# Cancers in Australia in 2010 attributable to modifiable factors: summary and conclusions 

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n Australia in 2010, 116,850 people were diagnosed with invasive cancers, not including keratinocyte cancers (basal cell carcinomas (BCC) or squamous cell carcinomas (SCC) of the skin). ${ }^{1}$ Epidemiologic and experimental evidence accrued over the past five decades has established that many factors to which humans are commonly exposed are causes of cancer. Because exposure to such factors is potentially modifiable, it is important to quantify the proportion of cancers that might be averted if exposure to those factors were removed entirely or reduced to the minimum compatible with optimum health.
We estimated the fraction and numbers of cancers occurring in the Australian population that were attributable to established causal factors that are, at least in theory, able to be modified. ${ }^{2}$ Using estimates of the prevalence of each exposure in the Australian population and estimates of the relative risk of cancer associated with different levels of exposure, we applied standard formulae to calculate the population attributable fraction (PAF) for cancers associated with exposure to each of 13 factors determined to cause cancer by independent, international cancer agencies (Table 1). The magnitudes of PAF estimates depend heavily on the definition of exposure; Table 1 lists the reference category for each factor that was used to estimate the PAF. For some factors (e.g. tobacco smoking, alcohol, infections), the reference category represents


#### Abstract

Objective: To estimate the numbers and proportions of cancers occurring in Australia in 2010 attributable to modifiable causal factors. Methods: We estimated the population attributable fraction (PAF) of cancers associated with exposure to 13 causal factors using standard formulae incorporating exposure prevalence and relative risk data. We also calculated the potential impact of changing exposure to some factors. Results: A total of $32 \%$ of all cancers diagnosed in Australia in 2010 (excluding keratinocyte cancers) were attributable to the 13 factors assessed (men 33\%; women $31 \%$ ). Leading factors were tobacco smoke (PAF all cancers: 13.4\%), solar radiation (6.2\%), inadequate diet (6.1\%) and overweight/obesity (3.4\%). Factors conferring highest PAFs differed by sex: highest PAFs for men were tobacco smoke (15.8\%), solar radiation (7.1\%) and alcohol (3.0\%); while highest PAFs for women were tobacco smoke (10.1\%), solar radiation (5.0\%) and overweight/obesity (4.5\%). Sites with the highest counts of potentially preventable cancers were lung $(8,569)$, colorectal $(7,404)$, melanoma of the skin $(7,220)$ and breast $(3,233)$.

Conclusions: At least one in three cancers in Australia is attributable to exposure to known modifiable factors. Implications: Up to 37,000 cancers could be prevented in Australia each year if the population avoided exposure to 13 common factors known or strongly suspected to cause cancer.


Key words: population attributable fraction, cancer, risk factor, potential impact fraction, prevented fraction
no exposure. Other factors are ubiquitous (e.g. body mass index, dietary factors, sunlight) and so the reference levels for those factors were widely accepted thresholds for optimum health.

## Proportions of cancers attributable to modifiable exposures

Table 2 summarises the PAF estimates for 26 cancer sites, and for all cancers collectively (excluding BCC and SCC of the skin), for
each of the 13 factors appraised. Detailed reports including numbers of cases of cancer attributed to these factors and sensitivity analyses have been reported in companion articles. ${ }^{3-14}$ In total, we found that $32 \%$ of all cancers (approximately 37,000 cases) diagnosed in Australia in 2010 (excluding BCC and SCC of the skin) could be attributed to the 13 factors ( $33 \%$ in men and $31 \%$ in women). Overall, tobacco smoke (PAF all cancers: $13.4 \%$ ) was the leading

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Table 2: Proportion (\%) of incident cancer cases diagnosed in 2010 in Australia attributable to lifestyle and environmental factors.

