Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low-and middle-income countries

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Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)


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Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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Analysis 3.5. Comparison 3 Unconditional cash transfers versus no unconditional cash transfers: social determinants of health, Outcome 5 Is extremely poor.

Analysis 4.1. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 1 Height for age (standard deviations) by rural-urban residency, currently.

Analysis 4.2. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 2 Height for age (standard deviations) by income poverty status, currently.

Analysis 4.3. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 3 Has had any illness in previous 2 weeks to 3 months.

Analysis 4.4. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 4 Food security index by gender.

Analysis 4.5. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 5 Dietary diversity (Household Dietary Diversity Score) in previous week by rural-urban residency.

Analysis 4.6. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 6 Level of dietary diversity (Household Dietary Diversity Score) in previous week by gender.

Analysis 4.7. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 7 Level of dietary diversity (Household Dietary Diversity Score) in previous week by income poverty status.

Analysis 4.8. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 8 Level of depression (Center for Epidemiologic Studies Depression Score) by rural-urban residency.

Analysis 4.9. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 9 Level of depression (Center for Epidemiologic Studies Depression Score) by gender.

Analysis 4.10. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 10 Level of depression (Center for Epidemiologic Studies Depression Score) by income poverty status.
Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries

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ABSTRACT

Background
Unconditional cash transfers (UCTs; provided without obligation) for reducing poverty and vulnerabilities (e.g. orphanhood, old age or HIV infection) are a type of social protection intervention that addresses a key social determinant of health (income) in low- and middle-income countries (LMICs). The relative effectiveness of UCTs compared with conditional cash transfers (CCTs; provided so long as the recipient engages in prescribed behaviours such as using a health service or attending school) is unknown.

Objectives
To assess the effects of UCTs for improving health services use and health outcomes in vulnerable children and adults in LMICs. Secondary objectives are to assess the effects of UCTs on social determinants of health and healthcare expenditure and to compare to effects of UCTs versus CCTs.

Search methods
We searched 17 electronic academic databases, including the Cochrane Public Health Group Specialised Register, the Cochrane Database of Systematic Reviews (the Cochrane Library 2017, Issue 5), MEDLINE and Embase, in May 2017. We also searched six electronic grey literature databases and websites of key organisations, handsearched key journals and included records, and sought expert advice.

Selection criteria
We included both parallel group and cluster-randomised controlled trials (RCTs), quasi-RCTs, cohort and controlled before-and-after (CBAs) studies, and interrupted time series studies of UCT interventions in children (0 to 17 years) and adults (18 years or older) in LMICs. Comparison groups received either no UCT or a smaller UCT. Our primary outcomes were any health services use or health outcome.
Data collection and analysis

Two reviewers independently screened potentially relevant records for inclusion criteria, extracted data and assessed the risk of bias. We tried to obtain missing data from study authors if feasible. For cluster-RCTs, we generally calculated risk ratios for dichotomous outcomes from crude frequency measures in approximately correct analyses. Meta-analyses applied the inverse variance or Mantel-Haenszel method with random effects. We assessed the quality of evidence using the GRADE approach.

Main results

We included 21 studies (16 cluster-RCTs, 4 CBAs and 1 cohort study) involving 1,092,877 participants (36,068 children and 1,056,809 adults) and 31,865 households in Africa, the Americas and South-East Asia in our meta-analyses and narrative synthesis. The 17 types of UCTs we identified, including one basic universal income intervention, were pilot or established government programmes or research experiments. The cash value was equivalent to 1.3% to 53.9% of the annualised gross domestic product per capita. All studies compared a UCT with no UCT, and three studies also compared a UCT with a CCT. Most studies carried an overall high risk of bias (i.e. often selection and/or performance bias). Most studies were funded by national governments and/or international organisations.

Throughout the review, we use the words ‘probably’ to indicate moderate-quality evidence, ‘may/maybe’ for low-quality evidence, and ‘uncertain’ for very low-quality evidence. UCTs may not have impacted the likelihood of having used any health service in the previous 1 to 12 months, when participants were followed up between 12 and 24 months into the intervention (risk ratio (RR) 1.04, 95% confidence interval (CI) 1.00 to 1.09, P = 0.07, 5 cluster-RCTs, N = 4972, I² = 2%, low-quality evidence). At one to two years, UCTs probably led to a clinically meaningful, very large reduction in the likelihood of having had any illness in the previous two weeks to three months (odds ratio (OR) 0.73, 95% CI 0.57 to 0.93, 5 cluster-RCTs, N = 8446, I² = 57%, moderate-quality evidence). Evidence from five cluster-RCTs on food security was too inconsistent to be combined in a meta-analysis, but it suggested that at 13 to 24 months’ follow-up, UCTs could increase the likelihood of having been food secure over the previous month (low-quality evidence). UCTs may have increased participants’ level of dietary diversity over the previous week, when assessed with the Household Dietary Diversity Score and followed up 24 months into the intervention (mean difference (MD) 0.59 food categories, 95% CI 0.18 to 1.01, 4 cluster-RCTs, N = 9347, I² = 79%, low-quality evidence). Despite several studies providing relevant evidence, the effects of UCTs on the likelihood of being moderately stunted and on the level of depression remain uncertain. No evidence was available on the effect of a UCT on the likelihood of having died. UCTs probably led to a clinically meaningful, moderate increase in the likelihood of currently attending school, when assessed at 12 to 24 months into the intervention (RR 1.06, 95% CI 1.03 to 1.09, 6 cluster-RCTs, N = 4800, I² = 0%, moderate-quality evidence). The evidence was uncertain for whether UCTs impacted livestock ownership, extreme poverty, participation in child labour, adult employment or parenting quality. Evidence from six cluster-RCTs on healthcare expenditure was too inconsistent to be combined in a meta-analysis, but it suggested that UCTs may have increased the amount of money spent on health care at 7 to 24 months into the intervention (low-quality evidence). The effects of UCTs on health equity (or unfair and remedial health inequalities) were very uncertain. We did not identify any harms from UCTs. Three cluster-RCTs compared UCTs versus CCTs with regard to the likelihood of having used any health services, the likelihood of having had any illness or the level of dietary diversity, but evidence was limited to one study per outcome and was very uncertain for all three.

Authors’ conclusions

This body of evidence suggests that unconditional cash transfers (UCTs) may not impact a summary measure of health service use in children and adults in LMICs. However, UCTs probably or may improve some health outcomes (i.e. the likelihood of having had any illness, the likelihood of having been food secure, and the level of dietary diversity), one social determinant of health (i.e. the likelihood of attending school), and healthcare expenditure. The evidence on the relative effectiveness of UCTs and CCTs remains very uncertain.

Plain language summary

Unconditional cash transfers for reducing poverty: effect on health services use and health outcomes in low- and middle-income countries

Review question

Some programmes provide cash transfers or grants for reducing poverty and vulnerabilities without imposing any obligations on the recipients (‘unconditional cash transfers’, or UCTs) in low- and middle-income countries (LMICs). Other times, people can only receive these cash transfers if they engage in required behaviours, such as using health services or sending their children to school (‘conditional cash transfers’, or CCTs). This review aimed to find out whether receiving UCTs would improve people’s use of health services and their health outcomes, compared with not receiving a UCT, receiving a smaller UCT amount or receiving a CCT. It also aimed to assess the effects of UCTs on daily living conditions that determine health and healthcare spending.

Background

UCTs are a type of social protection intervention that addresses income. It is unknown whether UCTs are more, less or equally as effective as CCTs. We reviewed the evidence on the effect of UCTs on health service use and health outcomes among children and adults in LMICs.

Study characteristics
The evidence is current to May 2017. We included experimental and selected non-experimental studies of UCTs in people of all ages in LMICs. We included studies that compared participants who received a UCT with those who received no UCT. We looked for studies that examined health services use and health outcomes.

We found 21 studies (16 experimental and 5 non-experimental ones) with 1,092,877 participants (36,068 children and 1,056,809 adults) and 31,865 households in Africa, the Americas and South-East Asia. The UCTs were government programmes or research experiments. Most studies were funded by national governments and/or international organisations.

**Key results**

We use the words 'probably' to indicate moderate-quality evidence, 'may/maybe' for low-quality evidence, and 'uncertain' for very low-quality evidence. A UCT may not impact the likelihood of having used any health service in the previous 1 to 12 months. UCTs probably led to a clinically meaningful, very large reduction in the risk of having had any illness in the previous two weeks to three months. They may increase the likelihood of having had secure access to food over the previous month. They may also increase the average number of different food groups consumed in the household over the previous week. Despite several studies providing relevant evidence, the effects of UCTs on the likelihood of stunting and on depression levels remain uncertain. No study estimated effects on dying. UCTs probably led to a clinically meaningful, moderate increase in the likelihood of currently attending school. The evidence was uncertain for whether UCTs impacted livestock ownership, extreme poverty, participation in child labour, adult employment and parenting quality. UCTs may increase the amount of money spent on health care. The effects of UCTs on differences in health were very uncertain. We did not identify any harms from UCTs. Three experimental studies reported evidence on the impact of a UCT compared with a CCT on the likelihood of having used any health services, the likelihood of having had any illness or the average number of food groups consumed in the household, but evidence was limited to one study per outcome and was very uncertain for all three.

**Quality of the evidence**

Of the seven prioritised primary outcomes, the body of evidence for one outcome was of moderate quality, for three outcomes of low quality, for two outcomes of very low quality, and for one outcome, there was no evidence at all.

**Conclusions**

This body of evidence suggests that unconditional cash transfer (UCTs) may not impact health services use among children and adults in LMICs. UCTs probably or may improve some health outcomes (i.e. the likelihood of having had any illness, the likelihood of having secure access to food, and diversity in one's diet), one social determinant of health (i.e. the likelihood of attending school), and healthcare expenditure. The evidence on the health effects of UCTs compared with those of CCTs is uncertain.
### Summary of findings: unconditional cash transfer versus no unconditional cash transfer: primary outcomes

**Patient or population:** children (0 to 17 years) and adults (≥ 18 years) or households  
**Settings:** low- and middle-income countries  
**Intervention:** an unconditional cash transfer for reducing poverty and/or vulnerabilities  
**Comparison:** no unconditional cash transfer

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Illustrative comparative risks* (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>Number of participants (number of studies)</th>
<th>Quality of the evidence (GRADE)</th>
<th>Comments</th>
</tr>
</thead>
</table>
| Has used any health service in previous 1 to 12 months  
Follow-up: 12 months to 24 months | 447 per 1000 (447 to 487) | RR 1.04 (1.00 to 1.09) | 4972 (5 RCTs) | ⊕⊕⊝ | Better indicated by a higher value. In conclusion, a UCT may not have an effect on this outcome. |
| Is moderately stunted  
Assessed with: height-for-age z-score ≤ −2 SD  
Follow-up: 24 months | 337 per 1000 (253 to 408) | RR 0.96 (0.75 to 1.21) | 551 (2 RCTs) | ⊕⊕⊕⊝ | Better indicated by a lower value. Additional evidence on the effect of a UCT on height-for-age scores from a second meta-analysis of 2 studies and on average height from an additional RCT was also very uncertain. In conclusion, we are very uncertain about the effect of a UCT on this outcome. |
| Has died | No evidence available on this outcome | | | | |
| Has had any illness in previous 2 weeks to 3 months  
Follow-up: 12 months to 24 months | 370 per 1000 (211 to 344) | OR 0.73 (0.57 to 0.93) | 8446 (5 RCTs) | ⊕⊕⊕| Better indicated by a lower value. One additional RCT reported that a UCT probably reduced the risk of having had an acute respiratory disease, and another additional RCT reported that a UCT led to a large, clinically meaningful reduction in the risk of illness or injury in the household. In conclusion, a
### Has been food secure in previous month

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Description</th>
<th>Count</th>
<th>GRADE</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not pooled</td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

**Follow-up:** range 13 months to 24 months

### Level of dietary diversity in previous week

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Description</th>
<th>Count</th>
<th>GRADE</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>The mean level of dietary diversity was 1.46 food categories consumed</td>
<td></td>
<td></td>
<td></td>
<td>Better indicated by a higher value.</td>
</tr>
<tr>
<td>0.59 food categories consumed higher (0.18 to 1.01 higher)</td>
<td></td>
<td></td>
<td>Low&lt;sup&gt;a,d&lt;/sup&gt;</td>
<td>Because of the very high level of statistical heterogeneity, we do not report totals from the meta-analysis. 2 RCTs reported a moderate, clinically meaningful increase in the likelihood of being food secure, whereas a third RCT reported that a UCT had perhaps not led to a change in this likelihood. A fourth RCT reported a moderate, probably clinically meaningful reduction in a summary measure of household food insecurity. In conclusion, a UCT may perhaps have had a beneficial effect on this outcome.</td>
</tr>
</tbody>
</table>

**Follow-up:** 24 months

### Level of depression

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Description</th>
<th>Count</th>
<th>GRADE</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>The mean level of depression was an unclear CES-D Score</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>0.06 of 1 SD of the CES-D score lower (0.25 of 1 SD lower to 0.13 of 1 SD higher)</td>
<td></td>
<td></td>
<td>Very low&lt;sup&gt;a,c&lt;/sup&gt;</td>
<td></td>
</tr>
</tbody>
</table>

**Follow-up:** range 15 months to 27 months

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*The basis for the assumed risk is the median control group risk across studies. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).**

**CI:** confidence interval; **OR:** odds ratio; **RR:** risk ratio; **SD:** standard deviation.

**GRADE Working Group grades of evidence**
High quality: further research is very unlikely to change our confidence in the estimate of effect.
Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.
Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.
Very low quality: we are very uncertain about the estimate.

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Summary of findings 2. Summary of findings: unconditional cash transfer versus no unconditional cash transfer: secondary outcomes

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Illustrative comparative risks* (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>Number of participants (number of studies)</th>
<th>Quality of the evidence (GRADE)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Owned livestock in previous year</td>
<td>Not pooled</td>
<td>Not pooled</td>
<td>Not pooled</td>
<td>1286 (2 RCTs)</td>
<td>⊕⊝⊝⊝ Very low\textsuperscript{a,b,c}</td>
</tr>
</tbody>
</table>

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\textsuperscript{a}Serious risk of bias indicated by no allocation concealment, no blinding, potential contamination, and/or confounding (minus one grade).
\textsuperscript{b}Serious imprecision indicated by the 95% confidence interval of the estimate or estimates ranging from no meaningful change to a meaningful benefit (minus one grade).
\textsuperscript{c}Very serious imprecision indicated by the 95% confidence estimate or estimates ranging from a meaningful benefit to a meaningful harm (minus two grades).
\textsuperscript{d}Serious inconsistency indicated by 3 studies reporting meaningful or probably meaningful benefits, but one outlier reporting no evidence for an effect (minus one grade).
\textsuperscript{e}Serious inconsistency indicated by 60% > I^2 < 90% (minus one grade).
### Unconditional Cash Transfers for Reducing Poverty and Vulnerabilities: Effect on Use of Health Services and Health Outcomes in Low- and Middle-Income Countries (Review)

<table>
<thead>
<tr>
<th>Outcome Description</th>
<th>Follow-up:</th>
<th>Control</th>
<th>Intervention</th>
<th>Risk Ratio (95% CI)</th>
<th>GRADE</th>
<th>Effect on Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Attends school</strong></td>
<td>range 12 months to 24 months</td>
<td>676 per 1000 (696 to 736)</td>
<td>716 per 1000 (706 to 726)</td>
<td>RR 1.06 (1.03 to 1.09)</td>
<td>Moderate</td>
<td>Better indicated by a higher value. In conclusion, a UCT probably led to a moderate, clinically meaningful, beneficial effect on this outcome.</td>
</tr>
<tr>
<td><strong>Engages in child labour</strong></td>
<td>24 months</td>
<td>299 per 1000 (236 to 305)</td>
<td>269 per 1000 (266 to 272)</td>
<td>RR 0.90 (0.79 to 1.02)</td>
<td>Very low</td>
<td>Better indicated by a lower value. In conclusion, we are very uncertain about the effect of a UCT on this outcome.</td>
</tr>
<tr>
<td><strong>Adult works</strong></td>
<td>Follow-up: 24 months</td>
<td>798 per 1000 (758 to 838)</td>
<td>798 per 1000 (798 to 838)</td>
<td>RR 1.00 (0.95 to 1.05)</td>
<td>Very low</td>
<td>Better indicated by no change or a higher value. In conclusion, we are very uncertain about the effect of a UCT on this outcome.</td>
</tr>
<tr>
<td><strong>Parenting quality</strong></td>
<td>Assessed with: Home Observation Measurement of the Environment Score</td>
<td>Follow-up: range 15 months to 27 months</td>
<td>The mean parenting quality was 2.40 HOME Scores</td>
<td>The mean parenting quality in the intervention group was 0.22 HOME Scores higher (0.60 lower to 1.01 higher)</td>
<td>Very low</td>
<td>Better indicated by a higher value. In conclusion, we are very uncertain about the effect of a UCT on this outcome.</td>
</tr>
<tr>
<td><strong>Is extremely poor</strong></td>
<td>Follow-up: 24 months</td>
<td>812 per 1000 (722 to 812)</td>
<td>771 per 1000 (722 to 812)</td>
<td>RR 0.95 (0.89 to 1.00)</td>
<td>Very low</td>
<td>Better indicated by a lower value. In conclusion, we are very uncertain about the effect of a UCT on this outcome.</td>
</tr>
<tr>
<td><strong>Amount of money spent on health care in last month</strong></td>
<td>Assessed with: various currencies</td>
<td>Follow-up: range 7 months to 24 months</td>
<td>Not pooled</td>
<td>Not pooled</td>
<td>Low</td>
<td>Better indicated by a higher value. Because of the very high level of heterogeneity, we did not combine the studies in a meta-analysis. 4 RCTs reported that a UCT may perhaps not have had an effect on this outcome, whereas 2 RCTs reported large, likely clinically meaningful, beneficial effects on this outcome. In conclusion, a UCT may increase the amount of money spent on health care.</td>
</tr>
</tbody>
</table>

**Notes:**
- The basis for the **assumed risk** is the median control group risk across studies. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI**: confidence interval; **RR**: risk ratio; and **SD**: standard deviation.

**GRADE Working Group grades of evidence**
- **High quality**: further research is very unlikely to change our confidence in the estimate of effect.
- **Moderate quality**: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.
- **Low quality**: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.
**Very low quality**: we are very uncertain about the estimate.

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- Serious risk of bias indicated by no allocation concealment, no blinding, potential contamination, and/or confounding (minus one grade).
- Very serious inconsistency indicated by $I^2 \geq 90\%$ (minus two grades).
- Serious imprecision indicated by the 95% confidence interval of the estimate or estimates ranging from a meaningful harm to no meaningful change (minus one grade).
- Very serious imprecision indicated by the 95% confidence estimate or estimates ranging from a meaningful benefit to a meaningful harm (minus two grades).
- Serious inconsistency indicated by $60\% > I^2 < 90\%$ (minus one grade).
- Serious inconsistency indicated by different measurement and estimates across studies.

### Summary of findings 3. Summary of findings: unconditional cash transfers versus conditional cash transfers: primary outcomes

**Unconditional cash transfers versus conditional cash transfers: primary outcomes**

**Patient or population**: children (0 to 17 years) and adults (≥ 18 years) or households

**Settings**: low- and middle-income countries

**Intervention**: an unconditional cash transfer for reducing poverty and/or vulnerabilities

**Comparison**: a conditional cash transfer

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Illustrative comparative risks* (95% CI)</th>
<th>Relative effect (95% CI)</th>
<th>Number of participants (number of studies)</th>
<th>Quality of the evidence (GRADE)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Has used any health service in previous 1 to 12 months</td>
<td>The mean number of routine preventive health services visits was 1.02 (0.83 to 0.19 lower)</td>
<td>—</td>
<td>2559 (1 RCT)</td>
<td>⊕⊕⊕⊕ Very low(^a,b)</td>
<td>Better indicated by a higher value. In conclusion, we are very uncertain about the effect of a UCT on this outcome.</td>
</tr>
<tr>
<td>Is moderately stunted</td>
<td>No evidence available on this outcome</td>
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<tr>
<td>Has died</td>
<td>No evidence available on this outcome</td>
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<td></td>
</tr>
<tr>
<td>Has had any illness in previous 2 weeks to 3 months</td>
<td>440 per 1000 (418 to 550)</td>
<td><strong>RR 1.11 (0.95 to 1.25)</strong></td>
<td>3896 (1 RCT)</td>
<td>⊕⊕⊕⊕ Very low(^a,c)</td>
<td>Better indicated by a lower value. In conclusion, we are very uncertain about the effect of a UCT on this outcome.</td>
</tr>
<tr>
<td>Outcome</td>
<td>No evidence available on this outcome</td>
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<td>------------------------------------------------------------------------</td>
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<tr>
<td><strong>Has been food secure</strong></td>
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<tr>
<td><strong>Level of dietary diversity in previous week</strong></td>
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<tr>
<td>Assessed with: number of times the participant ate protein-rich food,</td>
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<tr>
<td>last week</td>
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<tr>
<td>Follow-up: 12 months</td>
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</tr>
<tr>
<td>The mean level of dietary diversity was unclear</td>
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<tr>
<td>The mean number of times ate protein-rich food in the intervention</td>
<td>3896</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>group was 0.06 lower</td>
<td>(1 RCT)</td>
<td></td>
<td></td>
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<tr>
<td>(0.55 lower to 0.44 higher)</td>
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<td></td>
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<tr>
<td>Better indicated by a higher value. In conclusion, we are very</td>
<td></td>
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<tr>
<td>uncertain about the effect of a UCT on this outcome.</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td><strong>Level of depression</strong></td>
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<td>No evidence available on this outcome</td>
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</table>

*The basis for the assumed risk is the median control group risk across studies. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI). CI: confidence interval; RR: risk ratio; and SD: standard deviation.

**GRADE Working Group grades of evidence**

- **High quality**: further research is very unlikely to change our confidence in the estimate of effect.
- **Moderate quality**: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.
- **Low quality**: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.
- **Very low quality**: we are very uncertain about the estimate.

- **a** Serious risk of bias indicated by no allocation concealment, no blinding, potential contamination, and/or confounding (minus one grade).
- **b** Very serious indirectness (minus two grades).
- **c** Very serious imprecision indicated by the 95% confidence estimate or estimates ranging from a meaningful benefit to a meaningful harm (minus two grades).
UNCONDITIONAL CASH TRANSFERS FOR REDUCING POVERTY AND VULNERABILITIES: EFFECT ON USE OF HEALTH SERVICES AND HEALTH OUTCOMES IN LOW- AND MIDDLE-INCOME COUNTRIES (Review)

Description of the condition

This review focused on the effect of unconditional cash transfers (UCTs), an increasingly prominent type of social protection intervention, on the use of health services and health outcomes in low- and middle-income countries (LMICs). More specifically, we reviewed UCTs that principally aim to reduce poverty, vulnerabilities or both. This includes universal basic income interventions, where every citizen receives an unconditional basic income (Painter 2016). For national governments, international organisations, nongovernmental organisations and civil society, both poverty and vulnerabilities in LMICs remain central concerns (Alvaredo 2013). We have already reviewed the evidence on the effect of once-off or short-term UCTs for assistance in humanitarian disasters (Pega 2015a), including those that aim to bring immediate relief before, during or in the aftermath of climatic disasters such as storms, heat waves and droughts (Pega 2015b).

Poverty

Poverty (defined here as a daily income of USD 2.00 or less) affects more than 30% of the population in a typical LMIC (Alvaredo 2013), with an estimated 1.2 billion people living in extreme poverty (daily income of USD 1.25 or less) in 2010 ( Qin 2013). While overall extreme poverty has reduced considerably over the last two decades, partially driven by rapid advances in China, it remains at problematic levels in several LMICs (Alvaredo 2013). Poverty is an important social determinant of health (CSDH 2008; McDonough 2005). It is linked to ill health and causes (or exacerbates) both environmental and other social determinants of health, such as access to clean drinking water and sanitation, as well as education, labour force participation and housing (CSDH 2008; McDonough 2005).

Vulnerabilities

Vulnerabilities commonly tackled by UCTs include being an orphan, an older person, disabled or affected by HIV ( Arnold 2011; Garcia 2012). Over 100 million children in LMICs have lost one or both of their parents to conflict, HIV or other causes (Stover 2007). Many live in poverty or have other vulnerabilities, such as having to work to secure sufficient income or living with HIV (Stover 2007). The number of older people in LMICs has steadily increased, driven by lower fertility rates and increased life expectancy. Old age is associated with multiple vulnerabilities (including poverty and disability), especially in LMICs without universal old age pensions. Living with HIV (or in a family affected by HIV) is also associated with multiple vulnerabilities, including unemployment and poverty. These diverse and interlocked circumstances are central social determinants of health in LMICs (CSDH 2008).

Description of the intervention

Social protection

Social protection is defined as “protecting individuals and households during periods when they cannot engage in gainful employment or obtain enough income to secure their livelihoods – due to unemployment, sickness, chronic ill health or disability, old age or care responsibilities” (p 16, UNRISD 2010). In what has been called the “quiet revolution”, social protection policies have increasing-ly gained prominence on development agendas around the world (p 4, Barrientos 2008). These policies comprise three types of interventions, namely labour market, social insurance and social assis-
tance interventions ( Arnold 2011). Social assistance interventions are “noncontributory transfer programs targeted in some manner to the poor and those vulnerable to poverty and shocks” to ensure an adequate standard of living (p 4, Grosh 2008). Types of social assistance interventions include cash transfers, in-kind transfers, fee waivers, subsidies and public works programmes, amongst others.

The World Health Organization (WHO) Commission on Social Determinants of Health, together with other experts, have recommend-ed specific policies promoting social protection over the life course to policymakers as effective interventions for addressing the social determinants of health (e.g. poverty and vulnerabilities) and improving individual and population health and health equity in LMICs (CSDH 2008; Marmot 2010; Marmot 2012; WHO 2011). The Commission advised “[g]overnments, where necessary with help from donors and civil society organizations, and where appropriate in collaboration with employers, build universal social protection systems and increase their generosity towards a level that is sufficient for healthy living” (p 87, CSDH 2008). Development banks such as the World Bank have also expressed the opinion that “social protection programs can have a direct positive impact on poor families as they help build human capital and productivity as a result of better health, more schooling, and greater skills” (World Bank 2012). In the Sustainable Development Agenda 2030, the United Nations’ international development framework for 2015 to 2030, the 193 member states of the United Nations pledged under target 1.3 to “implement nationally appropriate social protection systems and measures for all, including floors, and by 2030 achieve substancial coverage of the poor and the vulnerable” (p 17, UNGA 2015), adding further health sector interest in cash transfers and their ef-fects on health.

Cash transfers for reducing poverty or vulnerabilities

Cash transfers are cash payments provided by formal institutions (governmental, international or nongovernmental organisations) to selected recipients, generally for meeting their minimum consumption needs (Garcia 2012). They first gained popularity during the 1990s as interventions used by several Latin American countries to counter the negative effects of the 1980s debt crises (Arnold 2011; Garcia 2012). However, they have proliferated in many LMICs around the world, especially since the early 2000s (Arnold 2011; Garcia 2012). Today, cash transfers are common in middle-income countries and in the WHO regions of the Americas (especially Latin America) and South-East Asia, but they have only more recently been introduced in low-income countries and in the WHO African, European, Eastern Mediterranean and Western Pacific regions (Garcia 2012). The primary funding agencies and administrators of cash transfers are national governments, international organisations (often development banks) and donors, as well as nongovernmental organisations (especially in Africa) (Garcia 2012). Between 2007 and 2010, development assistance spending on cash transfers more than sextupled (from USD 23 million to USD 150 million), mostly driven by increases in dedicated donor funding (Global Humanitarian Assistance 2012). An estimated total of 800 million to 1 billion people in LMICs received a cash transfer in 2011 (Arnold 2011).

The basic economic rationale for ongoing, regular cash transfers is that they provide a minimum income over an extended period of time. Such cash transfers aim to reduce poverty or vulnerabilities and promote wealth creation by enabling recipients to build human capital (including better health), accruing savings to pur-
chase productive assets and obtaining access to loans with better conditions (Arnold 2011). Moreover, the additional income from cash transfers also prevents recipients from adverse personal or systemic income shocks and protects their standard of living by enabling them to maintain their spending on essential goods (e.g. food and medicines) and services (e.g. health services) during financially lean times, without needing to sell their assets or accrue debt (Arnold 2011). Furthermore, by providing additional income to poor or otherwise vulnerable people, cash transfers may also change opinions, attitudes and relationships among citizens and between them and their government (Arnold 2011). For example, a cash transfer may increase the economic standing (and hence social status and inclusion) of the recipient group and may influence citizens’ electoral support for the government, depending on such factors as the transfer’s social acceptability and perceived fairness (Garcia 2012). Moreover, cash transfers may reduce poverty and vulnerabilities more effectively and cost-effectively than other public sector investments (Fiszbein 2009). Compared with in-kind transfers, cash transfers maximise utility by giving recipients greater flexibility to satisfy their specific needs rather than predetermining a commodity (Fiszbein 2009); they avert the high costs of storing and transporting goods (Lagarde 2009), and they are less prone to leakage through corruption (Lagarde 2009).

Cash transfer interventions have diverse objectives, designs and methods of implementation. However, they can be classified into two broad types based on their regularity and length. The first type, which this review focuses on, are regular transfers over extended periods of time to sustainably reduce income poverty and vulnerabilities (Arnold 2011; Garcia 2012). Most of these transfers primarily aim to reduce income poverty by addressing transitory poverty over the short term and, in turn, chronic and intergenerational poverty over the long term (Arnold 2011; Garcia 2012). Some cash transfers primarily (or as a second objective beside poverty reduction) aim to reduce vulnerabilities in target populations (Arnold 2011; Garcia 2012). The second general type of cash transfer, which is outside the scope of this review, are once-off, short-term payments, provided after natural or humanitarian disasters for immediate financial relief or to incentivise desirable actions such as repatriation of refugees or reintegration of former soldiers after an armed conflict (Arnold 2011; Garcia 2012; Global Humanitarian Assistance 2012). We have already systematically reviewed the effect of UCTs for assistance in humanitarian disasters on the use of health services and health outcomes in children and adults in LMICs (Pega 2015a).

Unconditional cash transfers for reducing poverty or vulnerabilities

Cash transfers for reducing poverty or vulnerabilities can also be differentiated by their degree of conditionality into UCTs and conditional cash transfers (CCTs). UCTs have no conditions beyond a broadly defined eligibility category that defines a segment of the population, such as poor people or orphans, as eligible (i.e. based on who one is) (Garcia 2012). They therefore include universal basic income interventions, which seek to provide a basic income universally to everybody without any targeting (Painter 2016). In contrast, CCTs are provided conditional on engaging in prescribed behaviours (sometimes called co-responsibilities), such as using certain health services or attending school (i.e. based on what one does) (Garcia 2012). Most UCTs define eligibility criteria, but UCTs have no conditions or co-responsibilities attached to their receipt (Garcia 2012).

‘Fuzzy’ cash transfers do not neatly fit into the traditional classification of UCTs versus CCTs (Baird 2013). For example, some cash transfers are designed to be conditional in theory, but because non-compliance is not monitored, enforced or penalised they are un-conditional in practice. This review focuses on all cash transfers for reducing poverty or vulnerabilities that are de facto unconditional, that is, both genuine UCTs and fuzzy cash transfers that are essentially unconditional.

The underlying theory for the use of UCTs understands people living in poverty as rational actors and assumes that providing them with additional income will result in them engaging in desired behaviours, through which they will eventually graduate from poverty and overcome their vulnerabilities (Arnold 2011). This theory expects UCTs to generate similar, beneficial behaviour change to CCTs, because recipients are motivated and able to engage in the behaviours that CCTs require. UCTs could also generate greater behaviour change, because they are more socially acceptable and less stigmatising for their recipients than CCTs. In contrast, the alternative theory underpinning the application of CCTs argues that “poor households lack full information on the long-term benefits of preventive health care and education” and that conditions are required to ensure that the cash transfer generates the desired behaviours among its recipients (p 49, Arnold 2011). This theory expects CCTs to generate greater behaviour change than UCTs, because CCTs incentivise desired behaviours not only through income effects, but also through (imposed) substitution effects (Fiszbein 2009; Garcia 2012). It is sometimes also argued that conditioning cash transfers may increase their political feasibility (Garcia 2012).

Some experts have made the case for using cash transfers as policy tools specifically for addressing key social determinants of health (poverty and vulnerabilities) to improve the health of socioeconomically disadvantaged populations and, in turn, health and health equity in the population in LMICs (Forde 2012). However, the extent to which UCTs for reducing poverty and vulnerabilities also improve the use of health services and ultimately, health outcomes, is unknown.

Furthermore, the relative effectiveness and cost-effectiveness of UCTs versus CCTs for improving the use of health services and health outcomes in LMICs is unclear (Baird 2012; Gaarder 2012; Robertson 2013). Some authors have hypothesised that UCTs, under certain conditions, are more effective (Schubert 2006). The reasons are that conditioning a cash transfer results in additional direct, indirect and opportunity costs to the recipients from having to comply with the conditions, as well as additional costs to the administrator for monitoring recipients’ compliance with the conditions. Costs to recipients are often higher in people with a lower socioeconomic position, with a potential perverse effect on health equity. Furthermore, conditioning a cash transfer on the use of health services will not confer any health benefits if health services are inaccessible or of insufficient quality. In addition, if use of health services increases due to a conditional cash transfer (CCT) without adjustment on the supply side, overall quality of care may suffer. Moreover, attaching conditions to a cash transfer could increase the social stigma attached to the transfer, which could reduce its positive health effects.

On the other hand, implementing UCTs may be less politically feasible, especially in middle-income countries, because of the perception that UCTs are merely a cash giveaway to the poor and vulnerable. For example, in the Philippines, policymakers decided to condi-
tion a cash transfer after deliberately considering the transfer’s political feasibility (Friedman 2014). There could also be savings from not paying people eligible for a CCT who do not comply with the required conditions, and if these savings more than compensate for the CCT’s additional administrative costs, then this would make the CCT more cost-effective than an equivalent UCT programme (Baird 2011). Therefore, if UCTs are equally as effective as CCTs (or marginally less effective, but effective nevertheless), they may be the preferred option in LMICs (as long as their implementation is politically feasible). The reasons are that CCTs additionally require an adequate supply of services to meet the transfer conditions, potentially carrying higher costs for both the recipients and the administrator; they also require adequate compliance monitoring systems.

**How the intervention might work**

Figure 1 presents a conceptual model of the causal relationship between an unconditional cash transfer (UCT) and a health outcome. The primary causal pathway through which UCTs impact health is through income. There is some evidence suggesting that cash transfer programmes reduce the depth or severity of income poverty in children and adults in LMICs (Arnold 2011; Barrientos 2006). This reduced risk of income poverty in the recipient household may improve health outcomes all by itself. More specifically, income from publicly funded cash transfers may impact health at the individual level through five types of causal effects (Borjas 2013; Lundberg 2010; Pega 2012; Pega 2013; Pega 2015a).

Figure 1. Conceptual framework of the causal relationship between an unconditional cash transfer for reducing poverty and vulnerabilities and the use of health services and health outcomes

1. Direct consumption effects (pathway A-B-C in Figure 1).
2. Direct status effects (pathway A-D-E).
3. Combined consumption and status effects (pathway A-B-F-E).
5. Reduced financial risk (arrow I).

In direct consumption effects, income influences material conditions, which determine health through physical mechanisms (Lundberg 2010). For example, if recipients of a UCT used the additional income from the transfer to purchase goods and services that benefit their health, such as health services or nutritious food, then the UCT would be expected to improve health outcomes in the recipients. However, if recipients used the income from a UCT to purchase goods and services that damage their health, such as tobacco or alcohol, then the UCT would be expected to negatively affect health outcomes. Another consumption effect would be differential investment behaviour on the part of the household and greater diversification of economic activities into those carrying a higher risk but also higher expected returns, which may influence health outcomes.

**Figure 1**

- **Material circumstances**
  - Income
  - Psychosocial factors
  - Employment

- **Unconditional cash transfer**
  - A

- **Use of health services and health outcomes**
  - B
  - C
  - D
  - E
  - F
  - G
  - H
  - I

With direct status effects, the additional income from a UCT impacts the health of recipients through psychosocial mechanisms associated with the recipients changing their relative income position (Lundberg 2010). For example, the additional income from a UCT could increase a recipient’s income position (relative to relevant individuals or comparison groups), enhancing their social status, reducing psychosocial stress and, ultimately, improving physical and mental health outcomes.

Combined consumption and status effects impact health through both physical and psychological mechanisms, namely material conditions and, in turn, social inclusion (Lundberg 2010). For example, if recipients used the additional income from a UCT to purchase goods and services that enhanced their inclusion in a social group (e.g. club membership), then this may positively impact their health. The level to which this social group promotes health is expected to mediate the level to which the additional income from the UCT increases health. So, social inclusion in groups promoting healthy behaviours (e.g. exercising and eating nutritious food) can have more positive health effects than social inclusion in groups promoting unhealthy behaviours (e.g. tobacco and alcohol use).
Employment effects impact health by enabling people to change employment (Borjas 2013). For example, assuming that leisure time is a normal good, additional income from a UCT would be expected to reduce whether and how many hours the recipient works, which, in turn, may impact health outcomes. Alternatively, recipients of a UCT could keep working or maintain their working hours but switch to an occupation with a lower wage, which could also impact health outcomes. The level to which the UCT would be expected to increase health would depend on the level to which a reduction in employment changed health, which likely depends on such factors as the status and condition of the employment (Benach 2010a; Benach 2010b). For example, a UCT might increase health more in recipients who reduced their working hours in a job with negative or hazardous working conditions (e.g. through exposure to hazardous substances) than in employment with positive and health-promoting working conditions (e.g. through increasing the recipients’ sense of self-efficacy and self-worth).

Finally, UCTs may also directly affect health through welfare security from reduced financial risk (Pega 2012; Sjöberg 2010). Welfare security is a sense of psychological security from knowing that specific (or combinations of) cash transfers ensure income supplementation in times of financial hardship (Pega 2012; Sjöberg 2010). A recent study demonstrated that high-income countries with cash transfers for the unemployed had higher levels of employment-related welfare security and subjective well-being than high-income countries without such transfers (Sjöberg 2010).

The theory of a minimum income for healthy living hypothesis—that income over a certain threshold is a prerequisite for good health (Morris 2000; Morris 2007). While minimum income thresholds have been calculated for selected populations in some high-income countries, they have not yet been established for LMICs (Gorman 2007). A UCT would be expected to have a more beneficial health effect in recipients whose income it lifts above the minimum threshold than in recipients whose income remains below it despite the transfer.

Why it is important to do this review

This review differs from previous reviews in that it specifically investigates the impact of UCTs whose primary aim is to reduce poverty and vulnerability on the use of health services and health outcomes in LMICs. It also synthesises existing evidence on the relative effectiveness of UCTs compared with CCTs for improving the use of health services and health outcomes in LMICs. Readers interested in the health- and healthcare-related effects of UCTs in the context of humanitarian assistance are referred to the parallel Cochrane Review on the topic (Pega 2015a); a similar systematic review has also since been published (Doocy 2016). The systematic review evidence presented in this review is particularly important, considering the relatively low costs and administrative ease of implementing UCTs.

Previous reviews have synthesised evidence on the effect of CCTs for use of health services and health outcomes in LMICs (Gaarder 2010; Lagarde 2009; Owusu-Addo 2014), while other research has assessed in-work tax credits (CCTs provisional on uptake or retention of employment) for health status improvements in adults (Pega 2013). However, these four reviews did not include UCTs. Eight reviews have assessed a combination of various financial credit interventions, including potentially UCTs, for health improvements. Bocca 2012 reviewed the effect of UCTs, CCTs and microfinance interventions on risk factors for tuberculosis. Bassani 2013 reviewed the effect of UCTs, CCTs, voucher programmes and removal of user fees on the use of health services and health outcomes in children. Manley 2013 reviewed the effect of UCTs, CCTs and public works programmes on nutrition. Three reviews have evaluated the effects of UCTs and CCTs on the incidence of HIV in LMICs (Adato 2009; Heise 2013; Pettifor 2012). Finally, two non-systematic reviews have assessed the effect of UCTs and CCTs on the use of several health services and health outcomes (Arnold 2011; Sridhar 2006), and an ongoing systematic review will synthesise the evidence on the effect of cash transfer interventions on the social determinants of health in Sub-Saharan Africa (Owusu-Addo 2016). UCTs, CCTs and other financial interventions may differ in their effect on health in LMICs (Baird 2012; Robertson 2013); therefore the evidence should be reviewed separately for each of these types of interventions.

National governments, international organisations, nongovernmental organisations, and civil society require systematic review evidence on the effectiveness of different types of cash transfers in improving the use of health services and health outcomes in LMICs. This information will enable them to prioritise, plan, cost and implement the most suitable and effective cash transfer type or types. This review provides such systematic review evidence for UCTs. It also provides evidence on the relative effectiveness of UCTs compared with CCTs.

OBJECTIVES

To assess the effects of UCTs for improving health services use and health outcomes in vulnerable children and adults in LMICs. Secondary objectives are to assess the effects of UCTs on social determinants of health and healthcare expenditure and to compare to effects of UCTs versus CCTs.

METHODS

Criteria for considering studies for this review

Types of studies

Before we commenced this review, we developed a detailed protocol that laid out our eligibility criteria and methods (Pega 2014). In terms of experimental and quasi-experimental studies, this review included parallel-group and cluster-randomised controlled trials (RCTs). Quasi-RCTs (allocating participants, for example, by means of alternation or date of birth) were also eligible, but we did not identify any. In terms of observational studies, we included controlled before-and-after studies (CBAs) and cohort studies. We would have also included interrupted time series studies but did not find any that were appropriate. We included only CBAs that met the minimum methodological criteria defined in the Cochrane Effective Practice and Organisation of Care (EPOC) Group guidelines (Cochrane EPOC 2012): two or more sites in each intervention arm; intervention and control group were collected contemporaneously; and intervention and control sites were comparable (for example, we would have excluded studies that compared two urban with two rural sites). We included only cohort studies that at a minimum: had three or more repeated measurements and controlled (or attempted control) for either or both confounders (for example, through standardisation, stratification or matching) and reverse causation (for example, through marginal structural modelling (Pega 2016a)). We excluded instrumental variable analytic
studies that used UCTs as instruments to estimate the effect of income on health (Pega 2016b).

To assess the effectiveness of UCTs (primary review objective), we included studies with two types of comparators. First, we included studies comparing a group receiving a UCT with a group not receiving the UCT. Second, we included studies comparing a group receiving a UCT with a group receiving a considerably smaller income amount from the UCT. If a study compared a UCT with both comparator types, then we prioritised comparisons with the group that received no UCT over those receiving a smaller amount of the UCT. The comparison with no intervention is more consistent with the objectives of the review to evaluate intervention effectiveness, because receipt of any UCT may be more important for health effects than the amount of a UCT received (Baird 2011; Filmer 2011). Only one study compared a UCT to a less generous UCT (Haushofer 2013), but this study also compared the same UCT to no UCT, so we prioritised the latter comparison.

To assess the relative effectiveness of UCTs versus CCTs (secondary review objective), we also included studies comparing a group receiving a UCT with a group receiving a CCT in a comparable context and setting.

Types of participants
This review included both children (0 to 17 years) and adults (18 years or older) residing in an LMIC as defined by the World Bank (World Bank 2014).

Types of interventions
This review included UCTs for reducing poverty or vulnerabilities, defined as:
- an in-hand cash payment (possibly disbursed directly into a bank account, paid directly onto a mobile phone or provided in the form of a value card);
- unconditional (i.e. the cash transfer may have certain eligibility criteria but does not have any de facto conditions attached to its receipt);
- noncontributory (i.e. the cash transfer is not a payment from a social insurance system that recipients have previously contributed to);
- provided by a formal institution (national governmental, international or nongovernmental organisation) or as part of a scientific study;
- provided with the goal of reducing poverty or vulnerability (e.g. orphanhood, old age or HIV infection);
- disbursed to an individual or household (i.e. communities do not receive the cash transfers); and
- provided regularly (i.e. twice or more over a one-year period) and over extended periods of time (i.e. eligible families in theory continue receiving the cash transfer over time until they become ineligible).

We included UCTs disbursed exclusively to women and those disbursed to all genders. We included fuzzy cash transfers as long as they were de facto unconditional (Baird 2013). For the included fuzzy cash transfers, we described the contexts that produced essentially no conditions, such as lack of monitoring, enforcement or penalisation of theoretical conditions. We excluded cash transfers designed to be unconditional but with de facto conditions attached to them due to contexts, such as clear messaging that implied conditions or administrative linking of enrolment in the cash transfer to certain conditions. We also excluded UCTs for assistance in humanitarian disasters (covered in Pega 2015a) because they address different causal pathways and therefore may have a different effect on use of health services and health outcomes. If we excluded a study due to the intervention being a CCT, a fuzzy cash transfer with de facto conditions or a UCT for assistance in humanitarian disasters, then we noted this as a reason in the Characteristics of excluded studies table.

