

Preventing severe asthma exacerbations in children: A randomised trial of mite impermeable bedcovers

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AUTHOR CONTRIBUTIONS

Clare Murray and Angela Simpson had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

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Acquisition, analysis, or interpretation of data: Clare Murray, Angela Simpson, Adnan Custovic, Helen Sumner, Phil Foden, Elizabeth Shepley

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Preventing exacerbations by avoiding mite allergen

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At a glance commentary

Scientific knowledge on the Subject: Asthma exacerbations in children are a leading cause of hospitalisation. Exposure in sensitised individuals in synergy with viral infections greatly increases hospital admission risk. In the developed world house dust mite is the commonest sensitising allergen. Studies to date have not investigated the effect of allergen avoidance on asthma exacerbations and hospital admissions in children.

What this study adds to the field: The use of mite impermeable bedding in mite sensitised asthmatic children can significantly reduce the risk of severe exacerbations resulting in emergency hospital attendance.

This article has an online data supplement, which is accessible from this issue's table of content online at www.atsjournals.org

ABSTRACT

Rationale: Allergen exposure in sensitised asthmatics interacts with viruses in increasing the risk of asthma exacerbation.

Objectives: To evaluate the use of house dust mite impermeable bedding on severe asthma exacerbations in children.

Methods: We randomized mite-sensitised asthmatic children (3-17 years), following an emergency hospital attendance with an asthma exacerbation, to receive mite-impermeable (Active) or control (Placebo) bed encasings.

Measurements and main results: Over a 12-month intervention period the occurrence of severe asthma exacerbations were investigated. Of 434 asthmatic children who consented, 286 (mean age 7.7 years, 65.8% male) were mite sensitised and 284 were randomised (146 Active; 138 Placebo). At 12 months, significantly fewer children in the Active group had attended hospital with an exacerbation compared to the Placebo group (36/123 [29.3%] versus 49/118 [41.5%], $p=0.047$). In the multivariable analysis, the risk of emergency hospital attendance was 45% lower in the Active group (Hazard Ratio 0.55 [95%CI, 0.36-0.85], $p=0.006$) compared with the Placebo group. The annual rate of emergency hospital attendance with exacerbations was 27% lower in the Active compared with the Placebo group, but this did not reach significance (Estimated marginal mean [95% CI]: Active 0.38 [0.26-0.56] vs Placebo 0.52 [0.35-0.76], $p=0.18$). No difference between the groups in the risk of prednisolone use for exacerbation was found (Hazard Ratio 0.82 [0.58-1.17], $p=0.28$).

Conclusions: Mite-impermeable encasings are effective in reducing the number of mite sensitised asthmatic children attending hospital with asthma exacerbations, but not the

Preventing exacerbations by avoiding mite allergen

number requiring oral prednisolone. This simple measure may reduce the health care burden of asthma exacerbations in children.

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INTRODUCTION

Asthma is the most common chronic disease in childhood. Although most children with asthma are well-controlled on pharmacotherapies, a significant number experience exacerbations which remain one of the commonest reasons for paediatric hospital admission in the developed world (1). This unscheduled care accounts for a large proportion of asthma costs, and a single exacerbation can increase annual costs more than 3-fold (2). Previous hospital admissions predict future hospitalizations (3). Respiratory virus infections are major risk factors for hospital admission (4-6), particularly amongst children who are exposed to allergens to which they are sensitised, where these factors act synergistically to markedly increase the risk of hospital admission (7, 8). However, disrupting this interaction in atopic asthmatics is challenging. There are currently no available vaccines for viruses which cause the majority of exacerbations. Allergen-specific immunotherapy is generally not recommended for patients with uncontrolled asthma (9). Anti-IgE monoclonal antibody (omalizumab) can reduce asthma exacerbations, but its use is limited to the most severe cases of asthma because of high cost and requirement for regular injections (10). Avoidance of allergen remains a potentially cost-effective intervention. However, no studies to date have investigated the effect of allergen avoidance on asthma exacerbations and hospital admissions, instead focussing on symptom scores, medication usage and lung function.

House dust mite (HDM) is a common allergen linked to expression of asthma, with ~65% of UK asthmatic children demonstrating sensitisation (7). Although high HDM exposure has been linked to asthma severity (11), a meta-analysis of 44 trials of mite avoidance was unable to demonstrate any clinical benefit of measures designed to reduce mite exposure, and concluded

Preventing exacerbations by avoiding mite allergen

that mite control measures could not be recommended for asthma (12). However, this meta-analysis included many studies where no exposure reduction was achieved, and did not distinguish between adult and paediatric studies. Indeed, most studies which suggest benefits of mite avoidance have been conducted in children (13-17). However, these studies were either small (13, 15, 17), used multifaceted interventions making blinding difficult (13, 17), targeted multiple allergens (14, 16), or were conducted in populations which have poor access to healthcare/medications (14, 16). Given the evidence of a synergism between viral infection and allergen exposure in increasing the risk of asthma exacerbations in sensitised individuals (7, 18), we hypothesized that effective reduction in mite exposure may reduce the risk of exacerbations in these patients.

