Nutritional impacts of a fruit and vegetable subsidy programme for disadvantaged Australian Aboriginal children

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Abstract

Healthy food subsidy programmes have not been widely implemented in high-income countries apart from the USA and the UK. There is, however, interest being expressed in the potential of healthy food subsidies to complement nutrition promotion initiatives and reduce the social disparities in healthy eating. Herein, we describe the impact of a fruit and vegetable (F&V) subsidy programme on the nutritional status of a cohort of disadvantaged Aboriginal children living in rural Australia. A before-and-after study was used to assess the nutritional impact in 174 children whose families received weekly boxes of subsidised F&V organised through three Aboriginal medical services. The nutritional impact was assessed by comparing 24 h dietary recalls and plasma carotenoid and vitamin C levels at baseline and after 12 months. A general linear model was used to assess the changes in biomarker levels and dietary intake, controlled for age, sex, community and baseline levels. Baseline assessment in 149 children showed low F&V consumption. Significant increases (P=0.05) in b-cryptoxanthin (28.9 nmol/l, 18%), vitamin C (10.1 μmol/l, 21%) and lutein–zeaxanthin (39.3 nmol/l, 11%) levels were observed at the 12-month follow-up in 115 children, although the self-reported F&V intake was unchanged. The improvements in the levels of biomarkers of F&V intake demonstrated in the present study are consistent with increased F&V intake. Such dietary improvements, if sustained, could reduce non-communicable disease rates. A controlled study of healthy food subsidies, together with an economic analysis, would facilitate a thorough assessment of the costs and benefits of subsidising healthy foods for disadvantaged Aboriginal Australians.

Key words: Fruit and vegetables: Subsidy programmes: Nutrition: Aboriginal children

Higher intake of fruit and vegetables (F&V) reduces the risk of chronic diseases including CVD(1,2), diabetes(3), stroke(4,5) and cancer(6). It is also linked to a lower risk of obesity(7) and hypertension(6). Despite this, low F&V intake is common in many countries(9). Among high-income countries, low F&V consumption is more prevalent in those of lower socio-economic status(10–12). Successful strategies to increase F&V intake offer the potential to reduce the economic costs and prevalence of chronic diseases. Strategies that are effective in those of low socio-economic status have the potential to also reduce health inequalities.

Extensive health promotion campaigns to promote F&V consumption have not been sufficient to reduce the high prevalence of less healthy dietary patterns(13). Pricing strategies, including both taxation of less healthy foods and price discounts or subsidies for healthier foods, have been proposed as alternative strategies that could complement health promotion(14). Economic modelling consistently indicates that price subsidies for healthy foods are likely to produce significant improvements in healthy food consumption, including F&V consumption(15,16), particularly in those on low incomes(17,18). In the USA and the UK, where long-standing

Abbreviations: F&V, fruit and vegetables; RCT, randomised controlled trial.

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food subsidy programmes are important components of social security for low-income families, there have been recent changes made to some of these programmes to better promote the consumption of F&V. Despite the large investment in food subsidy programmes, particularly in the USA, there is limited high-quality evidence of their effectiveness. A 12.5% price discount, but not tailored nutrition education, increased healthy food purchases by 11% in one randomised controlled trial (RCT) in a supermarket setting. Another RCT, currently underway, will enhance these findings by comparing skill building for healthy eating with or without price discounts.

In Australia, Aboriginal and Torres Strait Islander people have poorer health outcomes and are more likely to be of lower socio-economic status than the general population. Pricing strategies to promote healthy nutrition have not been widely implemented in Australia, either among Indigenous Australians or among those with low incomes. In this context, the present study reports on a F&V subsidy programme instituted at three rural Aboriginal community-controlled health services. The programme was initially instituted in 2005 at one service in response to poor nutrition among low-income families visiting the health service. The aims of the programme were to improve nutritional intake and engage participants in preventive health activities.

