SYSTEMATIC REVIEW

Health professional-delivered obesity prevention interventions during the first 1,000 days: A systematic review of external validity reporting [version 1; peer review: awaiting peer review]

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Abstract

Background: Childhood obesity prevention interventions delivered by health professionals during the first 1,000 days show some evidence of effectiveness, particularly in relation to behavioural outcomes. External validity refers to how generalisable interventions are to populations or settings beyond those in the original study. The degree to which external validity elements are reported in such studies is unclear however. This systematic review aimed to determine the extent to which childhood obesity interventions delivered by health professionals during the first 1,000 days report on elements that can be used to inform generalizability across settings and populations.

Methods: Eligible studies meeting study inclusion and exclusion criteria were identified through a systematic review of 11 databases and three trial registers. An assessment tool based on the RE-AIM (Reach, Effectiveness, Adoption, Implementation, Maintenance) framework was used to assess the external validity of included studies. It comprised five dimensions: reach and representativeness of individuals, reach and representativeness of settings, implementation and adaptation, outcomes for decision making maintenance and/or institutionalisation. Two authors independently assessed the external validity of 20% of included studies; discrepancies were resolved, and then one author completed assessments of the remaining studies.

Results: In total, 39 trials involving 46 interventions published between 1999 and 2019 were identified. The majority of studies were randomized controlled trials (n=24). Reporting varied within and between dimensions. External validity elements that were poorly described included: representativeness of individuals and settings, treatment receipt, intervention mechanisms and moderators, cost effectiveness, and intervention sustainability and acceptability.

Conclusions: Our review suggests that more emphasis is needed on research designs that consider generalisability, and the reporting of external validity elements in early life childhood obesity prevention.
interventions. Important gaps in external validity reporting were identified that could facilitate decisions around the translation and scale-up of interventions from research to practice.

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Keywords
External validity, childhood obesity, generalisability, intervention, implementation science, health professional, prevention, replication, systematic review

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Introduction
Effective, scalable, and affordable strategies that do not widen health inequities are needed to address childhood obesity. In addition, interventions that can be embedded into ongoing practice and existing systems are required, rather than implementing interventions that are resource-intensive and cannot be maintained in the long-term. This was echoed in a recent research prioritisation study in which ‘Implementation science’ and ‘How to integrate obesity prevention into existing service structures’ were the third and fourth ranked research priorities identified by researchers, policymakers and practitioners. To date, there has been limited scale-up of childhood obesity prevention interventions.

Appraising scalability prior to investment is vital. Greater consideration of the external validity of interventions (i.e. how generalisable the intervention is to populations and/or settings beyond those in the original study) is needed to inform decisions about whether interventions should be adopted elsewhere and/or scaled-up. Whether interventions can feasibility be generalised to other settings is important, but also a clear understanding of intervention implementation and causal mechanisms and how each of these might vary with context. Lack of attention to external validity can contribute to the failure to replicate findings. The poor reporting of external validity elements in childhood obesity prevention interventions also limits decision-makers’ ability around translation of interventions into practice.

As public health interventions are usually complex, and context dependent, it can be difficult to assess their generalisability to other contexts. There are many tools for assessing generalisability, however, there is no consensus on which should be used, or when. Indeed, Burchett and colleagues argue that such tools may not be the best method for generalisability assessments, instead advocating a focus on mechanisms of action through which an intervention exerts its effect – and which contextual elements underpin them, rather than solely on intervention characteristics. To improve reporting across behavioural interventions and enhance the translation of research into practice, Glasgow and colleagues developed the RE-AIM (Reach, Effectiveness, Adoption, Implementation, Maintenance) framework to evaluate the degree to which behavioural interventions report on external validity factors. Reach is the number, proportion of the intended target population, and the representativeness of participants compared with the intended target population. Effectiveness (or efficacy, depending on the study design) is the degree to which the intervention changes behaviour, quality of life, and participant satisfaction outcomes as well as physiologic endpoints, and includes attention to positive, unintended and negative results. Adoption is the number and proportion of settings and staff members that agree to initiate an intervention and how representative they are of the target setting and staff. Implementation is the degree to which settings and staff members deliver an intervention as intended, the adaptations made, and the related costs. Finally, maintenance is sustained effectiveness at the participant level and sustained (or adapted) delivery at the setting or staff level. At the individual level, it refers to the long-term results of intervention (defined as a minimum of six months following the last contact). RE-AIM is the most frequently applied framework in the translation of research evidence into policy and practice. It has been used to assess reports of external validity factors across a variety of areas, including weight loss maintenance interventions, behavioural interventions that target physical activity, mobile health physical activity promotion interventions, physical activity promotion in Latin American populations, behavioural intervention studies conducted in community settings, school health promotion studies, behaviour change interventions in healthcare settings, and housing improvement.