In this double-blind study, Preventing asthma exacerbations by avoiding mite allergen (PAXAMA), we compared the effect of mite-impermeable bed covers to that of placebo covers in reducing the risk of severe asthma exacerbations in mite-sensitised asthmatic children, who had recently attended hospital with an asthma exacerbation. Some of the results of these studies have been previously reported in the form of an abstract (19).

METHODS

STUDY DESIGN

This randomized, double-blind, placebo-controlled, parallel-group study of the effect of mite-impermeable bed covers on the risk of severe asthma exacerbations in mite-sensitised asthmatic children was conducted across 14 hospitals with acute pediatric secondary care services in North-West England. Children were recruited between November 2011 and May

Preventing exacerbations by avoiding mite allergen

2013 and were followed for 12 months. The protocol was approved by local research ethics committee (NRES Committee North-West/Lancaster, REC approval number 11/ NW/0262).

STUDY PARTICIPANTS

We screened children aged 3-17 years with physician-diagnosed asthma who had presented to hospital with an asthma exacerbation. Children were excluded if already using allergen-impermeable bedding, born prematurely (<36 weeks) or had another respiratory disease. Participants were skin-prick tested once their exacerbation had resolved, to HDM, cat, dog, pollen and other pet allergens if applicable (Stallergenes, Paris, France), and classed as sensitized if the weal diameter was at least 3mm greater than the negative control. Only children sensitised to HDM (+/- other allergens) were eligible for randomisation. Parents provided written informed consent and children assent.

Randomisation and Masking

Children were randomly assigned 1:1 to active or placebo encasings using a computer-based minimisation procedure by a researcher who was not otherwise involved in the study. Children were stratified for age (3-10 years; 11-17 years), household cigarette smoking, pet sensitisation/ownership and treatment level (GINA steps 1-2; ≥ 3 ; eMethods, Online Supplement) in a double-blind manner. To maintain blinding no other information on HDM avoidance was given. All participants received identical printed washing instructions for the supplied encasings (eMethods, Online Supplement). The active encasings (Astex Pristine, ACP solutions Ltd, Gloucestershire, UK) were selected because their mite-proof efficacy had been demonstrated previously (20). Placebo encasings (made from poly-cotton) were custom manufactured (Musbury Fabrics, Rossendale, UK), to match the active encasings (Figure E1).

Preventing exacerbations by avoiding mite allergen

Neither encasings contained a label. If more than one child from a family were allocated to the study, the second child was enrolled in the same arm as the index child, to avoid potential unblinding. Researchers fitting covers and collecting dust samples for allergen analysis were not involved in follow-up.

Procedures

Children had their inhaler technique checked and corrected if necessary. Encasings were fitted to the pillow, mattress and duvet of the child's bed. Other beds in the same room and beds in which participants spent >1 night per week were also encased.

STUDY ASSESSMENTS

Baseline evaluation included questionnaires on demographics, past medical/family history, sleeping arrangements, pet exposure and medication use. Interviewers masked to the child's group assignment conducted telephone interviews with the primary caregiver at one, four, eight and twelve months to collect data on exacerbations, unscheduled medical care and medication. Quality of life (QOL) was assessed using Pediatric Asthma Caregiver's Quality of Life Questionnaire PACQLQ(21) completed by parents, mini Pediatric Asthma Quality of Life Questionnaire (PAQLQ)(22) completed by children age ≥ 7 years, and asthma control by Asthma Control Questionnaire (ACQ)(23) completed in children aged 6 years or over.

Mite allergen (Der p 1) was measured in vacuumed dust samples collected from the child's mattress and lounge floor prior to fitting the encasings and at 12 months, using enzyme-linked immunosorbent assay (Indoor Biotechnologies, Cardiff, UK; eMethods, Online Supplement)

OUTCOME MEASURES

Preventing exacerbations by avoiding mite allergen

The primary outcome was the occurrence of severe asthma exacerbations during the 12-month intervention period. We used ATS/ERS definition of severe exacerbations (24), including:

(A) A hospitalization or emergency department (ED) visit because of asthma, requiring systemic corticosteroids (OCS) – abbreviated to emergency hospital attendance;

(B) Use of OCS or an increase from a stable maintenance dose, for ≥ 3 days (includes all OCS courses whether associated with an emergency hospital attendance or not).

Secondary endpoints included change in controller treatment from baseline to 12 months, PACQLQ(21), mini PAQLQ(22) and ACQ(23) scores. Compliance and acceptability of intervention was recorded.

STATISTICAL METHODS

Power calculation

Data from UK General Practice Research Database (GPRD; www.gprd.com) estimated that children who had ≥ 1 course of OCS in the previous 12 months had a mean exacerbation rate of 1.5/year (variance 1.02). For 90% power to detect a 30% reduction in exacerbation rate during the 12-month intervention period, 114 children per group were required, at a two-sided significance level of 0.05. Assuming 20% lost during follow-up, we aimed to randomise 284 children.

