Mortimer, K; Ndamala, CB; Naunje, AW; Malava, J; Katundu, C; Weston, W; Havens, D; Pope, D; Bruce, NG; Nyirenda, M; Wang, D; Crampin, A; Grigg, J; Balmes, J; Gordon, SB (2016) A cleaner burning biomass-fuelled cookstove intervention to prevent pneumonia in children under 5 years old in rural Malawi (the Cooking and Pneumonia Study): a cluster randomised controlled trial. Lancet, 389 (10065). pp. 167-175. ISSN 0140-6736 DOI: https://doi.org/10.1016/S0140-6736(16)32507-7

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A cleaner burning biomass-fuelled cookstove intervention to prevent pneumonia in children under 5 years old in rural Malawi (the Cooking and Pneumonia Study): a cluster randomised controlled trial

Kevin Mortimer, Chifundo B Ndamala, Andrew W Naunje, Jullita Malava, Cynthia Katundu, William Weston, Deborah Havens, Daniel Pope, Nigel G Bruce, Moffat Nyirenda, Duolao Wang, Amelia Crampin, Jonathan Grigg, John Balmes, Stephen B Gordon

Summary

Background WHO estimates exposure to air pollution from cooking with solid fuels is associated with over 4 million premature deaths worldwide every year including half a million children under the age of 5 years from pneumonia. We hypothesised that replacing open fires with cleaner burning biomass-fuelled cookstoves would reduce pneumonia incidence in young children.

Methods We did a community-level open cluster randomised controlled trial to compare the effects of a cleaner burning biomass-fuelled cookstove intervention to continuation of open fire cooking on pneumonia in children living in two rural districts, Chikhwawa and Karonga, of Malawi. Clusters were randomly allocated to intervention and control groups using a computer-generated randomisation schedule with stratification by site, distance from health centre, and size of cluster. Within clusters, households with a child under the age of 4.5 years were eligible. Intervention households received two biomass-fuelled cookstoves and a solar panel. The primary outcome was WHO Integrated Management of Childhood Illness (IMCI)-defined pneumonia episodes in children under 5 years of age. Efficacy and safety analyses were by intention to treat. The trial is registered with ISRCTN, number ISRCTN59448623.

Findings We enrolled 10 750 children from 8626 households across 150 clusters between Dec 9, 2013, and Feb 28, 2016. 10 543 children from 8470 households contributed 15 991 child-years of follow-up data to the intention-to-treat analysis. The IMCI pneumonia incidence rate in the intervention group was 15.76 (95% CI 14.89–16.63) per 100 child-years and in the control group 15.58 (95% CI 14.72–16.45) per 100 child-years, with an intervention versus control incidence rate ratio (IRR) of 1.01 (95% CI 0.91–1.13; p=0.80). Cooking-related serious adverse events (burns) were seen in 19 children; nine in the intervention and ten (one death) in the control group (IRR 0.91 [95% CI 0.37–2.23]; p=0.83).

Interpretation We found no evidence that an intervention comprising cleaner burning biomass-fuelled cookstoves reduced the risk of pneumonia in young children in rural Malawi. Effective strategies to reduce the adverse health effects of household air pollution are needed.

Funding Medical Research Council, UK Department for International Development, and Wellcome Trust.

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that reduce exposure mainly by venting emissions to the outdoor environment with a chimney rather than by improving combustion efficiency. Romieu and colleagues\(^{10}\) compared a Patsari stove intervention with traditional open fire on respiratory symptoms and lung function in 552 women in Mexico. Adherence to the intervention was poor but the chimney stove reduced respiratory symptoms and lung function decline in those who used the stove. The RESPIRE trial randomised 534 households with a pregnant woman or infant in highland Guatemala to a chimney stove (with good compliance) or open fire and assessed the impact on pneumonia in children younger than 19 months. In intention-to-treat analyses, there was a non-significant reduction in incidence of physician-diagnosed pneumonia (primary outcome) in the chimney stove group (rate ratio [RR] 0.84 [95% CI 0.63–1.13]; p=0.26) and a significant reduction in physician-diagnosed severe pneumonia (RR 0.67 [0.45–0.98]; p=0.04). Access to cleaner cooking fuels and technologies is limited by many factors in the developing world although poverty is a key underlying issue for most.\(^{11}\) The Global Alliance for Clean Cookstoves (GACC) was launched in 2010 to tackle the lack of access to clean affordable energy through public–private partnerships. A central aim of GACC is for 100 million homes to adopt clean cookstoves and fuels by 2020.\(^{12}\) However, there is limited evidence to assess the potential health benefits of such an approach: globally there are no data from clinical trials of cleaner burning biomass-fuelled interventions and in Africa there are no data from clinical trials of any cookstove interventions.\(^{13}\) We set out to determine whether a cleaner burning biomass-fuelled cookstove intervention would reduce pneumonia incidence in children under 5 years of age living in rural Malawi compared with continuation of traditional open fire cooking (see video).