| Cancer site |  | All Oesophagus (C15) ${ }^{\text {a }}$ | Stomach <br> (C16) ${ }^{\text {a }}$ | Colorectum$\left(C 18-(20)^{\mathrm{a}}\right.$ | Anus <br> (C21) ${ }^{\text {a }}$ | $\begin{aligned} & \text { Liver } \\ & \text { (C22) } \end{aligned}$ | Gall Bladder (C23) ${ }^{\text {a }}$ | Pancreas <br> (C25) ${ }^{\text {a }}$ | $\begin{aligned} & \text { Larynx } \\ & \text { (C32)a }^{a} \end{aligned}$ | $\begin{aligned} & \text { Lung } \\ & (C 34)^{a} \end{aligned}$ | \% of cancer (0-85+ years) |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Oral cavity and pharynx (COO-CO6; C09-(14) ${ }^{\text {a }}$ |  |  |  |  |  |  |  |  |  | Melanoma $(C 43)^{a}$ | Kaposi's <br> Sarcoma <br> (C46) ${ }^{\text {a }}$ |
| Males |  |  |  |  |  |  |  |  |  |  |  |  |
| Tobacco smoke | 71.0 | 71.4 | 23.6 | 5.9 | - | 24.3 | - | 23.7 | 76.9 | 84.5 | - | - |
| Alcohol | 24.7 | 9.9 | - | 12.9 | - | 15.4 | - | - | 22.1 | - | - | - |
| UV radiation | - | - | - | - | - | - | - | - | - | - | 69.7 | - |
| Infections | 16.9 | - | 32.8 | - | 74.5 | 34.4 | - | - | - | - | - | 100.0 |
| Overweight and obesity | - | 18.0 | - | 11.7 | - | - | 15.8 | 9.1 | - | - | - | - |
| Insufficient physical activity | - | - | - | 4.2 | - | - | - | - | - | - | - | - |
| Diet - insufficient fibre | - | - | - | 18.6 | - | - | - | - | - | - | - | - |
| Diet - insufficient vegetables | 6.0 | 2.6 | 1.5 | - | - | - | - | - | 5.9 | - | - | - |
| Diet - insufficient fruit | 6.0 | 12.0 | 6.5 | - | - | - | - | - | 5.9 | 6.9 | - | - |
| Diet - red and processed meat | - | - | - | 21.7 | - | - | - | - | - | - | - | - |
| PAF combined ${ }^{\text {c }}$ | 83.9 | 74.2 | 52.7 | 55.8 | 74.5 | 58.0 | 15.8 | 30.6 | 84.1 | 86.0 | 69.7 | 100.0 |
| Females |  |  |  |  |  |  |  |  |  |  |  |  |
| Tobacco smoke | 51.2 | 61.6 | 10.7 | 7.0 | - | 10.8 | - | 23.0 | 72.6 | 77.0 | - | - |
| Alcohol | 9.2 | 6.2 | - | 4.2 | - | 4.5 | - | - | 7.1 | - | - | - |
| UV radiation | - | - | - | - | - | - | - | - | - | - | 54.2 | - |
| Infections | 14.4 | - | 38.4 | - | 90.5 | 34.4 | - | - | - | - | - | 100.0 |
| Overweight and obesity | - | 7.7 | - | 5.5 | - | - | 13.3 | 6.1 | - | - | - | - |
| Insufficient physical activity | - | - | - | 5.4 | - | - | - | - | - | - | - | - |
| Diet - insufficient fibre | - | - | - | 16.2 | - | - | - | - | - | - | - | - |
| Diet - insufficient vegetables | 7.1 | 6.0 | 1.8 | - | - | - | - | - | 7.1 | - | - | - |
| Diet - insufficient fruit | 5.8 | 23.5 | 6.4 | - | - | - | - | - | 6.0 | 9.6 | - | - |
| Diet - red and processed meat | - | - | - | 12.5 | - | - | - | - | - | - | - | - |
| Hormones - MHT | - | - | - | - | - | - | - | - | - | - | - | - |
| Hormones - combined OCPs | - | - | - | - | - | - | - | - | - | - | - | - |
| Insufficient breast feeding | - | - | - | - | - | - | - | - | - | - | - | - |
| PAF combined ${ }^{\text {c }}$ | 66.8 | 76.1 | 49.4 | 41.6 | 90.5 | 44.1 | 13.3 | 27.8 | 77.8 | 79.3 | 54.2 | 100.0 |
| Persons |  |  |  |  |  |  |  |  |  |  |  |  |
| Tobacco smoke | 65.3 | 60.0 | 19.2 | 6.4 | - | 20.7 | - | 23.4 | 76.4 | 81.6 | - | - |
| Alcohol | 20.3 | 8.8 | - | 9.0 | - | 12.5 | - | - | 20.1 | - | - | - |
| UV radiation | - | - | - | - | - | - | - | - | - | - | 63.3 | - |
| Infections | 16.2 | - | 34.7 | - | 84.0 | 34.4 | - | - | - | - | - | 100.0 |
| Overweight and obesity | - | 15.0 | - | 9.0 | - | - | 14.1 | 7.7 | - | - | - | - |
| Insufficient physical activity | - | - | - | 4.8 | - | - | - | - | - | - | - | - |
| Diet - insufficient fibre | - | - | - | 17.6 | - | - | - | - | - | - | - | - |
| Diet - insufficient vegetables | 6.3 | 3.6 | 1.6 | - | - | - | - | - | 6.1 | - | - | - |
| Diet - insufficient fruit | 6.0 | 15.4 | 6.5 | - | - | - | - | - | 5.9 | 9.6 | - | - |
| Diet - red and processed meat | - | - | - | 17.6 | - | - | - | - | - | - | - | - |
| Hormones - MHT | - | - | - | - | - | - | - | - | - | - | - | - |
| Hormones - combined OCPs | - | - | - | - | - | - | - | - | - | - | - | - |
| Insufficient breast feeding | - | - | - | - | - | - | - | - | - | - | - | - |
| PAF combined ${ }^{\text {c }}$ | 79.6 | 74.7 | 51.4 | 49.8 | 84.0 | 54.4 | 14.1 | 29.3 | 83.3 | 83.3 | 63.3 | 100.0 |