Data Analysis

Baseline characteristics were compared between groups using t-tests, Mann-Whitney U and chi-squared tests as appropriate. Efficacy analysis was performed according to the intention-to-treat principle (per-protocol analysis in supplement). Outcomes were assessed between the groups using chi-squared tests and logistic regression for children who completed 12 months of

Preventing exacerbations by avoiding mite allergen

follow-up. Cox regression analysis assessed time to first emergency hospital attendance and prednisolone usage, and included all evaluable data with censoring for those who did not complete 12 months follow-up. Negative binomial generalized linear models analysed count data for outcomes; results expressed as estimated marginal means (EMM) and 95% confidence intervals (CI) for annual rates/participant (25). Multivariable models were adjusted for age, gender, ethnic group, maintenance asthma treatment, hospitalisations in the 12-months prior to randomisation, index of multiple deprivation and tobacco smoke exposure. General linear models with repeated measures were used to compare mite allergen levels and prescribed treatments across time between the groups. Der p 1 levels were $\log_e$ transformed to normalise the data prior to analysis; results presented as geometric means (GM). The conventional two-sided 5% significance level was used.

Exploratory subgroup analyses (not pre-specified) were conducted based on age, GINA step, sensitisation status, exposure to smoking and socioeconomic status (eMethods in Supplement) (26).

Analyses were carried out using SPSS 22 (IBM, New York, USA) and Stata 13 (StataCorp, Texas, USA).

RESULTS

PATIENTS

From November 2011 to May 2013, 434 children were screened to take part in the study. Of those, 286 were HDM-sensitised and 284 underwent randomization (146 Active; 138 Placebo; Figure 1). Baseline characteristics and Der p 1 levels were similar in both groups (Table 1;

Preventing exacerbations by avoiding mite allergen

Tables E1, E2). Twelve month follow-up was completed in 123 (84.2%) in the Active arm and 118 (85.5%) in the Placebo arm; overall, 208 children (73.2%) reported full compliance throughout the study (Per-protocol analysis).

Primary outcome

A) Hospitalization or ED visit because of asthma requiring systemic corticosteroids

Significantly fewer children in the Active group attended hospital with one or more severe asthma exacerbations compared to the Placebo group (36/123 [29.3%] versus 49/118 [41.5%]; OR 0.58 (0.34, 0.99), $p=0.047$; Figure 2A).

Time to first exacerbation requiring emergency hospital attendance was significantly longer in the Active group ($p=0.041$), and the risk of emergency hospital attendance was 45% lower in the Active group (Hazard ratio (HR) 0.55 [0.36-0.85], $p=0.006$), compared with the Placebo group (Figure 3; Table E3, multivariable model, adjusted for age, gender, GINA step, ethnicity, deprivation score and tobacco smoke exposure). Although, the annual rate of emergency hospital attendance was 27% lower in the Active compared with the Placebo group, this did not reach significance (EMM [95% CI]: Active 0.38 [0.26-0.56] vs Placebo 0.52 [0.35-0.76], $p=0.18$). The distribution of the numbers of attendances did not differ between the groups ($p=0.5$; Figure E2).

Per protocol analysis is presented in the Online Supplement (Figures E3-E6); the risk of emergency hospital attendance was 54% lower in the Active group compared to the Placebo group (HR 0.46 [0.28-0.76]; $p=0.002$).

B) Use of oral corticosteroids for ≥ 3 days

Preventing exacerbations by avoiding mite allergen

There was no difference in the numbers of children who received a course of OCS for an asthma exacerbation (whether associated with an unscheduled hospital or general practitioner attendance or by a home rescue pack) between the groups (Active 48.8% vs Placebo 50.0%, $p=0.85$, Figure 2B).

Investigating the time to first OCS use, there was no difference between groups in the univariable analysis ($p=0.67$) or in the multivariable model (HR 0.82 [0.58-1.17], $p=0.28$). The annual rate of OCS courses prescribed was not different between the groups (EMM [95% CI]; Active: 0.77 [0.55-1.06] vs Placebo: 0.85 [0.62-1.16], $p=0.57$). Per protocol analysis is presented in the Online Supplement (Figure E3b).

SECONDARY OUTCOMES

Mean values for PACQLQ and ACQ at each time are presented in Table E4 in the online supplement. Parents of children in both groups reported significantly improved PACQLQ between one and 12 months (mean difference [95% CI]; Active: 0.50 points [0.14-0.8], $p=0.007$; Placebo: 0.57 points [0.12-1.02], $p=0.01$), with no difference between the groups. Although significant improvement in ACQ score over time was observed only in Active children (-0.56 points, [-0.18,-0.93], $p=0.004$), and not in those with Placebo covers (-0.25 points, [-0.61, 0.11], $p=0.16$), there was no difference between the groups.

There was no difference in GINA step between the two groups at baseline (Table1, Table E1). At the end of the intervention period, GINA step had been increased in 10.7% of the Active group and 14.5% of Placebo group ($p=0.37$). Children who had any exacerbations during follow-up were more likely to have their GINA step increased by the end of follow-up compared to

Preventing exacerbations by avoiding mite allergen

children who did not suffer an exacerbation (27.1% vs 4.5% respectively, $p<0.001$), irrespective of group allocation.

