studies will continue in the future, pending further funding.
Publications and other outputs

Over 35 peer-reviewed papers have been published to date.

Access

Access is granted with permission of the data management agency.

Data management agency

Centre for Vision Research
Department of Ophthalmology,
Westmead Hospital, Hawkesbury Road
Westmead NSW 2145
<www.cvr.org.au>
(02) 9845 9077
paul_mitchell@wmi.usyd.edu.au
Sydney Older Persons Study

Description of data source
This study was conceived in 1988 as the Sydney Older Veterans Health and Services Project to investigate the health of community-dwelling veterans of World War II. The study was funded and commenced in 1991. At this stage, non-veteran members of the community were included as well.

Purpose(s)
The purpose of the Sydney Older Persons Study (SOPS) was to investigate the relationships between neurodegenerative ageing in people aged 75 and over and a range of other health factors including systemic diseases, disability, social supports and self-rated health and coping.

Collection methodology
Stage 1 of the study (which ran from 1991–1993) involved visiting 647 people in their homes and conducting medical and neuropsychological assessments. Participants were also asked questions concerning their health and lifestyle, and often a spouse, relative or friend was interviewed to provide further information.

The second wave of the study (SOPS Stage 2) involved interviews in the homes of the original study participants.

SOPS Stage 3 did not have the broad aims of the first two stages. Rather, it was based on the SOPS population and involved taking a blood sample and completing a medication history.

SOPS Stage 4 was a further follow-up of the original SOPS population using the same methodology and instruments used in Stages 1 and 2. This allowed a broader time frame over which to study the correlates of successful and unsuccessful ageing and the possible risk factors and protective factors for these. By this wave, 462 people remained in the sample. Of the people who participated in SOPS Stage 4, 128 took part in a further phase that consisted of a magnetic resonance imaging (MRI) scan of the brain and more detailed neuropsychological assessments.

In 2001, SOPS Stage 5 commenced and data collection was completed in 2003. This study focused on the subgroup of subjects who had a MRI scan at SOPS 4. These people were invited to participate in a further MRI scan, detailed neuropsychological assessment and medical review. In addition, we re-interviewed the informants (such as a family member or a close friend) to measure the presence of health changes and their impact over time. It is anticipated that these data will provide invaluable information about the correlates and predictors of cognitive decline, with the potential to assist in the diagnosis of dementia and its subtypes. Data analysis is in progress.

Data content
Over 1,000 variables including information collected on demographics, level of difficulty associated with activities of daily living, medical conditions and disabilities.

Data content related to eye health
The following information was collected:
• self-reported incidence of cataracts, glaucoma, macular degeneration

128
• need for glasses
• inability to read an eye chart
• clinical qualitative assessment of the degree of visual impairment (none, mild, moderate, severe)
• assessment of the degree to which visual impairment was contributing to functional impairment (none, mild, moderate, severe).

Eye health-related data are medically verified and self-reported.

**Indigenous identification**

No.

**Socioeconomic identification**

Income, education, occupation and retirement details.

**Scope/target population**

Community-dwelling people aged 75 years and over in the inner-west of metropolitan Sydney, NSW.

**Geographical coverage**

The inner-western municipalities of metropolitan Sydney including Concord, Drummoyne, Strathfield, Burwood, Ashfield, Leichhardt, Marrickville and South Sydney.

**Time period covered**


**Publications and other outputs**

A number of papers have been published, most recently:


**Access**

Access to SOPS data can be gained directly through the Prince of Wales Medical Research Institute/Ageing Research Centre or through the Australian Social Science Data Archive at the Australian National University.
Data management agency
The Prince of Wales Medical Research Institute
Cnr of Barker Street and Easy Street
Randwick NSW 2031
<www.powmri.edu.au>
(02) 9399 1000
Victorian Emergency Minimum Dataset injury surveillance collection

Description of data source

The Victorian Injury Surveillance Unit centralises and manages data from 38 Victorian public hospital emergency departments as part of the Victorian Emergency Minimum Dataset injury surveillance collection of emergency department presentations in Victoria.