Based on the RE-AIM framework, Green and Glasgow proposed a set of ratings to assess external validity. These were further adapted by Laws and colleagues and have been used to assess external validity in diabetes prevention research and obesity prevention in children aged 0–5 years. This adapted tool includes five main dimensions (defined in Table 1): 1) reach and representativeness (individuals); 2) reach and representativeness (settings); 3) implementation and adaptation (of intervention), which includes fidelity considerations; 4) outcomes for decision makers; 5) maintenance and institutionalisation (i.e. the potential for implementation of the intervention in routine service delivery).

Reviews of external validity reporting in childhood obesity interventions identify insufficient reporting of elements necessary to make decisions about generalisability. A review of external validity reporting in 19 long-term follow-up childhood obesity prevention trials (children aged 0–18 years) published between 1980 and 2004 found that all studies lacked full reporting on potential generalizability and dissemination elements; the most infrequent were reports of setting level inclusion and exclusion criteria and representativeness, characteristics regarding intervention staff, implementation of intervention content, costs, and program sustainability. A more recent review of external validity reporting in 32 trials of interventions to prevent obesity or improve obesity related behaviours in children aged 0–5 years from socioeconomically disadvantaged or Indigenous families found similar issues with reporting. Health professional-delivered interventions to prevent childhood obesity during the first 1,000 days have limited impacts on adiposity/weight outcomes, but have more positive impacts on behavioural outcomes. Despite the increasing numbers of trials to assess the impact of early life obesity prevention interventions, there is relatively little reporting on the potential for these interventions to be translated into routine practice. Furthermore, there is little evidence that interventions with demonstrated efficacy have been translated beyond the research setting and been broadly adopted. Given that it can take up to 17 years to translate evidence into practice, it is important to assess the extent to which trials report on factors that can provide additional explanation for variability in intervention outcomes. Insights into successful adaptations of interventions, inform generalizability across settings and populations, and help guide policy decisions.

This study aims to determine the extent to which childhood obesity interventions delivered by health professionals during the first 1,000 days report on factors that can be used inform...
generalizability across settings and populations, and to provide recommendations for researchers planning to conduct similar studies.

Methods
We conducted a systematic review of obesity prevention interventions delivered by health professionals targeting children in the first 1,000 days of life. A separate paper reports on the effectiveness of such interventions and what behaviour change theories and/or techniques are associated with more effective intervention outcomes. The review protocol was registered with the International Prospective Register for Systematic Reviews (PROSPERO) CRD42016050793 on 3rd November 2016. This paper adheres to the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) standardised reporting guidelines; the PRISMA checklist is available on OSF.

Search strategy
Key word searches, using combinations of key words and Medical Subject Headings (or equivalent), were used across six concepts using the AND Boolean operator: (1) child; (2) mother/parent; (3) BMI/obesity; (4) nutrition/physical activity/sleep/parenting; (5) intervention/prevention; (6) randomised controlled trial (RCT)/quasi-randomised trials. Within each of the categories, keywords were combined using the “OR” Boolean operator. The search strategy was purposefully broad enough to capture any study which might have assessed weight-related measures in children under the age of two. The search strategy was initially developed in Embase (see extended data), appropriately tailored for use within the other databases, and piloted before final searches were run.