Abbreviations: $P A F=$ Population attributable fraction; $M H T=$ menopausal hormone therapy; $O C P_{S}=$ oral contraceptive pills
a: International Classification of Diseases (ICD-10) code
b: Excluding cases of basal cell carcinoma and squamous cell carcinoma of the skin;
c: PAF Combined $=1-\left(1-P A F^{1}\right)^{*}\left(1-P A F^{2}\right)^{*}\left(1-P A F^{1}\right)$
Note: The proportions (\%) presented here are for the entire Australian population (0-85+ years) to ensure comparability across all exposures. The PAFs in the table therefore may differ from the PAFs presented in the exposure-specific papers in this series.

cause of cancer in 2010, followed by solar radiation (6.2\%), inadequate diet (6.1\%) and overweight and obesity (3.4\%), see Figure 1. The factors conferring the highest PAFs differed by sex: for men the highest proportions of cancer were attributable to tobacco smoke ${ }^{12}(15.8 \%)$, solar radiation ${ }^{10}$
(7.1\%) and alcohol consumption ${ }^{13}(3.0 \%)$; while for women tobacco smoke ${ }^{12}(10.1 \%)$, solar radiation ${ }^{10}(5.0 \%)$ and overweight and obesity $^{7}(4.5 \%)$ were the leading causes.
The cancer sites with the greatest fractions of potentially preventable cases were uterine cervix (100\%) and Kaposi's sarcoma


(100\%), for which infectious agents (human papillomarvirus [HPV] and Kaposi's sarcoma herpes virus [KSHV], respectively) are considered necessary causal factors (Figure 2). Other sites with very high fractions of potentially preventable cancers were oral cavity and pharynx (80\%), oesophagus (75\%), anus (84\%), larynx (83\%) and lung (83\%). In absolute terms, the sites with the greatest number of potentially preventable cancers were lung $(8,569)$, colorectal $(7,404)$, melanoma of the skin $(7,220)$ and breast $(3,233)$.