Children in the active group were more likely to complain about the encasings (Table E5).

Despite this, the number adhering to the intervention at 12 months was similar in both groups (101 Active, 107 Placebo, $p=0.11$). Amongst all those fully compliant with the bedding almost 90% reported they would continue to use the encasings after the study.

MITE ALLERGEN LEVELS

Der p 1 levels in dust from the child's mattress was reduced by 84% in the Active group following the intervention, with no change in the Placebo group ($p<0.001$; Figure 4). Der p 1 in the lounge floor was unchanged in both groups ($p=0.48$; Figure E7).

Post-hoc analyses

In a multivariable Cox regression analysis (Table E6) a reduction in risk of an emergency hospital attendance was seen for children in the Active group aged 3-10 years (HR 0.54 [0.33-0.87], $p=0.01$, Figure E8), in those sensitised only to mite ($p=0.04$), in those from non-smoking homes ($p=0.02$), in those on GINA treatment step ≥ 3 ($p=0.03$) and in those from the most deprived homes ($p=0.01$). None of the interaction terms were significant however. Also, in younger children (3-10years), a non-significant reduction in risk of OCS use was seen in those in the Active group (HR 0.69 [0.46-1.04], $p=0.08$, Figure E9).

DISCUSSION

In our study of HDM allergic children who had recently suffered a severe asthma exacerbation, the risk of further severe asthma exacerbations requiring an emergency hospital attendance

Preventing exacerbations by avoiding mite allergen

was reduced by 45% in those who had mite-impermeable encasings fitted to the mattress, pillow and duvet. This is the first study of the effect of such an intervention on exacerbations in children. The annual rate of emergency hospital attendance, though 27% lower in the active compared to the placebo group, was not significantly reduced ($p=0.18$). There was no difference in the proportion of children requiring courses of oral steroids for asthma exacerbations. The encasings were highly effective in reducing recoverable mite allergen.

Asthma exacerbations have been ranked highly by clinicians and parents as important outcomes for clinical trials in children (27). Although comparatively rare events, severe asthma exacerbations result in many hospital admissions which are particularly costly, emphasizing the relevance of this as an outcome (28). Using real data from UK GPRD to power the study, we estimated that the exacerbation rate would be 1.5/annum for children who had suffered an exacerbation in the previous year. As previous hospitalizations/exacerbations are amongst the best predictors of future risk (3,29), we recruited children when attending hospital with an exacerbation. However, our observed exacerbation rate during follow-up was materially lower (Placebo group: 0.85 oral corticosteroid courses/annum), reducing our power to detect significant differences between the groups. That we were able to detect a statistically significant difference between groups for numbers of children requiring hospital attendances for asthma exacerbations reflects the large effect size seen.

Although the number of children who experienced any emergency hospital attendance with an asthma exacerbation was significantly reduced following the active intervention, some children continued to have hospital attendances, a few of them having multiple attendances. However, the distribution of multiple attendances did not differ between treatment groups.

Preventing exacerbations by avoiding mite allergen

As with many treatments for asthma (e.g. long acting Beta-agonists, leukotriene receptor antagonists), it is clear that some individuals respond to the treatment and others do not (30), but predicting those who will respond is challenging. Although our trial was not powered to carry out subgroup analyses, we performed exploratory analyses in an attempt to identify the characteristics of the children who showed the best response; this indicated that younger children, those sensitised only to mite, those with more severe disease (GINA 3+) and those not exposed to smoking had fewer emergency hospital attendances. Similar subgroup analyses in those requiring OCS courses for asthma exacerbations also suggested that younger children may be more likely to respond to this intervention. Whilst recognising that the subgroup analysis is exploratory, we propose that allergen avoidance may be more effective in younger children, in whom the disease may have been present for a shorter time. This may be analogous to occupational asthma, where removal of allergen exposure is effective if done soon after the onset of disease (31), and may explain the differences between our results compared to large studies in adults (32). In addition, younger children may spend a higher proportion of time in bed, making this dust reservoir a potentially larger contributor to personal mite exposure than in older children or adults. Recent reports in adults suggested that mite exposure may be higher during the daytime, and may reflect lifestyle and clothing worn (33). We speculate that personal allergen exposure is different in young children, who generally wear clothes that can be hot washed, and undertake different activities. We have no evidence that the younger children were more compliant with the intervention.

Despite the risk of emergency hospital attendance being reduced in the Active group, the risk of receiving OCS was not significantly reduced, although a trend was seen in younger children. A

Preventing exacerbations by avoiding mite allergen

small number of OCS courses were administered by parents using home rescue packs without contemporaneous medical direction (eight, three day courses, in total in 6 children), some of which may have been unnecessary. Unfortunately we were unable to assess children to confirm the presence of an exacerbation; care was provided by their family doctor or by urgent care services as the parents saw appropriate. It may be that the study intervention genuinely reduced the severity of exacerbations resulting in fewer hospital attendances, but not fewer courses of OCS.