We included UCTs that were standalone interventions or had minor co-interventions, but we excluded UCTs provided in combination with or alongside major co-interventions. We judged a co-intervention as minor if we considered it to be very unlikely that the intervention could have a noteworthy impact on the outcome or outcomes included in this review, based on the best available evidence we retrieved on this co-intervention. For example, we would classify a short health educational intervention (e.g. one nutrition class) as minor, whereas a sustained, long-term nutritional education programme (e.g. eight weekly nutrition classes delivered over a period of two months) was major.

In this review, we report the amount of income from the UCT in USD. If the study record provided a UCT in a currency other than USD, we converted it to USD. To improve comparability in actual purchasing power across UCT amounts reported in this review, we adjusted for purchasing power parity. In line with economic theory, these adjustments approximate the total adjustment made on the currency exchange rate between countries that is required to allow the converted amount to have equal purchasing power in the currency across countries. Throughout the review, when we refer to amounts of UCT in USD, then these amounts were either provided as USD or provided in another currency but converted and adjusted for purchasing power parity.

Types of outcome measures
We chose outcomes to ensure comparability with the Lagarde 2009 review of the impact of CCTs on the use of health services and health outcomes in LMICs. Reporting at least one of our primary outcomes was an eligibility criterion. We excluded studies that only reported secondary outcomes. If a study reported measures for several included outcomes, then we included one measure for each of the reported outcomes in the review. If a study reported multiple measures for the same outcome, then we prioritised the most important measure, taking into consideration the need for consistency in measures across included studies. We prioritised measures that are more clinically important, such as the prevalence of a disease compared to the risk factors or behaviours for the disease. We prioritised measures that applied standard cut-offs to determine clinically relevant outcomes (e.g. moderate stunting, defined as a height for age of up to 2 standard deviations below the median (WHO 2016)) over measures of the variable from which the measure was derived (e.g. height for age), because the former are more informative for decision making. Moreover, for complex measurement concepts (e.g. dietary diversity), we prioritised established, standard composite measures (e.g. the Household Dietary Diversity Score, or HDDS (Kennedy 2011)) over measures of components of the composite index (e.g. ‘has eaten fruit’), and we prioritised these component measures over others that are less directly related to the prioritised standard composite measure (e.g. ‘level of protein intake’). We included studies reporting outcomes for any time period. If a
study reported multiple follow-up periods, then we prioritised the longest follow-up during the intervention. For example, if a study reported treatment effect estimates at 12 months and 24 months into the intervention (during) and at 8 months after a 24-month intervention, then we prioritised the follow-up at 24 months.

**Primary outcomes**

Eligible primary outcomes of the review were as follows.

- Use of health services, including but not limited to:
  - registered birth;
  - growth checks;
  - up-to-date in vaccination calendar;
  - treatment for parasites;
  - use of any health service;
- Health outcomes, including but not limited to:
  - anthropometric measures (stunting, height for age, weight for age);
  - death;
  - disease incidence or prevalence;
  - food security;
  - dietary diversity;
  - mental health outcomes.

Regarding the use of health services, the review included objective and subjective measures of the use of any health service. These measures were either administrative records or survey data of the use of health facilities or services, such as the number of routine preventive health clinic visits and the proportion of participants who were fully immunised or received parasite treatment. We considered neither the distance travelled, nor the travel time required to access the facilities or services in this review.

For health outcomes, we included both subjective measures as rated by a clinician, patient or caregiver (e.g. self-report of disease prevalence) and objective measures (e.g. clinical test for a specific disease). In the outcome domain of nutrition, for example, we prioritised standard composite indices of dietary diversity such as the HDSS (i.e. total number of food groups consumed) (Kennedy 2011) over measures of consumption of macronutrients (e.g. ate protein), and we prioritised the latter over micronutrients (e.g. intake of vitamins). We also included any potential harms that we identified. We would have included mortality, but we found no study reporting on this outcome.

**Measures of impact on equity in primary outcomes**

To measure the effect of a UCT on equity in a primary outcome, we included and prioritised direct measures of absolute or relative inequity in the primary outcome, but did not find any such prioritised measures in studies included in this review. We also included treatment effect estimates for two or more subgroups defined by population characteristics along the six PROGRESS categories (i.e. age, education, gender, rural-urban residency, income (or poverty status) and marital status), because these measures enabled us to indirectly draw conclusions on the effects of UCTs on equity in primary outcomes by these characteristics.

**Secondary outcomes**

The secondary outcomes of the review were:

- relevant social determinants of health (e.g. assets, education, labour force participation, parenting quality and extreme poverty); and
- healthcare expenditure (i.e. measures of direct and indirect costs borne by the healthcare recipient).

We defined extreme poverty according to the trial authors’ definitions.

**Search methods for identification of studies**

**Electronic searches**

**Academic databases**

Appendix 1 presents the search strategy for Ovid MEDLINE(R) 1946 to Present with Daily Updates. We developed this strategy based on the Lagarde 2009 and Pega 2013 systematic reviews of the effect of cash transfer interventions on health. We adapted the subject heading terminology and syntax of search terms to the requirements of the individual databases (Appendix 2 for the adapted search strategies). We sought records written in any language. Just before completion of the review (10 July 2017), we repeated the PubMed database search, this time for the most recent records published over the last six months (e.g. e-publications ahead of print).

We searched the following 17 databases initially in May 2015 and re-searched them in May 2017.

- Cochrane Public Health Group Specialised Register (because this registry has not been updated since 2014, we did not need to re-run the original search from 29 May 2015).
- Cochrane Central Register of Controlled Trial (CENTRAL; 2017, Issue 5) in the Cochrane Library (searched 2 May 2017).
- Ovid MEDLINE(R) 1946 to Present with Daily Update (1946 to 5 May 2017);
- Embase (1947 to 10 May 2017).
- CINAHL (1937 to 10 May 2017).
- 3IE database (1990 to 20 May 2017).
- PsycINFO (1920 to 10 May 2017).
- PubMed (1920 to 2 May 2017).
- Social Sciences Citation Index (1955 to 8 May 2017).
- The Campbell Library: the Campbell Collaboration (the Campbell Library, Volume 13; searched 20 May 2017.).
- TROPHI (1920 to 21 May 2017).
- WHOLIS (1948 to 20 May 2017).

**Grey literature databases**

We also searched the following six grey literature databases.

- EconPapers (www.econpapers.repec.org).
- ProQuest Dissertations & Theses Database.


For grey literature databases searches that returned more than 500 hits, we screened the first 100 hits only, after ordering the hits for relevance if the database permitted this.

**Internet search engines**

We screened the first 30 hits on the Internet search engines Google Scholar, Scirus and ReliefWeb.

**Targeted Internet searching of key organisational websites**

We searched the websites of the following eight key international, donor and nongovernmental organisations.

- European Bank for Reconstruction and Development (www.ebrd.com).
- Inter-American Development Bank (www.iadb.org).
- Cash Transfer Projects in Humanitarian Aid (www.sdc-cashprojects.ch).
- Save the Children (www.savethechildren.org.uk).

We did not conduct a targeted search of the WHO website because we searched WHOLIS, which comprehensively indexes publications from this organisation.

**Searching other resources**

**Previous reviews, academic journals and included records**

We handsearched for eligible studies and records:

- the eight previous reviews on cash transfers (potentially including unconditional ones) and health service use and/or health outcomes (Adato 2009; Arnold 2011; Bassani 2013; Boccia 2012; Heise 2013; Manley 2013; Pettifor 2012; Sridhar 2006);
- all issues published between May 2016 and June 2017 of the three journals with the highest number of included studies (Journal of Nutrition, Quarterly Journal of Economics, and The Lancet); and
- the reference lists of all included records.

**Expert advice**

During the synthesis stage, we sent a list of all eligible studies and records identified by our searches to the Review Advisory Group members and two additional researchers with expertise in cash transfers and health. We asked these experts to alert us to any other potentially eligible published or unpublished, completed or ongoing studies or records they knew of.

**Data collection and analysis**

**Selection of studies**

A research librarian (Dr Paul Bain) assisted the search for relevant literature in the database, which returned the titles and abstracts of each record. One author (out of: FP, SYL, SW and RP) initially screened the title and abstract of each identified record for relevance, eliminating obviously irrelevant records. We screened the full text of each record without an abstract to establish its relevance. We identified and excluded duplicate records.

At least two authors (out of: FP, SYL, SW, RP and SKL) then independently screened the abstract of each potentially relevant record in depth for eligibility. We retrieved records selected for full-text screening. We had records written in a language other than those spoken by the authors (Dutch, English, French, German, Italian and Spanish) translated into English.

Two authors (out of: FP, SYL, SW, RP and SKL) then independently established whether a record undergoing full-text screening met the inclusion criteria for the review. A third author (FP or SKL) resolved disagreements about the inclusion of controversial records. We documented the reasons for excluding the 30 studies that were closest to the inclusion criteria in the ‘Characteristics of excluded studies’ table.

**Data extraction and management**

Two data extractors (out of: FP, Carolin Henning and Tatjana Paec) independently extracted data for each included study, using the Cochrane Public Health Group’s data extraction form (Cochrane PHG 2011), expanded for the complex intervention perspective that we adopt in this review, with the Cochrane-Campbell Methods Group Equity Checklist added (Jeffing 2012). To ensure standardised data extraction, the data extractors first received training in data extraction, and they then piloted the dedicated form before commencing the extraction. One review author checked and resolved discrepancies between the data extraction forms of the two data extractors (FP or SKL), and a second author independently double-checked the extracted data (out of: SYL, SW, RP, RS and SKL).

At a minimum, we extracted data for the following categories: study eligibility (i.e. data required to assess eligibility along inclusion criteria); study details (including study objectives and methods); intervention groups (including group names and, for cluster-RCTs, all intervention arms); outcomes; and results (including for subgroups).

Where information was available from the record on the context, implementation, cost and sustainability of the UCT, we also extracted this information. Where this information was not available directly from the record, but where the record cited another source that described it, we extracted the data from this other source. The types of contextual information we extracted included design features of the UCT such as its generosity (e.g. as assessed by the percentage contribution of an average income from the UCT to the national average total income) and population coverage (e.g. as measured by the coverage rate of the UCT amongst the total population). We reported this information on the context, implementation, cost and sustainability of the UCT in the tables of ‘Characteristics of included studies’.
We also extracted data on key sociodemographic characteristics of participants at baseline and at the endpoint within the PROGRESS framework (Cochrane PHG 2011), for the purpose of assessing the interventions’ equity impact. The extracted sociodemographic characteristics included education, ethnicity, gender, gender identity, geographic residency, labour force participation, place of residency, sexual orientation, socioeconomic status, social status and religious affiliation. As noted above, we additionally incorporated the Cochrane-Campbell Methods Group Equity Checklist in our data extraction form (Jeffing 2012). We also recorded whether the intervention comprised dedicated strategies to support disadvantaged populations.

We extracted information on the comparator (i.e. definitions of the control group), again including contextual, implementation, cost and sustainability data. We extracted data on potential measured confounders and the methods for confounder control. We used Review Manager 5 (RevMan 5) software to enter, store and manage the extracted data (RevMan 2014).

Assessment of risk of bias in included studies

Two out of all authors independently assessed the risk of bias in the included studies. Where differences arose, a third author (generally FP) resolved these discrepancies.

To assess the risk of bias in the included cluster-RCTs, we applied the Cochrane ‘Risk of bias’ tool, including any special statistical considerations for this study design, such as risk of recruitment bias (Chapter 16.3, Higgins 2011b). To assess the risk of bias in the included CBAs, we used the EPOC ‘Risk of bias’ criteria (Cochrane EPOC 2012), adding an item assessing the risk of bias from confounding and reverse causation.

No credible, standardised tool for assessing the risk of bias in cohort studies currently exists (Sanderson 2007). However, as we have done previously (Pega 2013), we followed the best practice recommendation to assess the specific features of cohort studies and the extent to which these may introduce bias (Centre for Reviews and Dissemination 2009; Appendix 3 in Joyce 2010). At minimum, we assessed the risk of bias in the following features: sampling strategy; sample representativeness; participant allocation; initial survey response; attrition; exposure measurement; outcome measurement; missing data; reporting; and control of key confounders and of reverse causation.

We assessed and reported risk of bias at the outcome level, first for each outcome for each study (i.e. risk of bias of an individual study) and then for each outcome across all studies (i.e. risk of bias in the whole body of evidence).

Measures of treatment effect

For dichotomous outcomes

The included studies estimated treatment effects on dichotomous outcomes with an odds ratio (OR) or a coefficient from either a logistic regression model (i.e. an estimate of the log OR), a probit regression model (i.e. an estimate of the difference in log odds) or a difference-in-differences (DD) model.

In their calculation of treatment effect estimates, several included studies erroneously treated dichotomous data as if they were continuous data. For example, data from the question 'Have you had a growth check in the last six months?' with the two response categories 'yes' and 'no' are dichotomous, so treating the variable ‘percentage of participants who have had a growth check’ as continuous in a linear regression model is erroneous because it is based on the assumption that the variable is normally distributed. Cochrane does not accept these erroneous treatment effect estimates, and we therefore could not report these estimates in this review.

Coefficients of a DD model were the most commonly reported treatment effect estimate for dichotomous outcomes in the several cluster-RCTs included in this review. These treatment effect estimates were generally derived by first subtracting the proportion of participants in the intervention group who had the outcome (i.e. had received a growth check) before the intervention was implemented (e.g. at the baseline survey) from the proportion of participants in the intervention group who had the outcome after the intervention was implemented (e.g. at the prioritised follow up survey). In a second step, this before-and-after difference in the intervention group was subtracted from the equivalent before-and-after difference in the control group to adjust for underlying trends in the outcome. In addition, most DD estimators were also adjusted for potential confounders using regression analyses. These DD estimate can be interpreted as the average difference in the outcome in the intervention group from before and after the intervention, adjusted for underlying time trends in the outcome that occurred in the control group and adjusted for confounders. However, these DD estimates, which are common in economic research and increasingly present in epidemiological studies (Dimick 2014), are not preferred treatment effect estimates for Cochrane Reviews.

In this review, if possible we converted an odds ratio (OR) or coefficient from a logistic or probit regression model into a risk ratio (RR) estimate. If we were unable to convert an OR or a coefficient from a logistic or a probit regression model into an RR (i.e. where we could not retrieve the baseline risk in the control group before treatment with a UCT), we reported the OR that was provided in the study record or the OR that we calculated from the coefficients reported in the study record. If we could not retrieve the baseline risk from the same study but were able to retrieve a baseline risk for the outcome from another study from the same setting and context, then we used this baseline risk for our conversion and reported the source of the assumed baseline risk.

If a cluster-RCT reported a DD estimate only for a dichotomous outcome, as was common for econometric studies included in this review, and if we were able to retrieve the crude frequency measures for the outcome in the treatment and control groups from the study record or the principal study author, then we converted these crude frequencies into an RR. We calculated this RR using an approximately correct analysis for cluster-RCTs, as recommended in the Cochrane Handbook for Systematic Reviews of Interventions (Chapter 16.3, Higgins 2011b). In more detail, we calculated the effective sample sizes from: the crude frequencies of the outcome; the number of clusters in the cluster-RCT; and an intra-cluster correlation coefficient (ICC). We sourced the ICC from the only included study that reported such coefficients (Robertson 2012), and we used the median ICC across all included outcomes (i.e. ICC = 0.07). We calculated the RR by entering the effective sample sizes that we had calculated into analyses in RevMan 5 (RevMan 2014). If we were not able to calculate an RR for a study, we reported in the review that we were unable to extract or calculate an acceptable treatment effect estimate, and we did not report any treatment effect estimate for the outcome from that study.
Mean differences (MDs) of proportions, which Cochrane also does not accept, were reported in one included study, that is Baird 2010. For these measures, we sought and were granted access to the original micro-data for this study, and we re-analysed these data. Because the included outcomes from Baird 2010 were measured at three time points for each individual, nested within enumeration areas, we used a three-level multi-level model to estimate the effect of the UCT among participants in the UCT intervention group, compared with the comparator (i.e. the control group or the CCT group). Multilevel models are a generalisation of the linear model used in traditional regression analysis (Diez-Roux 2000; Raudenbush 2001). Several authors have shown that ignoring the hierarchical structure of a data set can lead to inferential errors and that estimating random-effects coefficients can more adequately model data structures typically obtained in field research (Diez-Roux 2000; Raudenbush 2001). We performed the analyses using HLM7.01 and Stata (Scientific Software International 2015; StataCorp 2015). To investigate the potential effect of exposure to the UCT treatment on the likelihood of the outcome, we adopted a step-up approach (Raudenbush 2002), conducting different sets of analyses. The first set of analyses investigated the crude relationship of the UCT in comparison to the control group and the likelihood of experiencing the outcome. We then added sociodemographic variables because they could potentially act as confounders of the relationship between the main exposure and outcome. The covariates added in the multilevel model were the same ones we adjusted for in the original analysis presented in the study record (i.e. student’s age, whether the father lived within the household, whether the girl previously had sex, and time point of data collection).

For continuous outcomes

All included studies reported a treatment effect on a continuous outcome variable as a mean difference (MD) between the intervention group and the control group or as the coefficient of a DD regression model. As with dichotomous outcomes, DD estimates were the before-and-after difference in the intervention group minus the before-and-after difference in the control group, and they can be interpreted as the average difference in the outcome in the intervention group from before and after the intervention, adjusted for underlying time trends in the outcome that occurred in the control group (see above). In this review, we reported the MD or DD estimates for studies with continuous outcomes. Several included studies reported MDs and DDs that were z-transformed (i.e., standardised by being divided by 1 standard deviation (SD)), but we did not consider these measures to be equivalent to what is referred to as standardised MDs in Cochrane, and therefore we report these treatment effect measures as MDs of 1 SD and DDs of 1 SD, respectively.

Prioritisation of treatment effect estimates

If two or more studies used the same data and outcome (for example, where two studies evaluated the same government programme), we prioritised for inclusion in the meta-analysis the study with the study design that carried a relatively lower risk of bias. If for an included outcome a study presented both a treatment effect estimate that was unadjusted for confounding and one that was adjusted for confounding, then we prioritised and reported the adjusted treatment effect estimate. If a study had presented only unadjusted treatment effect estimates, we would have adjusted the treatment effect measures for these variables as long as between-group differences in covariates at baseline and potential confounding variables were reported; however, this situation did not occur in this review. If a study reported multiple models, each of which adjusted for a different number or set of potential confounders, then we prioritised the model that we judged to have adjusted most appropriately for the largest number and most relevant set of potential confounders.

In econometric studies, authors routinely present several competing additional specifications of a main regression model as robustness checks. In this review, we prioritised the treatment effect estimate from the conservative or ‘baseline’ model that we judged to be most appropriately and fully adjusted. For example, if a study reported an unadjusted regression model (i.e. the baseline model), the same model with stronger methods of confounder control (i.e. more appropriately adjusted baseline model) and an alternative model that used an alternative exposure variable (i.e. a robustness check), then we prioritised the adjusted regression model.

If a study presented an intention-to-treat and another (e.g. average causal) treatment effect estimate, then we reported the intention-to-treat estimate. Related to this, we prioritised estimates of the effect of being eligible for or receiving a UCT (i.e. a ‘yes’ versus ‘no’ dichotomous exposure variable) over estimates of the effect of the specific dollar amount of the UCT that the recipient was eligible for or received (i.e. a continuous exposure variable). The reason is that the latter effect estimates carry a lower risk of certain biases. For example, violations of consistency in estimates of average treatment effects could occur whereby the dollar amount of the UCT is not irrelevant for treatment (VanderWeele 2009); for instance, USD 10 provided to a participant with an annual income of USD 15,000 is not equivalent to USD 10 provided to a participant with an annual income of USD 50,000.

We reported the 95% confidence intervals (CI) for each treatment effect measures, if feasible. If the study record(s) did not provide the 95% CI or the data required to calculate it (e.g. a standard error or a t-value), we requested either the 95% CI or the data to calculate it from the principal study author via email. If we could not retrieve the 95% CI or the data required to calculate it, then we reported in the review the information about the statistical significance that the study record provided (e.g. an exact P value or the reported P value threshold).

In this review we report several treatment effect estimates and/or their standard deviations (SDs) that differ from those reported in the included study records, generally because the previously published estimates were unadjusted for clustering in cluster-RCTs (see Unit of analysis issues). We also report several treatment effect estimates and/or their SDs that have not been reported in study records. We have retrieved these new estimates and/or SDs directly from the study authors (see Dealing with missing data).

Unit of analysis issues

We screened all studies for unit of analysis issues from randomisation (or non-random allocation) of participant clusters, treatment with multiple interventions, and multiple observations for the same outcome at different time points. If a study that randomised (or observed) participant clusters did not control for clustering effects in the analysis, we contacted the principal study author and requested treatment effects estimates and 95% CIs (or the stan-
Dealing with missing data

We requested all relevant missing information on the study methods, outcomes, and statistical measures required for this review from the principal study authors by email (using the contact details provided in the latest eligible study record or requesting current email addresses from the authors’ affiliated organisations). If a principal study author did not respond within a 14-day period, we contacted second or last study authors by email.

For all included studies, we requested detailed information on the following data if missing:

- Assumed risks (i.e. baseline risk in the control group).
- Numbers of participants.
- Standard deviations of continuous outcomes to be able to standardise treatment effect estimates.
- Treatment effect estimates acceptable to Cochrane (i.e. an OR or an RR for a dichotomous outcome and an MD for a continuous outcome) and fully adjusted for confounding.
- Standard errors that were fully adjusted for confounding and, if necessary, for unit of analysis issues (i.e. clustering).

We received the requested information, including the missing data, for the Baird 2010, Bazzi 2012, Bunha 2014, Fernald 2011, Galiani 2014, Leroy 2010, Luseno 2012, Miller 2008, Oxford Policy Management 2012, Pellerano 2014, Robertson 2012, Schady 2012, Seidenfeld 2013, and Ward 2010 studies. If we could not obtain missing information and data, we analysed only the available data and addressed the potential impact of the missing information and data on the findings of the review in the Discussion section.

Assessment of heterogeneity

We did not meta-analyse studies that differed considerably in their study designs (e.g. we did not combine a cluster-RCT with a CBA or a cohort study), outcomes (e.g. we did not combine a Center for Epidemiologic Studies Depression Score measure with a Geriatric Depression Scale measure) or participants (e.g. we did not combine individual participants with households), but otherwise we considered the included studies sufficiently homogenous across participants and interventions (including intervention design, context, and implementation, including the reporting period and the follow-up period) to potentially be combined in the same meta-analysis. For studies with the same outcome and study design, we calculated the I² statistic using RevMan 5 (RevMan 2014) to assess their statistical heterogeneity for the purpose of more formally establishing the feasibility of meta-analysis.

Assessment of reporting biases

Publication bias could have occurred if we failed to comprehensively identify all studies that were eligible for inclusion. For example, studies with unwelcome or null findings may not have progressed to publication in the academic literature and may therefore not have been indexed in the databases that we searched. To avoid missing eligible studies we employed a comprehensive search strategy. Moreover, in addition to several academic databases, we also searched the Cochrane Central Register of Controlled Trials (CENTRAL) and the Cochrane Public Health Group Specialised Register; several databases of grey literature, dissertations, theses, and conference proceedings; and the websites of seven key organisations. Additionally, we asked independent policy and research experts, including the Review Advisory Board, to identify unpublished studies. We found and included in the review many eligible studies published in non-academic, grey literature. Furthermore, the review also included articles written in any language to minimise the likelihood of language bias. Since the review did not identify 10 or more eligible studies reporting the same outcome, we did not produce a funnel plot and did not test for funnel plot asymmetry to assess the presence of publication bias for the outcome.

Data synthesis

Meta-analysis

We combined studies that we considered sufficiently homogenous across study design (including treatment effect estimate), intervention, outcome and participants in a meta-analyses using RevMan 2014. We combined only studies with the same study design. For example, we combined two or more cluster-RCTs but did not combine a cluster-RCT with another study design, such as a CBA or a cohort study. Similarly, we did not combine studies with different types of treatment effect estimates (e.g. we did not combine an RR with an OR or an MD or DD with a standardised MD). We pooled only the same type of treatment effect (e.g. RRs only), whether or not they were crude or adjusted for the same or different confounders.

For dichotomous outcomes, we did not combine RRs and ORs in the same meta-analysis. Rather, if feasible we converted ORs into RRs and then combined these converted RRs with the RRs extracted or calculated from other studies. If we were unable to convert ORs into RRs for the same dichotomous outcome for several studies, then we combined the ORs in the meta-analysis, and then converted the overall OR from the meta-analysis into an RR, if possible. For continuous outcomes, we assumed MDs and DAs to be sufficiently comparable to be combined, and we therefore combined MDs with DAs in the same meta-analysis.
We combined only studies that reported the same outcome in meta-analyses. If studies measured slightly different aspects of the same outcomes or measured the same outcome over slightly different reporting periods, we combined them in meta-analysis and noted major differences when we reported the results of the meta-analysis in the Effects of interventions section. We only combined all relevant studies of individual participants or of households with each other, and we did not combine individuals with households in the same meta-analysis. If a meta-analysis of individuals included both children and adults and if the effectiveness of the studied UCT was qualitatively different for children and for adults (e.g. for the outcome 'participation in the labour force', an increase in children engaging in child labour from a UCT would be a harm, whereas an increase in adults working from a UCT would be a benefit), then we displayed them as separate subgroups in the meta-analysis and did not report overall totals.

If a study reported treatment effect estimates for an outcome separately for different subsamples (e.g. one estimate for children aged up to 5 years and another estimate for children aged 6 to 17 years), and if these subgroup comparisons did not use the same comparison groups (e.g. treated young children were compared with untreated young children, and treated older children were compared with untreated older children), then we combined the treatment effect estimates for the subsamples in the same meta-analysis and defined the different subsamples when we reported the results of the meta-analysis in the Effects of interventions section.

If we combined crude frequencies in a meta-analysis to produce an RR for a dichotomous outcome (i.e. when we conducted approximately correct analyses of cluster-RCTs according to Chapter 16.3 of Higgins 2011b), we applied the Mantel-Haenszel method with random-effects models to address potential heterogeneity. In meta-analyses of dichotomous data with RRs and in meta-analyses of continuous outcomes with MD or DD effect estimates, we used the inverse variance method with random-effects models. We did not adjust any treatment effect estimate that we report in this review in any way.

We present each meta-analysis in a forest plot. For each study included in a meta-analysis, the forest plot presents the number of participants in the intervention group and the control group. If a study reported a different number of participants for a measure taken before the intervention was conducted than for the measure taken after the intervention had been provided, then we prioritised and report in the forest plot the numbers of participants measured after the intervention. If a study did not report the number of participants separately for the intervention group and the control group but only reported the total number of participants, then we reported the number of participants in the forest plot as if the total number of participants were equally split between the intervention and control groups.

If a meta-analysis was very highly statistically heterogenous (i.e. had an I² of 90% or higher), we turned totals in the meta-analysis off in the forest plots and instead synthesised the studies narratively, as recommended in Chapter 9.5 of the Cochrane Handbook for Systematic Reviews of Interventions (Deeks 2011).

**Narrative synthesis**

If we could not meta-analyse studies that reported the same outcome due to them using a different study design (e.g. cluster-RCT versus CBA) or them missing required statistical data (e.g. the standard error or data to calculate it), we reported these studies narratively, sometimes alongside the results from the meta-analysis. We narratively synthesised the results of studies that we judged to be too heterogeneous to permit meta-analysis (i.e. studies with considerably different study designs, interventions, outcomes, and/or participants, or those that had an I² of 90% or higher), reporting results separately for each outcome. If we could meta-analyse an outcome for some studies but could not include other studies of the same outcome in the meta-analysis, then we reported the results of the studies that could not be included in the meta-analysis alongside the results from the meta-analysis. To avoid introducing bias, we did not emphasise any one study in the review.

**Subgroup analysis and investigation of heterogeneity**

We planned to conduct subgroup analyses on the meta-analyses or narrative syntheses of the primary outcomes by age (comparing children with adults), gender (comparing girls or women with boys or men) and WHO region (comparing Africa, the Americas and South-East Asia). However, these subgroup analyses were infeasible because the review included an insufficient number of studies reporting intervention effects on primary outcomes among groups defined by these variables. If subgroup analyses in meta-analyses had included a sufficiently large number of studies to conduct meaningful statistical testing, we would have conducted t-tests and used the I² statistic to assess statistical significance using RevMan 2014.

**Sensitivity analysis**

We did not conduct any sensitivity analyses. The studies that we combined in meta-analyses were relatively methodologically homogenous and were generally of comparable quality, so there was no need to conduct sensitivity analyses to evaluate whether the sizes of the combined effect estimates were robust across studies. For the main comparison of UCTs with either no UCT or a UCT that provided a considerably smaller amount of income, all studies that we actually included in the review compared UCTs with no UCT. Consequently, there was also no need for sensitivity analyses to test for the effect of combining studies with both no UCT and a smaller UCT in meta-analysis.

**Summary of findings tables**

We assessed the quality of the evidence for each outcome. In following the Cochrane Public Health Group’s best practice guidelines, we applied the GRADE considerations, assessing quality based on study limitations, consistency of effect estimates, imprecision, indirectness, publication bias and strength of effect (Cochrane PHG 2011). We produced evidence profiles in the GRADE profiler Guideline Development Tool software for our GRADE assessments (GRADE Working Group 2015).

We presented results for the key measure of the seven most relevant primary outcomes of the review (i.e. use of health services and health outcomes) for the comparison of UCTs with no intervention in the main 'Summary of findings' table. In selecting the most important primary outcomes for presentation, we sought to ensure a range of outcomes covering the seven domains commonly regarded as central for improvements from UCTs: health services use, stunting, death, disease prevalence, food security, nutritional diversity, and mental health. Additionally, we also presented secondary outcomes measures (i.e. social determinants of health

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Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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and healthcare expenditure) and the relative effectiveness of UCTs compared with CCTs in additional, secondary 'Summary of findings' tables. These tables presented the number of included studies, the treatment effect estimate, and a GRADE assessment of the overall quality of the body of evidence for each outcome. We also developed the 'Summary of findings' tables with the GRADEprofiler Guideline Development Tool software (GRADE Working Group 2015).

**RESULTS**

**Description of studies**

**Results of the search**

Figure 2 presents a PRISMA flowchart of the study selection. Overall, our searches identified a total of 43,114 records. Of these, a total of 21 studies with 56 records met the inclusion criteria for the review.
Figure 2. Flowchart of study selection. Footnotes:

a Cochrane Public Health Group Specialised Register (N = 37); CENTRAL (N = 107); Ovid MEDLINE(R) (N = 6218); Embase (N = 9023); Academic Search Premier (N = 3687); Business Source Complete (N = 2430); CINAHL (N = 1255); EconLit (N = 1874); 3IE database (N = 16); PsychInfo (N = 1956); PubMed (excluding MEDLINE(R) records) (N = 1215); Scopus (N = 844); Social Science Citation Index (N = 3871); Sociological Abstracts (N = 2552); The Campbell Library (N = 107); TRoPHI (N = 33); WHOLIS (N = 6); Ovid MEDLINE(R) (N = 6218); Embase (N = 9023); Academic Search Premier (N = 3687); Business Source Complete (N = 2430); CINAHL (N = 1255); EconLit (N = 1874); 3IE database (N = 16); PsychInfo (N = 1956); PubMed (excluding MEDLINE(R) records) (N = 1215); Scopus (N = 844); Social Science Citation Index (N = 3871); Sociological Abstracts (N = 2552); The Campbell Library (N = 107); TRoPHI (N = 33); WHOLIS (N = 6).

b Grey literature databases (N = 863): ProQuest Dissertations & Theses Database (n = 87); Open-Grey (n = 357); OpenDOAR (n = 100); EconPapers (n = 100); Social Science Research Network eLibrary (n = 119) and National Bureau of Economic Research (n = 100).

c Google Scholar (N = 30).

d Organisational websites (N = 2359): African Development Bank (n = 838), Asian Development Bank (n = 197), European Bank for Reconstruction and Development (n = 88), Inter-American Development Bank (n = 191), World Bank (n = 527), and United Kingdom Department for International Development (n = 453), Cash Transfer Projects in Humanitarian Aid (n = 29), Save the Children (n = 36).

e Handsearching (N = 3752): Journal of Nutrition (n = 307), Quarterly Journal of Economics (n = 40), The Lancet (n = 1070), references of included studies (n = 1783), references of 8 previous reviews (n = 552).
Searching the 17 electronic academic databases identified a total of 36,110 records. After removing duplicates, 30,453 unique records remained. In-depth, full-text screening identified 15 studies with 21 records that fulfilled the inclusion criteria (Amarante 2011; Baird 2010, 3 records; Beck 2015; Cunha 2014; Fernald 2011; Galiani 2014; Haushofer 2013; Leroy 2010, 2 records; Luseno 2012; Miller 2008; Paxson 2007; Robertson 2012, 3 records; Salinas-Rodríguez 2014; Schady 2012; Seidenfeld 2013, 2 records).

Searching other sources yielded a total of 7004 additional records, namely 3252 records from additional database and Internet searches and 3752 records from handsearching. Of the 3252 records from additional database and Internet searches, 863 records came from the six electronic grey literature databases, 30 records came from the one Internet search machine, and 2359 records originated from the websites of eight key organisations. Full-text screening identified six additional eligible studies with 14 records (Agüero 2007, 2 records; Akresh 2012, 5 records; Bazzi 2012; Oxford Policy Management 2012, 2 records; Pellerano 2014, 2 records; Ward 2010, 2 records). It also identified 20 additional records of nine previously identified studies (Amarante 2011, 2 records; Baird 2010, 4 records; Galiani 2014; Luseno 2012; Miller 2008, 2 records; Paxson 2007; Robertson 2012; Schady 2012; Seidenfeld 2013, 7 records). We also found three ongoing studies (Galárraga 2014; O’Leary 2011; Oxford Policy Management 2013).

Fernald 2011 and Paxson 2007 analysed the same cluster-RCT, and when both studies reported the same outcome, we prioritised Paxson 2007 because it reported treatment effect estimates for the entire study sample, whereas Fernald 2011 reported results for a only a selection. Luseno 2012 and Miller 2008 analysed the same cluster-RCT but reported different outcomes, so we report analyses from both studies in this review.

Of the 3752 records from handsearches, 552 records came from the eight relevant previous reviews on cash transfers (potentially including UCTs) and health service use and/or health outcomes (Adato 2009; Arnold 2011; Bassani 2013; Boccia 2012; Heise 2013; Manley 2013; Pettifor 2012; Sridhar 2006), 1417 records came from all issues published over the year prior to finalising the review (May 2016 to June 2017) in the three academic journals with the largest number of records of included studies (Journal of Nutrition, Quarterly Journal of Economics and The Lancet) and 1783 records came from the reference lists of all included records. These handsearches identified no additional eligible study or record.

In the last search for this systematic review in May 2017, we identified 14 additional recently published or recently indexed studies, which may or may not fulfil the inclusion criteria of this review (Abdoulayi 2014; AIR 2014; Benedetti 2016; Brugh 2016; Cluver 2013; Davis 2016; Gangopahayay 2015; Grellety 2017; Handa 2014a; Hjelm 2017; Kilburn 2016; Lawlor 2015; Olajide 2016; Tiwari 2016). We describe the characteristics of these studies in the Studies awaiting classification table.

The experts we consulted did not identify any additional eligible study or record. Finally, searching the PubMed database for the most recent publications over the last six months near the end of the review identified no additional study or record that was published online ahead of print.

Included studies

We describe the characteristics of the included studies in the Characteristics of included studies table.

Type of study

Of the 21 studies included in this review, 16 were cluster-RCTs (Akresh 2012; Baird 2010; Beck 2015; Cunha 2014; Fernald 2011; Haushofer 2013; Leroy 2010; Luseno 2012; Miller 2008; Oxford Policy Management 2012; Paxson 2007; Pellerano 2014; Robertson 2012; Schady 2012; Seidenfeld 2013; Ward 2010), 4 were CBAs (Amarante 2011; Bazzi 2012; Galiani 2014; Salinas-Rodríguez 2014), and 1 was a cohort study (Agüero 2007). Cluster-RCTs were so common because – as some authors noted – by selecting clusters of individuals rather than individuals, there is less risk of bias from contamination. For each cluster-RCT included in this review, we report the number of clusters and the type of cluster that were randomised to the intervention and control groups in the Characteristics of included studies. Most included cluster-RCTs analysed data from their baseline survey and either one or two follow-up surveys.

Half (8 out of 16) of the included cluster-RCTs derived treatment effects using DD methods (Cunha 2014; Leroy 2010; Miller 2008; Oxford Policy Management 2012; Paxson 2007; Pellerano 2014; Seidenfeld 2013; Ward 2010). Difference-in-differences methods are common econometric methods for assessing the effect of a treatment on an outcome (Wooldridge 2010). In essence, as used in the included studies, they derive a treatment effect estimate by subtracting the before-and-after difference of the intervention group from that of the control group, thereby adjusting for underlying time trends of the outcome and for potential confounding that may have occurred despite random assignment or due to errors in random assignment of the intervention or interventions (see also Measures of treatment effect). The other half of the included cluster-RCTs derived treatment effects using regression analytic methods to control for potential confounding (Akresh 2012; Baird 2010; Beck 2015; Fernald 2011; Haushofer 2013, Luseno 2012; Robertson 2012; Schady 2012). All four CBAs used DD methods to estimate treatment effects (Amarante 2011; Bazzi 2012; Galiani 2014; Salinas-Rodríguez 2014). As is common in econometric studies, Amarante 2011 also used additional methods such as discontinuity regression analytic methods to derive alternative treatment effect estimates to check for robustness of results across methods. The cohort study used regression analysis to derive treatment effects (Agüero 2007). Most studies conducted intention-to-treat analyses by using eligibility for the UCT, as opposed to receipt of the UCT, as the exposure.

Participants

Overall, the included studies involved 1,092,877 participants (36,068 children and 1,056,809 adults) and 31,865 households in Africa, the Americas, and South-East Asia. Just over half of the included studies (11 out of 21) estimated the effect of a UCT on primary outcomes among children (Agüero 2007; Akresh 2012; Amarante 2011; Baird 2010; Cunha 2014; Fernald 2011; Luseno 2012; Paxson 2007; Pellerano 2014; Seidenfeld 2013; Ward 2010). In terms of age groups, almost all of these studies focused on children aged under (or just over) five years. The exceptions were Luseno 2012, examining children aged 6 to 17 years; Ward 2010, examining children aged 0 to 17 years; and Baird 2010, studying children or young adults aged 13 to 23 years. Just over a third of studies (8 out of 21)
examined treatment effects in either working-age adults (6 studies: Amarante 2011; Leroy 2010; Oxford Policy Management 2012; Paxson 2007; Schady 2012; Seidenfeld 2013) or older adults (2 studies: Galiani 2014; Salinas-Rodríguez 2014). And one third of the included studies (7 out of 21) examined households, either solely (Beck 2015; Haushofer 2013), or in addition to studying individual participants (Leroy 2010; Miller 2008; Oxford Policy Management 2012; Pellerano 2014; Ward 2010).

Most studies with individual participants (14 out of 17) included participants of both sexes (Agüero 2007; Akresh 2012; Beck 2015; Bazzi 2012; Cunha 2014; Fernald 2011; Galiani 2014; Luseno 2012; Oxford Policy Management 2012; Paxson 2007; Robertson 2012; Salinas-Rodríguez 2014; Seidenfeld 2013; Ward 2010). The other three studies exclusively examined either girls and young women (Baird 2010), or all women (Leroy 2010; Schady 2012). Two studies exclusively examined participants living in extreme poverty (Luseno 2012; Miller 2008), and one study involved only participants living below or just above the poverty line (Bazzi 2012).

About half of the included studies (11 out of 21) examined participants in countries of the WHO Africa region (predominantly Kenya and Malawi) (Agüero 2007; Akresh 2012; Baird 2010; Haushofer 2013; Luseno 2012; Miller 2008; Oxford Policy Management 2012; Pellerano 2014; Robertson 2012; Seidenfeld 2013; Ward 2010). Seven studies were located in Latin America (predominantly Ecuador and Mexico) (Amarante 2011; Cunha 2014; Fernald 2011; Galiani 2014; Leroy 2010; Paxson 2007; Schady 2012), and two studies took place in South-East Asia (India and Indonesia) (Bazzi 2012; Beck 2015).

Interventions
The review included 17 different UCTs, including one basic income intervention (Beck 2015). Nine UCTs were established government programmes, while four each were either pilot government programmes or experiments.

The government programmes were:
- Ecuador’s Bono de Desarrollo Humano (three studies: Fernald 2011; Paxson 2007; Schady 2012);
- Indonesia’s Direct Cash Transfer Program (Bazzi 2012);
- Lesotho’s Child Grants Programme (Pellerano 2014);
- Mexico’s Programa de Apoyo Alimentario (two studies: Cunha 2014; Leroy 2010);
- Mexico’s Programa de Atención a Adultos Mayores en Zonas Rurales (Galiani 2014);
- Mexico’s 70 y Más (Salinas-Rodríguez 2014);
- South Africa’s Child Support Grant (Agüero 2007);
- Uruguay’s Plan de Atención Nacional a la Emergencia Social (Amarante 2011); and
- Zambia’s Child Grant Program (Seidenfeld 2013).

The four pilot government programmes were:
- Burkina Faso’s Nahouri Cash Transfer Pilot Project (Akresh 2012);
- Kenya’s Hunger Safety Net Pilot Programme (Oxford Policy Management 2012);
- Kenya’s Cash Transfer Pilot Programme for Orphans and Vulnerable Children (Ward 2010); and
- Malawi’s Social Cash Transfer Pilot Scheme (Luseno 2012; Miller 2008).

The UCT experiments were conducted in:
- India, by a nongovernmental organisation (Beck 2015);
- Kenya, by a nongovernmental organisation (Haushofer 2013);
- Malawi, by a research organisation and an international organisation (Baird 2010); and
- Zimbabwe, by research organisations (Robertson 2012).

The duration of the interventions was most commonly 12 to 24 months, but studies collected outcomes at time points ranging from 7 months into the intervention in Haushofer 2013 to 57 months into the intervention in Schady 2012. The follow-up in most studies was undertaken during and at the end of the intervention, at 12 to 24 months. In some cases, investigators assessed persistence of effects with follow-up surveys after the intervention had ended (e.g. eight months after the intervention was completed in the Akresh 2012 study). However, as noted above, we prioritise the longest follow-up during the intervention in this review.

Some UCTs primarily aimed to reduce poverty and some vulnerabilities (generally by improving one or more of health, nutrition, food security and education), but most combined both of these objectives. Most UCTs were targeted to individuals, families or households living in poverty or at risk of it. Governments or communities generally applied targeting through various indicators (e.g. income poverty or residency in a low-income area) and using various mechanisms (including official surveying or selection through community committees). The amounts of cash transferred varied between 1.3% and 53.9% of the annual gross domestic product per capita. These total amounts were disbursed in regular payments made every month or every second month (except for every third month in Akresh 2012).

Two of the included interventions were fuzzy in that they had conditions attached to them in theory, but because programme administrators did not monitor or enforce compliance with conditions or penalise non-compliance, they were de facto unconditional (Baird 2013). First, the Plan de Atención Nacional a la Emergencia Social was conditional on pregnant women and children attending regular health check-ups and on children attending school regularly, but these conditions were not enforced (Amarante 2011). The Bono de Desarrollo Humano was conditional on children attending preventive health checks-ups and school but did not monitor compliance (Fernald 2011; Paxson 2007; Schady 2012). Moreover, the Direct Cash Transfer Program had no conditions, but eligible recipients may have understood that ongoing programme participation was contingent on reported level of household socioeconomic status (Bazzi 2012). However, we judged the risk of potential perceived conditionality as so low that we included the cash transfer in this review as unconditional.

Participants received minor co-interventions alongside three UCTs. UCT recipients received an electronic food card with a monthly value of approximately one-fourth to one-half of the value of the UCT in the Plan de Atención Nacional a la Emergencia Social (Amarante 2011). Workshops and social development activities were provided alongside the UCT in the Programa de Atención a Adultos Mayores en Zonas Rurales (Galiani 2014). And in Lesotho, participants received a UCT for assistance in humanitarian disasters, the Emer-
Programme uptake, when reported, was high, ranging between 78% and 100%. The included established government programmes often covered a considerable head count or proportion of the population. For example, the review included Indonesia’s Direct Cash Transfer Program, the world’s largest UCT programme with a population coverage of more than 19 million households (Bazzi 2012). Pilot government programmes and experiments often covered only fractions of the general population or smaller experimental samples. Studies rarely reported total costs of the included UCT interventions, but when they did, they were large for the established government programmes (e.g. approximately USD 250 million for the Plan de Atención Nacional a la Emergencia Social and USD 380 million for the Direct Cash Transfer Program).

All included studies compared a group eligible for or receiving a UCT with a group ineligible for or not receiving the UCT. The Haushofer 2013 was the only study that compared a group receiving a UCT with a group receiving a considerably smaller income amount from the UCT, but because this study also reported analyses of the UCT compared with no UCT, we prioritised the latter analyses. Three studies also compared both a UCT and a CCT with a control group and then tested for differences between the findings of these pair-wise comparisons (Akresh 2012; Baird 2010; Robertson 2012).