One reviewer (MH) searched the following databases from inception to 04 April 2019 using pre-specified search strategies: CINAHL Complete (EBSCOhost; 1994); Embase® (Elsevier; 1980); MEDLINE (Ovid®; 1966); PsycINFO (Ovid®; 1978); PubMed (1996); The Cochrane library databases: The Central Register of Controlled Trials; Database of Systematic Reviews; Database of Abstracts of Reviews of Effect (Wiley; 1996). Conference proceedings and other grey literature were searched on:

<table>
<thead>
<tr>
<th>Table 1. Study inclusion criteria.</th>
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<tbody>
<tr>
<td><strong>Design</strong></td>
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<td><strong>Participants</strong></td>
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<td><strong>Intervention</strong></td>
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<td><strong>Comparator</strong></td>
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<td><strong>Outcomes</strong></td>
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<td><strong>Publications</strong></td>
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</table>
Supplementary materials and trial registry protocols were also checked. No restrictions were applied to: language; date of publication; the length of follow-up of outcomes (given the diversity reported within systematic reviews to date); type of setting; mode of delivery. Records were de-duplicated in Endnote, imported into COVIDENCE and any remaining duplicates removed.

**Study inclusion and exclusion criteria**

Table 1 details the study inclusion and exclusion criteria. We included randomised controlled trials, including cluster-randomised controlled trials, or quasi-randomised trials comparing any behavioural intervention, delivered by health professionals, with ‘usual care’/active comparator which aimed to prevent overweight/obesity in children under the age of two that were born at term. Studies had to report at least one infant/child-related adiposity and/or weight outcome measure at follow-up, which could be immediately post-intervention, or at any time point thereafter; trials only reporting infant birth weight were excluded.

**Study selection**

MH and LT independently screened titles and abstracts against the inclusion criteria, and following the retrieval of full-texts, MH and LvR independently reviewed them for inclusion. Disagreements were resolved through discussion, with a third author (MB / CH / RL) where necessary.

**Data extraction**

All published papers and supplementary material related to the study (e.g. protocol papers and trial registry protocols, reference to websites with working hyperlinks, long-term follow-up studies) were used alongside the included article for data extraction. Data were extracted by one author (MH) using a pre-piloted data extraction tool (see extended data), with 20% double-checked by a second reviewer (HCW). Intervention descriptions were extracted following the criteria outlined in the TIDieR reporting guidelines. The external validity assessment tool previously developed by RL was used to assess the extent to which findings of included studies/trials could be generalised to populations or settings beyond those in the original study. Included studies were coded according to whether they met each criterion (yes, no, or not applicable). Initially, two authors (MH and RL) independently assessed the external validity of 20% of included studies. Any discrepancies were resolved through discussion, and then one author (MH) completed assessments of the remaining studies. We did not exclude any studies on the basis of the effectiveness and/or quality assessment.

**Results**

Electronic and hand searches identified 27,609 references (see Figure 1). Following duplicate removal and title and abstract screening, 230 references were selected for full text review. We identified 39 eligible studies with 46 unique intervention arms and a total of 180 eligible pairs. Five trials had more than one eligible intervention arm. Studies were mostly published from 2011 onwards (n=34), conducted in high-income countries (n=33), and targeted the period from birth to 2 years only (n=26). They focused on a range of behaviours and outcomes, including: multiple infant behaviours (n=13); infant feeding: formula feeding/breastfeeding/introduction to solids (n=10); maternal diet/physical activity/gestational weight gain (n=9); infant feeding: breastfeeding only (n=8). Only 16 of the 46 interventions were clearly delivered as part of routine care, with a further two partly delivered as such. Details of intervention descriptions and outcomes are available as extended data.

The external validity assessment of the 39 included studies is summarised in Table 2, with a summary by study available as extended data. Full details of these assessments, including supporting statements for each study, are available as extended data. The number and percentage of studies reporting all elements of each dimension of external validity are outlined in Table 3.