Purpose(s)

Deliver and maintain a Victorian injury surveillance program for injuries to all persons in Victoria with the following specific objectives to:

- promptly identify the emergence of new hazards, or the existence of previously unrecognised hazards
- give special focus to injuries and hazards aligned to state and nationally identified priority areas for action
- monitor secular trends in indicator injury rates
- provide support to primary care partnerships and local government supporting injury prevention activities
- assist in the evaluation of injury prevention interventions.

Collection methodology

Each hospital collects the required information for each presentation of an emergency department patient through a software package. If the presenting diagnosis is an injury, further related information is required. This information may be completed by the treating doctor, the triage nurse or clerical staff. Emergency department data are extracted monthly from the individual hospitals to Victoria’s Department of Human Services. Quarterly, the Department of Human Services extracts a subset of the Victorian Emergency Minimum Dataset, only presenting injury cases, and passes this to the Victorian Injury Surveillance Unit.

Data content

Variables collected include: age, sex, injury cause, location, activity, nature of main injury, body region, human intent, birthplace, preferred language, address postcode, departure status, referral on departure and a 250-character text description of the injury event.

Data content related to eye health

Injuries occurring to the eye, coded according to the National Data Standard for Injury Surveillance (NDS-IS).

Eye health-related data are medically verified.

Indigenous identification

Indigenous status is collected, but not utilised.

Socioeconomic identification

Postcode of usual residence.
Scope/target population
Patients presenting to monitored emergency departments with an injury.

Geographical coverage
Rural, remote and metropolitan Victoria, with 13 of the 38 hospitals in rural/remote Victoria.

Time period covered
Since October 1995 to present (ongoing collection).

Publications and other outputs
Short reports are available on the Victorian Injury Surveillance Unit (VISU) website.

Access
The VISU data and information request service is open to government and non-government organisations, the higher education and schools sector, industry and business and community members. The service is not available to primary and secondary school students.

Data management agency
Victorian Injury Surveillance Unit
Monash University Accident Research Centre
Building 70
Monash University VIC 3800
<www.monash.edu.au/muarc/VISU>
(03) 9905 1805
visu.enquire@general.monash.edu.au
**Victorian Population Health Survey**

**Description of data source**

The Victorian Population Health Survey is based on a core set of question modules that inform decisions about public health priorities. It measures the relevance and responsiveness of public health programs to current and emerging health issues.

**Purpose(s)**

The aim of the Victorian Population Health Survey is to provide quality, timely data to inform indicators of population health that are intended to have direct application to evidence-based policy development and strategic planning across the department and the wider community.

**Collection methodology**

The survey is conducted via CATI. The content of the survey was determined after reviewing the determinants of chronic disease states that are most likely to have an impact on Victorians. Priority has been given to areas in which a public health response is likely to be effective in improving health and reducing inequalities in health.

Interviews are conducted in the major non-English languages in Victoria to ensure people of culturally and linguistically diverse backgrounds are represented.

**Data content**

The survey collects data on areas including demographics, screening, health-care use, self-reported height and weight, alcohol, smoking, asthma, blood pressure, diabetes, social capital measures, physical activity, self-reported health status, Kessler 10 measure of psychological distress, health conditions and eye care.

**Data content related to eye health**

The following information is collected:

- change in vision over the last 12 months
- the use of eye protection
- consultation with an eye-care specialist or attendance at an eye clinic
- type of eye problem including cataract, glaucoma, diabetic eye disease or diabetic retinopathy and macular degeneration.

Eye health-related data are self-reported.

**Indigenous identification**

Yes.

**Socioeconomic identification**

Information is provided on a participant’s highest level of education, household income and area of residence.
Scope/target population
A representative state-wide sample of adults aged 18 years or over is randomly selected from households in each of the eight departmental health regions.

Geographical coverage
Victoria.

Time period covered
Since 1998 to present (ongoing collection).

Publications and other outputs
Annual report:

Access
Access can be granted to the Victorian Population Health Survey data via licensed deed with the Department of Human Services: contact Loretta Vaughan, Manager Victorian Population Health Surveys, (03) 9096 5286.