### Methods

#### Study design and participants

We did a cluster randomised controlled trial with two groups of equal size comparing the effects of a cleaner burning biomass-fuelled cookstove intervention with continued use of open fire cooking on pneumonia incidence in children under 5 years of age living in rural Malawi over a 2 year period (the Cooking and Pneumonia Study, CAPS). We defined 150 community-level clusters within villages across two districts of Malawi where we had established field research sites: Chikhwawa in the southern Shire river valley and Karonga on the northern

### Added value

This trial is the largest study of the effects of a biomass-fuelled cookstove intervention on health outcomes to date. The trial was done in very resource-poor populations in rural Malawi where pneumonia is one of the leading causes of morbidity and mortality in young children and almost all households burn biomass fuels for cooking. Data from 10 543 children from 8470 households across 150 community-level clusters contributed 15 991 child-years of follow-up data to the intention-to-treat analysis. The intervention comprising two biomass-fuelled cookstoves, a solar charger, repair and maintenance service, and user training had no effect on the risk of pneumonia in children under the age of 5 years in rural Malawi.

### Implications of all the available evidence

There is insufficient evidence on which to recommend replacing open fires with biomass-fuelled cookstoves for the prevention of pneumonia in young children in settings where biomass fuels are burned in open fires for cooking.
Malawi lakeshore. The Malawi College of Medicine Research Ethics Committee (ethics committee reference number P.11/12/1308) and the Liverpool School of Tropical Medicine Research Ethics Committee (ethics committee reference number 12.40) approved the protocol, which was peer reviewed, and a summary was published by The Lancet.15

Following community engagement exercises with village leaders and communities and the identification of a representative for each cluster, households with at least one child aged up to 4–5 years were invited to participate. Written informed consent (or witnessed thumbprint for those unable to read and write) was obtained at cluster and household level (parent or guardian of child) before participation. The trial was open to all consenting households with a usually resident child in the eligible age range. Households that became eligible for inclusion during the course of the trial (through birth, adoption, or in-migration) were recruited until 6 months before the trial ended.

Randomisation and masking
Clusters were randomly allocated to intervention and control groups using a computer-generated randomisation schedule with stratification by site, distance from health centre, and size of cluster. Randomisation was done by the trial statistician using dummy codes A and B to represent intervention and control groups; to ensure the statistician remained masked, the identity (allocation) of A and B was determined by a second, independent statistician. The trial statistician used a dummy randomisation code to develop statistical programs for data analyses before unblinding the trial data.

Procedures
Intervention households received two cleaner burning biomass-fuelled cookstoves (Philips HD4012LS; Philips South Africa, Johannesburg, South Africa), a solar panel to charge the battery for the stove fan, and user training. A fan incorporated into these cookstoves improves combustion efficiency; smoke emissions have been found to be reduced by around 90% compared with the open fire in laboratory testing.11 Cookstoves were repaired and replaced as needed. Stove use monitors were placed on one of the stoves in a randomly selected 10% sample of intervention households at baseline and at the 12 month visit to provide an objective measure of use, by recording temperature fluctuations. Control households continued using traditional cooking methods (typically open fires). At the start of the trial control households were informed they would receive the intervention at the end of the study period for equity and to maximise retention. Each household was visited every 3 months by fieldworkers although by the time the 21 month visit was due we were 3 months behind schedule and as such moved directly on to the final 24 month visit. Community engagement events were done regularly across all clusters to maintain engagement, facilitate discussion, and troubleshoot.

