The economic value of informal care in 2010

12 October 2010

Report by Access Economics Pty Limited for
Carers Australia
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Foreword

In 2005 Carers Australia commissioned Access Economics to undertake the first Australian study of the economic value of the informal care provided by unpaid family carers for people with disability, mental illness, chronic conditions, terminal illness and the frail aged. That report found that, five years ago, the annual ‘replacement value’ of informal care had passed the $30 billion mark.

Five years on, this report shows that – using the same method – the value of informal care has increased to exceed $40 billion per annum in 2010, 33% higher than in 2005.

The growth in the value of informal care derives largely from demographic ageing – which is increasing the number of Australians who require and receive care – together with growth in the replacement cost of care from $25 per hour on average in 2005 to $31 per hour today.

Even using the most conservative ‘opportunity cost’ method, the value of informal carer is around $6.5 billion per annum. This is about the same value as all high-end residential aged care paid for by the Australian Government (0.5% GDP).

Informal carers provide 1.32 billion hours of care each year, and represent a precious economic resource in an age of growing health and aged care workforce shortage.

The valuable contribution of Australia’s 2.9 million carers, which enables their loved ones to remain at home, comes at a health cost to carers themselves. The case study in this report illustrates the extent of the burden of caring-related conditions such as depression, stress-related illness, sleep deprivation and musculoskeletal problems.

Young carers also make enormous sacrifices, giving up what can be millions of dollars of lifetime earnings in order to provide the care that their parents need.

Yet informal family care remains one of the lowest subsidised forms of care, and family carers remain under-resourced in terms of education and information compared to paid care workers. Greater investment in training for informal carers could generate savings that are many times the value of the investment, as literature reviewed in this report shows.

Without some serious policy thinking, it will be challenging to address the emerging care shortfalls projected for the coming decade, as demand increasingly outstrips supply.

Carers Australia advocates strongly to enact the recommendations of this report to prioritise support for family carers that can prevent or delay institutionalisation – respite, income support, training and preventive health care.

I commend the report to you.

Tim Moore
President, Carers Australia
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<table>
<thead>
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<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ABS</td>
<td>Australian Bureau of Statistics</td>
</tr>
<tr>
<td>ADAT</td>
<td>Adult Disability Assessment Tool</td>
</tr>
<tr>
<td>AIHW</td>
<td>Australian Institute for Health and Welfare</td>
</tr>
<tr>
<td>CACP</td>
<td>Community Aged Care Packages (Federal program)</td>
</tr>
<tr>
<td>CDAP</td>
<td>Child Disability Assistance Payment</td>
</tr>
<tr>
<td>CPI</td>
<td>Consumer Price Index</td>
</tr>
<tr>
<td>CSTDA</td>
<td>Commonwealth State/Territory Disability Agreement</td>
</tr>
<tr>
<td>EACH</td>
<td>Extended Aged Care in the Home (Federal program)</td>
</tr>
<tr>
<td>EACH-D</td>
<td>Extended Aged Care in the Home Dementia (Federal program)</td>
</tr>
<tr>
<td>FaHCSIA</td>
<td>Family and Housing, Community Services and Indigenous Affairs (Australian Government Department of)</td>
</tr>
<tr>
<td>GDP</td>
<td>gross domestic product</td>
</tr>
<tr>
<td>GP</td>
<td>general practitioner</td>
</tr>
<tr>
<td>HACC</td>
<td>Home and Community Care (Federal program)</td>
</tr>
<tr>
<td>NATSEM</td>
<td>National Centre for Social and Economic Modelling, University of Canberra</td>
</tr>
<tr>
<td>RAC</td>
<td>residential aged care</td>
</tr>
<tr>
<td>REACH</td>
<td>Resources for Enhancing Alzheimer’s Caregiver Health</td>
</tr>
<tr>
<td>SDAC</td>
<td>Survey of Disability, Ageing and Carers (ABS)</td>
</tr>
<tr>
<td>UK</td>
<td>United Kingdom</td>
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</table>
Executive Summary

Carers provide informal care\(^1\) to people who need assistance due to disability, mental illness, chronic conditions, terminal illness or due to being frail aged. In this report, Access Economics examines the amount of informal care being provided in Australia today. Placing a dollar value on the work of informal carers is the first step in evaluating whether the current use of informal and formal care models is socially optimal, in terms of both efficiency and equity.

Section 1 identifies who are Australia’s carers, providing a profile of carers and the care needs of the people they care for.

In 2010, over 1 in 8 Australians (2.87 million people) are estimated to be providing informal care. This represents 13.1% of Australia’s population, a higher share than in 2003.

- **540,000 (1 in 40 Australians)** are ‘primary carers’, providing the majority of the recipient’s care.
- In absolute terms, there are around 310,000 more carers in 2010 compared to 2003, and 65,400 more primary carers. The number of carers, and primary carers, is increasing at some 2% per annum (around the same rate as population growth).

Informal carers together provide an estimated 1.32 billion hours of care in 2010.

- This is equivalent to each carer providing an average of 460 hours of care per year or 9 hours per week.
- However, care hours are in fact much more unevenly distributed, with primary carers providing 54% or 714 million hours annually, despite representing only 19% of all carers.

Section 2 looks at two measures of how this time spent by carers could be valued.

- In 2010, an estimated 129,900 carers will not be employed due to their caring responsibilities (1.1% of Australia’s workforce). The **opportunity cost** of time devoted to informal care, measured as reduction in paid employment due to caring, provides a ‘lower bound’ estimate of $6.5 billion (equivalent to 0.5% of GDP and 9.5% of the value of formal health care). This measure reveals the resources that are diverted each year from production in the formal economy to informal care.
  - Rates of employment and labour force participation among carers are substantially lower than the Australian average, even when standardised from differing age-gender profiles.

- The **replacement valuation** reveals the resources that would need to be diverted each year from the formal economy to replace the work done by informal carers, were their services no longer available. If all hours of informal care were replaced with services purchased from formal care providers and provided in the home, the **replacement value** would be $40.9 billion (equivalent to 3.2% of GDP and 60% of other formal health care).

Section 3 reviews, and quantifies where possible, other effects of informal care that should be included in a full account of the costs and benefits of alternative care models.

---

\(^1\) ‘Informal care’ refers to care provided by unpaid family carers.
The productivity losses associated with provision of informal care are borne mainly by the individuals themselves, whose wage income is reduced.

- However, the wider community is affected through reduced personal income tax collection and the payment of income support payments to carers.
- While the taxation and welfare payments are transfer payments not real economic costs, they have associated efficiency losses estimated to be $1.76 billion in 2010.

Also likely to be substantial, but not costed here, are the impacts of caring on the health and wellbeing of carers. Often the burden of pain and suffering associated with depression, musculoskeletal injuries and other problems dwarfs the financial costs.

- On a relative prevalence basis, the sleep impacts of caring alone may exceed $1 billion per annum, including a financial estimate of the reduced quality of life.
- One case study model presented in this report was of a hypothesised carer, ‘Karen’, with mild-moderate depression, stress-related angina (a cardiovascular condition), and chronic lower back pain. The total financial impact of all three conditions is over $33,000 in 2010. Adding in the cost of the loss of Karen’s healthy life (about 27% of a healthy life year is lost to the conditions), the total impact is nearly $79,000. Karen bears 56% of the financial costs (mainly lost income but also health costs over $1,000), while the Government bears 21%. Karen bears all the wellbeing loss.

Young carers, an often over-looked group, are at high risk of long term disadvantage as result of missed education opportunities and compromised high school performance. The ‘25-hour rule’ which limits recipients of Carer Payment to a maximum of 25 hours per week of study, work and travel, compounds the many difficulties faced by young carers in obtaining certain tertiary qualifications.

- A case study interview with Chantelle, 21, who has cared for her mother since she was 12, shows that by being unable to undertake a medical degree, for which she gained entry, due to the ‘25-hour rule’ Chantelle will earn less than 60% of what she could have as a doctor i.e. $3.1 million as a psychologist rather than $5.45 million over a lifetime, in real 2010 dollars. The difference is $2.35 million, comprising lost tax revenue of $0.5 million and a loss to Chantelle of $1.9 million.

Section 4 examines the level of public (Government) support for carers relative to other models of care.

Governments are increasingly recognising the contribution of family carers, and the costs they bear in providing informal care. The level of Government support, both through welfare support payments and service provision, has increased over the past few years.

- However, in contrast to the education and training provided to formal care workers, there is relatively little funded training for informal carers.
- A case study based on a randomised clinical trial of an informal carer training program in the UK showed that, if the same program was delivered in Australia, benefits would exceed costs by over $19,000 per annum per person – through reducing reliance on formal health sector and community services relative to a situation of no carer training. A similar randomised controlled trial should be conducted in Australia to confirm these modelled findings, potentially across a variety of care settings.
The economic value of informal care in 2010

The analysis conducted in this report based on average costs of care shows that combinations of informal care and community based formal care services provided together are generally lower cost than institutionalised care, using an opportunity cost valuation of carer’s time, excluding health costs to the carer and noting that the cost of residential care includes a housing component. For 2010 the estimated relativities per person on this basis are:

- Informal primary care with HACC services is lowest at $12,983; informal primary care with CACP costs around $23,425; EACH plus informal primary care is around $51,264; EACH-D plus informal primary care is around $53,831; RAC low care is around $44,319; while RAC high care is greatest at $69,178 per person in 2010.
- The relativities and public subsidy components are shown in Chart i.

![Chart i: Average cost of care models per recipient, Australia, 2010](chart)

Source: Access Economics calculations.

**Section 5 outlines the questions and challenges that face policy makers.** It describes how the analysis in this report can inform these questions, as well as further work that is needed.

The demand and supply of informal care will be influenced by many factors in the future, most notably:

- many chronic and disabling conditions are highly age-related suggesting that, with population ageing, more people may require care in the future (a greater demand for care); and
- there will be relatively fewer younger people, greater mobility and dispersion of families, increased female labour force participation, higher rates of relationship breakdown and single-person households, and potentially reduced propensity to care from Generation X and Y relative to previous generations, all of which may reduce the
desire or ability to provide informal care (a lower supply of informal care, or growth of supply lower than demand growth).

As Chart ii shows, the ‘carer ratio’, of primary carers to older people with a disability, is projected to fall from 60% now to under 40% by mid-century reflecting that demand for informal care will substantially outstrip supply.

Chart ii: Projected demand and supply of informal carers

The growing gap between demand and supply was not substantially affected by three scenario analyses.

- An overall decline in the propensity of people to care, represented by a 20% across-the-board decrease in carer rates, increased the base case deficit by 12.1% in 2050.
- A decline in the propensity of women to reduce paid employment in order to provide care, represented by a 20% decrease in the proportion of women reducing paid employment to care, increased the base case deficit by 5.5% in 2050.
- An increase in the availability of carers due to converging male and female life expectancy, represented by a 20% increase in carer rates in 65+ age group, reduced the base case deficit by 3.1% in 2050.

The future mix of care – between community and RAC, and between formal and informal services – is unclear – reflecting the changing pattern of disease (favouring RAC and formal care services), shifting social preferences (generally towards community care) and supply constraints (notably potentially fewer informal carers). The third InterGenerational Report
The economic value of informal care in 2010

(Treasury, 2010) concluded there will be a much larger relative increase in spending on RAC than on community care, with formal care expenditure increasing from 0.8% of GDP today to 1.8% of GDP in 2049-50.

Given that investments in informal carer support have been shown to be cost effective, there is a strong case to prioritise support for carers that can delay or prevent costly institutionalisation, including:

- respite care;
- income support;
- education and training; and
- preventive health care (e.g. lifting techniques, health checks and a personalised health record to help maintain carers’ health).

Such support for carers will help ensure and enhance future labour productivity across the formal and informal sectors of the economy, as well as meeting Australia’s obligations to optimise wellbeing for the frail aged those disadvantaged by disability, mental illness or chronic illness.

Access Economics
1 Who are Australia’s informal carers?

Carers are people who provide care to others in need of assistance or support. Informal carers provide this service free of charge and outside the boundaries of ‘formal’ government services provision. Typically, an informal carer is a family member or friend of the person receiving care. Much informal care is provided by people living in the same household as the recipient of the care. As such, many people receive informal care from more than one person. The person who provides the majority of informal care is known as the primary carer.

Although informal care can be defined to include parenting and other forms of unpaid child care, this report focuses solely on unpaid care provided to people with a disability, mental illness, chronic condition, terminal illness and the frail aged.

The most comprehensive profile of people receiving care and their carers is provided by the Survey of Disability, Ageing and Carers (SDAC). This national survey, conducted by the Australian Bureau of Statistics (ABS), covers people living in private dwellings in urban or rural areas. However, also within its scope are people living in non-private dwellings including care accommodation (nursing homes, hostels and other facilities). The most recent survey was conducted in 2009 but the results have not yet been published. As such this report relies on data from the 2003 survey, with those results presented first, followed by estimates for 2010.