Many factors influence consulting behaviours in parents with sick children; our recruitment strategy may have selected those more likely to present to hospital. Indeed, in our population, the majority of exacerbations resulted in an emergency hospital attendance (~70%). It is likely however, that those exacerbations requiring an emergency hospital attendance were more severe and regardless are certainly more expensive to the provider, and so we believe that the reduction seen is of clinical importance.

In order to establish that the reduction in exacerbations seen was not due to changes in controller medication we examined changes in prescribed medication during follow-up and found no difference between the groups. All treatments were prescribed by the participants' usual physicians who were blind to the treatment allocation and not influenced by the study team.

As there is no QOL score for asthma sufferers or caregivers validated for use in children under seven years, we used the PACQLQ for all participants (recognising that this has limitations), and the mini PAQLQ for children over seven years. There were no between-group differences in quality of life (both tending to show within-group improvements). Interestingly, despite recent

Preventing exacerbations by avoiding mite allergen

exacerbations, both groups reported good QOL and control at baseline, leaving little prospect of demonstrating significant between group differences.

Children who were in the Active group complained more about the bedding than those using the placebo covers. It is possible that this difference in perception led to unblinding of individuals (i.e. believing that they must have the real covers because they are uncomfortable). However, we believe this is unlikely to have affected the results of the study given the objective nature of our outcomes. It is important to note that compliance with the covers was not significantly different between the groups, and adherence (>70%) appeared to be at least as good, if not better than, with medications usually prescribed for asthma (e.g. inhaled corticosteroids ~50% (34)).

There are some other limitations to our study. All data on exacerbations and OCS use was reported by parents/carers and not confirmed by their primary care physician. However, we gathered information from parents on a 3 monthly basis and therefore recall should not be a significant issue and we would expect any bias to be similar across the groups. We were unable to measure adherence to prescribed treatment within this study, and although it is unlikely that one arm was more adherent than the other, we cannot exclude this from having occurred.

Evidence from previous studies suggest that viruses and allergens in sensitised individuals act synergistically to increase the risk of asthma exacerbation and hospitalisation. As we have no information on the trigger for individual exacerbations (viral or otherwise) we were unable to perform an analysis to identify whether the effectiveness of the intervention was dependent upon the trigger.

Preventing exacerbations by avoiding mite allergen

CONCLUSIONS

We found that the simple and relatively cheap intervention of mite allergen impermeable bed encasings, costing around £130/US \$200, is effective in reducing emergency hospital attendance with severe asthma exacerbations. In the population we have studied we estimate that approximately 8 children would need to be treated in order to prevent one child having any hospital attendances in the following year. It is likely that there is a subgroup of children in whom the intervention is more beneficial and although our subgroup analysis would suggest this might be younger, mono-sensitised children in non-smoking households, further research is required to clarify this.

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Preventing exacerbations by avoiding mite allergen

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CONFLICTS OF INTEREST DISCLOSURES

Dr Murray has received grants from NIHR, JP Moulton Charitable Foundation and from North West Lung Research Centre Charity. She has received lecture fees from GSK and Novartis and travel grants from Novartis. Professor Custovic has received grants from Medical Research Council, JP Moulton Charitable Foundation and from North West Lung Research Centre Charity. He receives personal fees from AstraZeneca, Novartis, ThermoFisher and Regeneron / Sanofi. Professor Simpson has received grants from Medical Research Council, NIHR and EU FP7. She has received lecture fees from GSK, Chiesi and Thermofisher Scientific. Philip Foden, Helen Sumner and Elizabeth Shepley have no conflicts of interest.

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Preventing exacerbations by avoiding mite allergen

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Preventing exacerbations by avoiding mite allergen