Data management agency
Health Surveillance and Evaluation Section
Rural and Regional Health and Aged Care Services Division
Department of Human Services
50 Lonsdale Street
Melbourne VIC 3000
(03) 9096 0344
public.health@dhs.vic.gov.au
Western Australian Aboriginal Child Health Survey

Description of data source

The Western Australian Aboriginal Child Health Survey (WAACHS), a large-scale epidemiological survey of the health and wellbeing of 5,289 WA Aboriginal and Torres Strait Islander children, was undertaken in 2001 and 2002 by the Telethon Institute for Child Health Research, under the direction of the project’s Aboriginal Steering Committee. The survey was designed to build an epidemiological knowledge base from which preventive strategies can be developed to promote and maintain healthy development and the social, emotional, academic and vocational wellbeing of young people.

Purpose(s)

The specific aims of the WAACHS were to:

• describe and define the health and wellbeing of WA Aboriginal and Torres Strait Islander children and young people aged 0–17 years
• estimate the prevalence and distribution of commonly occurring chronic medical conditions and disabilities (for example, asthma, cerebral palsy, visual and hearing impairments, intellectual disability) and describe how they may affect a child’s wellbeing and functioning
• estimate the prevalence, distribution and functional impact of common health and mental health problems in Aboriginal and Torres Strait Islander children and young people aged 0–17 years and their families
• estimate the prevalence and distribution of adverse health behaviours (for example, smoking, alcohol, drug and volatile substance misuse)
• estimate the prevalence and distribution of other psychosocial problems such as early school leaving, conduct problems and juvenile offending
• describe Aboriginal and Torres Strait Islander children, adolescents and their families’ access to, effective use of and satisfaction with health care, education, juvenile justice, housing and social services
• identify markers resulting in protection from, and amelioration of, poor health and wellbeing (mental health), adverse health behaviour(s) and other psychosocial problems
• develop estimates of risk and markers identifying Aboriginal and Torres Strait Islander children and young people at increased risk for various health, educational and vocational outcomes.

Collection methodology

Survey methodology and instrumentation were developed in consultation with Aboriginal leaders, key Aboriginal bodies (the Aboriginal and Torres Strait Islander Commission regional council, the Aboriginal Council of Elders, the Aboriginal Justice Council and the Western Australian Aboriginal Community Controlled Health Organisation) and through extensive community consultations throughout the state.

Dwellings were selected for screening using an area-based clustered multi-stage sample design. From 166,290 dwellings in 761 census collection districts, 139,000 dwellings were approached to determine if residents were eligible to participate in the survey. Using this method, a random sample of 2,386 families with 6,209 eligible children was identified throughout metropolitan, rural and remote regions of WA. A total of 1,999 of these families
(84%) with 5,513 eligible children consented to participate in the survey. Interviewers gathered usable data on 5,289 (96%) of these participating children.

Data were collected from interviews with the carers in the household who were the most knowledgeable about the survey children. In addition to the information collected on children, separate interviews were undertaken with up to two carers per child to gather information about the demographic and social characteristics of the household and family and to ask questions about the dwelling, neighbourhood and community.

In addition to the data gathered on children, data were also gathered on families.

**Data content**

Questionnaire content covered child and youth development; health and wellbeing; functional impairment and disability; use and access to health, education and social services; and a selected number of questions about diet; demographic and social characteristics of the household and family; information about the dwelling, neighbourhood and community.

The survey was subsequently linked to records of hospital admissions, diagnoses, procedures and transfers through the WA Record Linkage System.

**Data content related to eye health**

The following questions were asked:

- Does the respondent have normal vision in both eyes?
- Does the respondent use prescribed glasses or contact lenses?
- What access does the respondent have to vision services?

Eye health-related data are self-reported.

**Indigenous identification**

Yes.

**Socioeconomic identification**

Employment, education, remoteness.