Outcomes
The primary outcome was the incidence of WHO-defined pneumonia in children under 5 years of age diagnosed by physicians, medical officers, or other appropriately trained staff at local health-care facilities routinely accessed by trial participants, unaware of intervention allocation.16

Secondary outcomes were the incidence of all pneumonia including those not meeting IMCI criteria, severe pneumonia, and death in children under 5 years of age. Additional outcomes included: the association between exposure to household air pollution and the development of pneumonia in children under 5 years of age in rural Malawi; prevalence and determinants of obstructive lung disease in adults in rural Malawi and the relationship between exposure to household air pollution and the rate of decline in lung function in adults in rural Malawi; affordability and cost-effectiveness of the intervention from household, health-care system, and societal perspectives; and what can be learned from trial participants and non-participants about adoption of the intervention that could inform effective implementation of the trial findings in the future. The findings of these additional outcomes will be reported separately.

We provided training and equipment including respiratory rate timers, thermometers, and pulse oximeters to support local health services in delivering high quality clinical assessments and documentation of pneumonia diagnoses and severity assessments.17 A supply of antibiotics was provided when local health service stocks ran out.

The IMCI pneumonia assessment protocol was used because chest radiographs were not consistently available and were not part of the usual diagnostic work-up in rural Malawi. Non-severe IMCI pneumonia was defined as the presence of cough or difficulty breathing and fast breathing (60, 50, or 40 breaths per min or higher in those aged <2 months, 2–12 months, and 1–5 years, respectively). Severe IMCI pneumonia was defined additionally by chest in-drawing, stridor, or any general danger sign (inability to drink or breastfeed, vomiting, convulsions, lethargy, or unconsciousness). Oxygen saturation of less than 90% was included as an additional and objective marker of severity.

In Malawi there is a patient-held medical record system. Patients present their Malawi Ministry of Health and Population health passport whenever they seek medical care. All trial participants were provided with a new health passport if they did not already have one with sufficient blank pages. A sticker was inserted explaining that the child was in the trial; a brief summary of the IMCI pneumonia assessment protocol; and boxes to tick about clinical criteria and severity if the patient was diagnosed with pneumonia. Malaria was tested with a rapid
Trial profile

**Figure:** Intention to treat (ITT) = intention to treat.

**Intervention**
- Number of children: 5297
- Age (months): 24.5 (16.1)
- Male: 2641 (49.9%)
- Female: 2656 (50.1%)
- Vaccination status (course completed)*
  - Diphtheria, tetanus, pertussis, hepatitis B and Haemophilus influenzae B: 2893 (79.4%)
  - Pneumococcal conjugate: 2122 (58.3%)
  - Polio: 2726 (74.8%)
  - Rotavirus: 1723 (47.0%)
  - Measles: 2166 (59.5%)
- Had pneumonia at least once in the preceding 12 months: 762 (15.4%)
- Had a cooking-related burn in the preceding 3 months: 219 (4.1%)

**Control**
- Number of children: 5246
- Age (months): 23.8 (16.1)
- Male: 2561 (48.8%)
- Female: 2685 (51.2%)
- Vaccination status (course completed)*
  - Diphtheria, tetanus, pertussis, hepatitis B and Haemophilus influenzae B: 2561 (48.8%)
  - Pneumococcal conjugate: 1774 (47.3%)
  - Polio: 2857 (76.1%)
  - Rotavirus: 1774 (47.3%)
  - Measles: 2685 (51.2%)
- Had pneumonia at least once in the preceding 12 months: 844 (17.4%)
- Had a cooking-related burn in the preceding 3 months: 289 (5.5%)

**Table 1 continues on next page**

## Statistical analysis

We aimed to include 150 clusters representing an estimated 10600 eligible children. Clusters were randomised in equal numbers to two intervention groups to provide, over the whole study period, about 21200 child-years of follow-up and 90-3% power to detect a reduction in the rate of pneumonia from five per 100 child-years in the control group to four per 100 child-years in the intervention group (proportionally, a 20% reduction). This conservative estimate of incidence was used to allow for potential reductions in the rate of pneumonia following pneumococcal vaccine introduction. We assumed an effective intra-cluster correlation (ICC) of 0-10 and a coefficient of variation in cluster size of 30-35%. Full sample size considerations are available online.

