Lung Function Trajectory Types in Never-Smoking Adults With Asthma: Clinical Features and Inflammatory Patterns

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ABSTRACT

Purpose: Asthma is a heterogeneous disease that responds to medications to varying degrees. Cluster analyses have identified several phenotypes and variables related to fixed airway obstruction; however, few longitudinal studies of lung function have been performed on adult asthmatics. We investigated clinical, demographic, and inflammatory factors related to persistent airflow limitation based on lung function trajectories over 1 year.

Methods: Serial post-bronchodilator forced expiratory volume (FEV1) values were obtained from 1,679 asthmatics who were followed up every 3 months for 1 year. First, a hierarchical cluster analysis was performed using Ward's method to generate a dendrogram for the optimum number of clusters using the complete post-FEV1 sets from 448 subjects. Then, a trajectory cluster analysis of serial post-FEV1 sets was performed using the k-means clustering for the longitudinal data trajectory method. Next, trajectory clustering for the serial post-FEV1 sets of a total of 1,679 asthmatics was performed after imputation of missing post-FEV1 values using regression methods.

Results: Trajectories 1 and 2 were associated with normal lung function during the study period, and trajectory 3 was associated with a reversal to normal of the moderately decreased baseline FEV1 within 3 months. Trajectories 4 and 5 were associated with severe asthma with a marked reduction in baseline FEV1. However, the FEV1 associated with trajectory 4 was increased at 3 months, whereas the FEV1 associated with trajectory 5 was persistently disturbed over 1 year. Compared with trajectory 4, trajectory 5 was associated with older asthmatics with less atopy, a lower immunoglobulin E (IgE) level, sputum neutrophilia and higher dosages of oral steroids. In contrast, trajectory 4 was associated with higher sputum and blood eosinophil counts and more frequent exacerbations.

Conclusions: Trajectory clustering analysis of FEV1 identified 5 distinct types, representing well-preserved to severely decreased FEV1. Persistent airflow obstruction may be related to non-atopy, a low IgE level, and older age accompanied by neutrophilic inflammation and low baseline FEV1 levels.

Keywords: Adult; asthma; disease progression; forced expiratory volume; inflammation; phenotype
INTRODUCTION

Asthma is a heterogeneous airway disorder that includes distinct phenotypes with diverse etiologies, natural histories and treatment responses. One goal of asthma management is the maintenance of normal pulmonary function with the fewest anti-asthma medications and the control of risk factors such as acute exacerbation and persistent airway obstruction. Although the majority of asthmatics retain normal or near-normal lung function throughout life, a subset of individuals with asthma shows persistent declines in lung function despite high doses of inhaled (ICS) or systemic corticosteroids (SCSs). This group is categorized as a distinct phenotype of asthma (severe fixed airflow or remodeled asthma), and their novel clinical and molecular aspects have been highlighted by cluster analyses. One of the characteristic molecular features of this group is a diverse immune response with eosinophilic and/or neutrophilic airway inflammation. Eosinophilic inflammation, a well-established sub-phenotype, carries a risk of frequent exacerbations and decreased lung function, but responds well to corticosteroids or biological therapies targeted to T-helper-2 (Th2) inflammation. In contrast, chronic airflow obstruction with resistance to corticosteroid therapy is frequently observed in asthmatic patients with neutrophilic airway inflammation.

Most previous cluster analyses involved cross-sectional evaluations of clinical and laboratory parameters; thus, their results may have limited predictive power with regard to the persistence of airway dysfunction over time. Additionally, the levels of stability of the cluster-specific outcomes are imperfect after long-term follow up. These instabilities are perhaps unsurprising in asthma, which is characterized by temporal variability in lung function due to spontaneous or mediation-induced improvements and changes in triggers (e.g., allergen exposure and air pollution, viral infections, and new medications). Thus, it is necessary to understand the dynamics of the clinical and laboratory parameters of asthmatics with persistent airway obstruction using repeated assessments over time.

In our previous cluster analysis, the differences in FEV1 were consistent in 4 clusters. However, the severe obstructive asthma cluster showed persistently lower FEV1 values over the 12-month followup period. Although airway obstruction in this cluster improved over time, the mean final FEV1 value was subnormal, suggesting the presence of persistent airway obstruction in a certain number of asthmatics. Thus, to identify related clinical and demographic factors, an alternative method such as lung function trajectory analysis is needed. To date, several longitudinal lung function trajectory analyses of allergy sensitization, bronchial responsiveness and asthma symptoms in childhood have been performed. Lung function trajectories differ significantly according to comorbidity; considerable lung function impairment was observed from birth to 16 years of age among subjects with asthma, atopic dermatitis or allergic rhinitis. In a trajectory study, children who had lung function deficits as neonates showed persistent airway dysfunction at the age of 7 years. Another longitudinal population study from childhood to adulthood identified being male, airway hyperresponsiveness, low lung function and eosinophilia as risk factors for airway remodeling. However, few trajectory analyses of lung functions have identified risk factors associated with persistent airway obstruction in adult asthmatics.

This prompted us to evaluate clinical, demographic and inflammatory factors related to persistent airflow limitation using the lung function trajectories of adult asthmatics from baseline to 1 year.
MATERIALS AND METHODS

Selection of subjects
We included 1,679 non-smoking asthmatics drawn from a Korean Asthma Database cohort (n = 5,326) after excluding ex-smokers (n = 515), current smokers (n = 451), patients with less than 1 year of followup (n = 2,264), and those with fewer than 2 measurements of post-bronchodilator (BD) FEV1 over 4 scheduled visits at 3-month intervals during the 1 year followup (n = 417) (Fig. 1). Patients with parenchymal lung disease on chest X-ray — including bronchiectasis, pulmonary tuberculosis, chronic obstructive pulmonary disease (COPD) and interstitial lung diseases — were also excluded. Patients met the following criteria: physician diagnosis of asthma and presence of reversible airway obstruction (short-acting BD-induced FEV1 > 12% and 200 mL), airway hyper-reactivity (methacholine PC20 < 10 mg/mL), or improvement in FEV1 > 20% after 2 weeks of treatment with ICS or SCS.