Ongoing studies

We describe the characteristics of the three ongoing studies identified for this review in detail in the Characteristics of ongoing studies table. First, the Galárraga 2014 RCT estimates the effect of an experimental UCT on health outcomes (disease prevalence) among 267 male sex workers in Mexico City. Second, the O’Leary 2011 CBA estimates the effect of an established government programme called the Benazir Income Support Programme on health services and health outcomes (anthropometric measures, disease prevalence and nutrition) among an unclear number of participants in Pakistan. Third, the Oxford Policy Management 2013 CBA estimates the effects of two established government programmes, the Vulnerable Families Support Grant and the Senior Citizens Grant, on use of health services and health outcomes (food security and nutrition) among members of 3960 households in 48 subcounties of eight programme districts in Uganda.

Excluded studies

A total of 95 records of 86 studies underwent full-text screening but did not fulfil the inclusion criteria. We document the reasons for excluding the 30 studies that were closest to the inclusion criteria in the Characteristics of excluded studies table (a full list of the excluded studies is available from the principal study author on request). We excluded 32 studies because they did not examine an eligible UCT for reducing poverty and vulnerabilities. These comprised three studies of UCTs for assistance in humanitarian disasters (Aker 2011; Langendorf 2013; Macours 2008), which we synthesised in our previous review (Pega 2015a), as well as studies of UCTs with major co-interventions, such as the Livelihood Empowerment Against Poverty Program in Ghana, which provided a UCT together with health insurance coverage (Handa 2014b). We excluded 17 studies because they did not examine one or more primary outcomes of this review, four studies because they did not examine an eligible study population, 26 studies because they used an ineligible study type, and seven studies because they did not report any empirical data.

Risk of bias in included studies

For each included study, we describe the likelihood of each type of bias in detail in the study’s individual ‘Risk of bias’ table. Figure 3 presents a summary of the individual ‘Risk of bias’ assessments of each study included in the review. We judged the overall risk of bias in this review to be high, especially due to potential selection and performance bias. We considered most studies to carry a high risk of attrition bias, with just over half of all cluster-RCTs reporting balanced samples at baseline. Most of the included cluster-RCTs recruited participants after they had allocated clusters, leading to a high risk of recruitment bias. Almost all studies had a high risk of performance bias due to the infeasibility of blinding participants to cash transfer interventions (as is the case for most social interventions in general), with the risk of bias from contamination often unclear due to lack of assessment (e.g. spill-over control groups were not commonly included in cluster-RCTs). Most studies carried an unclear risk of bias from allocation concealment due to insufficient reporting, as well as an unclear risk of bias from selective reporting due to the lack of published study protocols. Most studies carried a low risk of selection bias from random sequence generation and a low risk of bias for other reasons such as misclassification, confounding and reverse causation. Most observational studies carried a high risk of confounding.
Figure 3. Risk of bias summary: review authors’ judgements about each risk of bias item for each included study.

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<tr>
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<th>Allocation concealment (selection bias)</th>
<th>Baseline outcome measurements similar</th>
<th>Baseline characteristics similar</th>
<th>Selective reporting (reporting bias)</th>
<th>Other bias</th>
<th>Binding of outcome assessment (detection bias)</th>
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Allocation

We considered most studies to carry a high risk of selection bias. We assessed selection bias based on the following four criteria: whether there was documented random sequence generator to select participants into the study and allocated to intervention or control arms; whether random allocation to the intervention group or the control group was appropriately concealed (e.g. in a sealed envelope); whether baseline differences existed between the intervention and the control group in outcome measurements; and whether there were baseline differences in population characteristics between the intervention and control group. For cluster-RCTs, we also assessed the risk of recruitment bias due to participants having been recruited after allocation of clusters, as detailed in Higgins 2011b, Chapter 16.3.

Based on the first criterion (i.e. random sequence generation documented), we judged six studies to have high risk of selection bias, 14 studies to have a low risk of selection bias from random sequence generation, and one study to have unclear selection bias.

Based on the second criterion of whether random allocation to the intervention group or the control group was appropriately concealed, we judged seven studies to be at low risk of selection bias due to inadequate or lack of allocation concealment, generally because they allocated the cash transfers in public lotteries, thereby protecting allocation concealment. Allocation concealment was unclear for the remaining 14 studies. Finally, of the 21 cluster-RCTs, we judged 9 to carry a high risk of recruitment bias, 9 to have a low risk, and 3 to have an unclear risk of this bias.

Based on the third criterion (i.e. no baseline differences between the intervention and the control group in outcome measurements), we judged 2 studies to be at high risk since the intervention and control groups exhibited baseline differences for outcome measures, 14 studies to carry low risk, and 5 studies to be at unclear risk of bias because they did not compare outcome measurements at baseline for the intervention group and/or the control group included in this review.

For the fourth criterion (i.e. no baseline differences between the intervention and the control group in population characteristics), seven studies were at high risk of bias because of documented baseline differences in population characteristics, nine studies were at low risk of bias, and five studies were at unclear risk because of a lack of information on baseline characteristics.

Blinding

We assessed the risk of performance bias in the included studies based on whether participants and study personnel were blinded to the intervention. In studies where participants allocated to the intervention group were given a cash transfer, blinding of participants was virtually impossible. Similarly, blinding of study personnel again is also not practical and was often reported ambiguously. Consequently, we judged the risk of performance bias to be high for all included experimental studies and for one observational study for all outcomes. For four observational studies, we judged the risk of performance bias to be low because these studies used secondary data collected for purposes other than an assessment of the UCT, and therefore we regarded performance bias to be unlikely in these circumstances.

Assessment of detection bias was based on a combination of whether or not outcome assessors (e.g. interviewers or medical study personnel) were blinded to participants’ intervention status and whether or not they used objective outcome measures. For self-reported outcome measures, even those collected through a structured interview by blinded study personnel, we considered the participants themselves to be outcome assessors. For these outcomes, we considered if the outcome was affected by lack of blinding of participants in two ways: whether it influenced the participants’ behaviour and expectations in a way that genuinely affected their outcomes, and if it led participants to report their outcomes in a way that over- or under-reported what actually happened. We considered that the nine included studies that neither blinded outcome assessors nor used objective measures carried a high risk of detection bias. It was unclear whether two studies blinded outcome assessors, and since they did not use objective measures either, we judged them to carry an unclear risk of detection bias. Finally, eight studies blinded outcome assessors and/or used objective measures, or they used self-reported measures in an way that we considered to neither influence the outcome itself nor its reporting, so we judged these studies to carry a low risk of detection bias.

We also assessed the risk of contamination as a result of performance bias, detection bias or both. About half of all studies (11 out of 21) failed to report investigations of the level of contamination (e.g. for cluster-RCTs, they did not include spill-over control groups), and we judged them to be at unclear risk of contamination bias. Based on reported levels of contamination, we assessed five studies each to be at high and low risk of bias from contamination.

Incomplete outcome data

We judged 13 studies to be at high risk of attrition bias, because: they lost a considerable percentage of participants to follow-up; the proportion of participants lost to follow-up in the intervention group differed considerably from that in the control group; and/or the report was missing a considerable percentage of clusters, participants and/or outcome values. We judged four studies to be at low risk of attrition bias because we considered the levels of loss to follow-up and missing data to be unlikely to have introduced noteworthy bias. Four studies were at an unclear risk of attrition bias due to insufficient reporting. Reporting of missing outcomes was relatively poor across the included studies. For example, only two studies reported the number or percentage of missing participants by outcome.

Selective reporting

We judged the risk of reporting bias to be unclear for virtually all studies. While several included studies (especially the large-scale cluster-RCTs) have comprehensive baseline reports, they generally did not seem to have pre-published study protocols that would have enabled us to check these studies rigorously for selective reporting. The only exception was Haushofer 2013, which had a published study protocol that we could assess; this study reported the outcomes and analyses that it had prespecified in the study protocol, suggesting a perhaps low risk of bias from selective reporting. However, the study protocol was only registered in the American Economic Association’s registry for randomised controlled trials on June 28, 2013 (RCT ID: AEArCTR-0000019), which is after data collection for the trial had occurred between May 1, 2011, and February 28, 2013.
Other potential sources of bias

Misclassification bias of the exposure variable

Three of the four CBAs and the cohort study also may potential-
ly have incurred a risk of misclassification bias of the exposure.
These studies used self-reported receipt of a UCT collected in sur-
veys as the exposure variable. Validation studies have shown that
survey data on receipt of publicly funded financial credits can suffer
from misclassification, at least in high-income countries (Hjollund
2007). However, overall, we judged this risk of bias to have been
low, considering that the study participants were likely to be aware
of whether they received a UCT.

Confounding

We judged all cluster-RCTs to carry a low risk of confounding. De-
spite some cluster-RCTs having baseline differences in outcome
measurements and/or population characteristics, they robustly
adjusted for these differences and several key confounders using
regression analyses, minimising the risk.

We judged three of the four included CBAs to be at high risk of
bias from confounding. The first CBA compared the before-and-aft-
derence in the outcomes among participants receiving the UCT
(exposed group) with the before-and-after differences in the
outcomes among participants not receiving the UCT (unexposed
group) (Amarante 2011). This DD approach adjusted for confound-
ing by underlying time trends in the outcome. However, if the un-
derlying trend in the unexposed group differed from that in
the exposed group, then the DD estimator is confounded. Because
we believe that this is conceivable, we judged the likelihood of con-
foundning from differences in underlying time trends in the outcome
to be high in this study. However, the study robustly controlled
for some confounders (i.e. children’s sex, mother’s age and edu-
cation, twindhood, number of previous pregnancies, and month
of the baseline survey and of enrolment into the UCT). It also in-
cluded individual fixed effects to adjust for time-invariant confound-
ing in maternal characteristics that may potentially have confound-
ed the cash transfer-health relationship in children. However, Ama-
rante 2011 did not adjust for several other potential time-invar-
iant confounders (e.g. caregiver’s motivation and cognitive abili-
ties) or time-varying confounders (e.g. changes in access to health
services, fertility and income over time). Therefore, we judged the
risk of bias from these confounders to be high.

Two other CBAs also determined a treatment effect estimate us-
ing similar DD methods and identification strategies as described
above for Amarante 2011 (Galiani 2014; Salinas-Rodriguez 2014).
Again, if the underlying time trends in the unexposed group dif-
fered from the exposed group (which we believe is plausible), then
the DD estimator was at a high risk of confounding. Galiani 2014
used individual fixed effects to adjust for time-invariant confound-
ing by participant’s time-invariant characteristics. However, it did
not adjust for time-varying confounders such as assets, income and
labour force participation, which we judged to carry a high risk of
confounding.

Finally, we judged the fourth CBA to be at low risk for confound-
ing (Bazzi 2012). This study also used DD methods to adjust for
confounding by underlying time trends in the outcome, which may
have conferred a risk of bias. However, the study also used inverse
probability of treatment weighting in addition to robustly adjusting
for a large number of relevant confounders (for a list, see p 48 of the
included study record) and province-level fixed effects to adjust for
time-invariant confounders of the provinces. We judged this level
of confounder adjustment to suggest a low risk of bias from popu-
lation characteristics in this study.

We judged the Agüero 2007 cohort study to be at high risk of con-
foundning. This study used regression analysis to robustly adjust
for several potential confounders (i.e. participant’s age, motivation
and sex; principal caregiver’s age, education, sex, marital status
and occupation), and it used village-level fixed effects to adjust for
time-invariant confounding by geographic residency (e.g. access to
and quality of health services). However, it did not adjust for sever-
al other potential time-invariant confounders (e.g. caregiver’s mo-
tivation and cognitive abilities) and time-varying confounders (e.g.
changes in access to health services, fertility and income over time).

Reverse causation

Reverse causation occurs in repeated measures studies when the
outcome variable at earlier time points influences the intervention
(or exposure) value at later time points. Because cluster-RCTs ran-
domly allocate clusters to the intervention or control group, re-
verse causation is generally not a concern for these study designs.
Observational studies, however, may be at risk of reverse causa-
tion because the researcher does not assign the intervention but
instead purely observes it. None of the included five observational
studies controlled for reverse causation. However, we judged the
risk of reverse causation to be so negligible that we appraised all
four observational studies to only carry a low risk.

Effects of interventions

See: Summary of findings for the main comparison

Summary of findings: unconditional cash transfer versus no unconditional cash
transfer: primary outcomes; Summary of findings 2 Summary of
findings: unconditional cash transfer versus no unconditional cash
transfer: secondary outcomes; Summary of findings 3 Summary of
findings: unconditional cash transfers versus conditional cash
transfers: primary outcomes

Unconditional cash transfer versus no unconditional cash

transfer

Use of health services

Registered birth

Four cluster-RCTs with an effective sample size of 2376 children
assessed the effect of a UCT versus no UCT on the likelihood of
having ever had one’s birth registered at the time of the interview
among participants, when followed up either 2 to 4 months after
12 months of the intervention, or at 24 months into the interven-
tion (Pellerano 2014; Robertson 2012; Seidenfeld 2013; Ward 2010).
The treatment effects for all four studies were DD estimates of pro-
portions, which are not accepted by Cochrane because they erro-
nosely treat a dichotomous outcome as a continuous outcome.
As recommended in the Cochrane Handbook for Systematic Reviews
of Interventions (Chapter 16.3, Higgins 2011b), we calculated an RR
for each study, conducting approximately correct analyses of clus-
ter-RCTs using the crude frequency measures reported in the study
records (see detailed description in Measures of treatment effect).
We considered the four studies to be sufficiently homogenous in
study design, treatment effect estimate, population, intervention,
comparator and outcome to be combined, and we therefore con-
ducted a meta-analysis using the Maentel-Haessler method with ran-


Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)
dom effects to adjust for heterogeneity. However, the meta-analysis (Analysis 1.1) suggested that the studies were highly statistically heterogeneous (i.e. $I^2 = 95\%$), and as recommended by Deeks 2011, we decided to not report totals from the meta-analysis and to synthesise the studies narratively. One study, the Pellerano 2014 cluster-RCT with an effective sample size of 666 participants, reported that in relative terms a UCT led to a very large increase in the likelihood of having ever had one's birth registered at the time of the intervention, when followed up after 24 months (Pellerano 2014: RR 3.02, 95% CI 2.36 to 3.86). In absolute terms, assuming a likelihood at baseline of 129 per 1000 participants (i.e. the baseline risk in the control group reported in the study record), after the intervention the likelihood was 390 per 1000 (95% CI 304 to 498). Although we are not aware of international standards for judging change in this outcome, we nevertheless judged this magnitude to be clinically meaningful. Three studies reported very imprecise and therefore very uncertain estimates of the effect of the UCT on the outcome at either 2 to 4 months after 12 months of the intervention, or at 24 months into the intervention (Robertson 2012: RR 0.92, 95% CI 0.69 to 1.23, N = 224 (effective sample size); Seidenfeld 2013: RR 0.94, 95% CI 0.77 to 1.16, N = 1112 (effective sample size); and Ward 2010: RR 0.97, 95% CI 0.71 to 1.32, N = 374 (effective sample size)). We applied the GRADE criteria to assess the quality of the body of evidence for this outcome, assessing study limitations (risk of bias), consistency, imprecision, indirectness, publication bias, strength of effect and evidence for a dose-response relationship. For this outcome we describe the assessment for each criterion, but for other outcomes we only describe in this section our assessment for the criteria for which we down- or upgraded the quality of the body of evidence. Regarding study limitations, because all four studies lacked allocation concealment and blinding and had potential contamination and/or confounding (see ‘Risk of bias’ tables in Characteristics of included studies), we downgraded the quality of evidence for serious risk of bias by one grade. Regarding consistency, as noted above, the percentage of the variability in effect estimates that was due to heterogeneity rather than sampling error was very high (i.e. an $I^2$ of 90% or more), suggesting that inconsistency across studies may have been important, and we therefore downgraded the quality of the evidence by two grades for very serious inconsistency. Regarding imprecision, for three of the included studies the 95% CI of the effect estimate suggested that the effect may range from a large reduction in the likelihood of having one’s birth registered (i.e. a harmful effect) to a large increase in the likelihood (i.e. a beneficial effect), and we therefore downgraded the evidence for very serious imprecision by two levels. We considered the measurement to capture the outcome well, did not identify any signs of publication bias, did not consider the strength of effect because we judged the study to carry a high risk of bias and did not find any evidence for a dose-response relationship. We consequently did not downgrade or upgrade the quality of the evidence for these criteria. In summary, we assessed the body of evidence on this outcome to be of very low quality (i.e. starting at very high for experimental evidence and downgrading by five grades in total). In conclusion, we are very uncertain about the effect of a UCT on the likelihood of having ever had one’s birth registered.

**Growth checks**

Four studies (of three cluster-RCTs) assessed the treatment effect of a UCT on the likelihood of having had a growth check in the previous 6 months, at a follow-up of 15 and 27 months into the intervention (Fernald 2011; Paxson 2007; Pellerano 2014; Ward 2010). Fernald 2011 and Paxson 2007 reported results from the same cluster-RCT, and we prioritised Paxson 2007 because it analysed the entire study sample, whereas Fernald 2011 analysed only a selection of the study sample, so we did not report any of its data for this outcome. Paxson 2007 reported a coefficient from a confounder-adjusted probit regression model that we converted into an OR. Pellerano 2014 and Ward 2010 reported treatment effect estimates that Cochrane does not accept (i.e. DD estimates of proportions), and to ensure comparability with the estimates of the other two studies we calculated an OR for each of these two studies, conducting an approximately correct analysis of cluster-RCTs using the crude frequency measures reported in the study records (Chapter 16.3, Higgins 2011b). We considered the three studies with an effective sample size of 2261 children to be sufficiently homogenous to be combined in a meta-analysis.

Paxson 2007 reported treatment effect estimates separately for children according to their household’s income. These analyses compared children from poor families in the intervention group with children from poor families in the control group and compared children from non-poor families in the intervention group with children from non-poor families in the control group. Because these two analyses used different control groups, we included them both in the meta-analysis, and for better transparency we added them as separate analyses in the forest plot.

In relative terms, the point estimate from the meta-analysis for the treatment effect was that a UCT led to moderately higher odds of having received a growth check, but the 95% CI allowed for both a non-meaningful change and a moderate increase in the odds (Analysis 1.2; OR 1.11, 95% CI 0.98 to 1.24, 3 cluster-RCTs, N = 2261 (effective sample size), $I^2 = 0\%$). In absolute terms, assuming the baseline likelihood of 450 per 1000 participants, after receiving the UCT an estimated 468 per 1000 participants (95% CI 446 to 491) had received a growth check.

We downgraded this body of evidence to low quality for serious risk of bias (minus one grade) and for serious imprecision (minus one grade). In conclusion, a UCT may not have had a meaningful effect on the likelihood of having received a growth check, but further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

**Up-to-date on vaccination calendar**

Four studies (all cluster-RCTs) assessed the effect of a UCT on the likelihood of being up-to-date for all vaccinations on the immunisation calendar at 8 to 24 months into the intervention. Three studies with an effective sample size of 563 children reported treatment effect estimates as DD estimates of proportions, so we calculated an RR for each study, conducting an approximately correct analysis for cluster-RCTs using the crude frequency measures reported in the study records (Chapter 16.3, Higgins 2011b; Pellerano 2014; Robertson 2012; Ward 2010). We considered the studies to be sufficiently homogenous to be combined in a meta-analysis. In relative terms, the point estimate from the meta-analysis was no meaningful change in the likelihood of being fully vaccinated, but the 95% CI suggested that the effect may lie between a moderate reduction and a moderate increase in the likelihood (Analysis 1.3; RR 1.02, 95% CI 0.90 to 1.15, 3 cluster-RCTs, N = 563 (effective sample size), $I^2 = 3\%$). In absolute terms, assuming a likelihood before the intervention of 646 per 1000 participants (i.e. the median risk in the control group before the intervention in the three studies), after receiving
the UCT an estimated 661 per 1000 participants (95% CI 583 to 745) were fully up-to-date on their vaccinations.

Beck 2015, a cluster-RCT with 2034 households (effective sample size could not be calculated due to missing frequency counts), assessed the effect of a UCT on the odds of all children in the household being completely vaccinated (i.e. with bacillus calmette guerin, diphtheria, pertussis, tetanus, polio, measles, mumps and rubella) at an age of 6 months to 5 years, when followed up 8 months into the intervention. We could not include this study in the meta-analysis because of the different participants (households, not individual participants). The point estimate was a small increase in the odds, with the 95% CI suggesting that the effect may lie between a large reduction and a large increase in the odds (OR 1.04, 95% CI 0.60 to 1.82). Because the likelihood at baseline was unclear, we could not convert the OR into an RR.

We downgraded this body of evidence to very low quality for serious risk of bias (minus one grade) and very serious imprecision (minus two grades). In conclusion, we are very uncertain about the effect of a UCT on the likelihood of having been fully vaccinated.

Treatement for parasites

Two studies (of the same cluster-RCT) assessed the effect of a UCT on the likelihood of having been given any parasite treatment in the previous year, at 15 to 27 months into the intervention (Fernald 2011; Paxson 2007). As above, because both studies reported results from the same cluster-RCT, we again included only the results from Paxson 2007, which analysed the entire study sample instead of only a selection. Paxson 2007 reported a coefficient from a confounder-adjusted probit regression model that we converted into an OR. We again included the two separate treatment effect estimates for children from poor and non-poor families reported in the study record as separate analyses in a meta-analysis. In relative terms, the point estimate for the treatment effect was a large increase in the odds, with the 95% CI suggesting that the effect may lie between a moderate and a large increase in the odds (OR 1.28, 95% CI 1.06 to 1.54, 1 cluster-RCT, N = 1478, I² = 35%). We assumed a baseline likelihood of 450 per 1000 participants (i.e. in the absence of baseline data from Paxson 2007 for this outcome, we here used the likelihood in the control group before the intervention in Fernald 2011). In absolute terms, after receiving the UCT an estimated 513 per 1000 participants (95% CI 463 to 558) had received parasite treatment in the last year. We are not aware of international standards for judging change for this outcome, but we did judge this level of change to probably be clinically meaningful. We downgraded the body of evidence to moderate quality for serious risk of bias (minus one grade). In conclusion, a UCT probably had a beneficial effect on the likelihood of receiving treatment for parasites. Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Use of any health service

Six studies assessed the effect of a UCT on a broad summary measure of the likelihood of having used any health service in the previous 1 to 12 months, at 12 to 24 months into the intervention: five cluster-RCTs with an effective sample size of 4972 participants (Luseno 2012; Oxford Policy Management 2012; Pellerano 2014; Seidenfeld 2013; Ward 2010), plus one CBA in 9034 households (Bazzi 2012).

Because the five cluster-RCTs reported treatment effect estimates as DD estimates of proportions, we calculated RRs for each study, conducting approximately correct analyses using the crude frequency measures reported in the study records (Chapter 16.3, Higgins 2011b). The studies differed somewhat in their outcomes (i.e. has used any health service for: any condition; worst illness; illness or injury; diarrhoea; and fever, cough or diarrhoea); reporting periods (i.e. one month, three months, and one year prior to the interview); and study population (i.e. children and adults). However, we nevertheless considered them sufficiently homogenous to combine in one meta-analysis. Pellerano 2014 reported separate treatment effect estimates for three age groups (i.e. children aged up to 17 years, adults aged 18 years to 59 years, and adults aged 60 years and over), which we included separately in the meta-analysis (Figure 4). In relative terms, the point estimate for the treatment effect was a small increase in the risk, with the 95% CI suggesting that the effect estimate may lie between no change and a small increase in the risk (RR 1.04, 95% CI 1.00 to 1.09, 5 cluster-RCTs, N = 4972, I² = 2%, Analysis 1.5). In absolute terms, assuming a risk before the intervention of 487 per 1000 participants (i.e. the median risk in the control group before the intervention reported in the five studies), after receiving the UCT an estimated 506 per 1000 participants (95% CI 487 to 531) had used any health service.
Bazzi 2012 reported an estimate, which we interpreted as an MD, of the effect of a UCT on the number of outpatient health services visits per household member in the month prior to the interview, at 12 months into the intervention. In relative terms, the point estimate from this study was a very small increase in the number of outpatient health services visits per household member, with the 95% CI suggesting that the effect may be anything from a large reduction to a small increase in the number (MD −0.06 visits, 95% CI −0.20 to 0.07). In absolute terms, assuming a risk before the intervention of 0.20 outpatient health services visits per household member (i.e. the risk in the control group prior to the intervention), after receiving the UCT a household used an estimated 0.14 outpatient health services visits (95% CI 0.00 to 0.27) per member.

We downgraded this body of evidence to low quality for serious risk of bias (minus one grade) and serious imprecision (minus one grades). In conclusion, a UCT may perhaps not have had a meaningful effect on the likelihood of using any health service, but further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Health outcomes

Stunting

Seven studies (six cluster-RCTs and the cohort study) assessed the effect of a UCT on the likelihood of being stunted or on a related measure (i.e. height for age or mean height). Two cluster-RCTs with an effective sample size of 551 children reported an estimate for the effect of a UCT on the risk of being moderately stunted at the time of the interview, at 24 months into the intervention (Oxford Policy Management 2012; Ward 2010). Both studies used UNICEF’s standard measure for moderate stunting (i.e. a height-for-age z-score of less than 2 standard deviations under the median height for age in the reference population) (UNICEF 2015). Because both studies reported treatment effects as DD estimates of proportions, we calculated RRs for each study, conducting approximately correct analyses using the crude frequency measures reported in the study records (Chapter 16.3, Higgins 2011b). We considered the two studies to be sufficiently homogenous to be combined in a meta-analysis. In relative terms, the point estimate was a small reduction in the risk, but the 95% CI suggested that the effect may be anywhere between a moderate reduction and a moderate increase in the risk (Analysis 2.1; RR 0.96, 95% CI 0.75 to 1.21, 2 cluster-RCTs, N = 551 (effective sample size), $I^2 = 2\%$). In absolute terms, assuming a risk before the intervention of 337 per 1000 participants, after receiving the UCT an estimated 324 per 1000 participants (95% CI 253 to 408) were moderately stunted.

Three studies (of two cluster-RCTs) assessed height for age (a relatively less preferable measure, because it is less direct) at 15 to 27 months into the UCT intervention (Fernald 2011; Paxson 2007; Seidenfeld 2013). As Fernald 2011 and Paxson 2007 reported results from the same cluster-RCT, we again only used data from Paxson 2007 because it analysed the entire study sample rather than only a selection. Paxson 2007 reported the treatment effect estimate as (we believe) an MD, and the Seidenfeld 2013 study reported a DD as the treatment effect estimate. We considered both types of treatment effect estimates to be sufficiently similar to be combined in one meta-analysis. Since the studies were also sufficiently homogenous in their other features, we combined them in a meta-analysis. In relative terms, the point estimate was an increase in the mean height-for-age score, but the 95% CI suggested that the effect may be anywhere from a slight reduction to a slight increase in the score (Analysis 2.2; MD 0.04 of 1 SD, 95% CI −0.05 to 0.13, 2 cluster-RCTs, N = 7545, $I^2 = 0\%$). Because the baseline height-for-age score was unclear and since we could not retrieve the value of 1 SD of the height-for-age score, we were unable to convert this relative treatment effect estimate into an absolute value and to calculate an absolute treatment effect estimate, such as a centimetre change. However, calculating an absolute value and an absolute treatment effect estimate for a standardised, z-transformed height-for-age score may not necessarily be desirable anyway, considering that the purpose of z-transformation of these measures is to give the reader a sense of deviation from ‘normality’. We are not aware of an internationally agreed standard on which level of change in the height-for-age score is sizeable or clinically meaningful, respectively, so we were unable to confidently judge the effect size and the clinical meaningfulness of this level of change in this outcome.
One additional cluster-RCT in 5190 children assessed the effect of a UCT on height at 24 months into the intervention (Cunha 2014). In relative terms, the point estimate was that the UCT led to no meaningful change in mean height, with the 95% CI suggesting that the effect may lie between a moderate reduction and a small increase in the mean height (MD −0.15 cm, 95% CI −0.99 to 0.61). In absolute terms, assuming a mean height before the intervention of 84.95 cm (i.e. the mean height in the control group before the intervention), after receiving a UCT the estimated mean height was 84.80 cm (95% CI 83.96 to 85.41).

A cohort study in 1606 participants (comprising all treated children and all non-treated beneficiary, applicant and non-applicant children) reported the height-for-age score at an unclear follow-up time point (Agüero 2007). We could not include the study in the meta-analysis because it was a different study type (i.e. not a cluster-RCT). The mean height-for-age score in the group that had received the UCT for two-thirds of the duration of the three-year nutritional window was 0.25 of 1 SD higher than the group that received a UCT for 1% of the duration of the three-year nutritional window (95% CI unclear, P < 0.05). The mean height-for-age score in the control group was −1.08 of 1 SD. As above, it is neither possible nor useful to convert this relative treatment effect estimate into an absolute treatment effect estimate, and in the absence of internationally agreed standards on which level of change is clinically meaningful, we could not judge the clinical meaningfulness of this level of change in this outcome.

We downgraded this body of evidence to very low quality for serious risk of bias (minus one grade), for serious indirectness due to data on the key outcome of interest (i.e. moderate stunting) only being available for two of the seven studies (minus one grade); and for very serious imprecision, especially for the effect of a UCT on the proportion of participants who were stunted. In conclusion, we are very uncertain about the effects of UCTs on the likelihood of being moderately stunted.

Underweight

Seven studies (6 cluster-RCTs, 1 CBA) assessed the effect of a UCT on the likelihood of being overweight or evaluated a related measure (i.e. weight for age, mean weight or the likelihood of having a low birth weight). Three cluster-RCTs with an effective sample size of 701 children reported the likelihood of participants being underweight at the time of the interview or at one year of age, when followed up 24 months into the intervention (Oxford Policy Management 2012; Pellerano 2014; Ward 2010). Because all three cluster-RCTs reported treatment effect estimates as DD estimates of proportions, we calculated an RR for each study, conducting approximately correct analyses using the crude numbers reported in the study records (Chapter 16.3, Higgins 2011b). The reporting period for two studies was at the time of the interview, whereas the third study measured the outcome when the child was one year old. Two studies measured the proportion of participants who were moderately underweight as per UNICEF standard definition (weight-for-age score less than 2 SDs under the median score in the reference population) (UNICEF 2015), whereas the third study did not specify the severity of underweight. Despite this slight heterogeneity in reporting period and outcome measurement, we considered the three studies to be sufficiently homogeneous to be combined in a meta-analysis. In relative terms, the point estimate was no change in the risk, with the 95% CI suggesting that the effect may have been anywhere from a very large reduction and a very large increase in the risk (Analysis 2.3; RR 1.00, 95% CI 0.75 to 1.32, 3 cluster-RCTs, N = 701, I² = 0%). In absolute terms, assuming a risk before the intervention in the control group of 337 per 1000, after receiving the UCT an estimated 337 per 1000 participants (95% CI 253 to 445) were moderately underweight.

Three other cluster-RCTs reported alternative weight measures in child participants, but these were too heterogenous to be combined in a meta-analysis. The Seidenfeld 2013 study in 6825 children reported the estimated weight-for-age score itself at 24 months into the UCT intervention. In relative terms, the point estimate was an increase in the weight-for-age score, with the 95% CI suggesting that the effect may be between no change and an increase (MD 0.13 of 1 SD of the score, 95% CI 0.00 to 0.26). In absolute terms, considering that the mean weight-for-age score in the control group was −0.90 of 1 SD, after receiving the UCT a participant would have an estimated weight-for-age score of −0.77 of 1 SD (95% CI −0.90 to −0.64). Because the value of 1 SD in the weight for age score was unclear, it was neither possible nor useful to convert this relative treatment effect estimate into an absolute treatment effect estimate. The Cunha 2014 cluster-RCT in 5277 children reported mean weight at 24 months into the intervention. The point estimate was no change in weight, but the 95% CI suggested that the effect may be anywhere between a small reduction and a small increase (MD −0.06 kg, 95% CI −0.39 to 0.27). In absolute terms, considering that the mean weight in the control group before the intervention was 12.19 kg, after receiving the UCT recipient children weighed 12.13 kg (95% CI 11.80 to 12.46). The Leroy 2010 cluster-RCT in 3010 adult mothers reported that a UCT had led to a small increase in maternal weight at 24 months, with the 95% CI suggesting that the effect was between no change and a moderate increase (MD 0.40 kg, 95% CI 0.01 to 0.79). In absolute terms, since the mean maternal weight in the control group before the UCT was 62.60 kg, the UCT would have increased it to 63.00 kg (95% CI 62.61 to 63.39). We judged this level of change to probably not be clinically meaningful.

Finally, one CBA in 68,858 children assessed the effect of a UCT for reducing the proportion of children with low birth weight (Amarante 2011). The study reported the treatment effect estimate as a DD estimate of a proportion, and we were not able to retrieve or calculate an accepted treatment effect estimate for this study for this outcome. However, for its 21,374 adult participants (all mothers), the study did report an acceptable estimate of maternal weight at week 35 of pregnancy, at 1 to 2 months into the intervention. The treatment effect estimate was a DD estimator of a continuous outcome, which subtracted the difference in the mean maternal weight among UCT recipients (intervention group) and UCT non-beneficiaries (control group) prior to the UCT intervention, from the same difference in the mean maternal weight among the two groups after the UCT intervention had been initiated, to adjust for changes in the outcome over time in the control group. Since DD estimators of continuous outcomes are perhaps comparable to confounder-adjusted MDs, Cochrane does accept them. The point estimate was a large increase in maternal weight among UCT recipients compared with non-UCT recipients, adjusted for changes in the outcome over time (DD estimator 0.97 kg, 95% CI 0.17 to 1.76). Considering that the mean weight in the control group before the intervention was 63.26 kg, the UCT would have increased mean maternal weight to 64.23 kg (95% CI 63.43 to 65.02). We judged this level of change to probably be clinically meaningful.
We downgraded the quality of the evidence to very low for serious risk of bias (minus one grade), very serious imprecision (minus two grades), and serious indirectness (minus one grade). We are very uncertain about the effect of UCTs on the likelihood of being underweight. Further research is very likely to have an important impact on our confidence in the estimates of effect and is likely to change the estimates.

Death
No evidence was available on the effect of a UCT on the likelihood of having died.

Disease or illness
Nine studies (all cluster-RCTs) assessed the effect of a UCT on the likelihood of having had any illness or the likelihood of having had a specific illness in the two weeks to three months prior to the interview. Five cluster-RCTs with an effective sample size of 1483 children and adults reported this outcome at 12 or 24 months into the intervention (Baird 2010; Cunha 2014; Luseno 2012; Oxford Policy Management 2012; Pellerano 2014). All five studies reported treatment effect estimates as DD estimates of proportions. For the Baird 2010 study we had access to micro-data, which we re-analysed to calculate an OR, adjusted for all the confounders that the original study considered (for details see Measures of treatment effect). For Cunha 2014, we received an OR estimate from the study author that was fully adjusted for all the confounders used in the effect estimate reported in the study record. For the other three cluster-RCTs, we conducted an approximate analysis with the crude numbers reported in the study records and estimated a crude OR (Chapter 16.3, Higgins 2011b). The reporting period for the outcome differed between the studies (two weeks, one month, and three months prior to the interview). Whereas four studies included children only, the fifth study included both children and young adults. However, we considered the studies to nevertheless be sufficiently homogenous to be combined in one meta-analysis. Figure 5 presents a forest plot of the meta-analysis. In relative terms, the point estimate represented a very large reduction in the odds of having had any illness, with the lower and upper limits of the 95% CI also suggesting that the effect was large (OR 0.73, 95% CI 0.57 to 0.93, 5 cluster-RCTs, N = 8446, I² = 57%). In absolute terms, assuming baseline risk in the control group of 370 per 1000 participants (i.e. the median risk in the control group before the intervention in the five studies), after receiving the UCT an estimated 300 per 1000 participants (95% CI 252 to 352) had had any illness. We judged this level of change to be clinically meaningful.

### Figure 5. Forest plot of comparison: 2 Unconditional cash transfer versus no unconditional cash transfer for improving health outcomes, outcome: 2.4 Has had any illness in previous 2 weeks to 3 months.

<table>
<thead>
<tr>
<th>Study or Subgroup</th>
<th>log(Odds Ratio)</th>
<th>UCT Total</th>
<th>SE</th>
<th>No UCT Total</th>
<th>Weight</th>
<th>Odds Ratio, 95% CI</th>
<th>Odds Ratio, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baird 2010</td>
<td>-0.09</td>
<td>0.12</td>
<td>263</td>
<td>2130</td>
<td>28.3%</td>
<td>0.81 [0.72, 1.16]</td>
<td></td>
</tr>
<tr>
<td>Cunha 2014</td>
<td>-0.48</td>
<td>0.14</td>
<td>2275</td>
<td>2275</td>
<td>26.2%</td>
<td>0.58 [0.48, 0.63]</td>
<td></td>
</tr>
<tr>
<td>Luseno 2012</td>
<td>-0.55</td>
<td>0.42</td>
<td>56</td>
<td>59</td>
<td>7.4%</td>
<td>0.58 [0.25, 1.31]</td>
<td></td>
</tr>
<tr>
<td>Oxford Policy Management 2012</td>
<td>0.04</td>
<td>0.27</td>
<td>30T</td>
<td>307</td>
<td>14.3%</td>
<td>0.94 [0.61, 1.77]</td>
<td></td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>-0.58</td>
<td>0.16</td>
<td>360</td>
<td>383</td>
<td>23.9%</td>
<td>0.58 [0.44, 0.77]</td>
<td></td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td></td>
<td></td>
<td>3295</td>
<td>5151</td>
<td>100.0%</td>
<td>0.73 [0.57, 0.93]</td>
<td></td>
</tr>
</tbody>
</table>

Heterogeneity: $\tau^2 = 0.04, \chi^2 = 9.36, df = 4 (P = 0.06), I^2 = 57%$

Test for overall effect: $Z = 2.52 (P = 0.01)$

Four other cluster-RCTs reported estimates of the effect on a related measure. The Haushefer 2013 cluster-RCT in 1327 households reported the proportion of household members (not individual participants) who had been sick or injured in the month prior to the interview, at between 7 months into a 9-month intervention and 10 months after completion of the intervention. The Schady 2012 cluster-RCT in 1196 children reported the likelihood of being anaemic at the time of the interview (not on the likelihood of having had any illness), at 50 to 57 months into the UCT intervention. However, both studies reported the treatment effect estimate as a DD of a proportion. Because we were not able to retrieve crude numbers for these studies, we could not conduct approximately correct analyses and cannot report estimates. The Seidenfeld 2013 cluster-RCT, with an effective sample size of 1104 children, did not report the likelihood of having had any illness, but it reported the likelihood of having had diarrhoea, fever or an acute respiratory illness in the two weeks prior to the interview, at 24 months into the intervention. Using the 2015 Global Burden of Disease Study estimates to judge relative importance of the outcomes in terms of burden of disability-adjusted life years attributable to the condition, we prioritised the estimate for the likelihood of having an acute respiratory illness, because upper and lower respiratory infections carry a larger burden of disease than diarrhoea, and fever does not adhere to any burden of disease category (GBD 2016). We could not include this study in the meta-analysis because we judged its outcome to be too different. Because the study reported treatment effect estimates as DD estimates of proportions, we calculated an RR with approximately correct analyses using the crude numbers reported in the study records (Chapter 16.3, Higgins 2011b). In relative terms, the point estimate represents a large reduction in the likelihood of having had an acute respiratory illness, with the 95% CI suggesting that the effect may lie between a very large and a small reduction in the risk (RR 0.61, 95% CI 0.39 to 0.96). In absolute terms, assuming a risk before the intervention of 200 per 1000 (i.e. the risk before the intervention in the control group), after the intervention 120 per 1000 participants (95% CI 80 to 190) had had an acute respiratory illness. We considered this considerable level of change to be clinically meaningful. The Beck 2015 cluster-RCT in 2034 households (effective sample size could not be calculated due to missing frequency counts) reported the likelihood of having cases of illness or injury in households that lasted more than 24 hours and needed treatment but not hospitalisation in the three months prior to the interview, at 8 months into the intervention. In relative terms, the point estimate represents a very large reduction in the odds, with the 95% CI suggesting that the effect may lie between a very large and a large reduction in the odds (OR 0.54, 95% CI 0.44 to 0.65).
In absolute terms, assuming the risk of 280 per 1000 participants before the intervention, after the intervention the risk was 174 per 1000 participants (95% CI 149 to 202). We judged this considerable level of change to be clinically meaningful.

We downgraded the body of evidence to moderate quality for serious risk of bias (minus one grade). In conclusion, the UCT probably led to a large, clinically meaningful reduction in the likelihood of having had any illness. Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Food security
Five studies (all cluster-RCTs) assessed the likelihood of being food secure or the level of food security at the time of the interview over the 1 month prior to the interview. Three cluster-RCTs with an effective sample size of 1386 households reported a single measure of the likelihood of being food secure, at 13 to 24 months into the intervention (Miller 2008; Oxford Policy Management 2012; Pellerano 2014). The measures used to capture food security were whether the household: had at least one member who had gone without adequate food for more than eight days per month (Miller 2008); and had been food insecure in the worst recent food shortage period (reverse coded in this review) (Oxford Policy Management 2012); and did not have enough food to meet its needs at least for 1 out of 12 months (reverse coded in this review) (Pellerano 2014). Despite the different measures and reporting periods we considered these studies to potentially be sufficiently homogenous to be combined in a meta-analysis. However, the meta-analysis (Analysis 2.5) suggested that the studies were highly statistically heterogeneous ($I^2 = 91\%$), and as recommended by Deeks 2011 (Chapter 9.5), we decided to not report totals from the meta-analysis and to synthesise the studies narratively. One study reported a possible small reduction in food security, but with the 95% CI suggesting that the effect may lie between a large reduction and a small increase (Oxford Policy Management 2012: RR 0.93, 95% CI 0.80 to 1.10), whereas two studies reported a large increase in the likelihood of being food secure (Miller 2008: RR 1.69, 95% CI 1.34 to 2.12; Pellerano 2014: RR 1.80, 95% CI 1.27 to 2.53).

Two cluster-RCTs examined the effect of a UCT on a composite index of food security among households. Hausoher 2013 used as the outcome measure a non-standard, non-validated household food security index measure (i.e. the weighted average of the proportions of household members going to sleep hungry and not eating protein in the week prior to the interview, with the score ranging from 0.00 to 1.00), reporting treatment effect as a DD of a weighted average of two proportions. Because we were not able to retrieve crude numbers for this study, we could not conduct approximate correct analyses and cannot report an estimate from this study. The Seidenfeld 2013 study in 2289 households reported a DD estimate of the effect of a UCT on the standard, validated Household Food Insecurity Access Scale in the month prior to the interview, when followed up 24 months in to the intervention (Coates 2007).

In relative terms, the DD estimate was a moderate increase in the score, with the upper and lower limits of the 95% CI also suggesting that the effect was moderate in size (DD 0.50, 95% CI 0.26 to 0.74). In absolute terms, assuming a score before the intervention of 15.10 (i.e. the average score in the study sample before the intervention), after receiving a UCT a household reported a score of 15.60 (95% CI 15.36 to 15.84). We are not aware of international standards for judging change in this score but nevertheless consider this level to probably be clinically meaningful.

