Asthma is one of the major health problems in industrialized countries. Although new drugs and evidence-based guidelines have been developed, there has been no major change in asthma morbidity and mortality. It is still important to determine whether pharmaceutical care (PC) influences health outcomes.

**OBJECTIVE:** To evaluate the effectiveness of PC with regard to clinical, humanistic, and economic outcomes in adults with asthma.

**METHODS:** An intervention study was conducted over 12 months. At baseline, 39 community/retail pharmacies, 84 primary care physicians (general practitioners, internal specialists, chest physicians), and 183 patients (aged 18–65 y) diagnosed with asthma were included. To evaluate economic outcomes, 2 German statutory health insurance funds provided 2 years of claims data for their insured patients (n = 55). A 1:10 matching was carried out to compare the data of this intervention subgroup with those of a control group (n = 550).

**RESULTS:** Significant improvements were found for all humanistic outcomes (eg, asthma-specific quality of life, self-efficacy, knowledge, medication adherence). In addition, asthma severity, self-reported symptoms, peak expiratory flow, and patients’ inhalation technique improved. Increases in forced expiratory volume in 1 second and vital capacity were not significant over time. Evaluation of the insurance claims data revealed a shift toward better adherence to evidence-based therapy.

**CONCLUSIONS:** The study shows that PC for people with asthma has a positive impact on humanistic and, to some extent, on clinical outcomes. To determine potential economic benefits, future research should focus on patients with more severe asthma.

**KEY WORDS:** asthma, community pharmacy, pharmaceutical care, pharmacy practice.


**Background:** Despite significant progress in asthma drug therapy in recent years, there has been no major change in asthma morbidity and mortality. It is still important to determine whether pharmaceutical care (PC) influences health outcomes.

**Objective:** To evaluate the effectiveness of PC with regard to clinical, humanistic, and economic outcomes in adults with asthma.

**Methods:** An intervention study was conducted over 12 months. At baseline, 39 community/retail pharmacies, 84 primary care physicians (general practitioners, internal specialists, chest physicians), and 183 patients (aged 18–65 y) diagnosed with asthma were included. To evaluate economic outcomes, 2 German statutory health insurance funds provided 2 years of claims data for their insured patients (n = 55). A 1:10 matching was carried out to compare the data of this intervention subgroup with those of a control group (n = 550).

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**Key Words:** asthma, community pharmacy, pharmaceutical care, pharmacy practice.
Based on intensified cooperation between local community pharmacists, general practitioners (GPs), and other physicians, their regulatory bodies, and GKV, we investigated the effectiveness of PC with regard to clinical, humanistic, and economic outcomes in adults with asthma. This was a pivotal study to evaluate the contributions of community pharmacies in disease management program and/or integrated care contracts with regard to outcomes. In this context, we also examined whether drug therapy shifted toward better adherence with evidence-based guidelines or a reduction of hospitalizations and absence from work, which could have positive cost implications for health insurance funds.

Methods

STUDY POPULATION

All of the 148 community/retail pharmacies in the region of Trier (part of the State of Rhineland–Palatino, population 512,000) were invited by mail to enter the study. The invitation letter was signed both by the president of the chamber of pharmacists in the State of Rhineland–Palatino and the principal investigator of the study (MS). Fifty-seven pharmacies agreed initially to participate and offer PC to their asthma patients for one year. In preparation for the study, at least one full-time pharmacist at each participating pharmacy was trained to provide asthma services. Thirteen hours of training was based on a nationally certified curriculum and a published manual/protocol and comprised the following: medical, pharmaceutic, and pharmacologic knowledge; communication skills; and the use of the study protocol and PC documentation forms.

In addition to the initial training provided, all pharmacies were monitored by a pharmacist based in the city of Trier and employed for this study. This pharmacist visited all practice sites regularly to check for compliance with the study protocol and the documentation forms for PC, minimize missing data, and enhance the documentation of drug-related problems detected and solved. In addition, counseling on-site and via phone/fax was offered from the first day of recruitment until the end of the study. Thus, assistance from the distant research center was limited to supervision of the support pharmacist.