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FIGURE LEGENDS

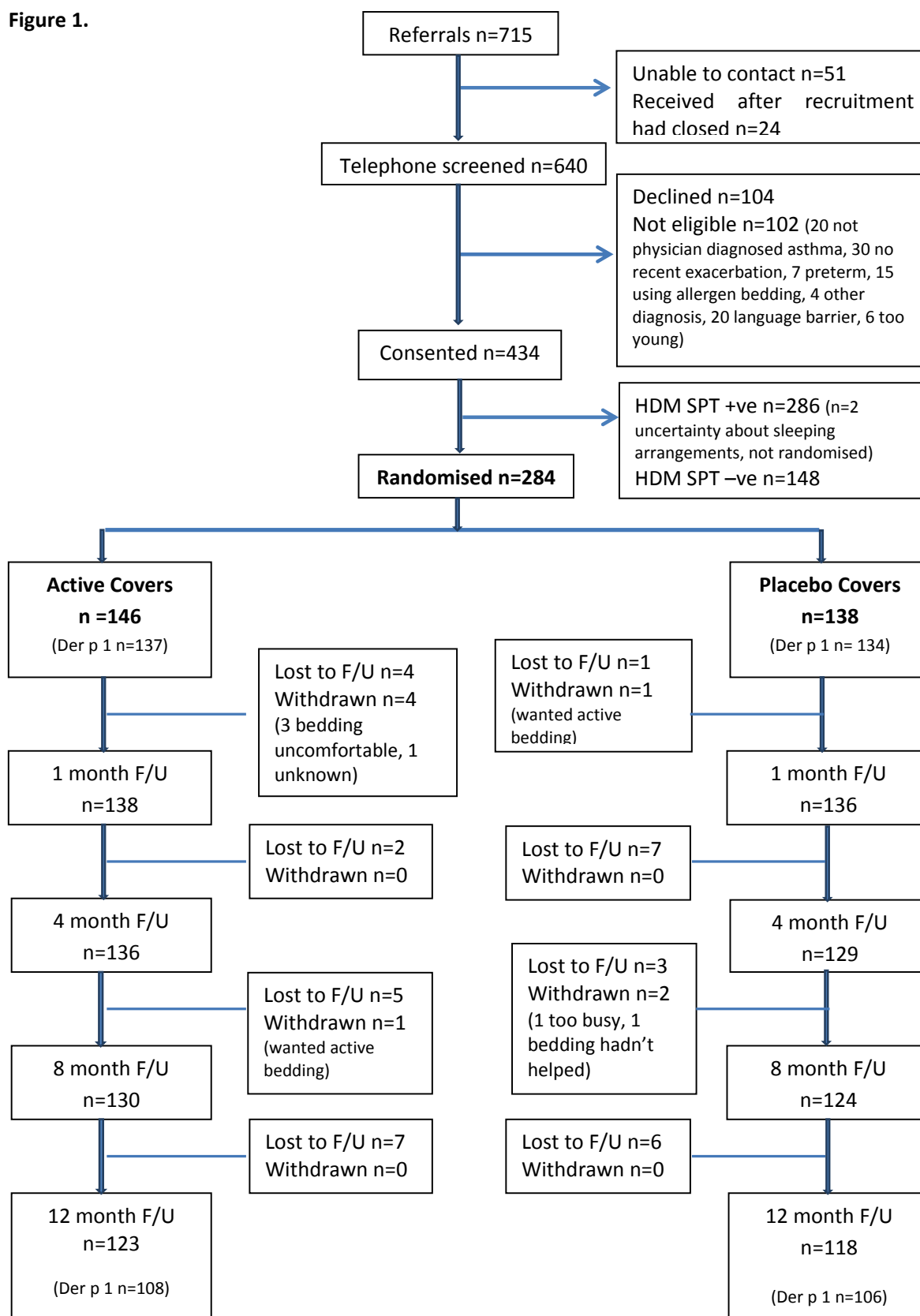
Figure 1. CONSORT (Consolidated Standards of Reporting Trials) flow diagram showing the participants' course during the study.

Figure 2. Proportion of children who suffered one or more severe exacerbation during the 12 month follow up period (for all children who completed 12 months Follow up, n=241). Results are shown for (A) one or more hospitalizations or ED visit requiring systemic corticosteroids because of an asthma exacerbation ($p=0.047$) and (B) the use of systemic corticosteroids for at least 3 days because of an asthma exacerbation ($p=0.85$).

Figure 3. Time to first hospitalizations or ED visit because of severe exacerbation of asthma. The model was adjusted for age, gender, ethnic group, maintenance asthma treatment, number of hospitalisations in the 12-month period prior to randomisation, index of multiple deprivation and tobacco smoke exposure. The risk was 45% lower in the Active compared with the placebo group ($p=0.006$).

Figure 4. Der p 1 levels in child's mattress (ng/m^2) at recruitment and 12 months after intervention. Results are shown as geometric mean and 95% confidence interval for Active covers (mite-impermeable) (dashed line) and Placebo covers (solid line).

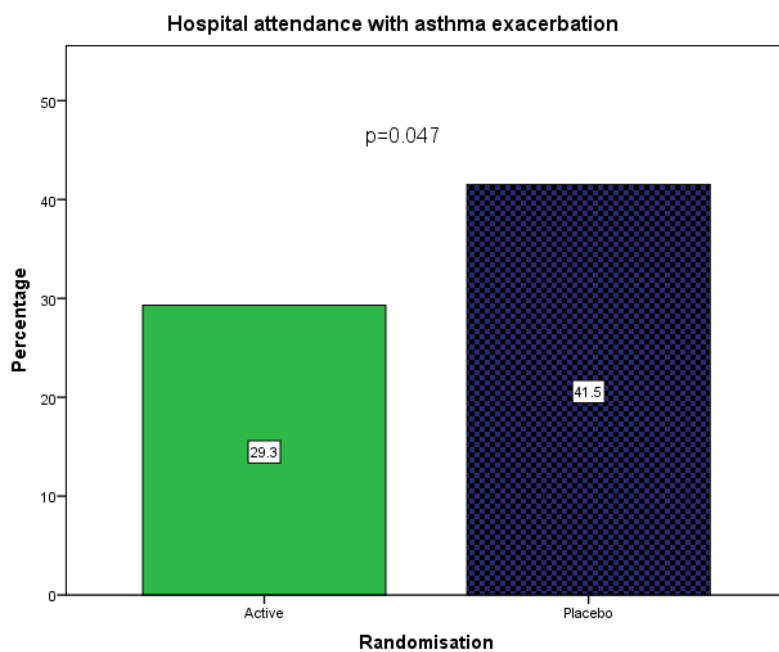
Figure 1.