Demographic information such as age at enrollment, sex, body mass index (BMI), age at asthma onset and asthma duration was collected at the baseline visit. All patients underwent standardized assessments, which included peripheral blood total and differential cell counts, serum total immunoglobulin E (IgE), chest radiography, and skin prick tests with 24 common inhalant allergens (Bencard Co., Brentford, UK). Atopy was defined as the presence of a wheal reaction to 1 or more of the allergens characterized by a ≥3 mm greater diameter than the normal saline control or equal to or greater than that induced by histamine (1 mg/mL).

Forced expiratory volume 1 second (FEV1), forced vital capacity (FVC), and pre- and post-BD FEV1 were measured before and after 2 puffs of albuterol (360 µg) were administered with a

Fig. 1. Flowchart of the study protocol.
FEV1, forced expiratory volume in 1 second.
metered-dose inhaler at baseline and at 3-month intervals. PC20 methacholine (mg/mL) and sputum cell analyses were performed in eligible cases at enrollment. Sputum was induced using isotonic saline containing a short-acting BD and processed within 2 hours as previously described. Slides were prepared using cytocentrifugation, and 500 cells were counted on each slide after staining with Diff-Quick (American Scientific Products, Chicago, IL, USA). The patients were treated according to the Guidelines for Asthma Treatment issued by the Korean Academy of Asthma Allergy and Clinical Immunology. The total amounts of ICS and SCS administered during the first year were determined based on the electronic medical records and expressed as equivalent dosages of fluticasone propionate per day and equivalent dosages of prednisolone per year, respectively. Exacerbation was defined as described by American Thoracic Society/European Respiratory Society Task Force on Asthma. The protocol was approved by the Ethics Committees of Soonchunhyang Bucheon Hospital (SCHBC_2014_07_028) and Hallym Sacred Heart Hospital (2016_I009).

Trajectory clustering analysis
We performed a trajectory clustering analysis on the longitudinal post-FEV1 data (baseline and months 3, 6, 9 and 12) from 1,679 patients as follows: 1) the first estimate of the number of likely trajectories using the non-missing data of 448 subjects and 2) the second estimate using the full data from the total of 1,679 subjects after imputation of missing values (Supplementary Table S1). In the first step, a hierarchical cluster analysis of the longitudinal post-FEV1 values was performed using Ward’s method to generate a dendrogram. Cluster boundaries were defined by large differences between successive fusion levels. Cluster quality was assessed by 2-step clustering using the Euclidean method with SPSS software version 18 (IBM Corp., Armonk, NY, USA) and then by trajectory clustering analysis using kml in the R package for k-means clustering of longitudinal data. In the second step, we imputed post-FEV1% missing values for 1,679 subjects using regression methods. The imputed data were subjected to trajectory clustering analysis as described above. The trajectory clustering analysis procedure is summarized in Fig. 1.

Statistical analysis
Analysis of variance (ANOVA) and Kruskal-Wallis and χ2 tests were used to compare parametric continuous, nonparametric continuous and categorical variables, respectively. Independent t tests and Mann-Whitney U tests were applied to compare continuous parametric and nonparametric variables between 2 trajectories. All statistical analyses were performed using SPSS.

RESULTS

Trajectory clustering analysis of post-BD FEV1% over 1 year in 448 asthmatics
Five clusters were optimal for discriminating the longitudinal values of post-BD FEV1 in the first 448 subjects and the second 1,679 subjects (Supplementary Fig. S1). The 5 trajectories yielded unique classification patterns for the 448 subjects (Tables 1 and 2, Supplementary Fig. S2): Trajectories 1 and 2 were characterized by normal baseline FEV1% values, which were maintained during the followup period. Trajectory 3 was associated with a mild decrease in baseline FEV1 with minimal, but significant, improvement over time (P < 0.001). Trajectory 4 was associated with moderate-to-severe derangement of baseline FEV1%, which dramatically increased to a normal level in the first 3 months (P < 0.001) and was maintained to the end of followup. Trajectory 5 was characterized by the lowest baseline FEV1%, which remained low to the end of the followup (P < 0.001).
Clinical characteristics associated with the trajectory clusters in the total 1,708 asthmatics

Among the total of 8,395 followup FEV1 values (1,679 subjects at 5 time points), 2,722 FEV1 values (32.3%) were missing (Supplementary Table S1). Thus, imputed FEV1 values were obtained by a regression analysis of the 5,673 measured FEV1 values. The trajectory clustering of the total of 8,395 FEV1 values yielded 5 trajectories (Tables 1 and 2, Fig. 2 and Supplementary Fig. S2), which was similar to the 5 trajectories for the 448 subjects. The clinical and demographic characteristics associated with the 5 trajectories are shown in Table 3. Trajectory 1 (15.4% of the total subjects) was associated with a supra-normal FEV1% from baseline to the 12-month follow up (112.75% ± 13.93% pred. and 116.07% ± 11.49% pred., respectively, Tables 1 and 2). The patients whose course followed this trajectory were middle-to-old-aged, predominantly female (84.6%) and had a higher BMI than those whose course followed the other trajectories. Among these subjects, 10% showed a short-acting BD-induced increase in FEV1 (85.7% for PC20 methacholine), and 10.4% exhibited improvement in FEV1 after treatment. The majority (87.6%) were on a low dose of inhaled ICS (<250 µg fluticasone equivalent/day) and had the lowest frequency of exacerbations over 1 year (0.2/year, Table 3).