We paid a maximum of €75 (about $100 US) per patient to pharmacists and physicians when data were provided at baseline and after 6 and 12 months. To be eligible for the study, patients had to be 18–65 years old, have a physician’s diagnosis of asthma (confirmed for the study), and give written informed consent. Recruitment was carried out by the participating pharmacists, as well as by the patients’ physicians (84 GPs, internal specialists, respiratory physicians). A list of participating community pharmacies was provided to all physicians’ offices. Eligible patients were asked during a regular visit to the pharmacy or physician whether they were interested in joining the program. In addition, an advertisement in the local newspaper helped with recruiting patients. The enrollment time period lasted from October 2001 until March 2002.

DATA COLLECTION AND INTERVENTIONS

In Germany, this type of research does not require an institutional review board approval. However, the work was conducted in compliance with the requirements of the data protection agencies of the health insurance funds involved.

Except for explicitly assessing readiness for change, all elements of PC, as described by Strand et al. and highlighted recently by McLean and MacKeigan, were included in our intervention. In cooperation with the patients’ physicians, 5 meetings between pharmacists and patients were scheduled over 12 months. These one-to-one counseling sessions took place in confidential areas or counseling rooms available within the pharmacy.

Patients were educated in asthma pathology, the use of asthma medication, inhalation technique, and self-management skills. Drug-related problems were detected and solved. At the beginning of the study, each patient was instructed to use a peak flow meter (Mini-Wright, Clement Clarke Int. Ltd., Essex, UK) and a well-recognized asthma diary (Atemwegsliga, Germany) twice daily.

At baseline and after 6 and 12 months, 4 questionnaires were administered to patients during an appointment in the pharmacy. The questionnaires comprised the following areas: disease-specific quality of life, generic self-efficacy, asthma knowledge, and adherence. All questionnaires were completed by the patients without the help of the pharmacist. At the same time, patients had to visit their physician to have their lung function (forced expiratory volume in 1 second [FEV1] and vital capacity [VC]) tested and for reassessment of asthma and dyspnea severity.

In addition to these study data, 2 German statutory health insurance funds (Allgemeine Ortskrankenkasse [AOK] in Rhineland–Palatino, and Barmer Ersatzkasse) provided 2 years of claims data for their insured patients (n = 55, 17 male) for the one-year period before the study and the study year. These patients signed a specific consent form to allow disclosure of their records. Statutory health insurance data about patients’ asthma-related hospital admissions (International Classification of Diseases [ICD] J 45 and J 46), absence from work, and drug consumption were evaluated in this subgroup. Control patients (n = 550) were identified by asthma diagnosis (ICD) and prescriptions of anti-asthmatics (Anatomical Therapeutic Chemical [ATC] code: R03) in a sample of the AOK in the Federal State of Hesse. Patients diagnosed with COPD were excluded. Control and intervention patients were matched by a ratio of 10:1 according to gender, age (± 5 y), date of recruitment, and amount (defined daily doses) of prescribed anti-asthmatics (ATC R03).

OUTCOME MEASURES

To monitor lung function, FEV1 and VC were measured by patients’ physicians at baseline and after 6 and 12 months. In case of insufficient, incorrect, or incomplete data, physicians were asked to provide flow volume curves. These data were reassessed independently by 2 experienced chest physicians. In addition, patients were asked to measure peak expiratory flow rates twice daily at home during the entire study year and on consecutive dates in the pharmacy. The peak flow measures under pharmacists’ supervision were recorded in the monitoring plan. To assess dyspnea severity, physicians used the Medical Research Council Dyspnoea Scale (Medical Research Centre of Great Britain, none = 0 to severe = 4). Asthma severity was classified according to German Asthma Guidelines (from intermittent = 1 to severe, persistent = 4).

A 7-point checklist was used to score patients’ inhalation technique. For each correct step, 1 point was assigned, and the total score of the inhalation technique was documented. The validated German version of the Living with Asthma Questionnaire was applied to measure asthma-specific quality of life. A self-constructed self-efficacy scale based in parts on a standardized generic self-efficacy questionnaire and supplemented by some disease-specific items was employed to investigate any changes in patients’ perceptions of their self-management skills and ability to deal with the disease.

The asthma knowledge questionnaire, which focused on basic information about the disease and drug therapy, was developed and tested in cooperation with respiratory physicians, clinical psychologists, and clinical pharmacists involved in the German PC efficacy study. To measure patients’ adherence, the validated 4-item Morisky medication adherence scale was utilized.