All primary analyses used intention-to-treat principles; the intention-to-treat population was all participants with at least one follow-up record before reaching 5 years of age. Secondary per-protocol analyses were also done; the per-protocol population included all participants 54 months of age or younger at recruitment and, in the intervention group, participants from households with at least one follow-up record at which stove use was reported. Generalised estimating equation (GEE)
modelling methods were used to assess the primary outcome, adjusting for clustering effects. Covariate adjusted analysis was done on the primary outcome by adding the following prespecified variables into the above GEE model: district, age of child at recruitment, sex, distance to nearest health centre, number of children younger than 5 years living in household, number of people in household who smoke regularly, other sources of fire or smoke (other than cooking) to which child was exposed on a daily (or almost daily) basis, socioeconomic status of household, number of previous pneumonia episodes experienced by child, and status of childhood vaccinations. Secondary outcomes were analysed similarly using GEE models; time to event data were analysed using Cox regression model with frailty for the cluster effect. One masked interim safety analysis was done and reviewed by the data monitoring committee to determine whether there were grounds to stop the trial for safety. Provision for an interim analysis for efficacy was included in the protocol although was not required by the data monitoring committee. SAS statistical package (version 9.3) was used. The full peer-reviewed statistical analysis plan is available online. The study is registered with ISRCTN, number ISRCTN59448623.

**Role of the funding source**
The funders had no role in the study design, data collection, analysis, interpretation, or writing of the report. The corresponding author had full access to all the study data and had final responsibility for the decision to submit for publication.

**Results**
We enrolled 10750 children from 8626 households across 150 community-level clusters in two sites in rural Malawi between Dec 9, 2013, and Feb 28, 2016. 207 children were lost to follow-up (103 from the intervention group and 104 from the control group) from 156 households (83 in the intervention group and 73 in the control group) between baseline and first follow-up visit (figure). Data were therefore available from 10471 children (5297 in the intervention group and 5220 in the control group) for at least one follow-up visit and form the intention-to-treat and safety dataset (figure). Baseline characteristics were balanced between intervention and control groups (table 1). There were 15 991 child-years of follow-up (7964 child-years from the intervention group and 8027 child-years from the control group). The last participant follow-up visit was on Sept 14, 2016.

We excluded 47 children (21 in the intervention group and 26 in the control group) because they were aged over 54 months at recruitment and 25 children from intervention households due to self-reported never use of the intervention to create the per-protocol population. The per-protocol population was therefore 10 471 children (5251 in the intervention group and 5220 in the control group) from 8429 households (4224 in the intervention group and 4205 in the control group).