The aim of the present study was to evaluate whether participation in this F&V subsidy programme for 12 months was associated with changes in nutritional biomarker levels and dietary behaviour in a cohort of Aboriginal children.

Subjects and methods

The fruit and vegetable subsidy programme

A F&V subsidy programme was established by the Bulgarr Ngarr Medical Aboriginal Corporation for the Aboriginal communities in rural towns in the Clarence Valley in New South Wales, Australia, in 2005. The health service invited low-income Aboriginal families with one or more young children to join the programme, which combined annual health assessments, including dental and hearing check-ups, with weekly receipt of a box of subsidised F&V. The boxes of seasonal F&V were organised by local F&V shops in each community at a family’s request. Each family paid five dollars for a box containing $40 worth of F&V (or $60 if five or more children in the household) and collected the box from the F&V shop. The F&V boxes contained a selection of commonly available F&V (with more vegetables than fruit) chosen by the F&V shops based on value for money and availability (see Supplementary file 1 for a list of options provided to guide the F&V shop staff). Families could exchange some unwanted F&V for alternative F&V. The programme was developed after the success of a nutrition programme in a remote Aboriginal community school in the Clarence Valley, described elsewhere.

This is an ongoing programme in the Clarence Valley; however, this evaluation study involved all new families who joined the programme at this time. At baseline (before receiving F&V) and after 12 months, all children in these families underwent assessments of dietary intake and nutritional status specifically as part of the evaluation. The recruitment and baseline assessments were undertaken between December 2008 and September 2009, with follow-up assessments being completed between December 2009 and September 2010.

Additional funding enabled the Galambila Aboriginal Health Service in Coffs Harbour and the Giingan Darrunday Marlaanggu Health Clinic in the Nambucca Valley to institute similar F&V subsidy programmes for the duration of this evaluation study. These health services decided to participate in this evaluation study, and all families at these locations participated in the nutritional status assessments for this evaluation. Although the demographics and health status are similar in each of these Aboriginal communities, the availability of and arrangements with the F&V shops varied in the communities. In Coffs Harbour, families received vouchers from the health service, which they redeemed at the F&V shop by selecting their own F&V. In the Nambucca Valley, the F&V shop was in a different town, hence the health service staff collected and delivered the boxes of F&V to families at their homes and collected the $5 contribution from them.

Nutrition promotion complemented the F&V boxes with seasonal recipe and practical cooking and nutrition education sessions conducted by dietitians (in the Clarence Valley and the Nambucca Valley) or trained nutrition health workers (in Coffs Harbour). There were three or four of these sessions conducted over 12 months in each of the towns in the Clarence Valley, although attending the sessions was not mandatory and less than half of the participating families attended the sessions. There were five sessions conducted over 5 weeks in the Nambucca Valley, which were well attended. No group education occurred in Coffs Harbour; however, individual families were given nutrition education after the children’s 24h dietary recalls by the nutrition health workers. Simple seasonal recipes were shared between sites, although the content of nutrition education was determined by the local staff.

Participants

The participants were low-income Aboriginal families (predominantly unemployed or receiving pensions) with one or more children aged ≤17 years (at commencement). Many of the children had identified nutrition risk (e.g. underweight or overweight and chronic or recurrent infections) or presented frequently with episodes of illness to the health service. Parents/carers provided consent and agreed for their children to undergo annual health assessments including the research evaluation assessments.

Data collection and laboratory methods

As part of the annual health assessments, each child’s nutritional status was assessed with a 24h dietary recall and a blood test before and after 12 months on the programme. These assessments were completed as close to 1 year apart as possible to minimise seasonal variation. Blood samples
were analysed for carotenoid, vitamin A, vitamin E, vitamin C, lipid and C-reactive protein levels. Individual results were discussed with the participant/carer at follow-up appointments, and overall summary results were presented in a non-technical language to the communities during focus groups.