**Reach and representativeness of participants**

Only 15% of studies reported on all elements of this external validity dimension (Table 3). While almost all studies outlined the target population for generalisability (97%) and inclusion and exclusion criteria (97%), less reported the recruitment method (77%), enrolment rate (67%), and recruitment rate (67%) (Table 2). Just over half (54%) reported all of the specified participant characteristics – gender, age, any socioeconomic indicators (education, employment status, or income) – and participation by racial or ethnic minority groups. Only one in four studies included comparisons between individuals who participated versus either (1) those who declined to participate or (2) target population.

**Reach and representativeness of settings**

One in four studies reported on all elements of ‘reach and representativeness of settings’ (Table 3). Almost all studies provided details of the target setting for intervention delivery (92%); however, the remaining criteria were poorly described: inclusion and exclusion criteria (21%), how settings were recruited/reached to participate in delivering the intervention (14%) (Table 2). Only one study reported the participation level among eligible sites (5%); this was also the case for the representativeness of setting(s) (4%).

**Implementation and adaptation**

No studies reported on all elements of this external validity dimension (Table 3). Most studies described the intervention characteristics (97%) and the characteristics and training of delivery agents (95%). Less described the time to deliver the intervention (65%), and intervention delivery and exposure (65%)
Figure 1. PRISMA flow diagram.
Table 2. Number and percentage of studies reporting external validity elements.