1.1 Demographic profile of carers

There were just under 2.6 million Australians in the 2003 SDAC providing informal care to those who needed help due to disability or age (ABS, 2004b), equivalent to 12.9% of the Australian population at that time (ABS, 2004c). Chart 1.1 shows the distribution of these carers by age and gender. The majority of carers are female (54.1%), with most carers being middle-aged (35 to 54 years) although over three quarters are of working age (18 to 64 years). There were also an estimated 169,900 minors who were carers.

Primary carers provide the majority of informal assistance to the care recipient and as such are usually living in the same household (78%). One in five informal carers or 474,600 Australians were primary carers. The age-gender distribution of primary carers (Chart 1.2) is different from that of all carers (both primary and non-primary). Women represent a far greater proportion of primary carers, at around 71.3% of the total.
Chart 1.1: Number of informal carers by age and gender, Australia

Notes: Data for the year 2003.

Chart 1.2: Primary carers by age and gender, Australia

Notes: Data for the year 2003.
1.1.2 Propensity to give care by demographic group

Chart 1.3 shows the ‘rate’ of informal care provision for each age-gender group, defined as the proportion of that population sub-group who provide some form of informal care. This chart shows that the rate of informal care provision is not uniformly distributed among the population. Proportionally, older age groups are more likely to provide care than younger people. For men, the likelihood of providing care increases steadily as they age. Women also experience an increased likelihood of providing care up to the age of 65, when the propensity to care begins to fall.

This change in the relative propensity to care among older males and females may be due in part to lower male life expectancy. The earlier loss of male partners means that older females are more likely to be widowed while older males are more likely to be married. In older age groups care is predominately provided to a partner, which makes surviving older men more likely to be carers than older women. Another explanation might be that older men are healthier than older women, with the rates of severe or profound disability lower for men than they are for women over the age of 65 (ABS, 2008a). However, this may be an artefact of the difference in life expectancy.

Chart 1.3: Rate of informal care provision by age and gender, Australia

![Chart 1.3: Rate of informal care provision by age and gender, Australia](image)


1.1.3 Regional and income inequalities in informal care provision

Compared to the Australian population as a whole, carers are slightly more likely to live outside major cities. Around 35.9% of all carers live in regional or remote areas (Chart 1.4), compared to 34.3% of all people. This difference was similar for primary carers, with 36.1% living in regional and remote areas.
Carers typically live in households with lower than average income per person, being over-represented in the lower quintiles and under-represented in the higher quintiles. This income inequality is more pronounced for primary carers. Chart 1.5 presents data for people aged between 15 and 64 years of age only, suggesting the differential is not entirely due to the larger proportion of elderly people in the carer population, who tend to have lower levels of (pension rather than wage) income than the general population. Instead, the income inequality is likely to be related to the reduced levels of paid employment among carers, which appears to be caused in part by the demands of their caring role (Section 2.1).

Notes: People aged 15-64 years only. 
If people aged 65 and over are included in the analysis the income inequality among primary carers increases further still. Over 25% or one quarter of all primary carers reported to the SDAC have gross household income in the lowest quintile.

1.1.4 Informal carers in 2010

As the latest publicly accessible SDAC was undertaken in the second half of 2003 the number of carers was extrapolated to 2010 to calculate the economic value of informal care. This was done by applying the rates of informal care provision (Table 1.1 and Chart 1.3) to the current demographic structure of the population. The best estimates of Australia’s current population are the ABS Series B or mid-level population projections, based on 2006 census data (ABS, 2008b). This application results in an increase in the total number of carers, based on the movement of some people to older age cohorts (with increased rates of informal care provision). In Section 5 of this report other factors that may affect the supply and demand for informal carers over a longer time horizon are considered.

<table>
<thead>
<tr>
<th>Age</th>
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<th>All carers</th>
<th>Female</th>
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<tr>
<td></td>
<td>Primary</td>
<td>Non-primary</td>
<td>Primary</td>
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<tr>
<td>0-17&lt;sup&gt;(a)&lt;/sup&gt;</td>
<td>0.1</td>
<td>3.5</td>
<td>3.6</td>
<td>0.1</td>
</tr>
<tr>
<td>18-24</td>
<td>0.3</td>
<td>8.9</td>
<td>9.2</td>
<td>1.3</td>
</tr>
<tr>
<td>25-34</td>
<td>0.6</td>
<td>9.1</td>
<td>9.8</td>
<td>2.4</td>
</tr>
<tr>
<td>35-44</td>
<td>1.2</td>
<td>12.0</td>
<td>13.2</td>
<td>4.4</td>
</tr>
<tr>
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<td>2.4</td>
<td>13.9</td>
<td>16.3</td>
<td>6.1</td>
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<td>16.7</td>
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<td>3.5</td>
<td>17.8</td>
<td>21.3</td>
<td>5.6</td>
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<tr>
<td>75+</td>
<td>5.2</td>
<td>16.8</td>
<td>22.0</td>
<td>3.7</td>
</tr>
</tbody>
</table>

Notes: (a) All primary carers in the SDAC data for this age group are at least 15 years old.

Based on these rates, there are a total of 2.87 million informal carers in Australia in 2010, of whom approximately 540,000 are providing primary care (Chart 1.6 and Table 1.2). This represents 13.1% of Australia’s population, a higher share than in 2003. One in eight Australians is a carer, and one in 40 is a primary carer.

In absolute terms, there are around 310,000 more carers in 2010 compared to 2003, and 65,400 more primary carers. The number of carers, and primary carers, is increasing at some 2% per annum (around the rate of population growth).
Chart 1.6: Number of carers, by type of carer, Australia 2010

Table 1.2: Estimated informal carers by age, gender and type, Australia, 2010

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Primary Carers</th>
<th>Non-Primary Carers</th>
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<td></td>
<td>M</td>
<td>F</td>
<td>P</td>
</tr>
<tr>
<td>&lt; 18 (a)</td>
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<td>1.6</td>
<td>3.0</td>
</tr>
<tr>
<td>18-24</td>
<td>3.5</td>
<td>13.9</td>
<td>17.3</td>
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<td>25-34</td>
<td>9.6</td>
<td>36.6</td>
<td>46.2</td>
</tr>
<tr>
<td>35-44</td>
<td>18.5</td>
<td>68.6</td>
<td>87.1</td>
</tr>
<tr>
<td>45-54</td>
<td>35.7</td>
<td>93.2</td>
<td>128.9</td>
</tr>
<tr>
<td>55-64</td>
<td>30.4</td>
<td>94.1</td>
<td>124.5</td>
</tr>
<tr>
<td>65-74</td>
<td>27.2</td>
<td>45.1</td>
<td>72.3</td>
</tr>
<tr>
<td>75+</td>
<td>30.0</td>
<td>30.0</td>
<td>60.1</td>
</tr>
</tbody>
</table>

Total | 156.4| 383.1| 539.4|1,165.0|1,160.9|2,325.9|1,321.4|1,544.0|2,865.4

Notes: (a) Access Economics estimate of the number of primary carers less than 15 years of age.
Source: Access Economics calculations.

1.2 Care needs

Informal care is most often provided by close family members. In the SDAC, 91% of primary carers were providing assistance to either a child, parent or partner (Chart 1.7).
Table 1.3 shows the range of assistance received, with 85% of people with a disability receiving some informal care and 57% receiving some formal care. Needs were higher for people with profound or severe disability relative to those with mild or moderate disability. Just under 5% of people with disabilities stated that their care needs were not fully met.

The more severe a person’s disability, the greater the amount of time devoted to caring by their primary carer. Nearly two thirds (64%) of primary carers for people with a profound core activity limitation provide 40 hours or more care a week, compared to 30.7% for primary carers whose main recipient of care has a severe core activity limitation and 19.7% for those with a mild, moderate or non-core activity limitation (Chart 1.8).
Physical disabilities are the main reason (57%) co-resident primary carers are providing care. One in five care recipients has a psychological or intellectual disability and one in six care recipients has a sensory or speech disability as their main disability (Chart 1.9).

Chart 1.9: Co-resident primary carers, main disability of the main recipient of care

2 Valuing carers’ time

Informal care is provided free of charge, but is not free in an economic sense, as time spent caring is time that cannot be directed to other activities such as paid work or leisure. As such, informal care can be valued as the opportunity cost associated with the loss of economic resources (labour) and the loss in leisure time valued by the carer.

There are three potential methodologies which can be used to place a dollar value on informal care.

- The opportunity cost method measures the formal sector productivity losses associated with caring, as time devoted to caring responsibilities is time which cannot be spent in the paid workforce.
- The self-valuation method measures how much carers themselves feel they should be paid.
- The replacement cost method – measures the cost of ‘buying’ an equivalent amount of care from the formal sector if the informal care were not supplied.

The self-valuation method is not commonly used, and there are no reliable Australian studies of the amount Australian carers feel they should be compensated. Interestingly, a 2000 Irish study of dementia carers provided a very low figure, of between £2 to £4 per hour (O’Shea, 2000).

Estimates of the value of informal care are sensitive to the estimation methodology used. In this study, the opportunity cost method is used to estimate a lower bound of the value of informal care, and the replacement cost method for the upper bound estimate.

2.1 Opportunity cost

Opportunity cost measures the productivity losses associated with caring, as time devoted to caring responsibilities is time that cannot be spent in the paid workforce. It attempts to measure the amount of production carers would have contributed to the economy, but for their caring responsibilities.

2.1.1 Employment status of carers

The SDAC found that carers are more likely to be unemployed or not participating in the paid workforce than those who are not carers (Table 2.1). Of carers aged 15 and over who are looking for work, 5.7% cannot find it, compared to 5.3% of non-carers. Only 56.1% of carers participate in the workforce (i.e. they are employed on either a part or full time basis or are looking for work) compared to 67.9% of non-carers. Primary carers have particularly low rates of labour force participation, only 39.0% in 2003.
Table 2.1: Unemployment and participation rates by carer status, Australia 2003

<table>
<thead>
<tr>
<th>% civilian population aged 15+</th>
<th>Primary carer</th>
<th>Non-primary carer</th>
<th>All carers</th>
<th>Not a carer</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unemployment rate</td>
<td>3.1</td>
<td>6.2</td>
<td>5.7</td>
<td>5.3</td>
<td>5.3</td>
</tr>
<tr>
<td>Participation rate</td>
<td>39.0</td>
<td>60.2</td>
<td>56.1</td>
<td>67.9</td>
<td>66.1</td>
</tr>
</tbody>
</table>

Source: ABS special data request.

Not all carers who are unemployed or not in the labour force would work if they did not have care obligations. Of working age primary carers not employed, 60% were not working prior to commencing their role as a carer, and less than 22% reported leaving work as a direct result of being carers (ABS, 2005).

To accurately measure differences in employment and participation rates between carers and the general community, it is necessary to standardise for differences in the demographic composition of the two groups. Carers are more likely to be female and of an older age than the general population. On average, both women and older people are less likely to be employed than the general population. Age and gender standardised rates of employment for carers are shown in Chart 2.1. These are the employment rates that would be observed if the age and gender profile of Australian carers was identical to that of the general population.

Chart 2.1: Age and gender standardised employment rates by carer status, Australia

Chart 2.1 shows that, even when the demographic differences between the carer population and the overall Australian population have been accounted for, there remain differences in the level and type of employment. However, it should be noted that there are other factors that
might affect employment rates that have not been controlled for here, including education and individual heterogeneity, which could affect the results (Leigh, 2010).

All carers are less likely to be working full-time than the Australian average (42.0%). In particular, the rate of full-time employment among primary carers is just 19.2%, less than half that of the general population. The rate for non-primary carers is also lower than the average, at 36.4%.

Conversely, carers are more likely to be in part-time work. The age-standardised rate of part-time employment is 22.8% for primary carers and 21.1% for non-primary carers, compared to 17.2% for the general population.

The substantial difference in the employment rate for primary carers compared to the Australian average (Table 2.2) is due to the much lower labour force participation rate among primary carers. The age-standardised rate of labour force participation for primary carers is 45.2%, compared to 62.7% of the Australian population. Labour force participation includes the unemployed (those looking for work) as well as the employed.

Table 2.2: Age and gender standardised employment rates by carer status, Australia

<table>
<thead>
<tr>
<th>%</th>
<th>Primary Carers</th>
<th>Non-primary carers</th>
<th>All Carers</th>
<th>Australian Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standardised employment rate, carers</td>
<td>42.1</td>
<td>57.5</td>
<td>54.9</td>
<td>59.1</td>
</tr>
<tr>
<td>Difference from Australian Average</td>
<td>17.0</td>
<td>1.6</td>
<td>4.3</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Source: Access Economics calculations.

2.1.2 Opportunity cost valuation

The opportunity cost of informal care is measured as the income forgone by the carer spending time providing care rather than undertaking paid work. The cost is calculated assuming that, in the absence of their caring responsibilities, carers would be employed at the same rate as members of the general population of the same age and gender. Hence the difference between the standardised employment rates for carers and those of the general population (Table 2.2) is a measure of the rate of forgone employment due to informal care. It is important to note that this method will not include the opportunity cost of forgone leisure time, as leisure (like informal care) is a non-market good (i.e. it is not formally traded or paid for in a market).