The incidence and attributable fraction of the 10 most common cancers in Australian men and women are displayed in Figure 3. Prostate cancer is the most commonly occurring cancer in Australian men and has no accepted causes. The PAF estimate for all cancers is clearly affected by the lack of modifiable factors for this very common cancer. Of the subset of cancers with known, modifiable causes, we estimated that $43 \%$ of those arising in the Australian population in 2010 could be attributed to exposure to the 13 causal factors listed. For some exposures, PAFs were much higher for men than women. This was most noticeable for cancers attributable to alcohol (e.g. oral cavity and pharynx: men $25 \%$ vs. women $9 \%$; colorectum: men $13 \%$ vs. women 4\%; liver: men $15 \%$ vs. women $5 \%$ ), red and processed meat intake (colorectal cancer: men $22 \%$ vs. women $13 \%$ ), and tobacco (oral cavity and pharynx: men 71\% vs. women 51\%; stomach: men $24 \%$ vs women $11 \%)$. As a result, a markedly higher proportion of those cancers were attributable to exposure to modifiable factors in men (52\%) than women (35\%), see Table 2. Knowledge about the causes of cancer continues to evolve. There is strengthening evidence that a number of the factors considered here also cause cancer at other sites, including overweight/ obesity, ${ }^{7,15}$ (cancers of the thyroid, nonHodgkin's lymphoma, multiple myeloma, melanoma and leukaemia), menopausal hormone therapy ${ }^{4}$ (cancers of the breast -oestrogen-only formulations - and ovary) and insufficient breast-feeding ${ }^{5}$ (ovarian cancer). Because no independent cancer agencies have declared these associations to be causal, we did not include them in our primary analyses. However, we did conduct supplementary analyses to investigate the potential magnitude of the additional cancer burden that might have accrued due to these exposures. In fact, including these 'supplementary' cancers made negligible
difference to the total PAFs for men (from $32 \%$ to $33 \%$ ) and women (from 31\% to 32\%).

The summaries reported above do not include estimates of the fraction of cancer attributable to occupational exposures. This is because there were no reliable data on past exposures to occupational carcinogens in Australia, although recent occupational exposures have been recorded with high quality. ${ }^{16}$ We considered using a 'lifetime risk' approach for occupational exposures, which estimates the proportion of future cancers that can be attributed to exposures received in a specific year. While similar in concept to the attributable risk approach used for all other exposures, the outputs from the two approaches cannot be combined. In future work, we plan to explore the burden of cancer in Australia attributable to occupation exposures.

## Prevented fractions

Four factors (daily sunscreen use, ${ }^{10}$ daily aspirin use, ${ }^{14}$ menopausal hormone therapy $[\mathrm{MHT}]^{4}$ and combined oral contraceptives ${ }^{6}$ ) were deemed likely to prevent certain cancers; for these factors, we estimated the prevented fraction (PF) of cancer associated with exposure (Table 3). The PF estimates the proportion of the hypothetical 'total load' of cancer that was prevented due to prevailing levels of exposure to the factor in the population. ${ }^{17}$ About $28 \%$ of Australians reported using sunscreen daily in 2010, and we estimated that this level of use reduced the population incidence of SCC of the skin by about $9 \%$ (and possibly melanoma of the skin by about 14\%). ${ }^{10}$ Similarly, the incidence of cancers of the colorectum and oesophagus were reduced by about $2 \%$ and $3 \%$, respectively, due to prevailing levels of
aspirin use in the population. ${ }^{14}$ Estimating the impact of exogenous hormones on cancer incidence presents some challenges, as certain formulations cause some cancers and prevent others. On balance, MHT caused more cancers than it prevented, whereas the oral contraceptive pill prevented more cancers than it caused. We estimated that exposure to the combined oral contraceptive pill prevented 1,032 endometrial (PF 31\%) and 308 ovarian (PF 19\%) cancers, but likely caused 105 breast (PAF $0.7 \%$ ) and 52 cervical (PAF 6.4\%) cancers. ${ }^{6}$ Similarly, oestrogenonly MHT was estimated to prevent 52 colorectal cancers, but post-menopausal MHT (combined and oestrogen-only) caused up to 670 cancers, mostly breast cancers. ${ }^{4}$