The 24 h dietary recalls were completed by the participants face to face with a dietitian or a trained Aboriginal health worker. For participants aged <10 years, these were completed together with a parent/carer. Data collection was undertaken by following the multi-pass method described in the 1995 Australian National Nutrition Survey(25). In addition, the participants were asked about their usual consumption of F&V using validated short dietary questions based on the Dietary Intake Assessment Tool developed in Queensland, Australia(26). Short dietary questions have been developed for use in children's dietary surveys(26) and were used in the present study to contrast with the estimates of self-reported F&V intake from the 24 h recalls.

Non-fasting venous blood samples were collected by clini- cal staff into tubes wrapped in a foil and stored on ice before transport to a local pathology laboratory. In the laboratory, they were separated, and plasma samples for the measurement of carotenoid, vitamin A and vitamin E levels were frozen at −20°C. The frozen plasma samples were trans- ported on dry ice and stored at −80°C before analysis. The samples were protected from light throughout processing to prevent carotenoid degradation.

Blood samples were analysed for the levels of lipids and C-reactive protein at either Grafton Hospital Pathology, on a Cobas Integra 800 chemistry analyser (Roche Diagnostics), or Symbion Laverty Pathology, Coffs Harbour, on a Cobas Integra 400 chemistry analyser (Roche Diagnostics). Frozen plasma samples were transported on dry ice to Royal Prince Alfred Hos- pital, Sydney, and were analysed for vitamin C level using HPLC with electrochemical detection. Due to logistical problems, samples for the measurement of vitamin C level were not obtained at baseline in Coffs Harbour or the Nambucca Valley.

The levels of carotenoids, retinol and tocopherols were ana- lysed using HPLC, as described by Su et al.(27) with minor changes. Briefly, after extraction and drying, 50 µl of a plasma mixture were analysed using HPLC (Shimadzu HPLC machine equipped with an SPD-M20A PDA detector and a NovoPak C18 column; Shimadzu Corporation) with the detection of absorbance at 292 nm for tocopherols, 325 nm for reti- nol and 450 nm for carotenoids. The samples were assayed using standardised techniques by laboratory staff who were unaware of the study design.

The dietary recall data were analysed using the FoodWorks Professional 2009 dietary software programme (Xyris) using the AusNut 2007 Australian food database. Data entry and analysis were completed by a post-graduate dietetics student with input from an experienced dietetics researcher.

Statistical analysis

The analysis sought to answer the question of whether the F&V programme was associated with changes in biomarker levels and dietary behaviour, and the primary outcomes analysed were the mean change in the levels of each carotenoid. The changes between the follow-up data and the baseline data for each participant were compared using paired t tests and the general linear model function in SPSS version 19 (IBM). The analysis was based on completed observations. Outcomes were assessed by comparing the change between the pre-levels and the post-levels, which increased the likelihood of these outcomes being normally distributed and checked for normality before undertaking parametric analyses. The crude changes in biomarker levels and dietary outcomes are presented as means and 95 % CI using paired t tests. These outcomes were adjusted by age, sex and community using a general linear model to account for differences due to age, sex or community. A 95 % CI of the mean of the difference for each outcome that did not cross zero indicated a statistically significant difference. The effect of additional covariates in the statistical model was assessed, including total cholesterol and baseline values for biomarkers and baseline values for dietary outcomes. Based on previous studies, the sample size required to detect the expected mean changes of 10–50 % in the levels of these carotenoids with a statistical significance level of 5 % and power of 80 % ranged from 9 for β-carotene to 108 for lycopene(28,29). Estimated sample sizes accounted for the family-level design effect, assuming an average family size of 2.6 with a family-level intra-class correlation coefficient of 0.5(30). Given the exploratory nature and hypothesis-generating intentions of the study, analyses were not adjusted for the design effect or for multiple comparisons. Not accounting for multiple comparisons was deemed to be justified on the basis of our intention to estimate the extent of changes in biomarker levels, consequent to the introduction of a fruit/vegetable intervention, rather than to formally test specific hypotheses as to whether changes occurred. Our statistical procedures reflect this focus on estimation rather than on hypothesis testing.