Preventing exacerbations by avoiding mite allergen

Figure 2

A.



B.

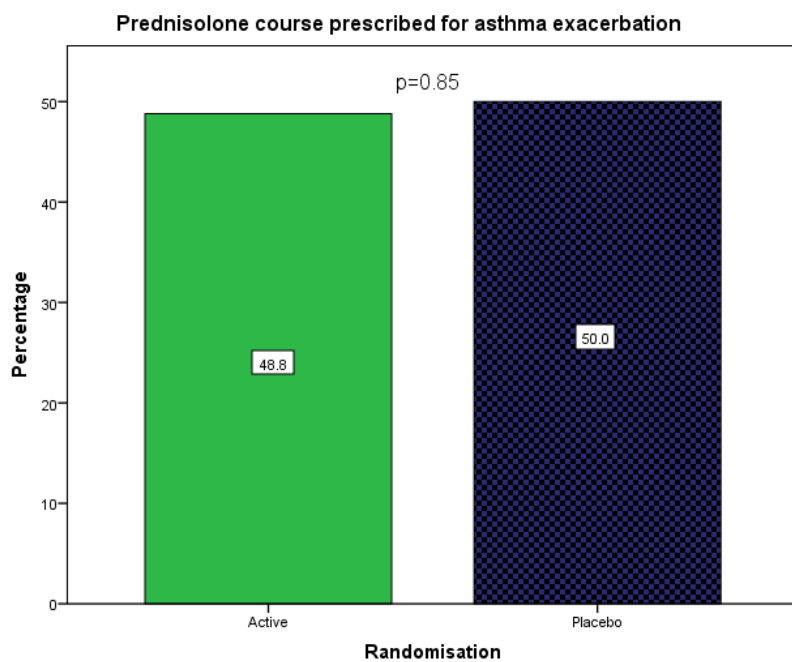
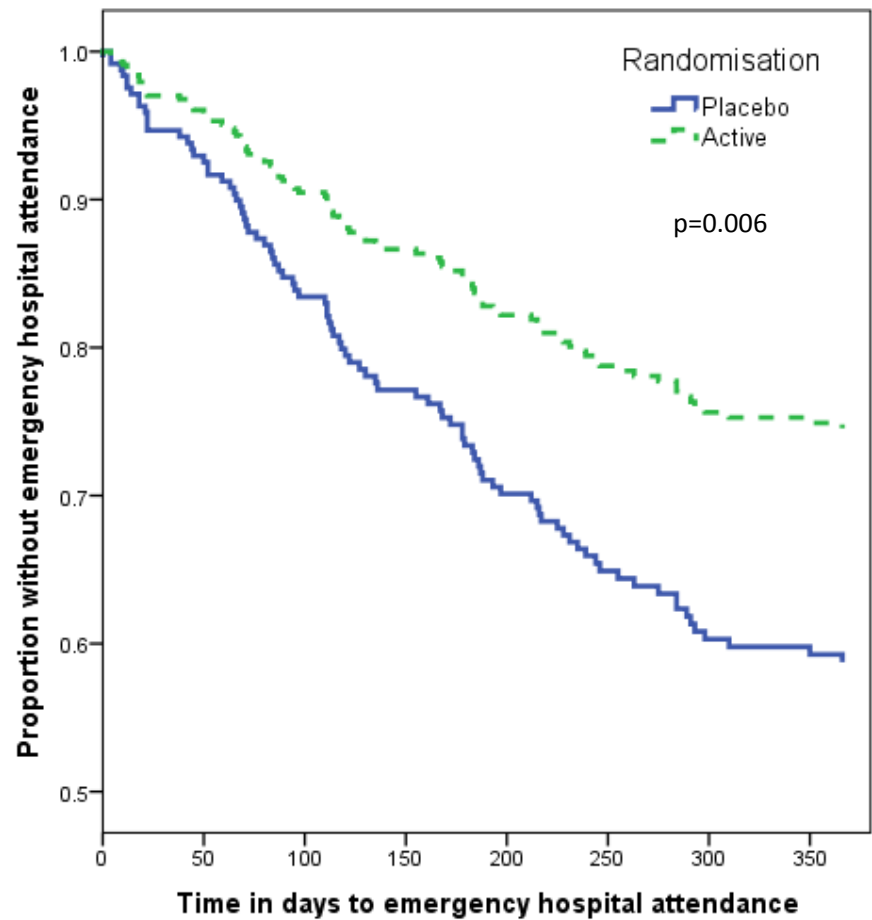
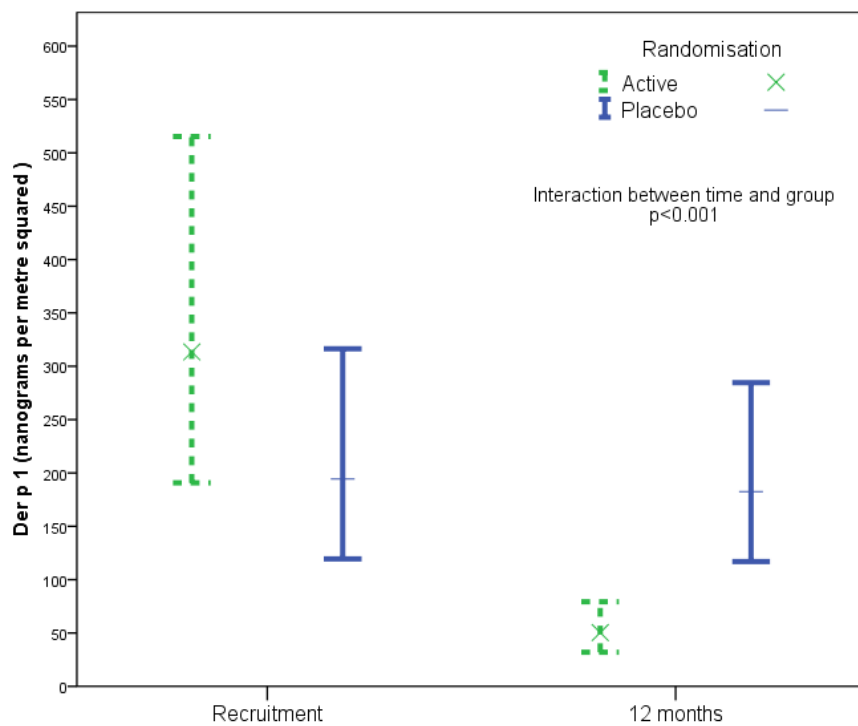