We downgraded the body of evidence to low quality for serious risk of bias (minus one grade) and serious inconsistency (minus one grade). Because the very high level of heterogeneity (i.e. $I^2 \geq 90\%$) in the meta-analysis seems to be due to one outlier (Oxford Policy Management 2012), and considering that the other three studies with estimates all report meaningful or probably meaningful benefits, we judged inconsistency to be serious, rather than very serious. In conclusion, a UCT may increase the likelihood of food security, and further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Dietary diversity
Eight studies (all cluster-RCTs) reported an estimate of the effect of a UCT on the level of dietary diversity or a related measure. Four cluster-RCTs in 9347 households estimated the effect of a UCT on the standard, validated Household Dietary Diversity Score (HDDS; Kennedy 2011), at 24 months into the intervention (Oxford Policy Management 2012; Pellerano 2014; Seidenfeld 2013; Ward 2010). An increase in the HDDS indicates an increase in dietary diversity. All four studies reported a DD estimate as the treatment effect estimate, but studies used somewhat different HDDS, and we standardised the score to ensure comparability across the four studies. We considered these studies to be sufficiently homogenous to be combined in a meta-analysis. Figure 6 presents the forest plot of the meta-analysis. The point estimate was a moderate increase in the HDDS score, with the 95% CI suggesting that the effect estimate may be between a small increase and a moderate increase in the score (DD 0.41 of 1 SD, 95% CI 0.12 to 0.69, 4 cluster-RCTs, 9347 households, $I^2 = 79\%$). In absolute terms, assuming an SD of 1.46 (i.e. the SD reported for the Pellerano 2014 study), then a UCT increased the score by an estimated 0.59 food categories (95% CI 0.18 to 1.01). The international guidelines for analysing the HDDS note that there is no international standard for judging change in the HDDS (Kennedy 2011), but considering that the pooled absolute treatment effect estimate suggests an average increase in food diversity by 0.59 food categories, we considered this moderate level of change to probably be clinically meaningful.
Four additional cluster-RCTs also assessed dietary diversity. One cluster-RCT in 819 households reported an estimate of the effect of a UCT on the HDDS at 13 months into the intervention (Miller 2008). Because we could not retrieve an SD to standardise this estimate, we could not include this study in the meta-analysis. The study records reported DD estimates that were unadjusted for clustering, but we received an estimate from the principal study author that was. The point estimate represents a small reduction in the score, with the 95% CI suggesting that effect may lie between a moderate reduction and a small increase in the score (DD −0.10, 95% CI −0.34 to 0.13). One cluster-RCT in 1169 children assessed the effect of a UCT on a non-standardised, non-validated composite index of level of individual dietary diversity (Fernald 2011). The authors constructed this composite index using principal components analysis on whether children had eaten any of a list of 11 food items. The relevant food items included both nutritious foods (e.g. liver, chicken, pasta and/or bread, spinach and/or chard, carrots, citrus fruits and non-citrus fruits) and those with less nutritious value (e.g. ice cream and/or soda, potato chips, cookies and/or crackers, and candy). While this measure has some commonalities with the standard HDDS, we considered it to be too different to be combined with the HDDS, and the study participants were children rather than households. Therefore, we considered this study to be too different to be combined with the other four cluster-RCTs that used the HDDS. The point estimate showed a moderate increase in the score, but the 95% CI suggested that the effect may lie between a moderate reduction and a moderate increase in the score (MD 0.06 of 1 SD, 95% CI −0.08 to 0.20). In absolute terms, assuming the mean score before the intervention was −0.10 of 1 SD (i.e. the median score in the control group before the intervention), after receiving the UCT the score was −0.04 of 1 SD (95% CI −0.28 to 0.10). Since the value of 1 SD was unclear, we were not able to convert this estimate into an absolute value, such as change in the HDDS score.

Moreover, two cluster-RCT studies examined a single measure of dietary diversity. Heterogeneity in the outcome and the participants prohibited their meta-analysis. Baird 2010 included 2080 children and adults, reporting a large increase in the mean number of days that children or adults had eaten protein-rich food in the week prior to the interview, at 24 months into the intervention (MD 0.59 days, 95% CI 0.15 to 1.02). In absolute terms, assuming the mean number of days before the intervention was 3.95 days per week (i.e. the mean number in the control group before the intervention), after the intervention a participant had on average eaten protein-rich food 4.54 days per week (95% CI 4.10 to 4.97). We considered this level of change to be a clinically significant benefit. Haushofer 2013 included 1372 households, reporting a large increase in the mean number of times that a household had eaten meat or fish in the week prior to the interview, at between 7 months into the 9-month intervention and 10 months after its completion (DD 0.73, 95% CI 0.30 to 1.16). In absolute terms, assuming the mean number before the intervention was 2.41 times per week (i.e. the mean number in the control group before the intervention), a household had eaten meat or fish 3.14 times per week (95% CI 2.71 to 3.57) after the intervention. We considered this level of change to also be a clinically significant benefit.

We downgraded the body of evidence to low quality for serious risk of bias (minus one grade) and serious inconsistency (minus one grade). A UCT may have increased the level of dietary diversity, and further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

### Depression

Five studies (three studies of two cluster-RCTs, and one study each of two CBAs) assessed the mean score achieved on a psychometric test of level of depression at the time of the interview (Fernald 2011; Haushofer 2013; Galani 2014; Paxson 2007; Salinas-Rodriguez 2014). Two studies (of the same cluster-RCT) reported an estimate of the effect of a UCT on the internationally standardised, validated Center for Epidemiologic Studies Depression Score (CES-D; Eaton 2004) at 15 to 27 months into the intervention (Fernald 2011; Paxson 2007). As with other outcomes, because the Fernald 2011 and Paxson 2007 studies reported results from the same cluster-RCT, we used the (more complete) data only from Paxson 2007. Paxson 2007 standardised the treatment effect estimates by dividing them through 1 SD (i.e. a z-transformation). We included the estimates for adults from poor families and those from non-poor families separately in the meta-analysis. A reduction in the CES-D indicates a reduction in depression. The point estimate was a very small reduction in the score, but the 95% CI suggested that the effect may lie between a small reduction and a small increase in the score (MD −0.06 of 1 SD, 95% CI −0.25 to 0.13, 1 cluster-RCT, N = 1044, I² = 24%). The study reported neither the baseline average in this outcome, nor the value of 1 SD, so we were unable to calculate and present an absolute treatment effect estimate.

The third study, a cluster-RCT with 667 households, reported an estimate of the effect of a UCT on the mean CES-D at 7 to 9 months into the 9-month intervention and at up to 10 months after its completion (Haushofer 2013). We could not include this study in the meta-analysis because it studied households. In relative terms, the point estimate was a moderate increase in the CES-D, but the 95% CI suggested that the effect may be between a small reduction and a large increase in the score (DD 1.26 points, 95% CI −0.27 to 2.79). In absolute terms, assuming the mean CES-D before the intervention was 26.48 points out of 60.00 points (i.e. the mean score in the control group before the intervention), after the intervention the score was 27.74 points (95% CI 26.21 to 29.27).

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**Figure 6. Forest plot of comparison: 2 Unconditional cash transfer versus no unconditional cash transfer for improving health outcomes, outcome: 2.6 Dietary diversity (Household Dietary Diversity Score) in previous week.**
Two observational studies also reported relevant estimates. One CBA in 1950 adults reported the mean Geriatric Depression Scale (GDS) score at the time of the interview at 7 to 9 months into the UCT intervention (Galiani 2014). We could not include the study in the meta-analysis due to its different study type and outcome measure. The GDS is a standard psychometric test for depression in older adults, and a reduction in the score indicates a reduction in depression (Yesavage 1982). The point estimate showed that the UCT led to a moderate reduction in the GDS score, with the 95% CI suggesting that the effect may be between a large and a small reduction (DD = −0.42 points, 95% CI = −0.76 to −0.09). In absolute terms, assuming the mean GDS score before the intervention was 3.82 points out of 30 points (i.e. the mean score in the control group before the intervention), after the UCT the score was 3.40 (95% CI = 3.06 to 3.73). Although were not aware of international standards to judge this level of change, we considered that the change, which exceeded 10% of the pre-intervention score, was likely to be clinically meaningful. Another CBA in 5465 older adults reported GDS at the time of the interview, when followed up to 24 months into the intervention (Salinas-Rodríguez 2014). The point estimate represents a small reduction in the GDS score, indicating a reduction in depressive symptoms, with the 95% CI suggesting that the effect may lie between a small reduction and non-meaningful change in the score (DD = −0.06, 95% CI = −0.12 to −0.01). We judged that this level of change was probably not clinically meaningful.

We downgraded the body of evidence to very low quality for serious risk of bias (minus one grade) and very serious imprecision (minus two grades). In conclusion, we are very uncertain about the effect of a UCT on the level of depression.

Social determinants of health

Livestock ownership

Two cluster-RCTs assessed livestock ownership at the time of the interview or in the year prior to the interview, at 24 months into the intervention (Oxford Policy Management 2012; Pellerano 2014). Because both studies reported treatment effect estimates as DD estimates of proportions, we calculated an RR for each study, conducting approximately correct analyses using the crude numbers reported in the study records (Chapter 16.3, Higgins 2011b). We considered these studies to be sufficiently homogenous to be combined in a meta-analysis. However, the meta-analysis (Analysis 3.1) suggested that the studies were highly statistically heterogeneous (I² = 93%), and as recommended in the Cochrane Handbook for Systematic Reviews of Interventions (Chapter 9.5, Deeks 2011), we decided to not report totals from the meta-analysis and to synthesise the studies narratively. Oxford Policy Management 2012 reported that a UCT led to a large reduction in the likelihood of owning any livestock (RR = 0.78, 95% CI = 0.69 to 0.89), indicating potential harm. Pellerano 2014 reported no evidence for an effect of a UCT on the outcome, but the estimate was seriously imprecise and therefore uncertain (RR = 1.06, 95% CI = 0.96 to 1.17). We downgraded this body of evidence to very low quality for serious risk of bias (minus one grade), very serious inconsistency (minus two grades), and serious imprecision (minus one grades). In conclusion, we are very uncertain about the effect of a UCT on livestock ownership.

School attendance

Seven studies (six cluster-RCTs with an effective sample size of 4800 children, plus one CBA) reported an estimate for the effect of a UCT on the likelihood of children attending school at the time of the interview. The cluster-RCTs reported this outcome at 12 to 24 months into the intervention (Baird 2010; Oxford Policy Management 2012; Pellerano 2014; Robertson 2012; Seidenfeld 2013; Ward 2010), using treatment effect estimates not accepted by Cochrane (i.e. an DD estimator or an MDs of a proportion), so we calculated RRs for these studies, conducting approximately correct analyses using the crude numbers reported in the study records (Chapter 16.3, Higgins 2011b). The outcome differed slightly across these six studies with regards to the measurement of attendance (e.g. currently attends, has ever attended, did not miss school last month) and regarding the education institution that the children were enrolled in (any school, preschool, primary school or secondary school). However, we nevertheless considered the studies to be sufficiently homogenous to be combined in one meta-analysis. Moreover, Robertson 2012 reported counts separately for children aged 6 to 12 years and 13 to 17 years, and Ward 2010 reported counts separately for children aged 4 to 5 years and 6 to 17 years. Because neither study counted the same people twice, we combined them in the same meta-analysis. However, for better transparency we report them separately in the forest plot (Figure 7). In relative terms, the point estimate showed a moderate increase in the likelihood of attending school, with the 95% CI suggesting that the effect was between a small and a large increase in the risk (RR = 1.06, 95% CI = 1.03 to 1.09, 6 cluster-RCTs, N = 4800, I² = 0%). In absolute terms, assuming a mean in the control group of 676 attenders per 1000 children (i.e. the median across the six studies), after receiving the UCT an estimated 716 per 1000 children (95% CI 696 to 736) attended school. We judged this considerable level of change to be clinically meaningful.
### Figure 7. Forest plot of comparison: 3 Unconditional cash transfers versus no unconditional cash transfers for improving social determinants of health, outcome: 3.2 Attends school.

<table>
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<th>Risk Ratio</th>
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<td>Overall</td>
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<td>2365</td>
<td>1.06</td>
<td>[1.03, 1.09]</td>
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</tr>
<tr>
<td></td>
<td>Total events</td>
<td></td>
<td>Heterogeneity: Tau² = 0.00, Chi² = 0.07, df = 7 (P = 0.40), I² = 0%</td>
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</tr>
<tr>
<td></td>
<td>Test for overall effect Z = -1.19 (P = 0.0001)</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

#### Estimates
1. Children aged 5-12 years
2. Children aged 13-17 years
3. Children aged 6-17 years
4. Children aged 4 or 5 years

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One CBA in 14,333 children reported an estimate of a UCT on the proportion of children who had dropped out of school (i.e. the inverse of those attending school) (Bazzi 2012). We could not include this study in meta-analysis because it was a CBA, not a cluster-RCT. Because the study reported treatment effects that Cochrane does not accept (i.e. DD estimates of proportions), and since we could not calculate an acceptable treatment effect estimate for this study, we do not report results from this study for this outcome.

We downgraded the body of evidence to moderate quality for serious risk of bias (minus one grade). In conclusion, a UCT probably led to a clinically meaningful, moderate increase in the likelihood of children attending school. Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

### Participation in the labour force

Eight studies (five cluster-RCTs and three CBAs) assessed the effect of a UCT on the likelihood of working at the time of the interview. We considered that three cluster-RCTs with an effective sample size of 4148 participants (2448 children and 1700 adults), reporting outcomes at 24 months into the intervention, were sufficiently homogeneous to be combined in a meta-analysis (Analysis 3.3: Oxford Policy Management 2012; Pellerano 2014; Ward 2010).

Considering that children engaging in child labour and adults working are qualitatively different, we conducted separate analyses by age for children and adults. Ward 2010 reported counts separately for children aged 4 to 5 years and 6 to 17 years. Because neither study counted the same people twice in their analyses, we combined them in the same meta-analysis, but for better transparency we report them separately in the forest plot.

Among children, the point estimate showed that the UCT led to a large reduction in the likelihood of children engaging in child labour, but the 95% CI suggested that the true effect may lie between a very large reduction and a very small increase in the likelihood (RR 0.91, 95% CI 0.80 to 1.03, 3 cluster-RCTs, N = 2449, I² = 0%). Among adults, the point estimate showed that the UCT led to no change in the likelihood of working, with the 95% CI suggesting that the true effect may lie between a small increase and a small reduction in the likelihood of working (RR 1.00, 95% CI 0.95 to 1.06, 2 cluster-RCTs, N = 1700, I² = 13%). Moreover, the two other cluster-RCTs (Haushofer 2013; Seidenfeld 2013), along with the three CBAs (Amarante 2011; Bazzi 2012; Galiani 2014), reported a treatment effect estimate that Cochrane does not accept (i.e. a DD of a proportion), and because we were unable to retrieve acceptable treatment effect estimates or the data to calculate these, we cannot report estimates from these studies. We downgraded this body of evidence for children and for adults to very low quality for serious risk of bias (minus one grade) and very serious imprecision (minus two grades). In conclusion, we are very uncertain about the effect of a UCT on the likelihood of children engaging in child labour and adults working.

### Parenting quality

Three studies (of two cluster-RCTs) assessed the effect of a UCT on the level of parenting quality. Two cluster-RCTs with a sample size of 2314 participants (all adult mothers) reported the standard, validated Home Observation Measurement of the Environment Inventory (HOME) score (see Bradley 1977) at 15 to 27 months into the intervention (Fernald 2011; Paxson 2007). Again, we only used data from Paxson 2007, which reported results for the entire study sample rather than only a selection. Paxson 2007 standardised treatment effect estimates by dividing through 1 SD. We again included the separate estimates for adults from poor families and those from non-poor families separately in the meta-analysis (Analysis 3.4). In relative terms, the point estimate showed a very small increase in the HOME score, but the 95% CI suggested that the effect may be between a small reduction and a small increase in the score (MD 0.09 of 1 SD of the score, 95% CI -0.25 to 0.42, 1 cluster-RCT, N = 1118, I² = 40%). Assuming an SD of 2.30 of the HOME score (i.e. the mean score in the control group before the intervention in the Fernald 2011 study, in the absence of the score from the Paxson 2007 study), the relative effect estimate was an increase in the score by...
0.22 (95% CI −0.60 to 1.01). In absolute terms, assuming a HOME score before the intervention of 2.40 (i.e. the score in the control group before the intervention in the Fernald 2011 study, given the lack of baseline data from Paxson 2007), after the intervention the score was 2.62 (95% CI 1.80 to 3.41). In addition, the Seidenfeld 2013 cluster-RCT of 5670 households reported the effect of a UCT on the likelihood of supporting their child’s learning at 24 months. However, the study reported a treatment effect estimate as a DD of a proportion, so we therefore cannot report an estimate from this study. We downgraded this body of evidence to very low quality for risk of bias (minus one grade) and very serious imprecision (minus two grades). In conclusion, we are very uncertain about the effect of a UCT on the level of parenting quality.

Extreme poverty

Four studies (all cluster-RCTs) with an effective sample size of 2048 households assessed the effect of a UCT on the likelihood of being extremely poor at 24 months into the intervention. Because all four cluster-RCTs reported treatment effect estimates as DD estimates of proportions, we calculated RRs for each study, conducting approximately correct analyses of cluster-RCTs (Chapter 16.3, Higgins 2011b). The included studies used different approaches and definitions of extreme poverty, ranging from ‘living on USD 1 or less per day’ and ‘living below the absolute poverty line’. We nevertheless considered the studies to be sufficiently homogeneous to be combined in one meta-analysis (Analysis 3.5). In relative terms, the point estimate showed that the UCT led to a small reduction in risk, with the 95% CI suggesting that the effect may be between a large reduction in the risk and no change in the risk (RR 0.94, 95% CI 0.89 to 1.00, P = 0.06, 4 cluster-RCTs, N = 2684, I² = 64%). In absolute terms, assuming a mean of 812 people living in extreme poverty per 1000 participants (the median baseline risk in the control group across included studies), after receiving the UCT an estimated 771 per 1000 participants (95% CI 722 to 812) were extremely poor. We downgraded the body of evidence to very low quality for serious risk of bias (minus one grade), serious inconsistency (minus one grade) and serious imprecision (minus one grade). In conclusion, we are very uncertain about the effect of a UCT on the likelihood of being extremely poor.

Healthcare expenditure

Expenditure on healthcare

Six studies (all cluster-RCTs) assessed the effect of a UCT on the amount of money spent on health care over the month prior to the interview, at 7 to 24 months into the intervention. We considered these studies to be too heterogeneous across outcomes to be combined in a meta-analysis. Four cluster-RCTs reported no evidence for an effect large enough to be meaningful, and they were relatively imprecise and uncertain (Cunha 2014; total amount of money spent on medicine and hygiene per month, DD MXN 14.60, 95% CI −5.12 to 34.32, 4923 households; Haushofer 2013; medical expenditure per month, DD USD 0.21, 95% CI −0.08 to 0.50, 1440 households; Oxford Policy Management 2012; mean monthly per capita health expenditure per household, DD KSH 12, 95% CI not reported, P > 0.05, 3107 households; Pellerano 2014; average monthly amount spent on health: DD LSL −1.03, 95% CI not reported, P > 0.05, 3102 households). Two studies reported that a UCT had led to a large increase in the amount of money spent monthly on health care. Seidenfeld 2013 reported that, in relative terms, a UCT had led to a large increase in healthcare expenditure (DD ZMW 1.08, 95% CI not reported, P = 0.05, 2515 households). In absolute terms, assuming a mean expenditure of ZMW 2.60 per month before the intervention (i.e. the amount that all study participants spent before the intervention), after the UCT the healthcare expenditure was ZMW 3.68 per month. We judged this to likely be a clinically meaningful effect. Ward 2010 included 9231 children and reported that in relative terms, a UCT led to a large increase in the mean monthly health expenditure per capita of KSH 17.16 (95% CI not reported, P < 0.05). In absolute terms, assuming a mean monthly expenditure of KSH 48.89 before the intervention (i.e. the amount that the control group spent before the intervention), after the UCT the monthly healthcare expenditure was KSH 66.05. We judged this to likely also be a clinically meaningful benefit. We downgraded this body of evidence to low quality for serious risk of bias (minus one grade) and serious inconsistency (minus one grade). In conclusion, UCT may lead to an increase in the amount of money spent on health care. The rationale for this conclusion was that the two studies that found clinically meaningful effects (out of the total of six studies) were relatively large and well conducted, and the effects were also relatively large. We note that further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Effects on equity in use of health services and health outcomes

None of the included studies estimated the effect of a UCT on measures of absolute or relative inequality in the use of health services or health outcomes, such as the MD or RR of the treatment effect in participants who are disadvantaged versus those who are not. However, several studies reported treatment effect estimates for two or more subgroups defined by population characteristics along the six PROGRESS categories of: age, education, gender, rural-urban residency, income (or poverty status) and marital status. We examined these analyses to indirectly draw conclusions on the effects of UCTs on equity in terms of health services use and health outcomes. If possible, we conducted formal tests of subgroup differences using RevMan 2014. We found that the effects of UCTs on health equity were very uncertain.

Height for age

Two cluster-RCTs reported treatment effect estimates on height for age, disaggregated by dimensions of inequality along PROGRESS. Fernald 2011 reported that a UCT had no effect on the height for age score in participants who resided in rural areas (MD −0.09 of 1 SD, 95% CI −0.36 to 0.18) or in urban area residents (MD 0.13 of 1 SD, 95% CI −0.11 to 0.37), and a test for subgroup differences likewise found no evidence for a statistically significant subgroup differences by rural-urban residency (P = 0.23) (Analysis 4.1). Similarly, Paxson 2007 also found no evidence for any differences (test for subgroup differences: P = 1.00) in treatment effect estimates on the outcome in participants living in poverty (MD 0.04 of 1 SD, 95% CI −0.12 to 0.20) versus those not living in poverty (MD 0.04 of 1 SD, 95% CI −0.06 to 0.14; Analysis 4.2). In summary, this evidence suggests that UCTs perhaps did not have a meaningful effect on health inequalities in height for age by rural/urban residency or income poverty status, but the evidence remains sparse and very uncertain.

Disease or illness

One cluster-RCT reported that a UCT reduced the risk of illness equally in both girls (RR 0.70, 95% CI 0.55 to 0.90) and boys (RR 0.69, 95% CI 0.54 to 0.88) in the two weeks to three months prior to the interview, with a test for subgroup differences finding no evidence...
for statistically significant subgroup differences in this outcome by gender (Pellarino 2014; P = 0.89; Analysis 4.3).

**Food security**

Three cluster-RCTs reported treatment effect estimates disaggregated by dimensions of inequality along PROGRESS. Haushofer 2013 reported very uncertain evidence for whether a UCT had an effect on the food security index among women (DD 0.27 of 1 SD, 95% CI −1.49 to 2.03) and men (DD 0.23 of 1 SD, 95% CI −1.53 to 1.99), and a test for subgroup differences found no evidence for a statistically significant subgroup difference by rural-urban residency (P = 0.97; Analysis 4.4). Oxford Policy Management 2012 and Pellarino 2014 reported separate treatment effect estimates for participants residing versus not residing in income poverty, but they used DD estimates of proportions, which are not acceptable to Cochrane. Moreover, we were not able to source the data required to calculate accepted treatment effect estimates ourselves. This evidence suggested that UCTs may not have impacted inequalities in food security by gender. All of this evidence is, however, very uncertain.

**Dietary diversity**

Four cluster-RCTs reported treatment effect estimates disaggregated by dimensions of inequality along PROGRESS. Fernald 2011 reported very uncertain evidence for whether a UCT had a differential effect on household dietary diversity in participants residing in rural areas (MD 0.20 of 1 SD, 95% CI −0.07 to 0.47) versus urban areas (MD −0.03 of 1 SD, 95% CI −0.30 to 0.24), and a test for subgroup differences found no evidence for a statistically significant subgroup differences by rural-urban residency (P = 0.25; Analysis 4.5). Haushofer 2013 reported that the impact of the UCT on the HDDS among women was a DD estimate of 0.60 food categories (95% CI 0.07 to 1.13), whereas that among men it was a DD estimate of 0.14 food categories (95% CI −0.37 to 0.65; Analysis 4.6). The test for subgroup differences found no statistically significant difference between the treatment effect estimates by gender (P = 0.22). Oxford Policy Management 2012 reported that the impact of the UCT on the HDDS in households living in poverty was a DD estimate of 0.71 food categories (95% CI unclear, P > 0.05), whereas that among households not living in poverty it was a DD estimate of 0.22 food categories (95% CI unclear, P > 0.05). Considering that the 95% CIs of the DD treatment effect estimates were unclear, we were not able to formally test for differences in effect by poverty status in this study. Ward 2010 reported that the impact of the UCT on the HDDS among households living in poverty was a DD estimate of 1.04 food categories (95% CI 1.04 to 1.04), whereas that among households not living in poverty it was a DD estimate of 0.56 food categories (95% CI 0.54 to 0.58; Analysis 4.7). The increase in the HDDS among households living in poverty was larger than that among those not living in poverty (test for subgroup differences: P < 0.001). The baseline scores of the HDDS by poverty status were unclear, but if we assume that the HDDS was lower among the group living in poverty at baseline, which seems like a reasonable assumption, then the UCT reduced inequities in dietary diversity measured using the HDDS. In summary, this body of evidence suggested that UCTs may have reduced inequalities in dietary diversity by income poverty status by improving the outcome more among those living in income poverty than among those not living in poverty. The evidence suggested that UCTs may perhaps not have impacted inequalities in the outcome by rural-urban residency or by gender. All of this evidence is, however, very uncertain.

**Depression**

Three cluster-RCTs assessed the level of depression, as measured with the 60-point CES-D and disaggregated by dimensions of inequality along PROGRESS. Fernald 2011 reported very uncertain evidence for whether a UCT had a differential effect on depression in participants residing in rural (MD 0.26 points, 95% CI −2.01 to 2.53) versus urban areas (MD 1.16 points, 95% CI −1.00 to 3.32), and a test for subgroup differences found no evidence for a statistically significant subgroup differences by rural-urban residency (P = 0.57; Analysis 4.8). Haushofer 2013 reported that the impact of the UCT among women was a DD estimate of −2.44 points on the CES-D (95% CI −4.20 to 0.68), whereas among men it was a DD estimate of −1.15 points (95% CI −2.72 to 0.42; Analysis 4.9). However, the test for subgroup differences found no statistically significant difference by gender (P = 0.28). Paxson 2007 reported evidence for whether a UCT had an effect on the CES-D score among participants living in poverty (MD −0.21 points, 95% CI −0.52 to 0.10) versus not living in poverty (MD 0.00 points, 95% CI −0.18 to 0.18), but a test for subgroup differences found no evidence for a statistically significant difference according to this variable (P = 0.25; Analysis 4.10). The evidence suggested that UCTs may not have impacted inequalities in the outcome by rural-urban residency, gender or income poverty status. This body of evidence, however, remains very uncertain.

**Unconditional cash transfers versus conditional cash transfers for improving use of health services and health outcomes**

None of the three included studies that measured the impact of UCTs versus CCTs reported the impact of UCTs compared to CCTs on growth, death, food security or depression.

**Use of health services**

**Registered birth**

One cluster-RCT with an effective sample size of 239 participants assessed the effectiveness of a UCT versus a CCT on the likelihood of having ever had one’s birth registered, at 2 to 4 months after a 12-month intervention (Robertson 2012). Because the study reported a treatment effect estimate that is not accepted by Cochrane (i.e. MD estimate of a proportion), we calculated an RR for this analysis using approximately correct analyses of cluster-RCTs, using frequency data extracted from the study record (Chapter 16.3, Higgins 2011b). The point estimate showed a large reduction in the likelihood of having had one’s birth registered, with the 95% CI suggesting that the effect may lie between a large reduction and a very small increase in the risk (RR 0.81, 95% CI 0.64 to 1.03). We downgraded the quality of this evidence to very low for serious risk of bias (minus one grade) and very serious impression (minus two grades). In conclusion, we are very uncertain about the effect of a UCT versus a CCT on the likelihood of having ever had one’s birth registered.

**Up-to-date on vaccination calendar**

One cluster-RCT with an effective sample size of 235 participants reported the effectiveness of a UCT versus a CCT on vaccination rates, at 2 to 4 months after a 12-month intervention (Robertson 2012). Because the study reported a treatment effect estimate as an MD estimate of a proportion, we calculated RRs for the study, conducting approximately correct analyses of cluster-RCTs using the crude numbers reported in the study records (Chapter 16.3, Higgins 2011b). The point estimate was no meaningful change in the likelihood of being fully vaccinated, with the 95% CI suggesting that the
effect may lie between a moderate reduction and a moderate increase in the risk (RR 0.99, 95% CI 0.86 to 1.14). We downgraded this evidence to very low quality of evidence for serious risk of bias (minus one grade) and very serious imprecision (minus two grades). In conclusion, we are very uncertain about the effect of a UCT compared with a CCT on the likelihood of being fully vaccinated.

**Use of any health service**

One cluster-RCT in 2559 children assessed the effect of a UCT versus a CCT on the number of routine preventive health service visits made over the previous two weeks to one month, at 8 months after a 24-month intervention (Akresh 2012). The CCT was provided conditional on children aged 0-6 years receiving one growth check at a local health clinic every 3 months and on children aged 7-15 years being enrolled at school and attending school for 90% of the time every quarter year. In relative terms, the point estimate showed a moderate reduction in the number, with the 95% CI suggesting that the effect may lie between a large and a small reduction (MD −0.51 visits, 95% CI −0.83 to −0.19). In absolute terms, assuming a number of 1.02 visits per two-week to one-month period before the intervention (i.e. the number in the group receiving the CCT before the intervention), the number after the intervention was 0.51 visits (95% CI 0.09 to 0.83). We judged this level of change to probably be clinically meaningful. We downgraded the quality of the body of evidence to low for serious risk of bias (minus one grade) and serious indirectness (minus one grade). In conclusion, UCTs may increase the likelihood of having used any health services less than CCTs. Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

**Health outcomes**

**Disease and illness**

One cluster-RCT in 3896 children and young adults assessed the effect of a UCT versus a CCT on the likelihood of having been ill in the previous two weeks, at 12 months into the intervention (Baird 2010). Because the study reported a treatment effect estimate as an MD estimate of proportions, and because we had access to the microdata from this study, we re-analysed these data, calculating an OR adjusted for all the confounders that were adjusted for in the original study (for details see the Methods). The point estimate showed a moderate increase in the odds of having been ill, with the 95% CI suggesting that the effect may lie between a moderate reduction and a large increase in the odds (OR 1.21, 95% CI 0.92 to 1.56). We downgraded this body of evidence to very low quality for serious risk of bias (minus one grade) and very serious imprecision (minus two grades). We are very uncertain about the effect of a UCT compared with a CCT on the likelihood of having had any illness.

**Dietary diversity**

One cluster-RCT in 3896 children and young adults assessed the effect of a UCT versus a CCT on the current level of dietary diversity as measured by the number of times participants ate protein-rich food in the week prior to the interview, at 12 months into the intervention (Baird 2010). The point estimate showed a very small reduction in the number, with the 95% CI suggesting that the effect may be between a large reduction and a large increase in the number (MD −0.06 times per week, 95% CI −0.55 to 0.44 times). We downgraded the quality of this body of evidence to very low for serious risk of bias (minus one grade), serious indirectness (minus one grade), and very serious imprecision (minus two grades). We are very uncertain of the effect of a UCT compared with a CCT on the level of dietary diversity.

**DISCUSSION**

**Summary of main results**

We included 21 studies (16 cluster-RCTs, four CBAs, and one cohort study) involving 1,092,877 participants (36,068 children and 1,056,809 adults) and 31,865 household in Africa, the Americas and South-East Asia and meta-analysed or narratively synthesised the results, although no single outcome was measured by all studies. The studied UCTs were pilot or established government programmes or research experiments. They provided cash of a value equivalent to 1.3% to 53.9% of the annualised gross domestic product per capita. The comparator in all studies was no UCT, and three studies additionally assessed a CCT. Most studies were funded by national governments and/or international organisations.

UCTs compared to no UCT may not have impacted the likelihood of having used any health service in the previous 1 to 12 months, at 12 to 24 months into the intervention. They probably led to a clinically meaningful, very large reduction in the likelihood of having had any illness in previous two weeks to three months, at 12 to 24 months into the intervention. UCTs may have increased the likelihood of having been food secure over the previous month, at 13 to 24 months into the intervention. UCTs may have increased the level of dietary diversity over the previous week, as assessed by the Household Dietary Diversity Score at 24 months into the intervention. Despite several studies providing relevant evidence, the effects of UCTs on the likelihood of being moderately stunted and on the level of depression remain uncertain. No evidence was available on the effect of a UCT on the likelihood of having died. UCTs probably led to a clinically meaningful, moderate increase in the likelihood of currently attending school, at 12 to 24 months into the intervention. The evidence was uncertain for whether UCTs impacted livestock ownership, extreme poverty, participation in child labour, adult employment and parenting quality. UCTs may have increased the amount of money spent on health care at 7 and 24 months into the intervention. The effects of UCTs on health equity were very uncertain. We did not identify any harms from UCTs.

Three cluster-RCTs also reported evidence on the impact of a UCT compared with a CCT on the likelihood of having used any health services, the likelihood of having had any illness, and/or the level of dietary diversity, but this evidence was very uncertain. None of these studies measured the likelihood of stunted growth, death, food security or depression.

**Overall completeness and applicability of evidence**

The current body of evidence is sufficient to address the principal review objective in only a few outcome domains. Existing evidence covers a large number of children and adults in 11 LMICs in three WHO regions. Although UCTs have been introduced in the WHO Africa region relatively recently (Garcia 2012), we positively note the several included studies that evaluate their health effects in this region and especially in Kenya and Malawi. However, more evidence is required for the effects of UCTs in the Eastern Mediterranean, South-East Asia and the Western Pacific. The included studies cover UCTs with a broad range of diverse designs. Evidence on the effects of UCTs on the use of health services should be expanded and improved in quality. This review found evidence that a UCT
may not have impacted the likelihood of a recipient having used any health services, but that, at the same time, a UCT lead to a large increase in the likelihood of a participant having had any illness; studies that estimated the effect of an UCT on health services use generally did not adjust for the prevalence of illnesses, and consequently the evidence may perhaps suggest an increase in health services use per illness. Evidence on health outcomes achieves better coverage of a more diverse set of relevant outcomes from several relevant outcome domains. However, no evidence was available on the outcome of mortality, and more evidence is required for important outcome domains with inconsistent or insufficient evidence, such as stunting, food security, dietary diversity and depression. Some studies have performed a few subgroup analyses along selected PROGRESS categories, predominantly education, gender, geographic residency, and level of income or poverty. More such subgroup analyses are required to more thoroughly and comprehensively determine equity impacts of UCTs, especially along less studied PROGRESS categories such as income. However, studies that estimate the effects of UCTs on measures of absolute and relative inequalities as primary outcomes are ultimately needed to strengthen the evidence on whether and how UCTs influence health equity in LMICs.

Quality of the evidence

For the seven most important prioritised primary outcomes (i.e. outcomes related to the use of health services and health outcomes) and prioritised comparisons of UCTs versus no UCTs, the body of evidence was of moderate quality for one outcome, of low quality for three outcomes, of very low quality for two outcomes, and completely absent for one outcome. For the seven prioritised secondary outcomes of comparisons of UCTs versus no UCTs (i.e. social determinants of health and health expenditure), the evidence was of moderate quality for one outcome, of low quality for one outcome and of very low quality for five outcomes. All three outcomes with data for comparisons of UCTs versus CCTs had very low-quality evidence. Therefore, the current body of evidence supports some conclusions regarding the principal review objective (i.e. for outcomes of moderate or low quality), but it is not possible to draw a conclusion for many other outcomes. Even where evidence is present, there is often still considerable uncertainty around it, and future studies may potentially change our conclusions.

This review included a large number of studies that covered over 1 million participants. The experimental design of most included studies (16 out of 21) was a methodological strength of this body of evidence. However, the review had an overall high risk of bias (especially selection and performance bias, but also attrition bias). In particular, because UCT interventions disburse a visible good (i.e. cash), studies cannot blind participants to these interventions. This is, however, a limitation for reviewing almost all social interventions, and especially those that are disbursed by governments or other public agencies. Studies should publish a priori study protocols so that future reviews can thoroughly assess bias from selective reporting.

The existing evidence was relatively consistent for some outcomes such as the likelihood of having had any illness and of having attended school, but it was highly inconsistent for many others, including the likelihood of having been food secure and the level of household dietary diversity. Estimated treatment effects showed acceptable or even good precision for the small number of outcomes that had evidence from several cluster-RCTs, relatively common events and relatively large (effective) sample sizes. However, for most outcomes, treatment effects were still relatively imprecise.

Potential biases in the review process

We applied the ROBIS tool to assess potential biases in our review process (Whiting 2016). One concern regarding study eligibility criteria was that we had to further specify some of our pre-defined eligibility criteria, and in some rare instances – make changes to our pre-defined criteria (see Differences between protocol and review). The reason was that when we conducted the review, we realised that we had insufficiently or incorrectly specified a small number of criteria in our protocol (Pega 2014). However, we are confident that the eligibility criteria that we applied were appropriate for the review question and unambiguous, and that restrictions in eligibility criteria based on study characteristics and information sources were also appropriate. While we acknowledge that the few and relatively minor differences in inclusion and exclusion criteria between protocol and review may have potentially introduced some bias in the review process, overall we judged this to be a minor concern, with low risk of having introduced noteworthy bias.

We have some confidence in our selection and identification of studies. Our search included an appropriate and broad range of databases for published and unpublished study records, and we employed additional methods to identify relevant records. The terms and structure of the search strategy should have retrieved as many eligible studies as feasible, and our search had no restriction based on date, publication format or language. An independent reference librarian, who is not an author of this review, conducted all academic and several of the grey literature database searches. However, we note that almost half of the studies included in this review were published in inaccessible grey literature – generally discussion or working papers written by economists or reports to governmental, international or nongovernmental organisations prepared by private consultancy firms. Considering the inaccessibility of this literature, it is possible and perhaps even likely that the review missed eligible studies. Another potential source of bias in the review is that we were unable to source some missing data for several studies, and if these missing data were not missing at random, then they may have introduced bias in this review. Also, we excluded studies that did not report a primary outcome in their study record but did not check whether the authors collected (but did not report) eligible outcomes. If authors systematically did not report treatment effect estimates for eligible outcomes

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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for which they did not find desirable effects, then our review would have missed these results, and this could perhaps have introduced bias in this review. Despite acknowledging these caveats, we nevertheless judged these concerns overall to be more likely indicative of low risk in our review process.

We believe that we made all possible efforts to minimise errors in our data collection and that sufficient study characteristics are probably available to us and to the readers to enable interpretation of the results of the review. Our risk of bias assessment applied standard Cochrane tools and relied on independent assessments from two or more review authors, and it should thus have been appropriate and reduced errors whenever possible. Given the very large number of potentially relevant study results reported in some study records, we cannot be absolutely certain that we were able to collect all relevant results for use in the synthesis, but this is probably of low concern.

Our synthesis included all appropriate studies, and we reported all pre-defined analyses or explained departures from them. Generally speaking, we believe the synthesis was appropriate given the nature and similarity in the research questions, study designs, and outcomes across included studies. One exception is that we combined UCTs in meta-analyses that varied considerably in their generosity (i.e. between 1.3% and 53.9% of GDP per capita), which may well have introduced heterogeneity in such meta-analyses. However, research suggests that higher amounts of cash transfer do not always lead to stronger effects on social outcomes (Baird 2011; Filmer 2011). Between-study variation (heterogeneity) was sometimes large or even very large, but we carefully addressed high levels of heterogeneity in the synthesis where they were observed, for example by not reporting totals from meta-analyses and instead synthesising the evidence narratively. The relatively small number of studies per meta-analysis prohibited us from being able to assess the robustness of our findings through funnel plotting, sensitivity analyses or similar other analyses. Because several of the included studies used erroneous analytic methods and treatment effects (e.g. DD estimates of dichotomous outcomes that are not defined in health research), we were forced to conduct approximately correct analyses from crude frequency measures and produce crude treatment effects for almost all cluster-RCTs. That we combined these crude treatment effects with each other or with adjusted treatment effects may have introduced risk of confounding in the pooled treatment effects. Several included studies had a high risk of bias; however, we attempted to address this in the synthesis wherever feasible. Overall, we nevertheless judged these potential and actual issues to probably cause low concerns, but we cannot fully preclude high concerns.

In judging the potential risk of bias in the review process based on the above described and quantified concerns, we believe that our interpretation of the findings probably addressed most, if not all, of the identified concerns. We believe that we considered the relevance of the included studies to the review’s research question appropriately, and we avoided emphasising results on the basis of their statistical significance. In conclusion, we therefore judge the potential risk of bias in the review process to probably be low overall, with a few caveats that we have here acknowledged openly.

Agreements and disagreements with other studies or reviews

We are not aware of any previous reviews that have synthesised evidence specifically on the effects of UCTs intended to reduce poverty and vulnerabilities on the use of health services and health outcomes. Our review findings confirm those of a previous systematic review that UCTs improve schooling outcomes in LMICs (Baird 2013). Previous reviews examining cash transfers and their effect on the use of health services and/or health outcomes in LMICs generally included either CCTs only (Gaarder 2010; Lagarde 2009; Pega 2013), or a broader set of cash transfers that combined UCTs with CCTs and sometimes even also with other financial interventions, such as microfinancing interventions (Adato 2009; Arnold 2011; Bassani 2013; Boccia 2012; Heise 2013; Manley 2013; Pettifor 2012; Sridhar 2006). They also generally included a broader set of study types, including designs that Cochrane Reviews generally exclude due to their high risk of bias, such as cross-sectional studies. For these reasons the findings of these previous reviews are not comparable with those of this review.

Authors’ conclusions

Implications for practice

The existing body of evidence, which is based on several cluster-RCTs, suggests that UCTs have probably had a large, clinical meaningful, beneficial effect on the likelihood of having had any illness and a moderate, probably clinically meaningful, beneficial effect on the likelihood of having attended school. UCTs may have also had a beneficial effect on food security, dietary diversity and the amount of money spent on healthcare, but they may not have impacted the likelihood of having used any health service. The evidence remains uncertain for the effect of UCTs on stunting, depression, livestock ownership, participation in child labour, adult employment, extreme poverty and parenting quality. We did not identify any harms of UCTs. The effect of UCTs on health equity are very uncertain. The relative effectiveness of UCTs compared with CCTs also remains uncertain with regard to the level of dietary diversity as well as the likelihood of having used any health service and of having had any illness, and no studies measured other relevant outcomes for this comparison.

Implications for research

More evidence from experimental studies is required to improve this currently still limited and overall still relatively uncertain body of evidence. RCTs of individual participants would be preferable over RCTs of clusters of individuals. However, we acknowledge that randomising individual participants may not always overcome the challenge of contamination, and cluster-RCTs are therefore likely to continue to be the dominant study design of choice. All future experimental studies of the impact of UCTs on health or health-related outcomes should always publish comprehensive a priori study protocols, both to reduce risk of bias from selective reporting in the study itself and to enable systematic reviewers to judge the risk of reporting bias. Future experimental studies should also comprehensively assess contamination (e.g. by using spill-over control groups) to reduce risk of bias from contamination. Similarly, study records should improve reporting of blinding of participants, study personnel and outcome assessors to enable robust assessment of likelihood of performance and detection bias.
The use of robust analytical methods is central to further advancing the existing body of evidence. If economists who conduct studies of the effect of UCTs on health services use, health outcomes, social determinants of health, healthcare expenditure or health equity want to contribute to health research and policy development, they should consider applying standard methods that are widely accepted amongst health researchers and epidemiologists, as defined by the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011a). They should avoid using DD estimates of proportions, which are not defined in health research and therefore not accepted by Cochrane but are currently very commonly used by economists conducting research on health effects of UCTs. Studies should focus on investigating impacts of UCTs on the most important outcomes, rather than on less important outcomes. Since the current body of evidence is primarily for the Americas and Africa, more research is particularly also needed for the Eastern Mediterranean, South-East Asian and Western Pacific regions. More high-quality experimental studies that determine the impacts of UCTs on equity in use of health services and health outcomes along PROGRESS categories are also needed. Strengthening the currently small body of evidence on the relative effectiveness of UCTs and CCTs requires more high-quality experimental studies that compare UCTs with CCTs in terms of their effects on health service use, health outcomes, social determinants of health, healthcare expenditure and health equity.

The effectiveness of UCTs for improving health service use and health outcomes may be related to the relative amount of the transfer, either in terms of mean population income, the income of the poor or a related measure; and/or in terms of the costs related to the outcome of interest, such as the costs of depression. Future systematic reviews should also seek to record this information on transfer magnitudes (as done in this review), which they could use as a mediator in the meta-analyses. Moreover, the relative effectiveness of UCT and CCT programmes may be, for select outcomes such as healthcare utilisation, related to the actual imposed conditions of the CCT programme. Future systematic reviews should therefore also always record the conditions of the included CCTs (as done in this review), and they could include this information in their evaluation of the relative effectiveness of UCTs versus CCTs.