<table>
<thead>
<tr>
<th>External validity dimension</th>
<th>Definition</th>
<th>Studies reporting Yes/Total</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Reach and representativeness of individuals</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Target population for generalizability</td>
<td>Is the intended target population acknowledged/stated (at the individual level) for which the findings intend to be generalised to?</td>
<td>38/39</td>
<td>97</td>
</tr>
<tr>
<td>Method to recruit target population</td>
<td>Was information provided about how the target population was recruited/reached (e.g., radio, newspaper, TV, school meeting)?</td>
<td>30/39</td>
<td>77</td>
</tr>
<tr>
<td>Inclusion or exclusion criteria</td>
<td>Were individual inclusion and exclusion criteria stated?</td>
<td>38/39</td>
<td>97</td>
</tr>
<tr>
<td>Enrolment rate</td>
<td>Is the enrolment rate or data needed to calculate the enrolment rate among individuals reported? Proportion of people who are eligible for participation who actually enrol in the study</td>
<td>26/39</td>
<td>67</td>
</tr>
<tr>
<td>Recruitment rate</td>
<td>Is the recruitment rate or data needed to calculate the recruitment rate among individuals reported? Proportion of potential participants (those invited or expressing interest) who actually enrol in the study</td>
<td>26/39</td>
<td>67</td>
</tr>
<tr>
<td>Representativeness of individuals</td>
<td>Are there comparisons between individuals who participated versus either (1) those who declined to participate or (2) target population?</td>
<td>10/39</td>
<td>26</td>
</tr>
<tr>
<td>Participant characteristics</td>
<td>Are all of the following reported: •Gender •Age •Any socioeconomic indicators (education, employment status, or income) •Participation by racial or ethnic minority groups</td>
<td>21/39</td>
<td>54</td>
</tr>
<tr>
<td><strong>Reach and representativeness of settings</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Target setting</td>
<td>Is the target setting for intervention delivery stated (such as workplace, general practice, outpatient facilities, churches, etc.)?</td>
<td>35/38</td>
<td>92</td>
</tr>
<tr>
<td>Method to recruit setting</td>
<td>Is information provided about how the site(s) within a given setting were recruited/reached to participate in delivering the intervention?</td>
<td>4/28</td>
<td>14</td>
</tr>
<tr>
<td>Inclusion or exclusion criteria</td>
<td>Were inclusion and exclusion criteria for selection of sites within a given setting stated? In the case of single sites, were the characteristics of the site described?</td>
<td>6/28</td>
<td>21</td>
</tr>
<tr>
<td>Participation rate</td>
<td>Is the participation level or data need to calculate the participation level among eligible sites reported (only applies to studies with more than one site)?</td>
<td>1/19</td>
<td>5</td>
</tr>
<tr>
<td>Representativeness of setting(s)</td>
<td>Are there comparisons between site(s) participating in the intervention and 1) those that decline to participate or 2) the target setting?</td>
<td>1/28</td>
<td>4</td>
</tr>
<tr>
<td><strong>Implementation and adaptation</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intervention characteristics</td>
<td>Were the intervention components described?</td>
<td>38/39</td>
<td>97</td>
</tr>
<tr>
<td>Intervention adaptation</td>
<td>Is information reported about how the study intervention is similar or different to original efficacy studies? Note: Only applicable to studies where an intervention is adapted from a previous trial</td>
<td>0/5</td>
<td>0</td>
</tr>
<tr>
<td>Time to deliver intervention described</td>
<td>Is the number and length of sessions or time required to deliver the intervention described?</td>
<td>24/37</td>
<td>65</td>
</tr>
<tr>
<td>Intervention delivery and exposure</td>
<td>Was the extent to which individuals were exposed to the intervention described? (e.g. proportion of planned intervention sessions actually attended (dose); content delivered as specified; provider adherence to intervention plan)</td>
<td>24/37</td>
<td>65</td>
</tr>
<tr>
<td>Delivery agents: characteristics and training</td>
<td>Is information provided on who delivered the intervention, such as the type of professional, or the amount of experience, skill or training required to deliver the intervention?</td>
<td>37/39</td>
<td>95</td>
</tr>
<tr>
<td>Methods to recruit delivery agents</td>
<td>Is information provided about how the delivery agents were identified/selected?</td>
<td>3/36</td>
<td>8</td>
</tr>
<tr>
<td>Delivery agents’ participation</td>
<td>Is the participation level amongst delivery agents reported (% of delivery agents agreeing to participate)?</td>
<td>4/35</td>
<td>11</td>
</tr>
<tr>
<td>Fidelity assessment: treatment receipt</td>
<td>Is information reported about whether the program was received as intended? (e.g. degree to which the participants understood the intervention and/or ability to perform the intervention skills)</td>
<td>4/39</td>
<td>10</td>
</tr>
<tr>
<td>Mechanisms for intervention effects</td>
<td>Was retrospective analysis conducted to identify the mediating variables through which the intervention achieved its effect?</td>
<td>2/39</td>
<td>5</td>
</tr>
<tr>
<td>External validity dimension</td>
<td>Definition</td>
<td>Studies reporting</td>
<td>Yes/Total²</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>----------------------------------------------------------------------------</td>
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<tr>
<td><strong>Outcomes for decision making</strong></td>
<td>Are outcomes (at least one) reported in a way that can be compared to either clinical targets or public health goals?</td>
<td>36/39</td>
<td>92</td>
</tr>
<tr>
<td>Adverse consequences</td>
<td>Does the article report whether they examined the occurrence of unintended consequences?</td>
<td>18/39</td>
<td>46</td>
</tr>
<tr>
<td>Effect moderators by participant characteristics</td>
<td>Are there any analyses of moderator effects by subgroups of participants</td>
<td>10/39</td>
<td>26</td>
</tr>
<tr>
<td>Effect moderator by delivery agent/setting</td>
<td>Are there any analyses of moderator effects by delivery agents or settings</td>
<td>0/37</td>
<td>0</td>
</tr>
<tr>
<td>Dose response effect of intervention (sensitivity)</td>
<td>Are there sensitivity analyses to assess dose-response effects of the intervention?</td>
<td>1/39</td>
<td>3</td>
</tr>
<tr>
<td>Total costs of intervention</td>
<td>Are total costs of the intervention presented?</td>
<td>6/39</td>
<td>15</td>
</tr>
<tr>
<td>Cost of intervention components</td>
<td>If costs are presented, were the costs itemized by intervention components (e.g., personnel, equipment)?</td>
<td>4/6</td>
<td>67</td>
</tr>
<tr>
<td>Cost effectiveness</td>
<td>If costs are presented, was there any analysis done to assess cost-effectiveness or cost-benefit of the program or policy?</td>
<td>3/6</td>
<td>50</td>
</tr>
<tr>
<td><strong>Maintenance / institutionalisation</strong></td>
<td>Are data reported on longer term effects on health-related outcomes, at least 12 months following program implementation, or environmental or policy change?</td>
<td>19/39</td>
<td>49</td>
</tr>
<tr>
<td>Long term effects (at least 12 months)³</td>
<td>Are data reported on the sustainability (or reinvention or evolution) or plans for sustainability of the intervention?</td>
<td>4/39</td>
<td>10</td>
</tr>
<tr>
<td>Institutionalization: sustainability / plans for sustainability</td>
<td>Are data reported on the number of individuals dropping out and/or lost to follow up</td>
<td>38/39</td>
<td>97</td>
</tr>
<tr>
<td>Attrition</td>
<td>Did the study report statistically significant differences in those that dropped out of treatment and those that finished?</td>
<td>19/38</td>
<td>50</td>
</tr>
<tr>
<td>Differential attrition (by condition or population sub-group)</td>
<td>Was information provided about acceptability of the intervention by stakeholders?</td>
<td>14/39</td>
<td>36</td>
</tr>
</tbody>
</table>