Figure 3.



Number at risk								
Placebo	136	119	105	94	89	77	71	70
Active	136	129	121	112	102	91	88	88

Figure 4.



Preventing exacerbations by avoiding mite allergen

Table 1. Characteristics of study participants at randomisation.

*All children were skin test positive to house dust mite, but not all children completed skin test to other allergens. **Ascertained based on SPT or on symptom reports from parents and pet ownership/exposure.

	Placebo covers	Mite-impermeable Covers (Active)	P value
	N=138	N=146	
Age (mean; SD)	7.45 (3.55)	7.11 (3.49)	0.42
Age 3-10 yrs	106 (76.8%)	117 (80.1%)	0.50
Age 11-17 yrs	32 (23.2%)	29 (19.9%)	
Gender; male	94 (68.1%)	93 (63.7%)	0.43
Ethnicity		N=143	0.96
White	89 (64.5%)	91 (63.6%)	
Asian	35 (25.4%)	36 (25.2%)	
Other	14 (10.1%)	16 (11.2%)	
Current hay fever	41/134 (30.6%)	46/129 (35.7%)	0.38
Current eczema	71 (51.8%)	57/140 (40.7%)	0.07
Food allergy	26/130 (20.0%)	40/138 (29.0%)	0.09
Maternal asthma	43 (31.2%)	39/142 (27.5%)	0.50
Paternal asthma	30/134 (22.4%)	40/142 (28.2%)	0.27
Maternal smoking	35 (25.4%)	34/145 (23.4%)	0.71
Paternal smoking	31/133 (23.3%)	43/141 (30.5%)	0.18
Smoking by a household member	57 (41.3%)	67 (45.9%)	0.44
Deprivation index (mean; SD)	34.16 (19.34)	34.74 (17.32)	0.79
Sensitized to*			
Mite	138/138 (100%)	146/146 (100%)	

Preventing exacerbations by avoiding mite allergen

Table 1. Characteristics of study participants at randomisation (“Continued”)

Mite only	50/125 (40%)	60/130 (46.1%)	0.28
Cat	46/125 (36.8%)	46/130 (35.4%)	0.81
Dog	45/125 (36.0%)	44/130 (33.8%)	0.72
Grass	49/129 (38.0%)	46/136 (33.8%)	0.48
Aspergillus	8/126 (6.3%)	3/136 (2.2%)	0.09
Tree pollen	7/125 (5.6%)	4/135 (3.0%)	0.29
Number of allergens sensitised to excluding HDM (median; IQR)	N=131 1 (0-2)	N=135 1 (0-2)	0.55
Pet contact	58/137 (42.3%)	64/145 (44.1%)	0.76
Cat owner	22/137 (16.1%)	21/145 (14.5%)	0.71
Dog owner	31/137 (22.6%)	36/145 (24.8%)	0.66
Sensitised and exposed to pet**	29 (21.0%)	31 (21.2%)	0.96
GINA Step			0.98
GINA step 1-2	72 (52.2%)	76 (52.1%)	
GINA step ≥ 3	66 (47.8%)	70 (47.9%)	