**Acknowledgements**

We thank: Jodie Doyle, Miranda Cumpston, Dr Rob Anderson, Patrick Condon, Dr Anke Rohwer, Dr Reza Yousefi-Nooraei and the late Dr Elizabeth Waters (all Cochrane Public Health Group) for editorial guidance and advice on this review; Dr Paul Bain (Harvard Medical School) for searching most electronic academic and some grey literature databases; Ruth Turley (Cochrane Public Health Group) for searching the Cochrane Public Health Specialised Register; Susan Hope (University of Otago) for contributing to handsearching of academic journals; Carolin Henning and Tatjana Paeck (both University of Bremen) for contributing to data extraction; Meggan Harris for copy-editing this review; Dr Rosangela Bando Grana (Inter-American Development Bank), Dr Samuel Bazzi (Boston University), Girija Bahety (Oxford Policy Management), Dr Jesse Cunha (Naval Postgraduate School), Dr Jeffrey Eaton (Imperial College London), Dr Lia C. Haskin Fernald (University of California, Berkeley), Dr Sebastian Galiani (University of Maryland), Martina Garcia (Oxford Policy Management), Dr Paul Gertler (University of California, Berkeley), Dr Simon Gregson (Imperial College London), Dr Sudhanshu Handa (University of North Carolina, Chapel Hill), Dr Melissa Hidrobo (International Food Policy Research Institute), Maja Jakobsen (Oxford Policy Management), Dr Winnie Luseno (Pacific Institute for Research and Evaluation), Dr Fred Mertens (Oxford Policy Management), Dr Candace Miller (Mathematica Policy Research), Dr Tia Palermo (United Nations Children’s Fund), Dr Luca Pellerano (Oxford Policy Management), Dr Norbert Schady (Inter-American Development Bank), Dr David Seidenfeld (American Institutes for Research) and Dr Patrick Ward (Oxford Policy Mangement) for providing missing data for included studies; Gordon Purdie and Dr James Stanley (both University of Otago) for statistical advice; Dr Sarah Baird (University of Washington), Aneil Jaswal (University of Oxford), Dr Vanessa Jordan (Cochrane New Zealand), Dr Santosh Mehrotra (Planning Commission, Government of India), and Dr Luca Pellerano (Oxford Policy Mangement) for their advice; and Dr Jed Friedman (World Bank), Kirti S Sahu (Public Health Foundation of India), and Dr Mesfin G Zbel (World Health Organization) for their external peer review.
References to studies included in this review

Agüero 2007 {published data only}


Akresh 2012 {published data only}


Amarante 2011 {published data only}


Baird 2010 {published data only}


Bazzi 2012 {published and unpublished data}


Beck 2015 {published data only}


Cunha 2014 {published and unpublished data}


Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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Fernald 2011 (published and unpublished data)


Galiani 2014 (published and unpublished data)


Haushofer 2013 (published data only)


Leroy 2010 (published and unpublished data)

Leroy JL, Gadsden P, Rodriguez-Ramirez S, de Cossio TG. Cash and in-kind transfers in poor rural communities in Mexico increase household fruit, vegetable, and micronutrient consumption but also lead to excess energy consumption. *Journal of Nutrition* 2010;140(3):612-7. [DOI: 10.3945/jn.110.116285]

* Leroy JL, Gadsden P, de Cossio TG, Gertler P. Cash and in-kind transfers lead to excess weight gain in a population of women with a high prevalence of overweight in rural Mexico. *Journal of Nutrition* 2013;143(3):378-83. [DOI: 10.3945/jn.112.167627]

Luseno 2012 (published and unpublished data)


Miller 2008 (published and unpublished data)


Oxford Policy Management 2012 (published and unpublished data)


Paxson 2007 (published data only)


Pellerano 2014 (published and unpublished data)

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

References to studies excluded from this review

Akee 2013 (published data only)


Aker 2011 (published data only)

Aker 2013 (published data only)


Angelucci 2009 (published data only)

Attanasio 2015 (published data only)

Ayuku 2013 (published data only)

Benhassine 2013 (published data only)


Buller 2016 (published data only)
Buller AM, Hidrobo M, Peterman A, Heise L. The way to a man's heart is through his stomach? A mixed methods study on causal mechanisms through which cash and in-kind food transfers decreased intimate partner violence. BMC Public Health 2016;16:488. [DOI: 10.1186/s12889-016-3129-3]

Buser 2014 (published data only)

Coetzee 2013 (published data only)

Doocy 2017 (published data only)

Fenn 2013 (published data only)


Hidrobo 2013 (published data only)

Holmquist 2011 (published data only)

Kenya CT-OVC Evaluation Team 2012 (published data only)

Langendorf 2013 (published data only)


Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

Macours 2008 *published data only*


Ndlovu 2013 *published data only*


Park 2013 *published data only*


Pereznieto 2014 *published data only*


Plagner 2011 *published data only*


Poulse 2011 *published data only*


Pratinidhi 2014 *published data only*


Rocha 2013 *published data only*


Santos 2011 *published data only*


Skoufias 2013 *published data only*


Skovdal 2012 *published data only*


Tadesse 2014 *published data only*


Tonguet-Papucci 2017 *published data only*


References to studies awaiting assessment

Abdoulayi 2014 *published data only*


AIR 2014 *published data only*


Benedetti 2016 *published data only*

Benedetti F, Ibarrarán P, McEwan PJ. Do education and health conditions matter in a large cash transfer? Evidence from a
Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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**Brugh 2016 (published data only)**


**Cluver 2013 (published data only)**


**Davis 2016 (published data only)**


**Gangophadyay 2015 (published data only)**


**Grellety 2017 (published data only)**


**Handa 2014a (published and unpublished data)**


**Huang 2014**


**Hjelm 2017 (published data only)**


**Killburn 2016 (published data only)**


**Lawlor 2015 (published data only)**

References to ongoing studies

Galárraga 2014 [published data only]


O’Leary 2011 [published data only]

Oxford Policy Management 2013 [published data only]

Additional references

Adato 2009

Alvaredo 2013

Arnold 2011

Baird 2011

Baird 2012

Baird 2013

Barrientos 2006

Barrientos 2008

Bassani 2013

Benach 2010a

Benach 2010b

Boccià 2012
Middle-income countries (Review)

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

Borjas 2013

Bradley 1977

Centre for Reviews and Dissemination 2009

Coates 2007

Cochrane EPOC 2012

Cochrane PHG 2011

CSDH 2008

Deeks 2011

Diez-Roux 2000

Dimick 2014

Doocy 2016

Eaton 2004

Filmer 2011

Fiszbein 2009

Forde 2012

Friedman 2014

Gaarder 2010

Gaarder 2012

Garcia 2012

GBD 2016

Global Humanitarian Assistance 2012
Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)
Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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Characteristics of included studies [ordered by study ID]

**Agüero 2007**

**Methods**
Cohort study, regression analysis with propensity score matching methods, 12 years: 1993-2004

**Participants**
720 children (aged 0-36 months) interviewed over 3 waves of the KwaZulu-Natal Income Dynamics Study (waves collected in 1993, 1998 and 2004), KwaZulu-Natal province, South Africa

**Interventions**

- **Exposure**

  1. Received an unconditional cash transfer (UCT): 245 participants
  2. Did not receive a UCT: 475 participants

  Duration: up to 36 months. Follow-up: up to 36 months into the intervention. Intervention design: aimed to reduce poverty among children in poor families; targeted to children living in poor house-
### Agüero 2007 (Continued)

holds; provided a total amount of up to USD 900 (USD 25 per month for 36 months; 4.8% of the annual gross domestic product (GDP) per capita).

### Outcomes

**Primary outcomes**

- Health outcomes
  - Moderate stunting (measure: current height for age)

### Notes


### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>High risk</td>
<td>Sampling strategy was random for the first wave, but a non-random sub-sample of the study sample was re-interviewed at waves 2 and 3. Nationally representative sample not achieved. High risk of selection bias from likely self-selection into the intervention by some eligible participants (e.g., those believing they may profit considerably from the intervention), but not other people (e.g., those not believing they may profit considerably from the intervention).</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation concealment among participants and study personnel unclear</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Low risk</td>
<td>Not a cluster-RCT</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Unclear risk</td>
<td>No overall P values, test statistics or SDs reported</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>Unclear risk</td>
<td>No overall P values, test statistics or SDs reported. Unexposed group potentially differed systematically from exposed group in terms of motivation of adopting the intervention.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Participants not allocated to the intervention by the researchers. Secondary analysis of survey data collected for a different purpose than estimating the effect of the UCT on the use of health services and health outcomes. Therefore, blinding of participants neither feasible nor necessary. Blinding of study personnel unclear.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>No subjectively measured outcome in this study</td>
</tr>
</tbody>
</table>

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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### Agüero 2007 (Continued)

<table>
<thead>
<tr>
<th>Source</th>
<th>Risk of Bias</th>
<th>Reason</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contamination</td>
<td>High risk</td>
<td>Allocation was by household, but additional income from the UCT provided to participants in the exposed group may have been transferred to participants in the comparator group (e.g. between family members)</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Survey non-response unclear. Attrition high (29%) and unclear whether differential by population characteristics and/or outcomes.</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>We did not identify a study protocol</td>
</tr>
<tr>
<td>Other bias</td>
<td>High risk</td>
<td>Misclassification bias of the exposure: used self-reported receipt of a UCT collected in surveys, but we considered the risk of this bias to be low. Confounding: this study did not adjust for several potential time-invariant and time-varying confounding variables, such as caregiver’s motivation, caregiver’s cognitive abilities, and changes over time in health services access, fertility and income. We judged this to carry a high risk of confounding. Reverse causation: the outcome may have impacted the exposure, but we considered the risk of this bias to be low.</td>
</tr>
</tbody>
</table>

### Akresh 2012

**Methods**

Cluster-randomised controlled trial (3 stages: first stage, 75 villages with a primary school were randomly assigned to 4 intervention groups and 1 control group in a public meeting; second stage, in the 60 intervention villages the cash transfer intervention assigned to the village was randomly assigned to poor households in a public meeting; third stage, in the 15 control villages poor households were randomly sampled), regression analytic methods, 24 months in 2008-2010

**Participants**

2559 children (aged 0-59 months) interviewed 3 times (baseline: June 2008; follow-up 1: June 2009; follow-up 2: June 2010); Nahour province, Burkina Faso

**Interventions**

4 intervention groups and 1 control group

1. An unconditional cash transfer (UCT) given to the mother: 15 villages, approximately 540 households, unclear number of participants (included in this review as intervention group, combined with the UCT given to the father)
2. UCT given to the father: 15 villages, approximately 540 households, unclear number of participants (included in this review as intervention group, combined with the UCT given to the mother);
3. Conditional cash transfer (CCT) given to the mother: 15 villages, approximately 540 households, unclear number of participants (included in this review as CCT comparison group, combined with the CCT given to the father)
4. CCT given to the father: 15 villages, approximately 540 households, unclear number of participants (included in this review as CCT comparison group, combined with the CCT given to the mother)
5. No cash transfer (i.e. pure control group): 15 villages, 615 households, unclear number of participants (included in this review as control group)

Intervention duration: 24 months. Follow-up: 8 months after 24 months of the intervention. Intervention design: aimed to improve health outcomes among children aged 0-6 years and education outcomes among children aged 7-15 years; targeted to children aged 0-15 years who resided in poor households; provided a total amount of USD 19.20 per child aged 0-6 years (USD 0.80 per month for 24 months; approximately 1.3% of the annual GDP per capita), USD 38.64 per child aged 7-10 years or in grade 1-4 (USD 1.61 per month for 24 months; approximately 1.5% of the annual GDP per capita) and USD 77.04 per child aged 11-15 years or in grade 5 or higher (USD 3.21 per month for 24 months; approximately 2.8% of the annual GDP per capita); paid in-hand to the mother or the father every 3 months; CCTs conditional on children aged 0-6 years receiving one growth check at a local health clinic every 3 months and on children aged 7-15 years being enrolled at school and attending school for 90%
Outcomes

Primary outcomes

- Health services use
  * Use of any health service in previous 2 weeks to 1 month (measure: number of routine preventive health clinic visits, last week)

Notes


<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Randomisation ensured</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Allocation was by public lottery and therefore concealed</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>High risk</td>
<td>Participants recruited after cluster allocation</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Unclear risk</td>
<td>No baseline data reported that could be used to test for baseline differences in the outcome measurement between the UCT intervention group, the pure control group, and the CCT comparison group. Baseline differences (if any) in the outcome measurement appear to not have been adjusted for.</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>Low risk</td>
<td>Baseline differences (P &lt; 0.05) reported between the UCT given to women intervention group and the pure control group in one characteristic. The households were larger in the UCT intervention group than in the control group (6.90 compared with 6.05, P &lt; 0.05). However, all baseline differences were comprehensively adjusted for using regression analytic methods. In addition, messaging to health administrators in villages in the UCT intervention group and especially the CCT comparison group may have occurred, but such messaging may not equally have occurred in villages in the control group, and this may have influenced the outcome. However, we judged the risk that this may have introduced confounding to be low.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>Blinding of participants not feasible, and blinding of personnel unclear</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All subjectively measured outcomes</td>
<td>High risk</td>
<td>Outcome assessors not blinded</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All objectively measured outcomes</td>
<td>Low risk</td>
<td>No objectively measured outcome in this study</td>
</tr>
</tbody>
</table>
### Akresh 2012 (Continued)

<table>
<thead>
<tr>
<th>Contamination</th>
<th>High risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assignment was by village, but additional income from UCTs provided to participants in the UCT intervention groups or the CCT comparison groups may have spilled over to participants in other UCT intervention groups, the control group, and/or the CCT comparison groups (e.g. between family members). Spill-over effects were not investigated (e.g. no spill-over control group in this study).</td>
<td></td>
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</table>

<table>
<thead>
<tr>
<th>Incomplete outcome data (attrition bias)</th>
<th>Low risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial survey non-response unclear. Attrition low (4%). No tests reported for differences in outcome measurements between participants lost to follow-up and those not lost to follow-up in the intervention groups, compared with those differences in the pure control group and the CCT comparison groups. Differences in 6 population characteristics reported between participants lost to follow-up and those not lost to follow-up in the intervention groups, compared with in the control group. Attritting households had fewer adults (P &lt; 0.01), were smaller (P &lt; 0.01), had younger household heads (P &lt; 0.05), were more likely to be Christian (P &lt; 0.01), and were less likely to be Animist (P &lt; 0.01). The numbers of missing participants per UCT intervention group, control group and CCT comparison group and for the outcome were unclear.</td>
<td></td>
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</table>

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<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Unclear risk</th>
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<tbody>
<tr>
<td>We did not identify a study protocol</td>
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</table>

<table>
<thead>
<tr>
<th>Other bias</th>
<th>Low risk</th>
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</thead>
<tbody>
<tr>
<td>None identified</td>
<td></td>
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</table>

### Amarante 2011

<table>
<thead>
<tr>
<th>Methods</th>
<th>Controlled before-and-after study, difference-in-differences methods, 59 months in 2003-2007</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>1,037,739 adults reported in 5 years of administrative data linked to vital statistics (before: January 2003-March 2005; after: April 2005-December 2007), women, Uruguay</td>
</tr>
<tr>
<td>Interventions</td>
<td>1 exposed group and 1 unexposed group</td>
</tr>
<tr>
<td>1. Received UCT: 50,939 participants (included in this review as the exposed group)</td>
<td></td>
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<tr>
<td>2. Did not receive a UCT: 20,872 participants (included in this review as unexposed group)</td>
<td></td>
</tr>
<tr>
<td>Duration: 1-32 months. Follow-up: 1-32 months into the intervention. Intervention design: aimed to reduce poverty; targeted to households with an income score predicted by government personnel to fall below a pre-determined level; provided a total amount of up to USD 1792 (USD 56 per month for 32 months; approximately 7.0% of the annual GDP per capita); co-intervention: from mid-2006 an electronic food card with a monthly value of USD 13.30 (1.3% of the annual GDP per capita) or one-fourth to one-half of the value of the UCT, depending on household size and demographic structure; and fuzzy design: conditional on pregnant women's and children's regular health checks and on children's school attendance (but the conditions were not enforced)</td>
<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Primary outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Underweight</td>
<td></td>
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<tr>
<td>Low birthweight</td>
<td></td>
</tr>
<tr>
<td>Maternal weight at week 35 of pregnancy</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Secondary outcomes</th>
<th>Employment (measure: adult worked during pregnancy)</th>
</tr>
</thead>
</table>

Alternative primary outcome measures not reported in this review
Notes

Intervention context: Plan de Atención Nacional a la Emergencia Social, implemented by the Government of Uruguay; covered 14% of the population; unclear population uptake; and the total annual cost of the UCT programme was approximately USD 250 million. Funder of the study: the Inter-American Development Bank. Conflict of interest: none identified.

Risk of bias

Bias | Authors' judgement | Support for judgement |
--- | --- | --- |
Random sequence generation (selection bias) | High risk | Sampling was not random. Rather, administrative records and vital statistics were used, the former of which were collected specifically to determine eligibility for the UCT. Multiple baseline differences between the exposed group and the unexposed group. |
Allocation concealment (selection bias) | Unclear risk | Allocation concealment among participants and study personnel unclear |
Participants recruited after cluster allocation (recruitment bias in cluster-RCTs) | Low risk | Not a cluster-RCT |
Baseline outcome measurements similar | Unclear risk | No confidence interval, P value or test statistic reported |
Baseline characteristics similar | Unclear risk | No confidence interval, P value or test statistic reported |
Blinding of participants and personnel (performance bias) All outcomes | Low risk | Participants not allocated to the intervention by the researchers. Secondary analysis of survey data collected for a different purpose than estimating the effect of the UCT on health services use and health outcomes. Therefore, blinding of outcome assessors was neither feasible nor necessary. |
Blinding of outcome assessment (detection bias) All subjectively measured outcomes | Low risk | Participants not allocated to the intervention by the researchers. Secondary analysis of survey data collected for a different purpose than estimating the effect of the UCT on health services use and health outcomes. Therefore, blinding of outcome assessors was neither feasible nor necessary. |
Blinding of outcome assessment (detection bias) All objectively measured outcomes | Low risk | Participants not allocated to the intervention by the researchers. Secondary analysis of survey data collected for a different purpose than estimating the effect of the UCT on health services use and health outcomes. Therefore, blinding of outcome assessors was neither feasible nor necessary. |
Contamination | Unclear risk | Allocation was by household, but additional income from UCTs provided to participants in the exposed group may have spilled over to participants in the comparator group (e.g. between family members) |
Incomplete outcome data (attrition bias) All outcomes | Unclear risk | The initial sample size and survey non-response were unclear. For the birthweight and social determinants of health outcomes, the complete sample was 71,811 participants. For mother’s weight, the sample size was 21,944 participants. Attrition was unclear, since the initial sample size was not reported. Numbers of missing participants per outcome were unclear. Number of obser-
vations (68,858) were reported only for low birth weight. Matching between data source could only be performed with a non-quantifiable error.

<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Unclear risk</th>
<th>We did not identify a study protocol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>High risk</td>
<td>Misclassification bias of the exposure: the study used self-reported receipt of a UCT collected in surveys, but we considered the risk of this bias to be low. Confounding: this study did not adjust comprehensively for all potential confounders, and it used difference-in-differences methods, which carry a risk of bias if the underlying time trends differ between the exposed group and the unexposed group. We therefore judged the risk of confounding to be high. Reverse causation: the outcome may have impacted the exposure, but we considered the risk of this bias to be low.</td>
</tr>
</tbody>
</table>

**Amarante 2011 (Continued)**

**Baird 2010**

**Methods**
Cluster-randomised controlled trial (3 stages: first stage, enumeration area were divided into 3 strata of geographic residency (i.e. within city, within 16 km radius of city, and far rural); second stage, enumeration areas were sampled for each strata (i.e. 29, 119, and 28, respectively); third stage, never-married females aged 13-22 years were randomly sampled from a list of all females within each strata), intention-to-treat analysis, difference-in-differences and regression analytic methods, 32 months in 2007-2010

**Participants**
3896 children or young adults (aged 13-22 years) who were never married (split into 2907 in school and 889 out of school/dropouts), interviewed 3 times (baseline: October 2007-January 2008, follow-up 1: October 2008-February 2009, follow-up 2: February-June 2010), all girls or young women, 176 enumeration areas, Zomba district, Malawi

**Interventions**
Girls and young women in school at baseline
2 intervention groups and 3 control groups
1. UCT: 27 enumeration areas, 283 participants (included in this review as intervention group)
2. No cash transfer (spill-over control group for UCT): 27 enumeration areas, 179 participants (excluded from this review)
3. CCT: 46 enumeration areas, 506 participants (included in this review as CCT comparison group)
4. No cash transfer (spill-over control group for CCT): 46 enumeration areas, 243 participants (excluded from this review)
5. No cash transfer (pure control group): 88 enumeration areas, 1495 participants (included in this review as control group)

Girls and young women out of school/dropouts at baseline
1 intervention group and 1 control group
1. CCT: 88 enumeration areas (including the 15 treatment areas where baseline schoolgirls were not given a cash transfer), 436 participants (included in this review as CCT comparison group)
2. Control group (pure control group): enumeration 88 areas, 456 participants (included in this review as control group)

Intervention duration: 24 months. Follow-up: 12 months into the intervention (alternative follow-up not reported in this review: 2-6 months after 24 months of the intervention). Intervention design: aimed to determine the effectiveness of UCT and CCT (in the baseline school girls group); UCT targeted to never married girls or young women aged 13-22 years from poor households; UCT provided a total amount of USD 96, USD 144, USD 192 or USD 240 to parents randomly by enumeration area so that all parents in the same enumeration area received the same amount (USD 4, USD 6, USD 8 or USD 10 per month for 24 months; approximately 8.4%, 12.5%, 16.7% or 20.9% of the annual GDP per capita) and USD 24,
USD 48, USD 72, USD 96 or USD 120 to girls or young women randomly by individual through an open public lottery, so that different girls and young women within the same enumeration area received different amounts (USD 1, USD 2, USD 3, USD 4 or USD 5 per month for 24 months; approximately 2.1%, 4.2%, 6.3%, 8.4% or 10.5% of the annual GDP per capita), and an amount equivalent to the average annual amount given to the CCT intervention group towards school fees; paid in-hand each month; CCT provided the same total amounts as the UCT to parents and girls using the same randomisation procedures; co-intervention for CCT: school fees were paid directly to the school for girls and young women enrolled in school; the CCT was conditional on regular school attendance (i.e. 80% or more of all school days attended); adherence with the condition was monitored, and non-adherence was punished (i.e. the CCT was for the following month was withheld).

### Outcomes

**Primary outcome**
- Has had any illness
- Level of dietary diversity (measure: number of days had eaten protein-rich food, last week)

**Secondary outcomes**
- School attendance (current)

**Alternative primary outcome measures not reported in this review**
- Has had any illness*
  - Has herpes simplex type 2
  - Has syphilis
  - Has HIV

### Notes

Intervention context: experiment by research organisations (the National Bureau of Economic Research), but also appears to be a programme of the Government of Malawi called the Zomba Cash Transfer; unclear who implemented the experiment; unclear population coverage, intervention uptake, and total programme costs. Funder of the study: Bill and Melinda Gates Foundation, Global Development Network, International Initiative for Impact Evaluation, National Bureau of Economic Research, and the World Bank. Conflict of interest: some study records were co-authored by staff of organisations that funded the study.

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Randomisation was ensured</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Allocation was by public lottery and therefore protected</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Low risk</td>
<td>Participants were recruited before clusters were allocated</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) reported between the UCT intervention group and the control groups (all 3 control groups among school girls pooled) in one outcome measurement (has had any illness). No baseline data reported that could be used to test for baseline differences between the UCT intervention group and the control groups in one outcome measurement (number of days had eaten protein rich food). No baseline differences reported between the UCT intervention group and the CCT comparison group in any of the outcome measurements. However, all baseline differences in outcomes measurements appear to have been comprehensively adjusted for using regression analytic methods.</td>
</tr>
</tbody>
</table>
### Baseline characteristics

Baseline differences ($P < 0.05$) reported between the UCT intervention group and the control groups (all 3 control groups pooled) in 5 characteristics. The highest grade attended by respondents in the UCT intervention group was higher than that in the control groups (7.90 compared with 7.48, $P < 0.05$). The proportion of participants in female-headed household in the UCT intervention group was lower than that in the control group (24% compared with 32%, no test reported). The proportion of participants in households that owned a radio in the UCT intervention group was higher than in the control groups (65% compared with 59%, no test reported). The proportion of participants in households that owned a television in the UCT intervention group was higher than in the control groups (34% compared with 24%, no test reported). The proportion of participants who had piped water available in their dwelling in the UCT intervention group was higher than that in the control groups (60% compared with 47%, no test reported). Baseline differences ($P < 0.05$) also reported between the UCT intervention group and the CCT comparison group in 2 characteristics. The age in the UCT intervention group was higher than in the CCT comparison group (15.43 compared with 14.95, $P < 0.01$). The highest grade attended by respondents in the UCT intervention group was higher than that in the CCT comparison group (7.90 compared with 7.25, $P < 0.01$). However, all baseline differences in characteristics appear to have been comprehensively adjusted for using regression analytic methods.

### Blinding of participants and personnel (performance bias)

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Low risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding of participants was not possible, and blinding of personnel was unclear. Qualitative interviews conducted as part of the study suggested that the UCT intervention group was aware of the existence of a CCT comparison group and that the purpose of the cash transfer programme was to improve education. We therefore considered the risk of performance bias to be high.</td>
<td></td>
</tr>
</tbody>
</table>

### Blinding of outcome assessment (detection bias)

<table>
<thead>
<tr>
<th>All subjectively measured outcomes</th>
<th>Unclear risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding for subjectively measured outcomes was unclear</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>All objectively measured outcomes</th>
<th>Low risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>There were no objectively measured outcome in this study</td>
<td></td>
</tr>
</tbody>
</table>

### Contamination

| Low risk |
| Allocation was by community and the study assessed, but we did not find any evidence of contamination |

### Incomplete outcome data (attrition bias)

| Low risk |
| Initial survey non-response was low (6%). Attrition rate was moderate (10%) and non-differential between the intervention group and the control groups. One area was lost amongst the baseline school girls for an unclear reason. For each group, the number of missing clusters and number and percentage of missing participants were: |

Girls and young women in school at baseline:

- 2 intervention groups and 3 control groups

1. UCT: 0 clusters missing, 23 participants missing (8%)
2. No cash transfer (spill-over control group for UCT): 0 clusters missing, 20 participants missing (11%)
3. CCT: 0 clusters missing, 38 participants missing (8%)
4. No cash transfer (spill-over control group for CCT) (excluded): 0 clusters missing, 23 participants missing (9%)
5. No cash transfer (pure control group): 0 clusters missing, 143 participants missing (10%) |
Girls and young women out of school/dropouts at baseline:

1 intervention group and 1 control group

1. CCT: 0 clusters missing, 67 participants missing (15%)

2. No cash transfer (pure control group): 0 clusters missing, 78 participants missing (17%)

The number of missing participants per outcome was unclear.

### Selective reporting (reporting bias)
- Unclear risk
- We did not identify a study protocol

### Other bias
- Low risk
- None identified

---

**Methods**

Controlled before-and-after study, difference-in-differences methods, 24 months in 2005-2007

**Participants**

10,574 households at baseline (7016 households at follow-up 2) interviewed 3 times in the National Socioeconomic Survey (SUSENAS) (baseline: February 2005; follow-up 1: February 2006; follow-up 2: February 2007); 31 provinces, Indonesia

**Interventions**

1 exposure and 1 control group

1. Received UCT: 1715 households at follow-up 2 (included in this review as exposure group)

2. Did not receive a UCT: 5301 households at follow-up 2 (included in this review as control group)

Intervention duration: 12 months. Follow-up: 12 months into the intervention (alternative follow-up not reported in this review: 3-6 months into the intervention). Intervention design: aimed to prevent poor households from having to reduce expenditures on essential commodities, health, and education during strong national inflation; targeted to poor, disadvantaged households (but targeting was poorly implemented, with many non-poor households receiving the UCT and many poor households not receiving it); provided a total amount of USD 120 (USD 30 every 3 months for 12 months; approximately 2.0% of the annual GDP per capita); paid mainly in-hand at a post office; and fuzzy design: minor messaging that UCT receipt may be conditional on reported level of household welfare.

**Outcomes**

**Primary outcomes**

- Use of any health service (measure: number of outpatient visits)

**Secondary outcomes**

- School attendance (current)
- Current adult employment
- Healthcare expenditure (measure: log expenditure per capita)

**Alternative primary outcome measures not reported in this review**

- Use of any health service (measure: number of inpatient visits)

**Notes**

Intervention context: the Direct Cash Transfer Program, implemented by the Government of Indonesia predominantly through village officials; programme uptake 100%; population coverage over 19 million households; and programme costs between October 2005 and September 2006 approximately USD 380 million. Funder of the study: International Initiative for Impact Evaluation. Conflict of interest: none identified.

---

**Risk of bias**

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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### Bazzi 2012 (Continued)

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>High risk</td>
<td>It was unclear whether the sampling strategy was random. A nationally representative sample was achieved.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation concealment among participants and study personnel was unclear.</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Low risk</td>
<td>The study was not a cluster-RCT.</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Unclear risk</td>
<td>No overall P values, test statistics or SDs reported.</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>Unclear risk</td>
<td>No overall P values or test statistics reported, only SDs for each mean. However, there were no apparent statistically significant imbalances.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>The participants were not allocated to the intervention by the researchers. A secondary analysis was conducted of survey data collected for a different purpose than estimating the effect of the UCT on health services use and health outcomes. Therefore, blinding of participants was neither feasible nor necessary. It was unclear whether study personnel were blinded.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All subjectively measured outcomes</td>
<td>Low risk</td>
<td>The participants were not allocated to the intervention by the researchers. A secondary analysis was conducted of survey data collected for a different purpose than estimating the effect of the UCT on health services use and health outcomes. Therefore, blinding of outcome assessors was neither feasible nor necessary.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All objectively measured outcomes</td>
<td>Low risk</td>
<td>There were no objectively measured outcomes in this study.</td>
</tr>
<tr>
<td>Contamination</td>
<td>High risk</td>
<td>Allocation was by household, but additional income from UCTs provided to participants in the intervention group may have spilled over to participants in the control group (e.g. between family members). The risk of spill-over effects was not investigated (e.g. no spill-over control groups in this study).</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>High risk</td>
<td>Initial survey non-response rate was unclear. Attrition rate was very high (34%), and it was unclear whether the attrition rate was differential between the intervention group and the control group. The number of missing participants per outcome was also unclear. We considered the missing data to potentially have impacted effect estimates because a very large percentage of participants had missing data (34% or more).</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>We did not identify a study protocol.</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None identified.</td>
</tr>
</tbody>
</table>
### Methods
Cluster-randomised controlled trial (2 stages: first stage, 20 villages were randomly selected; second stage, 2034 households were randomly selected), regression analytic methods with propensity score matching and multiple imputation, 12 months in 2011-2012

### Participants
2034 household interviewed 3 times (baseline: before the start of intervention, date unclear; follow-up 1: approximately 8 months into the intervention, February 2012; follow-up 2: after the end of the intervention, date unclear), Madhya Pradesh, India

### Interventions
1 intervention group and 1 control group

1. UCT; specifically, a basic income transfer: 8 villages, 938 households (included in this review as intervention group)
2. No UCT (pure control group): 8 villages, 1096 households (included in this review as control group)

Intervention duration: 12 months. Follow-up: 8 months into the intervention (alternative follow-up not reported in this review: unclear number of months after 12 months of intervention). Intervention design: aimed to reduce poverty and increase social protection; non-targeted (i.e. a basic universal income intervention (Painter 2016)); provided a total amount of approximately USD 160 to adults (USD 13.20 per month for 12 months; approximately 3.2% of the annual GDP per capita) and USD 80 to each child aged 0-18 years (USD 6.60 per month for 12 months; approximately 1.6% of the annual GDP per capita); total amount equivalent to approximately 25% to 30% of an average poor family’s income; and paid in-hand in the first 3 months and into a bank account in the remaining months to the mother.

### Outcomes

**Primary outcomes**

- Fully vaccinated (dichotomous; measure: households with complete vaccination coverage (bacillus calmette guerin, diphtheria, pertussis, tetanus, polio, measles, mumps and rubella) of all children in the household between 6 months and 5 years)
- Has had any illness (measure: households with cases of illness or injury in household in last 3 months lasting more than 24 hours and needing treatment but not hospitalisation)

Alternative primary outcomes not reported in this review:

- Has had any illness (measure: households with cases of illness or injury in household in last 3 months requiring hospitalisation)

### Notes
Intervention context: pilot programme conducted by non-governmental organisation (Self Employed Women’s Association); implemented by non-governmental organisation (Self Employed Women’s Association); and unclear population coverage, intervention uptake and intervention costs. Funder of the study: United Nations Children’s Fund. Potential conflict of interest: none identified.

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Cluster-randomisation was undertaken at the level of the villages. Propensity score matching was undertaken of the cases.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation concealment was unclear for participants and study personnel</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Unclear risk</td>
<td>It was unclear whether participants were recruited after clusters had been allocated</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) reported between the intervention group and the control group in outcome measurements</td>
</tr>
</tbody>
</table>
### Baseline characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Risk</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Similar</td>
<td>High</td>
<td>Differences (P &lt; 0.05) reported between the intervention group and the control group for 7 of 23 demographic characteristics. Caste, religion, household monetary income, and income sharing differed at P &lt; 0.01, and income sufficient for food needs, income sufficient for food needs (baseline survey), and income sufficient for other needs differed at P &lt; 0.05. It is unclear if these were differences at baseline, except for the one variable that has been clearly labelled as being at the baseline survey.</td>
</tr>
</tbody>
</table>

### Blinding of participants and personnel (performance bias)

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Risk</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not possible</td>
<td>High</td>
<td>Blinding of participants was not possible, and blinding of personnel was unclear</td>
</tr>
</tbody>
</table>

### Blinding of outcome assessment (detection bias)

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Risk</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unclear</td>
<td>Unclear</td>
<td>Blinding of outcome assessors was unclear</td>
</tr>
</tbody>
</table>

### Contamination

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Risk</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unclear</td>
<td>Unclear</td>
<td>Allocation was conducted by village, but additional income from UCTs provided to participants in the intervention group may have spilled over to participants in the control group (e.g. between family members). Spill-over effects were not formally investigated (e.g. through a spill-over control group).</td>
</tr>
</tbody>
</table>

### Incomplete outcome data (attrition bias)

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Risk</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>Problems with baseline surveying due to a lack of formal training and supervision of the enumerators led to loss of 29% of all responses and an unclear number and percentage of missing values for some unspecified variables. The number of missing clusters and households and the number of missing values per outcome are unclear. We considered the missing data to potentially have impacted effect estimates because a very large percentage of participants missed data at baseline (25%).</td>
<td></td>
</tr>
</tbody>
</table>

### Selective reporting (reporting bias)

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Risk</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unclear</td>
<td>We did not identify a study protocol</td>
<td></td>
</tr>
</tbody>
</table>

### Other bias

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Risk</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>None identified</td>
<td></td>
</tr>
</tbody>
</table>

### Beck 2015 (Continued)

### Cunha 2014

#### Methods

Cluster-randomised controlled trial (2 stages: first stage, 208 rural communities or villages were randomly sampled amongst those considered eligible for the government’s Programa de Apoyo Alimentario UCT programme; second stage, approximately 33 households were randomly sampled from each community or village), difference-in-differences methods, 26 months in 2003-2005

#### Participants

5028 households at baseline (4923 households at follow-up) and 4550 children (0-6 years) at baseline (4129 children at follow-up), interviewed twice (baseline: October 2003-April 2004; follow-up: October 2005-December 2005), 208 villages that were small (< 2500 inhabitants), highly marginalised (as classified by the Census Bureau), non-welfare (not currently receiving the subsidised milk programme Liconsa or the conditional cash transfer Oportunidades), Mexico

#### Interventions

3 intervention groups and 1 control group
1. UCT plus educational classes: unclear number of clusters, unclear number of households, and unclear number of participants (included in this review as intervention group)
2. In-kind transfer plus educational classes: unclear number of clusters, unclear number of households, and unclear number of participants (excluded from this review)
3. In-kind transfer without education classes: unclear number of clusters, unclear number of households, and unclear number of participants (excluded from this review)
4. No UCT (pure control group): unclear number of clusters, unclear number of households, and unclear number of participants (included in this review as control group)

Intervention duration: 24 months. Follow-up: 24 months into the intervention. Intervention design: aimed to improve food security, nutritional intake and health; targeted to persons in poor households in rural, poor villages; provided a total amount of approximately USD 360 (approximately USD 15 per month for 24 months; approximately 2.8% of the annual GDP per capita); paid in-hand to women (if possible); minor co-intervention for UCT was education classes that provided information on nutrition, hygiene, and health (but implemented with participants in the intervention groups not attending and participants in the control group attending); the 2 in-kind transfer intervention groups were combined in analyses because education classes were offered in both of these intervention groups, making them indistinguishable.

Outcomes

Primary outcomes
- Stunting (measure: current height)
- Underweight (measure: current weight)
- Has had any illness

Secondary outcomes:
- Amount of money spent on health care

Alternative primary outcome measures not reported in this review
- Has had any illness
  * Number of days sick
  * Has anaemia

Notes

Intervention context: experiment by the Government of Mexico along the government programme called the Programa de Apoyo Alimentario; implemented by a public-private company (Diconsa) that maintains subsidised general stores in each of the included communities or villages; programme uptake was > 97%; unclear population coverage; and estimated total programme costs were approximately 102% of the total amount of cash transferred. Funder of study: Stanford University. Conflict of interest: none identified.

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Randomisation ensured</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation concealment among participants and study personnel unclear</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>High risk</td>
<td>Participants recruited after cluster allocation</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>No baseline differences reported for any of the included outcomes</td>
</tr>
</tbody>
</table>
### Baseline characteristics

<table>
<thead>
<tr>
<th>Similar</th>
<th>Low risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low risk</td>
<td>Baseline differences (P &lt; 0.05) reported between the unconditional cash transfer (UCT) intervention group and the control group in 2 characteristics. The years of education of household heads in the UCT group was lower than that in the control group (3.96 compared with 4.50, P &lt; 0.05). The proportion of households who raised animals or farmed in the UCT group was higher than in the control group (0.30 compared with 0.43, P &lt; 0.05). However, all baseline differences in characteristics comprehensively adjusted for using regression analytic methods.</td>
</tr>
</tbody>
</table>

### Blinding of participants and personnel (performance bias)

<table>
<thead>
<tr>
<th>High risk</th>
<th>Blinding of participants not possible, and blinding of personnel unclear</th>
</tr>
</thead>
</table>

### Blinding of outcome assessment (detection bias)

<table>
<thead>
<tr>
<th>High risk</th>
<th>Outcome assessors not blinded</th>
</tr>
</thead>
</table>

#### All outcomes

### Blinding of outcome assessment (detection bias)

<table>
<thead>
<tr>
<th>Low risk</th>
<th>Outcome assessors not blinded. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured.</th>
</tr>
</thead>
</table>

### Contamination

<table>
<thead>
<tr>
<th>High risk</th>
<th>Allocation was by community, but additional income from UCTs provided to participants in the intervention groups may have been transferred to participants in other intervention and/or control groups (e.g. between family members)</th>
</tr>
</thead>
</table>

### Incomplete outcome data (attrition bias)

<table>
<thead>
<tr>
<th>High risk</th>
<th>Survey non-response unclear. 8 clusters with 306 baseline households and 216 follow-up households were excluded from the study because they could not be re-surveyed due to concerns for enumerator safety (2 clusters), received the intervention prior to the baseline survey (2 clusters), were ineligible due to receiving Oportunidades (2 clusters), and were geographically contiguous (2 clusters). For the sample of children, a total of 189 baseline households and 78 follow-up households were excluded because they missed more than half of the outcomes measurements (35 baseline households and 78 follow-up households), reported no individual level information (11 baseline households), or reported non-normal food consumption pattern (143 baseline households). For the sample of children, a total of 200 children were excluded because they reported age inconsistently across survey waves. Attrition high (12% to 17%) and differential between UCT intervention group and control group (17% compared with 12%). The study reports no differences in outcome measurements and characteristics between participants lost to follow-up and those not lost to follow-up between the UCT intervention group and the control group. The numbers of missing households and participants per intervention and control group were unclear. For the sample of households, the number and percentage of households missing per outcome were unclear. For each outcome, the number of missing participants for the sample of households was unclear. For each outcome, the number of missing participants for the sample of children was:</th>
</tr>
</thead>
</table>

1. height: 10 participants missing (i.e. a very small percentage, which is unlikely to have affected this continuous outcome);
2. weight: 11 participants missing (i.e. a very small percentage, which is unlikely to have affected this continuous outcome); and
3. was sick: unclear number of participants missing. |
Cunha 2014 (Continued)

We considered the missing data to potentially have impacted effect estimates because a non-random sample of clusters was excluded from the study and a large percentage of participants were lost to follow-up (12%-17%).

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>No study protocol identified</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None identified</td>
</tr>
</tbody>
</table>

Fernald 2011

Methods
Clustered-randomised controlled trial (1 stage: 378 parishes randomly assigned to intervention group and control group), regression analytic methods, 27 months in 2003-2006

Participants
786 children (aged 12-35 months) at baseline (1285 children at follow-up) and 786 adults (the included children’s mothers) at baseline (1285 adults at follow-up) interviewed twice (baseline: October 2003-March 2004; follow-up: September 2005-January 2006), 6 provinces, Ecuador

Interventions
1 intervention group and 1 control group
1. UCT: 79 parishes, 530 participants (included in this review as intervention group)
2. No UCT (pure control group): 39 parishes, 256 participants (included in this review as intervention group)

Intervention duration: 18-27 months. Follow-up: 18-27 months into the intervention. Intervention design: aimed to reduce poverty and promote human capital investments among poor families through the provision of direct monetary transfers and incentives for households to invest in human capital (World Bank 2006); targeted to mothers who lived in poverty and had children aged 0-16 years; provided a total amount of USD 270-405 (USD 15 per month for 18-27 months; equivalent to 2.7% to 4.1% of the annual GDP per capita); paid in-hand to mothers; and fuzzy design: conditional on preventive healthcare checks and school attendance among children (but compliance not monitored).

Outcomes
Primary outcomes
- Has had a growth check, previous 6 months
- Has been given any parasite treatment, previous year
- Moderate stunting (assessed with: height for age)
- Level of dietary diversity in previous week (assessed with: Food Index)
- Depression (assessed with: Center for Epidemiologic Studies Depression Score)

Secondary outcomes
- Level of parenting quality (assessed with: Harsh Parenting Subscale of the Home Observation Measurement of the Environment Score)

Notes
Intervention context: Bono de Desarrollo Humano, implemented by the Government of Ecuador; programme uptake 73% in the intervention group and 3% in the control group; and unclear population coverage and total programme cost. Funder of study: none stated. Conflict of interest: none identified.

Risk of bias
<table>
<thead>
<tr>
<th></th>
<th>Risk Level</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment</td>
<td>Unclear</td>
<td>Allocation concealment among participants and study personnel unclear</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Participants recruited</td>
<td>High</td>
<td>Participants recruited after cluster allocation</td>
</tr>
<tr>
<td>after cluster allocation</td>
<td></td>
<td>(recruitment bias in cluster-RCTs)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline outcome</td>
<td>Low</td>
<td>No baseline differences (P &lt; 0.05) reported between the unconditional cash transfer (UCT)</td>
</tr>
<tr>
<td>measurements similar</td>
<td></td>
<td>intervention group and the control group in all 8 outcome measurements</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline characteristics</td>
<td>Low</td>
<td>No baseline differences (P &lt; 0.05) reported between the UCT intervention group and the control</td>
</tr>
<tr>
<td>similar</td>
<td></td>
<td>group in characteristics</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of participants</td>
<td>High</td>
<td>Blinding of participants not possible, and blinding of personnel unclear</td>
</tr>
<tr>
<td>and personnel (performance bias)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
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<tr>
<td>Blinding of outcome as-</td>
<td>Unclear</td>
<td>Unclear whether outcome assessors blinded</td>
</tr>
<tr>
<td>ssessment (detection bias)</td>
<td></td>
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<tr>
<td>All subjectively measured outcomes</td>
<td></td>
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<tr>
<td>Blinding of outcome as-</td>
<td>Low</td>
<td>Unclear whether outcome assessors blinded. Outcome is unlikely to be influenced by lack of</td>
</tr>
<tr>
<td>ssessment (detection bias)</td>
<td></td>
<td>blinding because it is objectively measured.</td>
</tr>
<tr>
<td>All objectively measured outcomes</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Contamination</td>
<td>Low</td>
<td>Allocation was by community and little contamination reported (73% of the intervention group</td>
</tr>
<tr>
<td></td>
<td></td>
<td>and 3% of the control group received the intervention)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data</td>
<td>High</td>
<td>The study extracted and analysed only a small subsample of the total study sample at baseline</td>
</tr>
<tr>
<td>(attrition bias)</td>
<td></td>
<td>(i.e. 786 of 5547 children or 14.2% of the total study sample), without an explanation for this</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td>selection. Initial survey non-response unclear. Overall attrition moderate (11%). Attrition</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non-differential between treatment and control groups by outcome measurements and characteristics.</td>
</tr>
<tr>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Number of missing clusters and number and percentage of missing participants:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. UCT: 0 parishes, 63 participants missing (12%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. No UCT (pure control group): 0 parishes, 26 participants missing (10%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Number of missing participants for the included outcomes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. Has had a growth check: 0 parishes missing (0%), 89 participants missing (11%) (considering</td>
</tr>
<tr>
<td></td>
<td></td>
<td>that baseline risk for this dichotomous outcome was 0.52, we considered the risk of bias for</td>
</tr>
<tr>
<td></td>
<td></td>
<td>this dichotomous outcome to be low)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Has been given any parasite treatment: 0 parishes missing (0%), 89 participants missing (11%)</td>
</tr>
</tbody>
</table>
Fernald 2011 (Continued)

4. Level of dietary diversity: 0 parishes missing (0%), 89 participants missing (11%) (we considered the risk of bias from this level of missing data for this continuous outcome to be low)

5. Depression: 0 parishes missing (0%), 89 participants missing (11%) (we considered the risk of bias from this level of missing data for this continuous outcome to be low)

We considered the missing data to be unlikely to have impacted effect estimates.