Notes:
1. Laws et al. (adapted from Green et al.)
2. Total = the no. of overall studies (n=39) minus the no. of studies reporting not applicable to the relevant element
3. Long-term results of intervention are defined as a minimum of six months following the last contact; long-term is defined as a minimum of 12 months by Laws et al.

Table 3. Number and percentage of studies reporting all elements of each dimension of external validity¹.

<table>
<thead>
<tr>
<th>External validity dimension</th>
<th>No./Total²</th>
<th>%</th>
<th>Studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reach and representativeness of participants</td>
<td>6/39</td>
<td>15</td>
<td>NOURISH, PRIMROSE, Baby Milk Trial, INSIGHT, BLISS, POI</td>
</tr>
<tr>
<td>Reach and representativeness of settings</td>
<td>9/38</td>
<td>24</td>
<td>ProKind, Baby Milk Trial, Minding the Baby, SLIMTIME, INSIGHT, BLISS, POI, Healthy MOMS, Healthy Beginnings</td>
</tr>
<tr>
<td>Implementation and adaptation</td>
<td>0/39</td>
<td>0</td>
<td>None</td>
</tr>
<tr>
<td>Outcomes for decision making</td>
<td>0/39</td>
<td>0</td>
<td>None</td>
</tr>
<tr>
<td>Maintenance / institutionalisation</td>
<td>1/39</td>
<td>3</td>
<td>INSIGHT</td>
</tr>
</tbody>
</table>

Notes:
1. Laws et al. (adapted from Green et al.)
2. No. taken as sum of no. of studies reporting yes or not applicable to each of the element. Total excludes any studies for which the external validity criterion was not applicable (e.g. Grow2Gether was a social media intervention therefore ‘research and representativeness of settings’ criterion was not applicable).
(Table 2). Delivery agents’ participation (11%), fidelity assessment: treatment receipt (10%), methods to recruit delivery agents (8%), and mechanisms for intervention effects (5%) were very poorly reported. Only five of the studies tested an intervention that was adapted from a previous trial - none reported on how the study intervention was similar or different to original efficacy studies.

Outcomes for decision making
No studies reported on all elements of ‘outcomes for decision making’ (Table 3). Almost all studies reported outcomes in a way that could be compared to either clinical targets or public health goals (92%) (Table 2). Less than half of studies reported whether they examined the occurrence of unintended consequences (46%). Only six studies reported the total costs of the intervention (15%); of these, four studies reported the cost of intervention components (67%), and three examined cost effectiveness (50%). Ten studies (26%) examined effect moderators by participant characteristics; however, none reported effect moderators by delivery agent/setting. Only one study (3%) reported a sensitivity analyses to assess dose-response effects of the intervention.