Trajectory 2 (31.8% of the total subjects) was characterized by FEV1% values in the normal range from baseline (99.25% ± 9.64% pred.) to the end of follow up (100.98% ± 9.29% pred.), these values were lower than those associated with trajectory 1 (P < 0.001) (Tables 1 and 2, Fig. 2). This group was younger and predominantly females and had a high frequency of atopy. Among these subjects, 11.4% showed a short-acting BD-induced increase in FEV1 (81.7% for PC20 methacholine), and 10.4% exhibited improvement in FEV1 after treatment (Table 3). Most of these subjects (86.5%) received a low dose of ICS and had a low frequency of exacerbations over 1 year (0.3 ± 1.4). Thus, trajectories 1 and 2 represent mild asthma.

Trajectory 3 (32.0% of the total subjects) was associated with a mild decrease in basal lung function (86.09% ± 9.17% pred.), which was slightly improved at the 12-month followup (86.76% ± 9.60% pred., P < 0.001). These FEV1 levels were significantly lower than those of trajectories 1 and 2 (P < 0.001) (Tables 1 and 2, Fig. 2). This group comprised patients with a young age at onset with a longer disease duration. Among these subjects, 12.5% showed

### Table 1. Longitudinal changes in the complete data of post-BD (448 subjects)

| Variables                  | Trajectory 1 (n=104) | Trajectory 2 (n=164) | Trajectory 3 (n=103) | Trajectory 4 (n=39) | Trajectory 5 (n=38) | P value | Bonferroni post hoc P value (trajectory 4 vs. 5) |
|----------------------------|----------------------|----------------------|----------------------|---------------------|---------------------|---------|------------------------------------------------|
| Baseline post-BD FEV1 % pred. | 112.75 ± 13.93       | 99.25 ± 9.64         | 86.09 ± 9.17         | 61.25 ± 13.04       | 62.71 ± 15.97       | 1.25E-201 | 1.000                                           |
| Imputed baseline post-BD FEV1 % pred. | 115.88 ± 9.30    | 100.06 ± 7.81        | 85.24 ± 8.09         | 92.44 ± 11.17       | 66.53 ± 13.49       | 1.25E-223 | 6.12E-120                                       |
| Imputed 3rd month post-BD FEV1 % pred. | 119.34 ± 10.61   | 103.55 ± 8.43        | 87.29 ± 10.18        | 95.52 ± 10.37       | 66.84 ± 13.31       | 3.40E-218 | 2.74E-121                                       |
| Imputed 6th month post-BD FEV1 % pred. | 119.89 ± 14.00  | 102.09 ± 10.59       | 86.47 ± 10.16        | 99.51 ± 10.69       | 66.34 ± 14.09       | 1.6E-194  | 1.36E-126                                       |
| Imputed 12th month post-BD FEV1 % pred. | 116.07 ± 11.49 | 100.58 ± 9.29        | 86.76 ± 9.60         | 98.69 ± 10.53       | 69.21 ± 13.47       | 1.02E-191 | 7.80E-123                                       |

BD, bronchodilator; FEV1, forced expiratory volume in 1 second.

### Table 2. Longitudinal changes in the complete data of post-BD (1,679 subjects)

| Variables                  | Trajectory 1 (n=259) | Trajectory 2 (n=534) | Trajectory 3 (n=538) | Trajectory 4 (n=139) | Trajectory 5 (n=209) | P value | Bonferroni post hoc P value (trajectory 4 vs. 5) |
|----------------------------|----------------------|----------------------|----------------------|---------------------|---------------------|---------|------------------------------------------------|
| Baseline post-BD FEV1 % pred. | 108.19 ± 13.69       | 94.11 ± 9.06         | 77.07 ± 9.85         | 57.92 ± 13.48       | 52.34 ± 15.65       | 8.59E-118 | 0.338                                           |
| 3rd month post-BD FEV1 % pred. | 112.25 ± 9.51        | 94.13 ± 8.69         | 79.50 ± 7.24         | 94.90 ± 9.87        | 60.71 ± 14.99       | 1.07E-118 | 9.7E-45                                         |
| 6th month post-BD FEV1 % pred. | 116.04 ± 11.25       | 96.97 ± 10.44        | 80.27 ± 7.95         | 100.57 ± 9.27       | 62.80 ± 15.34       | 2.96E-111 | 1.41E-43                                        |
| 9th month post-BD FEV1 % pred. | 115.69 ± 11.29       | 95.99 ± 9.21         | 81.37 ± 8.91         | 101.27 ± 9.67       | 60.35 ± 15.91       | 6.45E-114 | 1.38E-50                                        |
| 12th month post-BD FEV1 % pred. | 113.66 ± 11.65       | 96.07 ± 8.76         | 82.01 ± 7.86         | 100.17 ± 10.22      | 62.25 ± 15.78       | 1.22E-106 | 2.89E-46                                        |

BD, bronchodilator; FEV1, forced expiratory volume in 1 second.

Clinical characteristics associated with the trajectory clusters in the total 1,708 asthmatics

