Small-group, interactive education and the effect on asthma control by children and their families

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Abstract

Background: Effective approaches to education about asthma need to be identified. We evaluated the impact on asthma control by children and their caregivers of an intervention involving small-group, interactive education about asthma.

Methods: We randomly assigned children who visited an emergency department for an exacerbation of asthma (n = 398) to either of 2 groups. Children assigned to the control group followed the usual care recommended by their primary care physician. Those assigned to the intervention group participated in a small-group, interactive program of education about asthma. We examined changes in the number of visits to the emergency department during the year after the intervention.

Results: During the year after enrolment, children in the intervention group made significantly fewer visits to the emergency department (0.45 visits per child) compared with those in the control group (0.75 visits per child) (p = 0.004). The likelihood of a child in the intervention group requiring emergency care was reduced by 38% (relative risk [RR] 0.62, 95% confidence interval CI 0.48–0.81, p = 0.004). Fewer courses of oral corticosteroids (0.63 per child) were required by children in the intervention group than by those in the control group (0.85 per child) (p = 0.006). We observed significant improvements in the symptom domain of the questionnaire on pediatric asthma quality-of-life (p = 0.03) and the activity domain of the questionnaire on caregivers’ quality of life (p = 0.05). Parents of children in the intervention group missed less work because of their child’s asthma after participating in the educational program (p = 0.04). No impact on hospital admissions was observed.

Interpretation: Education about asthma, especially in a small-group, interactive format, improved clinically important outcomes and overall care of children with asthma.

Management of asthma often focuses on “crisis intervention,” meaning that the disease is addressed only when a problem occurs. Shifting emphasis to a preventive health model, which includes guided self-management, has been shown to reduce costs related to hospital admissions and visits to emergency departments. Education about asthma is an integral part of the recommendations of most guidelines for the management of asthma.

There is no consensus on the best model for education about asthma. A review of publications from 1991 through 2004 showed that only half of the studies identified were randomized controlled trials and that many studies used patients as their own historic controls. Given that a regression to the mean is commonly observed in the data of such studies, with improvements in asthma over time, the use of historic controls is not an acceptable means of defining the impact of education. The studies reviewed also varied greatly in the demographic characteristics of the participants, the professional qualifications of the educators used, the nature of the interventions, the outcomes measured, the time frames for measurement and the inclusion of medical care and other services. These variations make it difficult to compare results and make firm recommendations.

Programs of education about asthma are typically directed toward the learner and conducted using either a one-on-one, large-group or small-group format. The intervention may consist of self-directed educational material, lectures (i.e., a teacher-focused format) or group interaction (i.e., a learner-focused format). Effective change in behaviour occurs when learners actively interact with the content to be learned, with the teacher and with each other. Small groups of fewer than 10 members allow for an ideal level of interaction. Westberg and Jason cite several compelling reasons for using small groups to promote learning. Learners are more likely to take ownership of their education and may be more engaged with.

From the Children’s Asthma Education Centre (Watson, Gillespie, Thomas, Filuk, McColm, Piwniuk, Becker), Children’s Hospital of Winnipeg, Winnipeg, Man.; the Division of Allergy (Watson), Dalhousie University, IWK Health Centre, Halifax, NS; and the Section of Allergy and Clinical Immunology, Department of Pediatrics and Child Health (Becker), University of Manitoba, Winnipeg, Man.

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the material. They can learn from each other in a supportive, nonjudgmental environment. They can both give and receive peer-oriented feedback. They can practise skills that can be applied later in real-life situations. Learners retain and transfer knowledge more effectively when they are able to practise what they have learned. For children with asthma and their families, such a model would help facilitate memory retention and a higher comfort level with future decisions related to management of asthma.

We conducted a randomized controlled trial to evaluate the impact on asthma control of an intervention involving small-group, interactive education of children with asthma and their caregivers.

**Methods**

**Participants**

We conducted the study at the Children’s Hospital of Winnipeg, located in a moderate-sized urban setting in the Canadian prairie province of Manitoba. The hospital’s emergency department had 40,000 visits per year, 3500 of which were due to acute asthma. We used emergency department records to identify children who had a diagnosis of asthma. We used letter and telephone-based follow-up to contact families of the children to invite them to participate in the study. Children were eligible if they were aged 3–16 years, had physician-diagnosed asthma and had required a visit to the emergency department for acute asthma during the recruitment phase. We excluded children who had other serious chronic illnesses, including respiratory illnesses other than asthma. We also excluded families for whom a significant language barrier existed. We did not offer incentives for participation. During the recruitment phase, we performed screening by contacting families by telephone after their visits to the emergency department. We closed recruitment after we reached our target for enrolment \( n = 400 \). We randomly assigned each child to either the control group or intervention group on their first visit to the study site.