GKV information about patients’ asthma-related hospital admissions (ICD J 45 and J 46), absence from work, and drug consumption (ATC R03) were evaluated in a subgroup of 55 patients of the intervention group for whom claims data were available and compared with a control group of 550 patients.

STATISTICAL ANALYSIS

Statistical computations concerning clinical and humanistic outcomes were performed using SAS version 8.2 (SAS Institute, Cary, NC). The medial trend of the intervention over time was evaluated using a pre/post/follow-up design (SAS, procedure “mixed” for repeated measure in a split-plot design). This type of ANOVA outperforms the standard General Linear Model approach in terms of power and flexibility of coding single contrasts. Furthermore, it allows for explicit modeling of covariance structure over time, thus giving appropriate standard error for determining statistical significance of effects. Every analysis was repli-
cated using its nonparametric counterpart (Friedman test), which displayed parallel results. Two-tailed $\alpha$ level was predetermined on $\alpha = 0.05$ with power $(1-\beta) = 0.8$.

All scales and subscales derived from the repeatedly administered questionnaires (quality of life, self-efficacy, knowledge) were linearly transformed to a percentage scale ranging from 0 to 100. Low values are associated with a low characteristic of the measured construct and vice versa.

GKV (AOK and Barmer) data were evaluated using a pre/post-design (1-y periods prior to and during the study) against a control group generated from AOK data in the Federal State of Hesse. There was a need to control for the amount of drugs prescribed before baseline and avoid potential bias with regard to anti-asthmatics prescribed to patients enrolled in the study. This was achieved by analyzing individual drug consumption in time frames of 180 days before and during the study (Figure 1, Table 1) instead of analyzing the entire data set.

Results

Over the enrollment period of 6 months, 39 of 57 pharmacies recruited a total of 183 patients. Females were slightly older than males (44.2 vs 41.1 y). Patients’ baseline characteristics are summarized in Table 2. During the study year, 6 (15%) pharmacies and 55 (30%) patients were eventually lost to follow-up. Reasons for patients’ dropout were lack of interest/lack of time (n = 17), patients diagnosed with COPD based on lung function tests and other patient data (n = 6), relocation (n = 4), physician related (n = 2), and pharmacy related (n = 4). No specific reason was filed for 22 patients. Pharmacy dropout was mainly due to change of working place of the responsible and trained pharmacist or when all recruited patients of the pharmacy were lost to follow-up.

**CLINICAL OUTCOMES**

Peak expiratory flow rates measured in the pharmacy improved significantly over time. In addition, the patients’ self-reported symptoms and asthma severity decreased significantly. Increases in lung function ($FEV_1$ and $VC$) were not significant over time (Table 3).

**HUMANISTIC OUTCOMES**

The inhalation technique improved significantly over time. Furthermore, patients’ knowledge, self-efficacy, and adherence were enhanced. Significant improvements could also be established in the summary score of the Living with Asthma Questionnaire and its 2 subscales: physical symptoms and psychological distress. The functional status subscale remained unchanged (Table 3).

**ECONOMIC OUTCOMES**

Equivalence tests revealed no statistically significant differences in any of the parameters of the subgroup of 55 patients for whom health insurance funds’ claims data were available compared with the entire intervention group. We therefore concluded that the subgroup is representative of the intervention group.

Evaluation of the claims data showed a higher increase in the number of recipients of long-acting $\beta_2$-agonists and inhaled steroids in the subgroup compared with the control.
group (Table 1). Concurrently, a larger decrease in prescribed short-acting inhaled β₂-agonists and oral steroids was observed. The number of prescriptions for theophylline remained unchanged in the control group and decreased in the study group.

Low overall asthma severity at/before baseline accompanied by low hospitalization rates in the preceding year and the small number of patients for whom health economic data (hospitalizations, absence from work) were available did not allow further analysis of economic outcomes.

Discussion

As the efficacy of PC has been proven before, we chose a more naturalistic pre/post-design with repeated measurement, despite known limitations in its internal validity. In contrast with other studies, this intervention focused not only on clinical and humanistic outcomes, but also on quality of medical care and economic outcomes.