<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Unclear risk</th>
<th>No study protocol identified</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None identified</td>
</tr>
</tbody>
</table>

Galiani 2014

Methods

Controlled before-and-after study, difference-in-differences methods with individual and year fixed effects, 15 months in 2007-2008

Participants

3477 households and 3556 adults (aged > 65 years) interviewed twice (baseline: September 2007-November 2007; follow-up: November 2008-December 2008), 463 localities, 7 states (Guerrero, Querétaro, Michoacán, San Luis Potosí, Puebla, Veracruz and Hidalgo), Mexico

Interventions

1 exposure group and 3 control groups

1. UCT: 724 households, 1144 participants (≥ 70 years old and residing in a small (< 2500 residents), rural village) (included in this review as exposure group)
2. No UCT: 693 households, 806 participants (≥ 70 years old and not residing in a small, rural village) (included in this review as control group)
3. No UCT: 605 households, 954 participants (< 70 years old and residing in a small, rural village) (excluded from this review)
4. No UCT: 555 households, 652 participants (< 70 years old and not residing in a small, rural village) (excluded from this review)

Intervention duration: 12 months. Follow-up: 12 months into the intervention. Intervention design: aimed to ensure food security and improve the living conditions and quality of life of older people residing in rural areas (ILO 2013); targeted to older adults aged ≥ 60 years who resided in small, non-welfare, rural communities (ILO 2013); eligibility determined through applicants providing proof of age and residence; provided a total amount of USD 540 (USD 45 per month for 12 months; 3.5% of the annual GDP per capita); paid in-hand to pensioner every second month; and minor co-interventions: workshops and social development activities.

Outcomes

Primary outcomes

- Depression (assessed with: Geriatric Depression Scale)

Secondary outcomes

- Current adult employment
- Current poverty

Notes

Intervention context: Programa de Atención a Adultos Mayores en Zonas Rurales, implemented by the Government of Mexico; unclear programme uptake; population coverage was 2.1 million people; and the total programme cost of the UCT was USD 683 million (approximately 0.1% of Mexico’s GDP). Funder of study: none stated. Conflict of interest: none identified.
**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>High risk</td>
<td>Unclear whether sampling strategy was random. Unclear whether nationally representative sample achieved. No baseline differences between treatment and control groups in outcomes, but minor differences in population characteristics.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation concealment among participants and study personnel unclear</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Low risk</td>
<td>Not a cluster-RCT</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) reported between the unconditional cash transfer exposure group and the control group in the outcome measurements</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>High risk</td>
<td>Baseline differences (P &lt; 0.05) reported between the UCT exposure group and the control group in 3 characteristics at the household level. The proportion of households with male household heads was higher in the exposure group than in the control group (0.74 compared with 0.57, P &lt; 0.01). Consumption per adult equivalents (i.e. the sum of food and non-food expenditures plus the value of home-produced food) was higher than that in the control group (270.72 compared with 422.91, P &lt; 0.01). The average household was larger in the UCT exposure group than in the control group (5.60 compared with 4.02, P &lt; 0.01). Baseline differences (P &lt; 0.05) reported between the UCT exposure group and the control group in 4 characteristics at the individual level. The proportion of males was higher in the exposure group than in the control group (0.50 compared with 0.35, P &lt; 0.01). The number of years of school was higher than that in the control group (1.86 compared with 1.39, P = 0.01). The proportion of participants who were married was higher in the UCT exposure group than in the control group (0.66 compared with 0.46, P &lt; 0.01). However, all baseline differences in characteristics comprehensively adjusted for using regression analytic methods.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>Participants not allocated to the intervention by the researchers. Secondary analysis of survey data collected for a different purpose than estimating the effect of the unconditional cash transfer on use of health services and health outcomes. Therefore, blinding of participants and of personnel was neither feasible nor necessary.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>No subjectively measured outcome in this study</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>Participants not allocated to the intervention by the researchers. Secondary analysis of survey data collected for a different purpose than estimating the effect of the unconditional cash transfer on use of health services and health outcomes, therefore blinding of outcome assessors was neither feasible nor necessary. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured.</td>
</tr>
</tbody>
</table>
Contamination

High risk

Allocation was by individual, and additional income from the UCT provided to participants in the intervention groups may have been transferred to participants in the control group (e.g. between family members).

Incomplete outcome data (attrition bias)

Low risk

Initial survey non-response unclear. Attrition moderate (9%), but unclear whether differential between the intervention and control groups. The number and percentage of missing values per UCT intervention group and control group for the outcome is unclear. We considered it unlikely that the missing data impacted effect estimates.

Selective reporting (reporting bias)

Unclear risk

No study protocol identified.

Other bias

High risk

Misclassification bias of the exposure: used self-reported receipt of a UCT collected in surveys, but we considered the risk of this bias to be low. Confounding: this study did not adjust comprehensively for all potential confounders, and it used difference-in-differences methods, which carry a risk of bias, if the underlying time trends differ between the exposed group and the non-exposed group. We therefore judged the risk of confounding to be high. Reverse causation: the outcome may have impacted the exposure, but we considered the risk of this bias to be low.

Hausloffer 2013

Methods

Cluster-randomised controlled trial (3 stages: first stage, villages were randomly selected; second stage, eligible households were randomly assigned to intervention group or control group; third stage, either the female or male head of the assigned household was randomly assigned to intervention group or control group), regression analytic methods, 19 months in 2011-2012

Participants

1440 poor (i.e. without a thatch roof) households in rural areas, 2140 primary household members and 1203 children (aged < 5 years) interviewed twice (baseline: May 2011-November 2011; follow-up: September 2012-December 2012), 62 villages, Rarieda region, Kenya

Interventions

2 intervention groups and 2 control groups

1. UCT paid monthly to head of assigned household: 63 villages at cluster level, 258 households, unclear number of participants (included in this review as intervention group)

2. UCT paid as once-off lump sum to head of assigned household: 63 villages at cluster level, 245 households, unclear number of participants (excluded from this review because the UCT did not fit the inclusion criteria)

3. No UCT to households in the same village as UCT receivers (spill-over control group): 63 villages at cluster level, 505 households, unclear number of participants (included in this review as control group)

4. No UCT to households in different villages from UCT recipients (pure control group): unclear number of clusters, 432 households, unclear number of participants (excluded from this review because households were sampled retrospectively)

Intervention duration: 9 months (for the included UCT intervention). Follow-up: 7-9 months into the intervention and up to 10 months after 9 months of intervention. Intervention design: aimed to alleviate poverty among poor households; in both intervention groups, the UCT was stratified into random assignment of either a small or large cash amount; provided a total amount of USD 404 for the small UCT (approximately USD 4.89 per month for 9 months; or approximately 14.3% of the annualised PPP-adjusted, per-capita GDP) and USD 1516 for the large UCT (USD 168.44 per month for 9 months; approximately 53.8% of the annual GDP per capita); provided via mobile money service; and minor co-intervention: participants were provided with a SIM card for their cell phone.

Outcomes

Primary outcomes
Hausofer 2013 (Continued)

- Has had any illness, previous 2 weeks to last 3 months
- Food security (assessed with: Food Security Index)
- Level of dietary diversity (assessed with: number of times ate meat or fish in previous week)
- Level of depression (assessed with: Center of Epidemiology Depression Scale)

Secondary outcomes

- Amount of money spent on health care (assessed with: medical expenditure per episode)

Alternative primary outcomes not reported in this review

- Food security
  * Number of meals skipped
  * Went a whole day without food
  * Ate less preferred/cheaper foods
  * Relied on help from others for food
  * Purchased food on credit
  * Hunted, gathered wild food or harvested prematurely
  * Begged because did not have enough food in the house
  * Household members usually eat 2 meals
  * Household members usually eat until content
  * Enough food in the house for tomorrow
  * Slept hungry
- Level of dietary diversity (assessed with: consumed protein)
- Level of depression
  * Level of psychological well-being
  * Level of log cortisol
  * Worries
  * Level of happiness
  * Level of life satisfaction
  * Level of trust
  * Locus of control
  * Level of optimism
  * Level of self-esteem

Notes

Intervention context: experiment implemented by a nongovernmental organisation (GiveDirectly); unclear programme uptake, population coverage and total programme cost of the UCT. Funder of study: National Institute of Health and Cogito Foundation. Potential conflict of interest: one study author co-founded and formerly directed the organisation implementing the studied UCT.

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Randomisation ensured</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Allocation concealment among participants unclear and among study personnel ensured</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Low risk</td>
<td>Participants recruited after cluster allocation, but they were randomly assigned to intervention or control group</td>
</tr>
</tbody>
</table>
### Baseline outcome measurements similar

<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Risk Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Unclear risk</td>
<td>The study reports &quot;largely insignificant&quot; (p 14) differences between the intervention groups (combined) and the control groups (combined). However, we found a significant baseline difference across index variables when comparing male to female recipients of the UCTs (P = 0.02).</td>
</tr>
</tbody>
</table>

### Baseline characteristics similar

<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Risk Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline characteristcs similar</td>
<td>Unclear risk</td>
<td>No baseline differences (P &lt; 0.05) were observed comparing the baseline characteristics between the UCT intervention groups (combined) and the control group included in this review.</td>
</tr>
</tbody>
</table>

### Blinding of participants and personnel (performance bias)

<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Risk Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>High risk</td>
<td>Blinding of participants not possible, and blinding of personnel unclear</td>
</tr>
</tbody>
</table>

### Blinding of outcome assessment (detection bias)

<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Risk Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>High risk</td>
<td>Outcome assessors not blinded</td>
</tr>
</tbody>
</table>

### Contamination

<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Risk Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contamination</td>
<td>Low risk</td>
<td>This study tested for spill-over effects, and it did not find any evidence for contamination for the outcomes included in this review</td>
</tr>
</tbody>
</table>

### Incomplete outcome data (attrition bias)

<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Risk Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>Initial survey response unclear. Attrition low (7%) and non-differential between treatment and control groups and by outcomes. No significant (P &lt; 0.05) baseline differences observed in index variables between people lost to follow-up and those remaining in the study. The number and percentage of missing values per UCT intervention group and control group and for each of the outcomes is unclear.</td>
</tr>
</tbody>
</table>

### Selective reporting (reporting bias)

<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Risk Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td>All outcomes reported in the study protocol were also reported in the study record. However, the study protocol was only registered in the American Economic Association’s registry for randomised controlled trials on June 28, 2013 (RCT ID: AEARCTR-0000019), which is after data collection for the trial had occurred between May 1, 2011, and February 28, 2013.</td>
</tr>
</tbody>
</table>

### Other bias

<table>
<thead>
<tr>
<th>Outcome Measure</th>
<th>Risk Assessment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None identified</td>
</tr>
</tbody>
</table>

### Leroy 2010

#### Methods

Cluster-randomised controlled trial (3 stages: first stage, 208 rural communities were randomly sampled; second stage, 33 households were randomly sampled from each community; and third stage, the households were randomly assigned to 3 intervention groups and 1 control group), difference-in-differences methods, 26 months in 2003-2005.

#### Participants

2876 households and 1509 women (18-49 years) from the households, who were not pregnant or lactating and had no missing data at baseline, interviewed twice (baseline: October 2003-April 2004; follow-up: October 2005-December 2005); 8 states (Chiapas, Guerrero, Oaxaca, Quintana Roo, Tabasco, Campeche, Yucatan, and Veracruz), Mexico

#### Interventions

3 intervention groups and one control group
1. UCT plus educational classes: 53 communities, 1492 households, 958 participants (extracted from only 735 households) (included in this review as intervention group)
2. In-kind transfer: 52 communities, unclear number of households, unclear number of participants (excluded from this review)
3. In-kind transfer plus educational classes: 52 communities, unclear number of households, unclear number of participants (excluded from this review)
4. No UCT (pure control group): 51 communities, 1385 households, 946 participants extracted from only 668 households (included in this review as control group)

Intervention duration: unclear, but 14 months on average. Follow-up: unclear, but 23 months after the intervention had started. Intervention design: aimed to reduce short-term household vulnerability and to invest in long-term human capital accumulation through interventions in health, nutrition and education; targeted to communities that did not receive benefits from other federal food aid programmes, had < 2500 inhabitants, and had a high level of marginalisation (but 37% of the communities receiving the UCT had a medium rather than a high level of marginalisation, suggesting that the community-level targeting was not implemented successfully); provided a total amount of approximately USD 168 (approximately USD 14 per month for 14 months, 1.3% of the annual GDP per capita); paid in-hand every second month; minor co-intervention of UCT: education classes (which recipients did not commonly attend); and the 2 in-kind transfer groups were joined in the analysis.

**Outcomes**

**Primary outcomes**
- Underweight (dichotomous, assessed with: weight)
- Level of nutritional diversity (assessed with: consumed any protein)

Alternative primary outcome measures not included in the review
- Underweight (dichotomous, assessed with: body mass index)
- Level of nutritional diversity
  - Total energy consumed
  - Energy consumed in fruits and vegetables
  - Energy consumed in grains and legumes
  - Energy consumed in animal-source foods
  - Energy consumed in processed foods
  - Consumed any fat
  - Consumed any fibre

**Notes**

Intervention context: Programa de Apoyo Alimentario, implemented by the Government of Mexico through its Ministry of Social Development; unclear programme uptake, population coverage and the total programme costs of the UCT. Funder of study: none stated. Conflict of interest: none identified.

**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Randomisation ensured (although exact mechanism unclear). No differences between treatment and control groups in outcomes at baseline.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Allocation concealment among participants unclear and among study personnel partially ensured</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Low risk</td>
<td>Participants recruited after cluster allocation, but they were randomly sampled from all households in the cluster</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>Among the sample of households, significant baseline differences were reported for one outcome. The UCT intervention group had a larger level of vitamin C</td>
</tr>
</tbody>
</table>
Baseline characteristics similar | Low risk | No significant differences (P < 0.05) were observed between the UCT intervention group and the control group. However, not many variables were assessed to show balance in baseline characteristics.

Blinding of participants and personnel (performance bias) | High risk | Blinding of participants not possible, personnel blinded at least to the study objectives

Blinding of outcome assessment (detection bias) | Low risk | Outcome assessors were partially blinded to treatment allocation

Blinding of outcome assessment (detection bias) | Low risk | Outcome assessors were partially blinded to treatment allocation. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured.

Contamination | Unclear risk | No spill-over control group was included, so that the risk of contamination is unclear

Incomplete outcome data (attrition bias) | High risk | Initial survey non-response unclear. Among the sample of households, attrition was large (13%) and differential between the UCT intervention group and the control group (12% compared with 17%, no test provided). Among the sample of communities (extracted from a subsample of households), attrition was small (2% of participants), but differential by the UCT intervention group compared with the control group (0% compared with 3%).

Number of missing clusters and number and percentage of missing participants per group

1. UCT + educational classes: 0 communities (0%), 195 households (12%), 172 participants (18%)
2. In-kind transfer: 0 communities (0%), 130 households (8%), unclear number of participants
3. In-kind transfer plus educational classes: 1 community (2%), 210 households (13%), unclear number of participants
4. No UCT or in-kind transfer (pure control group): 1 community (2%), 279 households (17%), 225 participants (24%)

The number and percentage of participants with missing values per outcome was unclear. We considered the missing data to potentially have impacted effect estimates because a large percentage of participants were missing in the UCT intervention group (18%) and in the control group (24%).

Selective reporting (reporting bias) | Unclear risk | No study protocol was identified

Other bias | Low risk | None identified
### Methods
Cluster-randomised controlled trial (2 stages: first stage, 8 villages and 100 households per village were selected by a community committee; second stage, 4 villages each were randomly assigned to the UCT intervention group and to the control group), regression analytic methods, 12 months in 2007-2008

### Participants
1649 children (aged 6-17 years) interviewed twice (baseline: March 2007; follow-up: April 2008); Mchinji district, Malawi

### Interventions
1 intervention group and 1 control group

1. **UCT**: 4 villages, 979 participants (included in this review as intervention group)
2. **no UCT (pure control group)**: 4 villages, 670 participants (included in this review as the control group).

Intervention duration: 12 months. Follow-up: 12 months into the intervention. Intervention design: aimed to alleviate poverty, reduce hunger and malnutrition, and improve school enrolment within ultra-poor households; targeted to ultra-poor (poorest 10% of the population) and/or labour-constrained households with one or more adults; eligibility determined by volunteer village committees; and provided an average total amount of USD 124 (USD 12 per month for 12 months; 25.1% of the annual GDP per capita), but amount depended on household size and number of school-aged children:

- USD 48 for a one-member household (USD 4 per month for 12 months; 8.4% of the annual GDP per capita)
- USD 156 for a household with ≥ 4 members (USD 13 per month for 12 months; 27.2% of the annual GDP per capita)

Plus

- USD 12 per primary school-aged child (1 USD per month for 12 months; 2.1% of the annual GDP per capita)
- USD 36 per secondary school-aged child in the household (3 USD per month for 12 months; 6.3% of the annual GDP per capita).

### Outcomes

#### Primary outcomes
- Use of any health service (assessed with: used a health service for child’s worst illness, previous year)
- Has had any illness

#### Secondary outcomes
- School attendance

Alternative primary outcome measures not reported in this review
- Has had any illness (assessed with: has had any illness that stopped normal activities)

### Notes

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Randomisation ensured. However, the exact randomisation procedures are unclear.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation concealment among participants and among study personnel unclear</td>
</tr>
<tr>
<td><strong>Luseno 2012</strong> (Continued)</td>
<td></td>
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<tr>
<td>-----------------------------</td>
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</tr>
<tr>
<td><strong>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</strong></td>
<td>High risk</td>
<td>Participants recruited after cluster allocation</td>
</tr>
<tr>
<td><strong>Baseline outcome measurements similar</strong></td>
<td>High risk</td>
<td>Baseline differences (P &lt; 0.05) were reported between the UCT intervention group and the control group in one outcome measurement. Health care used for child’s worst illness in the past year was lower in the UCT intervention group than in the control group (80% compared with 90%, P &lt; 0.01).</td>
</tr>
<tr>
<td><strong>Baseline characteristics similar</strong></td>
<td>High risk</td>
<td>At the individual level, baseline differences were reported between the UCT intervention group and the control group in one characteristic. The distribution of orphans (maternal, paternal, double) differed (P = 0.02). At the household level, baseline differences between the UCT intervention group and the control group were reported in 6 characteristics. The UCT intervention group had higher educational status of the household head (P = 0.02), a smaller number of working-age adults in household (P = 0.01), a smaller number of children aged 6-9 years (P &lt; 0.01), smaller number of children aged 10-14 years (P &lt; 0.01), a smaller number of children aged 14-17 years (P = 0.01), and a smaller overall household size (P &lt; 0.01).</td>
</tr>
<tr>
<td><strong>Blinding of participants and personnel (performance bias)</strong></td>
<td>High risk</td>
<td>Blinding of participants not possible and blinding of personnel unclear</td>
</tr>
<tr>
<td><strong>Blinding of outcome assessment (detection bias)</strong></td>
<td>High risk</td>
<td>Outcome assessors not blinded</td>
</tr>
<tr>
<td><strong>Blinding of outcome assessment (detection bias)</strong></td>
<td>Low risk</td>
<td>No objectively measured outcome in this study</td>
</tr>
<tr>
<td><strong>Contamination</strong></td>
<td>Unclear risk</td>
<td>No spill-over control group was included, so that the risk of contamination is unclear</td>
</tr>
<tr>
<td><strong>Incomplete outcome data (attrition bias)</strong></td>
<td>High risk</td>
<td>Initial survey non-response unclear. A large proportion of participants was lost to follow-up (15%), and a larger proportion of participants in the UCT intervention group was lost to follow-up (18%), compared with that in the control group (12%). For each group, the number of clusters and number and percentage of participants (i.e. those who did not drop out of the study) with missing values were: 1. UCT: 0 villages, 84 participants (9%) 2. No UCT (pure control group): 0 villages, 82 participants (12%) The number and percentage of missing values per outcome is unclear. We considered the missing data to potentially have impacted effect estimates because of the large and differential loss to follow-up and the large percentage of participants with missing values.</td>
</tr>
<tr>
<td><strong>Selective reporting (reporting bias)</strong></td>
<td>Unclear risk</td>
<td>No study protocol identified</td>
</tr>
</tbody>
</table>
### Luseno 2012 (Continued)

<table>
<thead>
<tr>
<th>Other bias</th>
<th>Low risk</th>
<th>None identified</th>
</tr>
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</table>

### Miller 2008

#### Methods
Cluster-randomised controlled trial (2 stages: stage 1, 800 households in 8 village groups with a total of 23 villages were selected by community committees and enrolled in the UCT; stage 2, 4 village groups each were randomly assigned to the intervention group and to the control group); difference-in-differences methods; 13 months in 2007-2008

#### Participants
819 ultra-poor, labour-constrained households interviewed 3 times (baseline: March 2007; follow-up 1: September 2007; follow-up 2: April 2008); Mchinji District, Malawi

#### Interventions
1 intervention group and 1 control group
1. UCT: 4 village groups, 408 households (included in this review as intervention group)
2. No UCT: 4 village groups, 411 households (included in this review as control group)

Duration: 13 months. Follow-up: 13 months into the intervention. Intervention design: aimed to alleviate poverty, reduce hunger and malnutrition, and improve school enrolment within ultra-poor households; was targeted to ultra-poor (poorest 10% of the population) and/or labour-constrained households with one or more adults; eligibility determined by volunteer community social protection committees; and provided an average total amount of USD 124 (USD 12 per month for 12 months; 25.1% of the annual GDP per capita), but amount depended on household size and number of school-aged children.

- USD 55.90 for a one-member household (USD 4.30 per month for 13 months; 9.0% of the annual GDP per capita)
- USD 167.05 for a household with ≥ 4 members (USD 12.85 per month for 13 months; 26.9% of the annual GDP per capita)

Plus
- USD 13 per primary school-aged child (USD 1 per month for 13 months; 2.1% of the annual GDP per capita)
- USD 39 per secondary school-aged child (USD 13 per month for 13 months; 6.3% of the annual GDP per capita)

#### Outcomes
Primary outcomes
- Food security
- Dietary diversity (assessed with: Household Food Diversity Score)

#### Notes
Intervention context: programme of the Government of Malawi called the Social Cash Transfer Scheme; unclear programme uptake; population coverage of over 83,000 households in 2010; and unclear total programme costs of the UCT. Funder of study: United Nations Children’s Fund and United States Agency for International Development. Conflict of interest: none identified.

### Risk of bias

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<tr>
<td><strong>Miller 2008</strong> (Continued)</td>
<td><strong>Unclear risk</strong></td>
<td><strong>Unclear whether participants recruited after cluster allocation</strong></td>
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<td>------------------------------------------------------------------------------------------------</td>
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<td>---------------------------------------------------------------</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td><strong>Low risk</strong></td>
<td>No baseline differences (P &lt; 0.05) in outcome measurements reported between the intervention group and the control group. However, formal statistical test are only reported for level of dietary diversity.</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td><strong>High risk</strong></td>
<td>Baseline differences (P &lt; 0.05) reported between the UCT intervention group and the control group for 5 population characteristics. The proportion of households headed by person with no schooling was lower in the UCT intervention group (44% compared with 65%, P &lt; 0.01). The household size was larger in the UCT intervention group (4.7 compared with 3.5, P &lt; 0.01). The proportion of elderly-only households was lower in the UCT intervention group (12% compared with 22%, P &lt; 0.01). The proportion of households in which one adult provides for more than 3 dependents was larger in the UCT intervention group (23% compared with 16%, P &lt; 0.01). The proportion of households with no healthy adult aged 19–64 years was lower in the UCT intervention group (55% compared with 62%, P &lt; 0.05). The proportion of households with their house's outer walls made from grass was higher in the UCT intervention group (4% compared with 2%, P &lt; 0.01).</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td><strong>Unclear risk</strong></td>
<td>Blinding of participants not possible, and blinding of personnel unclear</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td><strong>Low risk</strong></td>
<td>Outcome assessors not blinded. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured.</td>
</tr>
<tr>
<td>Contamination</td>
<td><strong>Unclear risk</strong></td>
<td>No spill-over control group was included, so that the risk of contamination is unclear. Resources (food buckets) were only given to the control group, but anticipation bias could have impacted household spending for the control group (they were told in March 2007 that they would receive money transfers in April 2008 and could have borrowed against the future transfer). The intervention group may have overestimated their food expenditures to make sure they would continue to get the intervention.</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td><strong>Unclear risk</strong></td>
<td>The proportion of participants who were lost to follow-up was moderate (9%), but the proportion of participants lost to follow-up in the UCT intervention group was larger than that in the control group (10% compared with 6%). The number and percentage of missing values per UCT intervention group and control group and for each of the outcomes is unclear. We were not able to judge the risk of attrition bias from incomplete outcome data.</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td><strong>Unclear risk</strong></td>
<td>No study protocol identified</td>
</tr>
<tr>
<td>Other bias</td>
<td><strong>Low risk</strong></td>
<td>Minor co-intervention (food transfer) may have affected the treatment effect estimate, but we judged this risk to be low.</td>
</tr>
</tbody>
</table>
**Methods**

Cluster-randomised controlled trial (2 stages: stage 1, 48 geographic areas were randomly selected and matched into pairs; stage 2, within each matched pair one geographic area was randomly assigned to the intervention group and the other to the control group), difference-in-differences methods, 39 months in 2009-2012

**Participants**

6800 children (aged 0-17 years) and 2440 adults (aged 18–54 years) interviewed 3 times (baseline: August 2009–November 2010, follow-up 1: November 2010–November 2011; follow-up 2: February–November 2012), 4 counties (Mandera, Marsabit, Turkana and Wajir), Kenya

**Interventions**

1 intervention group and 1 control group

1. UCT: 24 areas, 1571 households, unclear number of participants (included in this review as intervention group)
2. No UCT (pure control group): 24 areas, 1536 households, unclear number of participants (included in this review as control group)

Duration: 24 months. Follow-up: 24 months into the intervention. Intervention design: aimed to reduce poverty, food insecurity and malnutrition and to promote asset retention and accumulation for beneficiary households; targeted to poor households; eligibility determined through geographic residency in areas with a large proportion of the population living in poverty; provided a total amount of USD 37.40 to USD 74.8 per household (USD 3.40 to USD 6.80 per transfer per capita for an average of 11 transfers in 24 months; 15.6% of the annual GDP per capita); paid in-hand every second month; and number of transfers received varied considerably across households.

**Outcomes**

**Primary outcomes**

- Use of any health service (assessed with: did not consult formal healthcare provider when was ill/injured (inverse coded))
- Has had any illness (assessed with: was ill/injured (excluding chronic illness))
- Stunting
- Underweight
- Food security (assessed with: went entire days without eating solid foods, during worst recent period of food shortage)
- Level of dietary diversity (assessed with: Household Dietary Diversity Index)

**Secondary outcomes**

- Livestock ownership
- School attendance
- Adult employment (assessed with: main activity is productive work)
- Extreme poverty (assessed with: is below the absolute poverty line)
- Amount of money spent on health care (assessed with: mean monthly per capita health expenditure per household)

**Alternative primary outcome measures not reported in this review**

- Is food secure
  * Borrowed food or rely on help from family or relatives
  * Sold any animals to buy food
  * Sold other assets (not animals)
  * Bought food on credit from a shop
  * Collected and ate wild foods and/or animal
  * Reduced number of meals
  * Has eaten smaller meals
Notes

Intervention context: pilot programme of the Government of Kenya called the Hunger Safety Net Programme; implemented by the Government of Kenya through its Ministry of State for the Development of Northern Kenya and Other Arid Lands and service providers contracted to the ministry; unclear programme uptake; population coverage of 300,000 beneficiaries in 60,000 households; and unclear total programme cost of the UCT. Funder of study: United Kingdom Department for International Development. Conflict of interest: none identified.

Risk of bias

<table>
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<tr>
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<td>High risk</td>
<td>Randomisation was ensured. Several baseline differences between the treatment and control groups in outcomes and population characteristics.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Allocation determined by public lottery and therefore concealment protected</td>
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</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) between UCT intervention group and control group in outcome measurements</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) between UCT intervention group and control group in characteristics. However, the loss of 8 clusters at follow-up 2 reduced the balance in an unclear number of characteristics and seasonality of the UCT intervention group and the control group, compared with the balance at baseline of the original sample structure.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>Blinding of participants not possible, and blinding of personnel unclear</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All subjectively measured outcomes</td>
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<td>Outcome assessors not blinded</td>
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<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>High risk</td>
<td>Initial survey non-response unclear. Loss to follow-up was large, and the proportion of households lost to follow-up in the UCT intervention group was larger than that in the control group (18% compared with 13%, no test of statistical significance provided). Loss to follow-up was also differential by one population characteristic, i.e. by district.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Number of clusters and number and percentage of participants with missing values by group:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. UCT: 4 areas (17%), unclear number of participants</td>
</tr>
</tbody>
</table>
**Oxford Policy Management 2012 (Continued)**

2. No UCT (pure control group): 4 areas (17%), unclear number of participants

The number and percentage of missing values per outcome were unclear. Considering the large and likely differential loss to follow-up, we judged the risk of bias from attrition to be high.

<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
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<tbody>
<tr>
<td>Other bias</td>
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</tr>
</tbody>
</table>

**Paxson 2007**

**Methods**

Cluster-randomised controlled trial (3 stages: stage 1, 118 parishes were randomly selected from a total of 378 parishes; stage 2, the 118 parishes were randomly assigned to the intervention group and the control group; stage 3, 50 families per parish were selected into the study), difference-in-differences methods, 27 months in 2003-2006

**Participants**

2069 children (aged 36-84 months at follow-up) and 2069 mothers (aged 24 years on average) in poor, non-welfare (Bono Solidario) families with one or more children aged 0-72 months at baseline and no children aged > 72 months interviewed twice (baseline: October 2003-March 2004; follow-up: September 2005-January 2006), 6 rural provinces, Ecuador

**Interventions**

1 intervention group and 1 control group

1. UCT: 79 parishes, unclear number of participants (included in this review as intervention group); and
2. No UCT (pure control group): 39 parishes, unclear number of participants (included in this review as control group).

Duration: 15-19 months. Follow-up: 15-19 months. Intervention design: aimed to reduce poverty and promote human capital investments among poor families through the provision of direct monetary transfers and incentives for households to invest in human capital (World Bank 2006); targeted to low-income mothers of children aged 0-16 years; eligibility determined by a programme-specific poverty threshold; provided a amount of USD 225-285 in 2006 (USD 15 per month for 15-19 months; 2.7% of the annual GDP per capita); paid in-hand to mothers every month; and fuzzy design: compliance with conditions for preventive healthcare checks and school attendance for children existed but were not monitored.

**Outcomes**

Primary outcomes

- Has had a growth check, last 6 months
- Has been given any parasite treatment, last year
- Stunting (assessed with: height for age)
- Level of depression (assessed with: Center of Epidemiology Depression Scale)

Secondary outcomes

- Attends school (assessed with: is in a preschool or grade school)
- Level of parenting quality (assessed with: Home Observation Measurement of the Environment Score)

**Notes**

Intervention context: programme of the Government of Ecuador called the Bono de Desarrollo Humano; uptake was 73% in the intervention group and 3% in the control group; population coverage was 40%; and unclear total programme costs of the UCT. Funder of study: the World Bank, Government of Ecuador and Princeton University. Conflict of interest: none identified.

**Risk of bias**

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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### Paxson 2007 (Continued)

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</tr>
<tr>
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</tr>
<tr>
<td>Contamination</td>
<td>Low risk</td>
<td>3.7% of families in the control group received the UCT</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Initial survey non-response rate low (6%). Attrition low (6%) and reported as being non-differential by treatment versus control group. The number and percentage of missing values per UCT intervention group and control group is unclear. Approximately 33% of children have a missing value on one or more outcomes. Considering the large percentage of children with missing values, we judged the risk of attrition bias to be high.</td>
</tr>
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<tr>
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</tr>
</tbody>
</table>

### Pellerano 2014

Methods  
Cluster-randomised controlled trial (4 stages: stage 1, 96 electoral divisions were matched into 48 pairs; stage 2, 40 pairs were randomly selected from the 48 pairs; stage 3, secondary sampling units of clusters of villages in the pairs were constructed; stage 4, households were randomly selected from the secondary sampling units and randomly assigned to the intervention group and the control group), difference-in-differences methods, 26 months in 2011-2013.
**Participants**

3102 households interviewed twice (baseline: June-August 2011; follow-up: June-August 2013), 5 districts (Qacha’s Nek, Maseru, Leribe, Berea and Mafeteng), Lesotho

**Interventions**

1 intervention group and 1 control group

1. UCT: unclear number of electoral divisions, 1540 households (included in this review as intervention group)
2. No UCT (pure control group): unclear number of electoral divisions, 1562 participants (included in this review as control group)

Duration: 24 months. Follow-up: 24 months into the intervention. Intervention design: aimed to improve the living standards of orphans and other vulnerable children in order to reduce malnutrition, improve health status and increase school enrolment among these children; targeted to poor and vulnerable households with one or more children; eligibility determined through a combination of means-testing based on poverty, community validation and registration in the National Information System for Social Assistance; provided a total amount of USD 98 per household before April 2013 (approximately USD 4 per month for 24 months; 1.5% of the annual GDP per capita) and between USD 216 to households with ≤ 2 children (USD 9 per month for 24 months; 3.9% of the annual GDP per capita) and USD 450 to households with ≥ 5 children (USD 18.75 per month for 24 months; 8.1% of the annual GDP per capita) after April 2013; paid in-hand every 4 months; minor co-intervention: the Food Emergency Grant, a UCT for assistance in a humanitarian disaster (i.e. food insecurity from poor harvest) of USD 40 per month (17.4% of the annual GDP per capita), was provided alongside the UCT to recipients over 2012-2013; and fuzzy design: UCT was accompanied by instructions from social development officers at the pay point to spend the money on children.

**Outcomes**

**Primary outcomes**

- Registered birth (assessed with: proportion of children (aged 0-6 years) with a birth certificate)
- Has had a growth check, last 6 months (assessed with: child had any growth checks recorded in their Bukana health card)
- Up-to-date on vaccination calendar
- Has used any health services in previous 1 to 12 months (assessed with: consulted a healthcare provider)
- Is moderately underweight
- Has had any illness in previous 2 weeks to 3 months
- Has been food secure over previous month month (assessed with: had a food security index of ≥ 2)
- Level of dietary diversity, currently to last week (assessed with: household Dietary Diversity Score)

**Secondary outcomes**

- Livestock ownership in previous year
- School attendance (current)
- Current engagement in child labour
- Current adult employment
- Is extremely poor, currently
- Amount of money spent on health care (assessed with: average amount spent per child on healthcare in the last 3 months (across all children 0-5 years))

**Alternative primary outcome measures not reported in this review**

- Registered birth (assessed with: proportion of children (aged 0-6 years) in the process of getting a birth certificate)
- Has had a growth check, previous 6 months (assessed with: average number of growth monitoring checks recorded in Bukana health cards for children aged 0-36 months)
- Use of any health services, previous 1 to 12 months (assessed with: proportion of children living in household without enough money to spend on child (if needed) or child was not taken to consult a doctor if ill)
Pellerano 2014 (Continued)

- Has had any illness, last 2 weeks to last 3 months
  - Child aged 0-5 years had diarrhoea
  - Child aged 0-5 years had fever
  - Average number of days ill
  - Average number of days ill with flu or cold
  - Average number of days ill with fever
  - Average number of days ill with diarrhoea
  - Average number of days ill with persistent cough
  - Average number of days ill with stomach ache/vomit
  - Average number of days ill with skin rash
  - Average number of days ill with prolonged fever
  - Average number of days ill with unhealed sores
  - Average number of days ill with other disease
- Level of dietary diversity
  - Ate main staples
  - Ate pulses
  - Ate vegetables
  - Ate fruit
  - Ate meat, fish and egg
  - Ate dairy products

Notes

Intervention context: programme of the Government of Lesotho called the Lesotho Child Grants Programme, implemented through Ministry of Social Development; unclear programme uptake; population coverage was 20,000 households with 50,000 children by the end of 2013; and unclear total programme cost of the UCT. Funder of study: European Union. Conflict of interest: none identified.

Risk of bias

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<td>No baseline differences (P &lt; 0.05) between the UCT intervention group and the control group in outcome measurements</td>
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<tr>
<td>Baseline characteristics similar</td>
<td>High risk</td>
<td>Baseline differences (P &lt; 0.05) between the UCT intervention group and the control group in 5 characteristics, i.e. number of children aged 0-5 years (P &lt; 0.01), females aged 18-59 years (P &lt; 0.05), price of rubber boots (P &lt; 0.05), average daily wage for females (P &lt; 0.05) and proportion of households that borrowed or received support from other family members, friends or neighbours in (P &lt; 0.05).</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
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### Pellerano 2014 (Continued)

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<tr>
<th>Bias</th>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blinding of outcome assessment (detection bias)</strong></td>
<td><strong>Unclear risk</strong></td>
<td>Blinding of outcome assessors unclear.</td>
</tr>
<tr>
<td>All subjectively measured outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Blinding of outcome assessment (detection bias)</strong></td>
<td><strong>Low risk</strong></td>
<td>Blinding of outcome assessors unclear. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured.</td>
</tr>
<tr>
<td>All objectively measured outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Contamination</strong></td>
<td><strong>Unclear risk</strong></td>
<td>No spill-over control group was included, so that the risk of contamination is unclear. An unclear number of households included in the follow-up in the UCT intervention group might not have received the UCT.</td>
</tr>
<tr>
<td><strong>Incomplete outcome data (attrition bias)</strong></td>
<td><strong>High risk</strong></td>
<td>Initial survey non-response unclear. Attrition low (6%), but differential between the group eligible for the intervention and the group ineligible for it (9% versus 1%), as well as between the treatment and control groups among eligible participants (8% versus 12%).</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Selective reporting (reporting bias)</strong></td>
<td><strong>Unclear risk</strong></td>
<td>No study protocol identified</td>
</tr>
<tr>
<td><strong>Other bias</strong></td>
<td><strong>Low risk</strong></td>
<td>Minor co-intervention (UCT for assistance in humanitarian disaster) may have affected the treatment effect estimates, but we judged this risk to be low.</td>
</tr>
</tbody>
</table>

### Robertson 2012

#### Methods

Matched cluster-randomised controlled trial (3 stages: stage 1, each of 10 sites were divided into 3 clusters; stage 2, each cluster within each site was randomly assigned to a UCT group, CCT group and control group; stage 3, all eligible households in each cluster were included), regression analytic methods, 22 months in 2009-2011

#### Participants

2008 children (aged 0-5 years) in poor households that included one or more non-welfare (other cash transfers for orphans or children) children (aged 0-17 years at baseline) and were headed by a child (aged 0-17 years) or cared for one or more orphan children (aged 0-17 years), disabled persons or chronically ill persons interviewed twice (baseline: July 2009-September 2009; follow-up: March 2011-May 2011), 10 sites, Manicaland, Zimbabwe

#### Interventions

2 intervention groups and 1 control group

1. UCT: 10 sites, 763 children (included in this review as the intervention group)
2. CCT: 10 sites, 637 children (included in this review as CCT comparison group)
3. No cash transfer (pure control group): 10 sites, 608 children (included in this review as control group)

Duration: 12 months. Follow-up: 2-4 months after 12 months of the intervention. Intervention design: aimed to reduce poverty; targeted to poor households with one or more non-welfare (other cash transfers for orphans or children) children (aged 0-17 years at baseline) that were headed by a child (aged 0-17 years) or cared for one or more orphan children (aged 0-17 years), disabled persons or chronically ill persons; eligibility determined through population survey and community committees made up of a nongovernmental organisation (Diocese of Mutare Community Care Programme) and other local stakeholders (e.g. community health workers); UCT and CCT provided a total amount of USD 108 (USD 9 per month for 12 months; 7.8% of the annual GDP per capita) plus USD 24 per child (up to a maximum of 3 children) (USD 2 per month for 12 months; 1.7% of the annual GDP per capita); paid in-hand every 2 months; co-interventions for UCT and CCT: in-kind transfers of maize seeds and fertiliser were provided alongside the UCT twice (December 2009 and August 2010) and parenting skill training was provided from September 2010; and CCT was conditional on applying for a birth certificate within 3 months for all children younger than 18 years (including newborn babies) whose births had not been registered;
children younger than 5 years being fully vaccinated and attending growth monitoring clinics twice a year; children aged 6-17 years attending school at least 90% of the time each month; and a representative from every household attending two-thirds of local parenting skills classes.

### Outcomes

**Primary outcomes**
- Registered birth
- Fully vaccinated

**Secondary outcomes**
- School attendance (current)

### Notes

Intervention context: experiment conducted by research organisations; implemented by a nongovernmental organisation (Diocese of Mutare Community Care Programme); programme uptake was large, with 90% of eligible households reporting receiving the UCT; population coverage was 18% of the population (in the study sites); and unclear total programme cost of the UCT. Funder of study: World Bank, Programme of Support for the Zimbabwe National Action Plan for Orphans and Vulnerable Children and Wellcome Trust. Conflict of interest: none identified.

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Random sequence generation ensured. 2 control villages were accidentally enrolled into the UCT arm.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation determined by public lottery and therefore concealment protected</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Low risk</td>
<td>Participants recruited before cluster allocation</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>High risk</td>
<td>Baseline differences (P &lt; 0.05) between the UCT intervention group, CCT intervention group and the control group in one outcome measurement. The proportion of children who were fully vaccinated in the UCT intervention group was 65%, in the CCT comparison group was 66% and in the control group was 66% (k = 0.03).</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>Unclear risk</td>
<td>No baseline differences (P &lt; 0.05) between the UCT intervention group, CCT intervention group and the control group in population characteristics</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>Blinding of participants not possible, and blinding of personnel partially ensured (i.e. among data analysts)</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All subjectively measured outcomes</td>
<td>Low risk</td>
<td>Outcome assessors were blinded</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All objectively measured outcomes</td>
<td>Low risk</td>
<td>No objectively measured outcome in this study</td>
</tr>
</tbody>
</table>
### Robertson 2012 (Continued)

<table>
<thead>
<tr>
<th>Contamination</th>
<th>Unclear risk</th>
<th>No spill-over control group was included, so that the risk of contamination is unclear</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Initial survey response rate unclear. 2 of the original 12 eligible sites were not included in the study. The proportion of children lost to follow-up was very high (53%), and the proportion of participants lost to follow-up in the UCT intervention group (50%) differed from that in the CCT comparison group (56%) and the control group (55%). Number of clusters and number and percentage of participants with missing values per group: 1. UCT: 0 sites missing, at least 384 children missing (50% of original sample); 2. CCT: 0 sites missing, at least 354 children missing (56% of original sample); and 3. no cash transfer (pure control group): 0 sites missing, at least 361 children (55% of original sample). Number of missing participants for primary outcomes: 1. Registered birth: 0 sites missing (0%), 3 children missing (1%) (considering that the baseline risk for this dichotomous outcome was 0.43, we considered the risk of bias for this dichotomous outcome to be low) 2. Fully vaccinated: 0 sites missing (0%), 9 children missing (2%) (considering that the baseline risk for this dichotomous outcome was 0.63, we considered the risk of bias from this dichotomous outcome to be low). Due to the very high loss to follow-up (53%), we judged the risk of attrition bias to be high.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Unclear risk</th>
<th>No study protocol identified</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>The economic crisis may have affected the results, but we judged this risk to be low</td>
</tr>
</tbody>
</table>

### Salinas-Rodríguez 2014

<table>
<thead>
<tr>
<th>Methods</th>
<th>Controlled before-and-after study, difference-in-difference linear probability model with individual fixed effect, unclear number of months in 2007-2009</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>5465 older adults (≥ 70 years) residing in locations with ≤ 2500 residents at baseline (5270 older adults at follow-up) interviewed twice (baseline: October 2007-December 2007; follow-up: November 2008-December 2008), 516 rural locations, 7 districts, Mexico</td>
</tr>
<tr>
<td>Interventions</td>
<td>1 exposure group and 3 control groups 1. Received UCT: 1353 participants aged 70–74 years and residing in villages with ≤ 2500 inhabitants (included in this review as the intervention group) 2. Did not receive UCT (pure control group): 888 participants aged 70–74 years and residing in villages with 2501–2700 inhabitants (included in this review as the control group) 3. Did not receive UCT (pure control group): 1375 participants aged 65–69 years and residing in villages with ≤ 2500 inhabitants (excluded from this review) 4. Did not receive UCT (pure control group): 882 participants aged 65-69 years and residing in villages with 2501–2700 inhabitants (excluded from this review)</td>
</tr>
</tbody>
</table>
Intervention duration: unclear. Follow-up: up to 24 months into the intervention. Intervention design: aimed to improve the living conditions among adults aged ≥ 70 years by boosting their social protection through policy mechanisms; targeted to all older adults aged ≥ 70 years (i.e. universal UCT); and provided a total amount of up to USD 960 (USD 40 per month for 24 months).

**Outcomes**

**Primary outcome**

- Depression (assessed with: Geriatric Depression Scale)

**Notes**

Intervention context: programme of the Government of Mexico called the Programa 70 y más; population coverage was 1 million in 2007, 1.8 million in 2009 and 3.9 million in 2014; unclear intervention uptake; and total programme costs of the UCT were approximately USD 595 million in 2007, USD 1.4 billion in 2009 and USD 3.5 billion in 2014. Funders of the study: Mexican Ministry of Social Development and International Initiative for Impact Evaluation. Potential conflict of interest: none identified.