Ethics

The present study was conducted according to the guidelines laid down in the Declaration of Helsinki, and all procedures involving human subjects were approved by the human research ethics committees of the University of Melbourne, the University of South Australia and the Aboriginal Health and Medical Research Council (New South Wales). Community consent was obtained from the Boards of the three participating health services. The North Coast Area Health Service Research Ethics Committee approved the clinical audits of the participants' hospital records. Written informed consent was obtained from all subjects/parents or guardians.

Results

There were fifty-five Aboriginal families, which included 174 children, recruited to participate in the evaluation of the F&V programme at the three Aboriginal Health Services. All assessments were of these children aged 0–17 years at base- line, with the majority of the children being aged between 3 and 12 years. The demographic characteristics of the
participants at baseline are given in Table 1. As shown in Table 1, this was a cohort of low-income families with the majority receiving government benefits as their main source of income. The majority of adults in these households were smokers. There were differences between the cohorts of families at each location: the children were older and a small proportion of adults smoked in Coffs Harbour, while the children were younger and a greater proportion were smokers. There were differences between the cohorts of income. The majority of adults in these households were receiving government benefits as their main source of income.

Impact of the fruit and vegetable programme on biomarker levels and fruit and vegetable intake

The median period between baseline and follow-up assessments was 370 d (interquartile range 354–407 d). In the overall sample of children, mean β-cryptoxanthin, lutein–zeaxanthin and vitamin C levels were significantly higher after participation in the F&V programme for 12 months than the baseline levels (Table 2). These changes remained significant after adjustment for sex, age and community (Table 2). There were no significant changes in α-tocopherol, γ-tocopherol or retinol levels. Adjustment for total cholesterol did not alter the mean change in the levels of carotenoids overall or in each community. Adjustment for the baseline level of biomarkers did not alter the mean changes in the levels carotenoids or vitamin C overall. There were no significant differences in biomarker levels by sex at baseline, although girls did show a greater mean change in β-cryptoxanthin levels than boys at follow-up (girls: 43.6 nmol/l; boys: 2.4 nmol/l, \( F = 3.43, P = 0.07 \)). The use of vitamin and mineral supplements was reported by 13% of the children at baseline and 9% of the children at the follow-up assessments. The pattern of changes in biomarker levels was similar, except that the increase in β-cryptoxanthin level was not statistically significant, after excluding those participants who reported supplement use at follow-up.

At baseline, the F&V programme participants (and/or carers) reported that the children’s mean intake of fruit was

### Table 1. Baseline demographic characteristics of the participating children by individual community (n 174) (Mean values and standard deviations; number of participants and percentages)

<table>
<thead>
<tr>
<th></th>
<th>All communities</th>
<th>Clarence</th>
<th>Coffs Harbour</th>
<th>Nambucca</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( n )</td>
<td>%</td>
<td>( n )</td>
<td>%</td>
</tr>
<tr>
<td>Families</td>
<td>55</td>
<td>30</td>
<td>124</td>
<td>90</td>
</tr>
<tr>
<td>Children</td>
<td>174</td>
<td>90</td>
<td>36</td>
<td>48</td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>7.6</td>
<td>7.5</td>
<td>11.0</td>
<td>5.8</td>
</tr>
<tr>
<td>SD</td>
<td>4.2</td>
<td>3.8</td>
<td>3.3</td>
<td>4.3</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>82</td>
<td>47</td>
<td>51</td>
<td>18</td>
</tr>
<tr>
<td>Female</td>
<td>92</td>
<td>53</td>
<td>49</td>
<td>18</td>
</tr>
<tr>
<td>Children with smoker(s)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>in the household*</td>
<td>107</td>
<td>66</td>
<td>62</td>
<td>18</td>
</tr>
<tr>
<td>Families receiving</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>unemployment benefits</td>
<td>51</td>
<td>93</td>
<td>28</td>
<td>10</td>
</tr>
<tr>
<td>and pensions</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Percentage of those with a valid response to smokers in the household.