**Study protocol**

We assessed participating children and their families at a hospital-based site called the Children’s Asthma Education Centre. We collected the following data: a focused asthma history, height and weight, medications used, responses to the Pediatric Asthma Quality of Life Questionnaire for children aged 7 years or older, responses to the Pediatric Asthma Caregiver’s Quality of Life Questionnaire, the Classroom Productivity and Caregiver Work Productivity questionnaires related to asthma (adapted from an existing questionnaire on allergy-related impairment), and number of visits made to the emergency department during the year before enrolment. We performed spirometry for children aged 7 years and older using the Vitalograph-Compact Spirometer (Vitalograph Ltd., Buckingham, United Kingdom).

We randomly assigned families to either of 2 groups. Children assigned to the control group followed the usual care recommended by their primary care physician and were given a booklet with basic information on asthma (Appendix 1, available at www.cmaj.ca/cgi/content/full/cmaj.080947/DC2). The booklet had been developed previously for children admitted to hospital for an exacerbation of asthma. Children assigned to the intervention group participated in our program of education about asthma along with their usual care. At the time of our study, usual care did not include one-on-one education on asthma by an educator. We randomized the assignment of families based on a computer-generated allocation in blocks of 10. Because of the nature of the study, blinding was not possible. We assessed participants in both study groups at 12 months after enrolment using the same tools as in our initial assessment.

Our study was approved by the Faculty Committee on the Use of Human Subjects in Research, based at the University of Manitoba. Funding was provided by Manitoba Health, the Winnipeg Children’s Hospital Foundation and unrestricted educational grants from AstraZeneca, Merck Frosst and GlaxoSmithKline.

**Educational program**

The Children’s Asthma Education Centre developed a 4-week educational program using materials containing core content and supplemental resources that were based on recommendations from the Canadian Asthma Consensus Statement on education about asthma. These materials included a program-specific booklet for families enrolled in the educational program (Appendix 2, available at www.cmaj.ca/cgi/content/full/cmaj.080947/DC2) and personalized mailings to reinforce the program’s key messages (Appendix 3, available at www.cmaj.ca/cgi/content/full/cmaj.080947/DC2). Pamphlets on asthma-related topics, which were produced by the centre for a reader comprehension level of grade 8 or below, were also used.

The educational program was targeted to either the parent or the child, or both, depending on the age of the child, with core content adapted for age level as appropriate. The educational content was the same for each group. For children aged 3–6 years, parents were the participants in the intervention. For children aged 7–11 years, both parents and children participated in the education sessions, but did so in separate groups. For adolescents (aged 12–16 years), only the adolescent attended the program (Figure 1). The educational program was designed for a group of 6–8 participant families or adolescents and addressed different topics of importance in the management of asthma. The program was delivered by a nurse-educator experienced in asthma management and a respiratory therapist who were among the first nationally certified asthma educators in Canada. Groups met for about 1.5 hours per week over 4 weeks to discuss 4 themed topics (Appendix 4, available at www.cmaj.ca/cgi/content/full/cmaj.080947/DC2).

An important part of the program was the promotion of small-group interaction for children and their families. Parents actively discussed their successes and failures in the management of their child’s asthma. Children and their families learned from each other as well as from the educator. The educator acted as a resource and guide for the discussion. In addition, families were sent personalized mailings at 2, 4, 6 and 12 months after enrolment that reinforced key educational messages using the program’s main topics (Appendix 3, available at www.cmaj.ca/cgi/content/full/cmaj.080947/DC2).
Outcome measures
The primary outcome measure was change in the number of visits to the emergency department during the year after the intervention. Secondary outcome measures included the impact of asthma-related education on the number of admissions to hospital, quality of life, use of oral corticosteroid therapy for exacerbations of asthma, productivity in the workplace and pulmonary function. We calculated a sample size of 400 (200 in each group) to provide a power of 80% with an \( \alpha \) of 0.05 or less.

Statistical analysis
We performed an intention-to-treat analysis for number of emergency department visits. We analyzed results using repeated-measures analysis of variance, including the primary outcome measure. In addition, we performed a Poisson regression analysis to determine relative risks (RRs) for visits to the emergency department, hospital admissions and use of oral corticosteroids. For secondary outcomes, we used \( t \) tests, Fisher exact test and \( \chi^2 \) analysis. We considered a \( p \) value of 0.05 or less to be significant. Our primary hypothesis was that our intervention would result in a 25% reduction in visits to the emergency department by participants in the intervention group.