<table>
<thead>
<tr>
<th>(Continued from previous page)</th>
<th>Intervention</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Household-level data</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of households</td>
<td>4256</td>
<td>4214</td>
</tr>
<tr>
<td>Fuel used regularly for cooking</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wood</td>
<td>2494 (58 6%)</td>
<td>2390 (56 7%)</td>
</tr>
<tr>
<td>Crop residues</td>
<td>1339 (31 5%)</td>
<td>1320 (31 3%)</td>
</tr>
<tr>
<td>Charcoal</td>
<td>563 (12 2%)</td>
<td>723 (17 2%)</td>
</tr>
<tr>
<td>Dung</td>
<td>20 (0 5%)</td>
<td>25 (0 6%)</td>
</tr>
<tr>
<td>Electricity</td>
<td>5 (0 1%)</td>
<td>13 (0 3%)</td>
</tr>
<tr>
<td>Paraffin/kerosene</td>
<td>2 (0 0%)</td>
<td>0 (0 1%)</td>
</tr>
<tr>
<td>Gas</td>
<td>0 (0 0%)</td>
<td>2 (0 0%)</td>
</tr>
<tr>
<td>Other</td>
<td>2 (0 0%)</td>
<td>0</td>
</tr>
<tr>
<td>Tobacco smoker in the household</td>
<td></td>
<td></td>
</tr>
<tr>
<td>746 (17 5%)</td>
<td>634 (15 0%)</td>
<td></td>
</tr>
<tr>
<td>Daily or almost daily exposure to smoke from‡</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Burning rubbish</td>
<td>1747 (41 0%)</td>
<td>1677 (39 8%)</td>
</tr>
<tr>
<td>Cooking for business</td>
<td>554 (13 0%)</td>
<td>513 (12 2%)</td>
</tr>
<tr>
<td>Brick production</td>
<td>177 (4 2%)</td>
<td>214 (5 1%)</td>
</tr>
<tr>
<td>Paraffin/kerosene lamps</td>
<td>87 (2 0%)</td>
<td>105 (2 5%)</td>
</tr>
<tr>
<td>Mosquito coils</td>
<td>44 (1 0%)</td>
<td>82 (1 9%)</td>
</tr>
<tr>
<td>Beer production</td>
<td>54 (1 3%)</td>
<td>44 (1 0%)</td>
</tr>
<tr>
<td>Other sources</td>
<td>63 (1 5%)</td>
<td>47 (1 1%)</td>
</tr>
<tr>
<td>Source of drinking water†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bore hole</td>
<td>1900 (44 6%)</td>
<td>1973 (46 8%)</td>
</tr>
<tr>
<td>Tap to house</td>
<td>401 (9 4%)</td>
<td>378 (9 0%)</td>
</tr>
<tr>
<td>Shared communal tap</td>
<td>389 (9 1%)</td>
<td>324 (7 7%)</td>
</tr>
<tr>
<td>Covered well</td>
<td>254 (6 0%)</td>
<td>233 (5 5%)</td>
</tr>
<tr>
<td>Lake or river</td>
<td>255 (6 0%)</td>
<td>157 (3 7%)</td>
</tr>
<tr>
<td>Open well</td>
<td>191 (4 5%)</td>
<td>253 (6 0%)</td>
</tr>
<tr>
<td>Other</td>
<td>3 (0 1%)</td>
<td>1 (0 0%)</td>
</tr>
<tr>
<td>Toilet facilities</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Simple pit latrine</td>
<td>3448 (81 0%)</td>
<td>3553 (84 3%)</td>
</tr>
<tr>
<td>None</td>
<td>763 (17 9%)</td>
<td>612 (14 5%)</td>
</tr>
<tr>
<td>Water toilet</td>
<td>37 (0 9%)</td>
<td>23 (0 5%)</td>
</tr>
<tr>
<td>Ventilated improved pit</td>
<td>8 (0 2%)</td>
<td>26 (0 6%)</td>
</tr>
<tr>
<td>Experienced a time in the last year when there was not enough food for the household to have its normal meals</td>
<td>2131 (50 1%)</td>
<td>2113 (50 1%)</td>
</tr>
<tr>
<td>Experienced a time in the last year when the household did not have money to buy bathing soap</td>
<td>2608 (61 3%)</td>
<td>2597 (61 6%)</td>
</tr>
</tbody>
</table>

Data are mean (SD) or n (%). *Vaccination status available for 7395 children (3642 in the intervention group and 3753 in the control group). †Data available for 9798 children (4954 in the intervention group and 4844 in the control group). ‡Household could give multiple responses.

Table 1: Baseline characteristics of the intention-to-treat population
The estimated ICC was 0·03. After adjustment for baseline values, the IRR for IMCI pneumonia for the intervention versus control was 1·05 (95% CI 0·93–1·18; p=0·44). The incidence rate for IMCI pneumonia in the overall per-protocol population was 15·70 (95% CI 15·09–16·32) per 100 child-years with an intervention versus control IRR of 1·02 (95% CI 0·91–1·13; p=0·75). The hazard ratio for time from enrolment to first pneumonia episode in the intervention versus control group was 1·06 (95% CI 0·93–1·21; p=0·29). The incidence rate for all pneumonia episodes including those not meeting IMCI criteria was 17·21 (95% CI 16·57–17·85) per 100 child-years with an intervention versus control IRR of 1·02 (95% CI 0·91–1·13; p=0·75).

In the intervention versus control group the IRR for severe pneumonia episodes defined by IMCI criteria was 1·30 (0·99–1·71; p=0·06), for oxygen saturation less than 90% 1·56 (0·70–3·46; p=0·28), and for death 0·76 (0·17–3·37; p=0·71) during year 1. Over the course of the trial, repairs or replacements were done on 13 192 (3·1 per intervention household) occasions for cookstoves and 5259 (1·2 per intervention household) occasions for solar panels. Control households were provided with two advanced cookstoves (ACE-1; African Clean Energy Company, Maseru, Lesotho) with a solar charger and solar lamp (part of the standard cookstove package by this time) at the end of the trial.