Table 2. Plasma biomarker levels of the participating children (n 115) before and after the fruit and vegetable programme (Mean values and standard deviations; mean differences and 95% confidence intervals)

<table>
<thead>
<tr>
<th></th>
<th>Before (nmol/l)</th>
<th>After (nmol/l)</th>
<th>Mean difference† (nmol/l)</th>
<th>95% CI</th>
<th>Adjusted mean difference‡ (nmol/l)</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>β-Carotene</td>
<td>346.6 (173.0)</td>
<td>338.9 (195.4)</td>
<td>-7.7</td>
<td>-39.5, 24.0</td>
<td>-12.2</td>
<td>-48.4, 24.1</td>
</tr>
<tr>
<td>α-Carotene</td>
<td>66.1 (36.7)</td>
<td>73.7 (53.3)</td>
<td>7.6</td>
<td>-2.1, 17.3</td>
<td>4.3</td>
<td>-6.7, 15.2</td>
</tr>
<tr>
<td>Lycopene</td>
<td>637.7 (306.5)</td>
<td>586.9 (295.1)</td>
<td>-50.9</td>
<td>-106.8, 5.1</td>
<td>-42.9</td>
<td>-104.1, 18.3</td>
</tr>
<tr>
<td>β-Cryptoxanthin</td>
<td>159.0 (87.2)</td>
<td>183.9 (167.0)</td>
<td>25.0*</td>
<td>2.9, 47.1</td>
<td>28.9*</td>
<td>4.2, 53.6</td>
</tr>
<tr>
<td>Lutein–zeaxanthin</td>
<td>303.3 (163.5)</td>
<td>337.8 (170.4)</td>
<td>34.5*</td>
<td>4.3, 64.6</td>
<td>39.3*</td>
<td>6.2, 72.5</td>
</tr>
<tr>
<td>Vitamin C (n 57) (μmol/l)</td>
<td>49.1 (21.4)</td>
<td>59.6 (24.6)</td>
<td>10.5*</td>
<td>4.3, 18.7</td>
<td>10.5*</td>
<td>2.0, 18.1</td>
</tr>
<tr>
<td>α-Tocopherol (μmol/l)</td>
<td>23.6 (5.3)</td>
<td>32.2 (9.4)</td>
<td>-8.4</td>
<td>-13.0, 0.5</td>
<td>-8.4</td>
<td>-1.4, 0.7</td>
</tr>
<tr>
<td>γ-Tocopherol (μmol/l)</td>
<td>2.4 (1.2)</td>
<td>2.2 (1.1)</td>
<td>-0.2</td>
<td>-0.4, 0.1</td>
<td>-0.1</td>
<td>-0.4, 0.2</td>
</tr>
<tr>
<td>Retinol (μmol/l)</td>
<td>2.1 (0.7)</td>
<td>2.1 (0.7)</td>
<td>0.0</td>
<td>-0.1, 0.2</td>
<td>0.0</td>
<td>-0.1, 0.2</td>
</tr>
</tbody>
</table>

* Values were significantly different from zero (\( P < 0.05 \)).
† Unadjusted mean difference (after–before), paired \( t \) test.
‡ Adjusted for sex, age and community using the general linear model (univariate ANOVA).
Table 3. Fruit and vegetable intake of the participating children (n 121) before and after the fruit and vegetable programme

(Mean values and standard deviations; mean differences and 95 % confidence intervals)