Primary carers have much lower rates of employment than the national average, even when demographic differences have been accounted for (42.1% compared to 59.1% — see Table 2.2 above), suggesting that 17.0%² of primary carers are not in paid employment due to their caring role. In 2010, this is equivalent to 91,900 primary carers.

A much smaller percentage of non-primary carers are not employed due to their caring responsibilities. Of non-primary carers, 1.6% or around 37,200 people are not in paid employment due to their caring role in 2010. Combined with the 91,700 primary carers, this gives a total of **129,900 carers who are not in the paid workforce this year**, who would have

² This is higher than the proportion of primary carers who expressed a desire to return to work (14.1%).
been employed if not for their caring responsibilities. This is equivalent to 1.1% of Australia’s current workforce (ABS, 2010a).

If these people were employed and received, on average, the same rate of pay as the average weekly earnings of all Australian workers (seasonally adjusted) they would earn $968.10 per week\(^3\), equivalent to $49,818 per annum (ABS, 2010c). This average includes superannuation and overtime and penalty rates where earned.

For all primary carers, the earnings forgone are thus valued at $4.6 billion, and $1.9 billion for non-primary carers. Combined, the opportunity cost for all carers is $6.5 billion in 2010 (Table 2.3). This is equivalent to 0.5% of GDP and 9.5% of the value of formal health care.

### Table 2.3: Opportunity cost of informal care, primary and non-primary, Australia, 2010

<table>
<thead>
<tr>
<th></th>
<th>Primary carers</th>
<th>Non-primary carer</th>
<th>All carers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Difference in employment (%)</td>
<td>17.0</td>
<td>1.6</td>
<td>129.9</td>
</tr>
<tr>
<td>No. persons not employed due to caring responsibilities (000s)</td>
<td>91.7</td>
<td>37.2</td>
<td></td>
</tr>
<tr>
<td>Average weekly earnings ($)</td>
<td>968.10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lost earnings from lower workforce participation ($ millions)</td>
<td>4,616.2</td>
<td>1,873.5</td>
<td>6,489.7</td>
</tr>
</tbody>
</table>

Source: Access economics calculations.

Disaggregation into full-time and part-time employees would generate higher estimates due to the compositional effects – up to $12.3 billion for 2010. However this estimate must be used with caution, as the relatively small sample size for part-time caring employees reduces its robustness.

The relatively conservative $6.5 billion estimate is used in this report as the Access Economics estimate of the opportunity cost of informal care in 2010, both in order to achieve consistency with later disaggregation of costs and income support payments to primary and non-primary carers, and to avoid reliance on less robust estimates. The possibility of much higher estimates when compositional effects are accounted for, as well as the inability to calculate the value of lost leisure time, underscores the need to treat the opportunity cost estimate as a lower bound estimate of the value of informal care.

### 2.2 Replacement valuation

The replacement valuation method is based on the cost of buying the equivalent number of hours of informal care from the formal care sector. Section 1.2 above examined the average hours of care provided by primary carers depending on the level of disability of the main recipient of care. As would be expected, greater demands are placed on carers as a person’s level of disability rises (recall Chart 1.8).

These data are used as the basis for estimating the total hours of informal care provided by Australians in 2010. As data are reported in bands, and for primary carers only, it is necessary

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\(^3\) Most recent available at time of drafting.
to impute the average number of hours of care given per week by primary and non-primary carers. These imputed averages used for costing purposes are set out in Table 2.4.

**Table 2.4: Average hours of care provided per week by carer status**

<table>
<thead>
<tr>
<th>Carer Status</th>
<th>Average hours reported</th>
<th>Imputed average</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary carers</td>
<td>&lt;20 hours per week</td>
<td>10 hours per week</td>
</tr>
<tr>
<td></td>
<td>20 – 39 hours per week</td>
<td>29.5 hours per week</td>
</tr>
<tr>
<td></td>
<td>40+ hours per week</td>
<td>50 hours per week</td>
</tr>
<tr>
<td>Non-primary carers</td>
<td>n/a</td>
<td>5 hours per week</td>
</tr>
</tbody>
</table>

Source: Access Economics.

Applying these imputed average values to data from SDAC, it is possible to identify the average hours of informal care provided per week by primary carers, by the level of disability of the main recipient of care. As Table 2.6 shows, the more severe the level of disability, the greater the average number of informal care hours provided. The SDAC data also provide the proportion of primary carers caring for people with different levels of disability, and thus enable calculation of the replacement value of informal care provided by primary carers, when combined with data on average wage rates for workers from the formal care sector.

The estimate of the replacement value of care is sensitive to changes in the estimate of the wage parameter for alternate formal sector care. In this analysis, the unit cost used has been based on the wage of moderately skilled formal sector carers (supervised employees), based on the Australian and New Zealand Standard Classification of Occupations (ABS, 2006a). Full-time personal carers and assistants employed in the formal sector received an average wage of $22.30 per hour, or $869.60 for a 39 hour week in August 2008 (ABS, 2009). This is inclusive of personal income tax and superannuation, and includes payment of overtime for after hours work. However, the hourly rate received by employees does not account for on-costs such as the wages of supervisors, managers or administrative support staff or other capital overheads. Loadings are added for each of these additional costs, and for average wage growth between August 2008 (when the survey was last undertaken) and February 2010 (the most recent period for which estimates of average weekly earnings across all employees are available).

**Table 2.5: Replacement valuation of informal care, unit cost components**

<table>
<thead>
<tr>
<th>Component</th>
<th>% Loading</th>
<th>Hourly rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base rate per hour – August 2008</td>
<td></td>
<td>$22.30</td>
</tr>
<tr>
<td>Loading for growth in AWE May 2008 to February 2010</td>
<td>7.60%</td>
<td>$1.70</td>
</tr>
<tr>
<td>Loading for on-costs</td>
<td>6.70%</td>
<td>$1.61</td>
</tr>
<tr>
<td>Loading for capital</td>
<td>5.70%</td>
<td>$1.46</td>
</tr>
<tr>
<td>Loading for supervision &amp; administration</td>
<td>15.50%</td>
<td>$3.97</td>
</tr>
<tr>
<td><strong>Total hourly rate including overheads</strong></td>
<td></td>
<td><strong>$31.04</strong></td>
</tr>
</tbody>
</table>


The 6.7% loading for on-costs comprises of workers compensation, payroll and Fringe Benefits Taxation allowances (ABS, 2004a). Loadings for capital (5.7%) and administrative (15.5%)

---

4 Most recent available at time of drafting.
overheads are based on the relative shares of capital expenditure and administration costs to other areas of recurrent spending in Australia’s formal health sector (AIHW 2005, 2009a). When these loadings are added, the average hourly cost of employing a carer in the formal sector to replace an informal carer is estimated as $31.04 in 2010. These loadings reflect the fact that formal sector community care requires more than just labour inputs – there is a need to supervise and administer the care, often requiring a facility and vehicles. Travel costs have conservatively not been included, due to lack of data.

In 2010 the value of informal care provided by primary carers is estimated as $22.1 billion, of which $3.6 billion (16%) is informal care provided to people with a profound disability and $7.7 billion (35%) to people with a severe disability. The remaining $11.0 billion (49%) is informal care provided by primary carers to people with a mild or moderate disability, or a disability that does not affect core activities (including mobility and self-care). Again it is noteworthy that this is a conservative estimate of replacement valuation for primary carers. If compositional effects were ignored and the average hours of care provided by all primary carers were used instead (33.5 hours per week) then the estimate would be 940 million hours provided at a value of $29.2 billion per annum.

There are insufficient data in SDAC to determine the disability level of the recipients of care provided by non-primary carers or the average hours of care they provide. As non-primary carers are, by definition, not providing the majority of informal care, the replacement cost has been calculated based on an estimate that each of the 2.33 million non-primary carers in Australia in 2010 provide an average of five hours of care per week to people. On this basis the replacement value of care provided by non-primary carers is $18.8 billion per annum.

The total replacement cost of informal care in 2010 is thus estimated as $40.9 billion (equivalent to 3.2% of GDP and 60% of other formal health care). The table also shows that informal carers together provide an estimated 1.3 billion hours of care in 2010, providing nearly 460 hours of care per year or 9 hours per week, on average.

Table 2.6: Replacement cost of informal care, Australia 2010

<table>
<thead>
<tr>
<th>Level of Disability</th>
<th>Other</th>
<th>Severe</th>
<th>Profound</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary carers</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Av. hours of care per week</td>
<td>22</td>
<td>27</td>
<td>39</td>
<td></td>
</tr>
<tr>
<td>Number of primary carers (000s)</td>
<td>305.8</td>
<td>176.2</td>
<td>57.4</td>
<td>539.4</td>
</tr>
<tr>
<td>Total hours per annum (m)</td>
<td>350</td>
<td>247</td>
<td>116</td>
<td>714</td>
</tr>
<tr>
<td>Replacement cost ($m)</td>
<td>10,854</td>
<td>7,677</td>
<td>3,615</td>
<td>22,146</td>
</tr>
<tr>
<td><strong>Non-primary carers</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Av. hours of care per week</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Number of non-primary carers (000s)</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
<td>2,325.9</td>
</tr>
<tr>
<td>Total hours per annum (m)</td>
<td></td>
<td></td>
<td></td>
<td>605</td>
</tr>
<tr>
<td>Replacement cost ($m)</td>
<td></td>
<td></td>
<td></td>
<td>18,769</td>
</tr>
<tr>
<td><strong>Total replacement cost ($m)</strong></td>
<td></td>
<td></td>
<td></td>
<td>40,915</td>
</tr>
</tbody>
</table>

Source: Access Economics calculations.
2.3 Comparisons

Sections 2.1 and 2.2 above show how sensitive the estimated value of informal care is to the methodology. A replacement valuation approach yields an estimate of $40.9 billion in 2010, more than six times the estimate of $6.5 billion generated using the opportunity cost method. It is important to bear in mind that:

- the opportunity cost measure reveals the resources that are diverted each year from production in the formal economy to informal care; while
- the replacement cost measure reveals the additional care that would need to be provided each year by the formal economy to replace the work done by informal carers, were their services no longer available.

The replacement cost measure does not take into consideration differences in efficiency of providing care. Many informal carers only provide care to one person while a carer in the formal sector might be able to provide care to a number of people at the same time, if the care recipients were co-located. These economies of scale are the rationale behind residential care. However, many people prefer to stay in their own homes.

On the other hand, to provide the same number of hours of care as co-resident informal carers self-report, with the care recipient remaining at home, a formal carer might need to spend more hours overall, due to travel time. For other reasons also (e.g. inaccuracies in self-reporting, regulations in the formal sector requiring ‘round the clock’ care), the replacement cost estimate may not be a true reflection of the number of hours needed to provide the same level of care in the formal sector.

That said, both estimates are large numbers and this section provides a comparison with previous estimates of the value of informal care in Australia, and the estimated size of the other sectors in the Australian economy.

There have been previous attempts to estimate an imputed value for informal care. Chart 2.2 demonstrates the relative value, in 2010 dollars, placed on informal care in each of these studies. Care should be taken when comparing the studies, as there are significant methodological differences between them, as discussed below. Also, no allowance has been made for Australian population growth that has occurred since the earlier studies, which would increase the total stock of, and hence the value of, informal care.

A study by the Australian Institute of Family Studies (de Vaus et al, 2003) estimated the value of time spent by Australians in 1997 providing unpaid personal care to adults was $6.8 billion in 1997. A replacement valuation approach was used, combining the average hourly wage of a personal carer/nursing assistant with data from the ABS 1997 Time Use Survey (ABS, 1998). This estimate would not include other activities that informal carers may assist with, such as household chores.

Another study placed the value of all unpaid welfare work done during 2005-06, including informal care and volunteer work through organisations at $27.4 billion. The amount of informal care provided was determined from time-use surveys, and valued at the average hourly pay rate of an appropriately qualified person, in this case an adult employed as a full-time, community or personal services worker (AIHW, 2007).
Our opportunity cost valuation is lower than either of the other studies, which both use a replacement valuation approach. Both the previous studies, and the replacement valuation in this report, use the hourly wage of people employed as moderately skilled personal carers as the base for estimating the hourly value of care. However, the difference between previous estimates of replacement value, and that contained in this report is due to our use of:

- loadings for employee on-costs, supervision/administration and capital costs;
- alternative source data, which does not record time use quite as accurately, but is more representative in the sense that it has broad coverage of carers of the disabled and frail aged and
- a broader definition of care, including household tasks as well as personal care, which is appropriate in an economy operating at near full capacity, as Australia currently is.