The 30% patient dropout rate is comparable with the 28% rate found in our controlled study and less than that in the study by McLean et al. The small changes in FEV₁ and VC are also consistent with the results found in other studies. Obviously, the variable nature of clinical measurements, especially in multicenter studies without validation of the measurement procedure, is a limitation in evaluating clinical outcomes in a field study like this. Nevertheless, it is known that patients with asthma enrolled in a PC program perceive progress in symptoms and their well-being. This benefit of PC is also clearly supported by our findings. Self-reported symptoms and peak flow measurements (+30 L/min) under pharmacists’ supervision improved significantly over time. Most notably, the significant decrease in asthma severity rated by the patients’ physicians can be interpreted as a result of the intervention.

All humanistic parameters improved significantly. One of the most important conditions for patients’ ability to affect the course of their disease is a high level of self-efficacy and adherence. Therefore, the improved self-efficacy and adherence we noted can be regarded as the most fundamental results. The enhancement in self-management demonstrated by higher knowledge, self-efficacy, and inhalation technique might have led to the positive impact on adherence and quality of life. Patients who know more about the disease and drug therapy and perceive more control of their asthma are better prepared to cope with the burden of their disease. Given the importance of patient education in affecting the outcomes of persons with asthma, even small improvements in the knowledge score, as in our study, are relevant.

Although there is only a low correlation between increases in quality of life and lung function in patients with asthma, the subscales of the Living with Asthma Questionnaire show a clear relationship to asthma severity. Consequently, it is likely that the increase in quality of life indicates a reduction in asthma symptoms. The need for education by healthcare professionals has recently been highlighted again, showing similar deficits in patients’ inhalation techniques for both pressurized metered-dose and dry powder inhalers.

Concerning the evaluation of economic outcomes, GKV claims data from only 55 patients insured within the 2 cooperating statutory health insurance funds (AOK and Barmer) could be analyzed. Because of the rather low asthma severity of these patients and an unexpectedly low asthma-related hospitalization rate, an interpretation of changes in hospitalization and in days of disability/absence from work was not possible. Furthermore, information on total healthcare costs was not available.

An analysis of shifts in prescribed pharmacotherapy showed promising results. The higher treatment prevalence with long-acting β₂-agonists and inhaled steroids in the intervention group compared with the controls without an increase in asthma severity indicates an improvement in the quality of care.

Furthermore, the higher decrease in prescribed short-acting β₂-agonists and oral steroids in the intervention group emphasizes the achieved humanistic and clinical results of the study.

In this study, patient and healthcare practitioner satisfaction was not systematically explored. Not surprisingly, patient satisfaction in PC programs is usually high. Both pharmacists and physicians reported an overall high patient satisfaction with the service. This is despite the lack of time/lack of interest to comply with all scheduled appointments, completing questionnaires, self-monitoring, and documenting peak flow and symptoms, among others.

### Table 2. Baseline Data of Study Population

<table>
<thead>
<tr>
<th>Age, y, mean (SD)</th>
<th>43.1 (13.7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>range</td>
<td>18–65</td>
</tr>
<tr>
<td>Gender (%)</td>
<td></td>
</tr>
<tr>
<td>male</td>
<td>34.4</td>
</tr>
<tr>
<td>female</td>
<td>65.6</td>
</tr>
<tr>
<td>Employment status (%)</td>
<td></td>
</tr>
<tr>
<td>employed or self-employed</td>
<td>63.9</td>
</tr>
<tr>
<td>unemployed or retired</td>
<td>36.1</td>
</tr>
<tr>
<td>Smoking status (%)</td>
<td></td>
</tr>
<tr>
<td>current smoker</td>
<td>18.7</td>
</tr>
<tr>
<td>ex-smoker</td>
<td>58.8</td>
</tr>
<tr>
<td>non-smoker</td>
<td>22.5</td>
</tr>
<tr>
<td>Asthma severity (%)</td>
<td></td>
</tr>
<tr>
<td>4 (severe, persistent)</td>
<td>4.4</td>
</tr>
<tr>
<td>3 (moderate, persistent)</td>
<td>20.3</td>
</tr>
<tr>
<td>2 (mild, persistent)</td>
<td>33.0</td>
</tr>
<tr>
<td>1 (intermittent)</td>
<td>30.2</td>
</tr>
<tr>
<td>not specified</td>
<td>12.1</td>
</tr>
<tr>
<td>Asthma severity, mean (SD)</td>
<td>2 (0.9)</td>
</tr>
<tr>
<td>FEV₁, % VC (Tiffeneau), mean (SD)</td>
<td>75.7 (15.9)</td>
</tr>
<tr>
<td>Asthma etiology (%)</td>
<td></td>
</tr>
<tr>
<td>allergic</td>
<td>22.0</td>
</tr>
<tr>
<td>non-allergic</td>
<td>17.0</td>
</tr>
<tr>
<td>mixed form</td>
<td>45.6</td>
</tr>
<tr>
<td>unclear or not specified</td>
<td>15.4</td>
</tr>
</tbody>
</table>

FEV₁ = forced expiratory volume in 1 second; VC = vital capacity.