## Potential impact of changing the prevalence of exposures

The PAF for any causal factor represents the theoretical maximum proportion by which the incidence of cancer could have been reduced, assuming that everybody in the population had been exposed at the reference level. For many exposures, however, it is not feasible to expect population-wide changes to an optimum exposure level. We therefore measured the potential impact on cancer incidence in Australia in 2010 of changing the prevalence of some common factors to those levels recommended by Australian guidelines (for alcohol, ${ }^{18}$ red \& processed meat ${ }^{19}$ and physical activity ${ }^{20}$ ) or to historic levels (for overweight/obesity)


| Exposure | Colorectal$(\mathrm{C} 18-\mathrm{C} 20)^{\mathrm{a}}$ |  | Oesophagus (C15) ${ }^{\text {a }}$ |  | SCC of the Skin ${ }^{\text {b }}$ |  | Endometrium$(C 54, C 55)^{a}$ |  | Ovarian (C56) ${ }^{\text {a }}$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | PF\% | Cancers <br> Prevented | PF\% | Cancers <br> Prevented | PF\% | Cancers <br> Prevented | PF\% | Cancers <br> Prevented | PF\% | Cancers <br> Prevented |
| Males |  |  |  |  |  |  |  |  |  |  |
| Aspirin | 2.3 | 198 | 3.3 | 34 | - | - | - | - | - | - |
| Sunscreen | - | - | - | - | 7.5 | 6,793 | - | - | - | - |
| Females |  |  |  |  |  |  |  |  |  |  |
| Aspirin | 2.0 | 136 | 3.9 | 17 | - | - | - | - | - | - |
| Sunscreen | - | - | - | - | 12.1 | 7,399 | - | - | - | - |
| Hormones - MHT ${ }^{\text {c }}$ | 0.8 | 52 | - | - | - | - | - | - | - | - |
| Hormones - combined 0 CPd | - | - | - | - | - | - | 31.4 | 1,032 | 19.1 | 308 |
| a: International Classification of Diseases (IID-10) Code <br> b: SCC of the skin = Squamous cell carcinoma of the skin <br> c: MHT = menopausal hormone therapy <br> d: $O C P=$ combined oral contraceptive pill |  |  |  |  |  |  |  |  |  |  |

or otherwise achievable targets (for dietary fibre, ${ }^{8}$ breastfeeding ${ }^{5}$ and $\mathrm{MHT}^{4}$ ), see Table 4. Again, we used standard formulae to calculate the potential impact fraction (PIF) using either of two similar approaches, depending upon whether we modelled a shift in the prevalence distributions ${ }^{21}$ or a change in the risks of exposure. ${ }^{22}$ Briefly, for each cancer site, we calculated the number of cases that would have occurred in Australia in 2010 if the alternative scenario of exposure had prevailed. The PIF is then the proportional difference between the numbers of cancers actually observed and the numbers expected under the alternative
(counterfactual) exposure scenarios. Our analyses suggest that substantial gains in cancer prevention could be achieved through relatively modest reductions in alcohol ${ }^{13}$ and meat intake, ${ }^{9}$ and an increase in fibre intake simply by eating fruit ( 2 servings/day) and vegetables ( 5 servings/day). ${ }^{8}$

## Comparisons with international studies

The approach used in this Australian study was virtually identical to that taken for the recent UK PAF Project, ${ }^{23}$ and was greatly enabled by the UK investigator (DMP) making

Figure 3: The ten cancer sites with highest incidence in Australia in 2010: total number of cases and fraction attributable to modifiable factors for A. Males and B. Females.
A. Males


## B. Females


his background materials, including worked spreadsheets, freely available. Overall, the proportions of cancers attributable to modifiable exposures were remarkably consistent, with tobacco contributing by far the highest cancer burden in both countries (Table 5). Nonetheless, there were some differences between the Australian and UK studies in the magnitude of specific PAF estimates. In part, this was due to the use of updated (and typically lower) risk estimates for dietary factors, based on new metaanalyses of prospective cohort studies that were not available at the time of the UK study. However, given the overall uniformity of approaches, most of the differences between the UK and Australian PAF estimates largely reflect differences in the underlying prevalence of exposures between the two populations. Thus, the proportions of cancers attributable to smoking and alcohol were notably higher in the UK than Australia, due to higher exposure prevalences of both factors in the UK population in recent decades. Not surprisingly, the proportion of all cancers (except BCC and SCC) in Australia attributable to sunlight was almost twice that of the UK. While some caution is warranted in comparing the PAFs for sunlight, since different reference populations were required for the calculations in each country, the relative proportions of cancer attributable to sunlight in the two populations appear consistent with the relative amounts of solar radiation to which they are exposed.