**Scope/target population**

The survey was based on an area sample of dwellings. Families in selected dwellings who reported that there were ‘Aboriginal or Torres Strait Islander children or teenagers living at this address who are aged between 0 and 18 years’ were eligible to be in the survey.

Children living within group homes, institutions and non-private dwellings were not in the scope of the survey. However, where a selected household had a child temporarily living away from home (for example, in a boarding school or hostel), these children were included in the scope of the survey.

**Geographical coverage**

Western Australia.

**Time period covered**

Commenced in May 2000 and was completed in June 2002.
Publications and other outputs


Access

General enquiries about the WAACHS and enquiries seeking statistical clarification of any of the topics covered should be directed to the Telethon Institute for Child Health Research at waachs@ichr.uwa.edu.au.

Data management agency

Telethon Institute for Child Health Research (under the direction of the project’s Aboriginal Steering Committee)
PO Box 855
West Perth WA 6872
<www.ichr.uwa.edu.au>
(08) 9489 7777
enquire@ichr.uwa.edu.au
2.3 Analysis

There are a large number of data sources which can be used to indicate the prevalence and outcomes of eye diseases and injuries and eye health-care utilisation in Australia. Some of these data sources have been developed for the specific purposes of collecting information on eye health, while others have been developed with a broader purpose. Many of these data sources are based on nationally agreed standards that enable information to be compared across different service-delivery settings and sectors.

This section first provides an overview of how the data informs the National eye health framework (the Framework). It then provides information regarding how new data developments are complementing programs enacted under the Framework.

National Eye Health Framework

There is no comprehensive information framework that can assess the utility of these data sources. It is possible, however, to apply the analysis of data sources in this document to the information needs of the Framework.

The Framework outlines five key areas for action in preventing avoidable blindness and vision loss. These are:

1. reducing the risk
2. increasing early detection
3. improving access to eye health-care services
4. improving the systems and quality of care
5. improving the evidence base.

Reducing the risk

The objective of the first key action area is to prevent eye disease and vision loss where possible through addressing known modifiable risk factors. The first key action area is well informed by a number of the eye health data sources identified in Section 2.2. The monitoring of this key action area may be informed by data sources that measure prevalence and incidence of eye diseases, injuries and risk factors. The identified data sources which inform this area include population-based surveys, clinical-based studies and national monitoring systems. Many of these data sources also include risk factor-related data, Indigenous status, comorbidity and socioeconomic status.

Increasing early detection

The objective of the second key action area is to detect treatable eye conditions early so that interventions can be applied to preserve vision and prevent any further vision loss. Key action area 2 is only moderately informed by the key Australian eye health data sources identified. Data sources that may inform this area are restricted in availability to those that collect information on eye examinations. However, this area has improved more recently with some of the new data collections included in this guide, such as the National Indigenous Eye Health Survey and the Northern Territory Emergency Response data.
Improving access to eye health-care services

The objective of the third key action area is to ensure that all Australians have equitable access to appropriate eye health care when required. This is an area well informed by a number of the eye health data sources identified. The monitoring of this area may be informed by data sources that measure encounters with the health-care sector or those that collect information about the eye health workforce. The identified data sources which inform this area include national administrative data collections, surveys and professional registration collections.

Improving the systems and quality of care

The objective of the fourth key action area is to ensure eye health care is safe, affordable, well-coordinated, consumer-focused and consistent with internationally recognised good practice. Key action area 4 is the least informed of the five areas with few collections contributing to this area’s information needs. A small number of data sources have limited information about referral pathways.

Improving the evidence base

The objective of the fifth key action area concerns eye health-care policy, planning and programs which are supported by high quality research and data collection systems. Virtually by definition, key action area 5 is successfully informed by the identified data sources. To varying degrees, each data source informs this area.

In summary, the information needs of the Framework are informed by a number of data sources, as identified in Section 2.2. Key action area 5 (Improving the evidence base) is the area most informed by the identified data sources. Key action areas 1 (Reducing the risk) and 3 (Improving access to eye health-care services) are both well informed by the identified data sources, while key action area 2 (Increasing early detection) is only moderately informed and key action area 4 (Improving the systems and quality of care) is the least informed of the five areas.