Results
Participants
Over a 30-month period, we identified 2901 eligible children from emergency department records. The recruitment and flow of participants through the study is shown in Figure 2. We sent letters to all of the families of eligible children to invite them to participate and followed up by telephone. Of the 1120 families who responded, 41% declined to participate, 11% were unsure whether their child had asthma and 11% expressed interest in education but could not commit to it at that time. Of the 415 families who agreed to participate, 17 did not arrive for the initial assessment and 398 were enrolled. We carried out enrolment from June 1997 to January 2000.

The demographic characteristics of participants are shown in Table 1. In both the intervention and control groups, most children had made 1 visit to the emergency department in the year before enrolment (77% in each group). The groups also had similar proportions of participants who had made 2 visits (15% v. 12%) and who had made 3 or more visits (6.5% v. 5.9%). Use of inhaled corticosteroids, either intermittently or regularly, was not statistically different between the 2 groups. The distribution across socio-economic levels among the families in our study concorded generally with that of families seen in our emergency department. Ethnic background (accorded by self-identified maternal heritage) was similar in both groups. The groups also had similar proportions of live-in smokers and of families that allowed smoking inside the home. No significant differences existed in number of visits made to the emergency department, number of hospital admissions, pulmonary function or oral corticosteroid use during the year before enrolment. The level of attendance at the program’s classes was 65% in the preschooler group, 80% in the school-aged group and 79% in the adolescent group.

Primary outcome
The number of visits to the emergency department decreased in both the control and intervention groups (Table 2), but we observed a significantly greater reduction in the intervention group (\( p = 0.0037 \)). Compared with the control group, the intervention group had a lower likelihood of requiring emergency care. (RR 0.62, 95% confidence interval [CI] 0.48–0.81, \( p = 0.0004 \)).
Secondary outcomes
The number of admissions to hospital was low in both groups. Each group showed a significant drop in admissions but we observed no significant difference between the groups (Table 2). Compared with the control group, the intervention group used fewer courses of oral corticosteroid therapy per patient during the year after enrolment (p = 0.006). The likelihood of a child in the intervention group requiring oral corticosteroids was reduced by 36% (RR 0.64, 95% CI 0.50–0.82, p = 0.001).

For children who were able to undergo spirometry testing (control n = 79, intervention n = 70 at study completion), we analyzed results using percent-predicted forced expiratory volume in 1 second (FEV1). We found no difference in the percent-predicted FEV1 between groups. Asthma education had a significant impact on workplace productivity for primary caregivers who worked outside of the home (control n = 79, intervention n = 65). Primary caregivers in the intervention group missed significantly less work than those in the control group in the month before the final assessment at the end of the follow-up period (p = 0.04).

In both the control and intervention groups, we observed significant improvement in total scores on the Pediatric Asthma Quality of Life Questionnaire during the year after enrolment. In the symptom domain, a clinically important statistical difference (a change of 0.5 unit per question) was observed in favour of the intervention group (p = 0.03). Caregivers in both groups showed significant improvement in scores on the Pediatric Asthma Caregiver’s Quality of Life Questionnaire during the year after enrolment. This improvement was greater in the intervention group, but not significantly so (p = 0.10). In the domain of activities, we observed both a statistical (p = 0.05) and clinically important difference in the intervention group compared with the control group.

At completion of the study, 35% of the homes of participants had smokers living in them, with no significant difference between the intervention and control groups. Among families in the control group, 30% continued to allow smoking in the home, as compared with 18% of those in the intervention group (p = 0.019).

Interpretation
The purpose of education about asthma provided by the Children’s Asthma Education Centre significantly reduced the need for hospital-based emergency services among participating children with
asthma. Asthma-specific quality of life improved in the domains of the children’s assessments of their symptoms and in caregivers’ scores. Primary caregivers of participating children lost fewer workplace hours. This reduction implied that the asthma was under better control. It was consistent with studies showing that the number of workdays lost by caregivers was correlated highly with level of control of asthma.24 Fewer courses of oral corticosteroid therapy were required by participants in the control group made significantly fewer visits to the emergency department for acute asthma during the year after enrolment. Nevertheless, we observed a significantly greater reduction in visits among children in the intervention group.