**Chart 2.2: Value of informal care, Australia, selected studies, constant 2010 dollars**

<table>
<thead>
<tr>
<th></th>
<th>$ million 2010</th>
<th>%GDP</th>
<th>% formal health</th>
</tr>
</thead>
<tbody>
<tr>
<td>Opportunity cost – lower bound</td>
<td>6,490</td>
<td>0.5</td>
<td>9.5</td>
</tr>
<tr>
<td>Replacement cost – upper bound</td>
<td>40,919</td>
<td>3.2</td>
<td>59.7</td>
</tr>
<tr>
<td>Formal health sector(a)</td>
<td>68,541</td>
<td>5.4</td>
<td>100.0</td>
</tr>
<tr>
<td>GDP*</td>
<td>1,262,600</td>
<td>100.0</td>
<td></td>
</tr>
</tbody>
</table>

Notes: (a) 2009-10.
3 Other effects of informal care

The previous chapter showed how income is lower for carers, and associated with this there is also a reduction in taxation revenue. Moreover, in Australia, people with primary caring responsibilities may be eligible for income support payments from the Federal Government. In a budget-neutral setting these payments must also be funded by additional taxation. Taxation is a cost to Government and creates society-wide efficiency losses, which are estimated in this chapter.

3.1 Income support payments for carers

There are main two Federal Government assistance payments for carers – Carer Payment and Carer Allowance.

The Carer Payment is an income support payment for people who are unable to participate in the workforce full-time as a result of their caring responsibilities. The payment is means tested (with income and asset tests) and is paid at the same rate as other social security pensions. To be eligible, the carer must be providing care in the home of the person requiring care. There are two payments, paid at the same rate, but with different qualifying criteria.

1. To receive the Carer Payment (caring for a person 16 years or over) the carer must provide one of the following levels of care:
   - full-time care to an adult who has a disability or medical condition that is long term and severe and has a minimum level of care needs assessed by the Adult Disability Assessment Tool (ADAT); or
   - full-time care for an adult whose care requirements are less severe but also have a dependent child under 6 or a dependent child aged 6-16 with care needs that enable them to receive the Carer Allowance (caring for a person under 16 years).\(^5\)

2. The Carer Payment (caring for a person under 16 years) is paid to those who provide:
   - care for a child under 16 with a profound disability; or
   - care for between two and four children under 16 with a disability who, together, need a level of care that is at least equivalent to the level of care needed by a child with a profound disability; or
   - care to one or two children under 16 and an adult with a disability who, together, need a level of care that is at least equivalent to the level of care needed by a child with a profound disability; or
   - a significant amount of care to two or more children under 16 years of age each with a severe disability or a severe medical condition in an exchanged care arrangement (care is shared between two parents or legal guardians who are not in a relationship).\(^6\)

The Carer Allowance is an income supplement available to people who provide daily care and attention at home for a person who has a disability or severe medical condition or who is frail.


aged. Carer Allowance is not taxable or means tested. It can be paid in addition to wages or a social security income support payment. A carer is eligible to receive the Carer Allowance for up to two adults who meet the criteria. The care can be provided to people aged 16 years or over or to a child under 16 years who has a physical, intellectual or psychiatric disability (and hence the care is additional to what would be provided to a healthy child). Carer Allowance can also be paid to someone who cares for two children with disabilities and the children do not individually qualify but together create a substantial caring responsibility.

The adult being cared for must be likely to suffer from the disability permanently or for an extended period of at least six months (unless their condition is terminal or they are deemed to have substantial short-term care needs) assessed by a medical practitioner or other approved person meeting the medical eligibility criteria.

The maximum amount currently payable under each payment is set out in Table 3.1. Note that these amounts differ from the average payment rates set out in Table 3.4 for three reasons. First, Table 3.4 was based on an estimate of the total spending on each payment and the estimated number of recipients. The number of recipients is not constant throughout the year. Payment rates may be changed if the recipient’s circumstances change, such that the means test affects them differently. Second, a recipient can receive multiple Carer Allowances if they care for multiple people, and not all recipients of Carer Payment receive the maximum rate of payment. Third, the rate of payment is indexed to inflation and increased in March and September each year. This means that there are four rates of payment within each financial year.

**Table 3.1: Carer Payment and Allowance, rates, March 2010**

<table>
<thead>
<tr>
<th>Payment Type</th>
<th>$ per fortnight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carer Payment (single)</td>
<td>$644.20*</td>
</tr>
<tr>
<td>Carer Payment (couple)</td>
<td>$485.60 each*</td>
</tr>
<tr>
<td>Carer Allowance (adult)</td>
<td>$106.70 per adult cared for (max 2)</td>
</tr>
<tr>
<td>Carer Allowance (child)</td>
<td>$106.70 per child cared for</td>
</tr>
</tbody>
</table>


Recipients of the Carer Payment and Carer Allowance may also receive several additional support measures.

- **Carer Supplement** is a lump-sum payment of $600 paid at the beginning of each financial year. Those who receive both the Carer Payment and the Carer Allowance are eligible for a $1,200 payment. Carers receive a Carer Supplement for each person for whom primary care is provided.

- **Rent Assistance** is a fortnightly allowance that assists with the cost of accommodation. Eligibility depends on type and cost of accommodation and is means-tested.

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**Child Disability Assistance Payment (CDAP)** is a $1,000 annual lump-sum payment for carers of children who are receiving Carer Allowance.

In 2008-09, the Federal Government spent a total of $4.29 billion on income support for carers (Table 3.2). The budgeted cost of carer support payments is expected to increase around 6.9% per annum to $4.9 billion in 2010-11. The contraction of total carer income support in 2009-10 is a result of some 2009-10 payments being brought forward into the 2008-09 financial year.

Table 3.2: Carer Payment and Allowance, payments, 2008-09 to 2010-11 ($m)

<table>
<thead>
<tr>
<th>Carer Allowance</th>
<th>$1,801.01</th>
<th>$1,470.50</th>
<th>$1,569.34</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carer Payment</td>
<td>$1,938.83</td>
<td>$2,361.66</td>
<td>$2,738.02</td>
</tr>
<tr>
<td>Carer Supplement</td>
<td>$408.03</td>
<td>-</td>
<td>$443.00</td>
</tr>
<tr>
<td>CDAP</td>
<td>$142.79</td>
<td>$148.77</td>
<td>$152.76</td>
</tr>
<tr>
<td><strong>Total Carer Income Support</strong></td>
<td><strong>$4,290.65</strong></td>
<td><strong>$3,980.93</strong></td>
<td><strong>$4,903.12</strong></td>
</tr>
</tbody>
</table>

Source: FaHCSIA (2009b).

In 2008-09 a Carer Bonus was also distributed to recipients of the Carer Payment and Carer Allowance. The Commonwealth expects that payment of carer bonuses will cost a total of $428.1 million (FaHCSIA, 2009b). Due to the ‘once-off’ nature of these bonuses they are not included in Table 3.2.

In 2008-09 (the latest year for which data are available) there were 146,870 Carer Payment recipients who received a total of over $1.9 billion in benefits – an average payment of $258.85 per week indexed to 2010 (FaHCSIA, 2009a). Over the same period 461,023 Carer Allowance recipients received over $1.8 billion or $76.60 per week indexed to 2010. In June 2009, $408 million was paid through the Carer Supplement (FaHCSIA, 2009a). It is noted that 91% of people receiving a Carer Payment also receive a Carer Allowance (FaHCSIA, 2008). Table 3.3 summarises the estimated number of recipients of each payment in 2008-09, and expresses the percentage that are primary carers based on the estimated number of primary carers in 2008-09 (based on SDAC parameters).

Table 3.3: Recipients of income support for carers, 2008-09

<table>
<thead>
<tr>
<th>Number of recipients</th>
<th>% primary carers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carer Payment only</td>
<td>13,218</td>
</tr>
<tr>
<td>Carer Payment and Carer Allowance</td>
<td>133,652</td>
</tr>
<tr>
<td>Carer Allowance only</td>
<td>327,371</td>
</tr>
<tr>
<td><strong>Total Recipients</strong></td>
<td><strong>474,241</strong></td>
</tr>
</tbody>
</table>

Source: Access Economics estimates.

The expected number of recipients in 2010 is calculated by assuming the same ratio between the number of payment recipients and the total number of primary carers as calculated in the last column of Table 3.3. Since there were increases in Carer Payment, particularly for singles, that were outside the increase for consumer price inflation, the average weekly payment for 2008-09 is indexed by growth in the payment rate.
If the same proportion of primary carers in 2010 receive Carer Allowance, Carer Supplement and CDAP then the total value of scheduled welfare payments to carers for caring is estimated to be around $4.8 billion this year. The associated efficiency cost from the additional taxation required to finance these transfers is estimated as $1.4 billion (Table 3.4), with the 28.75% efficiency cost calculation described in more detail in the next section. The role of income support payments in distributing the burden of informal care between the individual and society (through government assistance) is outlined in more detail in Section 4.

<table>
<thead>
<tr>
<th>Table 3.4: Total cost of income support to carers, Australia, 2010</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Av. Weekly payment</strong></td>
</tr>
<tr>
<td>Carer Payment</td>
</tr>
<tr>
<td>Carer Allowance</td>
</tr>
<tr>
<td>Carer Supplement</td>
</tr>
<tr>
<td>CDAP</td>
</tr>
<tr>
<td><strong>Total</strong></td>
</tr>
</tbody>
</table>

Notes: (a) Average payment for 2008-09 (FAHCSIA 2009a) indexed for growth in average payment. (b) Assuming a payment of $600 per recipient of Carer Payment and Carer Allowance and $1,200 for recipients of both. (c) Assuming a payment of $1,000 per carer receiving Carer Allowance (Child).

Source: Access Economics calculations.

### 3.2 Forgone taxation and efficiency costs

The opportunity cost valuation of informal care essentially measures the productivity losses which flow from the direction of labour to informal care rather than other productive activities. In 2010 this productivity loss is estimated at $6.5 billion (recall Table 2.3). The reduced wage income of carers has other indirect or flow-on effects to the economy, through the taxation and welfare systems. People who are not working generally pay less tax and are more likely to receive an income support payment from the government.

Taxation and income support payments are transfer payments, not real economic costs. Payments transfer wealth, and hence consumption power, between individuals in society, but the total wealth or available resources in the community as a whole do not change. However, in reality transfers are not costless. Resources are consumed in the administration of the taxation and welfare systems. Administration of the taxation system costs around 1.25% of total revenue raised. More importantly, taxation induces distortions in relation to people’s work, leisure and consumption choices. The cost of these distortionary impacts is estimated to be equivalent to 27.5% of each additional taxation dollar raised (Lattimore, 1997; Productivity Commission, 2003a).

People who are not in paid employment due to their caring responsibilities will pay less personal income tax. There is also likely to be a fall in indirect (consumption) tax paid, as people on lower incomes generally purchase a lesser amount of goods and services (consumption is assumed to be 90% of after-tax income here). In 2010, a total of $1.29 billion of potential tax revenue is estimated to be forgone, due to the lower workforce participation of carers. Of this $1.24 billion (96%) is forgone personal tax and the remaining $45 million (4%) is forgone indirect tax; the latter calculation is based on 90% consumption of the difference between lost earnings after income tax and welfare payments (estimated in Table 3.4).
In a budget neutral setting, this forgone taxation revenue will need to obtained from other sources. As mentioned, each extra dollar of taxation imposes 28.75 cents of real efficiency costs on the economy. In 2010, $369 million of efficiency costs is estimated to be incurred due to the additional taxation required to replace revenue forgone due to the lost productivity of carers (Table 3.5).

<table>
<thead>
<tr>
<th>Table 3.5: Efficiency cost from additional taxation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value of earnings forgone</td>
</tr>
<tr>
<td>Average personal income tax rate</td>
</tr>
<tr>
<td>Potential personal income tax forgone</td>
</tr>
<tr>
<td>Average indirect tax rate</td>
</tr>
<tr>
<td>Potential indirect tax lost</td>
</tr>
<tr>
<td>Total potential tax revenue lost</td>
</tr>
<tr>
<td>Efficiency cost per dollar taxation</td>
</tr>
<tr>
<td>Efficiency cost from additional taxation</td>
</tr>
</tbody>
</table>


Together with the efficiency losses from welfare payments, the total efficiency costs from raising alternative taxation revenue are estimated to be 1.76 billion in 2010.

3.3 Health and wellbeing impacts on carers

Another important indirect cost is the impact of caring responsibilities on the health and wellbeing of the carer. ABS (2005) found that 72.4% of Australia’s primary carers experience some form of physical or emotional effect from providing care. There is ample evidence that this effect is predominately negative. Nevertheless, despite the well-documented negative impact on both mental and physical health many carers report a positive effect on their relationship with the care recipient.

Carers have a higher incidence of mental illness than the general population. Clinical studies have shown increased rates of depression among caregivers, compared to control populations (Spector and Tampi, 2005). The Australian Unity Project surveyed 3,766 carers in July 2007 and found that, on the Depression Anxiety Stress Scale, 56% of carers were classed with at least moderate depression compared with only 6% of non-carers. In ABS (2005) 29.2% of carers reported that they often feel worried or depressed. Depression is a particularly costly condition, accounting for the largest share of Australia’s disability burden measured by years of healthy life lost due to disability (Mathers et al, 1999).