New data and the Framework

Data sources which have been included since the first edition of this guide was published have further helped to inform the Framework.

There have been a number of new sources of data which will help improve our understanding of Indigenous eye health. These include the Northern Territory Emergency Response Child Health Check Initiative data collection, the National Indigenous Eye Health Survey, the National Trachoma Surveillance and Reporting Unit data, OATSIH service delivery data and the Longitudinal Study of Indigenous Children. These will be important for assessing programs provided under key action area 1 (Reducing the risk), most notably the need to target, treat and control trachoma in Indigenous communities and, under key action area 2 (Increasing early detection), the early detection of eye disease and childhood vision screening for Indigenous persons.

This edition has also included more information about health workforce data. This is a priority area under key action area 3 (Improving access to eye health-care services). The National Registration and Accreditation Scheme (beginning in 2010) should provide more accurate data regarding ophthalmologists and optometrists.
Since the Framework was released in November 2005, a number of programs have been initiated by both the Australian Government and state and territory governments in order to prevent avoidable blindness and vision loss. These programs cover a wide range of areas within the eye health sector. As such, although some key areas and programs are well informed by existing data, others (especially key action area 4, *Improving the systems and quality of care*) remain less well-informed by data sources in this guide.

Through the monitoring of data that inform the key areas for action, achievements over time can be measured. Monitoring of the prevalence and outcomes of eye diseases and injuries and eye health-care utilisation is necessary to improve Australia’s ability to make decisions for cost-effective allocation of resources, to plan preventive and treatment services and to target priority population groups. Monitoring of these data will also allow the impact of risk factors, health promotion and prevention strategies to be measured.
3 Terminology in eye health

Statistics on the prevalence of vision loss in Australia are available from a number of sources. Confident comparisons of these statistics both nationally and internationally rely on uniformity in the terms and definitions used to describe vision loss. Examination of these sources indicates that the most common terms used to refer to vision loss in Australia are vision impairment, low vision, blindness and legal blindness. Full details of the variety of definitions currently used to describe vision loss in Australia are outlined in the following pages.

3.1 Visual impairment and low vision

Definitions of visual impairment and low vision are less established than the definition of blindness. The terms are used regularly, but are frequently used interchangeably. While the International Statistical Classification of Diseases and Related Health Problems, Tenth Revision, Australian Modification (ICD-10-AM) definition of low vision relates to reduced vision excluding blindness, Vision Australia defines visual impairment according to the same criteria. Other versions of both visual impairment and low vision use the same definition—a significant reduction in vision, including blindness.

The visual acuity upper limit which constitutes a significant reduction in vision is also a point of difference among definitions. Visual acuity limits vary between less than 6/12 and less than 6/18. Definitions of visual impairment and low vision in use in Australia are presented on page 142.

3.2 Blindness and legal blindness

The most commonly adopted definition of blindness in Australia is classified in ICD-10-AM. This definition is based on the World Health Organization’s (WHO) standard definition of blindness which has been internationally accepted and incorporated into the International Statistical Classification of Diseases and Related Health Problems, Tenth Revision (ICD-10). The ICD-10-AM and ICD-10 classification systems are widely accepted by many Australian studies and data collections, including the Australian Corneal Graft Registry, the Florey Adelaide Male Ageing Study, the National Hospital Morbidity Database and the National Mortality Database.

Some definitions of blindness employ variants of the ICD-10-AM definition, including those used by the Blue Mountains Eye Study, the AIHW, the Department of Veterans’ Affairs and the Australian National Diabetes Information Audit and Benchmarking. Full details of variations of the definition of blindness are presented on page 143.