Maintenance / institutionalisation
Only one study – INSIGHT – reported on all elements of maintenance / institutionalisation (Table 3). Almost all studies (97%) reported on the number of individuals dropping out and/or lost to follow up (Table 2). Data on attrition by condition or population sub-group reported by 90% of studies (Note: we took condition to mean by intervention or control group). Only 50% of studies addressed the representativeness of completers/dropouts. Half of studies (49%) reported data on longer term effects on health-related outcomes (at least 12 months following program implementation, or environmental or policy change). Only 10% of studies reported on the sustainability (or reinvention or evolution) or plans for sustainability of the intervention. Only 36% reported on the acceptability of the intervention by stakeholders.

Discussion/ Conclusion
Early life interventions delivered by health professionals have the potential to influence important health behaviours, in addition to child weight. An understanding of the external validity of such interventions is vital to address their potential for translation and scalability, as well as replication efforts. In this systematic review we identified 39 studies, representing 46 interventions. External validity elements that were generally well described included target populations and settings, participant inclusion and exclusion criteria, intervention characteristics, delivery agents, outcomes, and attrition. Similar to other reviews of childhood obesity interventions33,34, however, we identified important gaps in the reporting of external validity elements within studies, and factors that could enhance translation and scale-up of interventions across all five external validity dimensions. External validity elements that were poorly described included: representativeness of individuals and settings, treatment receipt, intervention mechanisms and moderators, cost effectiveness, and intervention sustainability and acceptability.

Key gaps in informing the translation and scalability of health professional-delivered early life obesity prevention interventions were identified in this review. These included understanding the representativeness of settings, and whether these settings and delivery agents could be engaged to deliver these types of interventions in a sustained way, in a way that is acceptable to those involved. This is especially important given that only 16 of the 46 interventions (35%) in this review were clearly delivered as part of routine care, with a further two partly delivered as such, i.e. contacts as part of routine care but additional contacts also (Starting Early72 and STRIP86). The focus of the majority of studies was on establishing efficacy rather than effectiveness or how such interventions could be scaled up and translated into routine practice. This may account for the poor reporting of external validity in relation to settings and delivery agents.

Reporting of external validity elements considered important to inform decision makers was generally poor also. This included cost and cost-effectiveness measures, and an understanding of the intervention mechanisms and dose-response effects. While most interventions that are scaled up need to be adapted to fit the delivery context, knowing information about dose-response and the mechanism of intervention effects is essential in informing adaptions so that effectiveness of the intervention is not lost. The recent systematic review by McCrabb and colleagues highlights the decreased intervention effects when obesity interventions are scaled up – they found that effects on weight status, physical activity/sedentary behaviour, and nutrition reported in scaled-up interventions were typically 75% or less of the effects reported in pre–scale-up efficacy trials7. Reporting of fidelity components in our review was also varied – training (95%), delivery (65%), and receipt (10%). This has been noted in other childhood obesity-related reviews86,13, and has important implications for the interpretation, as well as the generalisability, of study findings.

Despite calls for greater attention to external validity for almost 40 years now12,13,40,41, we noted that problems with attention to generalisability persist. Only one trial within this review, the INSIGHT trial13, reported on all elements of the external validity assessment tool developed by RL29. Earlier this year, Huebschmann and colleagues made a further call for increased attention to external validity14. For trialists, there is a tension between internal validity and external validity, with preference historically for ensuring the former and minimising the risk of bias, at the expense of generalisability and applicability to real-world settings. Standard reporting guidelines such as the CONSORT statement for the reporting of randomized controlled trials32, the CONSORT extension for cluster trials33 and the CONSORT extension for pragmatic trials47 traditionally focus on internal validity elements, with limited focus and guidance around external validity. The TIDieR reporting guidelines for intervention description and replication somewhat address this gap35.