Stress, a risk factor for a number of diseases (e.g. cardiovascular disease) is also common among carers. Over 10% of primary carers have a stress related illness (ABS, 2005). High levels of stress can make it more difficult to deal with pain. Based on a wellbeing index Cummins et al (2007) found that pain has a greater effect on the wellbeing of a carer than it does on a non-carer.

As well as having an emotional impact, caring can also cause physical injury, especially musculoskeletal injuries from incorrectly lifting, lowering, carrying or moving the recipient of...
care. A 1999 survey into the wellbeing of carers conducted by Carers Australia found that 33% of all carers reported having being physically injured at least once in the course of providing care, of which over a half were due to lifting or carrying the patient or other objects (Carers Association of Australia, 2000). Similar results have been found in overseas studies (Carers UK, 2004).

Although health is an important component of wellbeing, family relationships and life satisfaction also play an important role. ABS (2005) found that more carers report feeling that giving care has brought them closer to the care recipient than those who report that it has strained their relationship (34% compared with 18%). Just over a quarter (25.7%) of primary carers reported that they felt satisfied with their caring role (implying that three quarters were not satisfied). Adverse effects on wellbeing were reported by 29% of carers (ABS, 2005).

Sleep deprivation can contribute to a number of health conditions and reduce overall wellbeing. Often feeling weary or lacking energy as a result of being a carer was reported by 33.7% of primary carers in ABS (2005). Sleep interruptions due to the requirements of their caring role were reported by 44.5% of carers. Of these, 46.4% reported that these interruptions were frequent. Hillman et al (2006) estimated that almost 1.2 million Australians experience sleep disorders with associated costs in excess of $10 billion in 2004, through the costs of sleep disturbance itself and its associated attributable costs from fatigue-related motor vehicle accidents, work-related injuries, depression, diabetes and cardiovascular disease. On a relative prevalence basis then, the sleep impacts of caring may exceed $1 billion per annum, including a financial estimate of the reduced quality of life.

**Case study model: health impacts of caring**

In this study, a hypotheticated female carer ‘Karen’ in her mid-forties has conditions typical of caring as shown in the literature above –mild-moderate depression, stress-related angina (a cardiovascular condition), and chronic lower back pain. As the primary carer for her disabled child, aged 19, Karen no longer works due to her caring responsibilities and her own health conditions.

The cost impacts of each of these health conditions have been previously estimated by Access Economics (2009a, 2007a, 2007b) – not just the health system expenditures, but also the productivity losses, other financial costs (such as efficiency losses and State Government programs), and the burden of disease.
The disease burden is measured in disability adjusted life years based on disease-specific weights published by the AIHW (Begg et al, 2007; Mathers et al, 1999). These are then converted to 2010 dollars using the value of a year of healthy life of $166,604 from the Department of Finance and Deregulation (2009).

The costs in 2010 dollars are shown in the table below, while the chart splits by who bears the cost – the Australian Government, Karen herself, or other entities in society (e.g. all Australians share the efficiency losses). The total financial impact of all three conditions is over $33,000 in 2010. Adding in the cost of the loss of Karen’s healthy life (about 27% of a healthy life year is lost to the conditions) the total impact is nearly $79,000.

<table>
<thead>
<tr>
<th>2010 $</th>
<th>Depression</th>
<th>Angina</th>
<th>Back pain</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Health costs</td>
<td>3,818</td>
<td>154</td>
<td>2,419</td>
<td>6,391</td>
</tr>
<tr>
<td>Productivity losses</td>
<td>17,992</td>
<td>286</td>
<td>4,043</td>
<td>22,321</td>
</tr>
<tr>
<td>Other financial costs</td>
<td>3,491</td>
<td>117</td>
<td>1,417</td>
<td>5,025</td>
</tr>
<tr>
<td><strong>Total financial costs</strong></td>
<td><strong>25,301</strong></td>
<td><strong>557</strong></td>
<td><strong>7,879</strong></td>
<td><strong>33,737</strong></td>
</tr>
<tr>
<td>Loss of wellbeing</td>
<td>41,162</td>
<td>83</td>
<td>3,974</td>
<td>45,219</td>
</tr>
<tr>
<td><strong>Total cost per annum</strong></td>
<td><strong>66,463</strong></td>
<td><strong>640</strong></td>
<td><strong>11,853</strong></td>
<td><strong>78,956</strong></td>
</tr>
</tbody>
</table>

Karen bears 56% of the financial costs (mainly lost income but also health costs over $1,000), while the Government bears 21%. Karen bears all the wellbeing loss.

Due to her caring responsibilities, Karen struggles to get the physical activity that her doctor recommends would assist in managing her health conditions. She finds she has little time or money to look after herself and her situation would be helped if appropriate respite care was more accessible in her rural town.
3.4 Young carers

Young carers are people under the age of 25 who provide informal care (Hill et al, 2009). Little formal research has been undertaken into the impact of providing care from a young age, in part due to the difficulties involved in identifying young carers. Consequently the indirect costs uniquely associated with providing care from a young age are often overlooked. The situation of young carers is distinctive because care is provided during the years in which a person would typically undertake education and establish themselves in the labour force. As such, providing informal care from a young age increases the risk of long term financial disadvantage.

Providing informal care can interfere with schooling and work because it is time intensive and stressful. Many young carers undertake the same tasks as an older carer would – including housework, assisting with bathing and toileting, administering medication and providing emotional support. Although many young people contribute to households tasks, young carers do so with a much greater intensity and over a wider range of tasks than other young people. A UK study revealed that 17% of young carers provide more than 20 hours per week of support to their family (Warren, 2007).

It is difficult to accurately estimate the number of young carers for several reasons. First, research suggests that a large proportion of young carers do not access support services remaining ‘hidden’ (Cass et al, 2009). Second, it is difficult to obtain survey data on people under the age of 18. In many surveys, including the SDAC, people under the age of 15 are not interviewed and those under 18 require parental consent to participate. Accordingly, information on minors is usually collected from the parents. However, parents may not be forthcoming with information about their child’s care giving responsibilities due to the social stigma attached. ABS (2005) showed that 62% of young carers in Australia were providing informal care to a parent. Finally, young people are less likely to identify themselves as carers than would be an older person providing the equivalent amount of care because they perceive their responsibilities as ‘normal’ (Cass et al, 2009).

Presently the best estimate of the number of young carers comes from the SDAC. In 2010 it is estimated that there are around 375,071 young people in Australia providing informal care to a relative or friend with a disability, of which 20,369 are primary carers (Table 1.2). Due to the sampling difficulties discussed above this is likely to be an underestimate. There has been some work done in estimating potential young carers. Hill et al (2009) defined potential young carers as those aged 15-24 who reside with someone who requires care but no one in the household has identified themselves as a carer. For children aged 12-14 the criteria is co-residence with a person requiring care regardless of whether or not another carer had been identified. Using data from the 2006 Census of Population and Housing, Hill et al (2009) identified 14,000 potential young carers aged 12-14 and 77,000 aged 15-24. This is lower than Cass et al (2009) who identified 717,000 potential young carers using Wave 5 of the HILDA survey, although the definition of a carer was less strict, only requiring a person to co-reside with someone likely to require care.

Young carers may be at greater risk of not completing high school, or poor achievement, than non carers. ABS (2008a) found that 66% of young carers aged 19-24 had completed high school compared to 73% of non-carers in this age group. This estimate should be interpreted
with caution as it suffers from some self acknowledged methodological issues. Although there is inconclusive statistical evidence on the educational difficulties faced by young carers, there is supportive anecdotal evidence. Dearden and Becker (2002) found that at least a quarter of young carers in the UK experienced educational difficulties including:

- worry about how the person that they care for is faring in their absence;
- tiredness, difficulty concentrating;
- difficulty completing homework on time and often being punished for it;
- social isolation and difficulty identifying with their peers;
- being bullied; and
- embarrassment about having a disability in their family or about being financially disadvantaged.

Many of these factors also reduce academic achievement, which is an important precursor to dropping out of education. Through interviews with 51 young carers, Moore et al (2006) confirmed that these factors also applied to Australia. However, they also found that being a carer could have a positive influence on education. Young carers develop valuable life skills through their caring role that are not obtained through formal education. They may also be motivated to persevere with education because they are financially aware and understand the need to secure employment in the future. Others did not want to allow caring to limit their future or they wanted their parents to be proud of them.

Poor school results are an obvious barrier to opportunity to undertake higher education. However, even those who are able to achieve in high school may face reduced opportunity as a result of their caring role for the following reasons.

- Young carers may be financially responsible for themselves and the person who they care for and may feel that they have no choice but to find paid employment.
- Young carers require flexibility in their education program and not all institutions or course convenors are willing to accommodate their needs (Carers Australia, 2002).
- Young carers who live in regional areas may not have access to the course of their choice and leaving or relocating the person who they care for might not be an option.

The lasting effect of missed education opportunities is a greater chance of financial disadvantage. It may also be difficult for young carers to establish themselves in the workforce, even if they do obtain a tertiary or vocational qualification. While non-carer counterparts are gaining valuable work experience and establishing their career path, this can be more difficult for young carers. Failure to gain work experience can make it more difficult to find successful employment in the future. Many young carers come from families who are

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9 This measure is an ex post measure, meaning that it does not imply causality. Since it does not control for whether or not people were young carers during high school, it could indicate that being a young carer increases the risk of not completely high school or that not completing high school increases the chance of being a young carer. This estimate is the most reliable available. Other estimates look at the proportion of young carers who are studying. None of these estimates adequately control for differences in the age distribution of the carer and non carer sample, which will affect the findings because the probability of being in education is different at different ages.
reliant on income support and without providing them with opportunity to change their circumstances, the cycle of poverty will continue.

Another important barrier to education for young carers is the 25 hour rule on the Carer Payment. Young carers who are eligible for the Carer Payment are not allowed to spend more than 25 hours per week in study or paid employment, including travel time. This means that they cannot study full-time, prolonging the length of time that they must juggle study and caring. By choosing to study, they may also have to forgo the income that they are allowed to earn to supplement their Carer Payment. The inclusion of travel time compounds the problem for young carers living in regional areas.
Case study: educational opportunities for young carers

Chantelle, now 21, has from the age of 12 been the primary carer for her mother (aged 50), who has congenital nephritis. Chantelle manages her mother’s many medical appointments, medication and personal care at home, and all the family’s domestic household work and finances (her older sister and estranged father are not co-resident). Her mother’s illness can be unpredictable, causing emergency hospitalisations, night-time sleep interruptions and periodic absences from education for Chantelle.

Chantelle did not identify herself as a carer until age 17. She simply did what she ‘had to do’ to make ends meet, including working after hours through high school to pay for rent, food and medical bills – which were particularly high after her mother’s two nephritis-related strokes (her mother’s DSP was inadequate to meet expenses and Chantelle was too young to receive any payments for caring). The strokes occurred when Chantelle was at critical points in her education, during Years 9 and Year 11.

Her mother’s illness complicated Chantelle’s exam and assignment deadlines, and teachers were not understanding of her missed classes, telling her she did not have her priorities right. Chantelle found this difficult as, through high school, she slept on average only four hours per night in order to discharge her study, work and caring duties. She developed an eating disorder at one point, which she now has under control.

Despite the obstacles, Chantelle’s resilience, optimism and diligence led to her being School Captain in Year 12 and being offered a place at the university of her choice to study Medicine. This had been her career goal since her mother’s diagnosis. Although it was difficult juggling lectures, laboratory pracs and her caring duties, she developed strategies to enable her to meet her study and caring schedules, as she had at high school with study and work.

However, after her first months in Medicine, Centrelink notified her that, due to studying more than 25 hours per week (essential for her course), she would lose her Carer Payment. Chantelle explored various options hoping to be able to continue her study, but the regulated financial constraints meant she was unable to complete her education and continue to care for her mother. She made the difficult decision to postpone her dream, possibly forever. Her family came first.

With regret, Chantelle transferred to a Psychology degree that she could undertake part time in order to meet Centrelink’s requirements. Now in her third year, Chantelle encourages policy-makers to reconsider current funding arrangements so that young carers can complete their chosen study, saying: ‘Education is everything – what 16-year old should have to work three jobs to finish high school?’ She also encourages greater awareness raising for teachers, as she says the statistics suggest there is a ‘carer in every class’, yet they are still an invisible minority whose needs should be mainstreamed like those of other disadvantaged groups.
Access Economics has calculated expected lost lifetime earnings for Chantelle, based on a combination of data sources. Psychology is a shorter degree than Medicine, and although Chantelle has to complete it part-time, the costing is conservatively based on the usual setting of qualifying as a psychologist three years earlier (at age 22) than as a doctor (age 25). For the initial years after graduation, NSW State Award rates are imputed as earnings, which increase based on years of service. These rates are used until they reach average earnings population-wide for psychologists and doctors, as estimated from ABS sources (the census data on weekly income by occupation, and more recent earnings data by broader occupational group). The chart below shows the income differential that Chantelle can expect, in real 2010 prices. As a psychologist she can expect to only earn less than 60% of what she could have as a doctor i.e. $3.1 million rather than $5.45 million over a lifetime, in real 2010 dollars. The difference is $2.35 million, comprising lost tax revenue of $0.5 million and a loss to Chantelle of $1.9 million.\(^{10}\)

\(^{10}\) With a 3% real discount rate, the present value of the difference in lifetime earnings (to age 65) is $1.1 million.
4 Who bears the cost of care?