Variations in definitions relate to the use of different levels of visual acuity loss. While the internationally aligned ICD-10-AM definition imposes a visual acuity of less than 3/60, a number of definitions of blindness maintain a visual acuity level of less than 6/60. Although a visual acuity of less than 3/60 aligns with internationally recognised definitions, it has been documented that the increased complexity of daily tasks requires a less severe level of visual acuity, that is, less than 6/60 (Dandona & Dandona 2006).
The concept of legal blindness arose with the introduction of social security systems. The term legal blindness is generally used by governments to determine the criteria by which a person is deemed eligible for government concessions, such as income assistance and disability support. In Australia the criteria for legal blindness is established under s. 95 of the Social Security Act 1991 (Commonwealth of Australia 2006). The definition of legal blindness is widely and consistently adopted by researchers and community organisations. Details of the definition for legal blindness and the organisations and studies which have adopted this definition are presented on page 144.

Definitions of visual impairment used in Australia

- The ICD-10-AM classification ‘visual impairment’ includes ‘blindness’ and ‘low vision’. Therefore a visual acuity with best possible correction of less than 6/18 and/or a corresponding visual field loss of less than 10 degrees around central fixation or no light perception (National Centre for Classification in Health 2006).
  
  Adopted by:
  - National Mortality Database (Dunn et al. 2007)
  - National Hospital Morbidity Database (AIHW 2006b).

- An individual has some degree of sight loss (Royal Blind Foundation Queensland 2006).

- Visual acuity <6/12 in both eyes, established by Eye Research Australia (Access Economics 2004).

- Visual acuity <6/18, established by the Katherine Region Diabetic Retinopathy Study.

- Best corrected visual acuity <6/18 and/or visual field constriction to within 20 degrees of fixation, established by the Melbourne Visual Impairment Project (Livingston et al. 1997).

- Best-corrected visual acuity of 6/12 or worse, established by Blue Mountains Eye Study (Wang et al. 2000).

- A person with visual acuity of less than 6/18 (0.3) but equal to or better than 6/60 (0.05) in the better eye with the best possible correction and/or a visual field of less than 20 degrees, referred to as ‘vision impaired’ (Vision Australia 2006c).

- People who are visually impaired include those who are blind, who have vision significantly less than normal (which is usually taken as acuity less than 6/18) but are not classified as blind (Fred Hollows Foundation 2008).

- ‘Visual impairment’ to be used when the condition of vision loss is characterised by a loss of visual functions (such as visual acuity, visual field etc.) at the organ level. Many of these functions can be measured quantitatively, established by International Council of Ophthalmology (ICO), as endorsed locally by the Royal Australian and New Zealand College of Ophthalmologist (ICO 2002).

Definitions of low vision used in Australia

- Visual acuity with best possible correction of less than 6/18, but equal to or greater than 3/60 (National Centre for Classification in Health 2006).

  Adopted by:
  - National Mortality Database (Dunn et al. 2006)
  - National Hospital Morbidity Database (AIHW 2006b).
• A person is said to have low vision when their eyesight is limited or impaired and cannot be corrected with conventional glasses or contact lenses (Vision Australia 2006b).

• Low vision is an impairment to vision that significantly interferes with the functioning of a person and cannot be adequately corrected with medical, surgical, therapy, conventional eyewear or contact lenses. It is often a loss of sharpness or acuity but may present as a loss of field of vision, light sensitivity, distorted vision or loss of contrast. Low vision may occur as a result of birth defects, injury or as a complication of disease (Macular Degeneration Foundation 2006).

• ‘Low vision’ is to be used for lesser degrees of vision loss, where individuals can be helped significantly by vision enhancement aids and devices, established by the International Council of Ophthalmology (ICO), as endorsed locally by the Royal Australian and New Zealand College of Ophthalmologist (ICO 2002).

Definitions of blindness used in Australia

• Visual acuity with best possible correction of less than 3/60, and/or a corresponding visual field loss of less than 10 degrees around central fixation or no light perception (National Centre for Classification in Health 2006).

  Adopted by:
  - Australian Corneal Graft Registry
  - Florey Adelaide Male Ageing Study
  - National Mortality Database (Dunn et al. 2006)
  - National Hospital Morbidity Database (AIHW 2006b).

• Presenting visual acuity less than 6/60 in the best eye, established by the Blue Mountains Eye Study and Melbourne Visual Impairment Project combined study (Taylor et al. 2005).