<table>
<thead>
<tr>
<th>Servings/d</th>
<th>Before Mean</th>
<th>Before SD</th>
<th>After Mean</th>
<th>After SD</th>
<th>Mean difference‡</th>
<th>95 % CI</th>
<th>Adjusted mean difference§</th>
<th>95 % CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>24 h dietary recall</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fruit</td>
<td>1·0</td>
<td>1·4</td>
<td>1·0</td>
<td>1·1</td>
<td>−0·03</td>
<td>−0·30, 0·24</td>
<td>−0·10</td>
<td>−0·42, 0·23</td>
</tr>
<tr>
<td>Vegetables</td>
<td>0·6</td>
<td>0·9</td>
<td>0·5</td>
<td>0·8</td>
<td>−0·09</td>
<td>−0·31, 0·14</td>
<td>−0·13</td>
<td>−0·40, 0·15</td>
</tr>
<tr>
<td>Vegetables (including potato)</td>
<td>0·8</td>
<td>1·2</td>
<td>0·7</td>
<td>0·9</td>
<td>−0·15</td>
<td>−0·42, 0·12</td>
<td>−0·20</td>
<td>−0·53, 0·12</td>
</tr>
<tr>
<td>Fruit and vegetables</td>
<td>1·8</td>
<td>1·8</td>
<td>1·6</td>
<td>1·6</td>
<td>−0·18</td>
<td>−0·56, 0·20</td>
<td>−0·30</td>
<td>−0·76, 0·15</td>
</tr>
<tr>
<td>Short questions§</td>
<td>2·2</td>
<td>1·0</td>
<td>2·5</td>
<td>1·4</td>
<td>0·23*</td>
<td>0·004, 0·46</td>
<td>0·11</td>
<td>−0·18, 0·39</td>
</tr>
<tr>
<td>Vegetables (n 118)</td>
<td>2·5</td>
<td>1·4</td>
<td>2·2</td>
<td>1·1</td>
<td>−0·27</td>
<td>−0·54, 0·00</td>
<td>−0·49*</td>
<td>−0·82, −0·16</td>
</tr>
</tbody>
</table>

* Values were significantly different from zero (P<0·05).
† Unadjusted mean difference (after–before), paired t test.
‡ Adjusted for sex, age and community using the general linear model (univariate ANOVA).
§ Questions about average servings of fruit and vegetables consumed, asked at the time of the 24 h dietary recall.

Changes in macronutrient intake

There was a decrease in the self-reported macronutrient intake in the follow-up dietary recalls than at baseline, particularly of carbohydrates and total sugars (Supplementary file 2, available online). The mean total energy intake at baseline was 7570 (SD 3156) kJ, while at follow-up, it was 6482 (SD 2197) kJ. To facilitate comparison with the large variation in intake among these children aged 0–17 years, macronutrient intake per MJ was also analysed. In the analysis of macronutrient intake per MJ, there were small but statistically significant decreases in total sugar (1·6 g/MJ) and added sugar (1·2 g/MJ) intake, while the intake of other macronutrients was unchanged (Table 4). However, after adjustment for sex, age and community.

Table 4. Change in macronutrient density (g/MJ) among the participants aged 0–17 years (n 121) before and after the fruit and vegetable programme

(Mean values and standard deviations; mean differences and 95 % confidence intervals)