Section 3.1 estimated the cost of income support to carers, estimated in total as $4.82 billion in 2010. In addition, there are payments made for education and training of carers, respite and other supportive program payments. This chapter assesses the relative contributions from government for informal and formal carers.

4.1 Support programs for carers

Support programs are generally aimed at reducing the amount of time a carer spends on providing care, enhancing quality of life for the carer and the care recipient, and delaying the need for institutionalisation. Services include respite care, and also interventions that reduce the dependence of care recipients such as memory clinics for patients with dementia and activity therapy programs designed to improve day-to-day activities (Sorenson, 2002).

Formal support programs are available to carers through programs such as:

- the Home and Community Care (HACC) program;
- Veterans’ Home Care;
- Community Aged Care Packages (CACP);
- Extended Aged Care in the Home (EACH) including dementia-specific places (EACH-D); and
- Commonwealth State/Territory Disability Agreement (CSTDA) services; and
- other government aged care and disability programs.

These are designed to assist carers through respite care, counselling, education, and care planning and coordination services.

The National Respite for Carers Program is the main Commonwealth government program that funds respite services for carers. Respite may be provided at the carer’s home, or in other residential settings such as day centres, residential overnight cottage-style accommodation or aged care facilities. It is estimated that around $202 million was spent in 2009-10 by the Australian Government through the National Respite for Carers Program, equivalent to an average of $374 per annum for each primary carer (Department of Health and Ageing, 2009). The different components of the program are summarised below.11

- Respite and Carelink Centres provide information and coordinate a national network of over 600 community-based respite services to assist carers in accessing general, short-term and emergency respite services.
- The Network of Carer Associations provides carers with professional counselling, specialist advice and information.

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The National Carer Counselling Program provides counselling on a sessional basis and aims to address issues specific to carers such as carer stress, grief and loss, coping skills and transition issues.

Only a relatively small number of carers make use of respite care. In 2007-08 13.7% of potential respite care users accessed a service and the average length of stay for residential respite care was 3.3 weeks in that year (AIHW, 2009b). On average, each user received 10.5 hours of respite care per week. Respite care can also be received in-home for short periods of time.

Carer Resource Centres provide information, support and advice to carers on a range of issues. Centres are located in each capital city and can be reached through a Freecall number.

In addition, over the period 2005-2009, the Australian Government allocated over $320 million to Dementia- a National Health Priority initiative, which included a number of programs that assist families and carers of people with dementia, as well as training programs and resources for care workers and health professionals working with people with dementia, such as:

3. the National Dementia Support Program (including a helpline and referral service, memory and community centres, early intervention and counselling, information awareness education and training, and support for people with special needs);
4. Dementia Behaviour Management Advisory Services, which assist informal and formal carers in the management of behavioural and psychological symptoms of dementia; and
5. four Dementia Training and Study Centres, which raise awareness of dementia issues in carer training.

While it is essential to provide a broad range of respite and support services, a particular area of focus of this report is in providing education and training to carers, which we review in the next section.

4.2 Education and training

As discussed in the previous chapters, the responsibilities associated with being a carer may impact on carers’ mental and physical wellbeing. According to a survey of over 1000 carers, 30% rated their health as fair or poor and 49% reported a long-standing physical health problem (Cormac and Tihanyi, 2006). As well, 13% had consulted a GP in the past year for anxiety, depression or emotional problems, with 71% reporting that their caring responsibilities made them feel worried. Kasuya et al (2000) has defined the carer burden as:

... a multidimensional response to physical, psychological, emotional, social, and financial stressors associated with the caregiving experience.

In light of this, it is important to improve accessibility to carer education and training programs to reduce the carer burden. Studies have shown that providing training for carers of stroke patients reduced costs of care while improving overall quality of life and psychosocial outcomes in carers at one year follow up (see for example, Patel et al, 2004; Kalra et al, 2004). Also, carer training and support may reduce the need for elderly care recipients to enter a nursing home, thereby reducing the cost of long-term care (Etters et al, 2008). According to the Resources for Enhancing Alzheimer’s Caregiver Health (REACH), a large randomised trial
studying the effectiveness of interventions for carers of people with dementia, carers in the intervention group had a significantly lower carer burden score.

In a cost effectiveness analysis of a behaviour intervention program using the REACH II study with carers of patients with Alzheimer’s, it was found that this educational intervention provided carers with an extra one hour per day not spent in caregiving, at a cost of $5 per day. This means the additional cost to bring about one additional unit of benefit (i.e. the reduction in hours per day of providing care) is approximately $5. This is significant as it suggests that, at a relatively low additional cost, the intervention provided carers with extra time that they did not have to spend providing care.

Education and training generally aim to enhance the wellbeing and coping skills of carers by providing information about the care recipient’s condition, as well as providing a network of resources to allow the carer to respond effectively to problems that arise. There is thus a close nexus with support interventions – such as professional counselling with a trained professional or peer group support groups that focus on building rapport among carers and creating a forum through which they can discuss their problems, successes and feelings – which may reduce the stress and relieve the psychological conflicts that caregiving can bring.

Naturally, some educational interventions have primarily domain-specific effects rather than a generalised global effect (Sorensen et al, 2002). For example, providing education and training for carers about the care recipient’s condition and symptoms may improve their caregiving ability, but not necessarily improve their ability to cope with stress and other negative impacts on their mental health, which requires more specific interventions such as counselling and psychoeducation.

Other training interventions aim to cover multiple educational aspects, and can involve new technologies. An example is a project developed in the European Union involving Sweden, England, Northern Ireland, the Republic of Ireland and Portugal (Assisting Carers Using Telematics Interventions to meet Older persons’ Needs). The project involved setting up technology in the homes of a number of family carers – such as a television and remote control, a small set-top box with CD-ROM and a videoconferencing card, a small video-camera lens and an ISDN telephone line. These services provided information and education using multi-media programs related to caregiving, and carers and care recipients could make direct contact with professional carers and other support networks via a video-telephone. The trial found that the use of the technology increased carers and care recipients’ sense of control and provided direct and, at times, immediate avenues for support in the provision of care (Magnusson et al, 2002).

In summary, the provision of training and support programs for carers has been shown to be beneficial for their health and wellbeing, and programs need to be well targeted (Sorensen et al, 2002) with consideration given to:

- the intensity of the program such as group or individual sessions and the number of sessions;
- the extent to which participants adhere to the program including drop-out rates and regularity of attendance; and
- the type of relationship between the carer and the care recipient such as spousal, adult or young carer.
Currently in Australia, there are a substantial investment in education and training for formal sector (paid) care workers, for example:

- **the Better Skills For Better Care Program**, which funds the education and training of RAC workers through courses including Certificate III in Aged Care, Certificate IV in Aged Care, Certificate IV in Aged Care Lifestyle and Diploma in Nursing (Enrolled Nurse Qualifications) at around $29 million per annum;

- **the Support for Aged Care Training** program, which provides training to upgrade skills of personal care workers in 600 smaller regional RAC services, cost $30 million over the four years to June 2010;

- **the Community Aged Workforce Program**, which supports training for personal care workers involved in the delivery of Australian Government-subsidised CACP, EACH and EACH-D packages, with some 2,000 to 3,000 places per annum offered in Certificate III in Home and Community Care and Certificate IV in Service Coordination; and

- **other programs specifically for education in dementia care** – such as Dementia Care Essentials.

Information on subsidies for the education of paid carers is presented in Table 4.1. The subsidies for a personal carer and enrolled nurse are based on the difference between the international\(^{12}\) and domestic fees\(^{13}\) for a Certificate III in Aged care and Diploma of nursing respectively at some randomly selected institutions. The subsidy for a RN is based on the difference between the average of the fee rates charged to international students at 13 Australian universities\(^{14}\) and the HECS-HELP contribution for 2010 (http://www.goingtouni.gov.au).

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\(^{14}\) Universities were included based on the share of their state or territory’s nursing graduates that they produced in 2008 (DEEWR, 2008). Institutions were included in order of magnitude (largest to smallest) until 70% of graduates the state or territory were accounted for. The universities included were Charles Sturt University (http://datasearch.uts.edu.au/international/prospective/studying/fees/course_fees.cfm), University of Technology Sydney (http://www.csu.edu.au/division/finserv/fees/future/internat_onshore.html), University of Western Sydney (http://www.uow.edu.au/future/international/apply/fees/index.html), Deakin University (http://www.deakin.edu.au/future-students/international/fees.php#tuition_fees___1), La Trobe University (http://www.latrobe.edu.au/coursefinder/international/2011/Bachelor-of-Nursing-%28Pre-registration%29.6584.html), Monash University (http://www.monash.edu.au/study/coursefinder/course/0727/), Victoria University (http://www.vu.edu.au/courses/international/bachelor-of-nursing-hbhn), Griffith University (http://www17.griffith.edu.au/cis/p_cat/fees.asp?ProgCode=1165&FeeType=Undergraduate), Queensland University of Technology (http://www.courses.qut.edu.au/cgi-bin/WebObjects/Courses.woa/wa/selectMajorFromMain?courseID=9631), The Flinders University of South Australia (http://www.flinders.edu.au/courses/undergrad/bngu/), University of Australia (http://www.unisanet.unisa.edu.au/programs/program.asp?Program=IBNU&Year=2010), University of Tasmania (http://www.international.utas.edu.au/static/HowtoApply/ApplicationsforBachelorofNursing.php), Australian Catholic University (http://www.acu.edu.au/courses/undergraduate/nursing/bachelor_of_nursing/). Information on full fees could not be found for Central Queensland University, The University of Newcastle, University of Southern Queensland, Curtin University of Technology, Edith Cowan University, University of Canberra and Charles Darwin University – all of which were eligible for inclusion.
The economic value of informal care in 2010

### Table 4.1: Subsidies for education of paid carers, 2010

<table>
<thead>
<tr>
<th>Paid carer</th>
<th>2010</th>
</tr>
</thead>
<tbody>
<tr>
<td>Personal carer (Certificate III)</td>
<td>$5,029</td>
</tr>
<tr>
<td>EN (Diploma)</td>
<td>$18,901</td>
</tr>
<tr>
<td>RN (Bachelors Degree)</td>
<td>$37,590</td>
</tr>
</tbody>
</table>

Source: Access Economics estimates.

In contrast to the education and training provided to formal care workers, there is relatively little funded training for informal carers. Data were insufficient to estimate the difference in expenditures, and some funded services do exist which support unpaid carers and provide education as one of their functions (such as the National Dementia Support Program), but these tend to be quite small and fragmented relative to the scale and national platform of investments in education for formal carers.

The box below provides a cost benefit analysis of an education and support program, based on the detailed information on the efficacy of such an intervention in the UK from Patel et al (2004), together with Australian unit cost data, as shown in Table 4.2.
Table 4.2: Input data for cost benefit analysis of a carer education and training interventions

<table>
<thead>
<tr>
<th>Services</th>
<th>Average number of services</th>
<th>% accessing service</th>
<th>Unit cost</th>
<th>Difference</th>
<th>Australian source</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Training</td>
<td>No training</td>
<td>Training</td>
<td>No training</td>
<td>A$2010</td>
</tr>
<tr>
<td>Stroke unit</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Therapy</td>
<td>30.8</td>
<td>43.2</td>
<td>100.0%</td>
<td>100.0%</td>
<td>1,141</td>
</tr>
<tr>
<td>Physiotherapist per personal interaction unit</td>
<td>115.1</td>
<td>145.3</td>
<td>100.0%</td>
<td>100.0%</td>
<td>59</td>
</tr>
<tr>
<td>Occupational therapist per personal interaction unit</td>
<td>9.3</td>
<td>12.4</td>
<td>99.3%</td>
<td>100.0%</td>
<td>59</td>
</tr>
<tr>
<td>Speech/language therapist per minute patient contact</td>
<td>6.7</td>
<td>5.3</td>
<td>57.6%</td>
<td>55.0%</td>
<td>3</td>
</tr>
<tr>
<td>Secondary care</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>General medical ward per day</td>
<td>10.9</td>
<td>12.8</td>
<td>8.2%</td>
<td>7.9%</td>
<td>1,141</td>
</tr>
<tr>
<td>Outpatient visit</td>
<td>2.2</td>
<td>2.3</td>
<td>42.5%</td>
<td>42.1%</td>
<td>261</td>
</tr>
<tr>
<td>Accident and emergency per visit</td>
<td>3</td>
<td>1</td>
<td>0.7%</td>
<td>2.4%</td>
<td>391</td>
</tr>
<tr>
<td>Day hospital per visit</td>
<td>5.6</td>
<td>7.2</td>
<td>28.4%</td>
<td>19.0%</td>
<td>81</td>
</tr>
<tr>
<td>Social services</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Personal care per hour</td>
<td>56</td>
<td>50</td>
<td>37.1%</td>
<td>33.6%</td>
<td>31</td>
</tr>
<tr>
<td>Domestic assistance per 1 hour contact</td>
<td>22</td>
<td>26</td>
<td>14.6%</td>
<td>17.4%</td>
<td>31</td>
</tr>
<tr>
<td>Laundry assistance per 1 hour contact</td>
<td>10</td>
<td>15</td>
<td>4.0%</td>
<td>6.0%</td>
<td>31</td>
</tr>
<tr>
<td>Shopping assistance per 1 hour contact</td>
<td>6</td>
<td>9</td>
<td>12.6%</td>
<td>11.4%</td>
<td>31</td>
</tr>
<tr>
<td>Meals on wheels per meal</td>
<td>19</td>
<td>17</td>
<td>10.6%</td>
<td>16.1%</td>
<td>8</td>
</tr>
<tr>
<td>Carelink per 15 minute contact</td>
<td>16</td>
<td>24</td>
<td>6.6%</td>
<td>10.1%</td>
<td>8</td>
</tr>
<tr>
<td>Social services day care centre per session</td>
<td>14</td>
<td>26</td>
<td>9.3%</td>
<td>17.4%</td>
<td>78</td>
</tr>
<tr>
<td>Other community based care</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>General practitioner per surgery visit</td>
<td>84</td>
<td>68</td>
<td>62.7%</td>
<td>54.4%</td>
<td>60</td>
</tr>
<tr>
<td>General practitioner per home visit</td>
<td>47</td>
<td>54</td>
<td>35.1%</td>
<td>43.2%</td>
<td>102</td>
</tr>
<tr>
<td>District nurse per minute of home visit</td>
<td>41</td>
<td>31</td>
<td>30.6%</td>
<td>24.4%</td>
<td>3</td>
</tr>
</tbody>
</table>