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| Variables                                      | Trajectory 1 (n=259) | Trajectory 2 (n=534) | Trajectory 3 (n=538) | Trajectory 4 (n=139) | Trajectory 5 (n=209) | P value (ANOVA) | P value (T4 vs. T5) |
|------------------------------------------------|----------------------|----------------------|----------------------|----------------------|----------------------|-----------------|-------------------|
| Age at enrollment (yr)                         | 50.0 ± 15.7          | 44.6 ± 14.8          | 43.3 ± 16.0          | 45.9 ± 15.7          | 49.9 ± 16.7          | 4.41E-10        | 0.022             |
| Age at onset (yr)                              | 45.9 ± 16.6          | 41.0 ± 15.9          | 39.3 ± 16.6          | 41.4 ± 17.1          | 44.3 ± 18.2          | 1.09E-06        | 0.105             |
| Male (%)                                       | 15.4                 | 16.7                 | 18.6                 | 23.3                 | 23.9                 | 0.840           | 0.726             |
| BMI (kg/m²)                                    | 24.4 ± 3.3           | 24.1 ± 3.6           | 23.8 ± 3.7           | 23.6 ± 3.8           | 23.6 ± 4.4           | 0.048           | 0.392             |
| Atopy (%)                                      | 41.3                 | 50.4                 | 49.6                 | 51.1                 | 40.7                 | 0.029           | 0.056             |
| Log₁₀ total IgE                                | 2.0 ± 0.6            | 2.0 ± 0.7            | 2.0 ± 0.6            | 2.2 ± 0.6            | 2.0 ± 0.7            | 0.007           | 0.008             |
| Asthma duration (yr)                           | 2.6 ± 6.7            | 2.5 ± 6.0            | 2.6 ± 6.9            | 3.0 ± 8.3            | 3.9 ± 8.6            | 0.154           | 0.265             |
| Subject experiencing exacerbation in the 1st one year (%) | 5.8                  | 9.9                  | 20.8                 | 61.9                 | 47.4                 | 1.26E-62        | 0.008             |
| Exacerbation frequency in the 1st one year     | 0.2 ± 1.1            | 0.3 ± 1.4            | 0.4 ± 1.4            | 0.6 ± 1.4            | 0.8 ± 1.6            | 4.27E-6         | 0.110             |
| Basal Post-BD FEV₁, % pred.                    | 112.7 ± 13.9         | 99.3 ± 9.6           | 86.1 ± 9.2           | 61.3 ± 13.0          | 62.7 ± 15.3          | 1.25E-201       | 0.241             |
| PC20 of methacholine, mean ± SD (No.)          | 7.2 ± 8.2 (239)      | 5.4 ± 7.4 (504)      | 4.4 ± 6.4 (481)      | 3.6 ± 4.9 (111)      | 3.9 ± 5.8 (132)      | 2.43E-7         | 0.716             |
| Patients with PC20 < 10 mg/ml (%)              | 85.7                 | 81.7                 | 85.9                 | 89.2                 | 87.9                 | 0.001           | 0.750             |
| Improvement of FEV₁ > 20% after medication (%) | 10.4                 | 11.2                 | 10.8                 | 97.1                 | 39.2                 | 5.40E-28        | 9.59E-28          |
| Dosage of ICS*a                                | 85.3 ± 176.9         | 90.0 ± 158.3         | 117.1 ± 204.3        | 140.5 ± 206.7        | 157.6 ± 247.5        | 2.36E-05        | 0.419             |
| Low dose ICS                                   | 2.27 (87.6)          | 462 (86.5)           | 440 (81.8)           | 112 (80.6)           | 157 (75.1)           | 2.68E-04        | 0.300             |
| Medium dose ICS                                | 26 (10.0)            | 50 (11.0)            | 64 (11.9)            | 15 (10.8)            | 35 (16.7)            | 2.36E-05        | 0.261             |
| High dose ICS                                  | 6 (2.3)              | 13 (2.4)             | 30 (6.3)             | 15 (10.8)            | 7 (3.7)              | 2.36E-05        | 0.125             |
| Dosage of systemic corticosteroid*b            | 59.3 ± 357.7         | 55.6 ± 294.5         | 106.2 ± 418.5        | 188.2 ± 377.7        | 313.1 ± 705.1        | 1.61E-13        | 0.006             |
| Blood eosinophils, % (No.)                     | 5.1 ± 5.3 (69)       | 5.6 ± 5.0 (201)      | 4.7 ± 4.0 (199)      | 7.3 ± 7.7 (65)       | 4.6 ± 4.4 (78)       | 0.003           | 0.001             |
| Blood neutrophils, % (No.)                     | 54.2 ± 10.3 (69)     | 54.8 ± 11.8 (201)    | 55.7 ± 12.0 (199)    | 55.5 ± 15.5 (65)     | 58.5 ± 12.6 (78)     | 0.192           | 0.141             |
| Sputum eosinophils, % (No.)                    | 7.1 ± 14.4 (102)     | 12.0 ± 20.9 (219)    | 9.9 ± 13.3 (237)     | 16.4 ± 21.5 (82)     | 9.3 ± 18.8 (117)     | 0.015           | 0.012             |
| Sputum neutrophils, % (No.)                    | 64.5 ± 28.6 (102)    | 62.7 ± 28.4 (219)    | 66.2 ± 26.9 (237)    | 63.1 ± 26.1 (82)     | 68.9 ± 27.6 (117)    | 0.330           | 0.151             |
| Newly enrolled subjects                       | 2.32 (89.6)          | 466 (87.3)           | 465 (86.4)           | 410 (82.0)           | 180 (86.3)           | 0.313           | 0.365             |
| Corticosteroid-naïve patients at enrollment    | 2.39 (100.0)         | 534 (100.0)          | 537 (99.8)           | 139 (100.0)          | 209 (100.1)          | 0.713           | 1.000             |

Data are presented as mean ± standard deviation or number (%). ANOVA, analysis of variance; BD, bronchodilator; BMI, body mass index; FEV₁, forced expiratory volume in 1 second; ICS, inhaled corticosteroids.

*aDosages are presented as equivalent of fluticasone (µg/day), Low dose ICS ≤ 250 µg/day, medium dose 250–500 µg/day, and high dose > 500 µg/day of fluticasone or equivalent; *bDosages are presented as equivalent of prednisolone (mg/year).
a short-acting BD-induced increase in FEV1 (85.9% for PC20 methacholine), and 10.8% showed improvement of FEV1 after treatment (Table 3). The exacerbation frequency over 1 year (0.4 ± 1.4) was slightly higher than that associated with trajectories 1 and 2 (P = 0.115) despite significantly higher doses of ICS and SCS (P = 0.007 and 0.020, respectively, Table 3).