<table>
<thead>
<tr>
<th>Macronutrients†</th>
<th>Before Mean</th>
<th>Before SD</th>
<th>After Mean</th>
<th>After SD</th>
<th>Mean difference‡</th>
<th>95 % CI</th>
<th>Adjusted mean difference§</th>
<th>95 % CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein</td>
<td>8·2</td>
<td>2·8</td>
<td>8·5</td>
<td>2·7</td>
<td>−0·2</td>
<td>−0·4, 0·9</td>
<td>0·1</td>
<td>−0·6, 0·9</td>
</tr>
<tr>
<td>Total fat</td>
<td>8·9</td>
<td>1·6</td>
<td>9·3</td>
<td>1·5</td>
<td>−0·4</td>
<td>−0·01, 0·8</td>
<td>0·5</td>
<td>0·1, 0·9</td>
</tr>
<tr>
<td>Saturated fat</td>
<td>4·1</td>
<td>1·2</td>
<td>4·2</td>
<td>1·0</td>
<td>−0·02</td>
<td>−0·2, 0·3</td>
<td>0·2</td>
<td>−0·1, 0·4</td>
</tr>
<tr>
<td>CHO</td>
<td>30·7</td>
<td>4·8</td>
<td>29·6</td>
<td>4·8</td>
<td>−1·1</td>
<td>−2·2, 0·02</td>
<td>−1·2</td>
<td>−2·5, 0·1</td>
</tr>
<tr>
<td>Total sugar</td>
<td>14·7</td>
<td>5·6</td>
<td>13·1</td>
<td>5·2</td>
<td>−1·6*</td>
<td>−2·8, −0·4</td>
<td>−1·0</td>
<td>−2·4, 0·4</td>
</tr>
<tr>
<td>Added sugar</td>
<td>10·6</td>
<td>5·9</td>
<td>9·4</td>
<td>5·2</td>
<td>−1·2*</td>
<td>−2·4, −0·001</td>
<td>0·03</td>
<td>−1·4, 1·5</td>
</tr>
<tr>
<td>Starch</td>
<td>15·8</td>
<td>4·4</td>
<td>16·3</td>
<td>4·7</td>
<td>0·5</td>
<td>−0·7, 1·6</td>
<td>−0·2</td>
<td>−1·6, 1·2</td>
</tr>
</tbody>
</table>

CHO, carbohydrate.
* Values were significantly different from zero (P<0·05).
† Alcohol intake was not reported in any 24 h dietary recall.
‡ Unadjusted mean difference (after–before), paired t test.
§ Adjusted for sex, age and community using the general linear model (univariate ANOVA).
community, there were no significant changes in the intake of any of the macronutrients per MJ overall (Table 4).

**Discussion**

The major findings of the present study of the impact of subsidised fresh F&V provided by three local Aboriginal community-controlled health services to disadvantaged Aboriginal families in rural communities were the significant increases in the levels of biomarkers of F&V intake including β-cryptoxanthin, vitamin C and lutein–zeaxanthin. In contrast to the changes in biomarker levels, there were no changes in the self-reported intake of F&V (when using either the 24 h dietary recall or the short dietary questions on usual intake). Furthermore, self-reported ‘usual intake’ was considerably higher than that reported in the 24 h dietary recalls.

These findings highlighted the well-known limitations of the self-reported dietary intake and the role of biomarkers as alternative more objective measures of F&V intake. Collecting self-reported dietary intake data is particularly challenging for children, as the dietary intake data are collected with inputs from both a parent/carer and a child, which may lead to a different likelihood of bias due to difficulties with the estimation of quantities and under-reporting. The decrease in self-reported energy intake at the 12-month follow-up (which would be expected to increase slightly in children) suggested that repeated self-reported dietary intake measurements may be susceptible to reporting bias in assessing the impact of nutrition interventions. The present study has also shown that it is feasible to obtain blood samples for the measurement of biomarker levels from children in a community-based intervention study with appropriate community engagement.

The baseline carotenoid levels in the present study were similar, except for lycopene level, which was higher, to those reported in a representative sample of 4231 American children aged 6–16 years and β-carotene level was also similar to that reported in 467 Australian preschool-aged children. While no comparable published data were obtained similar to that reported in 467 Australian preschool-aged children aged 6–16 years, and β-carotene level was also similar to that reported in 467 Australian preschool-aged children. While no comparable published data were obtained, the most challenging for children, as the dietary intake data are collected with inputs from both a parent/carer and a child, which may lead to a different likelihood of bias due to difficulties with the estimation of quantities and under-reporting. The decrease in self-reported energy intake at the 12-month follow-up (which would be expected to increase slightly in children) suggested that repeated self-reported dietary intake measurements may be susceptible to reporting bias in assessing the impact of nutrition interventions. The present study has also shown that it is feasible to obtain blood samples for the measurement of biomarker levels from children in a community-based intervention study with appropriate community engagement.