Note: For services marked with an asterisk (*), the cost is calculated based on the duration of the visit. For example, Meals on Wheels for 15 minutes is calculated at $31.04/4 for 15 minutes.
## The economic value of informal care in 2010

<table>
<thead>
<tr>
<th>Service Description</th>
<th>QTY</th>
<th>Price</th>
<th>Relativity to GP</th>
<th>Source of Data</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dentist per 20 minute visit</td>
<td>30</td>
<td>31</td>
<td>22.4%</td>
<td>MBS Item 85012 Benefit, relativity to GP</td>
<td></td>
</tr>
<tr>
<td>Optician per visit</td>
<td>35</td>
<td>35</td>
<td>26.1%</td>
<td>MBS Item 10900 Benefit, Schedule Fee</td>
<td></td>
</tr>
<tr>
<td>Chiropody per clinic visit</td>
<td>21</td>
<td>16</td>
<td>15.7%</td>
<td>MBS Item 10962 Benefit, Schedule Fee</td>
<td></td>
</tr>
<tr>
<td>Chiropody per home visit</td>
<td>7</td>
<td>8</td>
<td>5.2%</td>
<td>Home/clinic relativity from GP</td>
<td></td>
</tr>
<tr>
<td>Respite care per week</td>
<td>7</td>
<td>12</td>
<td>4.6%</td>
<td>-3,390 $31.04/hour*168 hours</td>
<td></td>
</tr>
<tr>
<td>Domestic assistance per 1 hour contact</td>
<td>1.3</td>
<td>1.4</td>
<td>100.0%</td>
<td>-3 As per domestic assistance in social services above</td>
<td></td>
</tr>
<tr>
<td>Australian minimum wage per hour</td>
<td>608.8</td>
<td>666.6</td>
<td>100.0%</td>
<td><a href="http://www.fairwork.gov.au/Fact-sheets-tools/Pages/FWO-fact-sheet-Minimum-wages.aspx">http://www.fairwork.gov.au/Fact-sheets-tools/Pages/FWO-fact-sheet-Minimum-wages.aspx</a></td>
<td></td>
</tr>
</tbody>
</table>

**Total**                                         |     |       |                   |                |       |

-870

Source: Patel et al (2004) and Access Economics. NHCDC = National Hospital Cost Data Collection (2009), for 2007-08, inflated to 2010 at 3% per annum. MBS Items as per May 2010. 
* Includes meal, transport, administration and capital, as per pers. comm. 27 July 2010 with the Australian Meals on Wheels Association.
Case study: Cost benefit analysis of a carer training program

There are few randomised clinical trials of informal carer training. One UK study randomised 300 stroke patients and their carers into two groups – one receiving training and the other receiving no training. The training comprised instructions in basic skills of moving and handling, facilitation of activities of daily living, and simple nursing tasks. The carers received the training over three to five sessions, lasting 30-45 minutes each, with a follow up session at home (Patel et al, 2004).

The UK study showed that total health and social care costs over one year for people whose carers received training were significantly lower than for those who did not receive training, saving £4,043 per person per annum on average (in 2000-01 prices) in this rehabilitative care setting.

We investigated what the impacts would be if Australian cost data were imputed to simulate a similar intervention here in 2010. Using data from the Medicare Benefits Schedule, National Hospital Cost Data Collection, this report and other sources, we estimated the costs saved from utilisation rates and unit cost data shown in Table 4.2 above.

The savings from the intervention were estimated as A$20,486 per person in 2010. Over two thirds (69%) of the savings were from fewer days in hospital, while 15% were savings from reduced use of other community health services such as GP home visits and respite care. A further 10% of the savings were from reduced use of physiotherapy and occupational therapy services. There was also a reduction in the informal care burden, which comprised 6% of the benefits (with the carer’s time valued at the minimum wage in this case, to match the UK method).

The cost of delivering the training ranged from A$342 (three 30-minute sessions and one home visit) to A$651 (five 45-minute sessions and one home visit) per trained carer in 2010 prices. Imputing a 100% loading for the cost of initial development of the training material, the benefit cost ratio ranged from 16:1 to 30:1.

The cost benefit analysis reveals that carer training can provide a substantial net benefit – over $19,000 per annum per person – through reducing reliance on formal health sector and community services relative to a situation of no carer training. A similar randomised controlled trial should be conducted in Australia to confirm these modelled findings, potentially across a variety of care settings.

4.3 Comparison of different care models

There is growing interest in the costs and benefits, from both a private and public perspective, of alternative means of providing care services. There is a continuum of care models currently provided in Australia ranging from reliance solely on an informal carer, ‘ageing in place’ with the assistance of an informal carer and some form of subsidised community care package (HACC or CACP), to higher levels of care provided either in the home (EACH, EACH-D) or in a
residential aged care (RAC) facility. RAC may be provided at either a low or high level of care intensity, depending on the needs of the resident.

To accurately determine the optimal balance and location of formal and informal care would require detailed modelling of the costs (both direct and other costs such as the opportunity cost of informal care) and benefits of different care models.\(^\text{15}\) Such an exercise is beyond the scope of this report. However, a simple comparison of the average cost per care recipient of various programs (Table 4.3) and the relative contributions of public and private funding sources, reveals a number of interesting relativities.

The comparison is based on calculations from Access Economics (2010c), which estimated the average cost to government of various care packages in 2008-09 – for HACC, CACP, EACH, EACH-D and RACS low care, high care and total –shown in the second column of Table 4.3 (indexed to 2010 at 3% per annum). To incorporate the costs paid by the individual, the public share was calculated based on updates to 2010 from Access Economics (2004). The average financial cost of informal care was calculated separately for primary carers on both an opportunity cost and replacement cost basis.

- The total opportunity cost valuation for informal carers includes their lost earnings together with efficiency costs from welfare payments and government funded direct support/respite services to carers.
- The total replacement cost valuation includes the replacement value of carers, without the efficiency costs from lost taxation or welfare payments since, in this case, the informal care services would be replaced with formal sector workers so the informal carers would not require payments or support services and would be able to participate in paid work at average levels.
- The welfare and taxation transfers themselves are not included in the real costs, but are taken into account in estimating the public sector shares of total costs (including tax payments of the replacement workers and tax losses from lost earnings using the opportunity cost approach).
- The public contribution towards the average cost of informal care is the sum of the welfare payments made to carers, the cost of carer support programs, and the lost taxation revenue. The cost of the carer’s time and the efficiency loss associated with welfare payments are assumed to be private costs (i.e. incurred by non-government entities in society).

<table>
<thead>
<tr>
<th>Care stream</th>
<th>Public cost $2010*</th>
<th>Public share</th>
<th>Total cost $2010</th>
</tr>
</thead>
<tbody>
<tr>
<td>HACC – Home and Community Care Program</td>
<td>1,992</td>
<td>98%</td>
<td>2,033</td>
</tr>
<tr>
<td>CACP – Community Aged Care Package</td>
<td>12,475</td>
<td>100%</td>
<td>12,475</td>
</tr>
<tr>
<td>EACH – Extended Aged Care in the Home</td>
<td>40,314</td>
<td>100%</td>
<td>40,314</td>
</tr>
<tr>
<td>EACH-D – Extended Aged Care in the Home (Dementia)</td>
<td>42,881</td>
<td>100%</td>
<td>42,881</td>
</tr>
<tr>
<td>Total – CACP, EACH and EACH-D</td>
<td>18,561</td>
<td>100%</td>
<td>18,561</td>
</tr>
<tr>
<td>RAC – low care</td>
<td>20,868</td>
<td>47%</td>
<td>44,319</td>
</tr>
</tbody>
</table>

\(^{15}\) For example, Canada recently completed a substantial study into the costs and benefits of home care compared to residential long term care of the elderly. See Hollander et al (2002).
The economic value of informal care in 2010

<table>
<thead>
<tr>
<th>Care stream</th>
<th>Public cost $2010*</th>
<th>Public share</th>
<th>Total cost $2010</th>
</tr>
</thead>
<tbody>
<tr>
<td>RAC – high care</td>
<td>59,227</td>
<td>86%</td>
<td>69,178</td>
</tr>
<tr>
<td>RAC – all</td>
<td>39,693</td>
<td>64%</td>
<td>62,219</td>
</tr>
<tr>
<td>Informal care, primary carers – opportunity cost</td>
<td>7,446</td>
<td>68%</td>
<td>10,950</td>
</tr>
<tr>
<td>Informal care, primary carers – replacement cost</td>
<td>33,076</td>
<td>78%</td>
<td>42,405</td>
</tr>
</tbody>
</table>


Table 4.3 and Chart 4.1 show that the average cost of informal care provided by primary carers is still relatively low ($10,950) when the carer’s time is valued at its opportunity cost. If the carer’s time is valued at replacement cost, then the average cost of a primary carer is much higher ($42,405), exceeding that of an EACH package ($40,314) or EACH-D package ($42,881) as would be expected, but still less than a high-care residential placement ($69,178).

![Chart 4.1: Average cost of care models per recipient, Australia, 2010](chart)

Source: Access Economics calculations.

Moreover, the average cost of combining informal primary care, valued at opportunity cost, with a HACC package or CACP is still lower than the cost of a low-care residential place (Chart 4.2).
Chart 4.2: Average cost of combined care packages per primary care recipient, 2010

Informal primary care with HACC services is lowest at $12,983; informal primary care with CACP costs around $23,425; EACH plus informal primary care is around $51,264; EACH-D plus informal primary care is around $53,831; RAC low care is around $44,319; while RAC high care is greatest at $69,178 per person in 2010.

It should be noted that these figures are average costs and so may mask important differences between people. In particular, a person in a high care RAC service may be receiving a greater intensity of care services than a person receiving some combination of informal care and a community care package, potentially due to higher disability and/or needs. If this is true, differences in cost would at least partially reflect the different level of services provided rather than the relative cost efficiency of providing the same level of service in a community or an institutionalised setting.

Moreover, it should be noted that residential options include the cost of the ‘housing’ component of care, whereas the at-home care cost does not. A final point is that, in residential care there may be economies of scale in service provision, particularly for ‘household tasks’ such as cooking, cleaning and maintaining facilities, that are not gained in individual community settings. The interaction of the welfare, taxation and residential aged care systems is complex and may possibly provide incentives for the frail aged (or their families) to maintain private care in the home beyond the point where there are net financial benefits in doing so, while recognising that other benefits are less tangible (e.g. the personal utility and quality of life derived from being at home, see Section 5.1).
5 Conclusions and implications for the future

The previous sections of this report have attempted to identify and place a financial value on current levels of informal care. It is clear that informal carers provide a considerable proportion of Australians’ care needs, in conjunction with formal care providers. However, estimates of the value of informal care may raise a number of related policy questions.

- Are current models of care the most appropriate method of care provision, both now and in the future?
  - What will happen to the demand and supply of informal care in future?
  - How cost effective are different models of care provision, when all the costs and benefits, including the cost of carer time are included?
  - Would it be more efficient for some informal carers to return to the paid workforce, with care needs met by the formal sector instead?

- Are carers appropriately remunerated for the value they provide?
  - How is the burden of care distributed between the carer, their family and the wider community?
  - What are the most effective means of supporting carers?