Trajectory 4 (8.3% of the total subjects) was associated with a very low FEV1% at baseline (61.25% ± 13.04% pred.), and this dramatically increased to normal in the first 3 months (92.44% ± 11.17% pred., P < 0.001) and was maintained over the followup period (98.69% ± 10.53% pred.). This trajectory was associated with the highest prevalence of atopy (51.1%) and the highest mean IgE level. Among these subjects, 12.9% showed a short-acting BD-induced increase in FEV1 (89.2% for PC20 methacholine), and 97.1% showed improvement in FEV1 after treatment. The exacerbation frequency over 1 year (0.6 ± 1.4) was significantly higher in those whose course followed this trajectory than in those whose course followed trajectories 1, 2, and 3 (P = 0.027). The PC20 methacholine value of trajectory 4 was the lowest among the 5 trajectories. Larger quantities of ICS and SCS were used for individuals in this trajectory than for those who courses followed trajectories 1, 2 and 3.

Trajectory 5 (12.4% of the total subjects) was associated with a marked decrease in baseline FEV1% values, similar to trajectory 4; however, the FEV1 improved over the followup period (69.21% ± 13.47% pred. at 12 months; P < 0.001), suggesting persistent airway obstruction. Trajectory 5 was associated with a lower FEV1% over the followup period than trajectory 4 (P < 0.001). Among these subjects, 14.8% showed a short-acting BD-induced increase in FEV1 (87.9% for PC20 methacholine), and 39.2% showed improvement in FEV1 after treatment. This group was associated with the lowest frequency of atopy, the longest duration of asthma, the highest exacerbation frequency over 1 year (0.8 ± 1.6), and the highest total ICS and SCS among the trajectories.

Compared with trajectory 4, trajectory 5 was associated with asthmatics with less atopy (P = 0.056), a lower IgE level (P = 0.007), a higher dosage of SCS (P = 0.006), and a similar dosage of ICS (Table 3). Interestingly, trajectory 4 was associated with a higher proportion of subjects experiencing exacerbations over 1 year than was trajectory 5 (69.1% vs. 47.4%, P = 0.008).
Medication other than ICS and SCS showed similar patterns with the corticosteroids; Trajectories 4 and 5 showed higher dose of aminophylline, doxofylline, indacaterol, pranlukast, tiotropium and acebrophylline than other trajectories during the study period (Supplementary Table S3). Among them, dose of doxofylline was higher for subjects in trajectory 5 than those in trajectory 4 ($P = 4.9 \times 10^{-5}$). Dose of other drugs were comparable between trajectories 4 and 5.

**Inflammatory cells in sputum and peripheral blood**
Sputum inflammatory cell patterns were stratified into eosinophilic, neutrophilic, mixed and paucigranulocytic. The subjects whose course followed trajectory 4 showed predominantly eosinophilic (28.0%) and mixed (32.9%) inflammation, and those whose course followed trajectory 5 predominantly showed neutrophilic inflammation (52.1%) ($P = 0.001$, Fig. 3, Supplementary Table S3). Unlike those whose course followed trajectory 5, subjects whose course followed trajectory 4 also had peripheral blood eosinophilia ($P = 0.001$) (Table 3).

**DISCUSSION**
In the present study, we identified 5 lung function trajectories based on post-BD FEV1 values over a 1 year followup and identified between-group differences in clinical and demographic parameters. Among the trajectories, 2 represented severe asthma: trajectories 4 and 5 (8.3% and 12.4% of the subjects, respectively) were associated with marked reductions in baseline FEV1; thus, about 20% of the subjects had severely deranged lung function at baseline. Interestingly, the baseline FEV1 value of those whose course followed trajectory 4 dramatically increased after 3 months of treatment, whereas those whose courses trajectory 5 showed persistent derangement of FEV1 despite continuous treatment with ICS. These data indicate that about 12% of asthmatics suffer from persistent airway obstruction.

Initial lung function is an important determinant of asthma outcomes; indeed, a low FEV1 value is a major risk factor for asthma exacerbations.$^{1-26}$ Interestingly, the 2 groups, trajectories 4 and 5 in the present study showed similar responses to short-acting BDs (18.7% vs. 23.9%, $P = 0.248$) and to methacholine (3.6 ± 4.9 vs. 3.9 ± 5.8 mg/mL, $P = 0.716$).
Therefore, airway constriction and bronchial smooth-muscle responsiveness were similar in the 2 groups because these 2 methods are related to the function of bronchial airway smooth muscle. Accordingly, other factors may be responsible for the variation in the outcomes of asthma treatment. Trajectory 4 was associated with atopic tendency, high IgE levels and high frequency of exacerbations, whereas trajectory 5 was associated with less atopic, lower IgE levels, and less frequent exacerbations. Additionally, differences in airway inflammation patterns may be related to the diverse responses to conventional treatments including ICS. In the present study, trajectories 4 and 5 were associated with contrasting patterns: eosinophilic inflammation in trajectory 4 and neutrophilic inflammation in trajectory 5. Eosinophilic inflammation is a pivotal trait of asthma and a marker of a good response to ICS and SCS. Although the FEV1 value for trajectory 4 at month 12 ranged from 77.7% to 135.0% of the predicted value, only 4 of 139 subjects had a subnormal FEV1 (< 80%), suggesting that, for those whose course follows this trajectory, airway obstruction with eosinophilic inflammation can be controlled using ICS.

Our finding of a high frequency of exacerbations in those whose course followed trajectory 4 is in agreement with previous reports that persistent eosinophilia in the peripheral blood and airways is associated with frequent exacerbations and fatal asthma attacks. Moreover, 15% of the subjects whose course followed trajectory 5 also had eosinophilic inflammation. Therefore, the eosinophilic inflammation in this group may be resistant to treatment with ICS and may require anti-IgE and -IL-5 treatment. Our patients did not receive any additional medication to control their asthma during the study period.