• Visual acuity of less than 3/60 or corresponding visual field loss in the better eye with best possible correction (Fred Hollows Foundation 2008).

• Whether the individual has become legally blind in either or both eyes. Blindness is less than 6/60 vision in the better eye with glasses. Vision 6/60 is the ability to see only at 6 metres what the normal eye can see at 60 metres. An indicator of the presence or development of a visual impairment or inability to see (AIHW 2005b).

• Visual acuity < 6/60 in both eyes established by the Australian National Diabetes Information Audit and Benchmarking (National Association of Diabetes Centres 2005).

• ‘Blindness’ to be used only for total vision loss (that is, no light perception) and for conditions where individuals have to rely predominantly on vision substitution skills, established by the International Council of Ophthalmology (ICO), as endorsed locally by the Royal Australian and New Zealand College of Ophthalmologist (ICO 2002).

• There is total loss of sight; or corrected visual acuity is less than or equal to 6/60 in both eyes; or where, in the written opinion of an ophthalmologist, the visual fields deficits; and/or combination of deficits results in a visual impairment which is the equivalent of a corrected visual acuity measure of less than or equal to 6/60 in both eyes established by the Australian Government for the purposes of determining permanent blindness pensions paid by the Department of Veterans’ Affairs (Australian Government, Department of Veterans Affairs 2006).
Definitions of legal blindness used in Australia

- Visual acuity on the Snellen scale after correction by suitable lenses must be less than 6/60 in both eyes; or constriction to within 10 degrees of fixation in the better eye irrespective of corrected visual acuity; or a combination of visual defects resulting in the same degree of visual impairment as that occurring in the above points, established by the Australian Government for the purposes of determining permanent blindness for Disability Support Pension or Age Pension—Blind under s. 95 of the Social Security Act 1991 (Australian Government 2006).

Adopted by:

- Blind Citizens Australia, used as determining criteria for full and junior membership (Blind Citizens Australia 2006)—referred to as ‘permanent blindness’
- Eye Research Australia (Access Economics 2004)
- Queensland Blind Association Inc., used as a basis for membership and purchase of white canes (Queensland Blind Association Inc. 2001)
- Royal Blind Foundation Queensland (2006)
- Retina Australia (2001)
- The Katherine Region Diabetic Retinopathy Study
- Vision Australia, as stated in the Constitution (Vision Australia 2006a).

3.3 Other definitional considerations

Presenting visual acuity versus corrected visual acuity

‘Corrected visual acuity’ is visual acuity measured according to the most appropriate refractive correction. ‘Presenting visual acuity’, however, is a measurement of an individual’s acuity with the refractive correction which is currently in use by the individual, for example, spectacles or contact lenses (Dandona & Dandona 2001).

An individual may qualify as ‘not blind’ because their vision improves with refractive correction. However, in their daily living, their vision may be poor enough, due to uncorrected or under-corrected refractive error, that they qualify as ‘blind’ when measured according to their presenting visual acuity.

Although easily managed, uncorrected refractive error remains a major cause of vision impairment (Vennewkirk et al. 2001). A study has determined that nearly 300,000 of the 480,000 Australians who are visually impaired are so because of under-corrected refractive error (Access Economics 2004). In 2006, WHO recognised uncorrected refractive error as an important cause of vision loss. Broadening the definition of visual impairment, to include uncorrected refractive errors, effectively doubled the estimated total number of visually impaired people worldwide (WHO 2007). Defining vision loss according to corrected visual acuity would therefore have a substantial statistical impact on measuring the burden of vision loss.
Visual field

Vision loss is generally measured according to visual acuity, visual field or both. Despite visual field loss being a major component of vision loss, second to visual acuity loss (ICO 2002), many of the definitions associated with vision loss do not utilise this measure. As visual field loss can exist independently of visual acuity loss (ICO 2002), prevalence estimates determined exclusively on visual acuity may underestimate the true prevalence rate of blindness.