Small-group interaction was the major learning tool of our educational intervention. This tool provided parents and children with opportunities to share their experiences with asthma, solve problems together and provide support to one another while allowing role-modelling to occur. From a theoretic perspective, small-group interaction is a highly desirable tool for enhancing education.36–38 Other investigators have observed a reduction in the severity of asthma only when knowledge was imparted in an interactive setting.25 Interactive learning may also increase the confidence of families in managing asthma, although we recognize that this method of learning will not be suitable for all people or all situations.

Individual studies undertaken through 2004 have shown that education about asthma improves knowledge about asthma,29–33 although it does not guarantee an improvement in other outcomes. Some studies have suggested a relation between educational intervention and improvement in attitudes or self-management skills.3,15,34–39 Educational interventions may be connected to improvements in quality of life40–42 and peak expiratory flow rates.41,43

The most important outcomes of providing education about asthma involve changes in the use of health care services. Post-education outcomes that have been identified by researchers include reductions in the numbers of visits to emergency departments, admissions to hospitals and unscheduled visits to physicians, and in costs related to health care.5,10,16,44–47 From 2005 onward, an increasing number of randomized control trials have been performed.48–55 Some have focused on schools with children as the target learners50–52 or included a multifaceted approach with education as one component.39,43,54 The results generally have been positive, showing increases in knowledge, fewer symptoms and fewer absences from school. Although studies that involved multifaceted programs showed fewer visits to emergency departments, it is difficult to conclude with certainty that such outcomes are attributable solely to the educational component in these studies.55,54 Our program remains unique in that it incorporates the principles of small-group learning for children or for both children and their parents and includes only an educational intervention.

**Limitations and strengths**

Our study has limitations. It did not have sufficient power performed.56–55 Some have focused on schools with children as the target learners50–52 or included a multifaceted approach with education as one component.39,43,54 The results generally have been positive, showing increases in knowledge, fewer symptoms and fewer absences from school. Although studies that involved multifaceted programs showed fewer visits to emergency departments, it is difficult to conclude with certainty that such outcomes are attributable solely to the educational component in these studies.55,54 Our program remains unique in that it incorporates the principles of small-group learning for children or for both children and their parents and includes only an educational intervention.

**Table 1:** Baseline characteristics of participants in a randomized controlled trial of a small-group, interactive education program about asthma

| Characteristic                  | Intervention n = 201 | Control n = 197 |
|--------------------------------|----------------------|-----------------|
| Age, yr, mean (SD)             | 7.4 (3.4)            | 7.4 (3.3)       |
| Sex, ratio of boys to girls    | 1.75:1               | 2.34:1          |
| Height, cm, mean (SD)          | 127.8 (20.2)         | 121.8 (20.6)    |
| Weight, kg, mean (SD)          | 30.9 (16.6)          | 28.3 (14.1)     |
| Ethnic background              |                      |                 |
| White                          | 68                   | 67              |
| Aboriginal                     | 6                    | 9               |
| Other (Asian, South Asian)     | 26                   | 24              |
| Socio-economic quintile        |                      |                 |
| 1 (lowest income)              | 18                   | 25              |
| 2                              | 33                   | 19              |
| 3                              | 15                   | 20              |
| 4                              | 19                   | 18              |
| 5 (highest income)             | 14                   | 18              |
| Homes with smokers             | 46                   | 40              |
| Homes allowing visitors to smoke| 29                 | 31              |
| Visits to emergency department in previous year, per child, mean (SEM) | 1.37 (0.07) | 1.34 (0.07) |
| Hospitalizations in previous year, per child, mean (SEM) | 0.21 (0.02) | 0.18 (0.02) |
| FEV1,% predicted,† mean (SEM)  | 87 (1.04)            | 88 (0.97)       |
| Using inhaled β-agonist at enrolment | 99.0     | 98.8           |
| Using inhaled corticosteroid at enrolment | 91.2     | 84.0           |
| Oral corticosteroid use in previous year, courses per child, mean (SEM) | 1.53 (0.08) | 1.33 (0.07) |
| Workplace productivity of primary caregiver,§ mean (SEM) | 18.7 (2.4) | 12.9 (2.2) |