Trajectories 1, 2 and 3 accounted for 79.3% of the study population. Trajectories 1 and 2 were associated normal to upper-normal lung function at baseline, whereas trajectory 3 was associated with a lower FEV1 at baseline, which remained evident at the end of the study (86.76%) (Tables 1 and 2). Additionally, trajectories 1 and 2 represent mild asthma with an annual frequency of exacerbations of less than 10%. In total, 20.8% of the subjects whose course followed trajectory 3 also had eosinophilic inflammation. Therefore, the eosinophilic inflammation in this group may be resistant to treatment with ICS and may require anti-IgE and -IL-5 treatment. Trajectories 1, 2, and 3 were categorized as mildly to moderately severe asthma. However, the demographic data were not consistent with previous cluster analyses of asthma phenotypes in Europe and the USA: The subjects in our study were less atopic, older at asthma onset, and had a normal BMI compared to asthmatics in Western countries. These parameters are in agreement with a previous study in Korea, suggesting that they are limited to adult Korean asthmatics.

Trajectory 3 showed distinct characteristics from the other groups; the final FEV1 was even worse than that in trajectory 4. When compared with other trajectory groups, trajectory 3 is more severe phenotypes than trajectories 1 and 2 regarding baseline lung function, final FEV1 value, the number of exacerbation and medication requirement during the study period. When compared with trajectory 4, trajectory 3 showed lower prevalence of BD reversibility, lower total IgE, and lower blood and sputum eosinophils. These observations suggest that these features might contribute to decreased lung function after 1 year of treatment in trajectory 3 compared with trajectory 4.

Among 1,679 asthmatics in this study, 1,457 (86.8%) were newly enrolled patients who were treatment-naive at baseline assessment, but 222 had been treated with asthma medications before the enrollment. The distributions of the 222 subjects in trajectory groups 15 were 10.4% (27 of 259 patients), 12.7% (68/534), 13.6% (73/538), 18.0% (25/139) and 13.9% (29/209),
respectively, which showed no statistical significance \((P = 0.313, \text{ Table 3})\). Among them, corticosteroid was administrated to only 1 patient in trajectory group 3, who was treated by a beclomethasone/formoterol inhaler (100/6 µg) at 28 days before the enrollment. Thus, our observation may be not biased by the previous treatment with inhaled corticosteroids.

When the same analysis was performed in 2,645 asthmatics including ex-smokers \((n = 515)\) and current smokers \((n = 451)\), the similar results between trajectories 4 and 5 were observed except smoking status; trajectory 4 was likely to be more eosinophilic, more atopic with higher levels of serum IgE, more frequent exacerbations, and more responsible to corticosteroids than those in trajectory 5 \((\text{Supplementary Fig. S3 and Supplementary Table S4})\). Interestingly, trajectory 5 had less proportion of ex-smokers and smokers compared to trajectory 4 \((18.0\% \text{ vs. } 39.3\%)\), but with greater amount of smoking \((15.4 \text{ vs. } 7.29 \text{ pack years})\) and longer periods of being ex-smoker \((4.09 \text{ vs. } 0.76 \text{ years})\). Thus, the different proportions of smoking and the difference of ex-smoking period were expected to have great confounding effects on interpretation of the longitudinal change of FEV1 trajectory: It was not clear whether the persistent airway obstruction of trajectory 5 was due to different airway dysfunctions or due to the different features of smoking and ex-smoking compared to trajectory 4. Accordingly, we excluded the effect of smoking on the 1-year trajectory of post-BD FEV1% changes to make the result to be clear. The effect of smoking on longitudinal changes in asthma symptoms should be assessed in further study with more precise design regarding balanced and matched duration and habits of cigarette smoking.

This study had several limitations. First, one-third of the longitudinal FEV1 data were missing; however, the missing data were imputed based on FEV1 values measured at least twice during the study period. Especially, the FEV1 values of 504 among 1,679 study subjects were restricted in the baseline and 3 months, which may introduce a bias in the imputation of late \((\text{over 6 months})\) values \((\text{Supplementary Table S1})\). However, as shown in \text{Tables 1 and 2, and Fig. 2}, the 5 FEV1% trajectories were mostly determined by values in the early stage of followup. When FEV1% trajectories were clustered using only the baseline and 3-month data \((1,578 \text{ subjects}, \text{Supplementary Table S1})\), the concordance rate of the trajectory with those using full-set data described in this study \((1,679 \text{ subjects})\) was 74.2\% for entire trajectory groups, and 92.0\% for trajectories 4 and 5. Thus, our observation, especially the difference between trajectory 4 and 5, could be acceptable. Furthermore, FEV1 values are unstable: in healthy individuals, the between-visit weekly variability in FEV1 is 12\% and the annual variability is 15\%; these values are higher in patients with airway disease.\(^{27}\) Secondly, we did not assess treatment adherence because we obtained medication information from the medical records. Thirdly, we did not collect longitudinal data on the blood and sputum levels of inflammatory markers, including eosinophils and neutrophils. Fourth, we excluded smokers from our asthma cohort to minimize the effect of smoking on lung function trajectories. As the prevalence of smoking in asthmatics is similar to that in the general population,\(^{38}\) our exclusion of smokers was unrealistic. However, this could be a strength of our study. Trajectory 5 was similar to the phenotype of severe asthmatics with fixed airflow obstruction or asthmaCOPD overlap, for which smoking is a major predisposing factor.\(^{39,40}\) Thus, our finding demonstrates that severe neutrophilic asthma is an important phenotype after controlling for the effect of smoking. Finally, we could not rule out the possibility of the over-diagnosis of asthma in patients whose course followed trajectories 1 and 2 because this occurs frequently \((25\%-35\%)\) in primary care patients.\(^{41,42}\) Additionally, blood and sputum cell counts were assessed in 612 \((36.5\%)\) and 757 \((45.1\%)\) of total 1,679 study subjects, respectively. Although the distributions of the subject assessed the inflammatory cell counts were statistically comparable between trajectories 4 and 5 \((46.8\% \text{ vs. } 37.3\%\); \(P = 0.095\) for blood cells, and 59.0\% and 56.0\%; \(P = 0.583\) for sputum cells), the
large proportion of subjects, who were not assessed, could introduce a bias in interpreting our observation about the different profiles of inflammatory cells between trajectory clusters.