596 serious adverse events were reported excluding pneumonia. With the exception of a borderline statistically significant increased risk of malaria in the intervention group (IRR 1·32 [95% CI 1·03–1·69]; p=0·03) there were no differences in incidence rates of serious adverse events between the intervention and control groups (table 4). Of these, 19 (nine in the intervention group and ten in the control group) were cooking-related burns (IRR 0·91 [95% CI 0·37–2·23]; p=0·83). Additionally, 1505 burns (549 in the intervention group and 956 in the control group) not meeting seriousness criteria were reported (IRR 0·58 [95% CI 0·31–0·92]; p=0·0001). There were 53 deaths (malaria [14], pneumonia [seven], sepsis [five], malnutrition [five], accidental injury [four], gastroenteritis [one], burns [one], and other or unknown [16]) from the time of enrolment, 44 (25 in the intervention group and 19 in the control group) of which occurred in the intention-to-treat population (IRR 1·32 [95% CI 0·73–2·40]; p=0·03). The data monitoring committee reviewed trial safety data in March, 2015, and found no grounds to stop the trial early.

### Discussion

This community-level cluster randomised controlled trial in Malawi is the largest trial of a cookstove intervention on health outcomes to date. Malawi is one of the world’s poorest countries and within particularly impoverished rural areas it is frequent for households to have insufficient money at times to buy basic items like soap or food for their normal meals. The intention-to-treat analysis included data from 10 543 children from 8470 households across 150 community-level clusters.
representing 15,991 child-years of follow-up data. The number of children included was close to our original sample size estimate of 10,600 although the total follow-up time was less than the estimated 21,200 child-years. The incidence rate of IMCI pneumonia was high (15·67 per 100 child-years) and about three times higher than the deliberately conservative assumption used for our sample size calculation (five per 100 child-years). The observed incidence rate was more in agreement with 2010 estimates of IMCI pneumonia incidence in Malawi (24·3 per 100 child-years). Baseline characteristics were well balanced between the intervention and control groups. The intention-to-treat and per-protocol analyses with and without adjustment for baseline data were consistent in finding no effect of the intervention on the incidence of IMCI pneumonia in the young children in the study. Although limited by a smaller number of cases, analyses restricted to severe pneumonia episodes similarly found no beneficial effect of the intervention.

The intervention comprised a package of two cleaner burning biomass-fuelled cookstoves to provide two cooking hobs, a solar charger for the battery in the cookstove, a repair and replacement service, and user training. The intervention cookstove was selected on the basis of being among the cleanest burning biomass-fuelled cookstoves to provide a technical problem requiring replacement of the batteries. This issue could have been largely avoided had the cookstoves had better quality batteries. Cookstove malfunction was the most common reason given by households for not using the cookstoves for all cooking needs. Although we did repairs and replacements as quickly as possible there were inevitably short periods when households only had one functioning cookstove. When this alone was not sufficient for all cooking needs the household would have had to revert to using the open fire. Nevertheless, and despite these challenges, self-reported use of the cookstoves was probably as high as could be realistically expected with three-quarters of intervention households reporting the use of the stoves for all their cooking needs at 3 months, reducing to about half of households after 2 years; as is true in kitchens around the world there is no single cooking device suitable for every task.

We included stove use monitors on one of two cookstoves in a subset of intervention households to provide an objective measure of use. Although these monitors generated less usable data than anticipated due to technical problems, and their presence on just one of the two stoves might have affected which stove was used, they similarly indicate a waning in use over time. The impact on pneumonia in children might have been greater had adoption of the intervention been higher but the levels of adoption we achieved are probably higher than would be the case outside the context of a trial, particularly if an intensive and free repair and replacement service was not available.

Baseline data showed that in addition to almost exclusive use of biomass fuels for cooking, exposure to smoke from other sources including burning of rubbish, tobacco, and income generation activities was a common day-to-day experience. Additionally, as cookstoves were only issued to households that had a resident child younger than 5 years, exposure to smoke from neighbours’ or relatives’ cooking fires is likely. It is thus possible that these other air pollution exposures would have negated any potential beneficial impact of the cookstove intervention and that the beneficial impact of the cookstove intervention and that other sources including burning of rubbish, tobacco, and income generation activities was a common day-to-day experience. Additionally, as cookstoves were only issued to households that had a resident child younger than 5 years, exposure to smoke from neighbours’ or relatives’ cooking fires is likely. It is thus possible that these other air pollution exposures would have negated any potential beneficial impact of the cookstove intervention and that other air pollution exposures would have negated any potential beneficial impact of the cookstove intervention and that had even cleaner burning cookstoves been used, they would be unlikely to have had an impact. Personal exposure and exposure-response data, to be published separately, will provide additional insights here.