*N = 183.

*bDetermined by patient’s physician.
For the first time in Germany, both physicians’ and pharmacists’ associations jointly and officially supported a PC program. Moreover, for the first time, health insurance funds provided claims data to evaluate such a program in more detail. Eventually, this cooperation was pivotal for institution of the first German integrated care contract signed in December 2004 between physicians’ and pharmacists’ associations and one of the participating health insurance funds (Barmer). This contract established the combined family pharmacy–family physician concept where patients choose and enroll in both a physician’s (GP) office and a community pharmacy.

Conclusions

The findings of our study show that an intensive cooperation between pharmacists, physicians, and patients with asthma within the concept of PC has a clear positive impact on humanistic and, to some extent, on clinical outcomes. In particular, the factors that affect patients’ self-management also improved. In cooperation with the prescribing physicians, drug therapy changed toward evidence-based guidelines. To evaluate potential economic benefits, future research should focus on patients with more severe or uncontrolled asthma, such as those with a significant number of emergency department visits or hospitalizations.

Table 3. Clinical and Humanistic Outcomes

<table>
<thead>
<tr>
<th>Outcome Measures</th>
<th>Baseline</th>
<th>6 mo</th>
<th>12 mo</th>
<th>p Value (global)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthma severity (1–4)</td>
<td>2.0 (0.9)</td>
<td>1.7 (0.8)</td>
<td>1.7 (0.8)</td>
<td>&lt;0.002</td>
</tr>
<tr>
<td>Dyspnea severity (0–4)</td>
<td>2.2 (0.8)</td>
<td>2.0 (0.9)</td>
<td>2.0 (0.9)</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>FEV1 (L)</td>
<td>2.8 (1.0)</td>
<td>2.9 (1.0)</td>
<td>2.9 (1.0)</td>
<td>0.48</td>
</tr>
<tr>
<td>VC (L)</td>
<td>3.8 (1.3)</td>
<td>3.8 (1.3)</td>
<td>3.8 (1.3)</td>
<td>0.26</td>
</tr>
<tr>
<td>FEV1 % VC (Tiffeneau)</td>
<td>75.7 (15.9)</td>
<td>75.6 (14.1)</td>
<td>76.2 (14.8)</td>
<td>0.49</td>
</tr>
<tr>
<td>Peak expiratory flow rate (L/min, measured in the pharmacy)</td>
<td>402.9 (114.9)</td>
<td>436.6 (116.0)</td>
<td>433.4 (110.3)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Asthma symptoms (0–3, rated by pt.)</td>
<td>3.1 (2.3)</td>
<td>2.2 (2.2)</td>
<td>2.5 (2.3)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Knowledge</td>
<td>18.9 (7.8)</td>
<td>23.4 (7.7)</td>
<td>24.2 (8.3)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Living with Asthma Questionnaire

- sum score                                      | 44.4 (14.3)| 48.3 (13.1)| 49.9 (13.5)| <0.001           |
- physical symptoms                             | 48.1 (19.8)| 52.2 (18.2)| 54.1 (18.4)| <0.001           |
- psychological distress                         | 48.1 (16.1)| 53.5 (14.7)| 55.4 (14.6)| <0.001           |
- functional status                              | 37.4 (14.9)| 39.4 (13.6)| 40.4 (13.7)| 0.61             |
- Self-efficacy                                  | 45.8 (9.8)| 52.3 (9.1)| 54.3 (9.5)| <0.001           |
- Adherence (Morisky, 4–8)*                      | 6.7 (1.8)| 7.1 (1.3)| 7.1 (1.4)| <0.001           |
- Inhalation technique (MDI, scale 0–7)          | 5.2 (1.5)| 6.4 (1.1)| 6.6 (1.0)| <0.001           |
- Inhalation technique (DPI, scale 0–7)          | 5.5 (1.3)| 6.6 (0.9)| 6.5 (1.0)| <0.001           |

DPI = dry powder inhaler; FEV1 = forced expiratory volume in 1 second; MDI = metered-dose inhaler; VC = vital capacity.