In conclusion, this longitudinal analysis of post-BD FEV1 values identified 5 distinct trajectories. Trajectories 1, 2 and 3 comprised 80% of subjects with well-preserved lung function, whereas trajectories 4 and 5 were associated with different clinical parameters. Additionally, eosinophilic inflammation in sputum, particularly in individuals whose course follows trajectory 4, may be predictive of the response to conventional treatment for asthma, whereas neutrophilic inflammation in those whose course follows trajectory 5 suggests persistent airway obstruction. Further research should identify other predictors of the clinical course of asthma to facilitate personalized and effective treatment.

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SUPPLEMENTARY MATERIALS

Supplementary Table S1
Number of missing data of post-bronchodilator FEV1 over the followup in the total 1,679 subjects

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Supplementary Table S2
Medication other than ICS and SCS during the study period

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Supplementary Table S3
Inflammatory cell patterns in sputum of the 5 trajectories

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Supplementary Table S4
Clinical characteristics of the 5 trajectories in total 2,645 study subjects including ex- and current smokers

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Supplementary Fig. S1
Hierarchical clustering using Ward's method of (A) complete post-bronchodilator FEV1% data from 448 subjects, and (B) imputed post-bronchodilator FEV1% values from 1,679 subjects calculated using regression methods.

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Supplementary Fig. S2
Trajectories using k-means cluster for longitudinal data of complete data of post-bronchodilator FEV1% in 448 patients (A) and imputed post-bronchodilator FEV1% values from 1,679 subjects calculated using regression methods (B). A black line indicates individual lung function trajectories, and the colored bold lines are the distinct 5 trajectories.

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Supplementary Fig. S3
Trajectory analysis of post-bronchodilator FEV1% values in 2,645 asthmatics including ex-smokers (n = 515) and current smokers (n = 451) and serial changes in the mean FEV1 values. Data are presented as mean ± standard deviation.

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REFERENCES

1. Reddel HK, Bateman ED, Becker A, Boulet LP, Cruz AA, Drazen JM, et al. A summary of the new GINA strategy: a roadmap to asthma control. Eur Respir J 2015;46:622-39.
2. Fajt ML, Wenzel SE. Development of new therapies for severe asthma. Allergy Asthma Immunol Res 2017;9:3-14.
3. Sears MR. Lung function decline in asthma. Eur Respir J 2007;30:411-3.
4. Newby C, Agbetile J, Hargadon B, Monteiro W, Green R, Pavord I, et al. Lung function decline and variable airway inflammatory pattern: longitudinal analysis of severe asthma. J Allergy Clin Immunol 2014;134:287-94.
5. Moore WC, Meyers DA, Wenzel SE, Teague WG, Li H, Li X, et al. Identification of asthma phenotypes using cluster analysis in the Severe Asthma Research Program. Am J Respir Crit Care Med 2010;181:315-23.
6. Shaw DE, Berry MA, Hargadon B, McKenna S, Shelley MJ, Green RH, et al. Association between neutrophilic airway inflammation and airflow limitation in adults with asthma. Chest 2007;132:1871-5.
7. Amelink M, de Groot JC, de Nijs SB, Lutter R, Zwinderman AH, Sterk PJ, et al. Severe adult-onset asthma: a distinct phenotype. J Allergy Clin Immunol 2013;132:336-41.
8. Chung KF. Neutrophilic asthma: a distinct target for treatment? Lancet Respir Med 2016;4:765-7.
9. Chang HS, Lee TH, Jun IA, Baek AR, Park JS, Koo SM, et al. Neutrophilic inflammation in asthma: mechanisms and therapeutic considerations. Expert Rev Respir Med 2017;11:29-40.
10. Newby C, Heaney LG, Menzies-Gow A, Niven RM, Mansur A, Bucknall C, et al. Statistical cluster analysis of the British Thoracic Society Severe refractory Asthma Registry: clinical outcomes and phenotype stability. PLoS One 2014;9:e102987.
11. Loza MJ, Djukanovic R, Chung KF, Horowitz D, Ma K, Branigan P, et al. Validated and longitudinally stable asthma phenotypes based on cluster analysis of the ADEPT study. Respir Res 2016;17:165.
12. Boudier A, Curjuric I, Basagaña X, Hazgui H, Anto JM, Bousquet J, et al. Ten-year follow-up of cluster-based asthma phenotypes in adults. A pooled analysis of three cohorts. Am J Respir Crit Care Med 2013;188:550-60.
13. Kim TB, Jang AS, Kwon HS, Park JS, Chang YS, Cho SH, et al. Identification of asthma clusters in two independent Korean adult asthma cohorts. Eur Respir J 2013;41:1308-44.

14. Park SY, Baek S, Kim S, Yoon SY, Kwon HS, Chang YS, et al. Clinical significance of asthma clusters by longitudinal analysis in Korean asthma cohort. PLoS One 2013;8:e83540.

15. Savenije OE, Granell R, Caudri D, Koppelman GH, Smit HA, Wijsma A, et al. Comparison of childhood wheezing phenotypes in 2 birth cohorts: ALSPAC and PIAMA. J Allergy Clin Immunol 2011;127:1505-1512.e14.

16. Panico L, Stuart B, Bartley M, Kelly Y. Asthma trajectories in early childhood: identifying modifiable factors. PLoS One 2014;9:e111922.