<table>
<thead>
<tr>
<th></th>
<th>Number of events</th>
<th>Incidence rate (cases per 100 child-years, 95% CI)</th>
<th>Number of events</th>
<th>Incidence rate (cases per 100 child-years, 95% CI)</th>
<th>Incidence rate ratio (95% CI)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Malaria</td>
<td>175</td>
<td>2.19 (1.87–2.52)</td>
<td>134</td>
<td>1.66 (1.38–1.95)</td>
<td>1.32 (1.03–1.69)</td>
<td>0.03</td>
</tr>
<tr>
<td>Sepsis</td>
<td>51</td>
<td>0.64 (0.46–0.82)</td>
<td>54</td>
<td>0.67 (0.49–0.85)</td>
<td>0.95 (0.64–1.41)</td>
<td>0.81</td>
</tr>
<tr>
<td>Gastroenteritis</td>
<td>24</td>
<td>0.30 (0.18–0.42)</td>
<td>28</td>
<td>0.25 (0.22–0.48)</td>
<td>0.86 (0.49–1.52)</td>
<td>0.62</td>
</tr>
<tr>
<td>Burns</td>
<td>9</td>
<td>0.11 (0.04–0.19)</td>
<td>10</td>
<td>0.12 (0.05–0.20)</td>
<td>0.91 (0.37–2.23)</td>
<td>0.83</td>
</tr>
<tr>
<td>Accidental injury</td>
<td>12</td>
<td>0.15 (0.07–0.24)</td>
<td>9</td>
<td>0.11 (0.04–0.18)</td>
<td>1.42 (0.58–3.45)</td>
<td>0.44</td>
</tr>
<tr>
<td>Malnutrition</td>
<td>5</td>
<td>0.06 (0.01–0.12)</td>
<td>5</td>
<td>0.06 (0.01–0.12)</td>
<td>1.01 (0.29–3.48)</td>
<td>0.99</td>
</tr>
<tr>
<td>Asthma</td>
<td>6</td>
<td>0.08 (0.02–0.14)</td>
<td>2</td>
<td>0.02 (0.01–0.06)</td>
<td>3.03 (0.51–18.11)</td>
<td>0.22</td>
</tr>
<tr>
<td>Other</td>
<td>36</td>
<td>0.45 (0.30–0.60)</td>
<td>36</td>
<td>0.45 (0.30–0.59)</td>
<td>1.02 (0.62–1.66)</td>
<td>0.94</td>
</tr>
<tr>
<td>Total</td>
<td>318</td>
<td>3.99 (3.55–4.43)</td>
<td>278</td>
<td>3.45 (3.05–3.86)</td>
<td>1.15 (0.95–1.38)</td>
<td>0.15</td>
</tr>
</tbody>
</table>

Table 6: Incidence rate and incidence rate ratio of serious adverse events excluding pneumonia.
There was a high incidence of serious adverse events unrelated to cooking across the trial population reflecting the expected range and frequency of serious childhood illnesses seen in rural Malawi. There was a borderline statistically significant increased risk of malaria in the intervention versus the control group, which might be a false positive finding reflecting multiplicity of secondary safety analyses. Data from elsewhere suggest smoke from open fire cooking is unlikely to protect against malaria vectors. Nevertheless, this is an area of uncertainty and this observation warrants further investigation. Serious adverse events related to cooking (all burns) resulted in considerable morbidity and one death in our study population. Less serious burns were a common occurrence with 508 (4.8%) of 10,543 children reporting a burn in the 3 months before study enrolment and 1505 burns not meeting seriousness criteria reported during the study. Although serious burns were evenly distributed across the trial groups, there was a substantial 42% reduction (IRR 0.58 [95% CI 0.51–0.65]; p<0.0001) in risk of non-serious burns in the intervention group, suggesting the intervention offers considerable safety advantages over the open fire.

The findings of our trial add substantially to the trials of chimney stoves in Mexico (552 women followed up for 10 months) and Guatemala (518 children from 534 households contributing 560 child-years of follow-up). Although none found a statistically significant effect on the primary health outcome of interest, all point to benefits of the interventions in secondary analyses—there was a reduction in burns in Malawi, a reduction in respiratory symptoms, and decline in lung function in adults in a per-protocol analysis of intervention adopters in Mexico, and a reduction in severe acute lower respiratory tract infections in Guatemala.