Policy issues such as these require consideration of both efficiency and equity criteria. Efficiency considers whether economic resources, including labour, are used in a manner which maximises the welfare of Australians (‘the size of the cake’). Equity, on the other hand, considers how costs and benefits of resource allocation are distributed among the community (‘the slices of the cake’).

The equity-efficiency trade off can be complex. This section of the report does not present a comprehensive policy prescription, but notes some of the questions and evidence that will need to be considered by Government.

5.1 Informal care demand and supply

A number of studies have pointed to the potential impact of long term demographic and social trends on the demand and supply of informal care in Australia. These trends include (Jenkins et al, 2003):

- demographic ageing of the Australian population;
- higher prevalence of chronic illness and associated disability among older age groups;
- deinstitutionalisation and greater emphasis on ‘ageing in place’;
- increased female labour force participation;
- increased rates of relationship breakdown;
- smaller families and increased childlessness (reduced fertility);
- older parent carers of adult children with disabilities;
- greater mobility and dispersion of families;
- increased number of single person households generally, and among elderly people in particular; and
- inter-generational propensities to care.

Exactly how these and other factors will impact overall on the level of informal care required and provided in the future is unclear.

Extrapolating current data to forecast trends in the need and provision of informal care over the coming decades is difficult. Reliable forecasts would need considerable information about the expected prevalence of medical conditions which require care, as well as demographic projections and assumptions about the disposition of people to care for others (e.g. will Generation X and Y be as willing to care for their parents as the baby boomers are?)

The AIHW (Jenkins et al, 2003) and the National Centre for Social and Economic Modelling at the University of Canberra (NATSEM, 2004) both produced projections of future demand and supply of informal care by extrapolating out from 1998 SDAC data and adjusting for possible social trends. A simplified version of this model is used to project the 2003 SDAC.

Chart 5.1 shows the increased need for care due to demographic ageing, simply measured as the ‘caretaker ratio’ - the number of women aged between 50 to 64 to people aged 80 years and over (a crude indicator of the number of daughters available to provide care to ageing parent(s)).

**Chart 5.1: Ratio of females 50 to 64 years of age to persons aged 80 years and over**

Notes: A, B & C represent the ABS high, medium and low population projections, respectively. Source: ABS (2008b).
The economic value of informal care in 2010

The caretaker ratio is currently quite favourable for Australia at around 2.5 ‘daughters per parent’. Under current population projections (high, medium and low) the ratio will start to decline substantially between 2010 and 2025, continuing to decline to 2050.

Demand for care was also explicitly modelled, based on the number of people with a profound or severe core-activity limitation over 65 living in the community. Age and gender-specific disability rates from the 2003 SDAC were applied to population projections to estimate the number of people demanding some form of care in future years.

The supply of formal care through residential placement was assumed to remain available for a constant proportion of all people with a disability. For other disabled people, informal care is the only alternative source of supply. The supply of informal care was based on the propensity to care of each different age and sex.

Chart 5.2: Projected demand and supply of informal carers

Note: The intersection of the curves in 2021 is a product of the scales used and has no particular implications for that year.
Source: Access Economics calculations.

As Chart 5.2 shows, the ‘carer ratio’ of primary carers to older people with a disability is projected to fall, from 60% now to under 40% by mid-century, reflecting that demand for informal care will substantially outstrip supply. Increasing demand for care due to the increased number of people aged 65 and over is the main factor behind this fall. The number of primary carers will also grow, but at a much slower rate. These projections also raise interesting, but as yet unanswered questions as to whether care from a number of non-primary carers can be a substitute for a single primary carer (Carers Australia, 2005).
Scenario analysis was conducted to show how the projected supply of carers would differ if social change reduced or increased the propensity of particular groups in society to provide care. Specifically, the three scenarios considered were as follows.

- An overall decline in the propensity of people to care, represented by a 20% across-the-board decrease in carer rates.
- A decline in the propensity of women to reduce paid employment in order to provide care, represented by a 20% decrease in the proportion of women reducing paid employment to care.
- An increase in the availability of carers due to converging male and female life expectancy, represented by a 20% increase in carer rates in 65+ age group.

The first scenario (grey line in Chart 5.3), a decline in across the board carer rates, has the largest impact on the deficit of informal carers. By construction, under this scenario there would be 20% less informal carers by 2050. There are many reasons why such a scenario could occur. Higher divorce rates, smaller family size and a decreasing propensity of generation X and Y to care for the elderly are all potential contributors (Access Economics, 2010b).

The second scenario (the dashed black line in Chart 5.3) results in a more moderate decline in the shortage of informal care than scenario 1. Working age females are an important source of informal care, so increasing female workforce participation reduces the hours available to provide informal care,. In addition, ageing of the population will result in a tighter labour market, raising wages and thereby increasing the opportunity cost of reducing paid employment to provide unpaid informal care. Carmichael et al (2010) found, using the British Household Panel Survey, that people who are already in paid employment when the decision to provide care arises are less likely to do so than those who are not. The probability of choosing not to provide unpaid care increases as wages increase. However, the use of part-time and other more flexible work arrangements has also increased, potentially providing greater opportunity to combine work and care roles (Howe and Sergeant, 1999). Female workforce participation and preference is likely to continue to change as family circumstances change (Glezer and Wolcott, 1997).

Scenario 3 shows the effect of an increasing male life expectancy, resulting in a small increase in the availability of care by those over 65. This scenario is based on the premise that spouses are an important source of informal care (although increasing rates of divorce may partially or fully offset such an effect). A higher male life expectancy could increase the supply of informal care, assuming this increase in longevity was associated with an increase in the number of years without or with minimal disablement.

In the base case there is already a deficit in 2010, which is projected to increase under all three scenarios by 2050. This deficit may be a result of several different, or a combination, of factors. First, there may already be unmet need for informal carers in the community. Second, the analysis only includes primary carers and an unknown proportion of people may be receiving care from one or more non-primary carers rather than a primary carer. Finally, it could be a result of disconnect between data sources as a result of reporting errors. Regardless of the accuracy of the initial deficit, the change in the deficit under the different scenarios is informative. The findings are summarised as follows.

- An overall decline in the propensity of people to care, represented by a 20% across-the-board decrease in carer rates, increased the base case deficit by 12.1% in 2050.
A decline in the propensity of women to reduce paid employment in order to provide care, represented by a 20% decrease in the proportion of women reducing paid employment to care, increased the base case deficit by 5.5% in 2050.

An increase in the availability of carers due to converging male and female life expectancy, represented by a 20% increase in carer rates in 65+ age group, reduced the base case deficit by 3.1% in 2050.

Chart 5.3: Scenario analysis of projected demand and supply of informal carers

Notes: Baseline refers to Chart 5.2. Scenario 1 corresponds to a 20% decrease across-the-board in carer rates. Scenario 2 is a 20% decrease in female carers aged 25-64. Scenario 3 is a 20% increase in carer rates in the 65+ age group.
Source: Access Economics calculations.

These simple scenarios illustrate the impact of several plausible changes in supply of informal care. However, there are many other factors that will also affect supply and demand, which have not been controlled for. In particular, the supply of formal care available will substantially impact on the demand for informal carers. McCallum (2003) cites evidence suggesting that 88% of Australians aged over 70 years would prefer to receive formal care. To the extent that informal and formal care are economic substitutes, a shortage of formal care services will increase demand for informal care.

16 The relationship between informal and formal care services is complex, and not yet fully understood. In some instances formal and informal care operate as substitutes; a person without access to informal care may instead seek assistance through formal care providers. In other scenarios the two types of care may be complementary, with specialised formal care (such as particular medical interventions) supplementing the provision of other personal care needs through informal carers (Jenkins et al, 2003).
The changing epidemiology of disease will also affect the demand for informal care. Increasing longevity is associated with higher prevalence of chronic disease, more prolonged duration of illness, and more complex comorbidities to manage. In particular, dementia prevalence will have a profound impact on the demand for care services (Access Economics, 2010b). Providing informal care to a person with dementia is time intensive and can result in negative health and well-being outcomes for the carer, in the absence of appropriate support services. Progression of dementia, including its behavioural and psychological symptoms, can precipitate greater need for formal care services.

5.2 Challenges and opportunities

Increasing longevity is associated with higher prevalence of chronic disease, more prolonged duration of illness, and more complex comorbidities to manage. Overall, the percentage of people with a profound or severe core-activity limitation living in the community is projected to increase. Access Economics (2009b) highlighted the challenges that the aged care system will face as the number of people with dementia increases. Neurodegenerative diseases such as dementia can precipitate the need for care services or elevate the level of care required.

In the context of the growing disability burden, movements towards community care and other models of integrated care have historically been both financially motivated as well as reflecting community attitudes and preferences. Most older Australians would prefer that, should they require care, they are able to receive it while remaining in their own homes (Rowland, 1991; McCallum and Mundy, 2002). However, this is usually only possible where a family member or friend is available to provide informal care. For example, there is evidence that elderly people who live alone are more likely to be recommended for admission to residential care than those living with a spouse or other people. Of people recommended for admission into residential care, people living alone are more likely to be assessed as needing low-level as opposed to high-level care. For people living with family members, the opposite is true. It is possible that living with a spouse or other family member may also allow people to remain living in the community until their level of disability requires high-level care (Lincoln Gerontology Centre, 2002).

However, the preferences of the carer as well as the care recipient must be considered. SDAC surveyed carers about their level of satisfaction with their role, and found that only 25.7% of primary carers felt satisfied. A significant number of carers also expressed the desire to return to paid work. A survey of Victorian carers found that younger carers were more likely to want to maintain or increase their workforce participation compared to carers aged over 50 (Schofield et al, 1998).

In terms of costs, older people and their carers meet a large proportion of the costs of care which would otherwise be borne by government (Productivity Commission, 2003b). Appropriate community-based interventions can also be more cost effective than institutionalised care; the analysis conducted in this report based on average costs of care appears to show that combinations of informal care and community based formal care are generally lower cost than institutionalised care (including a housing component), even when the opportunity cost of the carer’s time is included. However, the government subsidy for informal care is generally lower than for other community care services.

Overall, the future mix of care – between community and RAC, and between formal and informal services – is unclear – reflecting the changing pattern of disease (favouring RAC and
formal care services), shifting social preferences (generally towards community care) and supply constraints (notably potentially fewer informal carers). The third InterGenerational Report (Treasury, 2010) concludes that the proportion of services allocated to high care will need to increase over time and adapt to the special needs of people with dementia and other severe chronic disease. This is expected to result in a much larger relative increase in spending on RAC than on community care (Chart 5.4).

Chart 5.4: Projected Australian Government aged care spending

![](chart.png)

Source: Access Economics based on Treasury (2010).

It is not clear from the InterGenerational Report how increases in care costs in the future – from 0.8% of GDP today to 1.8% of GDP in 2049-50 – are to be funded. Rather, the projections show growing fiscal deficits resulting from aged care and health expenditures. Silence on funding deficits is accompanied by silence in the InterGenerational Report on other key issues such as how to meet emerging workforce shortages and sensitivity analysis of the projections if the propensity to provide informal care falls.

The analysis in this report emphasises that carers have their own needs, distinct from those of the care recipient. Clearly there is a need to consider these needs in policy planning for the coming decades. The SDAC showed that 37.2% of primary carers felt they needed more support in their caring activities. However, needs can vary considerably according to the type of disability of the person cared for. Carers of people with an intellectual (53%) or psychological (44%) disability were most likely to need additional support. Least likely to need more support were carers of people with a sensory or speech disability (24%) or a physical disability (35%).

Priority support for carers comprise:

- respite care;
- income support;
- education and training; and
- preventive health care.

Previous modelling by Access Economics found that increased respite support for family carers of people with dementia could generate a benefit cost ratio of around 7:1 if the respite care reduced the number of people moved into institutional care (Access Economics, 2003). American studies have found that respite care and other interventions aimed at reducing psychological distress in dementia caregivers can delay institutionalisation (Spector and Tampi, 2005). A lot has been accomplished in providing respite care over the past decade, although there remains a need to ensure respite care is uniformly accessible, flexible and appropriate across Australia in the coming years as demand burgeons.

This report has shown that expenditure on carer income support, counselling and training can also produce a net benefit. Almost all of the studies considered by Toseland and Smith (2004) found caregiver education and training programs have small or moderate effects on increasing knowledge about services and resources for carers, improving psychological and social wellbeing of carers and increasing problem solving abilities and reducing pressing problems related to caring. Section 4.2 highlighted the potential returns to carer education and training, together with the relative underinvestment in such programs for informal carers, compared to formal carers.

Training, information and support can assist carers’ capacity to fulfil their caring role in ways which also protect their own health (e.g. lifting techniques). Policy consideration should also be given to preventive health checks for informal carers and a personalised health record to help maintain carers’ health.

Such support for carers – through respite, income support, education and training and preventive health care – will help ensure and enhance future labour productivity across the formal and informal sectors of the economy, as well as meeting Australia’s obligations to optimise wellbeing for the frail aged those disadvantaged by disability.
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