*Mean (SD).  
*Transformed percentage scale 0–100 (see “Statistical Analysis”).  
*8 = best, 4 = worst adherence.

References


Evaluar la efectividad de PC con respecto a los resultados clínicos, humanísticos, y económicos en pacientes adultos con asma.

**MÉTODOS:** Se realizó un estudio donde se llevó a cabo una intervención por un período de 12 meses. Al comienzo del estudio, 39 farmacias de comunidad, 84 médicos primarios (medicina general, medicina interna, y neumólogos), y 183 pacientes entre las edades de 18 a 65 años con diagnóstico de asma fueron incluidos. Para evaluar los resultados económicos, 2 compañías alemanas aseguradoras de servicios de salud proveyeron datos de reclamaciones de 55 pacientes para un año antes de comenzar el estudio y el año subsiguiente. Se llevó a cabo un pareo de 1:1 para comparar los datos del grupo que recibió la intervención al de un grupo control de 550 pacientes.

**RESULTADOS:** Se detectaron mejoras significativas para todos los resultados humanísticos (calidad de vida específica para el asma, autoeficacia, conocimiento, y adherencia a medicamentos). Además, la severidad del asma, los síntomas reportados por los pacientes, el valor de flujo máximo de expiración, y la técnica de inhalación del paciente mejoraron. Aumentos en el FEV1 (volumen de expiración forzada en un segundo) y la VC (capacidad vital) no fueron significativos durante el transcurso del tiempo de la investigación. La evaluación de los datos de las compañías aseguradoras demostró un cambio hacia mejor adherencia y terapias basadas en evidencia.

**CONCLUSIONES:** El estudio demuestra que en pacientes con asma, el PC tiene un impacto positivo en resultados humanísticos y, en cierta medida, en los resultados clínicos. Para determinar los posibles beneficios económicos, investigaciones futuras deben enfocar pacientes con asma más severa.

Homero A Monsanto

**RESUMÉ**

**CONTEXT:** Malgré les progrès de la pharmacothérapie de l’asthme au cours des dernières années, aucun changement significatif de la morbidité et de la mortalité liés à cette maladie n’est survenu. Il demeure pertinent de déterminer si les soins pharmaceutiques (SP) peuvent influencer les retombées de santé.

**OBJECTIF:** Évaluer l’efficacité des SP sur les retombées cliniques, humanitaires, et économiques chez des adultes asthmatiques.

**MÉTHODES:** Une étude d’intervention a été réalisée sur une période de 12 mois. Au départ, cette étude comptait 39 pharmacies privées, 84 médecins de soins primaires ( omnipraticiens, spécialiste en médecine interne et en pneumologie), et 183 patients asthmatiques âgés de 18 à 65 ans. Afin d’évaluer les impacts économiques, 2 assureurs de soins de santé allemands ont donné accès aux réclamations de leurs patients (n = 55) pour une période de 2 ans. Un ratio d’appariement de 1:10 a été utilisé pour comparer les données du groupe intervention au groupe contrôle (n = 550).

**RÉSULTATS:** Des améliorations significatives ont été trouvées pour les toutes les variables humanitaires: qualité de vie liée à l’asthme, l’efficacité personnelle perçue, les connaissances, et l’observance au traitement. De plus, la sévérité de l’asthme, les symptômes rapportés par les patients, le débit expiratoire de pointe et la technique d’inhalation des médicaments ont été améliorés. Des améliorations du volume expiratoire forcé en un seconde (FEV1) et la capacité vitale n’étaient cependant pas significatives. L’analyse des réclamations aux assureurs a démontré une tendance à l’amélioration de l’observance et l’adhésion à des thérapies fondées sur les évidences médicales.

**CONCLUSIONS:** Cette étude démontre que les SP offerts aux patients asthmatiques ont des effets positifs sur les retombées humanitaires et, dans de moindres proportions sur les retombées cliniques. Pour déterminer les économies potentielles, les recherches futures devront cibler des patients plus sévèrement atteints.