Although the design of our trial was necessarily open, considerable effort was put into minimising the risk of bias associated with a non-blinded study. Specifically, we provided training to fieldworkers to ensure participants were treated in the same way irrespective of trial group; scheduled the same frequency of field visits to intervention and control villages; collected primary outcome data in the same way for intervention and control villages; provided training to local health-care workers about how to make an IMCI diagnosis of pneumonia, the importance of carrying out assessments, and treatment based on clinical and laboratory assessment; and provided independent source document verification for all pneumonia diagnoses. The trial was done in a challenging environment where poverty is widespread and health services are stretched. We used the widely adopted WHO IMCI diagnostic criteria for pneumonia implemented by independent health-care workers because this approach is most reflective of the reality on the ground where access to more sophisticated diagnostic tools such as chest radiographs are not consistently available. Although this pragmatic approach might lack specificity, it is the best measure of the entity reported as pneumonia and associated with morbidity and mortality among children under 5 years of age in Malawi and elsewhere in sub-Saharan Africa.

In conclusion, we found no evidence of a reduction in pneumonia in young Malawian children in a rigorously conducted clinical trial of a cleaner burning biomass-fuelled cookstove intervention. The lack of effect on pneumonia might be explained by exposure to other sources of air pollution, including rubbing burning and tobacco smoke, that could have overwhelmed any potential effects of the cookstoves. It is also possible that the cookstoves used simply did not reduce emissions sufficiently to have an effect. An important implication of these observations is that tackling any individual source of air pollution exposure in isolation is unlikely to be effective for improving health; an integrated approach to achieving clean air that tackles rubbish disposal, tobacco smoking, and other exposures, as well as robust cleaner cooking solutions (eg, cleaner stoves and fuels) that achieve a high rate of acceptance is probably needed to deliver health benefits.

Contributors
KM, DP, NGB, MN, JG, AC, JB, and SBG were responsible for the designing the study. KM, CBN, AW, JM, CK, WW, DW, DH, AC, JB, and SBG acquired the data. DW and KM analysed the data. KM, WW, DH, DP, NGB, MN, DW, AC, JG, JB, and SBG interpreted the data. All authors contributed to the writing of the manuscript, approved of the version to be published, and agreed to be accountable for all aspects of the work.

Acknowledgments
The trial was funded by a Joint Global Health Trials Grant from the Medical Research Council, UK Department for International Development, and Wellcome Trust (MR/K006533/1) and a New Investigator Research Grant from the Medical Research Council (MR/L002515/1). Additional support was provided by a MRC Partnership Grant (MR/L009242/1). JG is part-funded by the North Thames Collaboration for Leadership in Applied Health Research and Care. We thank the trial participants, village leaders and CAPS representatives, the study team in Chikhwawa (Patience Korera, Frank Mbalume, Emma T Nyirenda, Sarah Magombo Kamphandule, Aosenji Manyiska, Yamikani Dickson, Bright Mnesa, Caroline Kumbuyo, Martha Chipeta, Gloria Banda, Ernest Mwangoka, Suzgo Mkandawire, Wezze Nyapiyigot, Fletcher Nangupeta, Kondawani C Munsuli, Frank Jamesar, Steven Lankem, Nicholas Mwenda, James Chirambo, and others), the study team in Chilumba, Karonga (Alexander Nkosi, Andy Chisunda, Angellina Sakala, Bedgeman Chikweywa, Blessings Nyirenda, Calvin Mwalwanda, Desire Augustine, Dokko Chihana, Donaria Zgambo, Grecium Nkhono, Hawkins Ndiowe, Jimmy Mbawla, Levie Gondwe, John Mughogo, Kelvin Gondwe, Louis Chavula, Donald Kalua, Michael Nyirenda, Nathan Madise, Patrick Luhanza, Thembu Chirowa, Timothy Mwaungulu, Tyness Kandonga, Tugiwie Sikali, and Wazika Nyinu), the study team at Liverpool School of Tropical Medicine (Jane Ardyre, Debbie Jenkins, Rachael Lloyd, Andrew Bennett, James Smedley, Tao Chen, and Brian Faragher), the central support from Malawi Liverpool Wellcome Trust Programme (Moses Kamzazi, Asimenye Kayuni, Owen Mwenefumbo, and the work.

Authors contributed to the writing of the manuscript, approved of the version to be published, and agreed to be accountable for all aspects of the work.
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