17. Lødrup Carlsen KC, Mowinckel P, Hovland V, Håland G, Riiser A, Carlsen KH. Lung function trajectories from birth through puberty reflect asthma phenotypes with allergic comorbidity. J Allergy Clin Immunol 2014;134:917-923.e7.

18. Bisgaard H, Jensen SM, Bennelykke K. Interaction between asthma and lung function growth in early life. Am J Respir Crit Care Med 2012;185:1183-9.

19. Rasmussen F, Taylor DR, Flannery EM, Cowan JO, Greene JM, Herbertson GP, et al. Risk factors for airway remodeling in asthma manifested by a low postbronchodilator FEV1/vital capacity ratio: a longitudinal population study from childhood to adulthood. Am J Respir Crit Care Med 2002;165:1480-8.

20. Guerra S, Sherrill DL, Kurzius-Spencer M, Venker C, Halonen M, Quan SF, et al. The course of persistent airflow limitation in subjects with and without asthma. Respir Med 2008;102:1473-82.

21. Choi JS, Jang AS, Park JS, Park SW, Paik SH, Park JS, et al. Role of neutrophils in persistent airway obstruction due to refractory asthma. Respirology 2012;17:322-9.

22. Kim DK, Park YB, Oh YM, Jung KS, Yoo JH, Yoo KH, et al. Korean Asthma Guideline 2014: summary of major updates to the Korean Asthma Guideline 2014. Tuberc Respir Dis (Seoul) 2016;79:111-20.

23. Reddel HK, Taylor DR, Bateman ED, Boushey HA, Busse WW, et al. An official American Thoracic Society/European Respiratory Society statement: asthma control and exacerbations: standardizing endpoints for clinical asthma trials and clinical practice. Am J Respir Crit Care Med 2009;180:59-99.

24. Everitt BS. Cluster Analysis. 3rd ed. New York: John Wiley; 1993.

25. Genolini C, Falissard B. KmL: a package to cluster longitudinal data. Comput Methods Programs Biomed 2011;104:e112-21.

26. Tepper RS, Wise RS, Covar R, Irvin CG, Kercsmar CM, Kraft M, et al. Asthma outcomes: pulmonary physiology. J Allergy Clin Immunol 2012;129:865-87.

27. Pellegrino R, Vieggi G, Brusasco V, Crapo RO, Burgos F, Casaburi R, et al. Interpretative strategies for lung function tests. Eur Respir J 2005;26:948-68.

28. Chung KF, Wenzel SE, Brozek JJ, Bush A, Castro M, Sterk PJ, et al. International ERS/ATS guidelines on definition, evaluation and treatment of severe asthma. Eur Respir J 2014;43:343-73.

29. Engelkes M, Janssens HM, de Jongste JC, Sturkenboom MC, Verhamme KM. Medication adherence and the risk of severe asthma exacerbations: a systematic review. Eur Respir J 2015;45:396-407.

30. Tay TR, Hew M. Comorbid “treatable traits” in difficult asthma: current evidence and clinical evaluation. Allergy. Forthcoming 2017.

31. Fahy JV. Eosinophilic and neutrophilic inflammation in asthma: insights from clinical studies. Proc Am Thorac Soc 2009;6:256-9.
32. Kupczyk M, ten Brinke A, Sterk P, Bel EH, Papi A, Chanez P, et al. Frequent exacerbators—a distinct phenotype of severe asthma. Clin Exp Allergy 2014;44:212-21.
PUBMED | CROSSREF

33. van Veen IH, Ten Brinke A, Gauw SA, Sterk P, Rabe KF, Bel EH. Consistency of sputum eosinophilia in difficult-to-treat asthma: a 5-year follow-up study. J Allergy Clin Immunol 2009;124:615-7.
PUBMED | CROSSREF

34. Chung KF. Targeting the interleukin pathway in the treatment of asthma. Lancet 2015;386:1086-96.
PUBMED | CROSSREF

35. Shaw DE, Sousa AR, Fowler SI, Fleming LI, Roberts G, Corfield J, et al. Clinical and inflammatory characteristics of the European U-BIOPRED adult severe asthma cohort. Eur Respir J 2015;46:1308-21.
PUBMED | CROSSREF

36. Kim TB, Jang AS, Kwon HS, Park JS, Chang YS, Cho SH, et al. Identification of asthma clusters in two independent Korean adult asthma cohorts. Eur Respir J 2013;41:1308-14.
PUBMED | CROSSREF

37. Lee T, Lee YS, Bae YJ, Kim TB, Kim SO, Cho SH, et al. Smoking, longer disease duration and absence of rhinosinusitis are related to fixed airway obstruction in Koreans with severe asthma: findings from the COREA study. Respir Res 2011;12:1.
PUBMED | CROSSREF

38. Cerveri I, Cazzoletti L, Corsico AG, Marcon A, Niniano R, Grosso A, et al. The impact of cigarette smoking on asthma: a population-based international cohort study. Int Arch Allergy Immunol 2012;158:175-83.
PUBMED | CROSSREF

39. Thomson NC, Chaudhuri R, Heaney LG, Bucknall C, Niven RM, Brightling CE, et al. Clinical outcomes and inflammatory biomarkers in current smokers and exsmokers with severe asthma. J Allergy Clin Immunol 2013;131:1008-16.
PUBMED | CROSSREF

40. Gibson PG, McDonald VM. Asthma-COPD overlap 2015: now we are six. Thorax 2015;70:683-91.
PUBMED | CROSSREF

41. Montnémery P, Hansson L, Lanke J, Lindholm LH, Nyberg P, Löfdahl CG, et al. Accuracy of a first diagnosis of asthma in primary health care. Fam Pract 2002;19:365-8.
PUBMED | CROSSREF

42. Lucas AE, Smeenk FW, Smeele II, van Schayck CP. Overtreatment with inhaled corticosteroids and diagnostic problems in primary care patients, an exploratory study. Fam Pract 2008;25:86-91.
PUBMED | CROSSREF