**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>High risk</td>
<td>Randomisation not ensured</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation concealment not feasible among participants in the intervention group, unclear for control groups</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Low risk</td>
<td>Not a cluster-RCT</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>Baseline health outcomes were not statistically significant different from each other (P = 0.05)</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>High risk</td>
<td>Groups were not well balanced at baseline, no overall test reported</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>Blinding of participants not possible, and blinding of personnel unclear</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All subjectively measured outcomes</td>
<td>Low risk</td>
<td>No subjectively measured outcome in this study</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All objectively measured outcomes</td>
<td>Unclear risk</td>
<td>Blinding of outcome assessors unclear. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured.</td>
</tr>
<tr>
<td>Contamination</td>
<td>Unclear risk</td>
<td>Allocation was by locality, but risk of contamination is unclear</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>Unclear risk</td>
<td>Response rate of 91% (5465 out of 6000 participants), follow-up of 96% (5270 out of 5465 participants) and complete response of 4468 participants. Non-response, attrition, and incomplete response not reported by group, but 2 control groups have substantially lower complete responses than the intervention group and the third control group (59.2% and 58.8% vs 89.7% and 90.2%, respectively).</td>
</tr>
</tbody>
</table>
Salinas-Rodríguez 2014 (Continued)

Selective reporting (reporting bias)
- Unclear risk
- No study protocol found

Other bias
- High risk
  - Misclassification bias of the exposure: used self-reported receipt of a UCT collected in surveys, but we considered the risk of this bias to be low. Confounding: this study did not adjust comprehensively for all potential confounders, and it used difference-in-differences methods, which carry a risk of bias, if the underlying time trends differ between the exposed group and the non-exposed group. We therefore judged the risk of confounding to be high. Reverse causation: the outcome may have impacted the exposure, but we considered the risk of this bias to be low.

Schady 2012

Methods
- Cluster-randomised controlled trial (3 stages: stage 1, the 6 provinces in which the UCT had not yet been implemented were non-randomly selected; stage 2, all 77 rural parishes in the 6 provinces were randomly assigned to an intervention group and a control group; stage 3, all eligible families within the intervention were enrolled for the intervention), regression analytic methods, 57 months in 2003-2008.

Participants
- 1702 adults (aged 24 years on average) in poor households interviewed 3 times (baseline: October 2003-March 2004; follow-up 1: September 2005-January 2006; follow-up 2: May 2008-July 2008), all women, 77 rural parishes, 6 provinces, Ecuador

Interventions
- 1 intervention group and 1 control group
  1. UCT: 51 parishes, unclear number of participants (included in this review as intervention group)
  2. No UCT (pure control group): 26 parishes, unclear number of participants (included in this review as control group)

  Intervention duration: 50-57 months. Follow-up: 50-57 months into the intervention (alternative follow-up not reported in this review: 18/27 months into the intervention). Intervention design: aimed to reduce poverty and promote human capital investments among poor families through the provision of direct monetary transfers and incentives for households to invest in human capital (World Bank 2006); targeted to low-income mothers of children aged 0-16 years; eligibility determined by a programme-specific poverty threshold; provided a total amount of USD 750-855 in 2006 (USD 15 per month for 50/57 months; 2.7% of the annual GDP per capita); paid in-hand every month to mothers; and fuzzy design: compliance with conditions for preventive healthcare checks and school attendance among children existed, but they were not monitored.

Outcomes
- Primary outcome
  - Has had any illness, previous 1 to 12 months (assessed with: had anaemia)

Notes
- Intervention context: programme of the Government of Ecuador called the Bono de Desarrollo Humano; programme uptake was 84% of clusters in the intervention group and < 4% of clusters in the control group at follow-up 1, and 85% of clusters in the intervention group and 48% of clusters in the control group at follow-up 2; population coverage was approximately 40%; and unclear total programme cost of the UCT. Funder of study: none stated. Conflict of interest: none identified.

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Randomisation ensured. No baseline differences between treatment and control group in population characteristics and outcomes.</td>
</tr>
</tbody>
</table>
### Schady 2012 (Continued)

<table>
<thead>
<tr>
<th>Bias</th>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Allocation concealment among participants unclear and among study personnel ensured (at least among enumerators)</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>High risk</td>
<td>Participants recruited after cluster allocation</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) between the UCT intervention group and the control group in the outcome measurement</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) between the UCT intervention group and the control group in population characteristics</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>Blinding of participants not possible, and blinding of study personnel unclear</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All subjectively measured outcomes</td>
<td>Low risk</td>
<td>No subjectively measured outcome in this study</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All objectively measured outcomes</td>
<td>Low risk</td>
<td>Blinding of outcome assessors unclear. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured.</td>
</tr>
<tr>
<td>Contamination</td>
<td>Unclear risk</td>
<td>No spill-over control group was included, so that the risk of contamination is unclear.</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>High risk</td>
<td>Initial survey non-response unclear. The proportion of participants who were lost to follow-up was large (17.7%), and the proportion of participants who were lost to follow-up in the intervention group was similar to that in the control group. The number and percentage of clusters and participants missing in the intervention group and in the control group was unclear. The number and percentage of missing values per outcome was unclear. Considering the large proportion of participants who were lost to follow-up, we judged the risk of attrition bias to be high.</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>No study protocol identified</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None identified</td>
</tr>
</tbody>
</table>

### Seidenfeld 2013

<table>
<thead>
<tr>
<th>Methods</th>
<th>Cluster-randomised controlled trial (2 stage: stage 1, 90 communities were randomly assigned to the intervention group and the control group; stage 2, all eligible households in the intervention group were immediately enrolled into the UCT after having a newborn baby), difference-in-differences methods, 33 months during the years 2010–2013</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>2515 households with one or more children (aged &lt; 3 years) interviewed 3 times (baseline: October 2010–November 2010; follow-up 1: October 2012–November 2012; follow-up 2: June 2013–July 2013), 90 communities, 3 districts (Kalabo, Shangombo and Kaputa), Zambia</td>
</tr>
</tbody>
</table>
Interventions

1 intervention group and 1 control group

1. UCT: unclear number of communities, unclear number of households (included in this review as intervention group)
2. No UCT (pure control group): unclear number of communities, unclear number of households (included in this review as control group)

Intervention duration: 30 months. Follow-up: 24 months (for all primary outcomes and most secondary outcomes) or 30 months (for some secondary outcomes). Intervention design: aimed to reduce extreme poverty and the intergenerational transfer of poverty by increasing food security, young child nutrition and health and education for school-age children, as well as by strengthening livelihoods; provided a total amount of approximately USD 360 (approximately USD 12 per month for 30 months; 4.3% of the annual GDP per capita); and paid in-hand every second month.

Outcomes

Primary outcomes

- Registered birth
- Use of any health services, previous 1 to 12 months
- Moderate stunting (assessed with: current height for age)
- Moderate underweight (assessed with: current weight for age)
- Has had any illness, previous 2 weeks to 3 months (assessed with: had acute respiratory disease)
- Food security (assessed with: Nutrition Technical Assistance Project Food Security Score)
- Dietary diversity (assessed with: Household Dietary Diversity Score)

Secondary outcomes

- Livestock ownership, over previous year
- School attendance, currently
- Extremely poor, currently
- Amount of money spent on health care

Alternative primary outcome measures not reported in this review:

- Fully vaccinated, currently
  * Received tetanus vaccination during pregnancy
  * Received malaria preventative medication during pregnancy
  * Received VCT during pregnancy
- Use of any health service, previous 1 to 12 months
  * Received any antenatal care
  * Received antenatal care from doctor or nurse
  * Received antenatal care within first trimester
  * Received at least 4 antenatal care visits
  * Sought preventive care
  * Received care for diarrhoea
  * Received care for fever
  * Received care for acute respiratory disease
- Has had any illness, last 2 weeks to last 3 months
  * Had diarrhoea
  * Had fever
- Has been food secure, currently to last month
  * Is not severely food insecure
  * Ate more than one meal a day
- Level of dietary diversity
  * Ate meat or fish 5 or more times
  * Ate vegetables 5 or more times
**Notes**


**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Randomisation ensured (coin flip). No baseline differences between treatment and control group in outcomes and population characteristics.</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Allocation concealment among participants and study personnel unclear</td>
</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>Unclear risk</td>
<td>Unclear whether participants recruited after cluster allocation</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) between the UCT intervention group and the control group in the outcome measurements</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) between the UCT intervention group and the control group in population characteristics</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>High risk</td>
<td>Blinding of participants not possible, and blinding of personnel unclear</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>High risk</td>
<td>Outcome assessors not blinded</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
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<td>Outcome assessors not blinded. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured</td>
</tr>
<tr>
<td>Contamination</td>
<td>Unclear risk</td>
<td>No spill-over control group was included, so that the risk of contamination is unclear.</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>Survey nonresponse rates were unclear. The proportion of participants lost to follow-up was low (9%), and the proportion of participants lost to follow-up among the UCT intervention group were similar to those among the control group. The number and percentage of missing communities and households and the percentage of missing values per outcome were unclear. We are unable to judge the risk of attrition bias.</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>No study protocol identified</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>None identified</td>
</tr>
</tbody>
</table>
Methods
Cluster-randomised controlled trial (3 stages: stage 1, 4 locations in each of 7 districts were selected; stage 2, 2 locations each were randomly assigned to the intervention group and the control group; stage 3, households receiving the UCT were randomly selected from a list supplied by the UCT programme and other households were randomly selected from a household listing undertaken in a random sample of census enumeration areas), difference-in-differences methods, 28 months in 2007-2009

Participants
9231 children (0-17 years) in households with orphans or vulnerable (i.e. chronically ill or with a chronically ill caregiver) children interviewed twice (baseline: March 2007-August 2007; follow-up: March 2009-July 2009), 28 locations, 7 districts, Kenya

Interventions
1 intervention group and 3 control groups
1. UCT: 14 locations, 1540 participants (included in this review as intervention group)
2. No UCT (spill-over for intervention group): 14 locations (same locations as intervention group), 238 participants (excluded from this review)
3. No UCT (pure control group): 14 locations, 754 participants (included in this review as control group)
4. No UCT (spill-over for control group): 14 locations (same locations as control group), 227 participants (excluded from this review)

Intervention duration: 24 months. Follow-up: 24 months. Intervention design: aimed to provide a social protection system through regular and predictable cash transfers to families living with orphans or vulnerable children in order to encourage fostering and retention of orphans or vulnerable children within their families and communities, and to promote their human capital development; targeted to poor households with one or more non-welfare (i.e. not receiving any other cash transfers) orphans or vulnerable children; eligibility determined through screening geographically, by a community committee and a survey; provided a total amount of approximately USD 352.80 (USD 14.70 per month for 24 months; 29.6% of the annual GDP per capita); paid in-hand every second month; and fuzzy design: the cash transfer was conditional on attendance of a health facility for immunisations among children aged 0-1 year, growth monitoring and vitamin supplements among children aged 0–5 years, school enrolment among children aged 6-18 years and attendance of awareness sessions among adult parents or caregivers, but non-compliance was not penalised in 4 out of 7 clusters.

Outcomes
Primary outcomes
- Growth check in previous 6 months
- Fully vaccinated, currently
- Use of any health service, previous 1 to 12 months (assessed with: received treatment for child diarrhoea, acute respiratory infection or fever at a health facility)
- Moderate stunting
- Moderate underweight
- Level of dietary diversity

Secondary outcomes
- Attends school
- Child engages in child labour, currently (assessed with: worked for pay)
- Is extremely poor, currently

Notes
Intervention context: pilot programme of the Government of Kenya called the Cash Transfer Programme for Orphans and Vulnerable Children; implemented through Ministry of Gender, Children and Social Development; population coverage was approximately 15,000 recipients in July 2009; programme uptake was 97% among initial recipients (i.e. 3% of recipients dropped out of the programme); and the total programme cost of the UCT was USD 9.96 million in the 7 pilot districts between July 2006 and June 2009. Funder of studies: United Nations Children’s Fund, Government of Kenya and United Kingdom Department for International Development. Conflict of interest: none identified.
### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
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<td>Random sequence generation unclear</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
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</tr>
<tr>
<td>Participants recruited after cluster allocation (recruitment bias in cluster-RCTs)</td>
<td>High risk</td>
<td>Participants recruited after cluster allocation</td>
</tr>
<tr>
<td>Baseline outcome measurements similar</td>
<td>Low risk</td>
<td>No baseline differences (P &lt; 0.05) between the UCT intervention group and the control group in the outcome measurements</td>
</tr>
<tr>
<td>Baseline characteristics similar</td>
<td>High risk</td>
<td>Baseline differences (P &lt; 0.05) between the UCT intervention group and the control group in 4 characteristics outcome measurements. There were differences in the proportion of participants who were male (0.55 compared with 0.52, P &lt; 0.05), whose mother was dead (0.44 compared with 0.30, P &lt; 0.05) and whose mother is a caregiver (0.44 vs. 0.61, P &lt; 0.01), as well as the age of the participants’ caregiver (48.7 compared with 40.8, P &lt; 0.01).</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>Blinding of participants not possible, and blinding of personnel unclear</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All subjectively measured outcomes</td>
<td>High risk</td>
<td>Outcome assessors not blinded</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All objectively measured outcomes</td>
<td>Low risk</td>
<td>Outcome assessors not blinded. Outcome is unlikely to be influenced by lack of blinding because it is objectively measured.</td>
</tr>
<tr>
<td>Contamination</td>
<td>Low risk</td>
<td>4% of households in the UCT intervention group did not meet the eligibility criteria, with 3% of households containing no orphan or vulnerable child and 1% not being poor.</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>High risk</td>
<td>Initial survey non-response was high (13%). The proportion of participants lost to follow-up was large (18%), the proportion of participants lost to follow-up in the intervention group (14%) was smaller than that in the control group (24%; statistical significance not tested). Attrition in the intervention group and in the control group differed by several population characteristics, including by location. The number and percentage of missing communities and households and the percentage of missing values per outcome were unclear.</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Study protocol identified, but we were unable to locate a copy of it</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>Intervention occurred during a phase of postelection violence, which may have impacted the intervention’s effectiveness</td>
</tr>
</tbody>
</table>

Ward 2010 (Continued)

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
**CCT:** conditional cash transfer; **SD:** standard deviation; **UCT:** unconditional cash transfer; **VCT:** voluntary counselling and (HIV) testing.

### Characteristics of excluded studies [ordered by study ID]

<table>
<thead>
<tr>
<th>Study</th>
<th>Reason for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Akee 2013</td>
<td>No eligible study population studied</td>
</tr>
<tr>
<td>Aker 2011</td>
<td>UCT for assistance in humanitarian disasters studied (see Pega 2015a for systematic review of this type of UCT)</td>
</tr>
<tr>
<td>Aker 2013</td>
<td>UCT for assistance in humanitarian disasters studied</td>
</tr>
<tr>
<td>Angelucci 2009</td>
<td>No eligible UCT studied</td>
</tr>
<tr>
<td>Attanasio 2015</td>
<td>No eligible UCT studied</td>
</tr>
<tr>
<td>Ayuku 2013</td>
<td>No eligible UCT studied</td>
</tr>
<tr>
<td>Benhassine 2013</td>
<td>No eligible outcome studied</td>
</tr>
<tr>
<td>Buller 2016</td>
<td>No eligible UCT studied</td>
</tr>
<tr>
<td>Buser 2014</td>
<td>Ineligible study type used</td>
</tr>
<tr>
<td>Coetzee 2013</td>
<td>No eligible UCT studied</td>
</tr>
<tr>
<td>Doocey 2017</td>
<td>UCT for assistance in humanitarian disasters studied</td>
</tr>
<tr>
<td>Fenn 2013</td>
<td>UCT for assistance in humanitarian disasters studied</td>
</tr>
<tr>
<td>Fenn 2017</td>
<td>UCT for assistance in humanitarian disasters studied</td>
</tr>
<tr>
<td>Hidrobo 2013</td>
<td>No eligible outcome studied</td>
</tr>
<tr>
<td>Holmqvist 2011</td>
<td>Ineligible study type used</td>
</tr>
<tr>
<td>Kenya CT-OVC Evaluation Team 2012</td>
<td>No eligible outcome studied</td>
</tr>
<tr>
<td>Langendorf 2013</td>
<td>UCT for assistance in humanitarian disasters studied</td>
</tr>
<tr>
<td>Macours 2008</td>
<td>UCT for assistance in humanitarian disasters studied</td>
</tr>
<tr>
<td>Ndlovu 2013</td>
<td>No eligible outcome studied</td>
</tr>
<tr>
<td>Park 2013</td>
<td>No eligible UCT studied</td>
</tr>
<tr>
<td>Pereznieta 2014</td>
<td>Ineligible study type used</td>
</tr>
<tr>
<td>Plagerson 2011</td>
<td>Ineligible study type used</td>
</tr>
<tr>
<td>Poulsen 2011</td>
<td>Ineligible study type used</td>
</tr>
<tr>
<td>Pratinidhi 2014</td>
<td>Ineligible study type used</td>
</tr>
<tr>
<td>Rocha 2013</td>
<td>No eligible UCT studied</td>
</tr>
<tr>
<td>Study</td>
<td>Reason for exclusion</td>
</tr>
<tr>
<td>-----------------------</td>
<td>----------------------------------------------------------</td>
</tr>
<tr>
<td>Santos 2011</td>
<td>No eligible outcome studied</td>
</tr>
<tr>
<td>Skoufias 2013</td>
<td>No eligible outcome studied</td>
</tr>
<tr>
<td>Skovdal 2012</td>
<td>Ineligible study type used</td>
</tr>
<tr>
<td>Tadesse 2014</td>
<td>Ineligible study type used</td>
</tr>
<tr>
<td>Tonguet-Papucci 2017</td>
<td>UCT for assistance in humanitarian disasters studied</td>
</tr>
</tbody>
</table>

**UCT**: unconditional cash transfer.

**Characteristics of studies awaiting assessment** *(ordered by study ID)*

**Abdoulayi 2014**

<table>
<thead>
<tr>
<th>Methods</th>
<th>Cluster-randomised controlled trial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>4352 households</td>
</tr>
<tr>
<td>Interventions</td>
<td>Unconditional cash transfer (Social Cash Transfer Program, Malawi)</td>
</tr>
<tr>
<td>Outcomes</td>
<td>Health services use, stunting, disease prevalence, food security, dietary diversity, level of depression</td>
</tr>
<tr>
<td>Notes</td>
<td>—</td>
</tr>
</tbody>
</table>

**AIR 2014**

<table>
<thead>
<tr>
<th>Methods</th>
<th>Cluster-randomised controlled trial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>2630 households</td>
</tr>
<tr>
<td>Interventions</td>
<td>Unconditional cash transfer (Harmonised Social Cash Transfer Programme, Zimbabwe)</td>
</tr>
<tr>
<td>Outcomes</td>
<td>Food security, dietary diversity</td>
</tr>
<tr>
<td>Notes</td>
<td>—</td>
</tr>
</tbody>
</table>

**Benedetti 2016**

<table>
<thead>
<tr>
<th>Methods</th>
<th>Cluster-randomised controlled trial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>4416 households</td>
</tr>
<tr>
<td>Interventions</td>
<td>Fuzzy unconditional or conditional cash transfer (Bono 10,000, Honduras)</td>
</tr>
<tr>
<td>Outcomes</td>
<td>Health services use</td>
</tr>
<tr>
<td>Notes</td>
<td>—</td>
</tr>
<tr>
<td><strong>Methods</strong></td>
<td>Cluster-randomised controlled trial</td>
</tr>
<tr>
<td>------------</td>
<td>-----------------------------------</td>
</tr>
<tr>
<td><strong>Participants</strong></td>
<td>Unclear</td>
</tr>
<tr>
<td><strong>Interventions</strong></td>
<td>Unconditional cash transfer (Social Cash Transfer Program, Malawi)</td>
</tr>
<tr>
<td><strong>Outcomes</strong></td>
<td>Disease prevalence, food security, household dietary diversity</td>
</tr>
<tr>
<td><strong>Notes</strong></td>
<td>—</td>
</tr>
</tbody>
</table>

**Cluver 2013**

<table>
<thead>
<tr>
<th><strong>Methods</strong></th>
<th>Cohort study</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Participants</strong></td>
<td>3515 participants</td>
</tr>
<tr>
<td><strong>Interventions</strong></td>
<td>Unconditional cash transfer (Child Support Grant or Foster Child Grant, South Africa)</td>
</tr>
<tr>
<td><strong>Outcomes</strong></td>
<td>Disease prevalence</td>
</tr>
<tr>
<td><strong>Notes</strong></td>
<td>—</td>
</tr>
</tbody>
</table>

**Davis 2016**

<table>
<thead>
<tr>
<th><strong>Methods</strong></th>
<th>Cluster-randomised controlled trials or cohort studies</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Participants</strong></td>
<td>Various</td>
</tr>
<tr>
<td><strong>Interventions</strong></td>
<td>Unconditional cash transfers (Cash Transfer Program for Orphans and Vulnerable Children, Kenya; Child Grants Programme, Lesotho; Child Support Grant, South Africa; Harmonized Social Cash Transfer Programme, Zimbabwe; Social Cash Transfer Programme, Malawi; Social Cash Transfer Programme, Zambia; and Tigray Pilot Social Cash Transfer Programme, Ethiopia)</td>
</tr>
<tr>
<td><strong>Outcomes</strong></td>
<td>Health services use, anthropometric measures, disease prevalence, food security, household dietary diversity, mental health</td>
</tr>
<tr>
<td><strong>Notes</strong></td>
<td>—</td>
</tr>
</tbody>
</table>

**Gangophadyay 2015**

<table>
<thead>
<tr>
<th><strong>Methods</strong></th>
<th>Randomised controlled trial</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Participants</strong></td>
<td>450 households</td>
</tr>
<tr>
<td><strong>Interventions</strong></td>
<td>Unconditional cash transfer (experiment)</td>
</tr>
<tr>
<td><strong>Outcomes</strong></td>
<td>Food security</td>
</tr>
</tbody>
</table>
Gangopadhyay 2015

Methods
Cluster-randomised controlled trial

Participants
1481 participants

Interventions
Unconditional cash transfer (experiment)

Outcomes
Mortality, anthropometric measures, household dietary diversity

Notes
—

Grellety 2017

Methods
Cluster-randomised controlled trial

Participants
1481 participants

Interventions
Unconditional cash transfer (experiment)

Outcomes
Mortality, anthropometric measures, household dietary diversity

Notes
—

Handa 2014a

Methods
Cluster-randomised controlled trial

Participants
1549 children and young people

Interventions
Unconditional cash transfer (Cash Transfer Program for Orphans and Vulnerable Children, Kenya)

Outcomes
Sexual health risk behaviours

Notes
—

Hjelm 2017

Methods
Cluster-randomised controlled trials

Participants
14,565 participants and 15,630 participants, respectively

Interventions
Unconditional cash transfers (Child Grant Program, Zambia; and Multiple Category Cash Transfer Program, Zambia)

Outcomes
Food security

Notes
—

Kilburn 2016

Methods
Cluster-randomised controlled trial

Participants
1960 participants

Interventions
Unconditional cash transfer (Cash Transfer Program for orphans and Vulnerable Children, Kenya)

Outcomes
Mental health
### Kilburn 2016 (Continued)

**Notes** —

### Lawlor 2015

<table>
<thead>
<tr>
<th>Methods</th>
<th>Cluster-randomised controlled trial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>2455 participants</td>
</tr>
<tr>
<td>Interventions</td>
<td>(Child Grant Program, Zambia)</td>
</tr>
<tr>
<td>Outcomes</td>
<td>Food security, household dietary diversity</td>
</tr>
<tr>
<td>Notes</td>
<td>—</td>
</tr>
</tbody>
</table>

### Olajide 2016

<table>
<thead>
<tr>
<th>Methods</th>
<th>Cluster-randomised controlled trials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>6236 participants</td>
</tr>
<tr>
<td>Interventions</td>
<td>Unconditional cash transfers (Ekiti State Scheme, Nigeria)</td>
</tr>
<tr>
<td>Outcomes</td>
<td>Mental health</td>
</tr>
<tr>
<td>Notes</td>
<td>—</td>
</tr>
</tbody>
</table>

### Tiwari 2016

<table>
<thead>
<tr>
<th>Methods</th>
<th>Cluster-randomised controlled trials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>1540 participants; 739 participants; and 1256 participants, respectively</td>
</tr>
<tr>
<td>Interventions</td>
<td>Unconditional cash transfers (Cash Transfer for Orphans and Vulnerable Children, Kenya; Child Grants Program, Lesotho; and Social Cash Transfer Program, Zambia)</td>
</tr>
<tr>
<td>Outcomes</td>
<td>Food security, household dietary diversity</td>
</tr>
<tr>
<td>Notes</td>
<td>—</td>
</tr>
</tbody>
</table>

### Characteristics of ongoing studies [ordered by study ID]

#### Galárraga 2014

<table>
<thead>
<tr>
<th>Trial name or title</th>
<th>Conditional economic incentives to reduce HIV risk: a pilot in Mexico</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methods</td>
<td>Randomised controlled trial; methods unclear</td>
</tr>
</tbody>
</table>
### Galárraga 2014 (Continued)

| Participants | 267 adults (18-40 years); all men who had receptive or penetrative anal sex in exchange for money in the last 6 months; Mexico City, Mexico |
| Interventions | Unconditional cash transfer (experiment) |
| Outcomes | Primary outcomes: health outcomes (disease risk and prevalence) |
| Starting date | Unclear |
| Contact information | Dr Omar Galárraga  
Department of Health Services Policy and Practice  
Brown University School of Public Health  
121 South Main Street  
Box G-121S-7  
Providence, RI 02912  
USA |

### O'Leary 2011

| Trial name or title | Benazir Income Support Programme impact evaluation |
| Methods | Controlled before and after study; difference-in-differences methods |
| Participants | Number and type of participants unclear; Pakistan |
| Interventions | Unconditional cash transfer (Benazir Income Support Programme) |
| Outcomes | Primary outcomes: use of health services (preventive and other) and health outcomes (anthropometric measures, disease risk and prevalence and nutrition)  
Secondary outcomes: (assets, education, labour force participation and poverty) |
| Starting date | 2011 |
| Contact information | Mr Sean O'Leary  
Oxford Policy Management  
6 St Aldates Courtyard  
38 St Aldates  
Oxford OX1 1BN  
United Kingdom  
phone: +44 (0)1865 207 300 |

Notes —
Oxford Policy Management 2013

Trial name or title
Uganda Social Assistance Grants for Empowerment Programme impact evaluation

Methods
Controlled before and after study; regression discontinuity methods

Participants
Members of 3980 households; 48 subcounties, 8 programme districts, Uganda

Interventions
Unconditional cash transfers (Vulnerable Families Support Grant; and Senior Citizens Grant)

Outcomes
Primary outcomes: use of health services (other) and health outcomes (food security and nutrition)
Secondary outcomes: social determinants of health (education, housing, labour force participation and poverty) and healthcare expenditure

Starting date
2011

Contact information
Expanding Social Protection Programme
Ministry of Gender, Labour and Social Development
Plot 9, Lourdel Road
P.O. Box 28240 Kampala
Uganda
phone: +25 60414534202
email: esp@socialprotection.go.ug

Notes
—

DATA AND ANALYSES

Comparison 1. Unconditional cash transfer versus no unconditional cash transfer: health services use

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Has ever had birth registered</td>
<td>4</td>
<td></td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>2 Has had a growth check in previous 6 months</td>
<td>3</td>
<td>2261</td>
<td>Odds Ratio (Random, 95% CI)</td>
<td>1.11 [0.98, 1.24]</td>
</tr>
<tr>
<td>3 Is up-to-date on vaccination calendar</td>
<td>3</td>
<td>563</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>1.02 [0.90, 1.15]</td>
</tr>
<tr>
<td>4 Has been given any treatment for parasites in previous year</td>
<td>1</td>
<td>1478</td>
<td>Odds Ratio (Random, 95% CI)</td>
<td>1.28 [1.06, 1.54]</td>
</tr>
<tr>
<td>5 Has used any health service in previous 1 to 12 months</td>
<td>5</td>
<td>4972</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>1.04 [1.00, 1.09]</td>
</tr>
</tbody>
</table>
### Analysis 1.1. Comparison 1 Unconditional cash transfer versus no unconditional cash transfer: health services use, Outcome 1 Has ever had birth registered.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT n/N</th>
<th>No UCT n/N</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pellerano 2014</td>
<td>184/333</td>
<td>61/333</td>
<td>3.02 [2.36, 3.86]</td>
<td></td>
</tr>
<tr>
<td>Robertson 2012</td>
<td>41/92</td>
<td>64/132</td>
<td>0.92 [0.69, 1.23]</td>
<td></td>
</tr>
<tr>
<td>Seidenfeld 2013</td>
<td>132/556</td>
<td>140/556</td>
<td>0.94 [0.77, 1.16]</td>
<td></td>
</tr>
<tr>
<td>Ward 2010</td>
<td>81/251</td>
<td>41/123</td>
<td>0.97 [0.71, 1.32]</td>
<td></td>
</tr>
</tbody>
</table>

Favours no UCT: 0.2 0.5 1 2 5  Favours UCT

### Analysis 1.2. Comparison 1 Unconditional cash transfer versus no unconditional cash transfer: health services use, Outcome 2 Has had a growth check in previous 6 months.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT N</th>
<th>No UCT N</th>
<th>log(Odds Ratio) (SE)</th>
<th>Odds Ratio IV, Random, 95% CI</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paxson 2007</td>
<td>555</td>
<td>555</td>
<td>0.1 (0.08)</td>
<td>1.12 [0.95, 1.31]</td>
<td>55.74%</td>
</tr>
<tr>
<td>Paxson 2007</td>
<td>185</td>
<td>185</td>
<td>0.1 (0.11)</td>
<td>1.07 [0.86, 1.33]</td>
<td>29.48%</td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>238</td>
<td>238</td>
<td>0.1 (0.19)</td>
<td>1.09 [0.75, 1.59]</td>
<td>9.88%</td>
</tr>
<tr>
<td>Ward 2010</td>
<td>199</td>
<td>106</td>
<td>0.2 (0.27)</td>
<td>1.25 [0.73, 2.12]</td>
<td>4.89%</td>
</tr>
</tbody>
</table>

Total (95% CI): 1.11 [0.98, 1.24]

Heterogeneity: Tau²=0; Chi²=2.07, df=2 (P=0.36); I²=11%
Test for overall effect: Z=0.27 (P=0.79)

Favours no UCT: 0.2 0.5 1 2 5  Favours UCT

### Analysis 1.3. Comparison 1 Unconditional cash transfer versus no unconditional cash transfer: health services use, Outcome 3 Is up-to-date on vaccination calendar.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT n/N</th>
<th>No UCT n/N</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pellerano 2014</td>
<td>41/84</td>
<td>48/84</td>
<td>18.34%</td>
<td>0.85 [0.64, 1.14]</td>
</tr>
<tr>
<td>Robertson 2012</td>
<td>69/91</td>
<td>92/131</td>
<td>55.83%</td>
<td>1.08 [0.92, 1.27]</td>
</tr>
<tr>
<td>Ward 2010</td>
<td>76/118</td>
<td>35/55</td>
<td>25.83%</td>
<td>1.01 [0.81, 1.29]</td>
</tr>
</tbody>
</table>

Total (95% CI): 1.02 [0.91, 1.15]

Total events: 186 (UCT), 175 (No UCT)
Heterogeneity: Tau²=0; Chi²=2.67, df=2 (P=0.36); I²=13.2%
Test for overall effect: Z=0.27 (P=0.79)

Favours no UCT: 0.2 0.5 1 2 5  Favours UCT
### Analysis 1.4. Comparison 1 Unconditional cash transfer versus no unconditional cash transfer: health services use, Outcome 4 Has been given any treatment for parasites in previous year.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>log(Odds Ratio)</th>
<th>Odds Ratio</th>
<th>Weight</th>
<th>Odds Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>N</td>
<td>(SE)</td>
<td>IV, Random, 95% CI</td>
<td></td>
<td></td>
<td>IV, Random, 95% CI</td>
</tr>
<tr>
<td>Paxson 2007</td>
<td>554</td>
<td>554</td>
<td>0.2 (0.08)</td>
<td></td>
<td>66.5%</td>
<td>1.2 (1.02, 1.4)</td>
</tr>
<tr>
<td></td>
<td>185</td>
<td>185</td>
<td>0.4 (0.14)</td>
<td></td>
<td>33.5%</td>
<td>1.46 (1.11, 1.92)</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td></td>
<td></td>
<td>100%</td>
<td></td>
<td>1.28 (1.06, 1.54)</td>
<td></td>
</tr>
</tbody>
</table>

Heterogeneity: Tau^2=0.01; Chi^2=1.54, df=1(P=0.21); I^2=35%
Test for overall effect: Z=2.62(P=0.01)

### Analysis 1.5. Comparison 1 Unconditional cash transfer versus no unconditional cash transfer: health services use, Outcome 5 Has used any health service in previous 1 to 12 months.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>n/N</td>
<td>n/N</td>
<td></td>
<td>M-H, Random, 95% CI</td>
<td></td>
<td>M-H, Random, 95% CI</td>
</tr>
<tr>
<td>Luseno 2012</td>
<td>45/50</td>
<td>35/44</td>
<td></td>
<td>6.37%</td>
<td>1.13 (0.95, 1.35)</td>
</tr>
<tr>
<td>Oxford Policy Management 2012</td>
<td>130/154</td>
<td>122/154</td>
<td></td>
<td>17.31%</td>
<td>1.07 (0.96, 1.18)</td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>188/612</td>
<td>159/612</td>
<td></td>
<td>6.16%</td>
<td>1.18 (0.99, 1.41)</td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>148/730</td>
<td>141/730</td>
<td></td>
<td>4.64%</td>
<td>1.05 (0.85, 1.29)</td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>130/254</td>
<td>127/254</td>
<td></td>
<td>6.68%</td>
<td>1.02 (0.86, 1.22)</td>
</tr>
<tr>
<td>Seidenfeld 2013</td>
<td>434/551</td>
<td>436/551</td>
<td></td>
<td>48.47%</td>
<td>1.09 (0.84, 1.38)</td>
</tr>
<tr>
<td>Ward 2010</td>
<td>147/180</td>
<td>71/96</td>
<td></td>
<td>10.37%</td>
<td>1.10 (0.96, 1.27)</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>2531</td>
<td>2441</td>
<td>100%</td>
<td>1.04 (1.09)</td>
<td></td>
</tr>
</tbody>
</table>

Total events: 1222 (UCT), 1091 (No UCT)
Heterogeneity: Tau^2=0; Chi^2=6.15, df=6(P=0.41); I^2=2.45%
Test for overall effect: Z=1.8(P=0.07)

### Comparison 2. Unconditional cash transfer versus no unconditional cash transfer: health outcomes

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Is moderately stunted</td>
<td>2</td>
<td>551</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>0.96 [0.75, 1.21]</td>
</tr>
<tr>
<td>2 Height for age (standard deviations)</td>
<td>2</td>
<td>7545</td>
<td>MD or Difference-in-Differences (SDs) (Random, 95% CI)</td>
<td>0.04 [-0.05, 0.13]</td>
</tr>
<tr>
<td>3 Is moderately underweight</td>
<td>3</td>
<td>701</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>1.00 [0.75, 1.32]</td>
</tr>
<tr>
<td>4 Has had any illness in previous 2 weeks to 3 months</td>
<td>5</td>
<td>8446</td>
<td>Odds Ratio (Random, 95% CI)</td>
<td>0.73 [0.57, 0.93]</td>
</tr>
<tr>
<td>5 Has been food secure in previous month</td>
<td>3</td>
<td></td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>Totals not selected</td>
</tr>
</tbody>
</table>

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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### Analysis 2.1. Comparison 2 Unconditional cash transfer versus no unconditional cash transfer: health outcomes, Outcome 1 is moderately stunted.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxford Policy Management 2012</td>
<td>42/143</td>
<td>45/143</td>
<td>M-H, Random, 95% CI</td>
<td>46.15%</td>
<td>0.93 [0.66, 1.33]</td>
</tr>
<tr>
<td>Ward 2010</td>
<td>57/159</td>
<td>39/106</td>
<td>M-H, Random, 95% CI</td>
<td>53.85%</td>
<td>0.97 [0.71, 1.35]</td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td>302</td>
<td>249</td>
<td>M-H, Random, 95% CI</td>
<td>100%</td>
<td>0.96 [0.75, 1.21]</td>
</tr>
</tbody>
</table>

Total events: 99 (UCT), 84 (No UCT)
Heterogeneity: Tau²=0; Chi²=0.03, df=1(P=0.86); I²=0%
Test for overall effect: Z=0.38(P=0.71)

### Analysis 2.2. Comparison 2 Unconditional cash transfer versus no unconditional cash transfer: health outcomes, Outcome 2 Height for age (standard deviations).

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>MD or Difference-in-Differences (SDs)</th>
<th>Weight</th>
<th>MD or Difference-in-Differences (SDs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paxson 2007</td>
<td>695</td>
<td>695</td>
<td>0 (0.05)</td>
<td>M-H, Random, 95% CI</td>
<td>76.42%</td>
</tr>
<tr>
<td>Seidenfeld 2013</td>
<td>3078</td>
<td>3077</td>
<td>0.1 (0.09)</td>
<td>M-H, Random, 95% CI</td>
<td>23.58%</td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td></td>
<td></td>
<td>M-H, Random, 95% CI</td>
<td>100%</td>
<td>0.04 [-0.05, 0.13]</td>
</tr>
</tbody>
</table>

Heterogeneity: Tau²=0; Chi²=0.15, df=1(P=0.7); I²=0%
Test for overall effect: Z=0.9(P=0.37)

### Analysis 2.3. Comparison 2 Unconditional cash transfer versus no unconditional cash transfer: health outcomes, Outcome 3 Is moderately underweight.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxford Policy Management 2012</td>
<td>36/143</td>
<td>34/143</td>
<td>M-H, Random, 95% CI</td>
<td>48.7%</td>
<td>1.06 [0.71, 1.59]</td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>12/74</td>
<td>17/74</td>
<td>M-H, Random, 95% CI</td>
<td>18.28%</td>
<td>0.71 [0.36, 1.37]</td>
</tr>
</tbody>
</table>

Favours UCT

---

Cochrane Database of Systematic Reviews

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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## Analysis 2.4. Comparison 2 Unconditional cash transfer versus no unconditional cash transfer: health outcomes, Outcome 4 Has had any illness in previous 2 weeks to 3 months.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT n/N</th>
<th>No UCT n/N</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
<th>Weight %</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ward 2010</td>
<td>34/162</td>
<td>20/105</td>
<td>33.02%</td>
<td>1.1[0.67,1.81]</td>
<td></td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>379</td>
<td>322</td>
<td>100%</td>
<td>1[0.75,1.32]</td>
<td></td>
</tr>
</tbody>
</table>

Total events: 82 (UCT), 71 (No UCT)
Heterogeneity: Tau²=0.00; Chi²=1.28, df=2(P=0.53); I²=0% 
Test for overall effect: Z=0.03(P=0.98)

### Favours UCT

## Analysis 2.5. Comparison 2 Unconditional cash transfer versus no unconditional cash transfer: health outcomes, Outcome 5 Has been food secure in previous month.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT n/N</th>
<th>No UCT n/N</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
<th>Weight %</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baird 2010</td>
<td>283</td>
<td>2130</td>
<td>28.58%</td>
<td>0.91[0.72,1.16]</td>
<td></td>
</tr>
<tr>
<td>Cunha 2014</td>
<td>2275</td>
<td>2275</td>
<td>26.17%</td>
<td>0.63[0.48,0.83]</td>
<td></td>
</tr>
<tr>
<td>Luseno 2012</td>
<td>50</td>
<td>59</td>
<td>7.38%</td>
<td>0.58[0.25,1.31]</td>
<td></td>
</tr>
<tr>
<td>Oxford Policy Management 2012</td>
<td>307</td>
<td>307</td>
<td>14.03%</td>
<td>1.04[0.61,1.77]</td>
<td></td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>380</td>
<td>380</td>
<td>23.84%</td>
<td>0.56[0.41,0.77]</td>
<td></td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>100%</td>
<td></td>
<td>0.73[0.57,0.93]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Heterogeneity: Tau²=0.04; Chi²=9.36, df=4(P=0.05); I²=57.27% 
Test for overall effect: Z=2.52(P=0.01)

### Favours UCT

## Analysis 2.6. Comparison 2 Unconditional cash transfer versus no unconditional cash transfer: health outcomes, Outcome 6 Level of dietary diversity (Household Dietary Diversity Score) in previous week.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT n/N</th>
<th>No UCT n/N</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
<th>Weight %</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxford Policy Management 2012</td>
<td>1224</td>
<td>1212</td>
<td>19.31%</td>
<td>0.93[0.8,1.1]</td>
<td></td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>1344</td>
<td>1344</td>
<td>26.63%</td>
<td>0.11[0.13,0.35]</td>
<td></td>
</tr>
</tbody>
</table>

### Favours UCT
### Analysis 2.7. Comparison 2 Unconditional cash transfer versus no unconditional cash transfer: health outcomes, Outcome 7 Level of depression (Center for Epidemiologic Studies Depression Score).

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SE)</th>
<th>Weight</th>
<th>Mean Difference (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paxson 2007</td>
<td>262</td>
<td>261</td>
<td>-0.2 (0.16)</td>
<td>30.16%</td>
<td>-0.21 [-0.52, 0.1]</td>
</tr>
<tr>
<td>Paxson 2007</td>
<td>262</td>
<td>261</td>
<td>0 (0.09)</td>
<td>69.84%</td>
<td>0 [-0.18, 0.18]</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td></td>
<td></td>
<td></td>
<td>100%</td>
<td>-0.06 [-0.25, 0.13]</td>
</tr>
</tbody>
</table>

**Heterogeneity:** Tau²=0.01; Chi²=1.31, df=1(P=0.25); I²=23.58%

**Test for overall effect:** Z=0.66(P=0.51)

---

### Comparison 3. Unconditional cash transfers versus no unconditional cash transfers: social determinants of health

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Owns livestock in previous year</td>
<td>2</td>
<td></td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>Totals not selected</td>
</tr>
<tr>
<td>2 Attends school</td>
<td>6</td>
<td>4800</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>1.06 [1.03, 1.09]</td>
</tr>
<tr>
<td>3 Works</td>
<td>3</td>
<td></td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>3.1 Children</td>
<td>3</td>
<td>2448</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>0.91 [0.80, 1.03]</td>
</tr>
<tr>
<td>3.2 Adults</td>
<td>2</td>
<td>1700</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>1.00 [0.95, 1.06]</td>
</tr>
<tr>
<td>4 Level of parenting quality (Home Observation Measurement of the Environment Score) (standard deviations)</td>
<td>1</td>
<td>1118</td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>0.09 [-0.25, 0.42]</td>
</tr>
</tbody>
</table>
## Analysis 3.1. Comparison 3 Unconditional cash transfers versus no unconditional cash transfers: social determinants of health, Outcome 1 Owns livestock in previous year.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT n/N</th>
<th>No UCT n/N</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxford Policy Management 2012</td>
<td>127/199</td>
<td>162/199</td>
<td>0.78 [0.69, 0.89]</td>
<td>0.78 [0.69, 0.89]</td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>294/444</td>
<td>277/444</td>
<td>0.94 [0.89, 1.00]</td>
<td>0.94 [0.89, 1.00]</td>
</tr>
</tbody>
</table>

Heterogeneity: Tau^2=0; Chi^2=6.7, df=7(P=0.46); I^2=0%

Test for overall effect: Z =4.19(P <0.0001)

## Analysis 3.2. Comparison 3 Unconditional cash transfers versus no unconditional cash transfers: social determinants of health, Outcome 2 Attends school.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT n/N</th>
<th>No UCT n/N</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
<th>Weight</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baird 2010</td>
<td>314/348</td>
<td>484/577</td>
<td>30.28%</td>
<td>0.77 [0.56, 1.08]</td>
<td></td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>529/606</td>
<td>516/606</td>
<td>36.94%</td>
<td>1.03 [0.98, 1.07]</td>
<td></td>
</tr>
<tr>
<td>Robertson 2012</td>
<td>173/201</td>
<td>116/151</td>
<td>6.96%</td>
<td>1.12 [1.01, 1.24]</td>
<td></td>
</tr>
<tr>
<td>Robertson 2012</td>
<td>170/193</td>
<td>120/151</td>
<td>8.09%</td>
<td>1.11 [1.01, 1.22]</td>
<td></td>
</tr>
<tr>
<td>Seidenfeld 2013</td>
<td>70/452</td>
<td>61/452</td>
<td>0.74%</td>
<td>1.15 [0.84, 1.58]</td>
<td></td>
</tr>
<tr>
<td>Ward 2010</td>
<td>227/257</td>
<td>97/114</td>
<td>9.51%</td>
<td>1.14 [1.09]</td>
<td></td>
</tr>
<tr>
<td>Ward 2010</td>
<td>106/138</td>
<td>56/74</td>
<td>2.99%</td>
<td>1.03 [0.87, 1.19]</td>
<td></td>
</tr>
</tbody>
</table>

Total (95% CI) | 2435 | 2365 | 100% | 1.06 [1.03, 1.09] |

Total events: 1758 (UCT), 1598 (No UCT)

Heterogeneity: Tau^2=0; Chi^2=6.7, df=7(P=0.46); I^2=0%

Test for overall effect: Z =4.19(P <0.0001)
### Analysis 3.4. Comparison 3 Unconditional cash transfers versus no unconditional cash transfers: social determinants of health, Outcome 4 Level of parenting quality (Home Observation Measurement of the Environment Score) (standard deviations).