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thus have the most to gain from strategies that increase F&V intake.

An obvious limitation of the present study was the lack of a control group. This would have been difficult to implement given the clear economic benefit of the F&V subsidy, irrespective of any nutritional or health impacts that were the focus of the present study. However, a control group was contemplated as an equitable way to manage the waiting list of eligible families for this programme, but was ultimately considered impractical given the limited local research capacity in a busy Aboriginal community-controlled health service. Regression to the mean and physiological changes with age was addressed in the analysis by including age and baseline levels as covariates in the model. Another limitation of the present study was that the impact of the F&V subsidies could not be separated from that of other elements of the programme, including nutrition education and increased engagement with Aboriginal primary health care services. However, other studies have suggested that the benefits of this programme are likely to be primarily due to the F&V subsidies.

The use of non-fasting blood specimens was a potential source of error as serum vitamin C and β- and β’-carotene levels have been shown to increase acutely several hours post-ingestion of carotene- or ascorbate-rich foods. However, it was considered impractical to expect these children to arrive fasted for these assessments. In addition, the biological half-life of the serum carotenoids ranges from 26 d for lycopene to 76 d for lutein, while the biological half-life of vitamin C is 10–20 d. This suggests that the serum levels reflect intake in the previous several weeks, provided intake is relatively consistent. Thus, the significant increases in β-cryptoxanthin, lutein–zeaxanthin and vitamin C levels are likely to represent medium-term F&V intake, although the serum levels could have been influenced by recent ingestion of F&V, particularly vitamin C level by supplements or vitamin C-fortified foods.

The findings of this evaluation study of the nutritional improvements associated with F&V subsidies suggest that there is a need for a controlled study to verify these conclusions. Higher-quality longitudinal data would also provide the opportunity for an economic analysis that could also inform decisions about investing in food subsidies. The validity of biomarkers as indicators of F&V intake is well established in adults. International studies have established the normal range of plasma carotenoid concentrations in children. However, further studies are needed to confirm their validity and practicality as measures of F&V intake in interventions involving children. This is important given the discrepancies between self-reported dietary intake and biomarker levels observed in the present study.

The potential reduction in chronic disease risk due to increased F&V intake suggests that it would also be useful to assess the nutritional status of adults in family-focused nutrition intervention studies, given the long time lag between improving F&V intake in children and any manifestation of chronic diseases. Increased F&V intake is perhaps more likely to be sustained if it involves both adults and children in families. The sustainability of any change in F&V intake underpins any long-term health impacts. This programme provided a heavily subsidised supply of fresh F&V to these families, sufficient to meet the requirements of a family of five, which may be necessary to engage the most disadvantaged families. However, it would be useful to trial the impacts on dietary intake or nutritional status of F&V programmes that include lesser value subsidies or a larger participant contribution.

In conclusion, F&V subsidies improve the nutritional status of disadvantaged Indigenous Australian children. The measurement of plasma carotenoid and vitamin C levels as biomarkers are feasible to measure F&V intake in these children and may offer an objective alternative to self-reported dietary intake. Improving the F&V intake of these children is challenging and emphasises the need to link any subsidies with effective nutrition education to promote meal planning and preparation skills. The delivery of F&V subsidies in a community health setting strengthens the link between preventive health activities and improved nutrition. A controlled study of subsidised healthy foods is warranted to investigate the cost-effectiveness of this strategy to improve both the nutrition and health of the most disadvantaged families.

Supplementary material

To view supplementary material for this article, please visit http://dx.doi.org/10.1017/S0007114513001700

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References


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