Note: FEV1 = forced expiratory volume in 1 second, SD = standard deviation, SEM = standard error of mean.
*Unless stated otherwise.
†According to participant-reported maternal height.
‡Base on age and height. Control n = 94, intervention n = 99.
§Calculated based on percentage of total work hours missed in the month before assessment. Control n = 80, intervention n = 65.
to assess the differences in outcomes between different age groups. We encountered difficulties in recruiting participants. Only 37% of the families with whom we had direct contact were enrolled. We may not have reached the population most in need of education, despite the representation of a broad variety of different socio-economic backgrounds among the families of enrolled participants. Families who participated in our program were highly motivated. It would be difficult to assess the impact of our intervention in a less motivated group. Whereas this difficulty is true for any educational program, we uniquely used interactive strategies to motivate families to become active participants in the management of their children’s asthma. For families to participate in the interactive classes, time and commitment were required of them by our program. Although we believe that our program consisted of “need to know” information only, asking families in today’s hectic world to attend 4 classes (1.5 hours weekly for 4 weeks) may be asking too much. Further investigation is needed to assess the impact of a modified and shorter program.

We believe that the most important feature of our educational program was the interaction that it fostered between participants in their respective small-group settings. In the written feedback we received, parents and children reported that they felt less isolated when dealing with asthma and believed that they benefited from the group-oriented experience. Having other parents share their experiences may be a powerful learning tool. It may have a greater positive impact on strategies for the management of asthma than passively listening to someone teaching the same material. Opportunities to question, problem-solve and engage in activities with others who share similar challenges may serve to promote empowerment. These hypotheses will need to be tested using qualitative analysis.

**Conclusions**

Our small-group, interactive educational program for children with asthma was found to be effective in significantly reducing the need for hospital-based emergency services. During the year after enrolment, children of families who participated in the educational program required fewer courses of oral corticosteroid therapy and their primary caregivers lost fewer workplace hours. Quality of life for the children and their families improved. Education about asthma, especially in a small-group, interactive format, is an important aspect of overall care for children with asthma.

This article has been peer reviewed.

**Competing interests:** None declared.

**Contributors:** Wade Watson was the primary investigator of the study. Wade Watson, Cathy Gillespie, Michelle Piwniuk and Allan Becker were involved in the conception and design of the study protocol. Nicola Thomas, Shauna Filuk and Judy McColm

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**Table 2: Primary and secondary outcomes in the year after enrolment among participants in a randomized controlled trial of a small-group, interactive education program about asthma**

| Outcome                          | Trial group | RR (95% CI) | p value |
|---------------------------------|-------------|-------------|---------|
| **Primary**                     |             |             |         |
| Visits to emergency department, per child* | n = 190     | n = 190     |         |
| Mean (SEM)                      | 0.45 (0.07) | 0.75 (0.07) | 0.62 (0.48–0.81) | 0.004† |
| Change from baseline            | −0.92       | −0.59       |         |
| **Secondary**                   |             |             |         |
| Admissions to hospital, per child* | n = 190     | n = 190     |         |
| Mean (SEM)                      | 0.05 (0.02) | 0.06 (0.02) | 0.90 (0.38–2.1) | 0.80‡ |
| Change from baseline            | −0.16       | −0.12       |         |
| Oral corticosteroid use, courses per child | n = 129   | n = 170     |         |
| Mean (SEM)                      | 0.63 (0.08) | 0.85 (0.07) | 0.64 (0.50–0.82) | < 0.001† |
| Change from baseline            | −0.90       | −0.48       |         |
| FEV₁, % predicted               |             |             |         |
| Mean (SEM)                      | 90 (1.02)   | 88 (0.97)   | 0.36§   |
| Change from baseline            | 2           | 0           |         |
| Workplace productivity of caregiver† | n = 65     | n = 79      |         |
| Mean (SEM)                      | 0.8 (2.4)   | 5.0 (2.3)   | 0.04§   |
| Change from baseline            | −17.9       | −7.9        |         |
| Homes with smokers              |             |             |         |
| %                               | 34.6        | 34.5        | 0.99**  |
| Change from baseline            | −11.2       | −5.7        |         |
| Homes allowing visitors to smoke|             |             |         |
| %                               | 18          | 30          | 0.019** |
| Change from baseline            | −10.8       | −1.1        |         |

Note: FEV₁ = forced expiratory volume in 1 second, RR = relative risk, SEM = standard error of mean.

*Results were based on an intention-to-treat analysis.
†Calculated based on percentage of total work hours missed during the month before assessment.
‡Results were based on Poisson regression analysis.
§Results were based on repeated measures analysis of variance.
**Results were based on χ² test.
were involved in the acquisition of data. All of the authors were involved in the analysis and interpretation of the data. Wade Watson, Cathy Gillespie, Michelle Piwniuk and Allan Becker were involved in the drafting of the manuscript. All of the authors were involved in the critical revision of the manuscript and approved the final version submitted for publication.

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