We acknowledge the challenging context in which triallists work and that there are many positive activities in this area. We
have a number of suggestions for moving work in this area forward nevertheless. Trialists could plan their interventions with scalability and sustainability in mind, giving due consideration to the type of trial conducted as well as the intervention characteristics. Few researchers plan for the sustainability of their interventions. The aforementioned reporting guidelines can be used in combination to report on study findings, with additional materials published to enhance external validity assessment, including protocols and more detailed information made accessible via supplementary materials or open access repositories. Researchers could also use models such as RE-AIM for reporting. Glasgow and Estabrooks note the challenges in comprehensively reporting on all RE-AIM dimensions within community and clinical settings with limited resources, however, highlighting that even well-funded NIH grants and published research studies, stating use of the RE-AIM framework, only employ it partially, and inconsistently when they do so. Inconsistencies in the degree to which authors report each RE-AIM dimension in its entirety as well as inaccuracies in reporting elements within each dimension have been highlighted by other authors also. Further work is needed with researchers to embed such frameworks appropriately.

Funding bodies, review panels, journals/journal editorial boards, and policymakers could also take action to promote the integration of external validity considerations into the funding, design, conduct, reporting, synthesis and translation of research. This need not be at the expense of internal validity, and can help facilitate credible research and knowledge translation. The inclusion of a PRECIS-2 graphic when proposing or reporting on a study can also be undertaken to enable the assessment of external validity.

Strengths and limitations
The strengths of this work are the use of a comprehensive and rigorous methodology, including a broad search strategy and range of databases, no language restrictions, and the screening of trials and extraction of data by two independent review authors. A number of limitations, however, must be noted. While we included journal articles, protocols, grey literature and supplementary materials, it is possible that researchers of the reviewed studies may have collected some of the information required to complete the external validity assessment but did not report it in the articles published to date. Furthermore, the external validity tool only codes items as present, absent, or not applicable. The extent, or quality, to which the studies report on the various external validity elements, e.g. fidelity, is not assessed; this may result in an over-estimation of the reporting quality of some studies. While it is not necessary for all studies to be strong on all of the external validity criterion, researchers, decision-makers and others could use this information, if provided, to make judgments as to the applicability or generalisability of a study or review.

Conclusion
This review examined the reporting of external validity elements within 39 studies encompassing 46 early-life health professional-delivered interventions. While such interventions have the potential to influence important health behaviours, in addition to child weight, we identified important gaps in the reporting of external validity elements within studies, and factors that could enhance translation and scale-up of interventions across all five external validity dimensions. External validity elements that were poorly described included: representativeness of individuals and settings, treatment receipt, intervention mechanisms and moderators, cost effectiveness, and intervention sustainability and acceptability. More emphasis is needed on research designs that consider generalisability, and the reporting of external validity elements in early life childhood obesity prevention interventions.

Data availability
Underlying data
All data underlying the results are available as part of the article and no additional source data are required.

Extended data
Open Science Framework: Health professional-delivered obesity prevention interventions during the first 1,000 days: A systematic review of external validity reporting. https://doi.org/10.17605/OSF.IO/G2ZMY

This contains the following underlying data:
- SearchStrategy_v1_HealthProfessional-deliveredObesityPreventionF1D.pdf (Search strategy)
- DataExtractionForm_v3_HealthProfessional-deliveredObesityPreventionF1D.pdf (Data extraction form)
- InterventionDescriptions_v7_HealthProfessional-deliveredObesityPreventionF1D.xlsx (Intervention descriptions)
- InterventionOutcomes_v5_HealthProfessional-deliveredObesityPreventionF1D.xlsx (Intervention outcomes)
- FullDetailsofExternalValidityAssessments_v2_HealthProfessional-deliveredObesityPreventionF1D.pdf (Full details of external validity assessments)
- SummaryTable-ExternalValidity-ByStudy_v3_HealthProfessional-deliveredObesityPreventionF1D.xlsx (Summary table of external validity assessments by study)

Reporting guidelines
Open Science Framework: PRISMA checklist for ‘Health professional-delivered obesity prevention interventions during the first 1,000 days: A systematic review of external validity reporting’. https://doi.org/10.17605/OSF.IO/G2ZMY

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

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