Functional vision

One of the difficulties of limiting the measure (and hence definition) of vision-loss terms to ‘visual acuity’ and ‘visual field’ is that they are objective measurements that do not take into account an individual’s ability to use vision in activities of daily living.

The International Classification of Functioning, Disability and Health (ICF) provides a multi-perspective approach to the classification of functioning and disability as an interactive and evolutionary process (WHO 2001). Figure 3.1 illustrates the various components and interactions of the ICF. By qualifying vision loss according to the component of body functions and structures only, in terms of visual acuity and visual field, many other important components of measuring visual functionality are ignored.

In summary, there are a number of key definitional issues surrounding vision loss terminology. While the definition of legal blindness is well established, other terms associated with vision loss lack the same uniformity.
Glossary

ASCED  The Australian Standard Classification of Education (ASCED) was developed by the ABS for use in the collection, storage and dissemination of statistical and administrative data relating to educational activity undertaken in Australia. The classification system includes two component classifications: Level of Education and Field of Education.

CURF  Confidentialised unit record files are files of responses to ABS surveys that have had specific identifying information about persons and organisations confidentialised.

ICD-9-BPA  ICD-9-BPA is an adaptation of WHO's International Statistical Classification of Diseases, Ninth Revision by the British Paediatrics Association (the Royal College of Paediatrics and Child Health).


ICD-10  ICD-10 is a classification system in the WHO Family of International Classifications. It is the international standard diagnostic classification for all general epidemiological and many health management purposes. It is used to classify diseases and other health problems recorded on many types of health and vital records including death certificates and hospital records.

ICD-10-AM  ICD-10-AM is the Australian modification of the international standard ICD-10. It consists of a tabular list of diseases and accompanying index, a tabular list and index of procedures and Australian Coding Standards for the selection of codes.

The ICD-10-AM disease component is based on the WHO ICD-10. It uses an alphanumeric coding scheme for diseases. It is structured by body system and aetiology.

The ICD-10-AM procedures component is based on the MBS. The classification is structured by body system, site and procedure type. Procedures not currently listed in MBS have also been included (for example, allied health interventions, cosmetic surgery).

ICF  International Classification of Functioning, Disability and Health (ICF) is a classification system in the WHO Family of International Classifications. It describes how people live with their health condition. ICF is a classification of health and health-related domains that describe body functions and structures, activities and participation. The domains are classified from body, individual and societal perspectives. Since an individual's functioning and disability occurs in a context, ICF also includes a list of environmental factors.
<table>
<thead>
<tr>
<th>Acronym</th>
<th>Description</th>
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<tbody>
<tr>
<td>ICPC</td>
<td>Internal Classification of Primary Care (ICPC) has been incorporated into the WHO Family of International Classifications. It classifies patient data and clinical activity in the domains of general/family practice and primary care, taking into account the frequency distribution of problems seen in these domains. It allows classification of the patient’s reason for encounter (RFE), the problems/diagnosis managed, interventions and the ordering of these data in an episode of care structure.</td>
</tr>
<tr>
<td>K-10 measure</td>
<td>The Kessler measure is a 10-item self-report questionnaire intended to yield a global measure of ‘psychological distress’ based on questions about the level of anxiety and depressive symptoms in the most recent 4-week period.</td>
</tr>
<tr>
<td>MBS</td>
<td>Medicare Benefits Schedule (MBS) is a listing of the Medicare services subsidised by the Australian Government.</td>
</tr>
<tr>
<td>NDS-IS</td>
<td>National Data Standard for Injury Surveillance (NDS-IS) describes data items and classifications designed to support public health surveillance of injury.</td>
</tr>
<tr>
<td>PBS</td>
<td>Pharmaceutical Benefits Scheme (PBS) is listing of all the pharmaceuticals subsidised by the Australian Government.</td>
</tr>
<tr>
<td>Visual acuity</td>
<td>Measurement of the finest details that an eye can distinguish, often estimated using LogMar or Snellen charts.</td>
</tr>
<tr>
<td>Visual field</td>
<td>Measurement of the ability to detect objects to either side, or above or below, the direction of central fixation.</td>
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</tbody>
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References


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