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SDs)</th>
<th>Weight</th>
<th>Mean Difference (SDs)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>(SE)</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
</tr>
<tr>
<td>Paxson 2007</td>
<td>280</td>
<td>279</td>
<td>-0.14</td>
<td>64.66%</td>
<td>-0.04[-0.31,0.23]</td>
</tr>
<tr>
<td>Paxson 2007</td>
<td>280</td>
<td>279</td>
<td>0.24</td>
<td>35.34%</td>
<td>0.32[0.15,0.79]</td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td></td>
<td></td>
<td></td>
<td>100%</td>
<td>0.09[-0.25,0.42]</td>
</tr>
</tbody>
</table>

Heterogeneity: Tau²=0.03; Chi²=1.68, df=1(P=0.2); I²=40.43%
Test for overall effect: Z=0.51(P=0.61)

### Analysis 3.5. Comparison 3 Unconditional cash transfers versus no unconditional cash transfers: social determinants of health, Outcome 5 Is extremely poor.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H, Random, 95% CI</td>
<td></td>
<td>M-H, Random, 95% CI</td>
</tr>
<tr>
<td>Oxford Policy Management 2012</td>
<td>175/199</td>
<td>193/199</td>
<td>33.17%</td>
<td>0.91[0.86,0.96]</td>
<td></td>
</tr>
<tr>
<td>Pellerano 2014</td>
<td>307/444</td>
<td>299/444</td>
<td>23.31%</td>
<td>1.03[0.94,1.12]</td>
<td></td>
</tr>
<tr>
<td>Seidenfeld 2013</td>
<td>483/533</td>
<td>512/533</td>
<td>41.15%</td>
<td>0.94[0.91,0.97]</td>
<td></td>
</tr>
<tr>
<td>Ward 2010</td>
<td>49/234</td>
<td>30/98</td>
<td>2.37%</td>
<td>0.68[0.46,1.01]</td>
<td></td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td>1410</td>
<td>1274</td>
<td>100%</td>
<td>0.94[0.89,1]</td>
<td></td>
</tr>
</tbody>
</table>

Heterogeneity: Tau²=0; Chi²=8.37, df=3(P=0.04); I²=64.17%
Test for overall effect: Z=1.89(P=0.06)
### Comparison 4. Unconditional cash transfer versus no unconditional cash transfer: health equity

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Height for age (standard deviations) by rural-urban residency, currently</td>
<td>1</td>
<td></td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>1.1 Living in rural areas</td>
<td>1</td>
<td>654</td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>-0.09 [-0.36, 0.18]</td>
</tr>
<tr>
<td>1.2 Living in urban areas</td>
<td>1</td>
<td>542</td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>0.13 [-0.11, 0.37]</td>
</tr>
<tr>
<td>2 Height for age (standard deviations) by income poverty status, currently</td>
<td>1</td>
<td></td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>2.1 Living in income poverty</td>
<td>1</td>
<td>458</td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>0.04 [-0.12, 0.20]</td>
</tr>
<tr>
<td>2.2 Not living in income poverty</td>
<td>1</td>
<td>457</td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>0.04 [-0.06, 0.14]</td>
</tr>
<tr>
<td>3 Has had any illness in previous 2 weeks to 3 months</td>
<td>1</td>
<td></td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>3.1 Girls</td>
<td>1</td>
<td>428</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>0.70 [0.55, 0.90]</td>
</tr>
<tr>
<td>3.2 Boys</td>
<td>1</td>
<td>440</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>0.69 [0.54, 0.88]</td>
</tr>
<tr>
<td>4 Food security index by gender</td>
<td>1</td>
<td></td>
<td>Difference-in-Differences (SDs) (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>4.1 Women</td>
<td>1</td>
<td>686</td>
<td>Difference-in-Differences (SDs) (Random, 95% CI)</td>
<td>0.27 [-1.49, 2.03]</td>
</tr>
<tr>
<td>4.2 Men</td>
<td>1</td>
<td>686</td>
<td>Difference-in-Differences (SDs) (Random, 95% CI)</td>
<td>0.23 [-1.53, 1.99]</td>
</tr>
<tr>
<td>5 Dietary diversity (Household Dietary Diversity Score) in previous week by rural-urban residency</td>
<td>1</td>
<td></td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>5.1 Living in rural area</td>
<td>1</td>
<td>654</td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>0.2 [-0.07, 0.47]</td>
</tr>
<tr>
<td>5.2 Living in urban area</td>
<td>1</td>
<td>542</td>
<td>Mean Difference (SDs) (Random, 95% CI)</td>
<td>-0.03 [-0.30, 0.24]</td>
</tr>
<tr>
<td>6 Level of dietary diversity (Household Dietary Diversity Score) in previous week by gender</td>
<td>1</td>
<td></td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>6.1 Women</td>
<td>1</td>
<td>686</td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>0.6 [0.07, 1.13]</td>
</tr>
<tr>
<td>6.2 Men</td>
<td>1</td>
<td>686</td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>0.14 [-0.37, 0.65]</td>
</tr>
<tr>
<td>7 Level of dietary diversity (Household Dietary Diversity Score)</td>
<td>1</td>
<td></td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>Outcome or subgroup title</td>
<td>No. of studies</td>
<td>No. of participants</td>
<td>Statistical method</td>
<td>Effect size</td>
</tr>
<tr>
<td>-------------------------------------------------------------------------------------------</td>
<td>----------------</td>
<td>---------------------</td>
<td>---------------------------------------------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>Ty Score) in previous week by income poverty status</td>
<td></td>
<td></td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>1.04 [1.04, 1.04]</td>
</tr>
<tr>
<td>7.1 Living in income poverty</td>
<td>1</td>
<td>1774</td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>0.56 [0.54, 0.58]</td>
</tr>
<tr>
<td>8 Level of depression (Center for Epidemiologic Studies Depression Score) by rural-urban residency</td>
<td>1</td>
<td></td>
<td>Mean Difference (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>8.1 Living in rural areas</td>
<td>1</td>
<td>654</td>
<td>Mean Difference (Random, 95% CI)</td>
<td>0.26 [-2.01, 2.53]</td>
</tr>
<tr>
<td>8.2 Living in urban areas</td>
<td>1</td>
<td>542</td>
<td>Mean Difference (Random, 95% CI)</td>
<td>1.16 [1.00, 3.32]</td>
</tr>
<tr>
<td>9 Level of depression (Center for Epidemiologic Studies Depression Score) by gender</td>
<td>1</td>
<td></td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>9.1 Women</td>
<td>1</td>
<td>1070</td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>-2.44 [-4.20, -0.68]</td>
</tr>
<tr>
<td>9.2 Men</td>
<td>1</td>
<td>1070</td>
<td>Difference-in-Differences (Random, 95% CI)</td>
<td>-1.15 [-2.72, 0.42]</td>
</tr>
<tr>
<td>10 Level of depression (Center for Epidemiologic Studies Depression Score) by income poverty status</td>
<td>1</td>
<td></td>
<td>Mean Difference (Random, 95% CI)</td>
<td>Subtotals only</td>
</tr>
<tr>
<td>10.1 Living in income poverty</td>
<td>1</td>
<td>458</td>
<td>Mean Difference (Random, 95% CI)</td>
<td>-0.21 [-0.52, 0.10]</td>
</tr>
<tr>
<td>10.2 Not living in income poverty</td>
<td>1</td>
<td>457</td>
<td>Mean Difference (Random, 95% CI)</td>
<td>0.0 [-0.18, 0.18]</td>
</tr>
</tbody>
</table>

**Analysis 4.1. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 1 Height for age (standard deviations) by rural-urban residency, currently.**

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SDs)</th>
<th>Mean Difference (SDs)</th>
<th>Weight</th>
<th>Mean Difference (SDs)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td></td>
<td>IV, Random, 95% CI</td>
<td></td>
<td>IV, Random, 95% CI</td>
</tr>
<tr>
<td>4.1.1 Living in rural areas</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Fernald 2011</td>
<td>222</td>
<td>432</td>
<td>-0.1 (0.14)</td>
<td>-0.09 [-0.36, 0.18]</td>
<td>100%</td>
<td>-0.09 [-0.36, 0.18]</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td></td>
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<tr>
<td>Heterogeneity: Not applicable</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Test for overall effect: Z=0.64(P=0.52)</td>
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<tr>
<td>4.1.2 Living in urban areas</td>
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<td></td>
</tr>
<tr>
<td>Fernald 2011</td>
<td>177</td>
<td>365</td>
<td>0.1 (0.12)</td>
<td>0.13 [-0.11, 0.37]</td>
<td>100%</td>
<td>0.13 [-0.11, 0.37]</td>
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</table>
### Analysis 4.2. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 2 Height for age (standard deviations) by income poverty status, currently.

<table>
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<th>UCTs</th>
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<th>Mean Difference (SDs)</th>
<th>Weight</th>
<th>Mean Difference (SDs)</th>
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<tbody>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
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</tr>
<tr>
<td>Heterogeneity: Not applicable</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Test for overall effect: Z=1.08 (P=0.28)</td>
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<tr>
<td>Test for subgroup differences: Chi²=1.42, df=1 (P=0.23), I²=29.75%</td>
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</table>

<table>
<thead>
<tr>
<th>Favours no UCT</th>
<th>-5</th>
<th>-2.5</th>
<th>0</th>
<th>2.5</th>
<th>5</th>
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<tbody>
<tr>
<td>Favours UCT</td>
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</tbody>
</table>

### Analysis 4.3. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 3 Has had any illness in previous 2 weeks to 3 months.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
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<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
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</thead>
<tbody>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
<td></td>
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<tr>
<td>Heterogeneity: Not applicable</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Test for overall effect: Z=2.83 (P=0)</td>
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</table>

<table>
<thead>
<tr>
<th>Favours UCT</th>
<th>0.2</th>
<th>0.5</th>
<th>1</th>
<th>2</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Favours no UCT</td>
<td></td>
<td></td>
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</tbody>
</table>
Analysis 4.4. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 4 Food security index by gender.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Favours no UCT</th>
<th>No UCT</th>
<th>Difference-in-Differences (SDs) (SE)</th>
<th>Weight</th>
<th>Difference-in-Differences (SDs) (SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
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<td>IV, Random, 95% CI</td>
</tr>
<tr>
<td>4.4.1 Women</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haushofer 2013</td>
<td>343</td>
<td>343</td>
<td>0.3 (0.9)</td>
<td>100%</td>
<td>0.27[-1.49,2.03]</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>100%</td>
<td></td>
<td>0.27[-1.49,2.03]</td>
</tr>
</tbody>
</table>

Heterogeneity: Not applicable
Test for overall effect: Z=0.30 (P=0.76)

4.4.2 Men

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Favours no UCT</th>
<th>No UCT</th>
<th>Difference-in-Differences (SDs) (SE)</th>
<th>Weight</th>
<th>Difference-in-Differences (SDs) (SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
<td></td>
<td>IV, Random, 95% CI</td>
</tr>
<tr>
<td>Haushofer 2013</td>
<td>343</td>
<td>343</td>
<td>0.2 (0.9)</td>
<td>100%</td>
<td>0.23[-1.53,1.99]</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>100%</td>
<td></td>
<td>0.23[-1.53,1.99]</td>
</tr>
</tbody>
</table>

Heterogeneity: Not applicable
Test for overall effect: Z=0.26 (P=0.8)
Test for subgroup differences: Chi²=0, df=1 (P=0.97), I²=0%

4.5.1 Living in rural area

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SDs) (SE)</th>
<th>Weight</th>
<th>Mean Difference (SDs) (SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
<td></td>
<td>IV, Random, 95% CI</td>
</tr>
<tr>
<td>Fernald 2011</td>
<td>222</td>
<td>432</td>
<td>0.2 (0.14)</td>
<td>100%</td>
<td>0.20[-0.07,0.47]</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>100%</td>
<td></td>
<td>0.20[-0.07,0.47]</td>
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</tbody>
</table>

Heterogeneity: Not applicable
Test for overall effect: Z=1.43 (P=0.15)

4.5.2 Living in urban area

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SDs) (SE)</th>
<th>Weight</th>
<th>Mean Difference (SDs) (SE)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
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<td>IV, Random, 95% CI</td>
</tr>
<tr>
<td>Fernald 2011</td>
<td>177</td>
<td>365</td>
<td>-0.1 (0.14)</td>
<td>100%</td>
<td>-0.03[-0.3,0.24]</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>100%</td>
<td></td>
<td>-0.03[-0.3,0.24]</td>
</tr>
</tbody>
</table>

Heterogeneity: Tau²=0; Chi²=0, df=0 (P<0.0001); I²=100%
Test for overall effect: Z=0.21 (P=0.83)
Test for subgroup differences: Chi²=1.35, df=1 (P=0.25), I²=25.9%
### Analysis 4.6. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 6 Level of dietary diversity (Household Dietary Diversity Score) in previous week by gender.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Difference-in-Differences</th>
<th>Difference-in-Differences</th>
<th>Weight</th>
<th>Difference-in-Differences</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>(SE)</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
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</tr>
<tr>
<td><strong>4.6.1 Women</strong></td>
<td></td>
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</tr>
<tr>
<td>Haushofer 2013</td>
<td>343</td>
<td>343</td>
<td>0.6 (0.27)</td>
<td></td>
<td>100%</td>
<td>0.6 [0.07, 1.13]</td>
</tr>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
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<td></td>
<td></td>
<td>100%</td>
<td>0.6 [0.07, 1.13]</td>
</tr>
<tr>
<td>Heterogeneity: Not applicable</td>
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<tr>
<td>Test for overall effect: Z = 2.22 (P = 0.03)</td>
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<tr>
<td><strong>4.6.2 Men</strong></td>
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</tr>
<tr>
<td>Haushofer 2013</td>
<td>343</td>
<td>343</td>
<td>0.1 (0.26)</td>
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<td>100%</td>
<td>0.14 [0.37, 0.65]</td>
</tr>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
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<td></td>
<td></td>
<td>100%</td>
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<td>Heterogeneity: Not applicable</td>
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</tr>
<tr>
<td>Test for overall effect: Z = 0.54 (P = 0.59)</td>
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</tr>
<tr>
<td>Test for subgroup differences: Chi^2 = 1.51, df = 1 (P = 0.22), I^2 = 33.6%</td>
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### Analysis 4.7. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 7 Level of dietary diversity (Household Dietary Diversity Score) in previous week by income poverty status.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Difference-in-Differences</th>
<th>Difference-in-Differences</th>
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<td>N</td>
<td>N</td>
<td>(SE)</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
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<tr>
<td><strong>4.7.1 Living in income poverty</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Ward 2010</td>
<td>887</td>
<td>887</td>
<td>1 (0.001)</td>
<td></td>
<td>100%</td>
<td>1.04 [1.04, 1.04]</td>
</tr>
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<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
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<td>1.04 [1.04, 1.04]</td>
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<tr>
<td>Heterogeneity: Not applicable</td>
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<tr>
<td>Test for overall effect: Z = 1040 (P &lt; 0.0001)</td>
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<tr>
<td><strong>4.7.2 Not living in income poverty</strong></td>
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</tr>
<tr>
<td>Ward 2010</td>
<td>887</td>
<td>887</td>
<td>0.6 (0.01)</td>
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<td>100%</td>
<td>0.56 [0.54, 0.58]</td>
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<td><strong>Subtotal (95% CI)</strong></td>
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<td>Heterogeneity: Not applicable</td>
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<tr>
<td>Test for overall effect: Z = 56 (P &lt; 0.0001)</td>
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</tr>
<tr>
<td>Test for subgroup differences: Chi^2 = 2281.19, df = 1 (P &lt; 0.0001), I^2 = 99.96%</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Analysis 4.8. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 8 Level of depression (Center for Epidemiologic Studies Depression Score) by rural-urban residency.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>(SE)</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
<td></td>
</tr>
<tr>
<td><strong>4.8.1 Living in rural areas</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fernald 2011</td>
<td>222</td>
<td>432</td>
<td>0.3 [1.16]</td>
<td></td>
<td>100%</td>
<td>0.26 [-2.01, 2.53]</td>
</tr>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>100%</td>
<td>0.26 [-2.01, 2.53]</td>
</tr>
</tbody>
</table>

Favours no UCT -5 -2.5 0 2.5 5 Favours UCT

Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)
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### Analysis 4.8.2 Living in urban areas

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SE)</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fernald 2011</td>
<td>177</td>
<td>365</td>
<td>1.2 (1.1)</td>
<td></td>
<td>100%</td>
<td>1.16[-1,3.32]</td>
</tr>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
<td></td>
<td>IV, Random, 95% CI</td>
<td></td>
<td>100%</td>
<td>1.16[-1,3.32]</td>
</tr>
</tbody>
</table>

Heterogeneity: Not applicable
Test for overall effect: Z=1.05 (P=0.29)
Test for subgroup differences: Chi²=0.32, df=1 (P=0.57), I²=0%

### Analysis 4.9. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 9 Level of depression (Center for Epidemiologic Studies Depression Score) by gender.

#### 4.9.1 Women

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Difference-in-Differences (SE)</th>
<th>Difference-in-Differences</th>
<th>Weight</th>
<th>Difference-in-Differences</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haushofer 2013</td>
<td>535</td>
<td>535</td>
<td>-2.4 (0.9)</td>
<td></td>
<td>100%</td>
<td>-2.44[-4.2,-0.68]</td>
</tr>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
<td></td>
<td>IV, Random, 95% CI</td>
<td></td>
<td>100%</td>
<td>-2.44[-4.2,-0.68]</td>
</tr>
</tbody>
</table>

Heterogeneity: Not applicable
Test for overall effect: Z=2.71 (P=0.01)
Test for subgroup differences: Chi²=1.15, df=1 (P=0.28), I²=12.87%

#### 4.9.2 Men

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Difference-in-Differences (SE)</th>
<th>Difference-in-Differences</th>
<th>Weight</th>
<th>Difference-in-Differences</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haushofer 2013</td>
<td>535</td>
<td>535</td>
<td>-1.1 (0.8)</td>
<td></td>
<td>100%</td>
<td>-1.15[-2.72,0.42]</td>
</tr>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
<td></td>
<td>IV, Random, 95% CI</td>
<td></td>
<td>100%</td>
<td>-1.15[-2.72,0.42]</td>
</tr>
</tbody>
</table>

Heterogeneity: Not applicable
Test for overall effect: Z=1.44 (P=0.15)
Test for subgroup differences: Chi²=1.15, df=1 (P=0.28), I²=12.87%

### Analysis 4.10. Comparison 4 Unconditional cash transfer versus no unconditional cash transfer: health equity, Outcome 10 Level of depression (Center for Epidemiologic Studies Depression Score) by income poverty status.

#### 4.10.1 Living in income poverty

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SE)</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paxson 2007</td>
<td>229</td>
<td>229</td>
<td>-0.2 (0.16)</td>
<td></td>
<td>100%</td>
<td>-0.21[-0.52,0.1]</td>
</tr>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
<td></td>
<td>IV, Random, 95% CI</td>
<td></td>
<td>100%</td>
<td>-0.21[-0.52,0.1]</td>
</tr>
</tbody>
</table>

Heterogeneity: Not applicable
Test for overall effect: Z=1.31 (P=0.19)

#### 4.10.2 Not living in income poverty

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SE)</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>N</td>
<td>IV, Random, 95% CI</td>
<td>IV, Random, 95% CI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paxson 2007</td>
<td>229</td>
<td>228</td>
<td>0 (0.09)</td>
<td></td>
<td>100%</td>
<td>0[-0.18,0.18]</td>
</tr>
<tr>
<td><strong>Subtotal (95% CI)</strong></td>
<td></td>
<td></td>
<td>IV, Random, 95% CI</td>
<td></td>
<td>100%</td>
<td>0[-0.18,0.18]</td>
</tr>
</tbody>
</table>

Heterogeneity: Not applicable
Test for overall effect: Z=1.31 (P=0.19)
Test for subgroup differences: Chi²=1.15, df=1 (P=0.28), I²=12.87%
## APPENDICES

**Appendix 1: Appendix 1: Search strategy for Ovid MEDLINE(R) 1946 to Present with Daily Update**

### Intervention terms
1. maternal welfare/
2. public policy/
3. social welfare/
4. exp social security/
5. (social adj (assistance or polic$ or welfare or insurance$ or protection)).ti,ab.
6. public assistance.ti,ab.
7. family policy.mp.
8. ((financial or cash or pay$ or monetary or money) adj3 (transfer$ or measure$ or incentive$ or allowance$ or exclu$ or reform$ or gain $ or credit$1 or benefit$1)).ti,ab.
9. or/1-8

### Study terms
10. randomized controlled trial/
11. random$.ti,ab.
12. random allocation/
13. placebos/
14. placebo$.ti,ab.
15. single-blind method/
16. double-blind method/
17. ((single or double or triple or treble) adj blind$).ti,ab.
18. control groups/
19. exp clinical trial/
20. comparative Study/
21. intervention studies/
22. exp cohort studies/
23. evaluation studies/

### Table

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>UCT</th>
<th>No UCT</th>
<th>Mean Difference (SE)</th>
<th>Weight</th>
<th>Mean Difference</th>
<th>UCT</th>
<th>No UCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heterogeneity: Not applicable</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test for overall effect: Not applicable</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test for subgroup differences: ( \chi^2=1.31, \text{df}=1 (P=0.25), I^2=23.58% )</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Favours no UCT | -5 | -2.5 | 0 | 2.5 | 5 |
Favours UCT |
Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

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Appendix 2. Search strategies for electronic academic databases

Cochrane Central Register of Controlled Trial (CENTRAL)

29 May 2015 (This registry has not been updated since 2014, so the original search in 2015 was not required to be re-run.)

107 records

Intervention terms

TX (social N1 (assistance OR policy OR welfare OR insurance OR protection)) OR "public assistance" OR "family policy" OR ((financial OR cash OR pay OR monetary OR money) N3 (transfer OR measure OR incentive OR allowance OR exclu OR reform OR gain OR credit OR credits OR benefit OR benefits)))

Country terms

TX (Africa OR Asia OR "South America" OR "Central America" OR ((developing OR "less" developed) OR "third world" OR "under developed" OR "middle income" OR "low income" OR underserved OR "under served" OR deprived OR poor) N1 (count OR nation OR nations OR state OR states OR population OR populations)) IMic OR Imics

OR

Tl (Samoa* OR Argentin* OR Beliz* OR Botswana* OR Brazil* OR Bulgaria* OR Chile* OR Comoro* OR "Costa Rica"* OR Croatia* OR Dominica* OR Guinea* OR Gabon* OR Grenada* OR Hungary* OR Kazakh* OR Latvia* OR Lebanon* OR Libya* OR Lithuania* OR Malay* OR Maurit* OR Mexic* OR Micronesia* OR Monteneg* OR Oman* OR Palau* OR Panama* OR Poland* OR Polish* OR Romania* OR Russia* OR Seychelles* OR Slovak* OR "Saint Kitts" OR Nevis* OR "Saint Lucia"* OR "Saint Vincent" OR Grenadines OR Turk* OR Uganda* OR Venezuela* OR Yugoslavia* OR Liberia* OR Mayotte OR "Mariana Island"* OR Serbia* OR "St Kitts" OR "St Lucia"* OR "St Vincent"* OR Albania* OR Algeria* OR Angol* OR Armenia* OR Azerbaijan* OR Belarus* OR Bhutan* OR Bolivia* OR Bosnia* OR Herzegov* OR Cameroon* OR China OR Chinese OR Colombia* OR Congo* OR Cuba* OR Djibouti* OR Ecuador* OR Egypt* OR "El Salvador"* OR Fiji* OR Georgia* OR Guam* OR Guatemala* OR Guyana* OR Honduras* OR "Indian Ocean Island"* OR Indonesia* OR Iran* OR Iraq* OR Jamaica* OR Jordan* OR Lesotho OR Macedonia* OR "Marshall Island"* OR Micronesia* OR "Middle East"* OR Moldova* OR Morocco* OR Namibia* OR Nicaragua* OR Paraguay* OR Peru* OR Philippin* OR "Sri Lanka"* OR Suriname* OR Swaziland* OR Syria* OR Thai* OR Tonga* OR Tunisia* OR Turkmen* OR Ukrain* OR Vanuatu* OR "Cape Verde"* OR Gaza OR Kiribati* OR Maldives OR Palestin* OR "West Bank" OR Afghan* OR Banglades* OR Benin* OR "Burkina Faso"* OR Burundi* OR Cambodia* OR Central African Republic* OR Chad* OR Comoros OR Congo* OR "Cote d Ivoire" OR Eritrea* OR Ethiopia* OR Gambia* OR Ghana* OR "Guinea-Islaw" OR Haiti* OR India* OR Kenya* OR Korea* OR Kyrgyz* OR Laos OR Laos OR Liberia* OR Madagascar OR Malagasy OR Malawi* OR Mali* OR Mauritania* OR Melanesia* OR Mongolia* OR Mozambi* OR Myanmar OR Nepal* OR Niger* OR Nigeria* OR Pakistan* OR "Papua New Guinea"* OR Rwand* OR Senegal* OR "Sierra Leone"* OR Somalia* OR Sudan* OR Tajikistan* OR Tanzania* OR "East Timor"* OR Togo* OR Uganda* OR Uzbek* OR Vietnam* OR Yemen* OR Zambia* OR Zimbabwe* OR Burm* OR Lao OR "North Korea"* OR "Solomon Island"* OR "Sao Tome" OR Timor* OR "Viet Nam"* OR "ivory coast")

OR

AB (Samoa* OR Argentin* OR Beliz* OR Botswana* OR Brazil* OR Bulgaria* OR Chile* OR Comoro* OR "Costa Rica"* OR Croatia* OR Dominica* OR Guinea* OR Gabon* OR Grenada* OR Hungary* OR Kazakh* OR Latvia* OR Lebanon* OR Libya* OR Lithuania* OR Malay* OR Maurit* OR Mexic* OR Micronesia* OR Monteneg* OR Oman* OR Palau* OR Panama* OR Poland* OR Polish* OR Romania* OR Russia* OR Seychelles* OR Slovak* OR "Saint Kitts" OR Nevis* OR "Saint Lucia"* OR "Saint Vincent" OR Grenadines OR Turk* OR Uganda* OR Venezuela* OR Yugoslav* OR Liberia* OR Mayotte OR "Mariana Island"* OR Serbia* OR "St Kitts" OR "St Lucia"* OR "St Vincent"* OR Albania* OR Alge* OR Angol* OR Armenia* OR Azerbaijan* OR Belarus* OR Bhutan* OR Bolivia* OR Bosnia* OR Herzegov* OR Cameroon* OR China OR Chinese OR Colombia* OR Congo* OR Cuba* OR Djibouti* OR Ecuador* OR Egypt* OR "El Salvador"* OR Fiji* OR Georgia* OR Guam* OR Guatemala* OR Guyana* OR Hondur* OR "Indian Ocean Island"* OR Indonesia* OR Iran* OR Iraq* OR Jamaica* OR Jordan* OR Lesotho OR Macedonia* OR "Marshall Island"* OR Micronesia* OR "Middle East"* OR Moldova* OR Morocco* OR Namibia* OR Nicaragua* OR Paraguay* OR Peru* OR Philippin* OR "Sri Lanka"* OR Suriname* OR Swaziland* OR Syria* OR Thai* OR Tonga* OR Tunisia* OR Turkmen* OR Ukrain* OR Vanuatu* OR "Cape Verde"* OR Gaza OR Kiribati* OR Maldives OR Palestin* OR "West Bank" OR Afghan* OR Banglades* OR Benin* OR "Burkina Faso"* OR Burundi* OR Cambodia* OR Central African Republic* OR Chad* OR Comoros OR Congo* OR "Cote d Ivoire" OR Eritrea* OR Ethiopia* OR Gambia* OR Ghana* OR "Guinea-Islaw" OR Haiti* OR India* OR Kenya* OR Korea* OR Kyrgyz* OR Laos OR Laos OR Liberia* OR Madagascar OR Malagasy OR Malawi* OR Mali* OR Mauritania* OR Melanesia* OR Mongolia* OR Mozambi* OR Myanmar OR Nepal* OR Niger* OR Nigeria* OR Pakistan* OR "Papua New Guinea"* OR Rwand* OR Senegal* OR "Sierra Leone"* OR Somalia* OR Sudan* OR Tajikistan* OR Tanzania* OR "East Timor"* OR Togo* OR Uganda* OR Uzbek* OR Vietnam* OR Yemen* OR Zambia* OR Zimbabwe* OR Burm* OR Lao OR "North Korea"* OR "Solomon Island"* OR "Sao Tome" OR Timor* OR "Viet Name" OR "ivory coast")
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Somalia* OR Sudan* OR Tajikistan* OR Tanzania* OR "East Timor"* OR Togo* OR Uganda* OR Uzbek* OR Vietnam* OR Yemen* OR Zambia* OR Zimbabwe* OR Burm* OR Lao.tlw. OR "North Korea"* OR "Solomon Island"* OR "Sao Tome"* OR Timor* OR "Viet Nam"* OR "ivory coast")

OR

SU (Samoa* OR Argentin* OR Beliz* OR Botswana* OR Brazil* OR Bulgaria* OR Chile* OR Comoro* OR "Costa Rica"* OR Croatia* OR Dominica* OR Guinea* OR Gabon* OR Grenada* OR Hungary* OR Kazakh* OR Latvia* OR Lebanon* OR Libya* OR Lithuania* OR Malaysia* OR Mauritius* OR Mexico* OR Micronesia* OR Montenegro* OR Oman* OR Palau* OR Panama* OR Poland OR Polish OR Romania* OR Russia* OR Seychelles* OR Slovak* OR "Saint Kitts" OR Nevis OR "Saint Lucia"* OR "Saint Vincent" OR Grenadines OR Turk* OR Uragua* OR Venezuela* OR Yugoslavia* OR Libya* OR Mayotte OR Mariana Island* OR Serbia* OR "St Kitts" OR "St Lucia"* OR "St Vincent" OR Albania* OR Algeria* OR Angol* OR Armenia* OR Azerbaijan* OR Belarus* OR Bhutan* OR Bolivia* OR Bosnia* OR Herzegovin* OR Cameroon* OR China OR Chinese OR Colombia* OR Congo* OR Cuba* OR Djibouti* OR Ecuador* OR Egypt* OR "El Salvador"* OR Fiji* OR Georgia* OR Guam* OR Guatemala* OR Guyana* OR Honduras* OR "Indian Ocean Island"* OR Indonesia* OR Iran* OR Iraq* OR Jamaica* OR Jordan* OR Lesotho OR Macedonia* OR "Marshall Island"* OR Micronesia* OR "Middle East"* OR Moldova* OR Morocco* OR Namibia* OR Nicaragua* OR Paraguay* OR Peru* OR Philippin* OR "Sri Lanka"* OR Suriname* OR Swaziland* OR Syria* OR Thai* OR Tonga* OR Tunisia* OR Turkmen* OR Ukrain* OR Vanuatu* OR "Cape Verd"* OR Gaza OR Kiribati* OR Maldives OR Palestin* OR "West Bank" OR Afghan* OR Bangladesh* OR Benin* OR "Burkina Faso"* OR Burundi* OR Cambodia* OR "Central African Republic"* OR Chad* OR Comoros OR Congo* OR "Cote d'Ivoire"* OR Eritrea* OR Ethiopia* OR Gambia* OR Ghana* OR Guinea* OR "Guinea-Bissau" OR Haiti* OR India* OR Kenya* OR Korea* OR Kyrgyz* OR Laos OR Laot* OR Liberia* OR Madagascar OR Malagasy OR Malawi* OR Mali* OR Mauritania* OR Melaniesia* OR Mongolia* OR Mozamb* OR Myanmar OR Nepali* OR Niger* OR Nigeria* OR Pakistan* OR "Papua New Guinea"* OR Rwanda* OR Senegal* OR "Sierra Leone"* OR Somalia* OR Sudan* OR Tajikistan* OR Tanzania* OR "East Timor"* OR Togo* OR Uganda* OR Uzbek* OR Vietnam* OR Yemen* OR Zambia* OR Zimbabwe* OR Burm* OR Lao.tlw. OR "North Korea" OR "Solomon Island"* OR "Sao Tome"* OR Timor* OR "Viet Nam"* OR "ivory coast")

Ovid MEDLINE(R) 1946 to Present with Daily Update, plus Daily Update to 5 May 2017

5 May 2017

6281 records

See Appendix 1 for search strategy.

Embase

10 May 2017

9023 records

Intervention terms

'maternal welfare'/de OR 'policy'/de OR 'social welfare'/de OR 'social security'/exp OR (social NEAR/1 (assistance OR polic* OR welfare OR insurance* OR protection));ti,ab OR 'public assistance':ti,ab OR 'family policy':ti,ab OR ((financial OR cash OR pay* OR monetary OR money) NEAR/3 (transfer* OR measure* OR incentive* OR allowance* OR exclu* OR reform* OR gain* OR credit OR credits OR benefit OR benefit));ti,ab

Study terms

'clinical trial'/exp OR 'placebo'/de OR 'single blind procedure'/de OR 'double blind procedure'/de OR 'control group'/de OR 'comparative study'/de OR 'intervention study'/de OR 'cohort analysis'/exp OR 'evaluation study'/exp OR random*:ti,ab OR 'random allocation':ti,ab OR 'placebo':ti,ab OR ((single OR double OR triple OR treble) NEAR/1 blind):ti,ab OR (time NEAR/1 series):ti,ab OR (quasi NEXT/1 experiment):ti,ab OR 'controlled before':ti,ab OR 'controlled after':ti,ab OR 'controlled panel':ti,ab OR ((panel OR intervention* OR evaluate* OR compar*) NEXT/1 stud):ti,ab OR 'before and after':ti,ab OR 'repeat* NEXT/1 measure':ti,ab OR 'follow up* NEXT/1 assessment':ti,ab OR (group OR groups):ti,ab OR ((intervention OR observational OR process OR program) NEAR/8 (evaluate* OR effect* OR outcome)):ti,ab OR (program OR programmti,ab OR (secondary NEXT/1 analysis)):ti,ab OR 'intervention* OR treatment* AND (control* OR study OR program* OR comparison OR comparative));ti,ab

Country terms

'developing country'/exp OR 'Africa'/exp OR 'Asia'/exp OR 'South and Central America'/exp OR (Africa OR Asia OR 'South America' OR 'Latin America' OR 'Central America'):ti,ab OR ((developing OR 'less developed' OR 'third world' OR 'under developed' OR 'middle income' OR 'low income' OR underserved OR 'under served' OR deprived OR poor* NEXT/1 (count* OR nation* OR state* OR population)):ti,ab OR (Imic OR Imics):ti,ab

OR

(Samoa* OR Argentin* OR Beliz* OR Botswana* OR Brazil* OR Bulgaria* OR Chile* OR Comoro* OR (Costa NEXT/1 Rica*)) OR Croatia* OR Dominica* OR Guinea* OR Gabon* OR Grenada* OR Hungary* OR Kazakh* OR Latvia* OR Lebanon* OR Libya* OR Lithuania* OR Malaysia* OR Mauritius* OR Mexico* OR Micronesia* OR Montenegro* OR Oman* OR Palau* OR Panama* OR Poland OR Polish OR Romania* OR Russia* OR...
Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

Academic Search Premier
5 May 2017
3687 records
Unconditional cash transfers for reducing poverty and vulnerabilities: effect on use of health services and health outcomes in low- and middle-income countries (Review)

124

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Business Source Complete

11 May 2017

2420 records
**Study terms**

TI (random OR placebo OR ((single or double or triple or treble) N1 blind)) OR (time N1 series) OR "quasi-experiment" OR "pre test" OR pretest OR "pre-intervention" OR "post test" OR posttest OR "post-intervention" OR "controlled before" OR "independent panel" OR "panel study" OR "intervention" study** OR "before and after" OR "repeat" measure OR "evaluate" study OR compar* study** OR trial OR "follow up assessment" OR groups OR (intervention OR interventional OR process OR program) N8 (evaluate OR effect OR outcome)**

OR program OR programme OR "secondary analyses" OR ((evaluate OR intervention OR treatment)** and (control* OR study OR program OR comparison OR comparative))) OR AB (random OR placebo OR ((single or double or triple or treble) N1 blind)) OR (time N1 series) OR "quasi-experiment" OR "pre test" OR pretest OR "pre-intervention" OR "post test" OR posttest OR "post-intervention" OR "controlled before" OR "independent panel" OR "panel study" OR "intervention" study** OR "before and after" OR "repeat" measure OR "evaluate" study OR compar* study** OR trial OR "follow up assessment" OR groups OR (intervention OR interventional OR process OR program) N8 (evaluate OR effect OR outcome)** OR program OR programme OR "secondary analyses" OR ((evaluate OR intervention OR treatment)** and (control* OR study OR program OR comparison OR comparative)))

**Countries terms**

DE ("Africa" OR "Asia" OR "South America" OR "Central America" OR "Latin America") OR TI (Africa OR Asia OR "South America" OR "Latin America" OR "Central America" OR ((developing OR "less" developed OR "third world" OR "under developed" OR "middle income" OR "low income" OR underserved OR "under served" OR deprived OR poor)) N1 (count OR nation OR nations OR state OR states OR population OR populations)) OR Imic OR Imics) OR AB (Africa OR Asia OR "South America" OR "Latin America" OR "Central America" OR ((developing OR "less" developed OR "third world" OR "under developed" OR "middle income" OR "low income" OR underserved OR "under served" OR deprived OR poor)) N1 (count OR nation OR nations OR state OR states OR population OR populations)) OR Imic OR Imics)

OR

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**Study terms**

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CONTRIBUTIONS OF AUTHORS

Pega conceived of the review. Pega led and all authors contributed to the protocol development. Bain and Pega searched the electronic and grey literature databases. Pega led and Liu, Pabayo and Walter contributed to searches of key organisational websites. Walter and Pega led and all authors contributed to screening of records identified in the searches. Pega led and Henning, Paek and all authors contributed to the data extraction. Pega led and all authors contributed to the quality assessment of included studies, analysis and interpretation of data, and writing of the review.
DEclarations of interest

Pega: none known. Frank Pega is a technical officer for the World Health Organization but was a postdoctoral fellow for the University of Otago at the time of writing.

Liu: none known.

Walter: none known.

Pabayo: none known.

Saith: none known. Oxford Policy Management has been involved in the implementation and evaluation of a number of cash transfer schemes in low- and middle-income countries.

Lhachimi: none known.

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Internal sources

• Harvard Medical School, USA.
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• Oxford Policy Management, Asia (New Delhi Office), India.
  Oxford Policy Management, Asia (New Delhi Office) provided salary funding to Saith.

• University of Bremen & Leibniz Institute for Prevention Research and Epidemiology, Bremen, Germany.
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• University of Nevada, Reno, USA.
  The University of Nevada, Reno provided salary funding to Pabayo.

• University of Otago, New Zealand.
  The University of Otago provided salary funding through a Health Sciences Career Development Programme Postdoctoral Fellowship to Pega.

External sources

• Canadian Institutes of Health Research, Canada.
  The Canadian Institutes of Health Research provided a CIHR Postdoctoral Fellowship to Pabayo.

Differences between protocol and review

There are the following differences between the protocol and the review:

• Background: updated to reflect the most recent state of evidence.

• Types of studies: added that if a study compared a UCT with both no UCT and with a smaller amount of UCT, then we prioritised comparisons with the group who received no UCT over those receiving a smaller amount of the UCT. The comparison with no intervention is more consistent with the objectives of the review of evaluating intervention effectiveness, because receipt of any UCT may be more important for health effects than the amount of a UCT received (Baird 2011; Filmer 2011).

• Types of interventions: refined the definition of UCTs by excluding vouchers. Unlike cash transfers, transfers via vouchers restrict their recipients’ ability to spend the additional income, for example, by requiring recipients to only purchase certain goods and services from certain suppliers. Therefore, voucher transfers may impact health differently from genuine cash transfers, and may potentially act through different pathways.

• Types of interventions: refined the definition of UCTs by including payments via mobile phone, because these electronic payments may have another health effect than in-hand cash payments.

• Types of interventions: changed the inclusion/exclusion criteria for UCTs with co-intervention. In the review, we excluded UCTs with major co-interventions and included UCTs with minor co-interventions (defined as interventions that we anticipated to very likely be of relatively low or no impact, such as a minor educational co-intervention or very small once-off payment). We now believe that minor co-interventions, which are commonly provided alongside or in combination with UCTs, do not constitute a threat to causal inference.
• Types of interventions: changed the inclusion/exclusion of fuzzy UCTs. In the protocol, we included fuzzy UCTs if their intention was to be unconditional, and excluded (but noted) UCTs with any de facto conditions. In the review, we included fuzzy UCTs that were in practice unconditional, regardless of intention, and we excluded fuzzy UCTs with de facto conditions (e.g. major administrative linking of the cash transfer or major messaging around the cash transfer). We now believe that what matters for effects on use of health services and health outcomes is likely more so the actual, experienced conditionality of the cash transfer, rather than the cash transfer’s design as such.

• Types of outcomes measures: added criteria around selection of time points to be reported to ensure a systematic and consistent approach.

• Types of outcomes measures: refined morbidity outcomes included in the review to more specifically identify the most important health outcomes.

• Types of outcomes measures: refined the criteria for prioritising the types of outcomes measures to ensure prioritisation of the most relevant measures.

• Types of participants: In our protocol, we put the division between a child and an adult at 14 years. In the review, we put this division at 17 years. We now believe that children are more commonly defined as people under the age of 18 years, and that adults are more commonly defined as people aged 18 years or older.

• Search: added handsearches of previous reviews in the field as a search source.

• Search: did not search the Global Health, Web of Science database as planned.


• Search: in the protocol, we did not specify the number of hits from searches of grey literature databases we would screen for eligible records. In the review, we only screened the first 100 hits in grey literature database searches that exceeded 500 hits, after ordering hits for relevance, if possible. The reason was that some grey literature database searches returned very large numbers of hits, and it was not feasible to screen all of these hits.

• Search: also searched the websites of two additional organisations (i.e. the Cash Transfer Projects in Humanitarian Aid and Save the Children).

• Assessment of risk of bias in included studies: if the review had included interrupted time series studies, to assess risk of bias in interrupted time series studies, we would have used the Cochrane Effective Practice and Organisation of Care’s ‘Risk of bias’ criteria (EPOC 2012) plus an item assessing the risk of bias from confounding. Had the review included cohort studies, in the absence of credible standard tools for assessing risk of bias, we would have at a minimum assessed the risk of bias from sampling; low response rates; attrition; exposure measurement; outcome measurement; confounding; and reverse causation (as in our previous reviews: Pega 2013; Pega 2015a).

• Assessment of risk of bias: in the protocol, we planned to require all authors to agree on any discrepancy in risk of bias assessment. In this review, we resolved disputes between two authors through a third author.

• Measures of treatment effect: added a framework for selecting between multiple models of adjustment to ensure a systematic and consistent approach.

• Measures of treatment effect: added prioritisation of estimates of the effect of being eligible for or receiving the UCT over estimates of the effect of the specific dollar amount of the UCT that the recipient was eligible for or received. The reason was that we assume that the former treatment effect measures are more relevant than the latter for intervention effectiveness.

• Unit of analysis issues: In the protocol, we planned to request individual patient data for re-analysis for cluster-RCTs that did not adjust for clustering in their analysis and to exclude any studies for which individual data were not available. In the review, we requested adjusted data from the primary study authors. The reason is that it was not feasible to obtain individual-level data for all studies whose study records only provided effect estimates unadjusted for clustering, but we were able to obtain the cluster-adjusted effect estimates for all these studies.

• Meta-analysis: refined and added criteria for combining studies in meta-analysis for the purpose of ensuring consistent rules being applied on unanticipated issues, such as the question of whether or not to meta-analyse subgroups from the same study.

• Data synthesis: used RevMan 5.3, rather than RevMan 5.2.

• Subgroup analyses: If subgroup analyses had been feasible, we would have conducted such analyses by: age (children aged 0 to 17 years), adults (18 years or older); disaster type (natural, manmade); gender; level of income (e.g. total personal or household annual income after tax); and WHO region (Africa, Americas, Eastern Mediterranean, Europe, South-East Asia and Western Pacific).

INDEX TERMS

Medical Subject Headings (MeSH)

*Developing Countries; *Financial Support; *Financing, Government; Africa; Altruism; Americas; Asia, Southeastern; Cohort Studies; Controlled Before-After Studies; Depression [epidemiology]; Employment [statistics & numerical data]; Food Supply; Health Services...
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