Estimates of the burden of cancer attributable to modifiable factors have also been conducted for other populations recently, including France ${ }^{24}$ and the US. ${ }^{25}$ Those investigations reported generally higher fractions attributable to each factor than we have reported here. Again, some of this is explained by different exposure prevalences, but the methods used in those studies differed substantially from the recent UK and Australian studies. In particular, those studies tended to stratify the population into a limited number of exposure categories, an approach that does not account for the gradations in risk associated with continuous distributions of exposure. As such, they tend to inflate the estimates of attributable fractions and, for this reason, are not directly comparable to the analyses reported by the UK and Australian studies.

## Limitations

A number of issues have a bearing on the interpretation of the estimates reported
here and in companion papers. First, in terms of the overall proportion of cancers attributable to modifiable factors, we note that our investigation was limited to only 13 factors for which there is international consensus regarding causality for cancer. Other factors known to cause cancer were not included in our final analyses, either because they cannot be modified in any practical sense (such as early menarche or nulliparity) or because prevalence data in the Australian population are so poorly collated as to be unusable (such as exposure to ionising radiation from medical sources). The latter is especially disturbing; since one might expect that a register of exposures to a known carcinogen would be mandated and audited for safety purposes. ${ }^{26}$

Second, as mentioned above, the estimates of attributable fractions are sensitive to the magnitude of effect sizes and prevalence. For our primary analyses, we used the most recent summary risk estimates from independent agencies when available; if not, we used estimates from recent, highquality meta-analyses, preferably restricted to cohort studies. Prevalence data were sourced preferentially from national surveys, or else from population-based studies with large samples when national survey data were not available. A salutary finding from this exercise was the overall low quality of information regarding exposure to Group 1 carcinogens in the Australian population. For pharmacological factors in particular, nationally representative data describing past exposures across age and sex were not available, and so we constructed matrices of exposure using multiple data sources. Detailed data on population exposures to ionising radiation from medical sources are not collected, rendering analyses for these exposures incomplete. Similarly, data on exposures to infectious agents and dietary factors were incomplete or out of date. Given these constraints, we conducted sensitivity analyses for all factors incorporating a range of estimates of risk or exposure prevalence to assess the distribution within which a plausible estimate of attributable fraction might lie. We did not model interactions between causes owing to a scarcity of summary risk estimates and absence of suitably stratified prevalence data. As such, it is likely that estimates of the fraction of cancer attributable to certain factors will include a proportion that was caused only because of exposure to a co-factor.

| Table 4: The potential impact on cancer incidence of changing exposure to modifiable causal factors. |  |  |
| :--- | :--- | :--- |
| Modifiable factor | Alternative scenarios in population exposures | PIF (\%) |
| Alcohol | 1. All who consume $>4$ drinks/day reduce intake to $\leq 4$ drinks/day | $2.2 \%$ |
|  | 2. All who consume $>4$ drinks/day reduce intake to $\leq 2$ drinks/day |  |$\quad 4.3 \%$


| Exposure | Aust PAF males (\%) | Aust PAF females | Aust PAF persons | UK PAF males (\%) | UK PAF females <br> (\%) | UK PAF persons (\%) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Tobacco | 15.8 | 10.1 | 13.4 | 23.0 | 15.6 | 19.4 |
| Alcohol | 3.0 | 2.4 | 2.8 | 4.6 | 3.3 | 4.0 |
| Solar (UV) radiation | 7.1 | 5.0 | 6.2 | 3.5 | 3.6 | 3.5 |
| Infectious agents | 2.4 | 3.7 | 2.9 | 2.5 | 3.7 | 3.1 |
| Overweight and obesity | 2.5 | 4.5 | 3.4 | 4.1 | 6.9 | 5.5 |
| Insufficient physical activity | 0.5 | 2.9 | 1.6 | 0.4 | 1.7 | 1.0 |
| Diet - insufficient fibre | 2.3 | 2.1 | 2.2 | 1.4 | 1.7 | 1.5 |
| Diet - insufficient fruit | 1.5 | 1.2 | 1.3 | $6.1{ }^{\text {a }}$ | $3.4{ }^{\text {a }}$ | $4.7{ }^{\text {a }}$ |
| Diet - insufficient non-starchy vegetables | 0.3 | 0.2 | 0.3 |  |  |  |
| Diet - red and processed meat | 2.7 | 1.6 | 2.2 | 3.5 | 1.9 | 2.7 |
| Diet - salt | n/a | n/a | n/a | 0.9 | 0.2 | 0.5 |
| Hormones - MHT | - | 1.1 | 0.3 | - | $1.1{ }^{\text {b }}$ | $0.5{ }^{\text {b }}$ |
| Hormones - combined OCP | - | 0.3 | 0.1 | - | - | - |
| Breast feeding | - | 0.5 | 0.2 | - | 1.7 | 0.9 |
| Ionising radiation | n/a | n/a | n/a | 1.7 | 2.0 | 1.8 |
| Occupation | n/a | n/a | n/a | 4.9 | 2.4 | 3.7 |
| PAF Combined ${ }^{\text {c }}$ | 32.8 | 30.6 | 31.8 | 45.3 | 40.1 | 42.7 |

Abbreviations: PAF - population attributable fraction expressed as a percentage; Aust - Australia; UK - United Kingdom; UV - ultraviolet;
MHT - menopausal hormone therapy; OCP - oral contraceptive pills; n/a - not applicable
a: UK analysis provided PAF estimate for fruit and vegetables combined
b: UK analysis provided PAF estimate for post-menopausal hormones only c: PAF Combined $=1-(1-$ PAF1)*(1-PAF2)*(1-PAFn)

Thirdly, cancer risk assessments and causal associations are continually being reviewed. While we have anticipated some likely changes in assessments of causality for certain cancers (e.g. for obesity and thyroid cancer, non-Hodgkin's lymphoma, multiple myeloma and leukaemia), future assessments of cancer burden will need to consider newly classified associations. Notably, at the time of performing our analyses, obesity was not considered causally associated with prostate cancer; however, an update in November $2014{ }^{27}$ found that body fatness is "probably" a cause of advanced prostate cancer. We performed preliminary analyses to estimate
the ramifications of the WCRF declaration and found that about 353 advanced prostate cancer cases ( $6 \%$ of all advanced prostate cancers; $1.8 \%$ of all prostate cancers) diagnosed in 2010 might be attributed to overweight and obesity in the Australian adult population. ${ }^{7}$ Clearly, newly declared associations of this type will require further investigation as new data are generated.
Finally, consideration was given to the likely latent period between exposure and cancer for each factor, as this determined the choice of time period for obtaining prevalence data. For most factors, the latent period is unknown but is assumed to be many years; thus we
typically sought data for the prevalence of exposures about 10 years before cancer diagnosis, except for specific factors for which other patterns of exposure were considered relevant (e.g. exogenous hormones: MHT current use; breast feeding - total duration; OCP - ever exposed).

## Conclusions

Cancer imposes a heavy burden on Australians' health. In total, cancers are the leading cause of death and disability-adjusted life years (DALYs) in Australia and one of the most expensive disease categories in terms of health expenditure. ${ }^{28,29}$ Many cancers are known to be caused by factors that are prevalent in the environment and for which exposures can be modified.
Continued and better monitoring of exposure to causal factors is recommended to inform primary prevention efforts. While the National Health Surveys are an important tool for assessing the prevalence of healthrelated factors, they are necessarily limited in scope and cannot capture information on all causes of cancer. Moreover, there is no capacity presently to link the records from that survey to health outcomes data. Given the investment in collecting these prevalence data, serious consideration ought to be given to the enormous benefits that would be derived from linking them to health datasets in the future, particularly as techniques for preserving privacy are now well-established. It is greatly concerning, however, that nationally representative, person-based datasets are lacking for key factors known to cause cancer, especially medical exposure to ionising radiation and pharmacological factors. Establishing such repositories should be a national priority.
Conservatively, we estimate that about one-third of cancers arising in the Australian population are attributable to modifiable causes and could - in theory - be prevented if exposure to those factors were eliminated entirely. Of those cancers that have known causes, about half are potentially preventable in men, and more than one-third are preventable in women. Extreme interventions would be required to achieve such outcomes, however, and are unlikely ever to be realised. More modest targets can be identified by calculating potential impact fractions; our analyses suggest substantial gains by targeting smoking, diet, alcohol and sun exposure.

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## PAF Project

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